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Knockdown of legumain inhibits cleavage of annexin A2 in the mouse kidney

Takuya Yamane\textsuperscript{a*}, Rei Hachisu\textsuperscript{b}, Motoki Yuguchi\textsuperscript{b}, Keisuke Takeuchi\textsuperscript{c}, Sato Murao\textsuperscript{a}, Yoshio Yamamoto\textsuperscript{d}, Hisakazu Ogita\textsuperscript{c}, Toshihide Takasawa\textsuperscript{b}, Iwao Ohkubo\textsuperscript{e*} and Hiroyoshi Ariga\textsuperscript{a}

\textsuperscript{a}Graduate School of Pharmaceutical Sciences, Hokkaido University, Kita-ku, Sapporo 060-0812, Japan
\textsuperscript{b}Division of Bioscience, Hokkaido System Science Co. Ltd., Kita-ku, Sapporo 001-0932, Japan
\textsuperscript{c}Division of Molecular Medical Biochemistry, Department of Biochemistry and Molecular Biology, Shiga University of Medical Science, Seta, Otsu 520-2192, Japan
\textsuperscript{d}Department of Ecology and Molecular Biology, Mie University, Iga, Mie 518-0131, Japan
\textsuperscript{e}Department of Nutrition, School of Nursing and Nutrition, Tenshi College, Higashi-ku, Sapporo 065-0013, Japan

* Corresponding authors:

Takuya Yamane
Graduate School of Pharmaceutical Sciences, Hokkaido University, Sapporo 060-0812, Japan
Tel: +81-11-706-3711; Fax: +81-11-706-4988; E-mail: t-yamane@pharm.hokudai.ac.jp

Iwao Ohkubo
Department of Nutrition, School of Nursing and Nutrition, Tenshi College, Higashi-ku, Sapporo 065-0013, Japan
Tel: +81-11-741-1051 (ext. 223); E-mail: ohkubo@tenshi.ac.jp
Abstract

Legumain (EC 3.4.22.34) is an asparaginyl endopeptidase. Strong legumain activity was observed in the mouse kidney, and legumain was highly expressed in tumors. We previously reported that bovine kidney annexin A2 was co-purified with legumain and that legumain cleaved the N-terminal region of annexin A2 at an Asn residue in vitro. In this study, to determine whether annexin A2 is cleaved by legumain in vivo, siRNA-lipoplex targeting mouse legumain was injected into mouse tail veins. Mouse kidneys were then isolated and the effect of knockdown of legumain expression on annexin A2 cleavage was examined. The results showed that both legumain mRNA and protein expression levels were decreased in the siRNA-treated mouse kidneys and that legumain activity toward a synthetic substrate, Z-Ala-Ala-Asn-MCA, was decreased by about 40% in the kidney but not in the liver or spleen. Furthermore, cleavage of annexin A2 at the N-terminal region was decreased in the mouse kidney that had been treated with the legumain siRNA-lipoplex. These results suggest that legumain siRNA was delivered to the kidney by using LipoTrust and that the reduced legumain expression inhibited legumain-induced degradation of annexin A2 in vivo.

Key Words: Legumain, annexin A2, cationic liposome, siRNA, in vivo, mouse kidney
1. Introduction

Legumain (EC 3.4.22.34) is an asparaginyl endopeptidase belonging to the cysteine peptidase C13 family [1]. Legumain activity has been detected in a number of mammalian tissues, including the kidney, placenta, spleen, liver and testis, and the highest level of activity was detected in the kidney [2]. We previously reported that legumain is mainly expressed in proximal tubules of the rat kidney [3]. We further reported that legumain might have an important role in remodeling of the extracellular matrix through degradation of fibronectin in renal proximal tubular cells [4]. It has recently been suggested that legumain plays an important role in tumor growth/metastasis, carotid artery-atherosclerosis development [5-10], hemophagocytic syndrome [11] and formation of human unstable carotid plaque [12].

