Serum Tenascin-X Strongly Binds to Vascular Endothelial Growth Factor

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Interstitial extracellular matrix tenasin-X (iTNX) with about 450 kDa is prominently present in various tissues. Previously, we identified the serum form of TNX (sTNX) with 200 kDa in the mouse. In the present study, in order to investigate distinctive features and functions of sTNX, a plasmid encoding the recombinant mouse sTNX was constructed. As a control, we also constructed a plasmid encoding mouse 450-kDa iTNX and a plasmid encoding 250-kDa iTNX, which lacks the region of 200-kDa sTNX from 450-kDa iTNX. In cells stably expressing each recombinant TNX, a more than 7-fold larger amount of 200-kDa sTNX was released into conditioned medium than the amounts of 250-kDa iTNX and 450-kDa iTNX released into the medium. We previously reported that a splice isoform of iTNX (340-kDa iTNX) binds to vascular endothelial growth factor B (VEGF-B) as well as to VEGF-A. Therefore, the ability of VEGF-A and VEGF-B to bind to 200-kDa sTNX was examined by a co-immunoprecipitation assay in comparison with the binding abilities to 250-kDa iTNX and 450-kDa iTNX. It was found that sTNX strongly bound to VEGF-A and VEGF-B, compared with the binding abilities of other iTNX proteins. Based on the results of assays of incorporation of 5-ethyl-2'-deoxyuridine (EdU), we found that purified recombinant 200-kDa sTNX both alone and in combination with VEGF-A or basic fibroblast growth factor (bFGF) can weakly promote DNA synthesis in proliferating vascular endothelial cells (UV-2 cells). These results suggest that sTNX possesses weak activity for proliferation of endothelial cells.

Key words tenascin-X; extracellular matrix; serum form; vascular endothelial growth factor

The extracellular matrix (ECM) was originally thought to be merely a physical framework for cells in terms of mechanical strength. However, this concept has been revised in recent years. The ECM is now known to participate in cell proliferation, migration, differentiation, and survival.1) The ECM contains not only many adhesive proteins, including fibronectin, collagen and laminin, which generally promote cell attachment or migration, but also adhesive modulatory proteins such as tenasin, thrombospondin and SPARC/osteonectin/BM40, which regulate the interactions between cell receptors and adhesive proteins.

Among adhesive modulatory proteins, the tenasin family constitutes a group of ECM glycoproteins with a characteristic structure. The four members of this family identified so far [tenasin-C (TNC), restrictin/J1-160/180 (tenasin-R, TNR), tenasin-X (TNX), and tenasin-W/tenasin-N (TNW/TNN)] have been found in vertebrates.2—6) Tenasin family members are made up of the same types of structural domains, including a cysteine-rich segment at the amino terminus, epidermal growth factor (EGF)-like repeats, fibronectin type III (FNIII)-like repeats, and a fibrinogen-like domain at the carboxyl terminus.

TNX is the largest member of the tenasin family. Complete deficiency of TNX in humans leads to a rare recessive form of Ehlers–Danlos Syndrome (EDS), and TNX haploinsufficiency is associated with hypermobility type of EDS. The skin of TNX-deficient patients is markedly lax with poor recoil properties and shows easy bruising.7—9) In the skin of these patients, the collagen density is reduced in the dermis, and the elastic fibers are abnormal. Furthermore, recent genetic studies have revealed a strong association between the TNX locus and other diseases such as schizophrenia,10) arterial tortuosity syndrome (ATS),11,12) and systemic lupus erythematosus.13) Recent evidence has suggested that TNX is involved in collagen fibrillogenesis,13,14) collagen deposition,15) and development and maintenance of elastic fibers.16) As a biological feature of TNX, it has been shown that TNX interacts with types I, III and V fibrillar collagen and fibril-associated types XII and XIV collagen15,17) and with proteoglycan decorin.18)

Schalkwijk et al.19) reported that human TNX is also present in normal serum with an apparent molecular size of 140 kDa. Measurement of serum TNX revealed that high levels of serum TNX are a risk factor for abdominal aortic aneurysm.19) We previously identified the serum form of TNX (referred to as sTNX) with a molecular mass of 200 kDa in the mouse,20) and we gave the name iTNX to conventional interstitial TNX. The full-length iTNX has a molecular mass of about 450 kDa. The 200-kDa sTNX contains the last 15 FNIII repeats and a fibrinogen domain identical to the C-terminal portion of the 450-kDa iTNX. The N-terminus of sTNX is located in the juncture between the 16th FNIII and 17th FNIII repeats. Furthermore, 200-kDa sTNX is generated by proteolytic cleavage of the 450-kDa iTNX.

