Proteolytic processing of lysyl oxidase–like-2 in the extracellular matrix is required for crosslinking of basement membrane collagen IV

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Lysyl oxidase–like-2 (LOXL2) is an enzyme secreted into the extracellular matrix that crosslinks collagens by mediating oxidative deamination of lysine residues. Our previous work demonstrated that this enzyme crosslinks the 7S domain, a structural domain that stabilizes collagen IV scaffolds in the basement membrane. Despite its relevant role in extracellular matrix biosynthesis, little is known about the structural requirements of LOXL2 that enable collagen IV crosslinking. In this study, we demonstrate that LOXL2 is processed extracellularly by serine proteases, generating a 65-kDa form lacking the first two scavenger receptor cysteine-rich domains. Site-specific mutagenesis to prevent proteolytic processing generated a full-length enzyme that is active in vitro toward a soluble substrate, but fails to crosslink insoluble collagen IV within the extracellular matrix. In contrast, the processed form of LOXL2 binds to collagen IV and crosslinks the 7S domain. Together, our data demonstrate that proteolytic processing is an important event that allows LOXL2-mediated crosslinking of basement membrane collagen IV.

Lysyl oxidase–like-2 (LOXL2) is a member of the lysyl oxidase family of copper-dependent amine oxidases (1). Like the rest of the family members, LOXL2 catalyzes the oxidative deamination of peptidyl lysine residues, which promotes the formation of lysyl-derived crosslinks in collagens and elastin (1–5). LOXL2-catalyzed crosslinks are essential posttranslational modifications in the biosynthesis and maintenance of the extracellular matrix (ECM), contributing to the tensile strength and stability of tissues (6). We have recently shown that LOXL2 is the main lysyl oxidase of the glomerular basement membrane, a specialized form of ECM that is an important functional component of the kidney glomerulus (7). Furthermore, together with peroxidasin-catalyzed sulfilimine crosslinking of the NC1 domain (8, 9), LOXL2-catalyzed crosslinking of the 7S domain contributes to stabilizing collagen IV networks, the main structural component of basement membranes (10–12). Other proposed biological functions of LOXL2 include regulation of tumor growth (13), metastasis (14), and angiogenesis (15). Because it is up-regulated in fibrotic diseases, LOXL2 has been proposed as an attractive pharmacological target to inhibit the characteristic increase in ECM deposition in fibrotic tissues (16–20).

The amino acid sequence of human LOXL2 consists of 774 residues organized by an N-terminal signal peptide contiguous to four N-terminal scavenger receptor cysteine-rich (SRCR) domains followed by a highly conserved lysyl oxidase catalytic domain. SRCR domains, commonly found on secreted and cell surface–bound proteins, have been proposed to mediate protein–protein interactions, ligand binding, and more recently deacetylation and deactylimination activity over peptidyl-acetyl lysines (21, 22). Although the predicted molecular mass derived from the amino acid sequence is about 84 kDa, LOXL2 is found as a ~100-kDa protein in both the intracellular and the extracellular compartments, most likely because of posttranslational modifications such as N-glycosylation (23). Interestingly, several studies in cultured cells and tissues, including our own, have identified a smaller extracellular 65-kDa form (7, 24–26). The addition of a protease inhibitor mixture in the conditioned media of MCF-7 cells prevented the formation of the 65-kDa form, suggesting that it results from proteolytic processing of the large 100-kDa form (27). This observation is particularly relevant as other members of the LOX family such as LOX and LOXL1 become active only after they are proteolytically processed by bone morphogenetic protein-1 (28, 29). Although the crosslinking function of LOXL2 has been characterized in many biological systems, it is not known whether proteolytic processing regulates its enzymatic activity (30).

In this study, we investigated the role of LOXL2 processing on the crosslinking of the 7S domain of collagen IV, a key structural component of basement membranes. We identified the cleavage site in LOXL2 sequence by classical Edman microsequencing and site-directed mutagenesis. Furthermore, using CRISPR-Cas9 system, we developed a LOXL2-null cell line that allowed us to explore the effect of proteolytic processing on LOXL2 enzymatic activity. Our results suggest that LOXL2 proteolytic processing is a regulatory step on collagen IV and basement membranes biosynthesis.
LOXL2 is proteolytically processed by serine proteases

To purify sufficient 65-kDa extracellular form of LOXL2 for the identification of the site of proteolytic cleavage by Edman microsequencing, we stably expressed the LOXL2 human ORF (NM_002318) in HEK293 cells (7). Immunoblotting characterization of HEK293 cells expressing this construct using a polyclonal antibody raised against the C-terminal region of the LOXL2 sequence showed that the intracellular fraction produced the full-length form of LOXL2 (FL) without any sign of proteolytic cleavage (Fig. 1A). In contrast, the conditioned media of these cells showed the presence of both full-length and processed forms of LOXL2, similar to those observed in other cell types (24–26), and glomeruli (7). We purified LOXL2 forms from cell media by affinity chromatography and fractionated proteins by SDS-PAGE. The 65-kDa processed form was excised from the gel and prepared for N-terminal Edman microsequencing. The results revealed the first 11 amino acid residues of processed form N-terminal end starting at Ala-318 of human LOXL2 sequence, indicating that proteolytic processing occurred at Lys-317. This site resides in the linker region between the SRCR2 and SRCR3 domains (Fig. 1B).

