

anti-CD20 mAbs from Beckman Coulter (Brea, CA) were used to cross-link CD20. Secondary Abs, including enzyme-conjugated Abs and biotin-conjugated Abs, were purchased from Jackson, Dako and Vector Laboratories Inc. (Burlingame, CA). Retrovirus vectors for shRNA targeting *PERP* were obtained from Takara Bio Inc. (Shiga, Japan).

Cells and materials

The human B-cell lines BJAB and MD901 were used. Both were first reported as BL-derived cell lines [22, 23] but are later considered to be DLBCL-derived cell lines [24, 25] because of a lack of myc translocation [26]. Another human DLBCL-derived cell line SU-DHL-4 [27], BL-derived cell lines Daudi, Raji, P32/ISH (Japanese Cancer Research Resource Bank, JCRB, Tokyo, Japan), EB-3, NAMALWA (Institution of Fermentation, Osaka, Japan), CA-46 (Dainippon Pharmacology Co., Osaka, Japan), BALM-18 [28], BALM-24 [29], hairy cell leukemia-derived cell line MLMA (JCRB), multiple myeloma-derived cell line IM-9 [30], B-cell precursor acute lymphoblastic leukemia-derived cell line Reh (American Type Culture Collection), P30/OHK (JCRB), NALM-17, NALM-27 (Research Center Cell Biology Institute, Hayashibara Biochemical Laboratories, INC., Okayama, Japan), NALM-6 (Tohoku University Cell Bank, Sendai, Japan), LC4-1, T acute lymphoblastic leukemia-derived cell line CCRF-CEM, acute myeloid leukemia-derived cell line K562, and EOL-1 (JCRB) were used. Cells were cultured at 37°C in RPMI 1640 medium (Sigma-Aldrich Corp., St. Louis, MO), supplemented with 10% FCS (Thermo Fisher Scientific Inc., Waltham, MA), under a humidified 5% CO₂ atmosphere. The transformed embryonal kidney cell line 293FT (Invitrogen) was also used and cultured in DMEM (Sigma-Aldrich). For Tet-inducible system, Tet system-approved FBS (Clontech Laboratories, Inc., Mountain View, CA) was used.

Clinical specimens from pediatric patients, including nontumorous mononuclear cells separated from autopsied spleen of a patient with myelodysplastic syndrome, nontumorous tonsil, and formalin-fixed, paraffin-embedded tissue specimens from nontumorous reactive lymphadenitis, were used. The specimens have been collected between 1985 and 2001 at our laboratory and are now kept under conditions of anonymity. All of the experiments included in this study followed the tenets of the Declaration of Helsinki and were performed with the approval of the local ethics committee. In the case of splenic mononuclear cells, CD20⁺ B cells were sorted using EPICS ALTRA (Beckman) and then RNA was extracted.

MTC Multiple Tissue cDNA Panels were obtained from Clontech. MVPTM total RNA, human lymph node, was obtained from Stratagene Co. (La Jolla, CA).

Immunohistochemistry

For immunohistochemical staining, the formalin-fixed, paraffin-embedded tissue specimens were deparaffinized, treated using the

heat-induced epitope retrieval method in 10 mM citrate buffer, pH 6.0, or 1 mM EDTA, pH 8.0, stained with Abs using the avidin-biotin complex system and visualized with DAB. In the case of double staining with anti-ZNF385B and anti-CD20 or anti-cleaved caspase-3, visualizations with NovaRED and DAB-Ni (Vector Laboratories, inc., Burlingame, CA), respectively, were used as instructed by the manufacturer.

Plasmid construction

A gateway cassette (bases 1–1705) was amplified from pBLOCK-iT3-DEST (Invitrogen) with FLAG-tag by PCR, and the PCR product was inserted into the *EcoRV* site of pRetroX-Tight (Clontech) (termed pRetroX Tight-FLAG-DEST). Full-length (IF-1) and partially deleted (corresponding to AA93 to 471, IF-1/DEL) cDNA of *ZNF385B* IF-1 was amplified with primers shown in Table 1 from cDNA prepared from EB-3 cells by PCR using KOD plus ver.2 (Toyobo Co., Ltd., Osaka, Japan) and cloned into the *XmnI-EcoRV* sites of pENTR11 (Invitrogen). The resulting pENTR11-ZNF385B IF-1 and pENTR11-ZNF385B IF-1/DEL were recombined with pRetroX Tight-FLAG-DEST using an LR recombination reaction as instructed by the manufacturer (Invitrogen) to construct the tetracycline-inducible *ZNF385B* expression vector pRetroX Tight-FLAG-ZNF385B. Amplified *ZNF385B* IF-1 cDNA and *ZNF385B* IF-1/DEL with primers shown in Table 1 were also inserted into pGEMT easy vector (Promega Co., Fitchburg, WI) and subcloned into the *EcoRI* site of pAS2-1 (Clontech), termed pAS2-1-ZNF385B IF-1 and pAS2-1-ZNF385B IF-1/DEL, respectively. Amplified *p53* cDNA was inserted into pGEMT easy vector (Promega) and subcloned into the *EcoRI* site of pACT2, termed pACT2-p53.

Transfection and induction of ZNF385B

293FT cells were seeded at a density of 5×10^6 cells per 10-cm culture dish 1 day prior to transfection. Retroviral vectors (25 μg), together with pCL gag-pol (22.5 μg) and pCMV VSV-G (2.5 μg), were cotransfected into 293FT cells with Fugene HD (Roche, Penzberg, Germany) and incubated overnight as previously described [31]. After 24 h of incubation, the medium containing virus was collected, filtered through 0.45-μm filters (Millipore, Bedford, MA) and then centrifuged at $6000 \times g$ for 16 h. Supernatant was removed and viral pellet was diluted with 4 mL of fresh medium containing 5 ng/μL polybrene (Sigma-Aldrich). Typically, 5×10^5 cells in a well of a 6-well plate were infected with 2 mL of viral solution of each gene. After 24 h, the medium was replaced with fresh medium and selected in the medium containing appropriate antibiotics. The Tet-on advanced-introduced cells were further infected with retrovirus of pRetroX Tight FLAG-ZNF385B IF-1 and IF-1/DEL. The transfectants were treated with 1 μg/mL doxycycline (DOX) for the indicated period. Induced expression of ZNF385B in transfectants was confirmed by RT-PCR and immunoblotting. Knockdown of *PERP* in BJAB Tet-on

Table 1. The sequences of gene-specific primers for RT-PCR and real-time RT-PCR used in this study

