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# Identification of Equine Lactadherin-derived Peptides That Inhibit Rotavirus Infection via Integrin Receptor Competition\*

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Background: Human milk lactadherin protects breastfed infants against symptomatic rotavirus infections. **Results:** A 20-aa peptide (namely pDGE-RGD) derived from equine lactahderin inhibits human rotavirus infectivity. **Conclusion:** pDGE-RGD interacts specifically with  $\alpha 2\beta 1$  integrin, thus hindering the rotavirus-cell attachment process. Significance: The discovery of the pDGE-RGD peptide may prove useful in the development of inhibitors of receptor recognition by rotavirus or other integrin-using viruses.

Human rotavirus is the leading cause of severe gastroenteritis in infants and children under the age of 5 years in both developed and developing countries. Human lactadherin, a milk fat globule membrane glycoprotein, inhibits human rotavirus infection in vitro, whereas bovine lactadherin is not active. Moreover, it protects breastfed infants against symptomatic rotavirus infections. To explore the potential antiviral activity of lactadherin sourced by equines, we undertook a proteomic analysis of milk fat globule membrane proteins from donkey milk and elucidated its amino acid sequence. Alignment of the human, bovine, and donkey lactadherin sequences revealed the presence of an Asp-Gly-Glu (DGE)  $\alpha_2 \beta_1$  integrin-binding motif in the N-terminal domain of donkey sequence only. Because integrin  $\alpha_2\beta_1$  plays a critical role during early steps of rotavirus host cell adhesion, we tested a minilibrary of donkey lactadherin-derived peptides containing DGE sequence for anti-rotavirus activity. A 20-amino acid peptide containing both DGE and RGD motifs (named pDGE-RGD) showed the greatest activity, and its mechanism of antiviral action was characterized; pDGE-RGD binds to integrin  $\alpha_2 \beta_1$  by means of the DGE motif and inhibits rotavirus attachment to the cell surface. These findings suggest the potential anti-rotavirus activity of equine lactadherin and support the feasibility of developing an anti-rotavirus peptide that acts by hindering virus-receptor binding.

Human rotavirus is a double-stranded RNA virus belonging to the Reoviridae family. It is transmitted by the fecal-oral route, causing severe gastroenteritis in infants and children under the age of 5 years in both developed and developing countries, although mortality occurs mainly in developing countries. As a result of rotavirus infection, 114 million episodes of diarrhea, 25 million clinic visits, 2.4 million hospital admissions, and more than 500,000 deaths in children up to age 5 occur worldwide annually (1). Groups at increased risk for rotavirus infection include children who are hospitalized or in community care centers and undernourished and/or immunodeficient children (2). Moreover, because exclusive breastfeeding was found to be associated with a lower incidence of rotavirus gastroenteritis (3–6), non-exclusively breastfed children are considered an additional group more vulnerable to rotavirus infections. The mature virion is a triple-layered particle of about 100 nm in diameter; the most external layer is composed of two viral proteins (VPs), VP7 (~34 kDa) and VP4 (87 kDa) (7, 8), with VP4 being the major determinant of tropism and receptor binding (9-12). Trimeric spikes of VP4 are anchored into the intermediate VP6 layer, whereas the trimeric calciumbinding protein VP7 covers the virion surface, locking VP4 spikes into place. The proteolytic cleavage of VP4 by trypsin is essential for optimum rotavirus infectivity and produces two subunits, VP5\* (60 kDa) and VP8\* (28 kDa), which remain associated with the virion (13-15). Initial cell attachment by rotaviruses is mediated by VP8\* binding to host cell glycans (16). Infection of permissive cells by many rotaviruses, including human (e.g. Wa and K8), monkey (RRV and SA11), and bovine (NCDV) strains, also depends on virus binding to particular integrins, a family of cell surface proteins that recognize extracellular matrix proteins (e.g. collagen), cell surface ligands (e.g. vascular cell adhesion molecule-1) (17), growth factors (e.g. fibroblast growth factor-1) (18), and viral proteins (e.g. rotavirus). VP5\* recognition of the collagen-binding  $\alpha 2\beta 1$  integrin is

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<sup>&</sup>lt;sup>3</sup> The abbreviations used are: VP, viral protein; aa, amino acid(s); MFGM, milk fat globule membrane; MEM, minimum essential medium; MOI, multiplicity of infection; PMF, peptide mass fingerprinting; SI, selectivity index.

a key event in rotavirus binding and entry into cells, which is followed by the interaction of VP7 with integrins  $\alpha x \beta 2$ ,  $\alpha 4 \beta 1$ , and  $\alpha v \beta 3$  (9, 19–24). The VP5\* subunits of almost all group A rotaviruses contain the Asp-Gly-Glu (DGE) sequence at aa 308 – 310, a motif that has been implicated in  $\alpha 2\beta 1$  recognition by type I collagen (17). Mutation of the putative  $\alpha 2\beta 1$  ligand sequence DGE abrogates binding of truncated VP5\* to the integrin  $\alpha$ 2 subunit I domain ( $\alpha$ 2I) and VP5\* competition with RRV cell binding and infectivity (9, 25). In addition, DGE-containing peptides, such as Asp-Gly-Glu-Ala (DGEA), specifically inhibit rotavirus-cell binding and infection mediated by  $\alpha 2\beta 1$ (9, 20, 21, 25). Binding by infectious monkey (SA11 and RRV) and human (Wa) rotaviruses to recombinant  $\alpha 2\beta 1$  expressed on K562 cells was specifically inhibited by DG-containing peptides and a function-blocking antibody to the  $\alpha$ 2I domain (9, 21, 23). Therefore, the interaction of rotavirus with  $\alpha 2\beta 1$  integrin can be considered a target for the development of antiviral agents aimed at preventing or reducing rotavirus infection.

Bioactive components in milk are an important research focus (26). In vitro, lactadherin, a milk fat globule membrane (MFGM) glycoprotein, inhibits rotavirus infection when sourced from humans, whereas bovine lactadherin is not active against human rotavirus (27). Moreover, it has been reported that human milk lactadherin protects breastfed infants against symptomatic rotavirus infections (6). Lactadherins isolated from different animal species are characterized by two N-terminal epidermal growth factor (EGF)-like domains, the second of which contains an Arg-Gly-Asp (RGD) integrin-binding motif that engages  $\alpha v \beta 3/5$  integrins and two C-terminal discoidin/F5/8C domain repeats (28). Bovine lactadherin consists of 427 amino acid residues, including an N-terminal signal sequence of 18 amino acids that is cleaved during intracellular processing. In contrast, human lactadherin is a protein of 387 amino acids, including an N-terminal signal sequence of 23 residues. When separated by two-dimensional electrophoresis, each of these proteins shows an apparent molecular mass above the theoretical value, with the difference due to glycosylation (29-31). MFGM-associated proteins from water buffalo, sheep, and horse also have been analyzed (32–34). Comparative species-to-species structural studies among MFGM proteins showed high variation, especially for lactadherin (35). It is proposed that these differences could explain the divergent effects of bovine and human lactadherin on human rotavirus infection. In order to test if the lack of rotavirus inhibition by bovine lactadherin is atypical or represents a shared property among lactadherins of related domesticated animals, we undertook a proteomic analysis of MFGM proteins from equine milk and elucidated the amino acid sequence of donkey lactadherin. Alignment of the human, bovine, and donkey lactadherin sequences revealed the presence of a DGE integrin-binding sequence motif in the donkey sequence only, suggesting a potential anti-rotavirus activity. To test this hypothesis, a minilibrary of donkey lactadherin-derived peptides containing DGE sequence was tested for anti-rotavirus activity. A 20-amino acid peptide containing both DGE and RGD sequence showed the greatest activity, and its mechanism of antiviral action was characterized. These findings shed light on the mechanisms of equine lactadherin inhibition of rotavirus infection and support

the feasibility of developing an anti-rotavirus peptide that acts through inhibition of virus-receptor binding.

