The Laminin–Nidogen Complex is a Ligand for a Specific Splice Isoform of the Transmembrane Protein Tyrosine Phosphatase LAR

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Abstract. Leukocyte antigen–related protein (LAR) is a prototype for a family of transmembrane protein tyrosine phosphatases whose extracellular domain is composed of three Ig and several fibronectin type III (FnIII) domains. Complex alternative splicing of the LAR-FnIII domains 4–8 has been observed. The extracellular matrix laminin–nidogen complex was identified as a ligand for the LAR-FnIII domain 5 (Fn5) using a series of GST-LAR-FnIII domain fusion proteins and testing them in in vitro ligand-binding assays. LAR–laminin–nidogen binding was regulated by alternative splicing of a small exon within the LAR-Fn5 so that inclusion of this exon sequence resulted in disruption of the laminin–nidogen-binding activity. Long cellular processes were observed when HeLa cells were plated on laminin–nidogen, but not when plated on a fibronectin surface. Indirect immunofluorescent antibody staining revealed high expression of LAR in a punctate pattern, throughout the length of these cellular processes observed on laminin–nidogen. Antibody-induced crosslinking of LAR inhibited formation of these cellular processes, and inhibition was correlated with changes in cellular actin cytoskeletal structure. Thus, LAR–laminin–nidogen binding may play a role in regulating cell signaling induced by laminin–nidogen, resulting in cell morphological changes.

R egulation of protein tyrosine phosphorylation is a vital component of extracellular matrix (ECM)–induced signal transduction. Changes in the tyrosine phosphorylation status of specific proteins are implicated in the cytoskeletal reorganization required for dynamic regulation of focal adhesion sites in signal transduction at these sites, and in the transmission of guidance signals in motile cells (2). Protein tyrosine phosphorylation is regulated by both protein tyrosine kinases and protein tyrosine phosphatases (PTPases; 12). Nevertheless, the role of PTPases in ECM-induced signal transduction is not well understood. A number of studies have implicated PTPase activity in cell adhesion, cell spreading, neurite extension, disassembly of focal adhesion sites, and in signal transduction at the tips of neuronal growth cones (2, 9, 11). However, with the exception of the transmembrane PTPase CD45, which may regulate integrin-induced tyrosine phosphorylation in neutrophils (1), the identity and role of the transmembrane PTPase(s) involved have not been elucidated.

Leukocyte antigen–related protein (LAR) is a prototype for a family of transmembrane PTPases whose extracellular regions are composed of a combination of three Ig-like domains and several fibronectin-type III (FnIII) domains (see Fig. 1; 35, 37). Other members of this family include mammalian PTP8 and PTPα, Drosophila DLAR, and chicken CRYP-α (24, 33, 43). LAR family PTPases are prime candidates for regulating ECM-induced signal transduction for the following reasons: the extracellular region of these PTPases has a close similarity with the neural cell adhesion molecule (N-CAM) family of cell-adhesion molecules (35); furthermore, LAR is located at the disassembly side of focal adhesion sites, and in cadherin-mediated cell–cell adhesion sites (18, 31). Chicken CRYP-α is localized to the tips of neuronal growth cones, and Drosophila homolog of human LAR (DLAR) plays a role in neuronal pathfinding (17, 33). Phenotypes of LAR knockout mice suggest that LAR may also play a role in morphogenesis of the mammary gland in mice (30).

The LAR cytoplasmic domain is composed of two PTPase domains, the second of which appears to be catalytically inactive (36). Physiological substrates for the LAR...
PTPase are unknown. The LAR cytoplasmic domain binds to a coiled-coil protein LIP1, which might link LAR to cytoskeletal structures (31). The LAR C-terminus also associates with a multidomain protein called Trio, which contains a protein kinase domain and two guanine nucleotide exchange factor domains that are, respectively, specific to Rac1 and RhoA (5). These observations indicate a potential role for LAR in cytoskeletal reorganization induced by cell–ECM contact that is mediated by Rac and Rho-like molecules (39).