A number of fragments of ECM molecules have been identified as angiogenesis inhibitors. Angiostatin, a 38-kDa proteolytic fragment of plasminogen containing the first four kringle structures, inhibits proliferation, blocks migration, and increases apoptosis of endothelial cells.21) Likewise, endostatin, the 20-kDa C-terminal fragment of the NC1 domain of the α1 chain of type XVIII collagen is a potent inhibitor of angiogenesis.22) Endostatin specifically inhibits endothelial proliferation and potently inhibits angiogenesis and tumor growth. Canstatin23) and tumstatin24) derived from the NC1 domain of type IV collagen have also been shown to have anti-angiogenic activities. Besides the derivatives from collagen, anastellin, a 76-aa peptide derived from the first type III repeat in fibronectin,25) and endorepellin, the C terminus of
perlecan (a ubiquitous basement membrane heparan sulfate proteoglycan), have been reported to inhibit angiogenesis. These endogenous angiogenesis inhibitors have significant therapeutic potential.

We previously have demonstrated that a splice isoform of iTNX that lacks the 3rd FNIII repeat (M3) and the 15th FNIII to the 22nd FNIII repeat (M15—M22), 340-kDa iTNX, binds to vascular endothelial growth factor B (VEGF-B) as well as VEGF-A and that iTNX in combination with 340-kDa iTNX strongly promotes DNA synthesis in endothelial cells compared with the effect of VEGF-B alone. Furthermore, since 200-kDa STNX is derived from the C-terminal portion of 450-kDa iTNX, we were interested to clarify whether STNX is involved in the proliferation of endothelial cells. For this end, in this study we examined distinctive features of STNX such as susceptibility to be released into conditioned medium, binding capability to VEGF-A and VEGF-B, and proliferation activity for endothelial cells.

MATERIALS AND METHODS

Cell Cultures Human embryonic kidney 293T cells were cultured at 37°C in a 5% CO2 humidified atmosphere in Dulbecco’s modified Eagle’s medium (DMEM) (Nissui Pharmaceutical Co., Tokyo, Japan) supplemented with 0.15% NaHCO3, 10% bovine calf serum (JRH Biosciences, U.S.A.), penicillin (10 units/ml) and streptomycin (10 mg/ml). CHO cells (cell line derived from Chinese hamster ovarian carcinoma) for attached culture were grown in HAM’s F12 medium (Nissui). CHO cells in suspension culture were cultured at 37°C in a 5% CO2 humidified atmosphere in Dulbecco’s modified Eagle’s medium (DMEM) (Nissui). CHO cells in suspension culture were grown in a serum-free CHO-S-SFM II medium (Invitrogen, Carlsbad, California, U.S.A.) with 0.15% NaHCO3 for the production of recombinant proteins. To adapt CHO cells from the serum-supplemented medium to the serum-free medium, sequential adaptation was performed according to the instructions of the manufacturer (Invitrogen). The UV-sensitive cell line (mouse vascular endothelial cell line transformed by ultraviolet radiation) was purchased from RIKEN Bioresource Center (Tsukuba, Japan). UV-sensitive cells were grown in DMEM with 0.15% NaHCO3 and 10% fetal bovine serum (FBS) according to the provided protocol.

Plasmid Construction The following expression plasmids were constructed. pSec-F-sTNX (Fig. 1C), which encodes 200-kDa sTNX with 15 FNIII repeats and a fibrinogen domain identical to the C-terminal portion of 450-kDa iTNX, was constructed as follows. Spleens from adult ICR mice were dissected, frozen, and powdered in liquid nitrogen. Total RNA was isolated, and first-strand cDNA was synthesized by reverse transcription-polymerase chain reaction (RT-PCR) as described previously. First, pT7-M16j-19 was prepared by PCR using the prepared cDNA as a template and forward fM19 primer 5'-GGGAAATTCAGGCCCCCGAGCCCAGAG-3' and reverse del-mR25 primer 5'-GAGCCGTTGTCCTCCACCGAGAC-3'. PCR fragment was cloned into pGEM-T Easy vector, which encodes the 19th—25th FNIII repeats. Then a Nael site in the vector sequence of pGEM-M19-25 was changed to HindIII site by HindIII linker ligation. Subsequently, the EcoRI–SalI fragment of pT7-M16j-19 and the SalI–HindIII fragment of pGEM-M19-25 were ligated and then inserted into the respective sites of pGEM-T Easy vector, in which the Nael site in the vector sequence was changed to HindIII site, yielding pGEM-M16j-25. Finally, the EcoRI–ApalI fragment of pGEM-M16j-25 and the ApalI–Xhol fragment of the plasmid pB3CHX-18 encoding the 13th—15th and 25th—31st FNIII repeats followed by a fibrinogen domain were ligated and then inserted into the respective sites of pSec-F vector, a pSecTag2/HygroB vector (Invitrogen) containing a FLAG-tag, yielding FLAG-tagged pSec-F-sTNX.

