A novel M cell-specific carbohydrate-targeted mucosal vaccine effectively induces antigen-specific immune responses

Tomonori Nochi,1,3 Yoshikazu Yuki,1,3 Akiko Matsumura,1,3 Mio Mejima,1,3 Kazutaka Terahara,1,3 Dong-Young Kim,1,3 Satoshi Fukuyama,1,3 Kiyoko Iwatsuki-Horimoto,2,3 Yoshihiro Kawaoka,2,3 Tomoko Kohda,4 Shunji Kozaki,4 Osamu Igarashi,1,3 and Hiroshi Kiyono1,3

1Division of Mucosal Immunology, 2Division of Virology, The Institute of Medical Science, The University of Tokyo, Tokyo 108-8639, Japan
3Core Research for Evolutional Science and Technology, Japan Science and Technology Corporation, Saitama 332-0012, Japan
4Laboratory of Veterinary Epidemiology, Department of Veterinary Science, Graduate School of Life and Environmental Sciences, Osaka Prefecture University, Osaka 599-8531, Japan

Mucosally ingested and inhaled antigens are taken up by membranous or microfold cells (M cells) in the follicle-associated epithelium of Peyer’s patches or nasopharynx-associated lymphoid tissue. We established a novel M cell–specific monoclonal antibody (mAb NKM 16–2–4) as a carrier for M cell–targeted mucosal vaccine. mAb NKM 16–2–4 also reacted with the recently discovered villous M cells, but not with epithelial cells or goblet cells. Oral administration of tetanus toxoid (TT)– or botulinum toxoid (BT)–conjugated NKM 16–2–4, together with the mucosal adjuvant cholera toxin, induced high-level, antigen-specific serum immunoglobulin (Ig) G and mucosal IgA responses. In addition, an oral vaccine formulation of BT-conjugated NKM 16–2–4 induced protective immunity against lethal challenge with botulinum toxin. An epitope analysis of NKM 16–2–4 revealed specificity to an α(1,2)-fucose–containing carbohydrate moiety, and reactivity was enhanced under sialic acid–lacking conditions. This suggests that NKM 16–2–4 distinguishes α(1,2)-fucosylated M cells from goblet cells containing abundant sialic acids neighboring the α(1,2) fucose moiety and from non-α(1,2)-fucosylated epithelial cells. The use of NKM 16–2–4 to target vaccine antigens to the M cell–specific carbohydrate moiety is a new strategy for developing highly effective mucosal vaccines.
Currently used in humans because they have lower efficacy than the currently used injectable vaccines in inducing antigen-specific immune responses (6). Because M cells possess the ability to take up luminal antigens, it is logical and attractive to develop a system of delivery of vaccine antigen to both PP-associated and villous M cells to create an effective mucosal vaccine (7). In fact, *Ulex europaeus* agglutinin-1 (UEA-1)–conjugated (8, 9) or σ1 protein–conjugated nasal vaccination (10, 11) induce not only strong antigen-specific plasma IgG and mucosal IgA responses but also CTL immunity, because UEA-1 specific for α(1,2) fucose specifically reacts with murine PP–associated and villous M cells (4, 12), and σ1 protein derived from reovirus specifically binds to a carbohydrate structure containing α(2,3)–linked sialic acid on the membranes of M cells (13). However, because UEA-1 also reacts strongly with goblet cells and the mucus layer covering the intestinal epithelium (14), there have been no effective oral vaccines with UEA-1 as an M cell–targeting vehicle. To overcome this obstacle, we established an M cell–specific mAb and developed a novel strategy for oral vaccination with high efficacy.

