Trophoblast adhesion of the peri-implantation mouse blastocyst is regulated by integrin signaling that targets phospholipase C

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Abstract

Integrin signaling modulates trophoblast adhesion to extracellular matrices during blastocyst implantation. Fibronectin (FN)-binding activity on the apical surface of trophoblast cells is strengthened after elevation of intracellular Ca²⁺ downstream of integrin ligation by FN. We report here that phosphoinositide-specific phospholipase C (PLC) mediates Ca²⁺ signaling in response to FN. Pharmacological agents used to antagonize PLC (U73122) or the inositol phosphate receptor (Xestospongin C) inhibited FN-induced elevation of intracellular Ca²⁺ and prevented the upregulation of FN-binding activity. In contrast, inhibitors of Ca²⁺ influx through either voltage-gated or non-voltage-gated Ca²⁺ channels were without effect. Inhibition of protein tyrosine kinase activity by genistein, but not G-protein inhibition by suramin, blocked FN-induced intracellular Ca²⁺ signaling and upregulation of adhesion, consistent with involvement of PLC-γ. Confocal immunofluorescence imaging of peri-implantation blastocysts demonstrated that PLC-γ₂, but not PLC-γ₁ nor PLC-β₁, accumulated near the outer surface of the embryo. Phosphotyrosine site-directed antibodies revealed phosphorylation of PLC-γ₂, but not PLC-γ₁, upon integrin ligation by FN. These data suggest that integrin-mediated activation of PLC-γ to initiate phosphoinositide signaling and intracellular Ca²⁺ mobilization is required for blastocyst adhesion to FN. Signaling cascades regulating PLC-γ could, therefore, control a critical feature of trophoblast differentiation during peri-implantation development.

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Introduction

Blastocyst implantation in mice and humans depends on the interaction of differentiated, adhesion-competent trophoblast cells with extracellular matrix (ECM) components of the receptive uterus (Carson et al., 2000; Wang and Armant, 2002; Armant, 2005). Achievement of the adhesion-competent stage by mouse blastocysts correlates with trafficking of α₅β₁ integrins and the concomitant acquisition of fibronectin (FN)-binding activity on the apical surface of trophoblast cells (Schultz and Armant, 1995; Schultz et al., 1997). Blastocysts developing to the adhesion-competent stage exhibit strong FN-binding activity only after exposure to FN or an active fragment, FN-120, through a process dependent on energy, intracellular trafficking and actin microfilament integrity, but requiring no new protein synthesis (Schultz and Armant, 1995). The strengthening of trophoblast adhesion to FN is mediated by ligation of integrins that contain the α5, αv, β1 and β3 subunits and is strongly correlated with subsequent trafficking of the αIIb integrin subunit to the apical plasma membrane (Wang et al., 2002; Rout et al., 2004). Therefore, both “outside-in” and “inside-out” integrin signaling (Burridge and Chrzanowska-Wodnicka, 1996; Aplin et al., 1998; Giancotti and Ruoslahti, 1999) appear to be required by trophoblast cells in order to achieve strong adhesion to FN and progress through the peri-implantation developmental program (Armant, 2005). Evidence suggests that α5β1 is primarily responsible for outside-in signaling, while inside-out signaling positions αIIbβ3 on the surface of the blastocyst to strengthen adhesion (Rout et al., 2004).

Integrin ligation by FN transiently elevates intracellular Ca²⁺ levels in mouse blastocysts and interference with this Ca²⁺ signaling blocks the upregulation of FN-binding activity (Wang et al., 2002). Conversely, direct elevation of intracellular Ca²⁺ with ionomycin strengthens trophoblast adhesion to FN to the same
degree as integrin ligation. The role of intracellular Ca\(^{2+}\) in integrin-mediated signal transduction has been described in several cell types. Mobilization of intracellular Ca\(^{2+}\) followed by protein–tyrosine phosphorylation occurs when cells expressing αβ3 integrin adhere to FN or fibrinogen (Pelletier et al., 1992). Signal transduction stimulated in endothelial cells also involves an interaction between integrins present in focal adhesion complexes and the Ca\(^{2+}\)-dependent kinases, protein kinase C (PKC) and Ca\(^{2+}\)-calmodulin-dependent protein kinase (Berk et al., 1995). Ca\(^{2+}\)-related transducers of intracellular signaling, including inositol 1,4,5 trisphosphate (Ins(1,4,5)P\(_3\))-specific phospholipase C (PLC) and PKC, are recruited to the plasma membrane by α5β1 integrins when ligated by either FN or specific antibodies (Miyamoto et al., 1995). In adhesion-competent trophoblast cells, elevation of intracellular Ca\(^{2+}\) to strengthen trophoblast adhesion to FN appears to require the downstream factors, PKC and calmodulin (Wang et al., 2002). How intracellular Ca\(^{2+}\) signaling is initiated by integrin ligation in mouse blastocysts has not been delineated.