Name of gene	Forward primer	Reverse primer
ZNF385B (<i>common</i>)	ccaggccgaggccactaca	actcgggctccgctggagt
ZNF385B (<i>cloning, IF-1</i>)	atgaatatggcaaattttctacggggcttt	ttagtacggagcaaagaggatggaggcagg
ZNF385B (<i>Y2H cloning, IF-1</i>)	gatgaatatggcaaattttctacggggcttt	ttagtacggagcaaagaggatggaggcagg
ZNF385B (<i>cloning, DEL</i>)	atgcttctgctcttgtgcgcacacctacc	ttagtacggagcaaagaggatggaggcagg
ZNF385B (<i>Y2H cloning, DEL</i>)	gatgcttctgctcttgtgcgcacacctacc	ttagtacggagcaaagaggatggaggcagg
ZNF385B (<i>qPCR, IF-1</i>)	gggctaacctggaaccga	tcagctgacaggaatttggaca
ZNF385B (<i>qPCR, IF-2</i>)	ctgggtgctgtcacacaga	ggataatgcacagcgtacc
ZNF385B (<i>qPCR, IF-3</i>)	gggctaacctggaaccga	gcacaagagcaggaagggtga
ZNF385B (<i>qPCR control vector, IF-1</i>)	gggctaacctggaaccga	ctcagctcttctcctgtttgc
ZNF385B (<i>qPCR control vector, IF-2</i>)	agcggcactgatataacgtggtgc	gtgatggagcagctgggattagcct
ZNF385B (<i>qPCR control vector, IF-3</i>)	cgtgaacgtggactgtaaccttaaaaaacaaa ggcagtcacatgccacacta	gcacaagagcaggaagggtagtagtggcatg tactgcctttgtttta
p53 (<i>Full</i>)	gtggccctgtcatcttctgtccctccca	tcaggcagctcgtggtgagctccctttt
PERP	taccttaacggcgacattt	tggttctattatcaggctcgtg
BAX	gtggcagctgacatgttttc	caaagtagaaaaggcgacaa
CDKN1A	atgtgtctggtcccgtttc	cattgtgggaggagctgtga
14-3-3 σ	gacacagagtccggcattg	atggctctggggacacac
Fas/CD95	tgaaggacatggcttagaagtg	cacttgggtgttgcctggag
BCL2	ttgtggccttctttgattcgtg	ggtgccggttcaggctactcagta
BIM	tggcaaaagcaaccttctgatg	gcaggctgcaattgtctacct
PUMA	aaagcaaaatgagcacaacg	aaacgagccccactctctg
BCL6	agcaaggcattggtgaagaca	atggcgggtgaactggatac
P53TG1	gcaggtctggcttaccaca	gtgtaagtgttcctcgtgtg
NOXA	cagctgtccgaggtgctc	ccgccactcagctacag
GAPDH (<i>qPCR control vector</i>)	aaattgagcccgcagctcccgttcgctc	ggttgagcacagggtactttattgatgta
GAPDH (<i>RT-PCR</i>)	ccaccatggcaaatccatggca	tctagacggcaggtcaggtccacc
GAPDH (<i>qPCR</i>)	gctcagacacattggggaaaggt	gtggtgacaggagcattgctga

ZNF385B IF-1 cells using retroviral vector-mediated shRNA was performed similarly to that described above.

Detection of apoptosis

The incidences of apoptosis were quantified by using FITC-labeled Annexin V of MEBCYTO-Apoptosis kit (Medical & Biological Laboratories Co., Ltd., MBL, Nagoya, Japan). After each treatment indicated in the figure legends, 2.5×10^5 BJAB transfectants were collected and incubated with Annexin V-FITC in 500 μ L of binding buffer for 5 min at room temperature according to the manufacturer's protocol [32].

The incidences of apoptosis were also quantified by using MitoCapture™ Mitochondrial apoptosis detection kit (MBL). After each treatment indicated in the figure legends, 1.0×10^6 BJAB transfectants were collected and resuspended in 1 mL of the diluted MitoCapture solution and incubated at 37°C in a 5% CO₂ incubator for 15 min according to the manufacturer's protocol [32].

Activation of pancaspase, caspase-3 and caspase-8 were quantified using an APOPCYTO Intracellular Caspase Activity Detection Kit (MBL). After each treatment indicated in the figure legends, 2.5×10^5 BJAB transfectants were collected and incubated with

FITC-VAD-FMK, FITC-DEVD-FMK, and FITC-IETD-FMK, for 30–60 min at 37°C in CO₂ incubator.

Data were collected on a FC500 flow cytometer (Beckman) and were analyzed with CXP software (Beckman). Fluorescence-minus-one was used to assess background fluorescence for each population.

PCR

Total RNA was extracted from cultured cells with an Illustra RNAspin (GE Healthcare Bio-Sciences UK Ltd., Chalfont, UK), and cDNA was generated with a First Strand cDNA Synthesis Kit (GE) and ReverTra Ace- α (Toyobo). The set of primers used in this study is shown in Table 1.

Real-time RT-PCR was performed using power SYBR Green PCR Master Mix on an ABI PRISM 7900HT Sequence Detection System (Applied Biosystems, Carlsbad, CA) according to the manufacturer's instructions. The human GAPDH gene was used as an internal control for normalization. The set of primers used for real-time PCR is also shown in Table 1. The vector controls of ZNF385B isoforms and GAPDH were constructed by subcloning of PCR amplicons into pGEMT and pENTR11 (*Xmn*I and *Eco*RV site) vectors, respectively.

Immunoprecipitation and immunoblotting

Cell extracts were prepared and immunoprecipitations were performed using 1 mg cell extracts as described previously [33]. It was confirmed that the cell extracts contain nuclear p53 by immunoblotting (data not shown). Whole cell extracts were incubated with anti-p53 Ab (Santa Cruz) bound to Protein A Mag Sepharose Xtra (GE). The bound complexes were collected by magnet, washed and separated by SDS-PAGE.

Immunoblotting was performed as described previously [33]. Briefly, a 50- μ g sample of each cell lysate was electrophoretically separated on an SDS-polyacrylamide gel and transferred to a nitrocellulose membrane. The membrane was incubated with the appropriate combination of primary and secondary Abs, washed and examined with ECL Plus Western Blotting Detection System (GE).

Yeast two-hybrid assay

Yeast two-hybrid assay was performed following the manufacturer's protocol (Clontech). Briefly, pAS2-1-ZNF385B IF-1 vector and pAS2-1-ZNF385B IF-1/DEL vector were transformed into yeast strain Y187 with pACT2-p53 and then plated on SD/-Trp/-Leu/X-Gal plates. Vector set pAS2-1-SV40 large T antigen and p-ACT2-murine p53 were used as positive control. Representative colonies for the samples, positive control and negative controls were streaked onto an SD/-Trp/-Leu/X-Gal plate.

Statistical analysis

Data were analyzed using Student's *t*-test. Values of *p* less than 0.05 were considered statistically significant. All experiments were performed at least three times and means and SEM were calculated.

