Expression of Integrin- α_F by Mucosal Mast Cells in the Intestinal Epithelium and Its Absence in Nematode-Infected Mice Lacking the Transforming Growth Factor- β_1 -Activating Integrin $\alpha_{\text{v}}\beta_{6}$

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Peak intestinal mucosal mast cell (MMC) recruitment coincides with expulsion of *Trichinella spiralis***, at a time when the majority of the MMCs are located within the epithelium in BALB/c mice. Although ex**pression of integrin- $\alpha_{\rm E}\beta_7$ by MMCs has not been for**mally demonstrated, it has been proposed as a potential mechanism to account for the predominantly intraepithelial location of MMCs during nematode in**fection. Co-expression of integrin- $\alpha_{\rm E}\beta_{7}$ and the MMC **chymase mouse mast cell protease-1, by mouse bone marrow-derived mast cells, is strictly regulated by** transforming growth factor (TGF)- β_1 . However, $TGF-\beta_1$ is secreted as part of a latent complex *in vivo* **and subsequent extracellular modification is required to render it biologically active. We now show, for the first time, that intraepithelial MMCs express integrin-** $\alpha_{\rm E}\beta_{7}$ in *Trichinella* -infected BALB/c and S129 mice. In $S129$ mice that lack the gene for the integrin- β_6 sub**unit and, as consequence, do not express the epithe**lial integrin- $\alpha_{\rm v}\beta_6$, integrin- $\alpha_{\rm E}$ expression is virtually **abolished and recruitment of MMCs into the intestinal epithelium is dramatically reduced despite significant overall augmentation of the MMC population. Because a major function of integrin-v**-**⁶ is to activate latent** TGF- β_1 , these findings strongly support a role for TGF- β_1 in both the recruitment and differentiation of **murine MMCs during nematode infection.** *(Am J Pathol 2004, 165:95–106)*

Mast cell recruitment is a common feature of helminth infections and contributes to the immunological expulsion of some, but not all, gastrointestinal nematode parasites.^{1–3} Mast cells are, in addition, thought to play a significant role in allergic responses in the airways and gastrointestinal tract^{4,5} and in the immune response to bacterial pathogens.⁶ Although the role of mast cells in the immune expulsion of gastrointestinal nematodes remains incompletely resolved, experimental studies in rats and mice suggest that intestinal mucosal mast cell (MMC) -derived β -chymases increase intestinal epithelial permeability to macromolecules.^{7,8} MMCs may, therefore, contribute to the expulsion of gastrointestinal nematodes by facilitating the pathotopic translocation of effector proteins, including immunoglobulins, into the gut lumen.2 The predominantly intraepithelial location of mouse MMCs at the time of nematode expulsion is consistent with this hypothesis.⁹

Previous studies, by our group, have shown that expression of the MMC-specific β -chymase mouse mast cell protease-1 (mMCP-1) is strictly regulated by the multifunctional cytokine, transforming growth factor (TGF)- β_1 .¹⁰ Indeed, mouse bone marrow-derived mast cells (mBMMCs) cultured in TGF- β_1 together with interleukin (IL)-3, IL-9, and stem cell factor are highly homologous to MMCs.^{10–13} In agreement with studies on T lymphocytes,^{14,15} mBMMC expression of the $\alpha_{\rm E}$ chain of integrin- $\alpha_{\rm E}\beta_7$ is also TGF- β_1 -dependent.^{13,16} The presence of integrin- $\alpha_\mathsf{E}\beta_7$ has been demonstrated on the surface of intraepithelial T lymphocytes *in vivo*, 15,17 but its expression by MMCs has not been previously demonstrated, although antibody-blocking studies provide indirect evidence that α_F is required for recruitment and survival of MMCs within the epithelium.18 Given that the ligand for integrin- $\alpha_{\rm E}\beta_7$, E-cadherin,¹⁹ is expressed exclusively by epithelial cells,²⁰ TGF- β_1 -mediated expression of integrin- $\alpha_{\rm E}\beta_7$ by mMCP-1-positive mBMMCs¹³ is consistent with the predominantly intraepithelial location of mMCP-1-positive MMCs during the immune rejection of gastrointestinal nematodes.⁹ It therefore seems highly likely that, like intestinal epithelial T lymphocytes,¹⁵ intraepithelial MMCs express integrin- $\alpha_{\rm E}\beta_7$ through a TGF- β_1 dependent mechanism and that the interaction between integrin- $\alpha_{\rm E}\beta_7$ and E-cadherin contributes to their recruitment into the intestinal epithelium during nematode infection.

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TGF- β_1 is ubiquitously expressed in most tissue microenvironments but is secreted as a latent complex with latency-associated protein, derived from the N-terminal sequence of the TGF- β_1 proprotein.²¹ We hypothesize that the interaction of the TGF- β_1 -latency-associated protein complex with integrin- $\alpha_{\rm v}\beta_{\rm 6}$ and the subsequent presentation of active TGF- β_1 on the surface of epithelial cells²² is a mechanism of activating latent TGF- β_1 that is likely to be important for MMC recruitment and differentiation during gastrointestinal nematode infection. In support of this hypothesis, we have previously shown that integrin- β_6 -null mutation (β_6 ^{-/-}) mice²³ are severely compromised in their ability to mount a MMC response during infection with *Nippostrongylus brasiliensis*. ²⁴ However, the rat-adapted strain of *N. brasiliensis* used in these studies stimulates relatively poor MMC recruitment in the mouse (PA Knight, unpublished observation).²⁴ In the current study, $\beta_6^{-/-}$ and $\beta_6^{+/+}$ mice were infected with the epithelium-dwelling gastrointestinal nematode *Trichinella spiralis,* to generate a maximal MMC response. The specific aims of the current study were: 1) to formally demonstrate that MMCs express integrin-α_Εβ₇ in vivo; 2) to analyze the effects of the integrin- β_{6} -null mutation on the TGF- β_1 -dependent expression of the integrin- $\alpha_{\rm E}$ subunit; and 3) to determine whether MMC recruitment is attenuated by the integrin- β_{6} -null mutation.

Materials and Methods

Unless stated otherwise, reagents were purchased from Fisher Scientific UK, Loughborough, UK.