A number of observations suggest that the extracellular domain of LAR also has an important function. The LAR extracellular domain is proteolytically cleaved and undergoes controlled shedding (31, 34). Furthermore, the FnIII domains are alternatively spliced in a tissue-specific and developmentally regulated manner (25, 42). The entire region containing the FnIII domains 4–7 is alternatively spliced out in PTPs and PTP8 (24, 43). Although this form of alternative splicing has not been seen in LAR, we have previously demonstrated alternative splicing of the LAR FnIII domains 6 and 7, and of LAR FnIII domain 4 (25). We also reported the alternative splicing of a small exon (exon 13) within the LAR FnIII domain 5. Inclusion of exon 13 within FnIII domain 5 may have a particular function in neuronal tissue, since cells of neuronal origin express the highest amount of exon 13 containing mRNA. In all other cell types tested, the LAR isoform in which exon 13 is spliced out is the predominant form (25). Furthermore, alternative splicing of exon 13 is regulated in neuronal tissues during embryonic development (43). Perhaps more importantly, the alternatively spliced exon 13 is conserved among LAR-like PTPases (human LAR, rat LAR and PTP6; 25, 28). These observations suggest that alternative expression of FnIII domains 4–7 may have a conserved function.

To study the role of the LAR FnIII domains 4–7, we constructed a series of GST-LAR FnIII domain fusion proteins and tested them in in vitro ligand-binding assays. Using this approach, the ECM laminin-nidogen complex was identified as a ligand for the LAR FnIII domain 5. The laminin–nidogen complex is a major component of the ECM that modulates cell adhesion, cell migration, neurite outgrowth, cell proliferation, and cell differentiation (21). The ability of laminin–nidogen to modulate such a wide range of events is related to its large size and multi-domain structure that allows the interaction of laminin–nidogen with a number of different ligands. We found that alternative splicing of exon 13 within the LAR FnIII domain 5 regulated LAR–laminin–nidogen–binding activity. Furthermore, antibody-induced cross-linking of LAR in HeLa cells alters laminin-nidogen–induced cell morphology.

Materials and Methods

Cells

U373 MG (human glioblastoma) and HeLa (human epithelioid carcinoma) cells were obtained from the American Type Culture Collection (Rockville, MD). U373 MG were grown in Eagles media (GIBCO BRL, Gaithersburg, MD) containing 10% FCS. HeLa cells were grown in DMEM media (Bio-Whittaker, Walkersville, MD) containing 10% FCS and supplemented with 2 mM glutamine. Cells were maintained at 37°C in a humidified atmosphere with 10% CO₂.

Buffers

PBS: 10 mM sodium phosphate buffer (pH 7.4), 150 mM NaCl. HBS buffer: 10 mM Hepes (pH 7.4), 150 mM NaCl. SDS sample buffer: 0.125 M Tris (pH 6.8), 10% SDS, 25% glycerol, 0.01% bromophenol blue. Detachment buffer: 2 mM EDTA, PBS, 0.05% BSA. Wash media: serum-free DMEM media supplemented with 0.05% heat-inactivated BSA and 10 mM Hepes (pH 7.4). BSA was heat-inactivated at 80°C for 20 min.

Anti-LAR Monoclonal Antibodies

The anti-LAR monoclonal antibodies 75.3A, 11.1A, and 71.2E have been previously described (34). The anti-LAR monoclonal antibody 75.11.16 was raised by a similar approach using a GST-LAR Fn5-protein as antigen. The epitopes in the LAR extracellular domain recognized by these antibodies are as follows: mAb 75.3A, LAR Ig domain 1, 2, or 3; mAb 11.1A, between FnIII domain 5 and the transmembrane segment; mAb 71.2E, FnIII domain 6 or 7; mAb 75.11.16, LAR FnIII domain 5.

Other Antibodies

The antilaminin mAbs used were: antihuman laminin β1 chain (clone I) and antihuman laminin γ1 chain (clone II) from GIBCO BRL; and antihuman laminin (LAM-89) from Sigma Chemical Co. (St. Louis, MO). Affinity-purified rabbit polyclonal anti-Engelbreth-Holm-Swarm (EHS)-laminin was also from Sigma Chemical Co., and rabbit polyclonal anti-mouse laminin B1/B2 chain was from Upstate Biotechnology Inc. (Lake Placid, NY). Anti-human β2 microglobulin mAb MIG-B5 was from BioSource International (Camarillo, CA). Affini-pure rabbit anti-mouse IgG was from Jackson ImmunoResearch Laboratories, Inc. (West Grove, PA). Monoclonal anti-human integrin β1, anti-human talin, and anti-human vinculin antibodies were from Upstate Biotechnology Inc. HRP-conjugated rabbit anti-mouse IgG antibody and TRITC-conjugated goat anti-mouse IgG, were from Southern Biotechnology Associates Inc. (Birmingham, AL).