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Abbreviations: BL: Burkitt's lymphoma · DLBCL: diffuse large B-cell lymphoma · DOX: doxycycline · IF: isoform · ZF: zinc finger

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The human CD10 lacking an *N*-glycan at Asn₆₂₈ is deficient in surface expression and neutral endopeptidase activity

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ABSTRACT

Background: CD10, also known as neprilysin or enkephalinase exhibiting neutral endopeptidase (NEP) activity, is expressed by B-lineage hematopoietic cells as well as a variety of cells from normal tissues. It cleaves peptides such as cytokines to act for terminating inflammatory responses. Although CD10 molecules of the human pre-B-cell line NALM-6 have 6 consensus *N*-glycosylation sites, three of them are known to be *N*-glycosylated by X-ray crystallography.

Methods: In order to investigate the role of *N*-glycans in the full expression of NEP activity, we modified *N*-glycans by treatment of NALM6 cells with various glycosidases or alter each of the consensus *N*-glycosylation sites by generating site-directed mutagenesis and compared the NEP activities of the sugar-altered CD10 with those of intact CD10.

Results: CD10 of the human B-cell line NALM-6 was dominantly localized in raft microdomains and heterogeneously *N*-glycosylated. Although neither desialylation nor further degalactosylation caused defective NEP activity, removal of only a small part of *N*-glycans by treatment with glycopeptidase F under non-denaturing conditions decreased NEP activity completely. All of the three consensus sites of CD10 in HEK293 cells introduced with wild type-CD10 were confirmed to be *N*-glycosylated. Surface expression of *N*-glycan at Asn₆₂₈-deleted CD10 by HEK293 cells was greatly decreased as well as it lost entire NEP activities.

Conclusions: *N*-glycosylation at Asn₆₂₈ is essential not only for NEP activities, but also for surface expression.

General significance: Quality control system does not allow dysfunctional ecto-type proteases to express on plasma membrane.

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1. Introduction

CD10 was initially reported to be expressed by “common” acute lymphoblastic leukemia (ALL) cells [1] and thus termed as the common acute lymphoblastic leukemia antigen (CALLA), whereas it was later found to be expressed by a variety of cells from normal tissues. For example, CD10 was shown to be expressed on CD179a/b⁺ precursor-B-cells (a normal counterpart of “common”-ALL) in bone marrow [2] and early activated B-cells in the germinal center of peripheral lymphoid organs during normal B-cell development [3]. CD10 is transiently expressed during the development of not only hematopoietic lineages, but also multiple organs and stem cells in a variety of tissues, including mature granulocytes, renal epithelium, and myoepithelium [4,5].

Although the biological significance of CD10 is not fully clarified, it is known to be a type II transmembrane glycoprotein exhibiting NEP (EC 3.4.24.11) activity [6] and can cleave a number of biological active peptides such as substance P, endothelin, oxytocin, enkephalin, angiotensins [7], or amyloid- β -peptide [8] at the amino side of hydrophobic amino acids and reduces inflammatory responses or terminates signaling by degrading the ligands in peripheral tissues. In fact, CD10-targeted mice showed enhanced mortality upon galactosamine-sensitized endotoxin shock or treatment combining TNF- α and IL-1 β [9]. The protease activity of CD10 has also been implicated in maintenance of homeostasis such as fluid balance and nervous system function [10]. Furthermore, in regulating human mammary stem cells, CD10 is proposed to create a gradient of signaling proteins by degrading those nearest to the basement membrane [11].

Raft microdomains provide a platform for a variety of cell functions including signal transduction and play a crucial role in cell development and differentiation [12,13]. We previously investigated apoptotic signal transduction of B-cell precursor ALL cells [14] and Burkitt's lymphoma cells [15] via raft microdomains by cross-linking of B-cell receptor or

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GPI-anchored molecules. Since CD10 of B-cell precursor ALL cell lines is recovered in raft microdomain fractions, it might be involved in signal transduction via raft microdomains, and NEP activity of CD10 might play an important role by terminating cell signaling. In an attempt to investigate the functional role of CD10 in B-cell precursor ALL cells, we examined the localization of CD10 in raft microdomain and its biochemical characteristics and observed that NEP activities of their CD10 molecules were suppressed by glycopeptidase F treatment. It is well known that glycosylation regulates the function of glycoproteins [16], so we used oligonucleotide-directed mutagenesis to alter each of the consensus *N*-glycosylation sites in human CD10 and examined the NEP activities of the wild type (WT) and mutated CD10 transiently expressed in HEK-293 cells. Although a human CD10 molecule has 6 consensus *N*-glycosylation sites, X-ray crystallography has shown that only 3 of them are *N*-glycosylated [17] (Fig. 1). We confirmed that all of these sites were *N*-glycosylated and found that *N*-glycosylation at Asn₆₂₈ is essential not only for NEP activities, but also for surface expression.

2. Materials and methods

2.1. Cells and reagents

Pre-B ALL-derived cell line NALM-6, Pro-B ALL-derived cell line NALM-16 and Reh [2], and Burkitt's lymphoma-derived cell line RAMOS (Japanese Cancer Research Resource Bank, Tokyo, Japan) were cultured in RPMI 1640 supplemented with 10% FBS at 37 °C in a humidified 5% CO₂ atmosphere. The anti-human CD10 monoclonal antibodies (mAbs) used were IF6, developed in our laboratory [18], and NCL-CD10-270, purchased from Novocastra Laboratories Ltd. (Newcastle-upon-Tyne, UK). The second antibodies used were HRP-conjugated rabbit anti-mouse IgG + M (DAKO) and FITC-conjugated goat anti-mouse IgG + M (Jackson Immunoresearch, West Grove, PA). HRP-conjugated RCA lectin was purchased from J-OIL MILLS (Seikagaku Corp., Tokyo, Japan).

2.2. Preparation of cytoplasmic and membrane fractions and raft microdomains

Biochemical subcellular fractionation of NALM-6 cells was carried out as previously described [19]. Briefly, cells were homogenized in a low ionic cell lysis buffer (25 mM NaCl/0.5 mM MgCl₂/18 mM Tris-HCl buffer, pH7.5) and cell debris was removed by centrifugation at 200 ×g for 5 min at 4 °C. The supernatant was centrifuged in a microcentrifuge at 15,000 rpm (19,270 ×g) for 30 min at 4 °C to separate supernatant and pellet as the cytoplasmic fraction and the total membrane fraction,

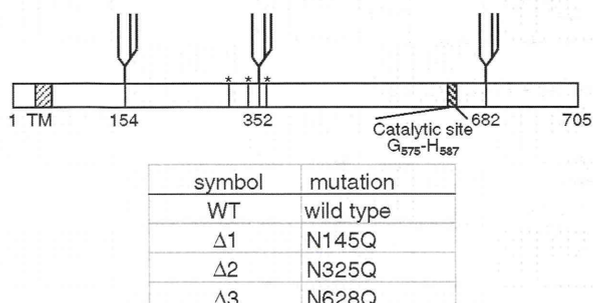


Fig. 1. Schematic diagram of *N*-glycosylation sites in human CD10 and site-directed mutagenesis. TM, transmembrane domain; *, putative glycosylation sites, but confirmed to be unglycosylated by X-ray crystallography; ¶ confirmed to be glycosylated. Symbols of mutants, in which Asn was substituted by Gln at the indicated sites, are shown.

respectively. The total membrane fraction was treated with 1% Triton X-100 in 0.15 M NaCl/25 mM Tris-HCl buffer, pH 7.5, for 30 min at 4 °C and centrifuged at 15,000 rpm for 30 min at 4 °C to separate supernatant and precipitate as the detergent-soluble and -insoluble fractions, respectively. Raft microdomains were obtained by ultracentrifugation of a suspension of the Triton-treated total membrane fraction in 40% sucrose as previously described [12]. Membranes or raft microdomains were solubilized in 40 μM heptylthioglucoiside (Dojin Laboratories, Kumamoto, Japan)/1% Triton X-100/0.15 M NaCl/25 mM Tris-HCl buffer, pH 7.5, for 30 min at 4 °C and centrifuged at 15,000 rpm for 30 min at 4 °C to separate membrane lysates.