Antibodies

Purified and fluorescein isothiocyanate (FITC)-conjugated monoclonal rat $I_{\text{g}}G_1$ anti-mouse IgE (clone 23G3; SouthernBiotech, Birmingham, AL) were purchased from Cambridge BioScience, Cambridge, UK. Purified monoclonal rat IgG_{2a} anti-human integrin- α_6 (clone GoH3), control rat $\log G_{2a}$ (clone R35-95), control rat $\log G_1$ (clone R3-34), and FITC-conjugated control rat \log_1 (clone R3-34) were purchased from BD Biosciences, Cowley, UK. Purified rat lgG_{2a} anti-integrin- $\alpha_{\rm E}$ (clone M290) and anti-integrin- β_7 hybridoma supernatant (clone M293) were provided by Dr. Peter Kilshaw (The Babraham Institute, Babraham, Cambridge, UK).^{14,25} Rat anti-mMCP-1 monoclonal $\lg G_1$ (clone RF6.1)⁹ and rabbit polyclonal anti-equine tryptase immunoglobulin (Ig)²⁶ were purified using affinity columns (Hi-Trap NHS-activated, 1 ml; Amersham Biosciences, Little Chalfont, UK) to which 1 mg of purified protein had been coupled according to the manufacturer's instructions. Alkaline phosphatase-conjugated mouse anti-rabbit IgG (clone RG-96) and purified control rabbit IgG were purchased from Sigma-Aldrich Company, Poole, UK. Proliferating cell nuclear antigen (PCNA)-specific and negative control DAKO EPOS conjugates were purchased from DakoCytomation Ltd., Ely, UK. Fluorophore conjugated monovalent goat antirat IgG and anti-rabbit IgG Fab fragments (Jackson ImmunoResearch Laboratories, West Grove, PA) were purchased from Stratech Scientific, Soham, UK.

Western Blot Analysis of Mouse Bone Marrow-Derived Mast Cells with Rabbit Anti-Equine Tryptase Immunoglobulin

Bone marrow cells, prepared from 10- to 12-week-old male BALB/c mice, were cultured for 12 days *ex vivo* in the presence of recombinant human TGF- β 1 (1 ng/ml; Sigma-Aldrich), recombinant mouse IL-3 (1 ng/ml; R&D Systems, Abingdon, UK), recombinant mouse IL-9 (5 ng/ml; R&D Systems), and recombinant mouse stem cell factor (50 ng/ml; PeproTech EC Ltd., London, UK).¹¹ The resultant mouse mBMMC cultures were extracted with 20 mmol/L of Tris-HCl, pH 7.5, containing 1 mol/L NaCl. Sodium dodecyl sulfate gels (12%) (Mini-Protean-II; Bio-Rad Laboratories, Hemel Hempstead, UK) were run containing molecular weight markers (Bio-Rad Laboratories) and mBMMC extract (equivalent of 5×10^4 cells/lane). Gels were either stained with Coomassie Brilliant Blue (Bio-Rad Laboratories), or transferred to polyvinylidene difluoride membrane (Immobilon-P; Millipore, Watford, UK) and probed with 1 μ g/ml of affinity-purified rabbit anti-equine tryptase \lg^{26} followed by alkaline phosphatase-conjugated mouse anti-rabbit IgG (1/20,000 dilution). Blots were developed using 5-bromo-4-chloro-3 indolyl phosphate/nitro blue tetrazolium (BCIP/NBT; Sigma-Aldrich).

Parasite Infections

All experiments involving laboratory animals were performed in accordance with the United Kingdom's Animals (Scientific Procedures) Act 1986. Integrin- β_{6} -null (S129 strain background; S129 $\beta_6^{-/-}$) mice²³ were originally obtained from Dr. Kairbaan Hodivala-Dilke (Cell Adhesion and Disease Laboratory, GKT School of Medicine, St. Thomas' Hospital, London, UK) and backcrossed with $S129 \beta_6$ ^{+/+} controls (B&K Universal, Hull, UK). Breeding colonies of S129 $\beta_6^{-/-}$, S129 $\beta_6^{+/+}$, and BALB/c mice were maintained at the Easter Bush Veterinary Centre animal facility. Maintenance, infection, and recovery of *Trichinella spiralis* larvae were performed using standard methods.27 Mice, 8- to 15-week-old, age- and sexmatched BALB/c (B&K Universal, Hull, UK), S129 $\beta_6^{+/+}$ (B&K), and S129 $\beta_6^{-/-}$, were infected by gavage with 250 muscle larvae in 0.2 ml of phosphate-buffered saline (PBS)/0.1% agar, freshly isolated from muscle cysts from 30- to 90-day infected BALB/c mice. Groups of mice (*n* 4) were killed on days 7 and 13 (S129) or 14 (BALB/c) after infection. Adult worms were isolated from the small intestine of S129 mice using a modified Baerman's technique¹ and samples of jejunum were prepared as described below. Samples were also prepared from agematched uninfected controls.

Isolated Jejunal Epithelial Whole Mounts

Intact sheets of jejunal epithelium that included both villi and crypts were isolated by vascular perfusion with ethylenediaminetetraacetic acid¹² from 8- to 15-week-old female BALB/c mice 14 days after infection with *T. spira-* *lis*, and from age- and sex-matched uninfected controls. Isolated epithelium was allowed to settle in ice-cold PBS before transfer into -20° C absolute methanol. After 20 minutes, small volumes of methanol/epithelium suspension were air-dried onto Snow Coat X-tra-charged slides (Surgipath Europe, Peterborough, UK). Samples were rehydrated and washed in three changes of PBS before proceeding with immunocytochemistry.

Jejunal Samples for Histochemistry, Immunocytochemistry, and RNA Analysis

Age- and sex-matched S129 $\beta_6^{-/-}$ and $\beta_6^{+/+}$ mice (n = 4) were killed on days 7 and 13 after infection with *T. spiralis*, along with uninfected controls. Freshly isolated samples of jejunum were collected into RNA-Later (Ambion, Huntingdon, UK) for RNA isolation (0.5 cm) and into Carnoy's fixative for histochemistry (3 cm) .¹ Samples of jejunum (3.0 cm) from mice killed on day 13 were embedded in OCT compound (BDH Laboratory Supplies, Dorset, UK) and snap-frozen in dry ice-chilled isopentane. Adult worms were isolated from the reminder of the small intestine.¹ Snap-frozen samples were also prepared from age-matched female BALB/c mice $(n = 4)$ 14 days after infection with *T. spiralis*, and from uninfected controls. For immunocytochemistry, 10 - μ m cryostat sections from frozen jejunal samples were mounted on Snow Coat X-tra-charged slides, air-dried for 10 minutes, and stored at -70° C. For histochemical mast cell evaluation, Carnoy's-fixed, paraffin-embedded $4-\mu m$ microtome sections of jejunum were stained overnight in 0.5% toluidine blue in 0.5 mol/L HCl, pH 0.5, and counterstained with 1% eosin solution. Toluidine blue-stained sections prepared from S129 $\beta_6^{-/-}$ and $\beta_6^{+/+}$ mice on day 13 were also washed thoroughly with PBS and probed with anti-PCNA or control EPOS antibodies as directed by the manufacturer (DakoCytomation Ltd.). PCNA-specific labeling was visualized with 3,3-diaminobenzidine substrate (Vector Laboratories Ltd., Peterborough, UK).