pSec-F-heptad-M16 encoding 250-kDa iTNX (Fig. 1B), which lacks the region of 200-kDa STNX from 450-kDa iTNX, was constructed as follows. First, pGEM-M13-16 was constructed by PCR using the cDNA prepared from mouse spleen as a template and forward fM13-14 primer 5'-GGGGATCCCGAGGGGGGAGCCAGAGCCAACAACA-3' and reverse rM16 primer 5'-GGGTGAGGAGACAGCAGCAGAGGCCACAGC-3'. The PCR product was cloned into pGEM-T Easy vector, yielding pGEM-M13-16, which encodes the 13th—16th FNIII repeats. Next, the HindIII–Xhol fragment of pGEM-M13-16 was inserted into HindIII/Xhol sites of pEGLNH-1 encoding the 10th—18th EGF repeats and the 1st—2nd and 4th—13th FNIII repeats, yielding pBlue-EGF10-M16. Finally, the NotI–Xhol fragment of pBlue-EGF10-M16 was inserted into NotI/Xhol sites of pSecF2-1.1EGF-1 encoding a cysteine-rich segment at the amino terminus and the 1st—10th EGF repeats, yielding FLAG-tagged pSec-F-heptad-M16.

pSec-F-TNX, which encodes the full-length 450-kDa iTNX (Fig. 1A), was constructed as follows. First, pGEM-M13-17 was constructed by PCR using the cDNA prepared from mouse spleen as a template and forward fM13-14 primer and reverse rM17 primer 5'-GGGTGAGGAGACAGCAGCAGAGGCCACAGC-3'. The PCR product was cloned into pGEM-T Easy vector, yielding pGEM-M13-17, which encodes the 13th—16th FNIII repeats. Then the Bhol–Xhol fragment of pGEM-M13-17 was replaced by the Bhol–Xhol fragment of pSec-F-sTNX, yielding pGEM-M13-FG. Next, the HindIII–Xhol fragment of pGEM-M13-FG was inserted into the HindIII/Xhol sites of pEGLNH-1, yielding pBlue-EGF10-FG. Finally, the NotI–Xhol fragment of pBlue-EGF10-FG was inserted into NotI/Xhol sites of pSecF2-1.1EGF-1, yielding FLAG-tagged pSec-F-TNX.

pSecFTNX-2 encoding a FLAG-tagged splice isoform of 340-kDa iTNX, which lacks M3 and M15—M22 fibronectin type III repeats, pSec-mVA65-HA encoding HA-tagged mouse VEGF-A146 and pSec-hVB186-HA encoding HA-tagged human VEGF-B186 have been described previously. Establishment of 200-kDa stTNX, 250-kDa iTNX and 450-kDa iTNX-Expressing Cell Lines Adherent CHO cells in a 10-cm dish were transfected with 5 μg of pSec-F-sTNX, pSec-F-heptad-M16 or pSec-F-TNX by the calcium phosphate precipitation technique and cultured in the presence...
ence of 800 μg/ml of hygromycin B (Wako, Osaka, Japan). About 2 weeks after transfection, hygromycin B-resistant colonies were selected and used as 200-kDa sTNX, 250-kDa iTNX or 450-kDa iTNX-expressing cell lines. As a control, mock cells carrying an empty pSecTag2/HygroB vector were also selected.