**RESULTS AND DISCUSSION**

**Establishment of an M cell–specific monoclonal antibody (NKM 16–2-4)**

To characterize the antigen-sampling M cells for development of an effective M cell–targeted mucosal vaccine, Sprague-Dawley (SD) rats were immunized 4 times at 2–wk intervals with highly purified (>95%) UEA-1–positive cells isolated from murine PPs to establish an M cell–specific mAb. A total of 1,000 hybridomas were generated and screened by immunohistochemical analysis of intestinal tissue sections containing PPs. On the basis of the initial screening, one clone (NKM 16–2-4; rat IgG2c), which possessed specificity to M cells located in the FAE of PPs (Fig. 1 A), was selected. Half of the hybridomas showed no specificity to tissue sections; ~40% of them showed strong reactivity to goblet cells and their secretions; and 10% showed reactivity to the microvilli in all parts of the intestinal epithelium, including M cells and neighboring columnar epithelial cells (unpublished data). These initial screening data indicated that the goblet cells contained in the immunized UEA-1–positive fraction, and their secretions, were vastly immunodominant compared with M cells. However, importantly, NKM 16–2-4 possessed no reactivity to UEA-1–positive goblet cells located in the intestinal villi (Fig. 1 A), indicating that NKM 16–2-4 is a novel mAb possessing high specificity to murine M cells. This is unlike the already known murine M cell–specific lectin UEA-1, which also reacts with goblet cells and their secretions (14). In addition, NKM 16–2-4 reacted very strongly with the apical surfaces of the M cells (Fig. 1 A), rather than the cytoplasm, suggesting that it might be able to be used as a carrier vehicle of M cell–targeted mucosal vaccine. In support of these results, flow cytometric and immunohistochemical analyses demonstrated that NKM 16–2-4 specifically reacted with the surfaces of UEA-1–positive M cells (Fig. S1, available at http://www.jem.org/cgi/content/full/jem.20070607/DC1), but not of UEA-1–negative epithelial cells. In addition, an electronmicroscopic analysis revealed that NKM 16–2-4 specifically reacted with typical M cells, which have short and irregular microvilli and a pocket structure containing lymphocytes and/or monocytes (Fig. 1 B). Furthermore, whole-mount staining analysis revealed that NKM 16–2-4 specifically reacted with villous M cells, in a manner similar to the reaction with PP-associated M cells (Fig. 1 C).

M cells recognized by UEA-1 in mice are also present in the FAE of NALT, as they are in PPs, and act as antigen-sampling cells for the induction of mucosal immunity (15), although our previous finding demonstrated that the mechanism of NALT organogenesis is distinct from that of PP organogenesis (16, 17). Recently, it was reported that group A streptococcus infects its hosts through M cells (15), meaning that M cells could be defined as a portal cell subset of mucosal infection in both the gastrointestinal and respiratory tracts. A subsequent immunohistochemical analysis of NALT tissue sections revealed that NKM 16–2-4 specifically reacted with UEA-1–positive M cells, but not UEA-1–positive, morphologically typical goblet cells with secretory granules (Fig. S2, available at http://www.jem.org/cgi/content/full/jem.20070607/DC1). These results further suggest the possibility of formulating an M cell–targeted nasal vaccine with NKM 16–2-4 for protection against infectious diseases entering through the respiratory tract. Thus, in summary, the novel mAb NKM 16–2-4 specifically reacted with all subsets of M cells, but not epithelial cells or goblet cells, located in PPs, NALT, and the intestinal villi; i.e., in both the gastrointestinal and respiratory tracts (Table I).

**Use of NKM 16–2-4 to develop an M cell–targeted mucosal vaccine**

Because it had been reported that the use of monoclonal antibodies to target injectable vaccine antigen to dendritic cells expressing endocytic receptor effectively initiated antigen–specific immunity (18, 19), we next addressed the characteristics of NKM 16–2-4 as a carrier vehicle of M cell–targeted mucosal vaccines. When we injected FITC-conjugated NKM 16–2-4 or FITC-conjugated control rat IgG into intestinal loops containing PPs, FITC-conjugated NKM 16–2-4 specifically attached to the apical surfaces of M cells in the dome regions of PPs within 10 min of inoculation, whereas FITC-conjugated control rat IgG did not (Fig. 2 A). Furthermore, FITC-conjugated NKM 16–2-4 was taken into the cytoplasmic regions of M cells within 30 min (Fig. 2 A) and reached the basal membrane of the M cells within 4 h, indicating that NKM 16–2-4 could likely be used as a carrier vehicle of orally administered vaccine antigen to M cells.