Annexin A2, also named annexin II, calpactin I, lipocortin II, chromobindin 8 and placental anticoagulant protein IV, was co-purified with legumain from the bovine kidney, and co-purified annexin A2 lacked the N-terminal region [3]. The N-terminal region of annexin A2 contains both p11 subunit-binding sites required for hetero-tetramer formation of annexin A2 and phosphorylation sites of pp60^src and protein kinase C (PKC) [13]. Annexin A2 is abundantly expressed in the receptor-recycling compartments of rat liver hepatocytes [14] and is required for aquaporin 2 (AQP2) trafficking in renal cells [15, 16]. Furthermore, it has been shown that the N-terminal region of annexin A2 has a role in endosomal membrane fusion [17] and that phosphorylation of Tyr-23 is essential for proper endosomal association and for annexin A2 function [18].

Cationic liposomes have been used in drug delivery systems, and LipoTrust, a novel cationic liposome, has been used as an efficient delivery system in vivo [19, 20]. In this study, to examine annexin A2 degradation by legumain in vivo, siRNA-lipoplex targeting mouse legumain was injected into the mouse tail veins. The results showed that both the expression levels of legumain mRNA and protein and the legumain activity toward a synthetic substrate

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were decreased and that degradation of annexin A2 at an N-terminal Asn residue was inhibited in legumain siRNA-lipoplex-treated mouse kidneys, indicating an important role of legumain in annexin A2 functions.

2. Materials and Methods

2.1. Materials

Sequences of siRNA targeting mouse legumain were designed by Hokkaido System Science (Sapporo, Japan) using the B-Algo™ algorithm. The sense and antisense strands of siRNAs and LipoTrust™ EX Oligo were also chemically synthesized by Hokkaido System Science. NIH3T3 cells were purchased from American Tissue Culture Collection (Manassas, VA, USA). RNAlater and RNeasy Mini were purchased from Qiagen (Venlo, Netherlands). SuperScript III, Bioanalyzer capillary chip and Z-Ala-Ala-Asn-MCA were obtained from Invitrogen (St. Louis, MO, USA), Agilent Technologies (Santa Clara, CA, USA) and Peptide Institute (Osaka, Japan), respectively. Recombinant human annexin A2 and human legumain were purchased from ProSpec-Tany TechnoGene (Ness Ziona, Israel) and R&D Systems (Minneapolis, USA), respectively. Balbc mice were purchased from Sankyo Lab. (Tokyo, Japan). All other chemicals were of analytical grade and purchased from Wako Pure Chemicals (Osaka, Japan).

2.2. Preparation of anti-annexin A2 antibodies

Annexin A2 purified from the bovine kidney was used as an immunogen to establish rabbit anti-annexin A2 polyclonal antisera. Briefly, one hundred fifty µg of purified annexin A2 dissolved in PBS was first injected into rabbits and then injected five times as booster injections at 10-day intervals. Antigen solutions were mixed with equal volumes of complete adjuvant and with incomplete adjuvant and were used for the initial and subsequent injections,
respectively. An IgG fraction of an anti-annexin A2 antibody was purified from antisera using a protein A-Sepharose column.

2.3. Western blotting and antibodies

To examine the degradation levels of annexin A2 and expression levels of legumain in the mouse kidney, proteins were extracted from cultured cells or from the mouse kidney in a solution containing 50 mM sodium citrate buffer pH 5.0. Proteins were then separated on a 12.5% polyacrylamide gel and subjected to Western blotting with respective antibodies. Proteins on the membrane were reacted with an IRDye 800- (Rockland, Philadelphia, PA, USA) or Alexa Fluor 680-conjugated secondary antibody (Molecular Probes, Eugene, OR, USA) and visualized by using an infrared imaging system (Odyssey, LI-COR, Lincoln, NE, USA). Antibodies used were anti-legumain (1:500, Santa Cruz, CA, USA) and anti-annexin II (1:1000) antibodies.