Further inspection of the sequence showed that a serine protease cleaves LOXL2 at Lys-317. To test this hypothesis, HEK293 cells stably expressing human LOXL2 were grown in media containing serine protease inhibitors including aprotinin, leupeptin, and PMSF (Fig. 1D). Immunoblotting analyses of the conditioned media from these cells showed that the formation of the 65-kDa processed form was significantly reduced by aprotinin and PMSF, but not leupeptin, suggesting that serine proteases may be responsible for LOXL2 cleavage. HEK293 cells were also cultured in the presence of decanoyl-RVKR-chloromethylketone (Dec-RVKR-CMK), an inhibitor that blocks the activity of members of the subtilisin-like family of proprotein convertases such as furin (31). As shown in Fig. 1D, Dec-RVKR-CMK inhibited the highest inhi-
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To evaluate the effect of proteolytic processing on amine oxidase activity, we performed site-directed mutagenesis to generate either full-length or processed forms of LOXL2. To block proteolytic processing and generate a full-length LOXL2 form, the two basic residues preceding the cleavage site were mutated from RK to GE. The introduction of a negatively charged glutamic acid at P1 was expected to block proteolytic processing by serine proteases (32–34). To evaluate the effect of proteolytic processing on amine oxidase activity, conditioned media of cells expressing LOXL2 wild type and mutants were assayed with Amplex Red assay (35). As shown in Fig. 2A, LOXL2BP, was used to describe this “blocked processing LOXL2 mutant” construct. As most serine proteases have a preference for Arg at P1 (33, 34), we generated a second mutant by changing Lys-317 to Arg-317 to potentially enhance proteolytic cleavage. The resulting K317R processing LOXL2 mutant was named LOXL2K317R.

To assess the effect of proteolytic processing on amine oxidase activity, conditioned media of cells expressing LOXL2 wild type and mutants were assayed with Amplex Red assay using 1,5-diaminopentane as substrate (35). As shown in Fig. 2A, initial studies in which only Lys-317 was mutated uncovered an anticipated atypical form of LOXL2 of about 70 kDa that is not observed in cells expressing wild-type LOXL2. A careful inspection of LOXL2 sequence identified Arg-256 as a potential cleavage site of amine oxidase activity (7). Interestingly, the nonreduced 7S dodecamers extracted after collagenase digestion of the ECM derived from LOXL2-null PFHR9 cells showed slightly faster electrophoretic mobility in comparison to the wild-type cells (Fig. 3A). Immunoblot analysis of the ECM-derived fraction, where both 100-kDa and 65-kDa forms are normally deposited in the wild-type cells, showed that both LOXL2 forms are absent in the knock-out cell line (Fig. 3B). In addition, no residual amine oxidase activity inhibited by BAPN was detected in LOXL2-null cell conditioned medium (Fig. 3C). Fig. 3D shows that 7S dodecamers extracted after collagenase digestion of the ECM derived from LOXL2-null PFHR9 cells are composed of only 7S monomeric subunits, confirming our previous crosslinking studies using pharmacological inhibitors of lysyl oxidase activity (7). Interestingly, the nonreduced 7S dodecamer of the LOXL2-null cells showed slightly faster electrophoretic mobility in comparison to the wild-type cells (Fig. 3D).
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These results show that LOXL2 is the sole source of lysyl oxidase activity for crosslinking the 7S dodecamer of collagen IV in PFHR9 cells. Furthermore, generation of a LOXL2-null PFHR9 cell line provides an excellent system to test the effect of LOXL2 proteolytic processing on collagen IV crosslinking.

To evaluate the 7S crosslinking activity of LOXL2 processing mutants, these constructs were expressed in the LOXL2-null PFHR9 cells. As expected, monomeric 7S subunits are predominant in the catalytically defective LOXL2Y689F mutant, indicating that the enzyme failed to reestablish the crosslinking profile observed in wild-type cells (Fig. 4). In contrast, the 7S dodecamer from wild-type LOXL2 and the enhanced processing mutant LOXL2K317R exhibited the typical crosslinking profile of oligomeric forms observed in wild-type cells. Surprisingly, the full-length, processing defective mutant LOXL2BP failed to crosslink the 7S dodecamer. These results demonstrate that, despite being catalytically active, the full-length processing mutant LOXL2 BP was unable to crosslink the 7S dodecamer. These results suggest that removal of the first two SRCR domains by proteolytic processing is required for effective collagen IV crosslinking in the insoluble matrix.