**Western Blot**
Proteins in cell lysates or conditioned medium were electrophoresed through 7.5% or 10% sodium dodecyl sulfate (SDS)-polyacrylamide gels and then electropholeted (PAGE) onto Hybond ECL nitrocellulose membranes (GE Healthcare, Buckinghamshire, England). The membranes were blocked at room temperature (RT) for 1 h with 5% nonfat dry milk in TBST [20 mM Tris–HCl (pH 7.5), 150 mM NaCl, 0.1% Tween-20] and then incubated with a primary antibody (10 μg/ml in TBST with 1% milk) for 1 h at RT. Membranes were washed three times with TBST, incubated with anti-mouse, anti-rabbit, anti-goat Alexa Flour 680-conjugated or IRDye 800-conjugated immunoglobulin (Ig)G (Molecular Probes, Eugene, OR, U.S.A.) followed by 680-conjugated or IRDye 800-conjugated immunoglobulin (Ig)G (Molecular Probes, Eugene, OR, U.S.A.) followed by visualization using an infrared imaging system (Odyssey, LI-COR, Lincoln, U.S.A.).

**Detection of 200-kDa sTNX, 250-kDa iTNX or 450-kDa iTNX in Conditioned Medium, Compared with That in Cell Lysates**
For the detection of FLAG-tagged 200-kDa sTNX, 250-kDa iTNX or 450-kDa iTNX in the conditioned medium of the cell line expressing each type of TNX cultivated for 48 h, the conditioned medium was clarified from debris by centrifugation. Subsequently, anti-FLAG M2 affinity resin (Sigma-Aldrich, Tokyo, Japan) equilibrated in phosphate-buffered saline (PBS)/0.5% NP-40 was added to the collected medium, followed by rotation at 4 °C for 3 h. The resin was pelleted by centrifugation and washed several times with wash buffer [10 mM Tris–HCl (pH 7.5), 50 mM NaCl, 0.1% Nonidet P-40 (NP-40)]. Afterwards, the resin was boiled in Laemmli buffer before being subjected to SDS-PAGE and Western blot analysis with anti-FLAG M2 monoclonal antibody (Sigma-Aldrich). On the other hand, for the detection of each TNX form in cells, cells were lysed in lysis buffer containing 50 mM Tris–HCl (pH 7.5), 120 mM NaCl, 0.5% NP-40, and protease inhibitors for 10 min on ice, and whole-cell lysates were obtained by subsequent centrifugation. The protein concentration was determined by a bicinchoninic acid (BCA) assay (Pierce, Rockford, U.S.A.). Cell lysates were electrophoresed and subjected to Western blot analysis.

**In Vivo Binding Assay**
pSec-mVA_{164}HA or pSec-hVB_{86}HA was transfected into 293T cells with each FLAG-tagged TNX form (200-kDa sTNX, 250-kDa iTNX and 450-kDa iTNX), and then 48 h after transfection the cell lysate was prepared as described in the above section. Approximately 3 mg of 293T cell proteins was first immunoprecipitated with mouse anti-FLAG M2-agarose affinity resin at 4 °C for 4 h. After centrifugation and washing of the resin five times with wash buffer, the precipitates were separated in a 7.5% SDS-polyacrylamide gel and electropholeted onto nitrocellulose filter membranes. Then the membranes were incubated with a specific antibody against HA (Bethyl Laboratories, Montgomery, TX, U.S.A.). After washing, the membranes were incubated with Alexa Fluor 680-conjugated anti-rabbit or IRDye800-conjugated anti-mouse antibodies followed by visualization using infrared imaging.

**Purification of Recombinant 200-kDa sTNX from Conditioned Medium**
To cultivate adherent 200-kDa sTNX-expressing cell lines in suspension culture, sequential adaptation of the 200-kDa sTNX-expressing cell line from serum-supplemented medium to serum-free CHO-S-SFM II medium was performed. To isolate the recombinant 200-kDa sTNX from the conditioned medium of sTNX-expressing cells in suspension culture, approximately 11 of the conditioned medium was collected and centrifuged to remove cell debris. Subsequently, the cleared conditioned medium was precipitated with 50% ammonium sulfate by stirring overnight at 4 °C. After centrifugation for 20 min at 12000 rpm, the pellet was dissolved in 10 ml of high-salt buffer containing 0.5 m NaCl and 20 mM Tris–HCl (pH 7.5) and dialyzed overnight against the same buffer. Sepharose 4B resin (Sigma-Aldrich) coupled with the anti-mouse tenascin-X antibody pAb10d21120 was resuspended in the dialyzed sample and incubated overnight at 4 °C. The resin was then washed extensively in the high-salt buffer. The antigen was eluted with 200 mM triethylamine in the high-salt buffer, and the eluate was immediately neutralized with 2 mM Tris–HCl (pH 7.0). The eluate containing sTNX was analyzed by 7.5% SDS-PAGE under reducing conditions and visualized by silver staining (Wako). Approximately 6 μg of the recombinant sTNX was obtained from 2 liters of the conditioned medium. As a negative control, the proteins were purified from conditioned medium of mock cells that carry an empty expression vector.