2.3. PAGE and Western blot analysis

Proteins were separated in 10% polyacrylamide gels under reducing conditions after dissolving in a loading buffer containing 1.7% SDS and 0.1 M dithiothreitol. Two dimensional (2D) PAGE was performed using the Mini-PROTEAN II 2-D system (Bio-Rad Laboratories, Hercules, CA) as previously described [13]. Gels were either silver-stained or transferred onto PVDF membranes (Millipore Corporation, Bedford, MS). The blots were blocked with 5% skimmed milk in PBS and probed with NCL-CD10-270, then with HRP-conjugated anti-mouse IgG + M antibodies, or blocked with 3% BSA, then probed with HRP-conjugated RCA lectin.

2.4. LC/MS/MS of *N*-glycans

The spots of CD10 in a 2D separated gel were excised and cut into pieces. The gel pieces were destained and dehydrated with 50% acetonitrile. The protein in the gel was reduced and carboxymethylated by incubation with dithiothreitol and sodium monoiodoacetate [20]. *N*-glycans were extracted from the gel pieces as reported previously [21] and reduced with NaBH₄. The borohydride-reduced glycans were desalted with Envi-carb (Aldrich). LC/MS/MS was carried out as previously described [22]. A full MS scan (*m/z* 450–2000) in positive and negative ion modes by FT-ICR MS was followed by data-dependent MS/MS for the most abundant ion performed in the positive ion mode.

2.5. Assay of NEP activity

Enzyme sources of NEP to be examined were intact cells of pro- and pre-B-cell lines, raft microdomains, and CD10 purified by immunoprecipitation with anti-CD10 mAb IF6 bound to Protein A/agarose beads (Bio-Rad). Cells were extensively washed in phosphate-buffered saline (PBS) and resuspended at 1 × 10⁵ cell/mL in the same buffer. Raft microdomains or purified CD10 bound to IF6/PA/agarose beads were suspended in PBS. NEP activity on intact cells or in raft microdomains and purified CD10 was fluorometrically assayed using an indirect coupled enzyme assay method [23]. 5 × 10⁴ cells, raft microdomains, or CD10 on PA/agarose beads were incubated with 0.2 mM glutaryl-Ala-Ala-Phe-4-methyl-coumarin-7-amides (MCA) (Peptide Institute, Osaka, Japan) at 37 °C for 30 min in 20 mM PBS (pH 7.0) in the presence or absence of 0.5 mM phosphoramidon (Peptide Institute), in a total volume of 50 μL. Subsequently, enzyme sources were removed from the reaction mixture by centrifugation. To the reaction mixture was then added 5 μL of a solution containing 0.1 mg (0.4 units equivalent)/mL leucine aminopeptidase (L-5006; Sigma, St. Louis, MO, USA) and 0.2 mM phosphoramidon, followed by further incubation for 30 min at 37 °C to remove a phenylalanine residue from Phe-MCA formed by NEP-catalyzed digestion. The intensity of the liberated 7-amino-4-methylcoumarin was measured at excitation 390 nm and emission 460 nm on a 96-well black plate (CORNING, Amsterdam, Netherlands) using a microplate spectrometer (ARVO, Perkin Elmer Inc., Waltham, MA). The NEP activity was determined on the basis of the decrease in the rate of digestion caused by 0.5 mM phosphoramidon. The relative

amount of CD10 was determined by Western blot analysis using LAS imaging analyzer (Fuji Film, Tokyo, Japan).

2.6. Glycosidase digestion

Raft microdomains or CD10 molecules on IF6-bound Protein A/agarose beads were incubated with 25 mU of sialidase (*Arthrobacter ureafaciens*, Nacalai Tesque, Kyoto, Japan) in 50 μ L of 0.1 M sodium acetate buffer, pH 5.0, for 7 h at 37 $^{\circ}$ C, and then/or 25 mU of β -galactosidase (*Escherichia coli*, Sigma) in 50 μ L of 0.1 M sodium phosphate buffer, pH 7.2/1 mM MgCl₂ for 13 h at 37 $^{\circ}$ C. Alternatively, they were incubated with 1 mU of recombinant glycopeptidase F

(*Escherichia coli*, Takara, Tokyo, Japan) in 50 μ L of native buffer or denaturing buffer according to the manufacturer's instructions overnight.

2.7. CD10 expression vector

The cDNA of the human CD10 was amplified by PCR from the reverse-transcribed product of total RNA from NALM-6 cells extracted using Illustra RNAspin Mini RNA Isolation Kit (GE Healthcare Bio-Science Corp.) and subcloned into the expression vector pcDNA3 (termed as pcDNA3-CD10). Mutations were introduced into the cDNA for CD10 using a QuickChange Site-Directed Mutagenesis kit (Stratagene, La Jolla, CA) according to the manufacturer's instructions. The coding regions of