Proteolytic processing enables the interaction of LOXL2 with insoluble collagen IV

Although full-length and processed forms of LOXL2 have been reported to localize in the ECM of different tissues (7, 15), our results showed that only the processed LOXL2 crosslinks the 7S dodecamer. Thus, we hypothesized that the processed, but not the full-length, form of LOXL2 interacts with collagen IV. To test this hypothesis, we performed a collagenase digestion to solubilize proteins interacting with the insoluble collagen IV scaffold. As we wanted to detect a potential LOXL2–7S dodecamer interaction, we first performed immunoblotting analyses of purified LOXL2 to characterize its electrophoretic mobility under reducing and nonreducing conditions. As shown in Fig. 5A, under nonreducing conditions the full-length and processed LOXL2 forms migrated with an apparent molecular mass of ~75 and ~45 kDa, respectively. The faster electrophoretic mobility of both LOXL2 forms is consistent with the presence of many disulfide bridges that restrain the polypeptide chain yielding a smaller apparent molecular mass in SDS-PAGE. Immunoblotting analyses of collagenase-soluble proteins from wild-type PFHR9 matrix show a 45-kDa band equivalent to that of processed LOXL2 (Fig. 5B, right panel). Surprisingly, the immunoblot also showed an unexpected high molecular mass form of LOXL2 with an electrophoretic mobility slightly slower than the 7S dodecamer shown in the SDS-PAGE (Fig. 5B). None of these two LOXL2 forms were observed in the collagenase digest of LOXL2-null PFHR9 matrix (Fig. 5B, right panel). Thus, the presence of the high molecular mass LOXL2 form suggests a stable interaction between processed
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LOXL2 and the 7S dodecamer that is resistant to the highly dissociative conditions of SDS-PAGE analysis.

To further characterize the interaction between 7S dodecamer and processed LOXL2, we performed immunoblotting analyses of purified 7S dodecamer, urea-soluble, and collagenase-soluble PFHR9 matrix proteins under reducing conditions of SDS-PAGE analysis.

Figure 4. Proteolytic processing of LOXL2 promotes crosslinking of 7S dodecamer. A, immunoblotting detection of 7S subunits in collagenase digested samples of ECM produced by WT LOXL2, LOXL2-null PFHR9 cells (KO), and LOXL2-null PFHR9 overexpressing different LOXL2 mutants described in Fig. 2. Protein samples were fractionated under reducing conditions (50 mM DTT). The nitrocellulose membrane stained with Ponceau S showing the NC1 monomers is presented as a surrogate for total protein loading control. M, D, T, TT, P, and H indicate the electrophoretic mobility of monomer, dimer, trimer, tetramer, pentamer, and hexamer 7S subunits, respectively. B, representation of the percentage of 7S oligomers relative to the total 7S subunits per lane generated in each set of experiments. Error bars represent S.D. from three independent experiments. Data were analyzed by Student’s t test. Statistical significance is shown above the bars. ***, p value ≤ 0.001; ns, not significant.

As shown in Fig. 5C, some bands are similar, but there are striking differences in the number and size of LOXL2 forms. Consistent with a LOXL2–7S dodecamer interaction, the unbound 65-kDa processed LOXL2 form was not detected in purified 7S dodecamer (Fig. 5C, left panel), as it is likely removed during chromatographic purification. However, a ladder of three or more additional bands, larger than the 65-kDa processed LOXL2, was observed in purified 7S dodecamer samples and collagenase digests of PFHR9 and glomerular matrices (Fig. 5C). Of note, the bands detected in the collagenase digest of bovine glomerular matrix showed a more complex pattern, which is probably a consequence of the highly heterogeneous collagenous material isolated from this tissue.

The striking differences between the electrophoretic profiles of LOXL2 forms present in urea-soluble and collagenase-soluble protein samples suggest that the enzyme interacts with collagen IV in different ways (Fig. 5C, right panel). For instance, the 100-kDa full-length LOXL2 band is detected in urea-extracted proteins, but not in collagenase digests, suggesting that it does not interact with collagen. In contrast, the 65-kDa processed LOXL2 (free form, equivalent to the 45-kDa band observed under nonreducing conditions) is present in both preparations, suggesting that this form associates with collagen IV in a noncovalent fashion. Lastly, the ladder of LOXL2 bands is only present in the collagenase digest, suggesting that processed LOXL2 forms a covalent complex with peptide fragments derived from 7S dodecamer that were not digested by collagenase under native conditions. Taken together, these results suggest that processed, but not full-length, LOXL2 is bound both covalently and noncovalently to collagen IV.