**Cell Proliferation Assay by 5-Ethynyl-2'-deoxyuridine (EdU) Labeling**
**UV** 2 cells were grown overnight in 8-well culture slides (BD Biosciences, San Jose, CA, U.S.A.) in DMEM supplemented with 10% fetal bovine serum (FBS), penicillin, and streptomycin. Then the conditioned medium was changed to fresh medium containing 0.5% FBS and VEGF_{65} (VEGF-A_{65}) (25 ng/ml) (PeproTech, Rocky Hill, NJ, U.S.A.), bFGF (25 ng/ml) (PeproTech) and/or recombinant 200-kDa sTNX (10, 50, 100 ng/ml). As a negative control, conditioned medium from mock cells was added. EdU labeling of proliferative cells was done according to the instruction manual for Click-iT™ EdU Alexa Fluor High-Through Put Imaging (HCS) Assay (Invitrogen). After 32 h, EdU was added to the culture medium at the concentration of 10 μM for 10 h. After labeling, cells were washed three times with phosphate-buffered saline (PBS) and were fixed with 4% paraformaldehyde for 15 min. After rinsing with 3% bovine serum albumin (BSA) in PBS, cells were permeabilized with 0.5% Triton X-100 in PBS for 20 min. Then cells were stained for 30 min with Click-iT™ reaction cocktails (Invitrogen). Nuclear DNA was visualized with 4',6-diamidino-2-phenylindole (DAPI) (1 μg/ml). The cells were then observed using an All-In-One (BZ-8000) fluorescent microscopy (Keyence, Osaka, Japan). Then stained cells were counted.

**RESULTS**

**Easier Release of sTNX into Conditioned Medium than That of 450-kDa iTNX and 250-kDa iTNX**
As shown in our previous study,^{20} 200-kDa sTNX is abundantly present in serum. Therefore, we speculated that 200-kDa sTNX is more easily released than 450-kDa iTNX from cells into ex-
In order to examine how 200-kDa sTNX is easily released to extracellular space, at first we attempted to construct an expression plasmid, pSec-F-sTNX, encoding FLAG-tagged 200-kDa sTNX as described in Materials and Methods. As a control, we also constructed an expression plasmid encoding FLAG-tagged 450-kDa iTNX and an expression plasmid encoding FLAG-tagged 250-kDa iTNX, which lacks the region of 200-kDa sTNX from 450-kDa iTNX (Figs. 1A, B, C). We transfected these plasmids into CHO cells and established stable cell lines expressing 200-kDa sTNX, 450-kDa iTNX and 250-kDa iTNX. To confirm the expression of each TNX in the stable cell lines, cell lysates were prepared and immunoblot analyses of each TNX were performed with the use of two polyclonal antibodies, pAbM11 and pAb10d211, specific for different FNIII repeats of mouse TNX. Since the pAbM11 and pAb10d211 antibodies recognize the 11th FNIII repeat (M11) and the 25th to the 30th FNIII repeat (M25—M30) of mouse TNX, respectively (Fig. 1D), it was expected that the pAbM11 (α-M11) antibody would detect 250-kDa iTNX and 450-kDa iTNX, while the pAb10d211 (α-10d211) antibody would detect 200-kDa sTNX and 450-kDa iTNX. As expected, 200-

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**Fig. 1. Establishment of 200-kDa sTNX, 250-kDa iTNX and 450-kDa iTNX-Expressing Cell Lines**