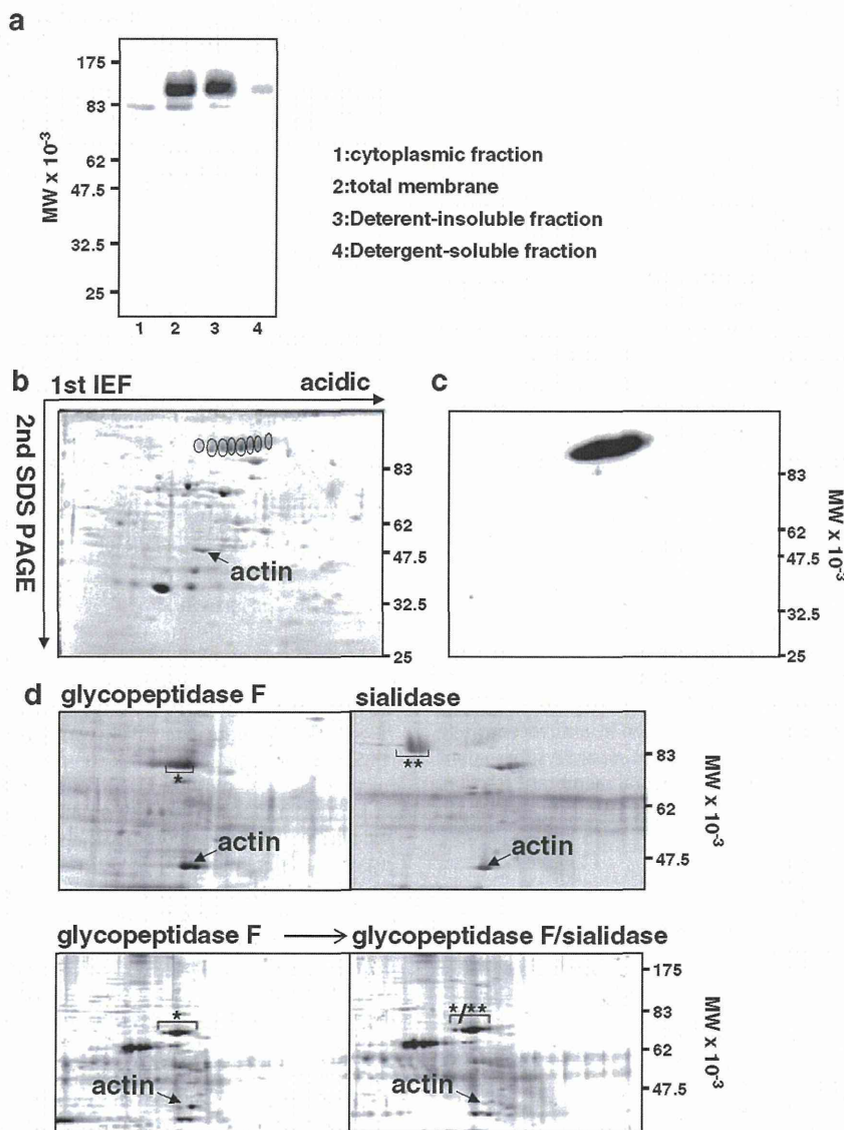


Fig. 2. Characterization of CD10 molecules expressed by human pre-B-cell line NALM-6. **a**, Distribution of CD10 molecules in cytoplasmic fraction, detergent-insoluble membrane fraction, and detergent-soluble membrane fraction of NALM-6 cells. Each lane contains proteins derived from 2.3×10^6 cells. **b**, 2D PAGE of NALM-6 raft microdomain. 60 μ g of protein was subjected to 2D PAGE, and the 2D gel was stained with CBB. **c**, Immunostaining of the 2D separated raft microdomain with anti-CD10. 12 μ g of protein was subjected to 2D PAGE. **d**, 2D PAGE of the glycosidase-treated raft microdomain. The raft microdomain (12 μ g each) was treated with glycopeptidase F under denaturing conditions (upper left) or sialidase (upper right), and the glycopeptidase F-treated raft microdomain (lower left) was further treated with sialidase (lower right). The raft microdomains thus obtained were subjected to 2D PAGE, and the 2D gels were silver-stained. * and ** indicate de-N-glycosylated and desialylated CD10, respectively.

all constructs of cDNA for CD10 were sequenced using an ABI PRISM 3130 sequencer (Applied Biosystems Japan Ltd., Tokyo, Japan). Primers for amplifying, sequencing, and mutagenesis are listed in Table 2. The constructed vectors were transiently transfected into HEK293 cells using Lipofectamine™ LTX (Invitrogen) according to the manufacturer's directions. Cells were cultured for up to 3 days in the presence of 1.5 mg of G418 disulfate (Nacalai Tesque).

2.8. Flow cytometry

Harvested cells were incubated with IF6 for 30 min on ice, and then with FITC-conjugated goat anti-mouse IgG + M for a further 30 min on ice. Stained cells were analyzed using a Beckman Coulter Gallios (Beckman Coulter, Miami Lakes, FL). Five thousand cells were analyzed using FlowJo software (Treestar, Inc., San Carlos, CA).

2.9. RT-PCR

Total RNA was isolated from the transfected cells as described above. cDNA was synthesized from 5 µg of total RNA using a First-Strand cDNA Synthesis Kit (GE Healthcare Bio-Science Corp.). The sequences of gene-specific primers for RT PCR were as follows: for CD10 (forward), 5'-GGAAGAAAGATTGCCCATCG-3'; for CD10 (reverse), 5'-GAGGCTGCTTCAAGATCCATTATG-3'; for GAPDH (forward), 5'-GCTCAGAACACCATGGGAAGGT-3'; and for GAPDH (reverse), 5'-GTGGTGCAGGAGGCAT TGCTGA-3'.

2.10. Protein transport and degradation inhibitors treatment

Cells were cultured in the presence or absence of 5 µg/mL brefeldin A (Sigma) or 25 µM chloroquine (Sigma) for 1 day and 10 µM lactacystin (Peptide Institute) for 2 days.

3. Results

3.1. Localization of NALM-6 CD10 in raft microdomains

The solubility of CD10 molecules for Triton X-100 detergent was examined by Western analysis. CD10 molecules having a molecular weight (MW) of 91 k~120 k were hardly recovered in the cytoplasmic fraction (Fig. 2a, lane 1), but were in the membrane fraction (lane 2), and 95% and 5% of membrane-bound CD10 were recovered in the detergent-insoluble fraction (lane 3) and detergent-soluble fraction (lane 4), respectively. A detergent-insoluble and low-density fraction, namely, raft microdomain, of NALM-6 cells was separated by 2D PAGE and stained with Coomassie brilliant blue (CBB) (Fig. 2b). Proteomic analysis of all spots enclosed by ovals revealed that the series of molecules with a wide range of isoelectric point (pI) from 4.9 to 5.3 was CD10. Anti-CD10 mAb specifically bound to these spots on a PVDF membrane to which 2D PAGE-separated raft molecules were transferred (Fig. 2c). Removal of *N*-glycans from CD10 molecules by digestion with glycopeptidase F under denaturing conditions reduced the molecular weight to 81 k and focused pI to the basic side (* in Fig. 2d, upper left), and sialidase treatment of CD10 molecules shifted the pI more to the basic side (** in Fig. 2d, upper right). Since further treatment of de-*N*-glycosylated CD10 with sialidase did not change the pI (Fig. 2d, lower left and right), all sialic acids are suggested to be on *N*-glycans. These results show that CD10 is a membrane glycoprotein having heterogeneously sialylated *N*-glycans.