Mechanisms responsible for elevation of intracellular Ca\(^{2+}\) include (1) release from intracellular stores through either the Ins(1,4,5)P\(_3\) or ryanodine receptors, (2) influx through voltage-gated Ca\(^{2+}\) channels and (3) capacitative Ca\(^{2+}\) entry, in which Ca\(^{2+}\) influx occurs subsequent to the release of intracellular Ca\(^{2+}\) stores (store-operated Ca\(^{2+}\) entry) (Putney et al., 2001). Integrin ligation can elevate intracellular Ca\(^{2+}\) levels through Ca\(^{2+}\) influx (Leavesley et al., 1993; Schwartz, 1993), activation of PLC to generate Ins(1,4,5)P\(_3\) and mobilize intracellular Ca\(^{2+}\) stores (Somogyi et al., 1994; Chan et al., 2001), or initiation of capacitative Ca\(^{2+}\) influx (Schottelndreier et al., 2001). The capacity for both Ca\(^{2+}\) influx and mobilization of intracellular Ca\(^{2+}\) stores exist in mouse preimplantation embryos. During early preimplantation development, the activities of PLC and the Ins(1,4,5)P\(_3\) receptor are operative during capacitative Ca\(^{2+}\) entry in response to platelet activation factor (Emerson et al., 2000). Evidence also exists for the presence of Ins(1,4,5)P\(_3\)-sensitive intracellular Ca\(^{2+}\) stores at the 8-cell and blastocyst stages (Stachecki and Arman, 1996; Liu and Arman, 2004), and Ca\(^{2+}\) influx through L- and N-type voltage-gated calcium channels after stimulation by cannabinoids or heparin-binding EGF-like growth factor (HBEGF), respectively, at the blastocyst stage (Wang et al., 2003; Wang et al., 2000b). In the present study, we have examined the mechanism underlying integrin-mediated Ca\(^{2+}\) mobilization in adhesion-competent trophoblast cells during a period when these cells differentiate from an epithelial phenotype into motile, invasive cells.

Materials and methods

Production and culture of mouse embryos

Female NSA or MF1 mice (Harlan Sprague–Dawley, Indianapolis, IN) 5–8 weeks old were superovulated, as previously described (Arman, 2006). Eggs were recovered by flushing oviducts with M2 medium (Sigma Chemical Company, St Louis, MO) the morning after injection of human chorionic gonadotropin. For generation of embryos, female mice were mated with B6SJF1/J males (Jackson Laboratory, Bar Harbor, ME) and embryos were collected by flushing the oviducts with M2 on gestation day (GD) 1–3 or the uterus on GD 4 (where GD1 is the day of the vaginal plug), as previously described (Arman, 2006). All experiments using adhesion-competent blas-
tocysts were conducted by first culturing blastocysts from GD 4 to GD 7 in Ham’s F10 medium containing 4 mg/ml BSA, 100 U/ml penicillin and 0.1 mg/ml streptomycin (all from Sigma) at 37°C in a 5% CO\(_2\)/air incubator.

Preparation of cell extracts

Monolayers of mouse B16-F10 melanoma cells (American Type Culture Collection, Rockville, MD), mouse trophoblast stem (TS) cells and human HTR-8/SVneo cytotrophoblast cells were grown as previously described (Yelian et al., 1995; Kibbun et al., 2000; Tanaka et al., 1998) and harvested in SDS sample buffer containing 1 mM sodium orthovanadate, and protease inhibitors (1 mM PMSF, 25 KIU/ml aprotinin, 2 μM leupeptin, 2 μM antipain, 10 μM benzamidn, 1 μM pepstatin and 1 μM chymostatin; all from Sigma). Mouse spleen was cut into small pieces, rinsed in ice-cold PBS three times and dissociated to prepare a single cell suspension using a Polytron (Omni International, Inc. Waterbury, CT) at 7500 rpm. Spleen cells were collected by centrifugation at 10000g and rinsed three times using cold PBS. Cells were lysed using cold NP-40 lysis buffer (137 mM NaCl, 20 mM Tris, 1% NP-40, 10% glycerol, pH 8 0) containing orthovanadate and protease inhibitors described above. After centrifuging at 13000g for 3 min, the supernatant was recovered and the protein concentration was determined using a Bio-Rad (Rockville Center, NY) DC protein assay kit.