Fixation and 3,3-Diaminobenzidine Pretreatment of Frozen Sections

Frozen sections were thawed for 10 minutes at 21°C, fixed in absolute methanol for 10 minutes at -20° C, and air-dried for a further 10 minutes under forced air. Sections were washed and rehydrated in pH 7.4 PBS containing 0.5% Tween 80 (Sigma-Aldrich). Eosinophil autofluorescence was quenched using a modified version of the procedure described by Kingston and Pearson.²⁸ Noneosinophil-derived endogenous peroxidase activity was quenched for 30 minutes with 1% H₂O₂ in pH 7.4 PBS containing 0.5% Tween 80. Sections were then washed with PBS and incubated for 3 minutes with 3,3 diaminobenzidine substrate. Sections were washed thoroughly with PBS before proceeding with immunocytochemistry.

Immunocytochemistry: Standard Conditions and Reagents

Unless stated otherwise, immunocytochemical procedures were performed at 21°C under humidified conditions in a Sequenza immunostaining center (Thermo Shandon, Runcorn, UK). Antibodies and blocking sera were diluted in staining buffer (pH 7.4 PBS containing 0.5 mol/L NaCl and 0.5% Tween 80). After immunocytochemical labeling, samples were washed in PBS and mounted with no. 1.5 (0.17-mm-thick) glass coverslips (BDH Laboratory Supplies) using Mowiol (pH 8.5) mounting media (EMD Biosciences, San Diego, CA).

IgE and mMCP-1 Dual-Immunofluorescent Labeling in Isolated Jejunal Epithelium

Nonspecific immunoglobulin interactions were blocked for 1 hour with staining buffer containing 10% heat-inactivated normal mouse serum (NMS). Samples were then incubated for 30 minutes with rat I_{QG_1} anti-mMCP-1 or control rat IgG₁, diluted to 5 μ g/ml in staining buffer and 10% NMS. After washing with PBS, slides were incubated for 30 minutes with Rhodamine Red-X (RRX)-conjugated goat anti-rat IgG Fab fragments, diluted to 4 μ g/ml in staining buffer and 10% NMS. Samples were washed in PBS, blocked for 1 hour in staining buffer containing 10% heat-inactivated normal rat serum (NRtS), and then incubated for 30 minutes with FITC-conjugated rat \log_1 antimouse I gE or FITC-conjugated control rat I g $G₁$, diluted to 5μ g/ml in staining buffer and 10% NRtS. Samples were then washed with PBS and counterstained for 10 minutes with DRAQ5 (Biostatus Limited, Shepshed, Leicestershire, UK), diluted to 5 μ mol/L in PBS.

Triple-Immunofluorescent Labeling of Frozen Sections

Nonspecific immunoglobulin interactions were blocked for 1 hour with staining buffer and 10% NMS. Sections were then incubated for 1 hour with staining buffer and 10% NMS containing rat anti-mouse integrin- α_{F} IgG_{2a} (2) μ g/ml), integrin- β_7 Ig $\rm{G_{2a}}$ (1/50 dilution of hybridoma supernatant), control rat \log_{2a} (2 μ g/ml), rabbit anti-equine tryptase Ig (2 μ g/ml), or control rabbit IgG (2 μ g/ml). Samples were washed with PBS and incubated for 30 minutes with staining buffer and 10% NMS containing Cyanine-5 (Cy5)-conjugated goat anti-rat or goat antirabbit Fab fragments (2 μ g/ml) as appropriate.

IgE and integrin- α_6 were labeled simultaneously using Fab fragment complexes. Samples were blocked for 1 hour with staining buffer and 10% NRtS. During this time, primary rat antibodies were incubated for 20 minutes at 21°C with fluorophore-conjugated anti-rat IgG Fab fragments in a small volume, 10 μ l per 1 μ g of primary antibody, of staining buffer in a microcentrifuge tube. Rat anti-mouse IgE and control $IgG₁$ were labeled with FITCconjugated Fab fragments at a ratio of 5 μ g of Fab fragments per μ g of primary rat IgG₁. Rat anti-human integrin- α_6 and control IgG_{2a} were labeled with RRXconjugated Fab fragments at a ratio of 5 μ g of Fab fragments per μ g of primary rat IgG_{2a}. The resultant Fab fragment complexes were diluted to 4 μ g/ml of primary antibody with staining buffer and 10% NRtS and incubated for 10 minutes at 21°C to block any unbound Fab fragment paratopes. Tissue sections were then incubated for 1 hour with the following combinations of Fab fragment complexes (mixed 1:1 to yield a final concentration of 2 μ g/ml of each primary/ control antibody): IgE-specific Fab FITC and integrin- α_6 -specific Fab RRX; IgE-specific Fab FITC and isotype-matched control Fab RRX; isotype-matched control-Fab FITC and integrin- α_6 -specific Fab RRX; and isotype-matched control Fab FITC and isotypematched control Fab RRX.

Microscopy and Imaging

Bright-field images were acquired with a Sony DXC-390P 3CCD color video camera (Scion Corp., Frederick, MD) mounted on an Axiovert 100 inverted microscope (Carl Zeiss, Welwyn Garden City, UK). The RGB video signal from the camera was digitized using Scion Image (Scion Corp.) installed in a G4 Macintosh computer (Apple Computer, Cupertino, CA) fitted with a CG-7 frame grabber (Scion Corp.).

Fluorescent images were acquired using an MRC-600 confocal laser-scanning microscope (CLSM; Bio-Rad Laboratories) mounted on an Axiovert 100 inverted microscope equipped with Plan-Apochromat objective lenses (Carl Zeiss). Fluorophores were excited and imaged sequentially using the 488-nm (FITC), 568-nm (RRX), and 647-nm (DRAQ5 and Cy5) lines from a 15 mW Kr/Ar laser (Bio-Rad Laboratories).