#### **EXPERIMENTAL PROCEDURES**

Preparation of Fat Globules from Donkey Milk—Milk was collected from several donkeys at a single farm. Protease inhibitors (Complete; Roche Applied Science) were added, and the milk was immediately stored at  $-20\,^{\circ}$ C. Thawed milk was centrifuged at  $5000\times g$  for 30 min at  $10\,^{\circ}$ C, and the pellet was discarded. The cream layer and skimmed milk were centrifuged at  $189,000\times g$  for 70 min at  $6\,^{\circ}$ C. Fat globules were recovered in the supernatant and washed three times with 0.9% (w/v) NaCl.

Sample Protein Preparation and Two-dimensional Electrophoresis-Washed fat globules were incubated at 4°C overnight in 20 mM Tris-HCl, pH 8.6, containing 1% (w/v) ASB-14, 1% (v/v) Triton X-100, 7 м urea, and 2 м thiourea to extract the proteins associated with fat globule membranes. After centrifugation at  $18,400 \times g$  for 10 min at 10 °C, the floating cream layer was discarded. Proteins were precipitated from the supernatant with methanol and chloroform, as described previously (36). Pellets containing proteins were solubilized in 20 mm Tris-HCl, pH 8.6, containing 7 M urea, 2 M thiourea, 1% (w/v) ASB-14, 1% (v/v) DTT, and 0.5% (v/v) IPG buffer. Total protein was quantified using the 2-D Quant Kit (GE Healthcare). Extracted proteins (200 μg) were loaded onto 13-cm pH 3-10 NLIPG strips (GE Healthcare). Isoelectric focusing was carried out on an IPGphor unit (GE Healthcare) at 20 °C and 8000 V for a total of 70,000 V-h. Strips were incubated at room temperature in 50 mм Tris-HCl, pH 8.6, containing 6 м urea, 30% (v/v) glycerol, 2% (w/v) SDS, enriched with 2% (w/v) DTT for 20 min and afterward with 4.5% (w/v) iodoacetamide for 20 min. SDS-polyacrylamide gel electrophoresis was carried out on homogeneous running gels with 11.7% total acrylamide concentration and a 2.6% grade of cross-linking (Ettan DALT II system, GE Healthcare) at 400 V and 50 mA per gel for ~3 h. Gels were stained using the Processor Plus (GE Healthcare) with Blue Coomassie Colloidal stain (37) and scanned with a GS-800 densitometer (Bio-Rad).

Enzymatic Digestion of Proteins—In-gel multiple enzymatic digestion on selected two-dimensional electrophoresis spots was performed according to the published method (38). Briefly, excised spots were destained for 2 h in 40% (v/v) ethanol containing 50 mm NH<sub>4</sub>HCO<sub>3</sub>, dehydrated three times (10 min each) in acetonitrile, and dried in a vacuum concentrator for 30 min. Rehydration was performed at room temperature for 5 min with 5 μl of 75 ng/μl trypsin in 25 mM NH<sub>4</sub>HCO<sub>3</sub> (Promega Corp., Madison, WI), 5 μl of 75 ng/μl Asp-N solution (Sigma), or 5  $\mu$ l of 75 ng/ $\mu$ l chymotrypsin in 25 mM NH<sub>4</sub>HCO<sub>3</sub> (Sigma). Digestion continued at 37 °C. Asp-N peptides were collected after 2 h of digestion and submitted to mass spectrometry (MS). Trypsin and chymotrypsin digestions were performed overnight after the addition of 15  $\mu$ l of 25 mm NH<sub>4</sub>HCO<sub>3</sub>, 10% (v/v) acetonitrile. These supernatants were collected and submitted for MS analysis on the following day.

Mass Spectrometric Analysis—For MALDI-TOF mass spectrometry, 0.5  $\mu$ l of each peptide mixture was applied to a target disk and allowed to air-dry. Subsequently, 0.5  $\mu$ l of matrix solution (1% (w/v)  $\alpha$ -cyano-4-hydroxycinnamic acid in 30% (v/v)

acetonitrile and 0.1% (v/v) trifluoroacetic acid were applied to the dried sample and dried under vacuum. Spectra were collected on an Ultraflex II MALDI-TOF-TOF (Bruker Daltonik GmbH, Germany). The reported mass spectra are the result of signals of ~800 laser shots. The spectra were externally calibrated using the standard calibration mixture from Bruker Daltonik. Manual/visual estimation of the mass spectra was performed using Flex Analysis software (Bruker Daltonik). For LC-MS/MS experiments, an Agilent 1100 series liquid chromatograph and LC/MSD XCT series ion trap mass spectrometer equipped with nano-electrospray ionization source were used. Peptide mixtures were separated on an RP C18 column (Zorbax 300SB-C18; 3.5  $\mu$ m 150  $\times$  0.075 mm) (CPS) equilibrated with 0.1% (v/v) formic acid in water. Peptides were eluted over 55 min with a linear gradient of 5–70% (v/v) of 0.5% water, 0.1% formic acid in acetonitrile. The flow rate was 300 nl/min. The mass spectrometer was operated in positive ion mode. Data were processed using DataAnalysis software (Agilent Technologies, Santa Clara, CA).

Spectra Interpretation—For peptide mass fingerprinting (PMF) identification, the MS-Fit software package in ProteinProspector version 5.10.9 was used for interpretation of spectra (39). Data were searched against the *Horse* database UniProtKB.2012.03.21.

The parameters used for the search were as follows: *S*-carb-amidomethyl derivate on cysteine as fixed modification, oxidation on methionine as variable modification, and two missed cleavage sites for trypsin digestion. The peptide mass tolerance was 30 ppm.

The LC-MS/MS and MALDI-TOF/TOF spectra were analyzed using the MASCOT MS/MS Ions search engine against the UniProtKB.2012.03.21 database. The parameters used for the search were as follows: S-carbamidomethyl derivate on cysteine as fixed modification, oxidation on methionine as variable modification, two missed cleavage sites for each enzymatic digestion, and default selection of charge states ions (+2, +3). The peptide mass tolerance was set to 0.6 Da, and the fragment mass tolerance was set to 0.8 Da. Protein hits were validated if the protein scores were above the MASCOT default significance threshold ( $p \le 0.05$ ) and a minimum of two unique peptides matched.

*N-terminal Sequencing*—Spots were passively eluted from gels as described elsewhere (40), and the membrane was analyzed by direct microsequencing on a Procise 492 protein sequencer (Applied Biosystems). All chemicals were purchased from Applied Biosystems.

Cell Lines and Viruses—African green monkey kidney epithelial cells (MA104) and human epithelial colorectal adenocarcinoma cells (Caco-2) were propagated, respectively, in Eagle's minimum essential medium (MEM) and Dulbecco's modified Eagle's medium (DMEM; Sigma). Both media were supplemented with 1% (v/v) Zell Shield (Minerva Biolabs, Berlin, Germany) and heat-inactivated, 10% (v/v) fetal bovine serum (Sigma).

Chinese hamster ovary (CHO-K1) cells were cultured in high glucose DMEM (PAA, Pasching, Austria) supplemented as above. The derivation of CHO K1 cell transfectants expressing human  $\alpha$ 2 integrin subunit (CHO  $\alpha$ 2) or the PBJ-1 empty vec-

tor (CHO K1 PBJ-1) has been described previously (41–45). CHO cell transfectants were grown as for the parental cell line, except that G418 sulfate (Sigma) at  $0.1–0.8~\mu g/ml$  was included in the growth medium.

Human rotavirus strain Wa was obtained from the ATCC; the origin of porcine rotavirus P9 G3 strain CRW-8 has been described previously (46). Wa and CRW-8 infectivity was activated with 5  $\mu$ g/ml porcine pancreatic trypsin type IX (Sigma) for 30 min at 37 °C and propagated in MA104 cells in MEM containing 0.5  $\mu$ g/ml trypsin as described previously (47).

Antibodies—Mouse monoclonal antibodies (mAbs) directed to human rotavirus VP6 (0036),  $\alpha 2\beta 1$  integrin (AK7), and  $\alpha v\beta 3$  integrin (LM609) were purchased from Covalab (Villeurbanne, France), Abcam (Cambridge, UK), and Millipore (Molsheim, France), respectively.