FN-binding activity

FN-binding activity was upregulated by exposing blastocysts to 50 μg/ml FN-120 for 1 h, as previously described (Wang et al., 2000b; Arman, 2006). In some experiments, the blastocysts were treated at 37°C for 1 h before and during exposure to FN-120 by addition of the following inhibitors to the culture medium: 10 μM U73122, 10 μM U73343, 1 μM Xestospongin C, 10 μM flunarine, 10 μM bepridil, 10 μM genistein, 10 μM daidzein or 10 μM suramin (all from EMD Biosciences, Inc., San Diego, CA). Similar treatments were also carried out using 2 μM FTX-3.3, 3 μM α-conotoxin GVIA, 1 μM α-conotoxin MVIIIC or 1 μM calcein (all from Alomone Labs, Jerusalem, Israel).

FN-binding activity was assayed, as previously described (Schultz and Arman, 1995; Arman, 2006), using 1.0 μM fluorescent-green polystyrene microspheres (Polyscience) coated with FN-120. The fluorescence intensity of the bound microspheres was quantified over the abembrionic pole of each blastocyst using computer-based image analysis, as described by Schultz and Arman (1995). Basal FN-binding activity, determined by exposing embryos to BSA in place of FN-120, was subtracted from all values obtained with embryos exposed for 1 h to 50 μg/ml FN-120. The derived change (Δ) in FN-binding activity value was normalized to the mean control values obtained in each experiment and reported as mean±SEM.

Intracellular Ca\(^{2+}\) measurements

For estimation of the intracellular Ca\(^{2+}\) concentration, blastocyst cells loaded for 1 h with 5 μM fluo-3-acetoxymethyl ester (fluo-3-AM) or fluo-4-AM (Molecular Probes, Inc., Eugene, OR) were subjected to epifluorescence microscopy and image analysis, as previously described (Wang et al., 1998). After dye loading, some embryos were treated for 15 min with 2 μM FTX-3.3, 3 μM α-conotoxin GVIA, 1 μM α-conotoxin MVIIIC, 1 μM calcein, 10 μM flunarine, 10 μM bepridil, 10 mM NiCl\(_2\), 1 μM caffeine (Sigma), 10 μM thapsigargin (EMD Biosciences), 10 μM U73122, 10 μM U73343, 1 μM Xestospongin C, 10 μM genistein or 10 μM daidzein. Embryos were imaged individually at 37°C in 5 μl drops of culture medium with or without 50 μg/ml FN-120 on Petri-dishes flooded with mineral oil. To monitor Ca\(^{2+}\) mobilization, embryos were briefly illuminated every 0.25 min for fluorescence imaging. All data presented depict single embryos that are representative of a minimum of 10 embryos for each treatment.

Immunological identification of PLC isoforms

Immunological procedures were conducted using polyclonal antibodies that recognize specific PLC isoforms. Rabbit IgGs recognizing PLC-γ1 (G-12), PLC-γ1 (530) or PLC-γ2 (Q-20) were obtained from Santa Cruz Biotechnology,
Inc. (Santa Cruz, CA). Tyrosine phosphorylation of the two PLC-γ isoforms was detected using antibodies that recognize specific tyrosine-phosphorylated epitopes. PLC-γ1 phosphorylated at tyrosine 783 was labeled with an anti-goat antibody (Tyr 783) from Santa Cruz or a rabbit antibody (Tyr783) from Cell Signaling Technologies, Inc. (Danvers, MA). PLC-γ2 phosphorylated at tyrosine 1217 was labeled with a rabbit antibody (Tyr1217) from Cell Signaling.

For immunofluorescence microscopy, blastocysts were fixed at room temperature for 30 min in 3% paraformaldehyde, washed through 2 drops of 150 mM glycine, pH 7.2 and permeabilized by treatment with 0.1% Triton × 100 (Sigma) for 15 min at room temperature. Five drops of PBS containing 10 mg/ml BSA (PBS/BSA) were used to rinse embryos after all incubations throughout the procedure. Blastocysts were incubated overnight at 4°C with primary antibodies prepared at 10 μg/ml in PBS/BSA. As primary antibody controls, some embryos were incubated with 10 μg/ml non-immune rabbit IgG (Jackson Immunoresearch Laboratories, West Grove, PA). Primary antibodies were detected using 10 μg/ml Texas-Red-conjugated goat anti-rabbit IgG (Jackson ImmunoResearch) incubated with embryos at room temperature for 1 h.