3.2. Glycosylation analysis of NALM-6 CD10

The 2D PAGE gel containing the CD10 molecules was excised and treated with glycopeptidase F. The released *N*-glycans were subjected to LC/MS/MSⁿ. Fig. 3 shows the oligosaccharide profile obtained by a full mass scan (*m/z* 450~2000) in the positive and negative ion

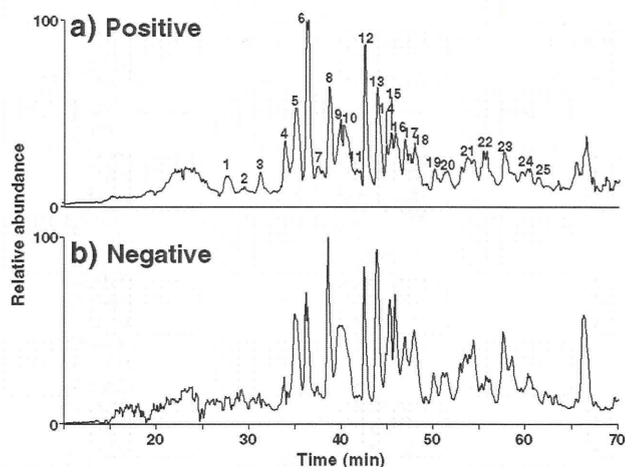


Fig. 3. Total ion current chromatograms of *N*-glycans released from NALM-6 CD10. a, Positive ion mode. b, Negative ion mode.

modes. The most abundant ion in each scan was automatically subjected to data-dependent collision-induced dissociation-MS/MS. Deduced oligosaccharide structures and compositions estimated on the basis of measured accurate masses are summarized in Table 1. Although only three sites of N₁₄₅, N₃₂₅, and N₆₂₈ of 6 potential *N*-glycosylation sites are known to be glycosylated [24], more than 46 oligosaccharide chains were detected. LC-MS analysis revealed that CD10 possesses extremely heterogeneous *N*-glycans.

3.3. NEP activities of surface-expressed CD10 of human B-cell lines

NEP activities of CD10 expressed by human pro-B-cell line Reh and NALM-16, pre-B-cell line NALM-6, and Burkitt's lymphoma RAMOS were measured using fluorogenic substrate, and are expressed as nmol MCA released in a 50 µL test volume for 30 min. Although NEP activities per 5×10^4 cells varied with cell lines, specific activities, which were defined as MCA released (nmol) per CD10 arbitrary unit, were almost the same (Fig. 4).

3.4. Effects of glycosidase treatment of CD10 on NEP activities

In order to investigate the role of oligosaccharide chains for the expression of NEP catalytic activity, we digested raft microdomains of NALM-6 cells or purified CD10 with glycopeptidase F under non-denaturing conditions, sialidase, and/or β -galactosidase, and measured the NEP activity.

The MW of CD10 of NALM-6 cells treated with sialidase was slightly reduced (Fig. 5a, lane 2). Treatment with glycopeptidase F under non-denaturing conditions did not reduce the MW of CD10 (lane 3), whereas extensive digestion under denaturing conditions reduced it to 81 k (lane 4). Disappearance of acidic and high MW area of glycopeptidase F-treated CD10 under non-denaturing conditions in a 2D-PAGE gel (indicated by the triangle in a lower panel of Fig. 5b) showed that this treatment released certain *N*-glycans from NALM-6 CD10. Desialylation and/or degalactosylation of CD10 molecules was confirmed by lectin blotting with RCA, which binds to lactose and *N*-acetylglucosamine. Binding of RCA to the raft microdomains was reduced by β -galactosidase treatment (Fig. 5c, lane 3), and enhanced by sialidase treatment (lane 4). The enhancement of RCA binding by desialylation was canceled by degalactosylation (lane 5). These results show that sialidase treatment successfully removed sialic acids at the non-reducing end and exposed LacNAc residues, and that the further

Table 1
Results of glycosylation analysis of *N*-glycans of NALM-6 CD10.