Peptides—Synthetic peptides, pRGD-Hu (EVRGDVFP), pRGD-Bo (SHRGDVFI) (derived from human and bovine lactadherin, respectively), pRGD-Eq (SHRGDVFT), pDGE (QND-GECHV) (derived from equine lactadherin), pRGE (SHR-GEVFT), and pAGE (QNAGECHV) were purchased from Eurogentec (Liege, Belgium). Synthetic peptides pDGE-RGDmut1 (NNDGECHVIDDSHRGDVFTQ), pDGE-RGDmut2 (QNDGECHVIDDSDRGDVFTQ), pDGE-RGDmut3 (QND-GECHVIDDSHRGDVFSQ), pAGE-RGD (QNAGECHVID-DSHRGDVFTQ), and pDGE-RGE (QNDGECHVIDDSHR-GEVFTQ) were purchased from Proteogenix (Valparc, France). pDGE-RGD19 (NDGECHVIDDSHRGDVFTQ) and pDGEA (DGEA) were purchased from the PolyPeptide Group (Strasbourg, France). pAGE-RGE (QNAGECHVIDDSHRGEVFQT) was purchased from the CASLO Laboratory (Technical University of Denmark, Lyngby, Denmark). pDGE-RGD peptide (QNDGECHVIDDSHRGDVFTQ), derived from equine lactadherin, was purchased from Eurogentec, PolyPeptide Group, Proteogenix, and CASLO Laboratory. Biotinylated peptides Biot-pDGE (Biotin-QNDGECHV), Biot-pAGE (Biotin-QNAGECHV), and Biot-pDGE-RGD (Biotin-QNDGECH-VIDDSHRGDVFTQ) were purchased from Proteogenix. MALDI-TOF mass spectrometry analysis confirmed that pDGE-RGD, the peptide of major interest for this study, was soluble and monomeric (data not shown).

Rotavirus Infectivity Assays—Peptide blockade of virus infectivity was determined by a focus reduction assay using confluent MA104 cell monolayers in 96-well trays, as described previously (21). Unless otherwise stated, cells were treated for 1 h at 37 °C with peptides at 0.01–500 μM in serum-free medium prior to the virus addition. Rotavirus infection was performed at a multiplicity of infection (MOI) of 0.02, unless otherwise indicated, for 1 h at 37 °C, in the presence of the peptide unless otherwise stated. Infected cells were washed with serum-free medium and incubated in this medium at 37 °C in a humidified incubator in 5% (v/v)  $CO_2$ , 95% (v/v) air. In some experiments (as indicated), peptide pDGE-RGD was included with these washed, infected cells for the duration of their incubation. After 16 h of incubation, infected cells were fixed with cold acetone/ methanol (50:50), and viral titers were determined by indirect immunostaining, by using the UltraTech HRP Streptavidin-Biotin detection system (Beckman Coulter) according to the manufacturer's instructions. Peptide blockade of viral infectiv-

ity is expressed as a mean percentage  $\pm$  S.D. Where possible, antiviral effective concentration (IC<sub>50</sub>) values were calculated by regression analysis using the dose-response curves generated from the experimental data, using PRISM 4 (GraphPad Software, San Diego, CA). IC<sub>50</sub> values were compared using the sum-of-squares F test.

Assay of Virus Yield—To test the ability of pDGE-RGD to inhibit multiple cycles of viral replication, confluent MA104 cells in 24-well trays were infected with trypsin-activated Wa rotavirus (MOI = 0.02) for 1 h at 37 °C and washed as above. Cells were incubated in medium supplemented with 0.5  $\mu$ g/ml porcine trypsin and 30, 100, or 300  $\mu$ M pDGE-RGD, with fresh pDGE-RGD added to peptide-treated wells at 16 h after infection and at later fixed intervals and an equal volume of medium added to untreated controls at these times. Infected cells and cell supernatants were harvested at 48 h postinfection, and virus titers were determined by indirect immunostaining of MA104 cell monolayers inoculated with serial dilutions of the samples. The assay was performed in triplicate.

Rotavirus-Cell Binding Assay—Confluent MA104 cell monolayers in 24-well trays were washed, incubated with peptides or antibodies for 1 h at 37 °C, and cooled on ice for 20 min in the continuing presence of peptides or antibodies. Trypsin-activated virus, which had been cooled to 4 °C, was allowed to attach to cells on ice for 1 h (MOI = 3) in the presence of peptides or antibodies. Cells were washed with cold MEM; cold MEM was added. Cells were subjected to two rounds of freezethawing and incubated at 37 °C for 30 min with 10  $\mu$ g/ml porcine trypsin to release bound virus, and the lysate was clarified by low speed centrifugation for 10 min. Cell-bound virus titers were determined by indirect immunostaining as above.

Rotavirus-Cell Entry Assay—Confluent MA104 cell monolayers in 96-well trays were washed twice with MEM and cooled on ice for 20 min. Trypsin-activated virus, which had been cooled to 4 °C, was adsorbed to cells on ice for 1 h at 4 °C (MOI = 0.02). Peptide pDGE-RGD was added to washed cells at 0.01–500  $\mu$ M for 1 h at 37 °C. Rotavirus entry was stopped by the removal of surface-bound virions with two quick washes with 3 mM EGTA in PBS, which removes the virion outer capsid (48, 49). Cells were incubated in MEM for 16 h at 37 °C, fixed, and stained, and titers were determined as described above for the infectivity assay.

*Virucidal Assay*—Trypsin-activated Wa rotavirus was reacted with 100  $\mu$ M pDGE-RGD peptide or diluent for 1 h at 37 °C and then titrated. Viral yields were determined by indirect immunostaining, by counting the number of infectious foci at 1:1,024, 1:2,048, and 1:4,096 dilutions.

Peptide-Cell Binding Assay—Confluent CHO K1 and CHO  $\alpha 2$  cells in 96-well trays were washed twice with DMEM containing 1% (w/v) BSA and incubated for 1 h at 37 °C with 0.3, 3, or 30  $\mu$ M Biotin-pDGE, Biotin-pAGE, or Biotin-pDGE-RGD peptides in the same DMEM diluent. Washed cells were incubated with streptavidin-HRP (Beckman Coulter) for 10 min at room temperature, washed twice with PBS, and reacted with ABTS reagent (ThermoScientific) for 30 min at room temperature. The reaction was stopped with PBS containing 5% (w/v) SDS, and the absorbance was determined at  $\lambda = 405$  nm.

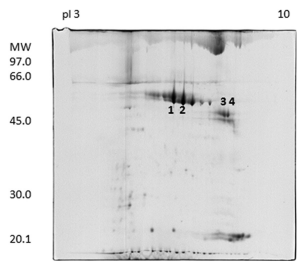


FIGURE 1. Two-dimensional electrophoresis map of donkey milk fat globule membrane proteins. Identified proteins are *marked* with *numbers*. The *spots numbered 1–4* were excised from the gel and analyzed by MALDI-TOF MS and PMF. All were identified as horse milk fat globule-EGF factor 8 splice variant, also known as horse lactadherin.

Cell Viability Assay—The cytotoxicity of peptide pDGE-RGD was assessed in MA104 and CaCo2 cells seeded in 96-well plates at a density of  $1.2\times10^4$  cells/well. After 24 h of culture, cells were treated with peptide at  $0.0044-880~\mu\text{M}$ . Cell viability was determined 48 h later with the CellTiter 96 proliferation assay kit (Promega) according to the manufacturer's instructions. Absorbances were measured using a model 680 microplate reader (Bio-Rad) at 490 nm. Viability in treated cells was expressed as a percentage relative to cells incubated with culture medium alone. The 50% cytotoxic concentration (CC<sub>50</sub>) was determined using logarithmic viability curves. Where possible, a selectivity index (SI) was calculated by dividing the CC<sub>50</sub> by the IC<sub>50</sub> value.

Statistical Analysis—One-way analysis of variance, in some cases followed by the Bonferroni test, was used to assess the statistical significance of differences in virus titers and absorbance data. Significance was set at the 95% level.