Laminin and ECM Proteins

The following preparations of ECM proteins were used: Matrigel™ (basement membrane matrix from the mouse EHS sarcoma; Becton Dickinson, Bedford, MA); purified laminin–nidogen complex from the EHS sarcoma (Sigma Chemical Co., St. Louis, MO); purified laminin was also from Sigma Chemical Co., and rabbit polyclonal anti-mouse laminin B1/B2 chain (clone II) from GIBCO BRL; and anti-human laminin (clone I) from Becton Dickinson. The antilaminin mAbs used were: anti-human laminin β1 chain (clone I) and anti-human laminin γ1 chain (clone II) from GIBCO BRL; and antihuman laminin (LAM-89) from Sigma Chemical Co. (St. Louis, MO). Affinity-purified rabbit polyclonal anti-Engelbreth-Holm-Swarm (EHS)-laminin was also from Sigma Chemical Co., and rabbit polyclonal anti-mouse laminin B1/B2 chain was from Upstate Biotechnology Inc. (Lake Placid, NY). Anti-human β2 microglobulin mAb MIG-B5 was from BioSource International (Camarillo, CA). Affini-pure rabbit anti-mouse IgG was from Jackson ImmunoResearch Laboratories, Inc. (West Grove, PA). Monoclonal anti-human integrin β1, anti-human talin, and anti-human vinculin antibodies were from Upstate Biotechnology Inc. HRP-conjugated rabbit anti-mouse IgG antibody and TRITC-conjugated goat anti-mouse IgG, were from Southern Biotechnology Associates Inc. (Birmingham, AL).

GST–LAR–FnIII Fusion Proteins

Fusion proteins of glutathione S-transferase (GST) with various LAR FnIII domains were constructed using the E. coli expression vector pGEX-2T (Pharmacia Biotech., Inc., Piscataway, NJ, 32). The GST-fusion proteins used in this study are shown schematically in Fig. 1. Various segments of LAR cDNA with appropriate restriction sites for subcloning were generated by PCR and inserted in-frame into the pGEX-2T vector downstream of the GST coding sequence. The resultant fusion proteins used in this study are shown schematically in Fig. 1. Various segments of LAR cDNA with appropriate restriction sites for subcloning were generated by PCR and inserted in-frame into the pGEX-2T vector downstream of the GST coding sequence. Their nucleotide sequences were confirmed by DNA sequencing. Fusion protein synthesis was induced in exponentially growing Esherichia coli cultures by adding 0.1 mM isopropyl-thio-galactoside, and continued for 2 h at 25°C. Fusion proteins were extracted by solubilizing the bacterial pellet with 1% Triton X-100/PBS, containing 1.0 mM PMSF and 1.0 mM benzamidine for 30 min at room temperature, followed by sonication for 30 s at 4°C. Fusion proteins were affinity-purified using glutathione Sepharose beads. Beads were washed five times with PBS. The purity and sizes of the fusion proteins were confirmed by SDS-PAGE followed by Coomassie blue staining.

35S-Labeling of Conditioned Media

U373 MG cells were grown to 80% confluence in 100-mm tissue culture dishes in Eagle’s media containing 10% FCS. Cells were then washed twice with serum-free media and metabolically labeled with [35S]methionine/cysteine (Dupont-NEF, Boston, MA). In brief, cells were incubated for 20 h in methionine-free media supplemented with [35S]methionine/cysteine (80 μCi/ml), 5% dialyzed FCS (1,000 D cut-off), and 10 mM Hepes buffer (pH 7.4). Labeled media was collected at 4°C, the protease inhibitors (1.0 mM PMSF and 1.0 mM benzamidine) were added, and the media was centrifuged at 1,500 rpm in a Beckman J-6M/E centrifuge at 4°C.
Ligand-binding Assays

The different GST-LAR-fusion proteins (20 μg) bound to glutathione Sepharose beads were incubated with [35S]methionine-labeled media (5 ml) collected from U373 MG cells. Incubation was carried out for 2 h at 4°C with rotation. The incubation was terminated by centrifuging the beads at 1,000 rpm in a Beckman J-6M/E centrifuge for 5 min at 4°C. The beads were washed four times with 5 ml of ice-cold HBS buffer. After the fourth wash, the beads were resuspended in SDS sample buffer, reduced with β-mercaptoethanol, boiled, and subjected to SDS-PAGE on a 5% gel. A sample of the media without incubation with the fusion proteins was run on the gel to assess the number and relative concentration of the labeled bands in the media. Gels were processed for autoradiography with Enhance™ (Dupont-NEC) according to the manufacturer's instructions. Dried gels were exposed to x-ray film in the presence of an enhancer screen.