Peak no	Observed mass (charge)	Monoisotopic mass (charge)	Composition ^a	Deduced structure ^b
1	866.32(2)	1730.624	[Hex]5[HexNAc]3[NeuAc]1	Hybrid
2	785.30(2)	1568.571	[Hex]4[HexNAc]3[NeuAc]1	Hybrid
3	1399.52(1), 700.26(2)	1560.555	[Hex]7[HexNAc]2	M7
3	1561.58(1), 781.29(2)	1398.502	[Hex]6[HexNAc]2	M6
4	947.35(2)	1892.677	[Hex]6[HexNAc]3[NeuAc]1	Hybrid
5	1113.41(2)	2224.799	[Hex]5[HexNAc]4[NeuAc]2	BiNA2
6	967.86(2)	1933.703	[Hex]5[HexNAc]4[NeuAc]1	BiNA
6	939.36(2)	1876.682	[Hex]5[HexNAc]3[NeuAc]1[Fuc]1	Hybrid
7	822.32(2)	1642.608	[Hex]5[HexNAc]4	Bi
8	1150.43(2)	2298.835	[Hex]6[HexNAc]5[NeuAc]1	TriNA1
9	1186.45(2)	2370.857	[Hex]5[HexNAc]4[NeuAc]2[Fuc]1	FBiNA2
10	1186.45(2)	2370.857	[Hex]5[HexNAc]4[NeuAc]2[Fuc]1	FBiNA2
11	1237.47(1)	1236.449	[Hex]5[HexNAc]2	M5
11	1040.90(2)	2079.761	[Hex]5[HexNAc]4[NeuAc]1[Fuc]1	FBiNA
12	1040.89(2)	2079.761	[Hex]5[HexNAc]4[NeuAc]1[Fuc]1	FBiNA
13	1223.47(2)	2444.893	[Hex]6[HexNAc]5[NeuAc]1[Fuc]1	FTriNA1
13	1113.42(2)	2224.799	[Hex]5[HexNAc]4[NeuAc]2	BiNA2
14	895.35(2)	1788.666	[Hex]5[HexNAc]4[Fuc]1	Fbi
14	1333.01(2), 889.00(3)	2663.968	[Hex]7[HexNAc]6[NeuAc]1	TetraNA1
14	1295.99(2), 864.32(3)	2589.931	[Hex]6[HexNAc]5[NeuAc]2	TriNA2
15	1369.02(2), 913.01(3)	2735.989	[Hex]6[HexNAc]5[NeuAc]2[Fuc]1	FTriNA2
15	1295.98(2), 864.32(3)	2589.931	[Hex]6[HexNAc]5[NeuAc]2	BiLac1NA2
15	1077.91(2)	2153.798	[Hex]6[HexNAc]5[Fuc]1	FBiLac1
15	1478.55(2), 986.04(3)	2955.063	[Hex]7[HexNAc]6[NeuAc]2	(MS only)
16	1333.01(2), 889.00(3)	2663.968	[Hex]7[HexNAc]6[NeuAc]1	TriLac1NA1
16	1514.56(2), 1010.04(3)	3027.084	[Hex]6[HexNAc]5[NeuAc]3[Fuc]1	FTriNA3
16	1295.98(2), 864.32(3)	2589.931	[Hex]6[HexNAc]5[NeuAc]2	(MS only)
16	1150.44(2)	2298.835	[Hex]6[HexNAc]5[NeuAc]1	BiLac1NA1
17	1186.45(2)	2370.857	[Hex]5[HexNAc]4[NeuAc]2[Fuc]1	FBiNA2
17	1223.47(2)	2444.893	[Hex]6[HexNAc]5[NeuAc]1[Fuc]1	FTriNA1
18	1551.59(2), 1034.72(3)	3101.121	[Hex]7[HexNAc]6[NeuAc]2[Fuc]1	FTetraNA2
18	1406.04(2), 937.69(3)	2810.026	[Hex]7[HexNAc]6[NeuAc]1[Fuc]1	FTetraNA1
18	1369.02(2), 913.01(3)	2735.989	[Hex]6[HexNAc]5[NeuAc]2[Fuc]1	FBiLac1NA2
18	1478.56(2), 986.04(3)	2955.063	[Hex]7[HexNAc]6[NeuAc]2	(MS only)
19	1333.01(2), 889.00(3)	2663.968	[Hex]7[HexNAc]6[NeuAc]1	TriLac1NA1
19	1223.47(2), 815.98(3)	2444.893	[Hex]6[HexNAc]5[NeuAc]1[Fuc]1	FBiLac1NA1
20	1406.04(2), 937.69(3)	2810.026	[Hex]7[HexNAc]6[NeuAc]1[Fuc]1	FTriLac1NA1 or FBiLac2NA1
20	1369.02(2), 913.01(3)	2735.989	[Hex]6[HexNAc]5[NeuAc]2[Fuc]1	FBiLac1NA2
21	1734.16(2), 1156.44(3)	3466.253	[Hex]8[HexNAc]7[NeuAc]2[Fuc]1	FTetraLac1NA2 or FTriLac2NA2
21	1697.14(2), 1131.75(3)	3392.216	[Hex]7[HexNAc]6[NeuAc]3[Fuc]1	FTriLac1NA3
21	1551.59(2), 1034.72(3)	3101.121	[Hex]7[HexNAc]6[NeuAc]2[Fuc]1	FTriLac1NA2
21	1478.55(2), 986.04(3)	2955.063	[Hex]7[HexNAc]6[NeuAc]2	(MS only)
21	1260.49(2), 840.66(3)	2518.930	[Hex]7[HexNAc]6[Fuc]1	FTriLac1 or FBiLac2
21	1406.04(2)	2810.026	[Hex]7[HexNAc]6[NeuAc]1[Fuc]1	FTriLac1NA1 or FBiLac2NA1
22	1588.61(2), 1059.40(3)	3175.158	[Hex]8[HexNAc]7[NeuAc]1[Fuc]1	FTetraLac1NA1 or FTriLac2NA1
22	1406.04(2)	2810.026	[Hex]7[HexNAc]6[NeuAc]1[Fuc]1	FTriLac1NA1 or FBiLac2NA1
23	1375.18(3)	4122.481	[Hex]9[HexNAc]8[NeuAc]3[Fuc]1	FTetraLac2NA3 or FTriLac3NA3
23	1734.16(2), 1156.43(3)	3466.253	[Hex]8[HexNAc]7[NeuAc]2[Fuc]1	FTetraLac1NA2 or FTriLac2NA2
23	1551.59(2), 1034.72(3)	3101.121	[Hex]7[HexNAc]6[NeuAc]2[Fuc]1	FTriLac1NA2
24	1278.15(3)	3831.385	[Hex]9[HexNAc]8[NeuAc]2[Fuc]1	FTetraLac2NA2 or FTriLac3NA2
24	1588.61(2), 1059.40(3)	3175.158	[Hex]8[HexNAc]7[NeuAc]1[Fuc]1	FTetraLac1NA1 or FTriLac2NA1
24	1406.03(2)	2810.026	[Hex]7[HexNAc]6[NeuAc]1[Fuc]1	(MS only)
25	1181.12(3)	3540.290	[Hex]9[HexNAc]8[NeuAc]1[Fuc]1	FTetraLac2NA1 or FTriLac3NA1
25	1375.19(3)	4122.481	[Hex]9[HexNAc]8[NeuAc]3[Fuc]1	(MS only)

^a Estimated from m/z values obtained by FTMS.

^b Deduced from MSⁿ spectra in positive ion mode; F, fucose; NA, *N*-acetylneuraminic acid; Lac, LacNAc; Bi, biantennary; Tri, triantennary; Tetra, tetraantennary.

treatment with β -galactosidase certainly removed β -galactose residues that had been exposed by the sialidase treatment.

Desialylation and/or degalactosylation of the raft microdomains reduced NEP activities to 80–90% of untreated NEP (Fig. 5d, lanes 2, 3, and 4), whereas glycopeptidase F treatment under non-denaturing conditions greatly reduced NEP activities, although this treatment did not change the MW of CD10 (Fig. 5d, lane 5). Treatment of purified CD10 of NALM-6 cells with glycopeptidase F under non-denaturing conditions also suppressed NEP activities (Fig. 5e). These results show that a certain *N*-glycan, which can be easily digested by glycopeptidase F even under non-denaturing conditions, is necessary for full expression of NEP activities.

3.5. Expression of CD10 lacking *N*-glycan in HEK 293 cells

To examine which *N*-glycans are required for expression of NEP activities, we constructed mutated CD10 lacking *N*-glycans as shown in Fig. 1, and expressed the mutated CD10 in HEK293 cells. Surface expressions of WT and mutated CD10 on HEK293 cells were examined by flow cytometry and Western analysis. WT, Δ 1, and Δ 2CD10 were fully expressed on the surface of HEK293 cells, but Δ 3CD10 was hardly expressed (Fig. 6a). Western analysis confirmed strong expression of WT, Δ 1, and Δ 2CD10 and weak expression of Δ 3CD10 (Fig. 6b). However, transcription of all of these 4 lines occurred at the same level (Fig. 6c). Reduction in molecular weight of WT and mutated CD10 by

Table 2

Nucleotide sequences of oligonucleotide primers used for amplifying, sequencing and site-directed mutagenesis of NALM-6 CD10.

Primer name	Primer sequence (5' to 3')
Cloning_sense	5'-gatgggcaagtcagaagtcagatgg-3'
Cloning_antisense	5'-caaaccggcactctttctgat-3'
Seq_sense	5'-gagatcaatgggaagcattcagctgg-3'
Seq_antisense	5'-gaggctgcttacaagatccattatg-3'
N145Q_sense	5'-agcaaaagcattgtacaggtctgtatacaggaatctgctattgatagcag-3'
N145Q_antisense	5'-ctgctatcaatagcagattcctgtatacagacctgtacaatgcttttgct-3'
N325Q_sense	5'-gggaagcattcagctggttcagttcacaatgaaatcatgtcaa-3'
N325Q_antisense	5'-ttgacatgattcattgtggaactgcaaccagctgaatggcttccc-3'
N628Q_sense	5'-catggtgtacagatggacagtttctctggacctggcag-3'
N628Q_antisense	5'-ctgccaggtccaggaaaactgtccatactgataccatg-3'

glycopeptidase F treatment indicates the presence of *N*-glycans in these CD10 molecules (Fig. 6d).