#### **RESULTS**

Proteomic Analysis and Protein Sequencing Indicate the Presence of Two Putative Integrin Ligand Motifs in Equine Lactadherin-The two-dimensional electrophoresis map of donkey milk fat globule membrane proteins was determined (Fig. 1). The two-dimensional electrophoresis map showed a pattern very similar to that previously published for horse MFGM proteins, where lactadherin was the major protein and was identified in several spots at different molecular masses (34). The spots 1-4, corresponding to the molecular masses of horse lactadherin, were excised, and the proteins were identified by MALDI-TOF MS and PMF as the horse milk fat globule-EGF factor 8 splice variant, also known as horse lactadherin (Table 1). To understand the nature of the difference between the  $\sim$ 45- and  $\sim$ 50-kDa proteins, their N-terminal amino acid sequences were determined (Table 2). The sequences of proteins 1 and 2 matched the N-terminal amino acid sequence of horse lactadherin (UniProt F7B0S3) with an identity of ≥95%. Proteins 3 and 4 corresponded to the same horse lactadherin

**TABLE 1**Identification of donkey milk fat globule membrane proteins by PMF

Spot no. (from Fig. 1)	Protein name	UniProt KB/TrEmbl entry	MOWSE score <sup>a</sup>	Experimental/nominal molecular mass	PMF number of signals m/z matched/number of signals m/z collected (% sequence coverage)
				Da	
1	Milk fat globule-EGF factor 8 splice variant	F5CEP2	5.25e + 8	50,000/43,308	24/78 (66.2%)
2	Milk fat globule-EGF factor 8 splice variant	F5CEP2	2.26e + 9	50,000/43,308	23/59 (61.1%)
3	Milk fat globule-EGF factor 8 splice variant	F5CEP2	4.74e + 6	45,000/43,308	17/60 (54.1%)
4	Milk fat globule-EGF factor 8 splice variant	F5CEP2	1.44e + 7	45,000/43,308	18/54 (54.1%)

<sup>&</sup>lt;sup>a</sup> MOWSE score is obtained by the MS-Fit search and concerns the highest scored protein.

**TABLE 2**N-terminal amino acid sequences and identification of donkey milk fat globule membrane proteins

Spot number (Fig. 1)	N-terminal amino acid sequence by Edman degradation	Comparison with horse lactadherin amino acid sequence (F7B0S3)	Identity	
			%	
1	ASGDFCDSSQCLNGGTCLLGQ	<sup>1</sup> AIAGDFCDSSQCLNGGTCLLGQ <sup>22</sup>	95%	
2	ASGDFCDSSQCLNGGTCLLGQDDL	<sup>1</sup> AIAGDFCDSSQCLNGGTCLLGQDDL <sup>25</sup>	96%	
3	ASGPCFPNPCQNDGECHVIDDSHR	44GPCFPNPCQNDGECHVIDDSHR <sup>65</sup>	96%	
4	ASGPCFPNPCQNDGECHVID	44GPCFPNPCQNDGECHVID <sup>62</sup>	90%	

sequence, commencing from the amino acid residue at position 44. Because the 45-kDa form was most similar to the human counterpart, its amino acid sequence was examined further. The results of multiple enzymatic digestions of this 45-kDa protein with trypsin, Asp-N, and chymotrypsin, followed by MALDI-TOF/TOF and nano-LC-electrospray ionization MS/MS analyses (Table 3), together with the N-terminal amino acid sequence, gave an (almost) complete coverage of the sequence of this form of donkey lactadherin (Fig. 2). The sequence was submitted to the UniProt database, and the accession number C0HJR4 was assigned.

The sequence alignment of this equine (*EQ*) lactadherin with the bovine (*BO*) and human (*HU*) lactadherins revealed some very interesting features in the N-terminal domains (Fig. 3). An RGD motif is conserved in the three proteins, being present at amino acid residues 23–25 in equine and human lactadherin and at residues 67–69 in bovine lactadherin. RGD is an integrin-binding motif that engages  $\alpha_{\rm v}\beta_3$ ,  $\alpha_{\rm v}\beta_5$ ,  $\alpha_3\beta_1$ , and  $\alpha_5\beta_1$  integrins to facilitate cell adhesion. Surprisingly, the equine lactadherin was shown to contain an additional integrin binding motif, the DGE sequence that can specifically bind the  $\alpha_2\beta_1$  integrin, which is present at residues 12–14.

Functional Roles of the Equine Lactadherin RGD and DGE Sequences in Inhibition of Rotavirus Infection-In order to assess the importance of the RGD sequence (shared by human, bovine, and equine lactadherin) and the DGE motif (present in equine lactadherin only) for inhibition of rotaviral infection, the effect on Wa infection of cellular treatment with the 8-amino acid peptides pRGD-Hu, pRGD-Bo, pRGD-Eq, and pDGE was determined. These human, bovine, and equine RGD peptides inhibited Wa infection of MA104 cells in a dose-dependent fashion to maxima at 500  $\mu$ M of 57, 50, and 47%, respectively (Fig. 4A). In contrast, MA104 cell treatment with an 8-amino acid RGE peptide harboring a single amino acid substitution  $(Asp \rightarrow Glu)$  known to inhibit the ability of the RGD triplet to bind integrin (pRGE) had no effect on virus infectivity even at high concentrations (Fig. 4A). Notably, the DGE-containing peptide pDGE derived from equine lactadherin inhibited Wa infectivity to a maximum of 84% (Fig. 4B), making it possible to

assess its IC<sub>50</sub> at 17  $\mu$ M (Table 4). As expected, an AGE peptide (pAGE), despite its ability to inhibit rotaviral infectivity to a maximum of 80%, showed a significantly higher IC<sub>50</sub> (257  $\mu$ M) than pDGE (p < 0.0001; Table 4). As a positive control, we tested the Type I collagen-derived peptide Asp-Gly-Glu-Ala (pDGEA); this  $\alpha_2\beta_1$  integrin ligand showed an antiviral activity comparable with the one previously reported (21), as depicted in Fig. 4B.

A 20-Amino Acid Region of Equine Lactadherin Has Elevated Specific Anti-rotaviral Activity in Epithelial Cell Cultures That Depends on Its RGD and DGE Sequences—Interestingly, the 20-amino acid peptide pDGE-RGD, corresponding to amino acids 10-29 of equine lactadherin and harboring both the RGD and DGE motifs, showed the highest anti-rotaviral efficacy  $(IC_{50} = 4.4 \mu M; Table 4)$ . pDGE-RGD was therefore selected as the lead compound for further studies. Of note, the pDGE-RGD peptide maintained a significant (p < 0.05) inhibitory activity against high MOIs of Wa (Fig. 4C). The stage of the rotavirus replication cycle inhibited by peptide pDGE-RGD was examined. Peptide addition 1 h before or immediately after infection did not inhibit Wa infectivity. The greatest infectivity inhibition was achieved when the peptide was added during infection (Fig. 5), as performed in the experiments described above and in the following studies. To assess the role played by each motif in inhibition of rotavirus infectivity, the antiviral activity of several mutant peptides derived from pRGD-DGE that harbored mutations in DGE (Asp  $\rightarrow$  Ala, pAGE-RGD), RGD (Asp  $\rightarrow$  Glu, pDGE-RGE), or both triplets (pAGE-RGE) was tested. These peptides failed to show any antiviral activity even at high concentrations, demonstrating the importance of these motifs for anti-rotavirus activity (Table 4). In contrast, peptides with single amino acid substitutions outside these functional groups, namely pDGE-RGDmut1, pDGE-RGDmut2, and pDGE-RGDmut3, retained the ability to inhibit Wa infection (Table 4), at levels comparable with those of pDGE-RGD. No significant difference was seen between the IC50 of pDGE-RGD and these peptides with amino acid substitutions outside the DGE and RGD functional groups (p > 0.05). The antiviral efficacy of pDGE-RGD and its AGE and RGE mutants was also evaluated

**TABLE 3**Identification of the 45-kDa donkey lactadherin short form by enzymatic digestions and mass spectrometry techniques