2.4. Inhibition of annexin A2 degradation in bovine kidney homogenates

To examine annexin A2 degradation by legumain under an acidic condition, 8 µg of bovine kidney homogenates was incubated in a solution containing 50 mM sodium citrate (pH 5.0) and 2 mM 2-mercaptoethanol in the presence or absence of 1 mM N-ethylmaleimide for 30 min at 37°C, separated on a 12.5% SDS-polyacrylamide gel, and subjected to Western blotting with an anti-annexin A2 antibody.

2.5. Preparation of siRNAs

Three siRNAs targeting mouse legumain (B-Bridge, Cat#: SMF27A-1537-1, SMF27A-1537-2 and SMF27A-1537-3) were used. Nucleotide sequences of the siRNAs used were as follows: LGMN (SMF27A-1537-1; siRNA1): sense,
5′-GGGCAAAGGGUCUGGAAAATT-3’ and antisense,
5′-UUUUCAGACCCUUGCCCTT-3′; LGMN (SMF27A-1537-2; siRNA2): sense,
5′-GGGAAACUGCUGAGAGACATT-3’ and antisense,
5′-UGUCUCUCAGCAGUUUCCCTT-3′; and LGMN (SMF27A-1537-3; siRNA3): sense, 5′-
GGGAAGGAAUCGUCUGAGATT-3’ and antisense, 5′-
UCUCAGACGAAUCCUCCCTT-3′. Nucleotide sequences of the negative control siRNA
were as follows: sense, 5′-CGUACGCGGAAUACUUCGATT-3′ and antisense,
5′-UCGAAGUAUUCGCGCUACCGTT-3′.

2.6. Formation of transfection complex
siRNAs were transfected into NIH3T3 cells by using LipoTrust™ EX Oligo (Hokkaido
System Science). A vial of LipoTrust was reconstituted of 1 ml of nuclease-free water, and 10
µl of LipoTrust solution was mixed with 100 pmol of siRNA. After incubation of the mixture
for 20 min at room temperature, legumain siRNA-lipoplex was formed. For in vivo study, a
vial of LipoTrust was reconstituted of 1 ml of nuclease-free water containing 1.6, 3.2 and 6.4
nmol siRNAs. After incubation of the mixture for 20 min at room temperature, legumain
siRNA-lipoplex was formed.

2.7. Characterization of legumain siRNA-lipoplex
The size dispersion of legumain siRNA-lipoplex was determined by N4 Plus (Beckman
Coulter, Miami, FL, USA) and the zeta potential was measured using a Zeta Potential
Analyzer Ver.3.47 (Brookhaven Instruments Corp., NY, USA).

2.8. Cell culture and knockdown of legumain expression in vitro
NIH3T3 cells were cultured in Dulbecco’s modified Eagle’s medium supplemented with 10%
calf serum at 37°C in a humidified atmosphere containing 5% CO2. The cells were transfected with Legumain siRNA-lipoplex. Forty-eight hours after transfection, cells were collected and their total RNAs were extracted.

2.9. Knockdown of mouse legumain expression in vivo

C57BL/6 mice were purchased from Sankyo Lab. (Tokyo, Japan). Eighty µl of each of the legumain siRNA-lipoplex solutions (1.6 and 3.2 and 6.4 nmol/ml siRNA) was injected into mouse tail veins. Lipoplex solution containing negative control siRNA was also injected. At 48 hours after injection, mouse kidneys were extracted and their total RNAs and proteins were extracted. All animal experiments were carried out in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals, and the protocols were approved by the Committee for Animal Research at Hokkaido University (permit number: 08-0484).

2.10. Quantification of mRNA level by reverse transcription polymerase chain reaction

Tissues were dissected and quickly frozen in an RNAlater solution. Total RNAs were prepared from approximately 20 mg of the kidney using an RNeasy mini kit, and quality of RNA was examined using Bioanalyzer. Reverse transcription was carried out in a mixture containing 500 ng of total RNAs, Superscript III and specific primers under the conditions of 96°C for 15 min, 32 cycles of 96°C for 30 sec, 60°C for 30 sec, 72°C for 30 sec and 72°C for 5 min by using a QuantiTect SYBER Green kit (Qiagen) and Opticon II real-time PCR analyzer (Bio-Rad, Hercules, CA, USA). β-actin (ACTB) mRNA was also amplified and used as an internal control.