Toluidine blue-based mast cell counts were determined from 20 jejunal villus-crypt units (VCU) per mouse. The proportion of mast cells expressing PCNA in samples from ${\beta_6}^{-/-}$ and ${\beta_6}^{+/+}$ S129 mice on day 13 after infection was determined by evaluation of 200 toluidine blue-positive mast cells per mouse. Manual cell counting and stereology measurements (Cavalieri method)²⁹ were performed on images of fluorescently labeled samples using custom software developed for Object-Image.30 Object-Image is a public domain software package, based on NIH Image,³¹ developed by Norbert Vischer (The University of Amsterdam, Amsterdam, The Netherlands) and is available via the Internet at *http://simon.bio.uva.nl/objectimage.html*. Images were prepared for publication using Object-Image and Photoshop [Adobe Systems (UK), Uxbridge, UK].

Detection of Integrin Transcripts by Semiquantitative Reverse Transcriptase-Polymerase Chain Reaction (RT-PCR)

Total RNA was extracted from RNA-Later (Ambion)-fixed jejunal samples using Tri-Reagent (Sigma-Aldrich) and contaminating DNA removed using DNA-free DNase (Ambion).1 One mg of RNA was reverse-transcribed using 2.5 mmol/L $(dT)_{15}$. A one-twentieth volume was amplified by PCR using gene-specific primers for α_F [AED (sense), CCCTTACTCTCCTTAGGACGATCAA; AEF (anti-sense), TATCAGTCATCAAAACGCATG] that give a RT-PCR product of 197 bp, and β_7 [B7D (sense), GCTCTCTGTGGAAATCTACGA, B7F (anti-sense), TCACTCTGAAAAATCTCAGCG) that give a RT-PCR product of 278 bp, or for the housekeeping gene $GAPDH₁¹$ with equivalent quantities of nonreversetranscribed RNA as negative controls. Reaction conditions were optimized to ensure the number of thermocycles used coincided with the amplification phase of the PCR. Amplifications were performed for 40 seconds at 94°C, 40 seconds at 55°C, and 120 seconds at 72°C for 34 thermocycles for $\alpha_{\rm E}$ and β_7 , and for 40 seconds at 94°C, 40 seconds at 60°C, and 120 seconds at 72°C for 32 thermocycles for GAPDH, in a final magnesium concentration of 1.5 mmol/L, pH 8.3. Resultant PCR products were visualized on ethidium bromide-stained 1.6% agarose gels. Images were acquired with a Kodak Digital Science Image Station 440CF (Eastman-Kodak, Rochester, NY) and analyzed using Kodak 1D Image Analysis software. PCR product identities were confirmed by Southern blotting following standard protocols and hybridized using digoxigenin-labeled gene-specific probes.³² Internal probe sequences for the $\alpha_\mathrm{E}/\beta_7$ RT-PCR products were as follows; α_{F} ; AEE (sense), AGTGCCTTTAATGAGCACG-GTT; β_7 ; B7E (sense), ACTGGAAGCAGGACAACAAT. Primers and internal complementary probes were designed using RawPrimer (The Virtual Genome Center, *http://alces.med.umn.edu/rawprimer.html*).

Results

Surface-Bound IgE as a Phenotypic Marker of Mast Cells in the Parasitized Intestine

Intraepithelial IgE^{+ve} Cells

mMCP-1 remains highly soluble and diffuses in frozen sections resulting in poor immunocytochemical labeling (data not shown). To address this problem and to confirm previous studies that identified MMCs on the basis of IgE staining,33,34 jejunal epithelial whole mounts from *T. spiralis*-infected (day 14) BALB/c mice $(n = 4)$ were probed with mMCP-1- and IgE-specific antibodies. The distribution of mMCP-1 and IgE staining within the jejunal epithelium confirmed that cell surface-bound IgE is exclusively restricted to mMCP-1^{+ve} MMCs (Figure 1, A to D; and Table 1). Although this method of sample preparation and fixation preserved the antigenicity of IgE and mMCP-1 (Figure 1; A to D) it was incompatible with integrin immunocytochemistry (data not shown). Subsequent analysis of MMC integrin- $\alpha_\mathsf{E}\beta_7$ expression was performed on 20°C methanol-fixed, snap-frozen sections of jejunum.

Figure 1. Immunocytochemical detection of MMCs in epithelial whole mounts and frozen sections of jejunum from *T. spiralis-*infected BALB/c mice. **A** to **D:** Whole mounts of jejunal epithelium were probed with anti-mMCP-1 (pseudo-colored red) or isotype-matched control antibodies together with anti-IgE (pseudo-colored green) or isotype-matched control antibodies and counterstained with DRAQ5 (pseudo-colored blue). IgE surface labeling was almost exclusively restricted to mast cells exhibiting strong intracellular mMCP-1 staining in samples probed with mMCP-1- and IgE-specific antibodies (**A** and **D**). Single-negative controls for IgE (**B**) and mMCP-1 (**C**) were used to confirm labeling specificity. Frozen sections of jejunum from uninfected (**E**) and *T. spiralis*-infected (**F**) BALB/c mice were probed with rabbit anti-tryptase (pseudo-colored green), anti-IgE (pseudo-colored red), and anti-integrin- α_6 antibodies (pseudo-colored blue). No IgE or tryptase-positive cel for tryptase (G), integrin- α_6 (H), and IgE (I), in which the two other antigen-specific antibodies were replaced with control IgG. Identical excitation and exposure settings were used to acquire images of multilabeled and control samples. Scale bar length (μm) is provided in the bottom left corner of each image.

IgE^{+ve} Cells in the Lamina Propria

IgE^{+ve} cells were enumerated in frozen sections of jejunum from *T. spiralis*-infected (day 14) BALB/c mice and uninfected controls. Although the majority of IgE^{+ve} cells within the jejunal mucosa of infected mice were located intraepithelially, substantial numbers of cells exhibiting IgE surface labeling were detected within the lamina propria (Table 2). To confirm that these cells were also mast cells, IgE staining was performed in conjunction with tryptase localization. Anti-equine tryptase²⁶ resulted in strong intragranular staining (Figure 1F) within $>$ 80% of IgE^{+ve} lamina propria cells and $>$ 60% of intra-

Exfoliated jejunal epithelium (20°C methanol-fixed) from *T. spiralis*infected BALB/c mice (day 14) was probed with anti-mMCP-1 and anti-IgE monoclonal antibodies (Figure 1; A to D). The relative frequencies of IgE single-positive (IgE only), mMCP-1 single-positive (mMCP-1 only), and IgE/mMCP-1 double-positive (IgE and mMCP-1) cells present in the epithelium were determined from a population of 500 cells/sample $(n = 4)$.

epithelial IgE^{+ve} cells (Table 2). Western blot analysis of lysates from mBMMCs¹¹ showed that this antibody recognizes a band of \sim 35 kd (Figure 2) consistent with the molecular weight (MW) of glycosylated mouse tryptases (mMCP-6 and -7).35 The relative frequencies of tryptase single-positive, IgE single-positive, and tryptase/IgE double-positive cells present in the epithelium and lamina propria were determined using integrin- α_6 -specific labeling to delineate the epithelium.