To directly confirm that M cell–targeted mucosal vaccination with NKM 16–2-4 is an effective strategy for inducing high-level, antigen–specific immune responses, tetanus toxoid (TT) was selected as a prototypical vaccine antigen, as TT has been extensively used in our previous experiments to elucidate the mechanism of the antigen–specific immune responses induced in both the mucosal and systemic compartments by mucosal immunization (20). A chimeric complex of...
IgG or 50 μg noncoupled TT induced, at best, very low TT-specific immune responses (Fig. 2B). In addition, the level of the TT-specific immune response induced by TT-conjugated UEA-1 was lower than that induced by TT-conjugated NKM 16–2-4. These data suggest that an M cell–targeted mucosal vaccine with UEA-1 might be insufficient for antigen delivery to M cells, because the UEA-1–based vaccine is trapped by goblet cells and their secreting mucus, as well as by M cells. Furthermore, 10 times more noncoupled TT (500 μg) induced a small TT-specific immune response compared with TT-conjugated NKM 16–2-4 containing 50 μg TT (Fig. 2B), perhaps because of the low efficacy of antigen delivery to M cells for the induction of antigen-specific immune responses. Although the levels of the antigen–specific antibody responses induced here by immunization with noncoupled TT and CT tended to

Figure 1. Immunohistochemical analysis for the specificity of NKM 16–2-4. (A) Immunohistochemical analysis of PPs revealed that NKM 16–2-4 specifically reacted with UEA-1–positive M cells (red arrows), but not UEA-1–positive goblet cells (yellow arrowheads). (B) Electronmicroscopic analysis revealed that typical M cells, which had short and irregular microvilli and pocket structures containing lymphocytes and/or monocytes, specifically reacted with NKM 16–2-4. Positive reactions are shown by gold particles (18 nm). IEC, intestinal epithelial cell. (C) Whole-mount staining of PPs and villous epithelium demonstrated that, in addition to PP-associated M cells, UEA-1–positive villous M cells were specifically recognized by NKM 16–2-4. Bars, 50 μm.
be lower than those in a previous study (20), this discrepancy might have been caused by differences in the mouse haplotype or the sources of TT and CT. Despite the discrepancy, our current findings emphasize the effectiveness of the newly established NKM 16–2–4 for the targeting of vaccine antigen to M cells to induce antigen-specific immune responses.

Moreover, when mice were orally immunized with botulinum toxoid (BT) conjugated with NKM 16–2–4 or control rat IgG (in total, each 200 μg contained 50 μg BT per mouse) in the presence of CT, brisk botulinum toxin–specific serum IgG and fecal IgA responses were induced in mice immunized with BT-conjugated NKM 16–2–4, but not in those immunized with BT-conjugated control rat IgG (Fig. 2 C). In addition, the mice immunized with BT–conjugated NKM 16–2–4 survived after challenge with 200 ng (10,000 × LD₅₀) of botulinum toxin, whereas the mice immunized with BT-conjugated control rat IgG died within 3 h (Fig. 2 D). These data strongly indicate that the M cell-targeted mucosal vaccine with NKM 16–2–4 can effectively induce protective immunity with the minimum dose of vaccine antigen.

To confirm the mechanism by which the NKM 16–2–4–based M cell–targeted mucosal vaccine induces brisk antigen–specific immune responses in the systemic and mucosal compartments, and its universality, OVA was then chosen as a prototype antigen with low antigenicity. An immunocytochemical analysis revealed that Alexa Fluor 647–labeled OVA conjugated with NKM 16–2–4 and FITC-conjugated avidin specifically reacted with UEA-1–positive isolated M cells in vitro (Fig. 3 A), and intestinal loop assay clearly demonstrated that it specifically attached to the apical surfaces of M cells and was subsequently taken up into the cytoplasmic regions of M cells in vivo (Fig. 3 B). Furthermore, brisk increases in the levels of OVA-specific serum IgG were induced in mice immunized with only 200 μg OVA-conjugated NKM 16–2–4 containing 50 μg OVA, but not with the same amount of OVA-conjugated control rat IgG (Fig. 3 C). Our previous study showed that amounts of OVA as high as 1 mg were required to induce OVA-specific immune responses (5); now, oral immunization with even small amounts of poorly immunogenic antigens (e.g., OVA) is possible by using the M cell–targeting concept with NKM 16–2–4.