For confocal imaging, Texas-Red-labeled blastocysts were mounted on slides with permanent mounting medium (Chemicon, Temecula, CA) and stored at 4°C. Embryos were viewed on a Zeiss (Thornwood, NY) 310 confocal scanning laser microscope, using identical contrast and gain settings to generate all images. Blastocysts labeled with anti-phospho-PLC antibodies were viewed using epifluorescence microscopy (Leica DM IRB; Wetzlar, Germany). Images captured with an Orca digital camera (Hammamatsu City, Japan) were deconvolved from 1 μm serial optical sections using SimplePCI (C-Imaging system, Cranberry Township, PA) imaging software and a “nearest neighbor” algorithm. The fluorescence intensities of labeled blastocysts were quantified from these images using SimplePCI software, which measured the total fluorescence (grey level) of an area at the outer edge of each embryo delineated within a digital image. The mean intensity of control embryos exposed to FN-120 for 30 min and labeled with non-immune IgG was subtracted from the intensity for each embryo to derive the specific phospho-PLC staining intensity.

Western blotting was used to validate all primary antibodies. Proteins extracted from B16, TS, HTR-8/Svneo and mouse spleen cells were diluted in SDS sample buffer (125 mM Tris, pH 6.8, 4% SDS, 10% glycerol, 0.006% bromophenol blue, 2% β-mercaptoethanol) and electrophoresed (30 μg/lane) on a 7.5% SDS polyacrylamide gel. Separated proteins were then transferred electrophoretically from the gel to polyvinylidene difluoride or nitrocellulose membranes (Millipore, Burlington, MA), and blocked overnight at room temperature with 5% (w/v) nonfat dry milk prepared in PBS. Membranes were incubated overnight at 4°C with 1 μg/ml primary antibody diluted in TTBS (20 mM Tris–HCl, pH 7.6, 145 mM NaCl and 0.1% Tween-20) containing 1% nonfat dry milk. After rinsing three times with TTBS, the membranes were incubated for 1 h at room temperature with 1:300 horseradish peroxidase-conjugated donkey anti-rabbit IgG (Jackson Immunoresearch) in TTBS/1% nonfat dry milk. After three rinses in TTBS, the probe was visualized using enhanced chemiluminescence, as previously described (Kilburn et al., 2000).

Statistical analysis

All experiments were repeated at least three times. Values reported for FN-binding activity were obtained using at least 15 blastocysts per treatment group. Differences between treatment groups in the FN-binding activity and PLC staining intensity were tested for significance using a one-way ANOVA and the Bonferroni/Dunn post hoc test.

Results

Upregulation of FN-binding activity does not depend on Ca2+ influx

To investigate whether FN-120 induces an intracellular Ca2+ transient in blastocysts by initiating Ca2+ influx through voltage-gated Ca2+ channels, we have examined the effects of Ca2+ channel (L-, N-, P-, Q- or T-type) blockers on FN-120-mediated Ca2+ signaling and upregulation of FN-binding activity. A similar approach was used previously for blastocysts treated with HBEGF to demonstrate that inhibitors of N-type Ca2+ channels significantly attenuate the acceleration of development (Wang et al., 2000b). None of the blockers tested significantly inhibited the ability of FN-120 to induce a Ca2+ transient or upregulate FN-binding activity (Table 1), suggesting that voltage-gated Ca2+ channels are not required for integrin signaling in trophoblast cells.

Integrins can induce Ca2+ influx through non-voltage gated Ca2+ channels (Schottelndreier et al., 2001). It was considered inadvisable to chelate extracellular Ca2+ because of possible interference with integrin binding to FN-120, as found by Schultz and Armant (1995) using EDTA. As an alternate approach, Ca2+ influx was inhibited by NiCl2. Blastocysts were first treated with 10 mM NiCl2 and then exposed to FN-120 in the continued presence of inhibitor prior to assessing intracellular Ca2+ or FN-binding activity. NiCl2 did not reduce FN-induced elevation of intracellular Ca2+ (Fig. 1) or upregulation of FN-binding activity (not shown), suggesting that influx of extracellular Ca2+ is not the primary source for elevation of intracellular Ca2+ levels during this process.

FN elevates cytoplasmic free Ca2+ through release from Ins(1,4,5)P3-sensitive Ca2+ stores

To examine whether FN-120 elevates intracellular Ca2+ through release from intracellular Ca2+ stores, FN-induced Ca2+ signaling was examined in blastocysts treated with either caffeine or thapsigargin to deplete ryanodine- or Ins(1,4,5)P3-sensitive intracellular Ca2+ stores, respectively. FN-120 induced an intracellular Ca2+ transient in caffeine-treated blastocysts, but not in thapsigargin-treated embryos (Fig. 1), suggesting that integrin signaling triggered by FN-120 primarily targets Ins(1,4,5)P3-sensitive intracellular Ca2+ stores.