IgE^{+ve} or tryptase^{+ve} cells were not detected in uninfected controls (Figure 1E). Similarly, few if any, MMCs were detected in control uninfected mice by toluidine blue staining (data not shown). Nonintraepithelial tryptase^{-ve} cells accounted for 1.4 \pm 0.5% of the total population of IgE^{+ve} cells in the jejunal mucosa of infected mice on day 14 (Table 2). Given that only 0.4 \pm 0.2% of intraepithelial IgE^{+ve} cells are mMCP-1-negative on day 14 (Table 1), IgE labeling can be used to identify mast cells within the jejunal mucosa of parasitized mice with a \geq 95% degree of confidence. This is consistent with previous observations in rats and mice in which different fixation and staining techniques were used to show that IgE-bearing cells in the mucosa were mast cells and not plasma cells, macrophages, or lymphocytes.33,34

Table 2. Surface-Bound IgE as a Phenotypic Marker of Mast Cells in the Parasitized Intestine

Location	Labeling	Mean (%)	Standard error of the mean (%)
Epithelium	IgE only	30.6	1.9
	Tryptase only	2.3	0.6
	Dual	49.8	27
Lamina Propria	IgE only	1.4	0.5
	Tryptase only	1.3	0.3
	Dual	14.6	19

Frozen sections of jejunum (20°C methanol-fixed) from *T. spiralis*infected BALB/c mice (day 14) were triple-labeled with rabbit antiequine tryptase Ig (cross-reactive with mMCP-6 and/or -7, see Figure 2), anti-IgE, and anti-integrin α_6 antibodies (Figure; 1; F to I). Using integrin α_6 staining to delineate the epithelium, the relative frequencies of IgE single-positive (IgE only), tryptase single-positive (tryptase only), and tryptase/IgE double-positive (IgE and tryptase) cells present in the epithelium and lamina propria were determined from five randomly selected 0.31 -mm² fields of view per sample ($n = 4$).

Figure 2. Detection of mouse tryptase(s) with rabbit anti-equine tryptase polyclonal antibody in Western blots of mBMMC extracts. Samples and molecular weight standards (**MW**) were run on 12% sodium dodecyl sulfate gels. Gels were stained with Coomassie Brilliant Blue (MW and **lane 1**), or transferred to Immobilon-P membrane and probed with affinity-purified rabbit anti-equine tryptase antibody (**lane 2**). Subsequent incubations with alkaline phosphatase-conjugated mouse anti-rabbit IgG and BCIP/NBT substrate resulted in the detection of a band of \sim 35 kd, consistent with the expected molecular weight for glycosylated murine tryptase(s), mouse mast cells protease-6, and/or protease-7.

The Integrin- α_F *Subunit Is Preferentially Expressed on Intraepithelial IgE*-*ve Cells*

Frozen sections of jejunum from *T. spiralis*-infected (day 14) BALB/c mice and uninfected controls $(n = 4)$ were labeled with integrin- α_{E} - or - β_{7} -specific antibodies together with IgE- and integrin- α_6 -specific Fab fragment complexes. Using integrin- α_6 to differentiate the epithelium and lamina propria, the relative frequencies of IgE single-positive cells and IgE/integrin double-positive cells present in each location were determined and stereology was used to calculate lgE^{+ve} cells mm⁻² (Figure 3). Although integrin- α_{E} - and - β_{7} -positive cells were present, IgE^{+ve} cells were not found in sections from uninfected mice (data not shown). Infection with *T. spiralis* was associated with a substantial IgE^{+ve} cell recruitment, with the majority (79.1 \pm 1.5%) of IgE^{+ve} cells located within the epithelium (Figure 3A). Nearly all of the IgE^{+ve} cells present within the jejunal mucosa expressed integrin- β_7 , regardless of their location (Figure 3B). Integrin- α_F was expressed by 17.1 \pm 1.3% of lamina propria IgE^{+ve} cells and was detected on a significantly ($P <$ 0.01; unpaired Student's *t*-test with Welch correction) higher proportion (54.5 \pm 2.9%) of intraepithelial IgE^{+ve} cells (Figure 3B).

Figure 3. IgE^{+ve} cells in the jejunal mucosa of *T. spiralis*-infected mice express integrin- $\alpha_{\rm E}\beta_{7}$. Frozen sections of jejunum from infected BALB/c mice were triple-labeled with anti-integrin- $\alpha_{\rm E}$ or anti-integrin- β_7 together with anti-IgE and anti-integrin- α_6 (see Figure 4). Using integrin- α_6 staining to differentiate the epithelium (**filled bars**) and lamina propria (**open bars**), IgE^{+ve} cells mm⁻² (**A**) and the relative frequencies o integrin- α_E or integrin- β_7 (**B**) were determined for each location using five randomly selected 0.31-mm² fields of view per sample. The mean percentage of intraepithelial IgE^{+ve} cells ± 1 SEM is provided in parentheses (**A**). Data are expressed as mean values \pm 1 SEM ($n = 4$). Differences observed between the epithelial and lamina propria compartments were significant at $P\leq$ 0.01 (**).

Deletion of the Integrin--*⁶ Gene Virtually* Abolishes Integrin- α_E Expression by IgE^{+ve} Cells *and Inhibits Their Recruitment into the Jejunal Epithelium*

Frozen sections of jejunum $(n = 3)$ from *T. spiralis*-infected (day 13) and uninfected control S129 $\beta_6^{+/-}$ and S129 β_6^{-1} mice ($n = 4$, frozen material from one of the S129 $\beta_6^{-/-}$ mice was damaged during processing) were labeled with integrin- $\alpha_{\rm E}$ - or - β_7 -specific antibodies together with IgE- and integrin- α_{ϵ} -specific Fab fragment complexes (Figure 4). IgE^{+ve} cells were not detected in sections from uninfected mice (Figure 4, A to D). Jejunum from uninfected S129 $\beta_6^{+/+}$ mice contained abundant intraepithelial integrin- α_{E}^{+ve} (Figure 4C) and integrin- β_7 ^{+ve} cells (Figure 4A). Integrin- β_7 ^{+ve} cells were also present in large numbers in the jejunum of uninfected S129 β_6 ^{-/-} mice but were predominantly restricted to the lamina propria (Figure 4B). Integrin- $\alpha_{E}^{+\rm ve}$ cells were extremely rare in samples from uninfected S129 $\beta_6^{\, -/-}$ mice and expressed substantially lower levels of integrin- α_F (Figure 4D).