	PMF by MALDI-TOF			LC-MS/MS			
Enzyme	Position	m/z, single charge	$Sequence^a$	Position	m/z, charge state	Sequence <sup>a</sup>	
Trypsin	127-134	838.4305	GGTAEYLK	89-102	770.3600 (+2)	TGIVNAWTASNYDK	
	138-144	866.4367	VAYSVDGR	289-300	706.2300 (+2)	DFGHIQYVAAYK	
	116-126	1074.5902	VTGVVTQGASR	25-37	801.7400 (+2)	GDVFTQYICSCPR	
	349-357	1105.6265	ILPVAWHNR	278-288	558.2200 (+2)	QVTGVVTQGAR	
	358-367	1173.6660	ITLRVELLGCam	160-171	624.6400 (+2)	VFVGNVDNSGLK	
	336-345	1225.6034	NMFETPFLAR	116-126	537.7100 (+2)	VTGVVTQGASR	
	336-345	1241.5983	NMoxFETPFLAR	199-224	959.2800 (+3)	FELLGCEVNAGCAEPLGLEDNSIPD:	
	289-300	1411.7005	DFGHIQYVAAYK	349-357	553.1500 (+2)	ILPVAWHNR	
	146-157	1411.7328	KFQFIRDAGDSK	173-185	854.8500 (+2)	VNMFDVPLEVQYVR	
	323-335	1483.7625	IFPGNLDNNSHKK	255-277	856.5900 (+3)	FNAWTAQSNSASEWLQVDLGSQK	
	186-198	1519.7620	LVPVACamHHGCamTLR	186-198	760.2300 (+2)	LVPVACHHGCTLR	
	25-37	1602.7039	GDVFTQYICamSCamPR	336-345	613.2900 (+2)	NMFETPFLAR	
	172-185	1708.8727	VNMFDVPLEVQYVR	323-334	678.1600 (+2)	IFPGNLDNNSHK	
	103-115	1717.8989	NPWIQVNLMRKMR	79-85	435.6300 (+2)	WVPELAR	
	172-185	1724.8676	VNMFDVPLEVQYVR				
	234-248	1878.8962	TWGLNAFSWYPFYAR				
	255-277	2567.2161	FNAWTAQSNSASEWLQVDLGSQK				
	199-224	2875.3237	FELLGCamEVNAGCamAEPLGLEDNSIPDR				
Chymotrypsin	347 - 354	953.5931	VRILPVAW				
	233-240	964.4999	RTWGLNAF				
	32-39	1012.4339	ICamSCamPRGY				
	87-95	1053.5588	HRTGIVNAW				
	71-79	1157.5561	FGFMGLQRW				
	246-255	1225.6688	YARLDKQGKF				
	76-86	1267.7270	QRWVPELARL				
	29-39	1404.6035	TQYICamSCamPRGY				
	246-258	1596.8281	YARLDKQGKFNAW				
	296-310	1618.7608	VAAYKVSHSNDGANW				
	184-197	1618.8305	VRLVPVACamHHGCamTL				
	148-161	1625.8282	QFIRDAGDSKDKVF				
	329-342	1708.7748	DNNSHKKNMFETPF				
	215-232	1966.9352	GLEDNSIPDRQITASSTY				
	162-183	2453.2017	VGNVDNSGLKVNMFDVPLEVQY				
	40-71	3393.4742	TGTHCamETTCamAMPLGMETGAIADAQISASSVYF				
	314-342	3393.6393	RDQRAADSKIFPGNLDNNSHKKNMFETPF				
A NT	101-132	3578.8319	DKNPWIQVNLMRKMRVTGVVTQGASRGGTAEY	272 200	072 2000 ( + 2)	DI CCOVOVECNITOCAD	
Asp-N	155–165	1207.6317	DSKDKVFVGNV	272-288	872.3800 (+2)	DLGSQKQVTGVVTQGAR	
	166–175	1140.5354	DNSGLKVNMF	289-305 180-198	968.8900 (+2)	DFGHIQYVAAYKVSHSN	
					814.3400 (+3)	EVQYVRLVPVACHHGCTLRF	
				131–142 166–175	724.3000 (+2)	EYLKTFKVAYSV	
				26-44	570.7000 (+2)	DNSGLKVNMoxF	
				26-44 158-165	774.8400 (+3)	DVFTQYICSCPRGYTGTHC DKVFVGNV	
				166–175	439.2100 (+2)	DNSGLKVNMF	
				320–328	562.6800 (+2)	DSKIFPGNL	
				320-328 45-60	495.7300 (+2) 842.7800 (+2)	ETTCAMoxPLGMoxETGAIA	
				45-60 143-151	842.7800 (+2)	DGRKFQFIR	
				13–151	583.7300 (+2)	DGECHVI	
				250–266	415.1600 (+2) 920.3900 (+2)	DKQGKFNAWTAQSNSAS	
				1-12	674.6700 (+2)	ASGPCFPNPCON	
				1-12	074.0700 (⊤2)	AGGECEENECQN	

<sup>&</sup>lt;sup>a</sup> Cam, carbamidomethylated cysteine; Mox, oxidized methionine.

F7B0S3 F5CEP2 C0HJR4	<sup>1</sup> AIAGDFCDSSQCLNGGTCLLGQDDLPFYCLCPEGFTGLICNETEKGPCFPNPCQNDGECHVIDDSHRGDVFTQYIC <sup>1</sup> GRPPFYCLCPEGFTGLICNETEKGPCFPNPCQNDGECHVIDDSHRGDVFTQYIC <sup>1</sup> <u>AS</u> GPCFPNPCQNDGECHVIDDSHRGDVFTQYIC
F7B0S3 F5CEP2 C0HJR4	77SCPRGYTGTHCETTCAMPLGMETGAIADAQISASSVYFGFMGLQRWVPELARLHRTGIVNAWTASNYDKNPWQ 55SCPRGYTGTHCETTCAMPLGMETGAIADAQISASSVYFGFMGLQRWVPELARLHRTGIVNAWTASNYDKNPWQ 34SCPRGYTGTHCETTCAMPLGMETGAIADAQISASSVYFGFMGLQRWVPELARLHRTGIVNAWTASNYDKNPWQ
F7B0S3 F5CEP2 C0HJR4	151VNLMRKMRVTGVVTQGASRGGTAEYLKTFKVAYSVDGRKFQFIRDAGDSKDKVFVGNVDNSGLKVNMFDVPLE 128VNLMRKMRVTGVVTQGASRGGTAEYLKTFKVAYSVDGRKFQFIRDAGDSKDKVFVGNVDNSGLKVNMFDVPLE 108VNLMRKMRVTGVVTQGASRGGTAEYLKTFKVAYSVDGRKFQFIRDAGDSKDKVFVGNVDNSGLKVNMFDVPLE
F7B0S3 F5CEP2 C0HJR4	<sup>224</sup> VQYVRLVPVACHHGCTLRFELLGCEVN -GCAEPLGLEDNSIPDRQITASSTYRTWGLNAFSWYPFYARLDKQGKFN <sup>202</sup> VQYVRLVPVACHHGCTLRFELLGCEVNAGCAEPLGLEDNSIPDRQITASSTYRTWGLNAFSWYPFYARLDKQGKFN <sup>181</sup> VQYVRLVPVACHHGCTLRFELLGCEVNAGCAEPLGLEDNSIPDRQITASSTYRTWGLNAFSWYPFYARLDKQGKFN
F7B0S3 F5CEP2 C0HJR4	<sup>288</sup> AWTAQSNSASEWLQVDLGSQKQVTGVITQGARDFGHIQYVAAYKVSHSNDGANWTEYRDQRAADSKIFPGNLDNN <sup>277</sup> AWTAQSNSASEWLQVDLGSQKQVTGVITQGARDFGHIQYVAAYKVSHSNDGANWTEYRDQRAADSKIFPGNLDNN <sup>288</sup> AWTAQSNSASEWLQVDLGSQKQVTGV <u>V</u> TQGARDFGHIQYVAAYKVSHSNDGANW <u>XX</u> RDQRAADSKIFPGNLDNN
F7B0S3 F5CEP2 C0HJR4	<sup>373</sup> SHKKNMFETPFLARFVRILPVAWHNRITLRVELLGC <sup>409</sup> <sup>302</sup> SHKKNMFETPFLARFVRILPVAWHNRITLRVELLGC <sup>308</sup> <sup>202</sup> SHKKNMFETPFLARXVRILPVAWHNRITLRVELLGC <sup>328</sup>

FIGURE 2. Amino acid sequence of the 45-kDa form of donkey lactadherin obtained by N-terminal sequencing, PMF by MALDI-TOF, and LC-MS/MS. The sequence of the 45-kDa form of donkey lactadherin obtained (UniProt entry COHJR4) is aligned with two isoforms of Equus caballus lactadherin (UniProt entries F7B0S3 and F5CEP2). Amino acids of the 45-kDa form of donkey lactadherin that differ from the other lactadherins are given in boldface type and underlined.