Schematic diagrams of the domain structure of each recombinant TNX, recombinant 450-kDa iTNX (A), 250-kDa iTNX (B), and 200-kDa sTNX (C). The recombinant protein is depicted as a linear array from the N-terminus to C-terminus showing the central domain (segment of circle), heptad repeats (wavy line), 18.5 EGF-like repeats (diamonds), 31 FNIII-like repeats (M1—M31) (rectangles) and the fibrinogen-like domain (circle). The horizontal boxes under the diagrams of each recombinant TNX show the encompassed regions of the plasmid DNA clones used for the construction of the recombinant TNX with its designation of each plasmid. The encompassed domain region of each plasmid used is indicated in parenthesis. ^ indicates contiguity. (D) FNIII domains in which pAbM11 (α-M11) and pAb10d211 (α-10d211) antibodies are able to recognize each recombinant TNX. The pAbM11 antibody recognizes the 11th FNIII (M11) repeat, whereas the pAb10d211 antibody recognizes the 25th to the 30th FNIII repeats of mouse TNX. (E) Expression of 200-kDa sTNX, 250-kDa iTNX and 450-kDa iTNX in corresponding cell lines. Cell lysates from mock cells (lane 1), 250-kDa iTNX-expressing cells (lane 2), 200-kDa sTNX-expressing cells (lane 3), and 450-kDa iTNX-expressing cells (lane 4) cell lines were immunoblotted with anti-FLAG (α-FLAG), pAbM11 (α-M11), or pAb10d211 (α-10d211) antibody.
kDa sTNX was only detected by the pAb10d211 antibody, whereas 250-kDa iTNX was only detected by the pAbM11 antibody (Fig. 1E). Furthermore, 450-kDa iTNX was detected by both antibodies, and the anti-FLAG antibody detected all of three recombinant TNX forms. However, neither of these antibodies detected the proteins from lysates of mock cells carrying an empty pSecTag2/HygroB vector. These results indicated that the cell lines bearing the 200-kDa sTNX, 250-kDa iTNX and 450-kDa iTNX recombinant plasmids, express the corresponding proteins precisely.

Next, in order to compare the amounts of 200-kDa sTNX, 250-kDa iTNX, and 450-kDa iTNX released into conditioned medium from these established cell lines for 48 h, we detected each TNX both in cell lysates that have a certain fixed amount of total proteins and in its corresponding conditioned medium. (Lower panel, Cell) Various amounts of cell lysates from 250-kDa iTNX (lanes 1—3), 200-kDa sTNX (lanes 4—6), and 450-kDa iTNX (lanes 7—9) were immunoblotted with anti-FLAG antibody. Ten (lanes 1, 4, 7), fifty (lanes 2, 5, 8), and seventy-five (lanes 3, 6, 9) micrograms of cell lysates from each cell line were applied to SDS/PAGE. (Upper panel, CM) The conditioned medium corresponding to the amount of each cell lysate was immunoprecipitated with anti-FLAG M2 monoclonal antibody and the precipitates were blotted with the anti-FLAG antibody. (B) Quantification of the percentage of the released TNX into the conditioned medium versus TNX in the cell lysate for 250-kDa iTNX, 200-kDa sTNX and 450-kDa iTNX-expressing cell lines. Bands in (A) were scanned with an Odyssey apparatus, and their relative intensities are represented by histograms.

200-kDa sTNX Strongly Binds to VEGF-A and VEGF-B

We have reported that a splice isoform of 340-kDa iTNX that lacks M3 and M15—M22 FNIII repeats binds to VEGF-A and VEGF-B. Since we were interested in knowing whether 200-kDa sTNX also interacts with VEGF-A and VEGF-B, the ability of VEGF-A and VEGF-B to bind to 200-kDa sTNX was investigated by a co-immunoprecipitation assay. As a comparison, the abilities of 250-kDa iTNX and 450-kDa iTNX to bind to VEGF-A and VEGF-B were also investigated. To observe the interaction of each form of TNX with VEGF-A and VEGF-B in vivo, expression vectors for each FLAG-tagged form of TNX and HA-tagged VEGF-A or HA-tagged VEGF-B were together transfected into human 293T cells. Forty-eight hours after transfection, the cell extract was prepared and the proteins in the extract were first immunoprecipitated with an anti-HA antibody [upper panel, VEGF-A-HA (A) or VEGF-B-HA (B)]. To confirm the expression of each TNX, 1/100 amount of the proteins (input) used for the immunoprecipitation reactions was run in parallel and immunoblotted with an anti-FLAG antibody [middle panel, F-TNX (A and B)]. To confirm equivalent levels of the expression of HA-tagged VEGF-A (A) and VEGF-B (B), 1/50 amount of the proteins (input) used for the immunoprecipitation reactions was run in parallel and immunoblotted with an anti-HA antibody [lower panel, VEGF-A-HA (A) or VEGF-B-HA (B)]. IP, immunoprecipitation; IB, immunoblot. Relative binding abilities of each TNX to VEGF-A (A) and VEGF-B (B) are also shown. The band intensities of 450-kDa iTNX [lane 4 in upper panels in both (A) and (B)] was considered as 1.0.
cell extracts (Fig. 3). The relative binding ability of each TNX form to VEGF-A or VEGF-B was compared by measuring the intensity of each band. It was found that the abilities of 200-kDa sTNX to VEGF-A and VEGF-B were approximately 8-fold and 38-fold greater, respectively, than those of 450-kDa iTNX. These results indicate that 200-kDa sTNX strongly binds to VEGF-A and VEGF-B, compared to the binding abilities of 250-kDa iTNX and 450-kDa iTNX.