Binding of GST-LAR Fn5 (with or without the exon 13 sequence) to Matrigel™ was assayed as follows. Glutathione bead-bound GST-LAR Fn5 or Fn5-2 (20-μl beads coated with 40 μg protein) was diluted into 10 ml HBS containing 2 mM EDTA, 1.0 mM PMSF, and 1.0 mM benzamidine, and mixed with various amounts of Matrigel (150-750 μg) for 2 h at 4°C with rotation. The beads were then precipitated by centrifugation and washed four times with 5 ml of ice-cold HBS buffer each. After the final centrifugation, the beads were resuspended in SDS sample buffer, reduced with β-mercaptoethanol, boiled, and subjected to SDS-PAGE on a 6.5% gel and immunoblotting. Blots were probed with the rabbit polyclonal anti-mouse laminin B1/B2 chain antibody, followed by HRP-conjugated goat anti-rabbit antibody. Bound antibodies were visualized by the enhanced chemiluminescence reagent (Amersham Life Science Inc., Arlington Heights, IL).

Binding of GST-LAR Fn5 proteins to the purified laminin-nidogen complex was assayed in a similar manner, except that the concentration of laminin-nidogen was kept constant (150 μg in 10 ml), while the amount of GST-LAR Fn5 protein bound to glutathione beads varied (50-150 μg of protein per 40 μl beads), and the mixtures were incubated for 3.5 h at 4°C with rotation. Samples were analyzed on a 5% SDS gel, and blots were probed with an affinity-purified monoclonal anti-laminin antibody.

Immunoblotting with Anti-laminin Antibodies

Conditioned media collected from U373 MG cells was incubated with the GST-LAR fusion proteins as described in the binding assay above. Electrophoresis was carried out on a 3–12% gradient SDS-polyacrylamide gel under nonreducing conditions, or on a 5% SDS-polyacrylamide gel under reducing conditions. After electrophoresis, the gels were blotted to nitrocellulose and probed with monoclonal antibodies specific to the laminin β1 chain, laminin γ1 chain, or laminin (LAM-89). Bound anti-laminin antibodies were detected with HRP-conjugated rabbit anti–mouse IgG antibodies and the enhanced chemiluminescence reagent (Amersham Life Science Inc., Arlington Heights, IL). Akt was visualized by incubation of the cells with 2U rhodamine-conjugated phalloidin (Molecular Probes Inc., Eugene, OR) in blocking solution for 30 min according to the manufacturer's instructions. For all stainings, the specificity of staining was determined in control experiments in which either the primary antibody or both the primary and secondary antibodies were omitted.

After labeling, all slides were washed three times with PBS and mounted in Fluorsave (Calbiochem-Novabiochem Corp., La Jolla, CA). Slides were viewed on an Axiophot™ microscope (Carl Zeiss Inc.) equipped for epifluorescence. Photographs were taken on Tri-X400 black and white film (Eastman Kodak Co.).

Results

Ligand Binding of GST–LAR–FnIII Domain Fusion Proteins

The complex alternative splicing of FnIII domains 4–7 among LAR family members suggests that these domains may have an important function. One possibility is that this region of LAR encodes a ligand-binding site. To test this possibility, a series of GST fusion proteins containing various GST-FnIII domains were constructed for testing in ligand-binding assays. The GST–LAR–FnIII constructs used in this study are shown schematically in Fig. 1 b. Most of the constructs that include the LAR-FnIII domain 5 were made in two different versions: with or without the alternatively spliced exon 13, which encodes nine amino acids. These versions are referred to as Fn5+ and Fn5− forms, respectively, throughout the text.