3.6. NEP activities of WT and mutated CD10

The amounts of MCA (nmol) released by 5×10^4 HEK293 cells transfected with WT-, $\Delta 1$ -, $\Delta 2$ -, and $\Delta 3$ -CD10 were 2.34, 0.39, 1.10, and 0.06, respectively (Fig. 7a). Since the expression level of CD10 varied among transfectants, we examined the NEP activities of CD10 purified from transfectants by immunoprecipitation. The amount of CD10 was determined by Western analysis of the immunoprecipitates, which were separated by 2D PAGE (Fig. 7b). We detected NEP activities in the immunoprecipitates from WT-, $\Delta 1$ -, and $\Delta 2$ -CD10 transfectants, but not any activities in that of $\Delta 3$ -CD10 transfectant (Fig. 7c). These results show that CD10 lacking an *N*-glycan at N₆₂₈ is defective not only in surface expression, but also in expression of NEP catalytic activities.

3.7. Effects of inhibition of intracellular traffic on the protein expression of CD10

In order to elucidate in which step the protein expression of $\Delta 3$ CD10 is suppressed, we cultured human B-cell lines of NALM-6 and Reh and transfectants of WT- and $\Delta 3$ -CD10 in the presence of brefeldin A or lactacystin, and examined their expression by Western analysis. Brefeldin A inhibits transport from ER to Golgi apparatus, resulting in avoidance of degradation in a lysosome [25], and lactacystin inhibits proteasome activity [26]. CD10 molecules having an MW of more than a few hundred thousand were detected in NALM-6 cells cultured in the presence of lactacystin (open arrow in Fig. 8a). The amount of

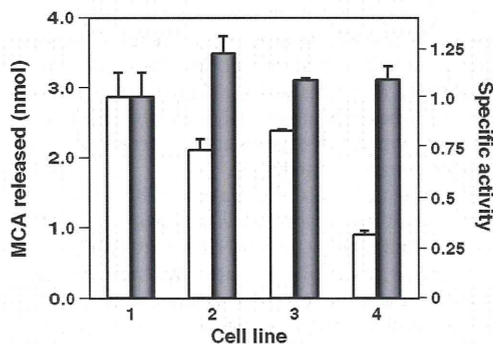


Fig. 4. NEP activities of human B-cell lines. The open column represents nmol MCA released by 5×10^4 cells into 50 μ l of the assay buffer for 30 min and the solid column represents the specific activity. The specific activity was calculated as follows: nmol MCA released was divided by the arbitrary unit of CD10 in a test tube, and the value for Reh was set to 1.0. 1, Reh; 2, NALM-16; 3, NALM-6; 4, RAMOS. Error bars mean SD values of triplicate measurements.

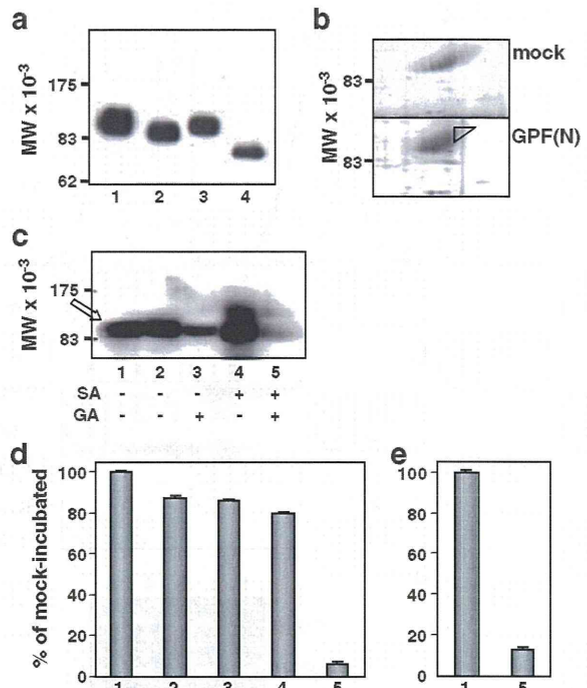


Fig. 5. Effects of glycosidase treatment on NEP activities of NALM-6 cells. **a**, Reduction in MW of CD10 of the glycosidase-treated raft microdomain. The raft microdomain (4 μ g each) was mock-incubated or incubated with glycosidases for 16 h, and then subjected to Western blot analysis with anti-CD10. 1, Mock-incubated; 2, sialidase; 3, glycopeptidase F under non-denaturing conditions; 4, glycopeptidase F under denaturing conditions. **b**, 2D PAGE of the glycosidase-treated CD10. 12 μ g of protein of mock-incubated (mock, upper panel) and glycopeptidase F-treated raft under non-denaturing conditions (GPF(N), lower panel) were subjected to 2D PAGE. 2D gels were silver-stained. The triangle indicates an area which disappeared after glycopeptidase F treatment under non-denaturing conditions. **c**, RCA lectin binding to the glycosidase-treated raft microdomain. The raft microdomain (4 μ g each) was mock-incubated or incubated with glycosidases, and then subjected to lectin-blot analysis with RCA lectin. 1, Mock-incubated for 7 h; 2, mock-incubated for 20 h; 3, β -galactosidase (GA) for 7 h; 4, sialidase (SA) for 13 h; 5, sialidase for 13 h and then β -galactosidase for 7 h; open arrow indicates CD10. **d**, NEP activities of the glycosidase-treated raft microdomains. **e**, NEP activities of the glycosidase-treated CD10. Percentages of NEP activities of the glycosidase-treated to that of the mock-incubated are shown. 1, Mock-incubated; 2, sialidase; 3, β -galactosidase; 4, sialidase and then β -galactosidase; 5, glycopeptidase F under non-denaturing conditions.

CD10 decreased and increased in Reh cells cultured in the presence of brefeldin A and lactacystin (closed and open arrowheads in Fig. 8a), respectively. Aggregated CD10 of NALM-6 or misfolded CD10 of Reh seems to be digested in proteasome, and inhibition of transport from ER to Golgi apparatus suppresses the expression of CD10 of Reh. HEK293 cells transfected with WT-CD10 decreased and increased the expression during the culture in the presence of brefeldin A and lactacystin, respectively (closed and open arrowheads in Fig. 8b), whereas those with $\Delta 3$ -CD10 increased the expression during the culture in the presence of brefeldin A (asterisk in Fig. 8b) and did not increase the expression during the culture in the presence of lactacystin. Next, we used chloroquine to inhibit lysosomal proteases [27]. The amount of $\Delta 3$ -CD10 was not increased by the treatment with either lactacystin or chloroquine (Fig. 8c). These results indicate that the low level of expression of $\Delta 3$ -CD10 in HEK293 cells may not be due to degradation either in proteasome or lysosomes.

4. Discussion

As we presented in this study, CD10 of human BCP-ALL cell line is one of the major raft microdomain glycoproteins and heterogeneously