2.11. Proteolytic activity of legumain
Enzyme activity of legumain was examined by measuring the fluorometrical number (excitation, 380 nm; emission, 440 nm) of liberation of 7-amino-4-methylcoumarin (AMC) in a mixture containing 10 µl of 10 mM Z-Ala-Ala-Asn-MCA, 100 µl of 0.5 M sodium citrate buffer pH 5.0, 5 µl of 1 M 2-mercaptoethanol, 20 µl of enzyme solution and water (18 mΩ) in a total volume of 1 ml. After incubation of the mixture at 37°C for 30 min, 2 ml of 0.2 M acetic acid was added to the mixture to stop the reaction. One unit of activity was defined as the amount of enzyme that hydrolyzed 1 µmol of the substrate per min.

2.12. Statistical analyses

Data are expressed as means ± S.D for mouse experiments. Statistical analyses were performed using analysis of variance (one-way ANOVA) followed by unpaired Student’s t-test. For comparison of multiple samples, the Tukey-Kramer test was used.

3. Results

3.1. Effect of legumain siRNAs on legumain expression in NIH3T3 cells

To examine the effect of legumain siRNAs on the expression of legumain mRNA, three siRNA-lipoplexes targeting legumain were transfected into NIH3T3 cells and the expression of legumain mRNA was examined by real-time PCR. As shown in Fig. 1A, the expression level of legumain mRNA was decreased by legumain siRNA-lipoplexes compared to that in control siRNA-transfected cells. The degrees of knockdown of legumain mRNA by legumain siRNA1, 2 and 3-lipoplexes were 42, 55 and 66%, respectively. The siRNA #3 having the strongest knockdown activity toward legumain expression was therefore used for further studies.

3.2. Effect of legumain siRNAs on legumain expression and activity in the mouse kidney
To examine the effect of legumain siRNAs on legumain expression in vivo, legumain siRNA3-lipoplex and negative siRNA-lipoplex were injected into the mouse tail veins and the expression level of legumain in the mouse kidney was examined by real-time PCR. As shown in Fig. 1B, legumain expression level was decreased by legumain siRNA-lipoplex containing 3.2 nmol/ml siRNA3 to 80% of that by negative siRNA-lipoplex, but legumain siRNA-lipoplex containing 6.4 nmol/ml siRNA3 did not show a knockdown effect. To determine the factor(s) that affects knockdown efficiency of legumain siRNA-lipoplexes, zeta potential and size dispersion of legumain siRNA-lipoplexes containing 1.6, 3.2 and 6.4 nmol of siRNA3 were measured using a Zeta Potential Analyzer and N4 Plus, respectively. The results showed that the size and zeta potential of legumain siRNA-lipoplex containing 6.4 nmol/ml of siRNA3 were increased more than 2-fold and decreased by 20%, respectively, compared with those of legumain siRNA-lipoplex complex containing 3.2 nmol/ml of siRNA3 (Figs. 1C and 1D).

The effect of legumain siRNA3-lipoplex on legumain activity in mouse tissues was then examined. As shown in Fig. 1E, legumain activity in the kidney, but not that in the liver or spleen, was decreased by injection of legumain siRNA-lipoplex containing 3.2 nmol/ml of siRNA3 to about 40% of that in negative siRNA-lipoplex-injected mice.