Table 1

<table>
<thead>
<tr>
<th>Ca2+ channel blockers</th>
<th>Channel type</th>
<th>ΔFN-binding activity</th>
<th>Ca2+ elevated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td></td>
<td>99.9 ± 5</td>
<td>Yes</td>
</tr>
<tr>
<td>2 μM FTX-3.3</td>
<td>P- / Q-</td>
<td>90.0 ± 15</td>
<td>Yes</td>
</tr>
<tr>
<td>1 μM α-conotoxin MVII</td>
<td>Q-</td>
<td>102 ± 18</td>
<td>Yes</td>
</tr>
<tr>
<td>3 μM α-conotoxin GVIA</td>
<td>N-</td>
<td>110 ± 15</td>
<td>Yes</td>
</tr>
<tr>
<td>1 μM calcisentepine</td>
<td>L-</td>
<td>71.3 ± 16</td>
<td>Yes</td>
</tr>
<tr>
<td>10 μM flunarizine</td>
<td>T-</td>
<td>73.4 ± 10</td>
<td>Yes</td>
</tr>
<tr>
<td>10 μM bepridil</td>
<td>T-</td>
<td>73.1 ± 20</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Blastocysts were treated with 50 μg/ml FN-120 in the absence (Control) or presence of the indicated inhibitors and assayed for ΔFN-binding activity and Ca2+ signaling, as detailed in Materials and methods.

a Values (mean±SEM) were not significantly different according to ANOVA.

b Intracellular Ca2+ increased from about 200 nM to at least 365 nM after adding FN-120 where scored as elevated. At least 10 embryos all tested positively for each treatment.
Integrin ligation mobilizes intracellular Ca\textsuperscript{2+} stores through activation of PLC

Ins(1,4,5)P\textsubscript{3} is produced by activation of phosphoinositide-specific PLC and the subsequent hydrolysis of phosphatidylinositol 4,5-bisphosphate (Rhee and Bae, 1997). To examine the role of PLC activity and Ins(1,4,5)P\textsubscript{3} signaling after ligation of integrins by FN, blastocysts were treated before exposure to FN-120 with 10 \( \mu \)M U73122, a potent inhibitor of phosphoinositide-specific PLC, or 1 \( \mu \)M Xestospongin C, an Ins(1,4,5)P\textsubscript{3} receptor antagonist (Gafni et al., 1997). Both inhibitors prevented the appearance of a Ca\textsuperscript{2+} transient (Fig. 2A) and blocked the upregulation of FN-binding activity (Fig. 2B), suggesting that activation of the PLC/Ins(1,4,5)P\textsubscript{3} signaling pathway is required for both events. U73343, a less active structural analog of U73122, inhibited neither the upregulation of FN-binding activity nor induction of intracellular Ca\textsuperscript{2+} transients (Figs. 2A, B). It can be concluded that PLC operates upstream to Ca\textsuperscript{2+} mobilization in the integrin signaling pathway initiated by FN-120.

PLC-\( \gamma \) mediates outside-in signaling during upregulation of FN-binding activity

Six major families of PLC isoforms (PLC-\( \beta,-\gamma,-\delta,-\varepsilon,-\zeta \) and \(-\eta\)) have been identified in mammalian tissues that include at least 13 different isozymes (Rebecchi and Pentyala, 2000; Harden and Sondek, 2006). PLC-\( \gamma \) isoforms are activated by protein tyrosine kinases (PTK), while PLC-\( \beta \), PLC-\( \delta \) and the other isoforms are activated through pathways dependent on heterotrimeric G proteins or small GTPases of the Ras family (Rebecchi and Pentyala, 2000; Harden and Sondek, 2006). We have examined the expression of three PLC isozymes in mouse embryos using antibodies that recognize proteins of the expected molecular weights and tissue distribution. As expected, PLC-\( \beta 1 \) and PLC-\( \gamma 1 \) were detected by western blot in melanoma and trophoblast cells, while PLC-\( \gamma 2 \) was only detected in spleen (Figs. 3A–C). Trophoblast cells in adhesion-competent mouse blastocysts expressed both PLC-\( \beta \) and PLC-\( \gamma \) isoforms (Fig. 4), indicating that either a tyrosine kinase or G protein-based PLC-
activation pathway could link integrin ligation to Ca\(^{2+}\) mobilization.