To validate that LOXL2 is bound to 7S polypeptides, we subjected collagenase-soluble PFHR9 proteins to pepsin digestion under nondenaturing conditions. As expected, the 7S dodecamer was resistant to pepsin digestion (Fig. 5D, left panel). However, LOXL2 was completely degraded under the same conditions (Fig. 5D, right panel), indicating that the LOXL2 signal detected by the antibody is specific despite the presence of excessively abundant 7S oligomeric subunits. In contrast, because it is possible that processed LOXL2 may have protected the attached 7S peptide fragments from collagenase digestion under native conditions, collagenase-soluble ECM proteins were subjected to disulfide reduction and denaturation followed by a second collagenase digestion. Fig. 5E shows that the 7S dodecamer was almost completely degraded by collagenase after reduction/alkylation and denaturation. At the same time, the ladder above processed LOXL2 was also completely degraded by collagenase followed by an evident increase of the 65-kDa processed LOXL2 (Fig. 5E, right panel). Collectively, these results clearly show that the ladder of LOXL2 forms observed in collagenase digests results from the covalent interaction of processed LOXL2 with 7S collagenous peptides.

To further characterize these findings, we analyzed the presence of LOXL2 in purified 7S from different biological samples. Although the bacterial collagenase preparation used for purification of 7S dodecamers is of the highest commercially available quality, to rule out LOXL2 cleavage by nonspecific proteases, we incubated recombinant LOXL2 with collagenase and monitored the ratio of processed and full-length forms of
LOXL2 by immunoblotting. As shown in Fig. 6A, bacterial collagenase did not alter the relative abundance of the two forms of LOXL2, indicating that collagenase does not process LOXL2. Subsequently, we performed LC-MS/MS analyses of purified 7S dodecamers from PFHR9, bovine glomerular and placental matrices. In all samples, only peptides derived from processed LOXL2 sequences were detected (Fig. 6B). As expected, 7S dodecamer purified from the LOXL2-null PFHR9 cell line did not yield any LOXL2 peptides in the LC-MS/MS analyses. All together, these results revealed that only processed LOXL2 is solubilized from ECM by collagen degradation, and that a fraction of the enzyme is covalently bound to 7S dodecamer.

To gain insight about the type of covalent interaction between processed LOXL2 and collagen IV, we examined the distribution of the different LOXL2 forms in the PFHR9–LOXL2-null rescue system. Processed LOXL2 and ladder bands were present in the collagenase-soluble proteins from PFHR9 cells transfected with wild-type LOXL2 and enhanced processing LOXL2K317R constructs (Fig. 7). As expected, none of the LOXL2 forms were detected in the collagenase digest of cells transfected with the full-length LOXL2BP, further supporting that proteolytic processing is required for LOXL2 localization to the insoluble matrix and interaction with collagen IV. However, although the catalytically defective LOXL2Y689F mutant undergoes proteolytic processing and it interacts noncovalently with collagen IV, the absence of LOXL2 ladder bands indicated that lysyl oxidase catalytic activity is required for the covalent interaction between processed LOXL2 and collagen IV. These findings demonstrate that covalent interaction between LOXL2 and collagen IV is mediated by lysyl oxidase enzymatic activity.

Discussion

Our previous work identified LOXL2 as the main lysyl oxidase expressed in the glomerulus that crosslinks the 7S domain of collagen IV (7). This study provided the first direct evidence
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A

Recombinant LOXL2

Collagenase

+     -

kDa 150

175

150

LOXL2 human
LOXL2 mouse
LOXL2 bovine

B

Signal peptide

LOXL2 human
LOXL2 mouse
LOXL2 bovine

SRC1

LOXL2 human
LOXL2 mouse
LOXL2 bovine

SRC2

LOXL2 human
LOXL2 mouse
LOXL2 bovine

SRC3

LOXL2 human
LOXL2 mouse
LOXL2 bovine

SRC4

LOXL2 human
LOXL2 mouse
LOXL2 bovine

LOX domain

LOXL2 human
LOXL2 mouse
LOXL2 bovine

PHFR9 (mouse)

GBM (bovine)

PBM (bovine)
linking LOXL2 enzymatic activity and collagen IV crosslinking, an important step for basement membrane stabilization (10–12). Here, we extended those studies by investigating the role of LOXL2 proteolytic processing in 7S dodecamer crosslinking. Our results demonstrate that LOXL2 is cleaved by serine proteases at Lys-317, generating a processed form lacking the first two SRCR domains, without an apparent effect over the amine oxidase enzymatic activity. However, proteolytic processing enables the interaction of processed LOXL2 with collagen IV in the insoluble matrix promoting the formation of intermolecular crosslinks in the 7S dodecamer. Fig. 8 summarizes a proposed mechanism based on our findings.