3.3. Degradation of annexin A2 by legumain in the mouse kidney

To examine degradation of annexin A2 by cysteine proteases under an acidic condition, bovine kidney homogenates were incubated in a mixture containing 1 mM N-ethylmaleimide (NEM), an inhibitor of cysteine proteases, and proteins in the mixture were separated on a polyacrylamide gel and subjected to Western blotting with an anti-annexin A2 antibody. Purified annexin A2, which lacks the N-terminal region, was also loaded on the gel as a positive control. As shown in Fig. 2A, annexin A2 degradation was observed in the bovine
kidney homogenate but was inhibited by the presence of NEM, indicating that annexin A2 degradation was carried out by a cysteine protease(s). Since legumain is a cysteine protease, the possibility that legumain degrades annexin A2 in vivo was examined. To do that, legumain siRNA3-lipoplex and negative siRNA-lipoplex were injected into mouse tail veins. Forty-eight hrs after injection, proteins in the kidney were analyzed by Western blotting with anti-legumain, anti-annexin A2 and anti-actin antibodies. Two bands of annexin A2 corresponding to full-sized and cleaved annexin A2 were obtained (Fig. 2B), and the levels of cleaved and full-sized annexin A2 in legumain siRNA3-lipoplex-injected mice were decreased and increased to about 60% and 30% of those in negative siRNA-lipoplex-injected mice, respectively (Figs. 2C and 2D), suggesting that legumain directly cleaves annexin A2 or affects cleavage of annexin A2.

3.4. Direct cleavage of annexin A2 by legumain in vitro

To examine degradation of annexin A2 by legumain, 6 nmol of recombinant human annexin A2 was reacted with 0.12, 0.24 and 1.2 nmol of recombinant human legumain and subjected to Western blotting. As shown in Figs. 3A and 3B, annexin A2 was cleaved by recombinant legumain in a dose-dependent manner, indicating that legumain directly cleaved annexin A2.

4. Discussion

In this study, we showed that legumain siRNA-lipoplex containing 3.2 nmol/ml siRNA3 knocked down legumain mRNA expression by about 60% and 20% in NIH3T3 cells and in the mouse kidney, respectively (Figs. 1A and 1B). These results suggest that legumain siRNA was delivered to the kidney and that more than 40% of legumain siRNA3-lipoplex was degraded during a period of 48 hours after injection into mouse tail veins. Legumain siRNA-lipoplex containing 6.4 nmol/ml siRNA3, on the other hand, did not affect the
expression level of legumain (Fig. 1B). The size and zeta potential of legumain siRNA-lipoplex containing 6.4 nmol/ml of siRNA3 were increased more than 2-fold and decreased by 20% compared to those of legumain siRNA-lipoplex complex containing 3.2 nmol/ml of siRNA3 (Fig. 1C, D). These results indicate that the size and zeta potential of legumain siRNA3-lipoplex are important factors to obtain an efficient knockdown effect in the mouse kidney. We also found that legumain activity toward the synthetic substrate Z-Ala-Ala-Asn-MCA in the mouse kidney was decreased by about 40% of that in negative siRNA-lipoplex-injected mice, while legumain activities were not changed in the liver and spleen (Fig. 1E), suggesting that the knockdown effect of legumain siRNA3-lipoplex is tissue-specific.

Furthermore, we found that the cleavage level of annexin A2 was decreased in the legumain siRNA-treated mouse kidney (Fig. 2) and that legumain directly degrades annexin A2 (Fig. 3). These results suggest that legumain siRNA inhibited annexin A2 degradation under an acidic condition through the reduced expression of legumain in vivo. It has been reported that the N-terminal region of annexin A2 was cleaved by plasmin under the condition of neutral pH [21]. We previously suggested that legumain cleaves annexin A2 under an acidic pH condition, which resembles conditions in the endosome or lysosome [3]. Annexin A2 was identified as one of the AQP2-associated proteins [15]. The N-terminal region of annexin A2 is required for cAMP-induced AQP2 exocytosis in renal cells [16], and phosphorylation of the N-terminal region is essential for proper endosomal association [18]. It is therefore thought that legumain regulates functions of annexin A2, including AQP2 trafficking and endosomal association-mediated cleavage of annexin A2 in the endosome.

In conclusion, we found that legumain siRNA3-lipoplex decreased the expression of legumain mRNA and protein and its proteolytic activity in vivo, thereby inhibiting annexin A2 degradation by legumain under an acidic condition. Although legumain was suggested to
regulate the function of annexin A2 in the endosome and lysosome in the kidney, future studies are needed to reveal the molecular mechanism underlying the regulation by legumain of annexin A2 functions such as fusion and trafficking of membrane proteins.