200-kDa sTNX Possesses Weak Activity for Proliferation of Endothelial Cells  We previously showed that 340-kDa iTNX alone has weak activity for proliferation of endothelial cells, whereas 340-kDa iTNX in combination with VEGF-B186 strongly promotes endothelial cell proliferation compared with the effect of VEGF-B186 alone.27) It is known that VEGF-A and basic fibroblast growth factor (bFGF), also known as FGF-2, stimulate endothelial cell proliferation.30) To investigate the effect of purified 200-kDa sTNX with VEGF-A165 on endothelial cell proliferation, UV2 cells were stimulated by purified recombinant 200-kDa sTNX alone and in combination with VEGF-A165 and bFGF. Then cell proliferation was examined by analysis of EdU incorporation into UV2 cells. As shown in Fig. 4A, VEGF-A165 increased EdU incorporation into DNA of UV2 cells (1.6-fold induction of that of non-treated cells). Interestingly, 200-kDa sTNX alone also stimulated endothelial cell proliferation to an extent similar to that induced by VEGF-A165 (1.4-fold induction of that of non-treated cells at 100 ng/ml). Furthermore, 200-kDa sTNX together with VEGF-A165 increased EdU incorporation into DNA by 1.9 fold compared with that of non-treated cells. These results demonstrate that the effect of purified 200-kDa sTNX with VEGF-A165 on endothelial cell proliferation is greater than that of the 200-kDa sTNX alone.

We also examined the effect of 200-kDa sTNX together with bFGF on endothelial cell proliferation. As shown in Fig. 4C, bFGF increased EdU incorporation to 1.5 fold compared with that of non-treated cells. Moreover, 200-kDa sTNX together with bFGF increased the 2.0-fold incorporation compared with that of non-treated cells, demonstrating again that the effect of purified 200-kDa sTNX with bFGF on endothelial cell proliferation is greater than that of 200-kDa sTNX alone. These results indicate that 200-kDa sTNX alone has weak activity for endothelial cell proliferation and that 200-kDa sTNX in combination with growth factors such as VEGF-A165 and bFGF shows slightly stronger endothelial cell proliferation activity than that of 200-kDa sTNX alone.

DISCUSSION

This is the first report on characterization of 200-kDa sTNX. The amount of 200-kDa sTNX released into conditioned medium is 7-fold larger than that of 250-kDa iTNX or 450-kDa iTNX. In addition, sTNX strongly binds to VEGF-A and VEGF-B compared with the binding abilities of other iTNX proteins. The purified 200-kDa sTNX both alone and in combination with VEGF-A or bFGF possesses weak activity for endothelial cell proliferation.

The binding sites of mouse 450-kDa iTNX to cells have not yet been determined. However, for bovine TNX it has been reported that the contiguous b9 and b10 FNIII repeats and the fibrinogen domain participate in cell adhesion via αvβ3 integrin and β1-containing integrin receptors, respectively.31) Furthermore, the same group31 identified two FNIII repeats, b10 and b11, of bovine TNX in which heparin-binding sites exist. They also showed that the b10 and b11 repeats are involved in cell adhesion mediated by heparan sulfate cell
surface receptors. We previously performed phylogenetic analysis for 94 FNIII repeats derived from human, mouse and bovine TNX. The results of this analysis showed that the b9, b10 and b11 FNIII repeats of bovine TNX have high similarity to the 10th (M10), 11th (M11) and 12th (M12) FNIII repeats in the 31 FNIII repeats of mouse TNX, respectively. Therefore, it is possible that mouse M11 and M12 FNIII repeats of 250-kDa iTNX and 450-kDa iTNX possess heparin-binding activity. Since heparan sulfate/heparin (e.g., heparan sulfate cell surface receptors) is prominently present on the cell membrane, 250-kDa iTNX and 450-kDa iTNX with M11 and M12 repeats might interact with cell membranes and adhere to cells. Actually, in this study (Fig. 2), we showed that 200-kDa sTNX, which lacks the 250-kDa N-terminal region (including the M11 and M12 FNIII repeats) of 450-kDa iTNX, is released into conditioned medium and is not easily maintained on the cell surface compared with 250-kDa iTNX and 450-kDa iTNX.