In an effort to resolve the principal signaling pathway upstream to PLC, blastocysts were treated with suramin to inhibit G-protein activity, as well as genistein to block activation of PTK. Genistein effectively blocked the elevation of intracellular Ca\(^{2+}\) induced by FN-120 (Fig. 5A) and the upregulation of FN-binding activity (Fig. 5B), while the G-protein inhibitor had no effect on the ability of FN-120 to upregulate FN-binding activity (Fig. 5C). As a positive control, suramin blocked the upregulation of FN-binding activity by lysophosphatidic acid (data not shown), an inducer of Ca\(^{2+}\) mobilization in blastocysts (Liu and Armant, 2004) that binds to a G-protein-coupled receptor (van Corven et al., 1989). The less active structural analogue of genistein, daidzein, did not inhibit the effects of FN-120 treatment (Figs. 5A, B). These findings suggest that the PTK-dependent PLC-γ isoforms, but not PLC-β1 or other G protein-dependent isozymes, are responsible for Ca\(^{2+}\) signaling during trophoblast interaction with FN.

Expression of PLC-γ during preimplantation development

To delineate the expression of the two PLC-γ isoforms during mouse preimplantation development, unfertilized eggs and embryos at the 1-cell stage to the adhesive blastocyst stage (cultured to GD 7) were assessed by immunofluorescence labeling and scanning laser confocal microscopy. PLC-γ1, a ubiquitously expressed isoform (Rebecchi and Pentyala, 2000), was detected at a relatively high level in unfertilized eggs and zygotes (Fig. 6A). PLC-γ1 declined during preimplantation development, with only trace levels observed in adhesion-competent blastocysts. In contrast, PLC-γ2, which is primarily expressed in the lymphatic system (Rebecchi and Pentyala, 2000), was highly expressed in preimplantation mouse embryos (Fig. 6B). Similar to PLC-γ1, high levels of PLC-γ2 were found in unfertilized eggs and zygotes with decreasing expression after blastocyst formation. However, a relatively high level of PLC-γ2 appeared in adhesive blastocysts on GD7 localized primarily at the apical surface of trophoblast cells, making it accessible to integrin signaling complexes.

PLC-γ phosphorylation during integrin signaling

Activation of PLC-γ is accomplished through tyrosine phosphorylation (Harden and Sondek, 2006) and can be monitored using phosphorylation-specific antibodies. We examined PLC-γ1 and PLC-γ2 phosphorylation in adhesion-competent blastocysts during the period immediately following exposure to FN-120 using antibodies directed against phosphorylated tyrosine residues at positions 783 and 1217 in the respective proteins. Western blotting of mouse TS cell extracts demonstrated that antibodies against both phosphorylated proteins specifically detected bands at the expected molecular sizes, matching antibodies against PLC-γ1 and PLC-γ2 (Fig. 3D). The phosphorylation-specific antibodies were then used to
detect PLC activation in blastocysts by immunofluorescence (Fig. 7A). Five minutes after exposure to FN-120, phospho-PLC-γ1 staining remained at or near background levels, while phospho-PLC-γ2 labeling became intense near the apical surface of the trophoblast. Quantification of the fluorescence intensity indicated no change in PLC-γ1 phosphorylation over a 30-min period; however, phospho-PLC-γ2 labeling increased significantly within 5 min of FN-120 treatment and continued to rise over the next 25 min (Fig. 7B). Specific phospho-PLC staining shown in Fig. 7B was calculated by subtracting the background staining of non-immune IgG, which was low (6.0±0.8). While not ruling out a role for PLC-γ1 or other PLC isozymes, these findings provide strong evidence that PLC-γ2 has a major role in transducing integrin-initiated signals in mouse trophoblast cells.

Discussion

We previously reported that ligation of integrins on the apical surface of adhesion-competent trophoblast cells by FN initiates intracellular Ca\(^{2+}\) signaling and upregulates FN-binding activity (Wang et al., 2002). These biochemical events could be critical for blastocyst adhesion to endometrial ECM during implantation and may modulate adhesive interactions with ECM components as trophoblast cells pass through the basal lamina, invade the decidua and remodel the uterine vascular system. We now present evidence demonstrating that PLC transduces outside-in signaling by integrins upstream to Ca\(^{2+}\) mobilization, and that the responsible isozyme could be PLC-γ2.