Initially we tested binding of the GST–LAR–FnIII fusion proteins to components of the conditioned media from cultured cells. Conditioned media is a potential source of ligands such as ECM proteins. Incubation of 20 μg GST–LAR–FnIII fusion proteins with [35S]methionine-labeled conditioned media from U373 MG (human glioblastoma) cells revealed specific binding of ~350-, 220-, and ~200-kD proteins to the GST-fusion constructs #3, #5, #7, and #8 (encoding Fn5−, Fn5−, 6, 7, 8; Fn3,4,5−, and Fn5−, 8, respectively; Fig. 2). The bound proteins are highly enriched by the LAR constructs when compared with their concentration in the original media. Specificity of binding is indicated by the observation that the ligand-binding constructs all contained the Fn5− domain. All of
The Fn5\(^{-}\)-containing isoforms showed only background binding similar to the GST protein alone. The Fn5\(^{-}\) isoform bound the same three labeled bands whether the Fn5\(^{-}\) domain was present as a single domain (construct #3), or was flanked by other FnIII domains (constructs #5, #7, and #8). The binding was also independent of the distance from the Fn5\(^{-}\) domain to the GST protein, indicating that the presence of the GST domain did not influence binding. Construct 8, in which Fn5\(^{-}\) is joined directly to Fn8, shows a weaker binding than the other Fn5\(^{-}\) constructs, suggesting that the presence of Fn6 and Fn7 may influence the ligand-binding activity of Fn5\(^{-}\). Fn4 or Fn6 alone do not show specific ligand binding (constructs #9 and #10). Thus, the Fn5\(^{-}\) domain has a specific ligand-binding capacity, while inserting nine amino acids in this domain completely abolishes ligand binding.

**The LAR-binding Protein Contains Laminin-1**

The molecular masses of the Fn5\(^{-}\)-binding proteins are similar to those of the ECM protein laminin-1. Laminin-1 is a disulfide-bonded heterotrimer of \(~350-, 220-, and \(~200-kD subunits (\(\alpha\), \(\beta\), and \(\gamma\) chains, respectively), and is reported to be synthesized and secreted into the media by U373 MG cells (19). Indeed, laminin chains immunoprecipitated from \(^{35}S\)labeled media of U373 MG cells migrated at the same positions as the Fn5\(^{-}\) binding proteins (data not shown). We therefore tested if the Fn5\(^{-}\)-binding protein is laminin by determining its reactivity to antilaminin antibodies.

First, the Fn5\(^{+}\) and Fn5\(^{-}\) isoforms (constructs #2 and #3) were incubated with unlabeled conditioned media from U373 MG cells, and the bound proteins were separated by SDS-PAGE under nonreducing conditions followed by immunoblotting. Under nonreducing conditions, the laminin-1 trimer migrates as a protein of \(~850\) kD. When probed with an antilaminin antibody, a high–molecular weight protein was detected bound to the Fn5\(^{+}\), but not to the Fn5\(^{-}\) isoform. This protein migrated at the same molecular weight as the purified laminin-1, which was run in parallel (Fig. 3 a). Next, the reactivity of antibodies specific to the laminin-\(\beta\) and -\(\gamma\) chains with the Fn5\(^{-}\)-bound ligands was tested in a similar assay, except that electrophoresis was carried out under reducing conditions. The anti-\(\beta\) and anti-\(\gamma\) laminin antibodies detected bands of \(~220\) kD and \(~200\) kD, respectively, in the proteins bound to the Fn5\(^{-}\), but not to the Fn5\(^{+}\), isoform. These bands migrated at the same position as the laminin-\(\beta\) and -\(\gamma\) chains in purified laminin run in parallel (Fig. 3, b and c).