Infection with T. spiralis was associated with IgE^{+ve} cell recruitment in both S129 $\beta_6^{+/-}$ and S129 $\beta_6^{--/-}$ mice (Figure 4E and Figure 5H). However, the frequency and distribution of IgE^{+ve} cells differed markedly in the two genotypes (Figure 5A). As observed in BALB/c mice (Figure 3A), IgE^{+ve} cell counts per unit area did not differ significantly between the epithelium and lamina propria in S129 $\beta_6^{+/-}$ mice (Figure 5A), despite the fact that the 5129 p_6 more (rigule 5A), despite the fact that the majority of the IgE^{+ve} cell population (74.1 \pm 0.6%) was located intraepithelially (Figure 5B). IgE^{+ve} cell hyperplasia also occurred in S129 $\beta_6^{-/-}$ mice. However, only 18.0 \pm 2.5% of IgE^{+ve} cells were located intraepithelially (Figure 5B), and there was a significant reduction (*P* 0.01, unpaired Student's *t*-test with Welch correction) in the population density of intraepithelial IgE^{+ve} cells compared to that in S129 β_6 ^{+/+} mice (Figure 5A). By contrast, the population density of IgE^{+ve} cells, when compared with infected $\beta 6^{+/+}$ controls, increased by more than fivefold in the lamina propria (Figure 5A) and by more than 2.5-fold in the jejunal mucosa as a whole $(P < 0.01$, unpaired Student's *t*-test with Welch correction).

The pattern of integrin- $\alpha_{\rm E}$ expression in S129 $\beta_{\rm 6}^{\,++}$ mice was similar to that observed in BALB/c mice (Figure 3B), with 64.4 \pm 8.0% of epithelial and 25.2 \pm 7.2% of lamina propria IgE^{+ve} cells exhibiting strong integrin- $\alpha_{\rm E}$ staining (Figure 5C). Integrin- α_{F} -positive cells were rare in sections from S129 $\beta_6^{-/-}$ mice and were typically IgE^{-ve} (Figure 4H). Only 0.2 ± 0.2 % of epithelial and $1.2 \pm 0.3\%$ of lamina propria IgE^{+ve} cells showed evidence of integrin- $\alpha_{\rm E}$ staining in S129 ${\beta_{\rm 6}}^{-/-}$ mice (Figure 5C). Virtually all IgE^{+ve} cells, regardless of their location, expressed integrin- β_7 in both S129 $\beta_6^{+/+}$ and S129 $\beta_6^{-/-}$ mice (Figure 5D).

Deletion of the Integrin--*⁶ Gene Is Associated with Reduced Levels of Integrin-* α_F *Transcripts and Aberrant Mast Cell Distribution in the Murine Jejunum*

Levels of integrin- α_E and integrin- β_7 transcripts were assessed by RT-PCR in jejunal RNA from *T. spiralis*-infected (day 13) and uninfected control S129 $\beta_6^{+/-}$ and S129 β_6 ^{-/-} mice ($n = 4$). Although there was no evidence for a reduction in the levels of integrin- β_7 or GAPDH control transcripts in S129 $\beta_6^{-/-}$ mice, integrin- $\alpha_{\rm E}$ transcripts appeared to be less abundant in S129 $\beta_6^{-/-}$ RNA than in wild-type samples (Figure 6A), confirming the protein expression data described earlier. Densitometry confirmed that there was a significant ($P < 0.05$, unpaired Student's *t*-test with Welch correction) reduction in the net intensity of integrin- α_{E} -specific RT-PCR products from S129 $\beta_6^{-/-}$ samples (Figure 6B).

Toluidine blue staining of Carnoy's-fixed sections (Figure 6, C and D) produced a similar pattern of staining to that observed using IgE-specific antibodies (Figure 4). Intestinal MMCs appeared to be primarily restricted to the lamina propria of S129 $\beta_6^{-/-}$ mice and were present in greater numbers (Figure 6D) than in S129 $\beta_6^{+/+}$ mice (Figure 6C). The jejunal mucosa of S129 $\beta_6^{-/-}$ mice

Figure 5. IgE^{+ve} cell recruitment and integrin- $\alpha_{\rm E}\beta_7$ expression in the jejunal mucosa of *T. spiralis*-infected $\beta_6^{-/-}$ and $\beta_6^{-+/+}$ S129 mice. Frozen sections of jejunum from infected $\beta_6^{-/-}$ (**open bars**) and $\beta_6^{+/+}$ (**filled bars**) S129 mice were triple-labeled with anti-integrin- $\alpha_{\rm E}$ or anti-integrin- β_7 together with anti-IgE and anti-integrin- α_6 (see Figure 4). Data are presented as mean values \pm 1 SEM ($n = 3$) for IgE^{+ve} cells mm⁻² (**A**); the percentage of IgE^{+ve} cells located intraepithelially (B) ; and the relative frequencies of IgE^{+ve} cells expressing integrin- α_E (**C**) or integrin- β_7 (**D**). Differences observed between β_7 (– and β_7 + (+ 5120 mice were significant at $P \le 0.05$ (*) and $P \le 0.01$ (**) β_{6} $\frac{1}{2}$ and β_6 ⁺ $+$ S129 mice were significant at $P \le 0.05$ (*) and $P \le 0.01$ (**).

contained significantly $(P < 0.01$, unpaired Student's *t*-test with Welch correction) more toluidine blue-positive mast cells on day 13 after infection but there was no evidence for any difference in mast cell numbers between S129 $\beta_6^{-/-}$ and S129 $\beta_6^{-+/+}$ mice on day 7 after infection or in the uninfected controls (Table 3). PCNAspecific labeling of toluidine blue-stained sections (Figure 4, O and P) did not reveal any differences in the proportion of actively proliferating (S phase) mast cells on day 13 and was difficult to interpret at earlier time points (Table 3). Similarly, small intestinal worm burdens did not differ significantly between S129 $\beta_6^{-/-}$ and S129 $\beta_6^{+/+}$ mice at either time point. However, there was a trend toward increased worm burdens in the S129 $\beta_6^{\, -/-}$ mice on day 13, indicating that the integrin- $\beta_{\rm 6}$ -null mutation may be associated with a delay in *T. spiralis* expulsion (Table 3). The toluidine blue-based mast cell counts have been repeated in an independent study in S129 β_6 ^{-/-} and S129 $\beta_6^{+/+}$ mice with similar findings (data not shown).