Activation of integrins by ECM components frequently initiates intracellular Ca\(^{2+}\) signaling (Ng-Sikorski et al., 1991; Pardi et al., 1989; Smith et al., 1991; Pelletier et al., 1992; Schwartz, 1993; Berk et al., 1995; Sjaastad et al., 1996; Ng-Sikorski et al., 1991; Smith et al., 1991; Pelletier et al., 1992; Schwartz, 1993; Berk et al., 1995; Sjaastad et al., 1996). We have previously reported that RGD-containing peptides and ECM components, as well as activating antibodies against \(\alpha_v\), \(\alpha_5\), \(\beta_1\) and \(\beta_3\) integrins, upregulate trophoblast adhesion to FN (Wang et al., 2002). Since intracellular Ca\(^{2+}\) signaling is required for this process, FN-120 could, in binding to these integrins, initiate Ca\(^{2+}\) influx through specific Ca\(^{2+}\) channels located on the apical surface of adhesive trophoblast cells. Ligation of \(\alpha_\nu\beta_3\) and \(\alpha_5\beta_1\) integrins by RGD-containing ECM components produces Ca\(^{2+}\) influx through L-type Ca\(^{2+}\) channels (Wu et al., 1998). However,
antagonists of most known voltage-dependent Ca\textsuperscript{2+} channels were unable to abolish the ability of FN-120 to elevate either intracellular Ca\textsuperscript{2+} or FN-binding activity of peri-implantation mouse blastocysts. Treatment with all channel blockers simultaneously was also without inhibitory effect (data not shown), and general inhibition of Ca\textsuperscript{2+} influx by NiCl\textsubscript{2} was ineffective, arguing against the involvement of multiple channels that compensate for one another. Therefore, Ca\textsuperscript{2+} influx does not appear to participate in integrin-mediated Ca\textsuperscript{2+} signaling that leads to strong adhesion of mouse blastocysts to FN.

Thapsigargin, which depletes the Ins(1,4,5)P\textsubscript{3}-sensitive intracellular Ca\textsuperscript{2+} stores (Lewis and Cahalan, 1995), abolished the ability of FN-120 to elevate intracellular Ca\textsuperscript{2+}, while depletion of stores regulated by the ryanodine receptor with caffeine did not interfere. These findings indicate that trophoblast adhesion to FN induces Ca\textsuperscript{2+} release from Ins(1,4,5)P\textsubscript{3} receptor-operated Ca\textsuperscript{2+} stores, with or without subsequent capacitative Ca\textsuperscript{2+} influx. The lack of inhibition by NiCl\textsubscript{2} suggests that capacitative Ca\textsuperscript{2+} influx is not a requirement for integrin signaling in this case, although it could facilitate subsequent trophoblast development. A role for the Ins(1,4,5)P\textsubscript{3} receptor in mobilizing intracellular Ca\textsuperscript{2+} was supported by the observed inhibition by Xestospongion C. Production of Ins(1,4,5)P\textsubscript{3} depends on hydrolysis of phosphatidylinositol 4,5-bisphosphate by PLC (Rhee and Bae, 1997; Rebecchi and Pentyala, 2000; Rebecchi and Pentyala, 2000), which is activated by integrins in several cell types (Kanner et al., 1993; Cybulsky et al., 1993). PLC activity was required for Ca\textsuperscript{2+} signaling and the upregulation of FN-binding activity in mouse trophoblast cells, as demonstrated by specific inhibition with U73122.

Adhesion of fibroblasts to FN or treatment with monoclonal antibodies against FN-binding integrins recruits PLC-\gamma into a membrane-associated complex in a PTK-dependent manner (Miyamoto et al., 1995). Activation of PLC-\gamma through tyrosine phosphorylation occurs when platelets bind to collagen (Blake et al., 1994; Ichinohe et al., 1995) and tyrosine phosphorylation of PLC-\gamma is widely reported during integrin-mediated outside-in signaling (Kanner et al., 1993; Blake et al., 1994; Ichinohe et al., 1995; Blake et al., 1994; Ichinohe et al., 1995). When activation of PLC is blocked using PTK inhibitors, both intracellular Ca\textsuperscript{2+} signaling and strengthening of cell adhesion are prevented (Ichinohe et al., 1995; Melford et al., 1997). In peri-implantation blastocysts, inhibition of PTK activity blocked the ability of FN-120 to elevate intracellular Ca\textsuperscript{2+} and upregulate FN-binding activity, which strongly favors a role for PLC-\gamma in mediating integrin signaling by trophoblast cells adhering to FN. Indeed, recent investigations have established a central role for PLC-\gamma in the tyrosine kinase-based signaling cascades that regulate cell adhesion via integrins (Jones et al., 2005; Tvorogov et al., 2005; Watson et al., 2005). Depending on cell type and the initiating signaling source, various PTKs were found to participate in PLC-\gamma phosphorylation, as well as upstream and downstream transduction of signaling. The cytoplasmic PTK, Src, is required for activation of PLC-\gamma1 in fibroblasts (Tvorogov et al., 2005) and a variety of other cell types (Jones et al., 2005) cultured on fibronectin, laminin, collagen or complex basement membrane, as well as in platelets adhering to collagen or fibrinogen (Watson et al., 2005). Cell and platelet spreading in response to integrin engagement, which involves formation of lamellipodia, fails in the absence of PLC-\gamma activation (Jones et al., 2005; Tvorogov et al., 2005;
McCarty et al., 2004; Wonerow et al., 2003). Strong adhesion to FN at the apical surface of trophoblast cells appears to require both PTK and PLC activities, and subsequently leads to cell spreading and migration on the ECM. Our findings suggest that invasive trophoblast differentiation requires the acquisition of integrin signaling pathways that share many elements with other cell types, particularly PLC-γ activation.