Thus, laminin is specifically bound to the Fn5\(^{-}\) fusion...
However, it cannot be concluded from these experiments alone whether laminin is bound directly to LAR, or indirectly via a molecule present in the conditioned media that acts as a bridge between laminin and LAR. Laminin-1 is generally found in a very tight 1:1 complex with the extracellular matrix protein nidogen (also known as entactin; 6, 40). The laminin–nidogen complex has binding sites for a number of different ligands, including collagen, perlecán, fibulin-1, heparin sulfate, and other ECM components (3, 27, 40). To examine if the Fn5\(^2\) domain binds the laminin–nidogen complex directly or not, the Fn5\(^2\) and Fn5\(^1\) isoforms were incubated with either Matrigel™ or the purified mouse laminin–nidogen complex. Matrigel™ is a preparation of ECM from the mouse EHS sarcoma, which is highly enriched for the laminin–nidogen complex (15). Incubation of the Fn5\(^2\) and Fn5\(^1\) isoforms with increasing concentrations of Matrigel™, followed by immunoblotting with a polyclonal anti-laminin antibody, indicated a corresponding increase in binding of laminin to the Fn5\(^2\), but not the Fn5\(^1\) isoform (Fig. 4a). In another experiment, the purified laminin–nidogen complex (constant amount) was incubated with increasing amounts of the Fn5\(^2\) and Fn5\(^1\) proteins. The purified laminin–nidogen complex can also bind Fn5\(^2\) in a dose-dependent manner, suggesting that LAR binding to the laminin–nidogen complex does not require any other ECM component (Fig. 4b). At high concentrations of the Fn5\(^2\) protein, some laminin–nidogen binding is detected, suggesting that the Fn5\(^2\) may retain low-affinity binding to the laminin–nidogen complex. These data therefore indicate that the laminin–nidogen complex is a ligand for the LAR–Fn5\(^2\) domain, although we cannot specify the exact binding site(s) on the laminin–nidogen complex.

## Role of LAR–Laminin–Nidogen Binding in Cell Morphology

The above finding suggests a role for LAR in laminin–nidogen–mediated cellular events. Laminin–nidogen bind-
LAR plays a role in the cell morphology induced by laminin–nidogen, long, thin cellular processes are expressed in HeLa cells where actin is present mainly as numerous thin stress fibers (Fig. 7, c). As a control, we examined localization of the integrin β1 subunit after LAR cross-linking. The integrin β1 subunit is a component of a number of integrin cell surface laminin receptors (α1β1, α2β1, α3β1, α6β1, and α7β1). There were no significant changes in the distribution of the integrin β1 subunit (data not shown), indicating that LAR cross-linking probably does not alter cell morphology through cointernalization of integrin laminin receptors.

Formation of cellular processes on laminin requires dynamic cytoskeletal reorganization. Actin staining of cells 1 h after plating on a laminin–nidogen substrate indicates a relatively disordered, meshwork actin cytoskeleton typical of spreading cells (Fig. 7 a). As the cell extends long processes, the actin skeleton becomes more organized, and a long, thick actin fiber is seen to extend from the cell body along the length of these processes (Fig. 7 b). This thick central actin fiber is absent from the LAR cross-linked cells where actin is present mainly as numerous thin stress fibers (Fig. 7, c and d). Thus, correlating with the effect on cell morphology, LAR cross-linking also affected the cellular cytoskeletal structure.

Migration of cellular processes along an ECM requires that cell morphology be modulated to allow movement along the substrate. In the spreading cells, LAR cross-linking was found to cause a dramatic change in the morphology of the spreading cells. In LAR cross-linked cells, formation of the laminin–nidogen-induced cellular processes is inhibited by 97%, although the cells do remain attached to the substrate (Table I; compare Fig. 5 a with Fig. 5, c and d). Similar inhibition of the cellular processes was observed using four different anti-LAR mAbs with distinct epitopes in the LAR extracellular domain (data not shown). Neither treatment of the cells with secondary antibody alone or a control cross-linking using anti-β2 microglobulin antibody affected the laminin–nidogen-induced cell morphology (Fig. 5, e and f, respectively). HeLa cells express surface β2 microglobulin (26), which has been shown to be sequestered after antibody-induced cross-linking (8).

Antibody-induced LAR cross-linking has been shown to induce patching and ultimately internalization of cell surface LAR (31). The extent of LAR internalization induced by cross-linking over the time period of our study was determined by immunofluorescence. Before LAR cross-linking, (after plating on laminin–nidogen for 1 h), a high concentration of cell surface LAR was visualized (Fig. 6 a). After antibody-induced LAR cross-linking, however, cell surface LAR was barely detectable (Fig. 6 b). Permeabilization of the cross-linked cells followed by LAR staining revealed high concentrations of aggregated LAR within the cell, indicating internalization of LAR had occurred after cross-linking (Fig. 6 c).

As a control, we examined localization of the integrin β1 subunit after LAR cross-linking. The integrin β1 subunit is a component of a number of integrin cell surface laminin receptors (α1β1, α2β1, α3β1, α6β1, and α7β1). There were no significant changes in the distribution of the integrin β1 subunit (data not shown), indicating that LAR cross-linking probably does not alter cell morphology through cointernalization of integrin laminin receptors.