We could not directly compare the efficacy of NKM 16–2–4–based mucosal vaccine with those of already published σ1-based mucosal vaccines (10, 11) because the latter systems have been used for nasal, but not oral, vaccines and no information is currently available on whether σ1 possesses
BRIEF DEFINITIVE REPORT

Identification of antigens recognized by NKM 16–2-4

In attempts to elucidate the antigen-sampling mechanism of M cells for the induction of antigen-specific immune responses, a major drawback has been the lack of knowledge of the specific genes and the corresponding molecules expressed by M cells. In addition, no information regarding which murine M cell-specific glycoproteins are recognized by UEA-1 is currently available, although UEA-1 is used extensively as a specific marker of M cells in mice. Therefore, we tried to identify the membrane antigen recognized by NKM 16–2-4 by using a proteomics approach with liquid chromatography-tandem mass spectrometry (LC-MS/MS) after immunoprecipitation of an M cell lysate with NKM 16–2-4. 4 major bands (3 bands >250 kD and 1 band of ~150 kD) were precipitated by NKM 16–2-4 (Fig. 4 A), and these were identified by LC-MS/MS as maltase glucoamylase (top three bands) and alanyl (membrane) aminopeptidase (bottom band). These two molecules, which have been reported as intestinal enzymes of 410, 275, and 260 kD (21), and 150 kD, respectively, (22) under denatured conditions, are distributed at the brush borders of epithelial cells for the final digestion of dietary nutrients (21, 22). Because they were not homologous specificity for villous M cells. However, our strategy for using NKM 16–2-4 as an M cell-targeting vehicle might be superior, because NKM 16–2-4 possesses specificity for both villous M cells and PP-associated M cells. In support of our hypothesis, our previous data showed that villous M cells are capable of taking up orally administered antigens for the induction of PP-independent, antigen-specific immune responses (4). However, it should be noted that TT- or OVA-specific immune responses were not effectively induced without the presence of the mucosal adjuvant CT, even if the antigen was targeted to M cells by using NKM 16–2-4. This finding could be explained by the observation that the gastrointestinal immune system generally operates via a sophisticated mucosal regulatory network to avoid unnecessary hyperimmune responses to the numerous orally encountered antigens in the harsh environment of the intestinal tract (3). Therefore, it is essential to use the mucosal adjuvant, which temporarily breaks the mucosal regulatory network system, to activate gastrointestinal immunity. In practical terms, further studies are needed to develop a safe mucosal adjuvant and take advantage of M cell-targeted mucosal vaccines with NKM 16–2-4.

Figure 3. Effective uptake and universality of the M cell–targeted mucosal vaccine. (A) Immunocytochemical analysis showed that an M cell–targeted OVA vaccine composed of Alexa Fluor 647–conjugated OVA, FITC-conjugated avidin, and NKM 16–2-4 specifically reacted with isolated UEA-1-positive M cells. (B) In an intestinal loop assay, the M cell-targeted OVA specifically attached to the apical surfaces of M cells (red arrows) and was immediately taken up into the cytoplasmic regions of M cells. Bars, 10 μm. (C) Orally administered OVA-conjugated NKM 16–2-4 effectively induced an OVA-specific serum IgG response, whereas an OVA-conjugated control rat IgG did not. Data are expressed as the mean ± the SD.
with each other, and subsequent in situ hybridization analysis demonstrated that their mRNAs were ubiquitously and abundantly expressed in the intestinal epithelium, including in M cells (Fig. S3, available at http://www.jem.org/cgi/content/full/jem.20070607/DC1), we hypothesized that NKM 16–2–4 possesses specificity for the M cell–specific carbohydrate moiety containing α(1,2) fucose, as the precipitated antigens were commonly recognized by UEA-1 (Fig. 4 B).