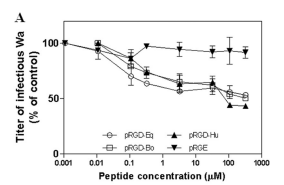
in the Caco-2 intestinal cell line (Table 5). Whereas pDGE-RGD showed antiviral activity in Caco-2 cells (IC $_{50}=8.8~\mu\text{M}$ ) comparable with that in MA104 cells, pDGE-RGE, pAGE-RGD, and pAGE-RGE exhibited a significant loss of activity

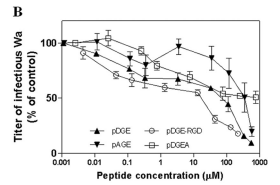
(p < 0.0001). Interestingly, these Caco-2 cell studies identified significant differences in antiviral activity between these three peptides (0.002  $, with pDGE-RGE showing a lower IC<math display="inline">_{50}$  (46  $\mu\rm_{M}$ ) than pAGE-RGD (IC $_{50} = 101~\mu\rm_{M})$  and



EQ Lact45 BO Lactad Hu Lactad	ASGCFPNPCQN <b>DGE</b> CH 16 ASGDFCDSSLCLHGGTCLLNEDRTPPFYCLCPEGFTGLLCNETEHGPCFPNPCHNDAECQ 60LDICSKNPCHNGGLCE 16
EQ Lact45	17 VIDDSH <b>rgd</b> /FTQYICSCPRGYTG-HCETTCAMPLGMETG 55
BO Lactad	61 VTDDSH <b>RGD</b> /FIQYICKCPLGYVGIHCETTCTSPLGMQTG 100
Hu Lactad	17 EISQEV <b>RGD</b> /FPSYTCTCLKGYAGNHCETKCVEPLGMENG 56

FIGURE 3. Sequence alignment of the equine (EQ Lact45), bovine (BO Lactad), and human (HU Lactad) lactadherins. The integrin-binding motifs RGD and DGE are given in boldface type and boxed.





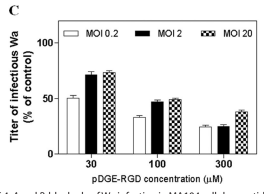


FIGURE 4. *A* and *B*, blockade of Wa infection in MA104 cells by peptides pRGD-Eq, pRGD-Hu, pRGD-Bo, and pRGE (*A*) and pDGE-RGD, pDGE, pAGE, and pDGEA (*B*). *C*, blockade of Wa infection in MA104 cells by pDGE-RGD at different MOIs. The infectivity titer of virus in the treated sample is expressed as a percentage of the titer obtained in the absence of peptide (untreated control). *Error bars*, S.D. of three independent experiments.

pAGE-RGE (IC $_{50}=152~\mu \rm M$ ). This suggests that the DGE sequence plays a major role in the process leading to rotavirus inhibition. In order to rule out any nonspecific effect due to loss of cell viability, MTS cell viability assays were performed after incubation of Wa-infected and control MA104 and Caco-2 cell cultures for 72 h with a range of peptide concentrations. The tested peptides did not impair cell viability at any concentration tested on MA104 cells, showing a 50% cytotoxic concentration

 $(CC_{50})$  of >880  $\mu$ M (Tables 4 and 5). In Caco-2 cells, the pDGE-RGD  $CC_{50}$  was 945  $\mu$ M.  $CC_{50}$  values were used to calculate the SI of pDGE-RGD, which was >100 for both cell lines (Tables 4 and 5).

Infectious Rotavirus Yield Is Reduced by the 20-Amino Acid Lactadherin Peptide—The antiviral activity of pDGE-RGD was evaluated in a virus yield assay, a more stringent test that allows multiple cycles of viral replication to occur before measuring the titer of infectious virus produced. Wa rotavirus yield was reduced in a dose-dependent manner in cells treated with 100 or 300  $\mu$ M pDGE-RGD (p < 0.05), showing 50 and 84% inhibition, respectively (Fig. 6).

The 20-Amino Acid Lactadherin Peptide Binds  $\alpha_2\beta_1$  Integrin in a DGE-dependent Manner-The results described above suggested that pDGE-RGD may inhibit rotavirus infectivity by binding the  $\alpha_2\beta_1$  integrin, thereby blocking viral binding to this cellular receptor. The ability of equine lactadherin peptides to specifically bind  $\alpha_2\beta_1$  was tested by treating CHO cells expressing surface  $\alpha_2\beta_1$  with biotinylated pDGE-RGD, pDGE, and pAGE peptides. Peptides pDGE-RGD and pDGE bound to the surface of  $\alpha_2\beta_1$ -expressing CHO cells in a dose-dependent manner, whereas pAGE did not show this ability (Fig. 7A). Peptide pDGE-RGD bound to a greater extent than pDGE, because pDGE-RGD binding was significant greater than that of pDGE at 3 and 30  $\mu$ M (0.01 0.02). In contrast, pDGE binding tothe CHO parental cell line (CHO-K1) (Fig. 7B) was reduced compared with  $\alpha_2\beta_1$ -expressing CHO cells at all concentrations tested (p < 0.0006). No difference was detected between pDGE and pAGE binding to CHO-K1 cells (p > 0.05). Interestingly, substantial binding of pDGE-RGD to CHO-K1 cells was evident, indicating binding to a cell surface component(s) other than  $\alpha_2\beta_1$ . Consistently, at 30  $\mu$ M, pDGE-RGD binding to the surface of  $\alpha_2\beta_1$ -expressing CHO cells was significantly greater than to the parental cell line (p < 0.05).

The 20-Amino Acid Lactadherin Peptide Preferentially Inhibits Rotavirus-Cell Attachment Rather than Entry in a DGE-dependent Manner—In order to evaluate the ability of pDGE-RGD to inhibit rotavirus-cell attachment, we measured the titer of infectious Wa rotavirus particles attached to pDGE-RGD-treated and untreated MA104 cells. pDGE-RGD significantly reduced Wa binding by a maximum of 67% (p < 0.0001), whereas pAGE-RGE did not show any inhibitory activity (p > 0.05; Fig. 8A). As found previously (9), the AK7 mAb to integrin  $\alpha_2\beta_1$  reduced virus-cell binding to a maximum of 51% (p < 0.05), whereas mAb LM609 to integrin  $\alpha_v\beta_3$  had no significant effect (p > 0.05). In a control experiment, we tested the antiviral efficacy of pDGE-RGD against the integrin-independent rotavirus strain CRW-8. As expected, pDGE-RGD did not inhibit CRW-8 infectivity at any concentration tested (Fig. 8B). The

**TABLE 4**Effect of lactadherin-derived peptides on cell viability and Wa rotavirus infectivity in MA104 cells

			95% confidence		
Peptides	Features	$IC_{50}$	Interval	$CC_{50}$	SI
		μм		$\mu$ M	
pDGE	peptide DGE, 8 aa, equine	17.3	6.9-43.5	>880	>50.9
pAGE	aa substitution in position 3	256.6	177.7-370.3	>880	>3.4
pDGE-RGD	Wild type peptide	4.4	2.7-7.1	>880	>200
pAGE-RGD	aa substitution in position 3	>220	$NA^a$	>880	>4
pDGE-RGE	aa substitution in position 16	>220	NA	>880	>4
pAGE-RGE	aa substitutions in position 3 and 16	>220	NA	>880	>4
pDGE-RGDmut1	aa substitution in position 1	13.5	5.1-35.7	>880	>65.2
pDGE-RGDmut2	aa substitution in position 13	13.1	7.5-22.9	>880	>67.2
pDGE-RGDmut3	aa substitution in position 19	6.7	2.2-20.4	>880	>131.3

a n.a., not assessable.