In adhesion-competent trophoblast cells, both PLC-β and PLC-γ isoforms were expressed. The finding that suramin did not interfere with upregulation of FN-binding activity diminishes the likelihood of a role in this pathway for PLC-β or other PLC isoforms activated by heterotrimeric G proteins. Suramin inhibits GDP-GTP exchange, the rate limiting step in the activation of Go-subunits, making it a fairly broad inhibitor of heterotrimeric G proteins (Freissmuth et al., 1996). Suramin has poor specificity in that it can inhibit other signaling molecules, but its lack of effect on FN-binding activity supports the interpretation that Go proteins are not involved. However, we cannot exclude the possible involvement of suramin-resistant PLC isoforms that are activated by small GTPases. Due to complex crosstalk, it is conceivable, for example, that PLC-ε could be activated by Ras family GTPases downstream of PTK (Harden and Sonde, 2006). Rho expression and function have been reported as early as the 8-cell stage during mouse development (Clayton et al., 1999). The small GTPases Rho, Rac and Cdc42 are all highly expressed by rat Rhc-1 choriocarcinoma cells and mouse secondary trophoblast giant cells, and appear to be developmentally regulated during trophoblast differentiation (Parast et al., 2001; El Hashash and Kimber, 2006). At this time, the expression of additional PLC isoforms by mouse blastocysts and the role of small GTPases in the upregulation of FN-binding activity are unknown.

The pattern of the expression of both PLC-γ1 and PLC-γ2 during preimplantation development has not previously been investigated, although it is known that both isoforms are expressed in mouse eggs (Mehlmann et al., 1998). We report that the relative levels of both isoforms decrease at the blastocyst stage, a pattern suggesting that both are initially maternal gene products. In adhesion-competent trophoblast cells of late stage blastocysts, levels of PLC-γ2 specifically increased, although both isoforms were detectable. Unlike PLC-γ1, which was distributed primarily within the cytoplasm, and PLC-β1, which localized near cell–cell junctions, PLC-γ2 was located near the apical surface of trophoblast cells. Therefore, PLC-γ2 was strongly expressed in the vicinity of integrins that mediate blastocyst interaction with maternal ECM. Activation of PLC-γ2 immediately after exposure to FN-120 was established by monitoring its phosphorylation at tyrosine 1217. No evidence of PLC-γ1 phosphorylation was found using antibodies from two commercial sources that recognize an epitope containing phosphorylated tyrosine at position 783. We conclude that PLC-γ2 is a prime candidate to mediate FN-induced Ca²⁺ signaling required for strong adhesion to FN during blastocyst implantation.

Knockout of the PLC-γ1 gene in mice causes significant embryonic growth retardation, although no specific systemic defects were noted (Ji et al., 1997). Interestingly, developmental defects, including embryonic death, occur only after GD 8.5, and no placental defects were described. PLC-γ2 deficient mice are viable and apparently have a normal implantation phenotype (Wang et al., 2000a). The possibility that the two PLC-γ isozymes are able to compensate for one another during implantation has not been investigated and no information is yet available regarding a double knockout. Signaling molecules that regulate integrin-mediated adhesion are numerous and frequently have overlapping functions, reducing the likelihood that deletion of a single component will generate a phenotype (Watson et al., 2005). Within the endometrium, the blastocyst encounters a complex matrix that contains not just FN, but numerous components. We have examined FN as an archetypical ECM component, demonstrating its ability to precipitate intracellular signaling that directs invasive trophoblast differentiation. It remains to be determined whether this response is typical of interaction with other ECM components or if there is diversity among the effects of each element to fine tune trophoblast phenotype.

Our study strongly suggests a critical role for PLC-γ in FN-mediated intracellular Ca²⁺ signaling that leads to strengthening of trophoblast adhesion to FN. A deeper understanding of the developmental mechanism that controls blastocyst implantation awaits elaboration of the molecular cascade that regulates PLC-γ activity in trophoblast cells. The biochemical data presented here clearly indicate that PLC-γ activity plays a major role in early trophoblast differentiation and its conversion to an invasive phenotype as directed through interactions with endometrial ECM components.

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