Figure 4. Identification of the antigen recognized by NKM 16–2–4. (A) Immunoprecipitation and Western blot analysis with NKM 16–2–4 were performed with an M cell lysate. 4 major bands (3 bands >250 kD and 1 band of ~150 kD) were precipitated. A subsequent LC-MS/MS analysis identified the three top bands as maltase glucoamylase and the bottom band as alanyl (membrane) aminopeptidase. (B) Lectin blot analysis performed after immunoprecipitation with NKM 16–2–4 showed that the precipitated antigens were all recognized by UEA-1. (C) mFUT1 and mFUT2 genes were transfected into original CHO cells and CHO-derived mutant transfectants were all recognized by UEA-1. (1,2) fucose, as the precipitated antigens specifically reacted with CHO cells expressing mFUT1 or mFUT2, but neither NKM 16–2–4 nor UEA-1 showed specificity for original or empty vector–transfected CHO cells (Fig. 4 C). In addition, a blocking analysis showed that pretreatment of NKM 16–2–4 with α-1-fucose did not completely abolish reactivity to mFUT1- or mFUT2-expressing transfectants, although UEA-1 reactivity to these transfectants was dramatically decreased (Fig. S4, available at http://www.jem.org/cgi/content/full/jem.20070607/DC1), indicating that the epitope recognized by NKM 16–2–4 is an mFUT1- or mFUT2-mediated carbohydrate complex containing α(1,2) fucose that is different from the UEA-1–reactive portion of α(1,2) fucose.

Because immunohistochemical analysis demonstrated that UEA-1, but not NKM 16–2–4, recognized goblet cells in the intestinal villi (Fig. 1 A), we turned to examining the differences in recognition patterns between NKM 16–2–4 and UEA-1 by using a mutant line of CHO cells to elucidate the importance of the glycosylation of M cells and goblet cells in the mucosal immune system. When the mFUT1 or mFUT2 gene was introduced into a mutant line of CHO cells (Lec2) with an inactivated CMP-sialic acid transporter (24), the reactivity of NKM 16–2–4, but not of UEA-1, was higher in these transfectants than in the mFUT1- or mFUT2-expressing original CHO cells; however, mFUT1- or mFUT2-expressing Lec1 cells with inactivated GlcNAc transporter I (i.e., a lack of N-glycans) (25) or Lec8 cells with inactivated UDP-galactose transporter (26) were not recognized at all by NKM 16–2–4. On the other hand, we observed very low reactivity of UEA-1 to mFUT1- or mFUT2-expressing Lec8 cells, although mFUT1- or mFUT2-expressing Lec1 cells were not recognized by UEA-1. This is because UEA-1 might recognize α(1,2) fucose, which is linked to very low levels of galactose on N glycans in mFUT1- or mFUT2-expressing Lec8 cells because it has been reported that Lec8 cells retain 10–20% of their galactosylation (26), and no information is currently available on whether α(1,2) fucose links to anything other than galactose. These data suggest that sialic acid might be useful in distinguishing the reactivity of NKM 16–2–4, but not UEA-1 to galactose-binding α(1,2) fucose on N-glycans, although the reactivity to α(1,2) fucose regulated by O-glycans remains unclear. Thus, our initial immunohistochemical analyses demonstrated that the specificity of NKM 16–2–4 to UEA-1–positive M cells, but not UEA-1–positive goblet cells, is attributable to the existence of abundant sialic acids neighboring the α(1,2) fucose–containing carbohydrate moiety on goblet cells, but not on M cells. With the exception of their expression patterns at the tissue level, there is currently little reliable information available on the glycobiochemical and molecular biological differences between mFUT1 and mFUT2 as α(1,2) fucosyltransferases (23). Therefore, further studies, especially in terms of in situ expression patterns at a cellular level at inductive
sites, such as in PPs, are needed to elucidate the role of the carbohydrate moiety containing α(1,2) fucose in the mucosal immune system.

In summary, we established a novel M cell–specific mAb (NKM 16–2-4; rat IgG2c) that selectively recognizes M cells, but not goblet cells or epithelial cells, and we characterized the M cell–specific carbohydrate moiety containing α(1,2) fucose. Our strategy for M cell–targeted vaccination with NKM 16–2-4 is attractive for the development of mucosal vaccines.

**MATERIALS AND METHODS**

**Animals.** Female BALB/c mice, Crlj: CD1–Foxn1<sup>nu</sup> mice, and SD rats between 6 and 8 wk old were obtained from CREA and Charles River Laboratories. All of them were maintained in the experimental animal facility at the Institute of Medical Science, the University of Tokyo, and experiments were performed according to the guidelines provided by the Animal Care and Use Committee of the University of Tokyo.