Discussion

The results described here, show, for the first time, that a significant proportion of MMCs recruited during nema-

Figure 6. RT-PCR analysis of integrin- $\alpha_{\rm E}$ and - β_7 transcript abundance and toluidine blue analysis of MMC distribution in the jejunal mucosa of *T.* spiralis-infected $\beta_6^{+/-}$ and $\beta_6^{-/-}$ S129 mice. A: RT-PCR products for integrin- α_E , integrin- β_7 , and the housekeeping gene GAPDH from reverse-
transcribed total jejunal RNA from infected $\beta_6^{-/-}$ (**lanes 1** to **4**) and β_6^{++} (**lanes 5** to **8**) mice. **B:** RT-PCR product signal intensity expressed as a percentage of GAPDH signal intensity. Differences in PCR product abundance between $\beta_6^{+/+}$ and $\beta_6^{-/-}$ S129 mice were significant at $P < 0.05$ (*). Mast cells were identified using toluidine blue staining (pH 0.5) in Carnoy'sfixed/paraffin-embedded sections from $\beta_6^{+/-}$ (**C**) and $\beta_6^{-/-}$ (**D**) S129 mice. Note that mast cells are present in greater numbers in $\beta_6^{-/-}$ mice but are predominantly restricted to the lamina propria. Scale bars, 50 μ m.

tode infection express the integrin- $\alpha_\mathsf{E}\beta_7$, that these are predominantly intraepithelial mast cells, and that their intraepithelial location is highly dependent on the integrin- $\alpha_{\lor}\beta_{\odot}$. A major difference between this and our previous study, showing a general reduction in mast cell recruitment in integrin- β_6 ^{-/-} mice when compared with controls infected with rat-adapted *N. brasiliensis*, ²⁴ is that with *T. spiralis* there was substantially enhanced recruitment of mast cells in the ${\beta_6}^{-/-}$ mice compared with ${\beta_6}^{+/+}$ controls. The intestinal nematode *T. spiralis* induces a particularly potent MMC response³⁶ when compared with *N. brasiliensis²⁴* and this is strikingly evident from the present study.

Because we were unable to co-localize mMCP-1 and integrin- α_F in frozen sections or in whole mounts of exfoliated epithelium, we used surface-bound IgE to identify MMCs, as described by others (Figure 1 and Tables 1 and 2).33,34 Our preliminary co-localization of mMCP-1 and IgE in exfoliated epithelium and of IgE and tryptase in lamina propria mast cells is in agreement with previous studies in which Carnoy's fixation and toluidine blue staining showing that virtually all IgE-bearing cells are mast cells in parasitized rodent intestines.^{33,34} Using surface IgE as a marker, we have established that the α_{F} and β ₇-integrin subunits are expressed on the surface of

Figure 4. Immunofluorescent detection of integrin- α_E and - β_7 expression by IgE^{+ve} cells and PCNA-specific staining of toluidine blue-positive mast cells in the jejunal mucosa of S129 $\beta_6^{-/-}$ and $\beta_6^{+/+}$ mi (pseudo-colored blue), IgE (pseudo-colored green), and integrin- β_7 or - α_E (pseudo-colored red). Representative images are shown from uninfected β_6 ^{+/+} (pseudo-colored blue), IgE (pseudo-colored green), and integrin- β_7 or - α_E (pseudo-colored red). Representative images are shown from uninfected $\beta_6^{+/-}$ (A) and $\beta_6^{-/-}$ (B) mice stained for IgE, integrin- α_6 IgE, integrin- $\alpha_{\rm c}$, and integrin- $\alpha_{\rm E}$. Arrowheads in **D** and **H** highlight occasional cells that exhibit low-level integrin- $\alpha_{\rm E}$ immunoreactivity in sections from $\beta_{\rm 6}$ mice (**inset** in **D** has been contrast enhanced). Higher magnification images from *T. spiralis*-infected $\beta_6^{+/-}$ (**I**) and $\tilde{\beta}_6^{-/-}$ (**J**) mice stained for IgE, integrin- α_6 , mice (**inset** in *D* has been contrast enhanced). Higher magnification images from *1*. *spiralis*-infected β_6 (γ) and β_6 (γ) mice stained for get, integrin- α_6 , and integrin- α_E are annotated with **a** mice for integrin- $\beta_7(\mathbf{K})$, integrin- $\alpha_E(\mathbf{L})$, IgE (M), and integrin- $\alpha_6(\mathbf{N})$, in which two of the three antigen-specific antibodies were substituted with isotype-matched controls, were used to confirm staining specificity. Identical excitation and exposure settings were used to acquire images **A** to **D**. Constant excitation and exposure settings were also maintained during acquisition of images **E** to **N**. Bright-field images of toluidine blue-labeled mast cells probed with anti-PCNA antibodies from *T. spiralis*-infected $\beta_6^{+/-}$ (**O**) and β_6 \sim (P) mice, **arrow** indicates a PCNA-positive mast cell in **O**. Scale bars, 25 μ m.

Total adult worm burdens and numbers of Toluidine Blue-positive mast cells per villus crypt unit were determined for $\beta_6^{-/-}$ and $\beta_6^{+/+}$
S129 mice on days 7 and 13 after infection with *T. spiralis*. Toluidine Blue-positive cells counts were also performed on uninfected β_6 ^{-/-} blue-positive cells counts were also performed on unimected p_6 .

and $\beta_6^{+/+}$ S129 mice (day 0). The proportion of mast cells in the S

phase of cell proliferation (PCNA^{+ve}) was determined on day 13 after infection using PCNA immunohistochemistry and Toluidine Blue staining (Figure 4, O and P). Data are presented as mean values \pm 1 SEM $(n = 4)$

*Differences observed between $\beta_6^{-/-}$ S129 mice and $\beta_6^{+/+}$ S129 controls were significant at $P < 0.01$ (unpaired Student's *t*-test with Welch correction).

MMCs *in vivo* (Figure 4). Furthermore, our data suggest that integrin- α_{F} expression is up-regulated on intraepithelial MMCs, compared to lamina propria MMCs (Figures 3B and 5C).