Inhibition experiments with aprotinin, PMSF, and Dec-RVKR-CMK indicated that a serine protease is likely to be responsible for the proteolytic processing of LOXL2. Interestingly, it has recently been shown that a proprotein convertase (PC) activates peroxidasin, the enzyme responsible for the formation of sulfilimine crosslinks in the C-terminal noncollagenous domain (NC1) of collagen IV (36). Although the blockade of LOXL2 proteolytic processing by Dec-RVKR-CMK inhibitor suggested the involvement of a PC, this family of proteases has a strong preference for Arg in P1 in the motif R-X-R/K-R. In contrast, the cleavage site of LOXL2 exhibits a Lys in P1, suggesting that other serine proteases may cleave LOXL2 (33). Interestingly, Dec-RVKR-CMK has been shown to also inhibit members of the trypsin-like (S1) family of membrane-anchored serine proteases. Unlike the PC family, trypsin-like proteases may cleave substrates encompassing either Arg or Lys at P1 (34, 37), suggesting that these proteases are more likely candidates to process LOXL2 at Lys-317. The importance of proteolytic cleavage as a regulatory event controlling LOXL2 crosslinking activity may be appreciated by the evolutionary conservation of Lys in the P1 site of LOXL2 sequences of many different organisms (Fig. 1C). Further studies are required to reveal the identity of the specific protease responsible for LOXL2 processing.

Our results using 1,5-diaminopentane as the substrate for in vitro activity assays demonstrated that both full-length and processed forms of LOXL2 are active. More importantly, proteolytic processing did not increase LOXL2 enzymatic activity toward a small substrate, indicating that LOXL2 crosslinking function is different to LOX and LOXL1, which requires proteolytic cleavage as a regulatory event.
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teolytic cleavage before becoming active. However, our results are in agreement with previous reports showing that LOXL4 exhibits amine oxidase activity in vitro regardless of the presence or absence of SRCR domains (38). The role of processing in collagen IV crosslinking became apparent in rescue experiments using LOXL2-null PFHR9 cells. The results indicated that removal of SRCR1 and SRCR2 domains by proteolytic processing was required before LOXL2 could effectively crosslink collagen IV deposited in the insoluble ECM. Indeed, our immunoblotting analyses of collagenase digests and purified 7S dodecamers from different tissues demonstrated that only the processed form of LOXL2 interacts with collagen IV. Conceivably, the presence of SRCR1 and SRCR2 domains blocks the proper interaction between LOXL2 and collagen IV by steric hindrance. These results suggest that, rather than an activation of enzymatic activity, LOXL2 proteolytic processing is a regulatory event that is required for an effective enzyme-substrate interaction to achieve oxidation of lysine residues and promote crosslinking of the 7S dodecamer.

Strikingly, we have found that a significant fraction of the processed LOXL2 is covalently attached to collagenase-sensitive 7S peptide fragments in a catalytically dependent manner, not only in the PFHR9-ECM but also in animal tissues, indicating that covalent crosslinking of LOXL2 to collagen IV can take place not only in matrix produced by cultured cells, but also in vivo. Although the nature of the collagenous sequences bound to LOXL2 were not identified, the evidence suggests that such fragments may be derived from the 7S dodecamer. These findings are the first report of a stable covalent interaction between a lysyl oxidase and its substrate. Although the biological implications of this finding are unclear at this time, one possibility is that the LOXL2-collagen IV interaction is the result of an arrested enzymatic intermediate in the enzyme's catalytic cycle. The chemical mechanism of lysyl oxidases proposed by Kagan (39) involves the formation of a Schiff base between the lysyl-tyrolyl quinone (LTQ) cofactor and the ε-amino group of Lys in the substrate. The covalent intermediate spontaneously breaks off from the enzyme's catalytic site and molecular oxygen regenerates the LTQ cofactor. Thus, it is possible that a Lys residue from the 7S dodecamer substrate is trapped in the active site of LOXL2 (Fig. 8). Obviously, covalent attachment of LOXL2 through its LTQ cofactor may render the enzyme inactive. It is not known whether this event is reversible, but perhaps this could be a mechanism to quench enzyme activity and avoid excessive lysine oxidation and collagen crosslinking. This new finding deserves further investigation to better understand the molecular mechanisms controlling LOXL2-catalyzed crosslinking of collagen.

LOXL2 proteolytic processing may also play a role as a regulatory mechanism for the crosslinking of other ECM substrates. Although our studies did not address whether proteolytic processing of LOXL2 is required for crosslinking of other ECM proteins that are also known substrates of LOXL2, such as type I and III collagens (40, 41), there is no reason to believe that this proteolytic event is a specific requirement for collagen IV crosslinking. Furthermore, because LOXL2 has been recognized as an important pharmacological target for fibrosis and cancer, it would be interesting to test if the proteolytic processing could regulate crosslinking and deposition of other ECM substrates.