**Establishment of an M cell–specific mAb.** The M cell–enriched fraction was prepared from murine PPs as previously described, with some modifications, by using UEA-1 (4). In brief, cells isolated from murine PPs were fixed in 4% paraformaldehyde (Wako) and stained with 500 ng/ml PE-conjugated UEA-1 (Biogenesis). UEA-1–positive cells were sorted by a FACSaria cell sorter (Becton Dickinson) and injected into the footpads of SD rats (10<sup>6</sup> cells/rat) 4 times at 2-wk intervals, with TiterMax Gold (TiterMax) as an adjuvant. 4 d after the final immunization, lymphocytes isolated from the spleen and inguinal lymph nodes of the immunized rats were fused with P3X63-AG8.653 myeloma cells (CRL-1580; American Type Culture Collection) in the presence of 50% (wt/vol) polyethylene glycol 1500 (Roche). Established hybridomas were injected into Crlj: CD1–Foxn1<sup>nu</sup> mice, and mAbs were then purified from ascitic fluids by using protein G–Sepharose (GE Healthcare) and labeled with EZ-Link Sulfo-NHS-LC-biotin (Thermo Fisher Scientific), FITC (Sigma-Aldrich), or Alexa Fluor 647 (Invitrogen).

**Immunohistochemical analysis.** One monoclonal antibody (NKM 16–2-4; rat IgG2c) was selected on the basis of the initial screening and its specificity to M cells determined by immunohistochemical and whole-mount staining analyses, as described previously, with some modification (4). In brief, after a blocking step with 1% BSA, 7-μm fixed frozen sections or fixed tissues containing PPs were stained with 5 μg/ml FITC-conjugated NKM 16–2-4 or FITC-conjugated isotype control (FITC-conjugated rat IgG2c; MBL International) and 1 μg/ml tetramethylrhodamine isothiocyanate–conjugated UEA-1 (Vector Laboratories). The sections were then counterstained with 400 ng/ml DAPI (Sigma-Aldrich) for histochemical analysis and analyzed under a confocal laser-scanning microscope (TCS SP2; Leica). For electron-microscopic analysis, ultrathin sections (100 nm) were incubated with 1 μg/ml purified NKM 16-2-4 after blocking with 1% BSA, followed by 18-nm gold particle–conjugated goat anti-rat IgG (Jackson ImmunoResearch Laboratories) diluted 1:10. Finally, the sections were stained with 4% uranyl acetate and analyzed under a transmission electron microscope (JEM100S; JEOL).

**Uptake of NKM 16–2-4 by M cells.** After the mice were anesthetized with 2 mg ketamine (Sigma-Aldrich), we injected 100 μg of FITC-conjugated NKM 16–2-4 or FITC-conjugated control rat IgG (Sigma-Aldrich) into intestinal loops containing PPs, in accordance with our previous study (4). The mice were killed 10 or 30 min, or 4 h, after the inoculation, and frozen sections (7 μm) of intestinal loop were prepared and analyzed under a confocal laser-scanning microscope after counterstaining with DAPI.

**M cell–targeted vaccination.** TT (provided by the Research Foundation for Microbial Diseases, Osaka University, Osaka, Japan) and type A BT (prepared according to a previous study; reference [27]) were first treated with EZ-Link Sulfo-NHS-LC-biotin. Next, biotinylated TT or BT at 1 mg/ml was incubated with the same volume of avidin (1 mg/ml; Sigma-Aldrich). The complexes were then incubated with twice the volume of biotinylated NKM 16–2-4, biotinylated control rat IgG (Sigma-Aldrich), or biotinylated UEA-1 (Vector Laboratories; each 1 μg/ml). Mice were orally immunized with the complexes (in total, each 200 μg contained 50 μg TT or BT per mouse), noncoupled TT (50 or 500 μg per mouse), or PBS alone 3 times (once a week), together with 10 μg CT (List Biological Laboratories) as a mucosal adjuvant. 7 d after the final immunization, serum and fecal extracts were collected and analyzed for TT- or type A botulinum toxin–specific serum IgG and fecal IgA responses by ELISA, as previously described (5, 27).