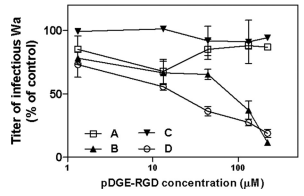


FIGURE 5. A–C, time-of-addition experiments were performed by treating cells with pDGE-RGD for 1 h before infection (A) or for 1 h during infection (B) or by adding the peptide immediately after infection (C). Control experiments were performed by treating cells both before and during viral adsorption (D). The infectivity titers of virus in the treated samples are expressed as a percentage of the titer obtained in the absence of peptide (untreated control). Error bars, S.D.

effect of pDGE-RGD on viral entry also was investigated. This peptide significantly inhibited Wa rotavirus-cell entry only when added to cells at very high concentrations (i.e. 300 or 500  $\mu$ M) (p < 0.05; Fig. 8C). In order to rule out any virucidal effect of pDGE-RGD on rotavirus, Wa was exposed for 1 h to a pDGE-RGD concentration (100  $\mu$ M) able to inhibit 80% of Wa infectivity. The infectious titer of the pDGE-RGD-treated viral suspension was indistinguishable from that of the untreated control (p > 0.05; Fig. 8D).

## **DISCUSSION**

Many human and animal rotaviruses contain the Asp-Gly-Glu (DGE) sequence in VP8\*, which is necessary for their usage of the  $\alpha 2\beta 1$  integrin to aid infection. In this study, we have identified the putative integrin ligand motifs RGD and DGE in the N-terminal domain of donkey lactadherin. Whereas the RGD motif is common to human, bovine, and donkey lactadherins, the DGE motif is found only in the donkey sequence. Direct evaluation of the antiviral activity of donkey lactadherin would be of interest but has proven to be difficult. Difficulties arise when experiments to purify donkey lactadherin with the same methods used for the human and bovine proteins are attempted, because human and bovine milk contain a much higher amount of fat ( $\sim$ 3.5%) than donkey milk ( $\sim$ 0.3%) from which only a very small amount of lactadherin could be recovered. For this reason, we turned to the recombinant protein. We achieved bacterial expression of full-length donkey lactadherin,

but the recombinant His-tagged protein was insoluble and could not be refolded. This is consistent with the insolubility of milk-derived lactadherins, which are associated with the milk fat globule membrane (50). In any case, our main focus is on the development of rotavirus inhibitors based on derivative peptides rather than analyzing the full-length protein; one main reason is that it is very unlikely that lactadherin molecules pass through the digestive tract intact, at least not in a relevant amount (e.g. we have easily digested donkey lactadherin with trypsin to solve their primary structure). Moreover, the ability of the DGE integrin binding domain to inhibit rotavirus replication can be addressed much more clearly working with small peptides rather than intact proteins, where the tertiary and eventually quaternary structure of the molecule could mask the biological activity of the inner part of some protein segment. To this end, we have demonstrated that peptides derived from donkey lactadherin, which contain the intact RGD and DGE sequences, specifically and effectively inhibit rotavirus infection at low micromolar levels. We selected a 20-amino acid peptide derived from donkey lactadherin harboring the RGD and a DGE motif (pDGE-RGD), depending on its favorable SI and its ability to limit the production of viral progeny resulting from multiple cycles of viral replication. The differences between MA104 and Caco-2 cells in the effectiveness of pDGE-RGD in preventing rotavirus cell infection parallel the data documented in a previous work (21), showing that a DGEA peptide was a less effective inhibitor of virus binding in Caco-2 cells than in MA104 cells. The higher surface expression of  $\alpha 2\beta 1$  on Caco-2 cells than on MA104 cells (51) could explain these findings, because higher levels of peptide would be needed to block all available integrin sites that could bind virus.

During our investigation of the functional roles of the equine lactadherin RGD and DGE sequences in inhibition of rotavirus infection, we found that pAGE-RGD, pDGE-RGE, and pAGE-RGE mutant peptides lack antiviral activity on MA104 cells. Interestingly, we were able to discriminate a different contribution from the two motifs using CaCo2 cells, with the DGE triplet being more critical than RGD in limiting Wa infectivity (IC $_{50}$  of pAGE-RGD > IC $_{50}$  of pDGE-RGE). These data, together with the fact that peptides harboring single amino acid substitutions outside these functional motifs retain their antiviral activity, suggest that both triplets are important for the

<sup>&</sup>lt;sup>4</sup> A. Civra, M. G. Giuffrida, M. Donalisio, L. Napolitano, Y. Takada, B. S. Coulson, A. Conti, and D. Lembo, unpublished observations.



**TABLE 5** Effect of lactadherin-derived peptides on cell viability and Wa rotavirus infectivity in Caco-2 cells

			95% confidence		
Peptides	Features	IC <sub>50</sub>	interval	CC <sub>50</sub>	SI
		$\mu_M$		$\mu$ M	
pDGE-RGD	Wild type peptide	8.8	6.1-12.5	945.1	102.7
pDGE-RGE	aa substitution in position 16	46.0	24.9-84.9	>880	19.1
pAGE-RGD	aa substitution in position 3	101.7	78.6-131.6	>880	8.7
pAGE-RGE	aa substitutions in position 3 and 16	152.5	123.3-188.6	>880	5.8

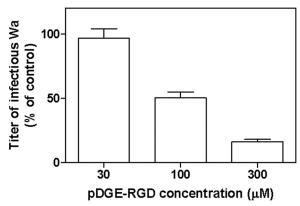
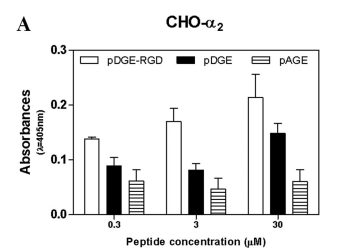


FIGURE 6. Effect of peptide pDGE-RGD on multiple cycles of Wa replication. Cells were treated after virus infection at the given peptide concentrations. The titer of rotavirus in the treated samples is expressed as a percentageof the titer in the untreated control. Error bars, S.D.

inhibition of rotavirus infectivity. These specific substitutions (*i.e.* Asp  $\rightarrow$  Ala for DGE and Asp  $\rightarrow$  Glu for RGD) were chosen because an aspartate or glutamate residue is a critical feature of all integrin recognition sites (52). The amino acid substitution in the DGE motif (i.e. Asp  $\rightarrow$  Ala) is known to alter the integrin  $\alpha 2\beta 1$  binding ability of VP5\* (21, 24, 32), with Asp-308 in VP5\* being the major requirement for  $\alpha 2I$  binding by rotavirus. Overall, these results suggest that the antiviral activity of pDGE-RGD is dependent on its ability to interact with  $\alpha 2\beta 1$ integrin. This notion is supported further by the finding that infection by integrin-independent virus CRW-8 was unaffected by pDGE-RGD.