Inhibition of the enzymatic activity of LOXL2 to suppress crosslinking of ECM proteins is a primary focus of anti-fibrotic therapeutic strategies (16–20). So far, a few LOXL2 inhibitors of different types have been developed for the treatment of several fibrotic and metastatic diseases (20, 41, 42). For instance, simtuzumab, a humanized monoclonal antibody described as an allosteric inhibitor of LOXL2, was shown to be an effective inhibitor in mouse models of lung and liver fibrosis (17, 20), but many of the clinical trials using this humanized antibody have failed to show improvement in the development of fibrosis (43). Although simtuzumab inhibitory characteristics were derived from preclinical studies using a mixture of full-length and processed LOXL2 (41), it is not clear whether processing may alter the antibody’s inhibitory allosteric effect. LOXL2 has also been shown to be up-regulated in different nephropathies (44, 45) and, because of its role in collagen crosslinking, it has been suggested as a potential target to prevent increased deposition of collagen IV and other ECM proteins in kidney fibrosis (46). Although there are no published studies addressing this particular hypothesis, it would be interesting to see if any of the currently available LOXL2 inhibitors might indeed ameliorate kidney fibrosis. However, in light of our results, a better understanding of LOXL2 structural determinants mediating collagen crosslinking will better assist the development of more effective treatments targeting LOXL2 activity.

In summary, our study proposes proteolytic processing by serine protease(s) as an important regulatory mechanism of LOXL2-mediated crosslinking of the 7S domain of collagen IV. It is demonstrated that rather than increasing the enzyme catalytic activity, removal of SRCR1 and SRCR2 domains allows the interaction between LOXL2 and collagen IV, resulting in effective oxidation of lysine residues during basement membrane biosynthesis.

Experimental procedures

Cell culture

PFHR9 cells were grown in Dulbecco’s modified Eagle’s medium supplemented with 5% bovine growth serum from HyClone Laboratories (Logan, UT) and 1% penicillin-streptomycin and incubated at 37 °C in 10% CO2. When ECM production was intended, PFHR9 were grown after confluency for 7 days and media were supplemented with 50 μg/ml ascorbic acid. HEK293 cells were cultured in Dulbecco’s modified Eagle’s medium and Ham’s F-12 medium (1:1) with 10% bovine growth serum from HyClone Laboratories and 1% penicillin-streptomycin and incubated at 37 °C in 5% CO2.

LOXL2 purification

LOXL2-His6 was generated and purified as previously reported (7). HEK293 cells stably expressing LOXL2-His6 were expanded, and secreted LOXL2-His6 was purified with a nickel nitritotriacetic acid column to 95% purity as judged by SDS-PAGE. The lysyl oxidase activity of the purified recombinant protein was assayed by Amplex Red assay with 5 mM diamino-pentane as substrate.
**7S dodecamer purification**

The collagen IV 7S dodecamer from ECM produced by PFHR9 was purified as previously published (7). Briefly, cells were homogenized in 1% (w/v) deoxycholate with sonication, and the insoluble material was isolated after centrifugation at 20,000 × g for 15 min. The pellet was then extracted with 1 M NaCl plus 50 mM Tris-Cl, pH 7.5, and 10 mM Tris-Cl, pH 7.5, and then digested in 50 mM Tris-Cl, pH 7.5, 5 mM CaCl₂, 5 mM benzamidine, 25 mM 6-aminocaproic acid, 0.4 mM phenylmethylsulfonyl fluoride, and 0.1 mg/ml bacterial collagenase (Worthington Biochemical Corp.). Collagenase-solubilized material was dialyzed against 50 mM Tris-Cl, pH 7.5 and then subjected to DEAE-cellulose and gel filtration chromatography.

**SDS-PAGE and immunoblotting**

For immunoblotting assays, protein samples were loaded for 4–15% (w/v) SDS-PAGE and electrophoresed onto nitrocellulose membranes. Blots were incubated with polyclonal anti-7S collagen IV (Assay Biotech) or LOXL2 (Abcam) antibodies and visualized with horseradish peroxidase–conjugated donkey anti-rabbit as secondary antibody (Jackson Immunoresearch Laboratories). Detection of HRP activity from secondary antibodies was revealed using Pierce enhanced chemiluminescence reagents (Thermo Scientific) following the instructions of the manufacturer. Quantification of immunoblots was performed using ImageJ software (47).