To examine the protective immunity, the mice were challenged via the i.p. route with 200 ng type A botulinum toxin (10,000× LD<sub>50</sub> i.p.) diluted in 100 μl of 0.2% gelatin/PBS (27). To confirm the universality of M cell–targeted mucosal vaccine with NKM 16–2-4, OVA (Sigma-Aldrich) was conjugated with NKM 16–2-4 or control rat IgG and orally immunized together with 10 μg CT. In addition, intestinal loop assay was performed by using M cell–targeted OVA composed of Alexa Fluor 647–conjugated OVA (Invitrogen), FITC-conjugated avidin (Sigma-Aldrich), and NKM 16–2-4 or control rat IgG. Conjugation of NKM 16–2-4 or control rat IgG and the protein antigen was confirmed by sandwich ELISA (unpublished data).

**Analysis of antigen recognized by NKM 16–2-4.** To identify the antigen recognized by NKM 16–2-4, we performed an immunoprecipitation assay with NKM 16–2-4 followed by an LC-MS/MS analysis. In brief, a lysate of M cells was incubated with 10 μg/ml NKM 16–2-4 or an isotype control antibody (rat IgG2c; BD Biosciences) followed by protein G–Sepharose (GE Healthcare). Immune complexes were analyzed by SDS-PAGE and Western or lectin blot with 5 μg/ml biotinylated NKM 16–2-4, 5 μg/ml biotinylated isotype control antibody (biotin–conjugated rat IgG2c; BD Biosciences), or 5 μg/ml biotinylated UEA-1 (Vector Laboratories) and ABC–AP complex (Vector Laboratories). To identify the precipitated antigen, LC-MS/MS analysis was performed after digestion with 50 nM trypsin gold (Promega).

**Transfection of cells.** mFUT1 and mFUT2 genes were synthesized from mRNAs from intestinal tissue, including PPs, using specific primers (mFUT1: sense, 5′-TCTAAGGCAGCATGTCAGCAATCCGGCGAGCAGC-3′; antisense, 5′-GCTAAGCGATCCATGGCGAGTGCCCAGGTAC-3′; mFUT2: sense, 5′-ATCTAGGCTATGGCAGTGGCCACCTCAGCTTCTTG-3′; antisense, 5′-TGAGGAGATTCTGAGTAAAGCTTGAGGAGGACAC-3′; Nilel and Banh [mFUT1] and Nilel and EoKiri [mFUT2] restriction enzyme sites are shown by underlining) by RT–PCR and inserted into pIRE2-EGFP vector (BD Biosciences). These plasmids were then transfected into CHO-K1 cells (CCL-61; American Type Culture Collection) and three CHO–cell–derived mutant lines (Lec1, CRL-1735 [reference 25]; Lec2, CRL-1736 [reference 24]; and Lec8, CRL-1737 [reference 26]). 2 d after transfection, the cells were stained with 500 ng/ml Alexa Fluor 647–conjugated NKM 16–2-4 and 500 ng/ml PE-conjugated UEA-1, followed by the application of 10 μl/test VIA-PROVE (BD Biosciences). They were then analyzed by flow cytometry with FACSCalibur (Becton Dickinson). For blocking analysis, 500 ng/ml Alexa Fluor 647–conjugated NKM 16–2-4 or 500 ng/ml PE-conjugated UEA-1 was first pretreated with 0.5 M α-L-fucose (Wako).

**Data analysis.** Data are expressed as the mean ± the SD. All analyses for statistically significant differences were performed by Tukey’s t test, with P < 0.01 considered significant (denoted in the figures with an asterisk).

**Online supplemental material.** Fig. S1 shows the specificity of NKM 16–2-4 to isolated UEA-1–positive M cells. Fig. S2 shows that NKM 16–2-4 specifically reacts with M cells in NALT, similar to its reaction with PP–associated M cells. Fig. S3 shows the expression of malate glycoamylase and d- and l-aminopeptidase mRNAs in PPs. Fig. S4 shows that NKM 16–2-4 reacts with different form of UEA-1–reactive portion of α(1,2) fucose. The online version of this article is available at http://www.jem.org/cgi/content/full/jem.20070607/DC1.
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