The addition of TGF- β_1 to mBMMCs induces the expression of integrin- $\alpha_{E}\beta_{7}^{-16}$ and mMCP-1.¹³ In our previous study of *N. brasiliensis*-infected β_6 ^{-/-} mice,²⁴ we established that there was little or no mMCP-1 expression in the few MMCs that were recruited to the gut. We hypothesized that the absence of integrin- $\alpha_{\lor}\beta_{6}$, which is known to activate latent $TGF-\beta_1$ -latency-associated protein,²² would result in an absence of activated TGF- β_1 within the epithelium and, hence, lack of expression of mMCP-1.²⁴ The present results, in which there is a virtual absence of integrin- α_{F} -expressing MMCs and a markedly reduced level of transcription of α_F integrin, is consistent with the hypothesis that integrin- $\alpha_{\lor}\beta_{6}$ plays a significant role in the activation of TGF- β_1 within the epithelial compartment. In addition to integrin- $\alpha_{\nu}\beta_{6}$, 22 plasmin, thrombospondin-1, reactive oxygen species, matrix metalloproteinases, and various other integrin heterodimers have also been implicated in latent TGF- β activation.37 Although the low-level expression of integrin- $\alpha_{\rm E}$ observed in $\beta_{\rm 6}^{\rm -/-}$ mice may be accounted for by one or more of these additional mechanisms, it would seem that, as is the case in lung, 22.23 integrin- $\alpha_{\vee}\beta_{6}$ is the predominant factor in regulating TGF- β_1 activity in the murine intestinal epithelium.

Mast cells derive from hematopoietic progenitor cells of, as yet, an incompletely defined phenotype.^{38,39} Despite the massive mast cell hyperplasia induced by intestinal nematode infection, there is little evidence of mast cell progenitor (MCp) proliferation in the bone marrow or circulation, with the main site of proliferation and differentiation occurring in the jejunum.⁴⁰⁻⁴² Differentiation of MCps into mature mast cells occurs after the cells leave the blood stream and is regulated by factors in their local tissue microenvironment.³⁸ MCp homing to the small intestine has been shown to be dependent on

expression of integrin- $\alpha_4\beta_7$.⁴³ Intraepithelial MCps can be readily isolated from normal intestinal epithelium and the numbers increase within 4 days of infection with *T. spiralis*. ⁴⁰ The microenvironment that permits the intraepithelial expansion of MMCs includes expression of stem cell factor⁴⁴ and TGF- β_1 ⁴⁵ by epithelial cells and of IL-3 by intraepithelial T cells.⁴⁶ There are also reports that IL-9-expressing cells can be located within allergic airway epithelium,⁴⁷ suggesting that enteric epithelium could, similarly, support the presence of IL-9-secreting lymphocytes. These observations support the hypothesis that the conditions for MMC differentiation and proliferation exist within the jejunal epithelium.

There was striking difference between infection with *T. spiralis*, in which mast cells were very abundant in the lamina propria of ${\beta_6}^{-/-}$ mice, and the result of a comparable experiment with the rat-adapted strain of *N. brasiliensis*, in which very few MMCs were recruited.²⁴ It is, however, recognized that *T. spiralis* promotes a particularly potent MMC response⁴⁸ and the stimulus is presumably strong enough to overcome the apparent inhibitory effect on MMC recruitment of a lack of $\alpha_{\nu}\beta_6$ integrin.²⁴ The reason why there were more MMCs in the $\beta_6{}^{-/-}$ mice than in ${\beta_6}^{+/+}$ controls is less obvious. It could be that the slightly larger worm burden on day 13 (Table 3) provides a more potent stimulus. The preliminary analysis on day 13 using PCNA (Table 3) suggests that MMCs were proliferating at the same rate in both groups. An alternative explanation is that the failure of the cells to enter the epithelium and the probable lack of expression of mMCP-1, as described for infection with *N. brasiliensis*, 24 might result in greater retention of MMCs in $\beta_6{}^{-/-}$ mice. This is analogous to the augmented accumulation of MMCs in mMCP-1^{$-/-$} mice.¹ It is possible, for example, that some intraepithelial MMCs can migrate directly into the gut lumen as a consequence of the proteolysis of tight junctions by secreted mMCP-1.⁷ Alternatively, the presence of mMCP-1 in the extracellular space may promote cleavage of epithelial stem cell factor and subsequent apoptosis of MMCs⁴⁹ as they are carried up into the villus. Any loss of autoregulatory function because of the absence, or to very low levels, of mMCP-1, may disturb the rate of apoptosis and/or luminal shedding of MMCs. This is further compounded by the fact that the MMCs were unable to migrate into the epithelium.

Because the most likely defect associated with the absence of the β_{6} -integrin gene is a lack of activated TGF- β_1 on the basolateral membranes of enterocytes, it is possible that the reduction in intraepithelial MMCs in $\beta_6^{-/-}$ mice occurs primarily because MCps fail to express α_F , do not bind to epithelially expressed E-cadherin, and thus have a reduced ability to remain intraepithelial. If the MCps enter the lamina propria of $\beta_6{}^{-/-}$ mice and expand and differentiate in the relative absence of TGF- β_1 , then not only would numbers in the lamina propria be increased, but the phenotype of the differentiated MMC population would also differ from that described in β_6 ^{+/+} controls. As mentioned above, in our study with N. *brasiliensis*, the few MMCs that were recruited did not express mMCP-1, 24 and this is consistent with an altered MMC phenotype.

Studies using integrin-deficient mice and, more specifically, antibody neutralization of integrin- α_{F} in *T spiralis*infected mice¹⁸ suggest that integrin- $\alpha_{E}\beta_{7}^{-}$ is not only required for the retention of MCp and MMCs within the intestinal epithelium but may also be essential for the recruitment of MCps to the gut mucosa because treated mice lacked MMCs both in epithelium and lamina propria. The present results suggest that the reduced expression of integrin- α_{F} as a consequence of defective processing of extracellular TGF- β_1 latency-associated protein is associated with reduced recruitment of intraepithelial MMCs but, paradoxically, results in a threefold expansion of the lamina propria MMCs.

This report is the first to demonstrate, unequivocally, that MMCs express integrin- $\alpha_{\rm E}\beta_7$. Furthermore, the absence of epithelially expressed integrin- $\alpha_{\vee}\beta_{6}$ is associated with depletion of intraepithelial MMCs, substantial down-regulation of $\alpha_{\rm E}\beta_7$ expression, and a paradoxical increase in the frequency of lamina propria MMCs.

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