200-kDa sTNX strongly bound to VEGFs compared with the binding abilities of 250-kDa iTNX and 400-kDa iTNX. We previously examined the interaction of 340-kDa iTNX with VEGF-B and found that the M11—M31 FNIII repeats and a fibrinogen-like domain with VEGF-B might be localized within M23, M24 and M25 FNIII repeats. Although 250-kDa iTNX does not include the M23, M24 and M25 repeats, it binds to VEGF-B weakly compared with 200-kDa sTNX. This might be due to the fact that 250-kDa iTNX has the 10th (M10) and 15th (M15) FNIII repeats with high similarity to the M23, M24 and M25 repeats. On the other hand, for the reason why 450-kDa iTNX having M23, M24 and M25 repeats binds to VEGFs very faintly, we speculated that the M23, M24 and M25 repeats are masked by N-terminal regions of 450-kDa iTNX which 200-kDa sTNX lacks and consequently VEGFs cannot approach the M23, M24 and M25 repeats of 450-kDa iTNX.

The 200-kDa sTNX alone displays weak endothelial cell proliferation activity (Fig. 4). The reason for this is not clear at present. This proliferation might be mediated by unidentified receptor(s) for 200-kDa sTNX. However, it is also possible that 200-kDa sTNX directly interacts with VEGF receptor (VEGFR-1 or VEGFR-2) and transmits signals from the receptor to regulate gene expression, leading to endothelial cell proliferation. We performed preliminary experiments on the interaction of 200-kDa sTNX with VEGFR-1, and we found that the sTNX interacts with VEGFR-1 (data not shown). However, it has not been determined whether the interaction results in activation of VEGFR-1 and transmits signals to its downstream targets.

Previously, we showed that 340-kDa iTNX, a small splice isoform that lacks M3 and M15—M22 FNIII repeats of 450-kDa iTNX, in combination with VEGF-B increased endothelial cell proliferation by 3.9 fold compared with that induced by VEGF-B alone. On the other hand, the present study demonstrated that 200-kDa sTNX in combination with VEGF-A increased endothelial cell proliferation by only 1.2 fold compared with that induced by VEGF-A alone. The major reason why 340-kDa iTNX with VEGF-B possesses a high level of endothelial cell proliferation activity, but 200-kDa sTNX with VEGF-A has a little such activity, might be due to the difference of cell proliferation assay system and cells used between the previous study and this study. Namely, in previous study the cell proliferation was examined through an analysis of [3H]thymidine incorporation in mouse endothelial UV2 cells. Other conceivable reason might be as follows. Since 340-kDa iTNX has M11 and M12 FNIII repeats that bear possible heparin/heparan sulfate-binding sites, as in the case of 450-kDa iTNX 340-kDa iTNX might interact with heparan sulfate cell surface receptors and be maintained on cell membranes. The 340-kDa iTNX associated with cell membranes might capture VEGF-B close to the cell membrane and facilitate presentation of VEGF-B to its receptor VEGFR-1. On the other hand, as mentioned above, the rate of association of the 200-kDa sTNX with the cell membrane is lower than that of the 340-kDa iTNX and the sTNX is easily released into the extracellular environment. Due to the strong binding of VEGF-A and 200-kDa sTNX, the released sTNX might catch VEGF-A away from the cell membrane where VEGF-A receptors such as VEGFR-2 and VEGFR-1 are localized and transmit signals for cell proliferation. However, even though VEGF-A to some extent was trapped by the released sTNX in the extracellular environment, sTNX with VEGF-A would also stay on the cell membrane to some extent since 79.2% of 200-kDa sTNX exist in the cell and on the cell (Fig. 2B). Therefore, the left VEGF-A with sTNX associated with cell membranes might lead to the 1.2-fold induction of cell proliferation compared with that induced VEGF-A alone.

Further studies are needed to determine the endothelial cell proliferation activity of 450-kDa iTNX alone, 200-kDa sTNX with 450-kDa iTNX and also that of 200-kDa sTNX and 450-kDa iTNX with VEGF-A or VEGF-B treated at the same time. It would be of great interest to investigate whether 200-kDa sTNX gives influence to the endothelial cell proliferation activity that 450-kDa iTNX might possess.

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