**CRISPR-mediated LOXL2 knockout**

The gRNA sequence for LOXL2 exon 2 (forward 5′-CACC-GCCGCGGTCTACTACGA-3′; reverse 5′-AAACTC-GTATGAGACTCCAGGC-3′) was ligated into the BbsI sites of pSpCas9(2B)-A-Puro (PX459) (gift from Dr. Feng Zhang, Addgene plasmid no. 48139) (48). PFHR9 cells were cotransfected with Lipofectamine 2000 (Invitrogen) with PX459 plasmid and a hygromycin resistance homologous recombination vector, with 500 base pairs’ homology arms surrounding the targeted sequence of the exon 2. Transfection in absence of the homology recombination vector was performed to address the effectiveness of the designed sgRNA by Surveyor assay (Integrated DNA Technologies (IDT)). Transfected PFHR9 cells were initially treated with 4 µg/ml puromycin to select cells containing the PX459 plasmid expressing the Cas9 protease and the specific gRNA, after a first round of selection cells was selected with 250 µg/ml hygromycin B for the homologous recombination vector. Resistant clones were isolated and screened for the presence of the insertion cassette via homologous recombination with the following primers: forward/external 5′-CAGTCTGAGTCTGCAAAGG-3′ and reverse/internal 5′-GCTTGTGCTGTTGGAACGTAGGG-3′. Positive clones for the insertion were subjected to analysis by Western blot using two different anti-LOXL2 antibodies: rabbit polyclonal to LOXL2 (ab96233 and ab113960, Abcam).

**Cloning, plasmid constructs, and transient transfection**

TrueORF Gold cDNA for human LOXL2 (C-terminal Myc-DDK–tagged) construct cloned into the pCMV6-Entry vector (LOXL2) was purchased from Origene (Rockville, MD). The catalytically inactive mutant (LOXL2K317R) was described in a previous publication (7). The following primers were designed to introduce the point mutations K317R, and K317E/R316G/R257G into the construct pCMV6-LOXL2-DDK-Myc: LOXL2-K317R forward 5′-GGACCTCTAAATCCGGGAGGCGGTAACAGGACGACAC-3′; LOXL2-K317E reverse 5′-GTTGCTCTGGTTGTAAGCTGCTTCCGGAGATTTGGAATGGCCAGTG-3′; LOXL2-K317R forward 5′-GGACCTCTAAATCCGGGAGGCGGTAACAGGACGACAC-3′; LOXL2-K317E reverse 5′-GTTGCTCTGGTTGTAAGCTGCTTCCGGAGATTTGGAATGGCCAGTG-3′; LOXL2-K317R forward 5′-GGACCTCTAAATCCGGGAGGCGGTAACAGGACGACAC-3′; LOXL2-K317E reverse 5′-GTTGCTCTGGTTGTAAGCTGCTTCCGGAGATTTGGAATGGCCAGTG-3′. Clones containing LOXL2-K317R (LOXL2K317R) and LOXL2-K317E/R316G/R257G (LOXL2KP) were verified by DNA sequencing. LOXL2, LOXL2K317R, LOXL2K317E, and LOXL2KP constructs were transiently transfected in HEK293 cells using Lipofectamine 2000 reagent (Invitrogen).

**Amine oxidase activity assay**

LOXL2 amine oxidase enzymatic activity was measured in concentrated serum-free conditioned cell culture medium or 4 mM urea-soluble PFHR9 matrix proteins using 1,5-diaminopen tane as substrate and monitoring hydrogen peroxide production by an Amplex Red fluorescence assay (Invitrogen) (35). The enzymatic reaction was performed in 0.05 M sodium borate, pH 8.2, buffer containing 1 unit/ml of HRP, 10 mM Amplex Red, and 5 mM 1,5-diaminopentane in the presence or absence of 1 mM BAPN. Samples from 4 mM urea-extracted ECM were diluted in a final 1 mM urea concentration before measurement of amine oxidase activity. After 120-min incubation at 37 °C, fluorescence was measured on a SpectraMax M5 plate reader using excitation and emission wavelengths of 540 and 590 nm, respectively.

**7S crosslinking by LOXL2 processing mutants**

LOXL2-null PFHR9 cells were transfected with the previously described constructs LOXL2, LOXL2K689N, and LOXL2KP. Forty-eight h after transfection the cells were selected with 200 µg/ml G418 (Geneticin). Resistant clones were expanded and grown for 7 days after confluency in the presence of 50 µg/ml ascorbic acid. The cells and underlying ECM were scraped in deoxycholate buffer (1% sodium deoxycholate, 25 mM Tris-HCl pH 7.5, 1 mM EDTA-Na, 0.4 mM phenylmethylsulfonyl fluoride, 10 µg/ml leupeptin, 10 µg/ml aprotinin and 1 µg/ml pepstatin) and sonicated until homogeneous and centrifuged for 30 min, 14,000 × g. The insoluble pellet was subsequently washed with 1 M NaCl in 25 mM Tris-HCl pH 7.5. For some experiments urea-soluble proteins were extracted by incubating the insoluble pellet with 4 mM urea in 25 mM Tris-HCl pH 7.5 for 1 h at room temperature. The remaining pellet was digested overnight with 50 µg/ml collagenase (Worthington Biochemical Corp.) in collagenase buffer (50 mM HEPES, pH 7.5, 5 mM CaCl₂, 5 mM benzamidine, 25 mM 6-aminocaproic acid, 150 mM NaCl, 25 mM 1,5-diaminopentane, and 25 mM benzamidine) and the lysate dialyzed against 50 mM Tris-Cl, pH 7.5 and then subjected to DEAE-cellulose and gel filtration chromatography.
acid, 0.4 mM phenylmethylsulfonyl fluoride). The supernatant fraction was redigested overnight with 10 μg/ml collagenase. Collagenase-soluble proteins were analyzed by immunoblot using anti-7S polyclonal antibody. The ratio between cross-linked 7S subunits and uncrosslinked 7S monomers was calculated as percentage of the total protein per lane as quantified with Image J (47) gel analysis plugin.