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Migration of cellular processes along an ECM requires...
continuous assembly and disassembly of focal adhesions in order to provide traction during migration (13). Localization of LAR to the disassembly side of focal adhesions (31) suggests a mechanism by which LAR might influence the laminin-induced morphological changes in HeLa cells. To determine if LAR is localized to the cellular extensions, and thus might be involved in focal adhesion turnover, we visualized cellular localization of LAR as well as that of the focal adhesion–associated cytoskeletal proteins talin and vinculin, using immunofluorescence microscopy. LAR was highly enriched along the length of the cellular extensions, and was present in a punctate pattern (Fig. 8, a and b). Talin and vinculin were also present in these extensions, suggesting that focal adhesions are important for the

formation or functioning of these cellular extensions (Fig. 8, c and d). Talin and vinculin are also abundant in the cell proper, where LAR was present in lower abundance. Predominant localization of LAR to the extended cell process suggests a specific role for LAR in this location. Antibody-induced internalization of LAR might inhibit formation of the cellular extensions by affecting the assembly–disassembly cycle of focal adhesion sites.
We have previously shown that the FnIII domain 4, and FnIII domains 6 and 7, are also alternatively spliced. However, no specific ligands were found in our assays for the LAR-FnIII domains 6 and 7 or for the LAR-FnIII domain 4. It is possible that splicing of these FnIII domains functions in fine tuning the laminin–nidogen-binding properties of the FnIII domain 5. The lower laminin–nidogen-binding capacity of construct #8, in which Fn5 is directly linked to Fn8 without the intervening Fn6 and Fn7, lends support to this possibility (Fig. 2). Modulation of ligand binding of the RGD sequence of fibronectin by alternative splicing of a neighboring FnIII domain has been reported (22).

The relatively high level of expression of the non-laminin–nidogen binding LAR Fn5+ isoform in cells of neuronal origin, and upregulation of the LAR-Fn5+ isoform in confluent fibroblasts (25, 42) suggest that the LAR extracellular domain may have other ligands. Such ligand-binding activity may be mediated by the FnIII domains 1–3 or by the NH2-terminal Ig domains of LAR. Ig domains of other cell surface proteins are often involved in homophilic and heterophilic interactions (29).

The LAR-Fn5− isoform is the predominant LAR isoform in most other cell types. Localization of both laminin–nidogen and LAR at focal adhesion sites is consistent with a role for the laminin–nidogen complex as an LAR ligand. However, assessment of the laminin–nidogen-binding capacity of cell surface–expressed LAR is complicated by coexpression of other laminin-binding cell surface molecules such as integrins, heparin sulfate, and β1,4 galactosyltransferase (3, 27). Nevertheless, a role for LAR in modulating laminin-nidogen-induced cell morphological changes is indicated by the LAR antibody crosslinking studies (Fig. 5). The mechanism by which LAR cross-linking inhibits laminin–nidogen-induced cell spreading may be twofold. The first possibility is that laminin–nidogen-LAR binding and subsequent signaling is inhibited by antibody-mediated downregulation of LAR from the cell surface. LAR cross-linking does induce internalization of LAR over the time period used in our studies (31; Fig. 6).

A second possibility is that antibody-induced LAR clustering may interfere with laminin–nidogen-induced LAR cellular localization, or may directly influence LAR PTPase activity, generating a signal antagonistic to laminin–nidogen-induced cell signaling. At the present time, our data cannot distinguish between these possible effects of LAR antibody cross-linking. Localization of both vinculin and talin to the cellular extensions, and high enrichment of LAR in these extensions (Fig. 8) suggest a role for LAR in modulating the assembly/disassembly of focal adhesion sites in these cellular extensions. This role for LAR would be consistent with the previous localization of LAR to the disassembly side of focal adhesions (31).

A crucial factor for regulating laminin-induced cell spreading is modulation of cell signaling through tyrosine phosphorylation (2). Definition of the role of the LAR PTPase activity requires elucidation of the physiological LAR substrate(s) as well as determination of the effect of laminin–nidogen–LAR binding on LAR PTPase activity. Few physiological substrates have been identified for any of the transmembrane PTPases. A number of potential LAR substrates, the tyrosine phosphorylation of which are implicated in cell spreading, are generated by laminin–
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