**Abbreviations:** Z, benzyloxycarbonyl; MCA, methylcumarinamide; siRNA-lipoplex, siRNA-cationic liposome complex; NEM, N-ethyl maleimide; 2-ME, 2-mercaptoethanol; PBS, phosphate buffered saline.

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**References**


**Figure legends**

**Fig. 1.** Effect of legumain siRNAs on legumain expression and activity

A. Lipoplexes containing three kinds of legumain siRNAs or negative control siRNA were transfected into NIH3T3 cells. At 48 hours after transfection, the expression levels of legumain and β-actin mRNA were examined by real-time PCR. Proteins extracted from NIH3T3 cells were also analyzed by Western blotting with anti-legumain and β-actin antibodies. Statistical analyses were carried out using the Tukey-Kramer test. Significance: *p<0.05, **p<0.01 versus negative control siRNA.

B. Legumain siRNA-lipoplex solutions containing 1.6, 3.2 and 6.4 nmol/ml siRNA or 1.6 nmol/ml control siRNA were injected into mouse tail veins. At 48 hours after injection, the mouse kidneys were extracted and their expression levels of mRNA and protein of legumain were examined by real-time PCR and by Western blotting.

C and D. Size dispersion of legumain siRNA-lipoplex was determined by N4 Plus, and the zeta potential was measured using a Zeta Potential Analyzer Ver.3.47.

E. Legumain siRNA-lipoplex solutions containing 3.2 nmol/ml siRNA or 1.6 nmol/ml negative siRNA-lipoplex were injected to mouse tail veins. At 48 hours after injection, the mouse liver, spleen and kidney were extracted and their proteins were extracted. Their legumain activities were then measured by fluorometrical assays against Z-Ala-Ala-Asn-MCA. Statistical analyses were carried out using the Tukey-Kramer test. **p<0.01, *p<0.05, N.S., not significant

**Fig. 2.** Effect of knockdown of mouse legumain expression on annexin A2 degradation *in vivo.*

A. Eight µg of bovine kidney homogenate was incubated with or without 1 mM NEM in a solution containing 50 mM sodium citrate buffer (pH 5.0) and 2 mM 2-ME. Annexin A2 was
analyzed by Western blotting with an anti-legumain antibody. Lane C indicates a band corresponding to purified bovine kidney annexin A2 lacking the N-terminal region.

B. Lipoplex solutions containing 3.2 nmol/ml legumain and negative control siRNAs were injected into mouse tail veins. At 48 hours after injection, the mouse kidneys were extracted and the expression levels of legumain, annexin A2 and β-actin were examined by Western blotting. Lane C indicates a band corresponding to purified bovine kidney annexin A2 lacking the N-terminal region.

C and D. Intensity of cleaved and non-cleaved annexin A2 bands shown in Fig. 2B was quantified. Statistical analyses were carried out using Student’s t-test. Significance: *p<0.05, **p<0.01

**Fig. 3.** Direct cleavage of annexin A2 by legumain

A. Six nmol of recombinant human annexin A2 was reacted with 0.12, 0.24 and 1.2 nmol of recombinant human legumain in a solution containing 50 mM sodium citrate buffer (pH 5.0) and 2 mM 2-ME at 37°C for 30 min. Annexin A2 was analyzed by Western blotting with an anti-annexin A2 antibody.

B. Intensity of the cleaved annexin A2 band shown in Fig. 3A was quantified. Statistical analyses were carried out using the Tukey-Kramer test. Significance: *p<0.05
Fig. 2

A

C

NEM

+  -

kDa

90

48

36

29

19

B

Legumain siRNA

Legumain

Annexin A2

Actin

Full-sized

cleaved

C

D

Relative protein expression

Legumain/Actin

%  %

0  100

40

80

20

Legumain siRNA

-  +

[Graph showing relative protein expression with Legumain siRNA]
A

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Annexin A2

B

Cleaved Annexin A2

Legumain (nmol)

Fig. 3