**Enzymatic digestion of 7S dodecamers**

Ten micrograms of purified 7S dodecamer from PFHR9 cells, bovine glomerulus or bovine placenta were each mixed with an equal volume of 0.1 M Tris-HCl, pH 7.5, buffer containing 8 M guanidine-HCl, and 50 mM dithiothreitol to denature proteins and reduce disulfide bonds. Samples were heated in a boiling water bath for 10 min. After reaching room temperature, proteins were alkylated with 100 mM iodoacetamide in the dark for 45 min. Following ethanol precipitation at −20 °C overnight, the pellets were resuspended in 0.1 M ammonium bicarbonate pH 7.8 and digested with LysC (1:20 enzyme to protein ratio) for 4 h at 37 °C. The samples were heated at 95 °C for 7 min to inactivate LysC. Samples were cooled down to room temperature before adding trypsin at an enzyme to substrate ratio of 1:20 (w/w) for overnight at 37 °C. After trypsin inactivation, peptide samples were deglycosylated with PNGase F at 37 °C overnight. Peptide samples were cleaned using OMIX C18 tips (Agilent) before mass spectrometry analyses.

**LC-MS/MS analyses and database searching**

7S peptides were analyzed on a Q Exactive mass spectrometer equipped with an Easy nLC-1000 system (Thermo Scientific) as described earlier (49). Approximately a total of 1 μg of peptide was reconstituted in water containing 0.1% formic acid (solvent A) and separated on a PicoFrit (New Objective, Woburn, MA) column (75 μm inner diameter × 110 mm, 10 μm inner diameter tip) packed with Reprosil-Pur C18-AQ resin (3 μm particle size and 120 Å pore size) at a flow rate of 300 nl/min. Peptide mixture was fractionated by a 70-min gradient consisting of the following steps: 0–5 min, increase to 2% B (acetonitrile containing 0.1% formic acid); 5–55 min, 5–35% B; 55–60 min, 90% B, and held at 90% B for 10 min before returning to the initial conditions of 2% B. Data were acquired using a data-dependent method with a scanning window of 300–1800 m/z and dynamic exclusion enabled. Full scans were acquired at a resolution of 70,000, an automatic gain control (AGC) target of 3 × 106, and a 64 ms maximum injection time. The top 20 most abundant ions were selected for fragmentation with higher energy C-trap dissociation (HCD) at a resolution of 17,500, an automatic gain control target of 2 × 105, 100 ms maximum injection time, 2 m/z isolation width, and 27% normalized collision energy. The data were collected with unassigned and +1 charge excluded in the charge state settings. Dynamic exclusion was set to 20 s.

Thermo (.raw) files were independently searched by MyriMatch algorithm run on Windows server maintained by the Vanderbilt University Center for Structural Biology (50). Files were searched against full proteome-level mouse and bovine FASTA databases (NCBI), which also included the reverse sequences for each protein to compute the false discovery rate with a maximum of two missed cleavages and one missed terminus cleavage (semitryptic peptides), with a minimum peptide length of 5 residues and a maximum length of 50 residues. The search was configured to identify peptides with the following amino acid modifications: variable oxidation of methionine (+15.9949) as well as fixed carbamidomethylation of cysteine (+57.021). Precursor and fragment mass tolerance were set to 10 and 20 ppm, respectively. The pepXML files obtained from the search were imported into IDEmpicker (version 3.1) for the parsimonious grouping of identified proteins with a maximum Q value of 2% for peptide spectrum match (51, 52).

**Author contributions**—R. M. V. and A. J. L.-J. conceived and coordinated the study. A. J. L.-J., T. B., and R. M. V. designed experiments and acquired and analyzed data. R. M. V. and A. J. L.-J. interpreted the data. A. J. L.-J. and R. M. V. wrote the paper. All authors reviewed the results and approved the final version of the manuscript.

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