The demonstration of the antiviral efficacy of pDGE (an 8-aa peptide harboring only the DGE triplet) extends data reported previously (21), which showed that individual peptide polymers containing the collagen-binding motif DGEA inhibit rotavirus infectivity by more than 60% in Caco-2 and MA104 cells and provided additional evidence of the importance of  $\alpha 2\beta 1$  in the rotavirus-cell attachment and entry process. We also tested the antiviral efficacy of DGEA peptide in our experimental systems, confirming the efficacy previously reported against the Wa strain (21) and showing that pDGE has a greater inhibitory capacity. Notably, an 8-aa mutant peptide (pAGE) showed a significant anti-rotaviral activity only at high concentrations. Taken together, these results confirm that the antiviral activity of pDGE-RGD peptide is mostly dependent on the integrity of the DGE triplet. Nevertheless, pDGE shows a less effective antiviral activity than pDGE-RGD, suggesting that a second region of this peptide provides an additional contribution in inhibiting rotavirus infectivity. To this end, we showed that 8-aa RGD peptides (namely pRGD-Hu, pRGD-Bo, and pRGD-Eq derived from human, bovine, or equine lactadherin, respectively) are endowed with a slight antiviral efficacy. Published studies pro-



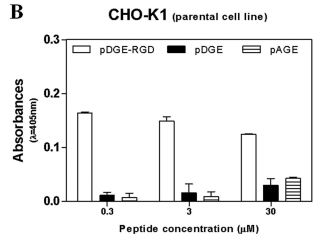


FIGURE 7. Evaluation of the ability of biotinylated pDGE-RGD, pDGE, and **pAGE to interact with the**  $\alpha_2\beta_1$  **integrin.** Experiments were performed with CHO cells expressing  $\alpha_2\beta_1$  (CHO- $\alpha_2$ ; A) and the parental CHO-K1 cells lacking this integrin (B). Levels of peptide-cell binding are expressed in absorbance units ( $\lambda = 405$  nm). *Error bars*, S.D. of three independent experiments.

vide evidence that rotavirus infectivity is not inhibited by RGD peptide, and function-blocking antibodies to RGD-binding integrins do not inhibit rotavirus infection (20, 23, 53). No evidence for Wa binding to integrins via RGD is available. We speculate that the slight antiviral efficacy of RGD peptides may be due to an indirect effect on rotavirus cell binding, perhaps by anchoring pDGE-RGD peptide to the cell surface and sterically blocking Wa virus access to receptors.

The results of the virucidal assay rule out the possibility of a direct effect of peptide treatment on rotavirus particles, whereas the time of addition assays suggest that pDGE-RGD might function by hindering the attachment of the virus to the host cell. The results of rotavirus cell-binding experiments confirm this conclusion; the titer of infective particles attached on

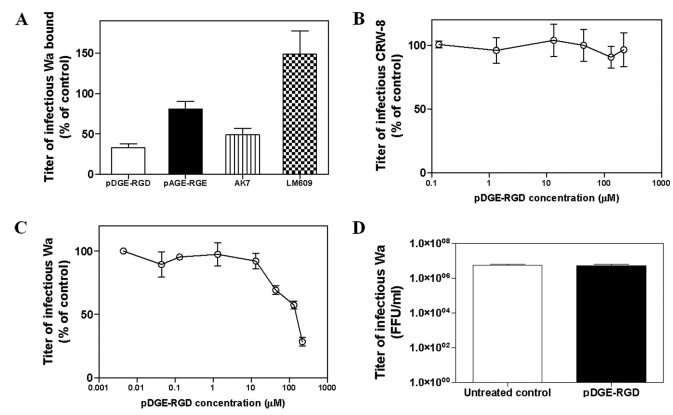


FIGURE 8. **Evaluation of the mechanism of action of pDGE-RGD.** A, effect of pDGE-RGD on Wa binding to the MA104 cell surface; control experiments were performed by treating cells with pAGE-RGE, anti- $\alpha_2$  antibody AK7 or anti- $\alpha_v\beta_3$  antibody LM609. On the y axis, the infectious titer of Wa bound to cells is expressed as a percentage of the titer bound to control MA104 cells in the absence of peptides or antibodies. B, effect of pDGE-RGD on CRW-8 infection in MA104 cells. The infectivity titer of virus in the treated sample is expressed as a percentage of the titer obtained in the absence of peptide (untreated control). C, effect of pDGE-RGD on Wa rotavirus entry into MA104 cells; the rotavirus titer measured in the treated samples is expressed as a percentage of the titer obtained in the untreated control. Evaluation of the virucidal effect of pDGE-RGD on infectious rotavirus particles is shown in D. On the y axis, the Wa infectious titers are expressed as focus-forming units/ml (*FFU/ml*). in the three graphs, *error bars* represent the S.D. values of three independent experiments.

the surface of pDGE-RGD-treated cells is significantly reduced compared with the untreated control. Moreover, pAGE-RGE peptide does not reduce the number of infectious particles attached to the cell surface, showing that this ability is strictly dependent on the integrity of the DGE or RGD sequences. Consistent with these considerations, we demonstrated that both pDGE and pDGE-RGD bind  $\alpha_2\beta_1$  integrin in a DGE-dependent manner, because pAGE does not interact with  $\alpha 2\beta 1$ -expressing CHO-K1 cells. The pDGE and pDGE-RGD peptides interact in a dose-dependent manner with  $\alpha 2\beta 1$  expressed on the surface of CHO-K1 cells. Importantly, pDGE-RGD bound the cellular surface to a greater extent than pDGE, which parallels the higher antiviral activity of the longer peptide. Moreover, we show that pDGE-RGD (unlike pDGE) is able to interact with the surface of the CHO-K1 parental cell line that does not express  $\alpha 2\beta 1$  on the cellular surface. This result suggests that pDGE-RGD interacts with a second surface determinant, a clue that could explain the greater antiviral activity of pDGE-RGD compared with pDGE. Of the known RGD-binding integrins, CHO-K1 cells express  $\alpha 5\beta 1$  and  $\alpha v\beta 5$  but not  $\alpha 3\beta 1$  or  $\alpha v\beta 3$ (54, 55). It is likely that pDGE-RGD binds  $\alpha$ 5 $\beta$ 1 or  $\alpha$ v $\beta$ 5 on CHO-K1 cells and  $\alpha 2\beta 1$ -expressing CHO-K1 cells as well as binding  $\alpha 2\beta 1$  on  $\alpha 2\beta 1$ -expressing CHO-K1 cells. The close proximity of the RGD and DGE motifs in pDGE-RGD is expected to preclude binding to both  $\alpha 2\beta 1$  and an RGD-recognizing integrin simultaneously by a single peptide molecule.

However, peptide anchorage to an RGD integrin like  $\alpha 5\beta 1$ , which is abundant on CHO-K1 cells, could prevent virus access to  $\alpha 2\beta 1$  and possibly other receptors. Interestingly, rotaviruscell entry experiments showed that pDGE-RGD is able to inhibit rotavirus infectivity when added after virus-cell attachment, even if only slightly and at high concentrations. One possibility is that high concentrations of pDGE-RGD could interfere with virus-cell attachment, even after it took place, by mediating detachment of part of the viral population from the cell surface. It is also possible that integrin-bound pDGE-RGD is endocytosed and inhibits virion conversion into the transcriptionally active, double-layered particle, which for Wa has been proposed to occur during exit from late endosomes (56). Interestingly, Wa VP8\* binds the GM1 ganglioside, which can associate with integrins such as  $\alpha 5\beta 1$  and  $\alpha 2\beta 1$  in membrane microdomains (57, 58) and has been proposed to classify cargo for uptake and trafficking in late endosomes (59). Binding of pDGE-RGD to β1 integrins internalized in endosomes also might interfere with Wa access to the cytoplasm, which is essential for replication (60).

Overall, the information obtained in the present study indicates that a specific 20-aa peptide from the N-terminal domain of donkey lactadherin is able to block rotavirus cell adhesion and infectivity in a DGE-dependent manner. Previously reported data (21) show that polymerization increases the effectiveness of peptide polymers containing the collagen-bind-

ing motif DGEA as rotavirus inhibitors by up to 10-fold, suggesting that it might be possible to develop more effective inhibitors of rotavirus-integrin interactions. In our experimental system, we also tested the in vitro antiviral activity of a dendrimeric peptide harboring multiple DGE motifs; this molecule displays an IC<sub>50</sub> (3.6  $\mu$ M) that is not significantly different from the pDGE-RGD IC<sub>50</sub> (p > 0.05; data not shown). Currently, no antivirals are commercially available for rotavirus. The discovery of the pDGE-RGD peptide may prove useful in the development of inhibitors of receptor recognition by rotavirus or other viruses that use  $\alpha 2\beta 1$  integrin during cell attachment and entry like human echovirus 1 (61, 62).

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