An Endothelial Storage Granule for Tissue-Type Plasminogen Activator

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Abstract. In previous studies we have shown that, after stimulation by a receptor ligand such as thrombin, tissue-type plasminogen activator (tPA) and von Willebrand factor (vWF) will be acutely released from human umbilical vein endothelial cells (HUVEC). However, the mechanisms involved in the secretion of these two proteins differ in some respects, suggesting that the two proteins may be stored in different secretory granules.

By density gradient centrifugation of rat lung homogenates, a particle was identified that contained nearly all tPA activity and antigen. This particle had an average density of 1.11–1.12 g/ml, both in Nycodenz density gradients and in sucrose density gradients. A similar density distribution of tPA was found for a rat endothelial cell line and for HUVEC. After thrombin stimulation of HUVEC to induce tPA secretion, the amount of tPA present in high-density fractions decreased, concomitant with the release of tPA into the culture medium and a shift in the density distribution of P-selectin. vWF, known to be stored in Weibel-Palade bodies, showed an identical distribution to tPA in Nycodenz gradients. In contrast, the distribution in sucrose gradients of vWF from both rat and human lung was very different from that of tPA, suggesting that tPA and vWF were not present in the same particle.

Using double-immunofluorescence staining of HUVEC, tPA- and vWF-containing particles showed a different distribution by confocal microscopy. The distribution of tPA also differed from the distribution of tissue factor pathway inhibitor, endothelin-1, and caveolin. By immunoelectronmicroscopy, immunoreactive tPA could be demonstrated in small vesicles morphologically different from the larger Weibel-Palade bodies. It is concluded that tPA in endothelial cells is stored in a not-previously-described, small and dense (d = 1.11–1.12 g/ml) vesicle, which is different from a Weibel-Palade body.

O f the physiologically occurring plasminogen activators, tissue-type plasminogen activator (tPA) is the most important one in triggering physiological fibrinolysis and thrombolysis. Transgenic mice in which the tPA gene has been functionally disrupted (Carmeliet and Collen, 1996a,b) and that consequently do not have tPA circulating in their blood, show a reduced thrombotic potential and an increased thrombogenic tendency. Also, clots from which tPA has been removed (functionally by inactivation, or immunologically by immunosorption) will not lyse (Wun and Capuano, 1985, 1987). Circulating tPA is derived from vascular endothelial cells, which have been shown to synthesize tPA both in vivo (Levin and del Zoppo, 1994; Lupu et al., 1995b; Padró et al., 1995; Schneiderman et al., 1995) and in vitro (for review see van Hinsbergh et al., 1991; Emeis et al., 1996; Kooistra and Emeis, 1997). In vivo, tPA can also be demonstrated in vascular endothelial cells by functional and immunocytochemical techniques, although the precise localization of tPA in endothelial cells is still not settled (for review see Emeis et al., 1996). In culture, endothelial cells constitutively secrete tPA into the medium. Regulated secretion from endothelial cells has been demonstrated as well, both in vivo (for review see Emeis, 1988, 1996; Emeis et al., 1996) and recently in vitro (van den Eijnden-Schrauwen et al., 1995, 1997).

A similar situation pertains to von Willebrand factor (vWF), another endothelial cell protein that is synthesized by, and stored in, the endothelium and is secreted from endothelial cells by both constitutive and regulated secretion (for review see Reinders et al., 1988; Wagner, 1990, 1993; Mayadas and Wagner, 1991; Hop and Pannekoek, 1996). In most experimental and clinical situations, tPA and vWF are secreted simultaneously upon stimulation (Tranquille and Emeis, 1990; van den Eijnden-Schrauwen et al., 1995;
for review see Eimeis, 1996). Recent data, however, suggest that the regulated secretion of tPA and vWF are differentially regulated, so that the release of the two proteins can be induced separately (van den Eijnden-Schrauwen et al., 1997). For this to be possible, tPA should be stored in a secretory granule different from the Weibel-Palade body in which vWF is stored in endothelial cells (for review see Reinders et al., 1988; Wagner, 1990, 1993). No storage granule for tPA has yet been described, and the possibility is still open that tPA is also stored in Weibel-Palade bodies. The present study was therefore designed to isolate and describe the endothelial storage particle for tPA and to investigate whether this particle was identical to the Weibel-Palade body, the storage granule for vWF.

Materials and Methods

Materials

Nycodenz was obtained from Nycomed AS (Oslo, Norway); sucrose for density gradients from BDH (Poole, UK). Human α-thrombin, BSA, cycloheximide, N-[3-(2-furyl)acryloyl]-t-phenylalanylglucylglycine and kits for the determination of lactate dehydrogenase (LDH: No. LD-L 10) and 5′-nucleotidase (No. 253-3) were purchased from Sigma Chemical Co. (St. Louis, MO). Recombinant human tPA (Activase) was from Genentech (So. San Francisco, CA), 4-chloro-1-naphthol from Merck (Darmstadt, Germany), and sodiumpentobarbital (Nembutal) from Sanofi (Paris, France).

Murine monoclonal antibodies against human tPA (clones ESP-4, ESP-5, ESP-6, PAM-3) and rabbit polyclonal anti-recombinant human IgA (AD135R) were purchased from American Diagnostica (Greenwich, CT). The murine monoclonal anti-tPA antibodies 8C11 and 2B5 were from Celsus Laboratories (Cincinnati, OH). Rabbit polyclonal anti-mouse tPA was a gift from Dr P. Carmeliet (Leuven, Belgium). Rabbit anti-vWF, murine monoclonal anti-vWF IgG, peroxidase-conjugated swine anti-rabbit IgG, and antibody diluent were from DAKO (Glostrup, Denmark); rabbit anti-caveolin was from Transduction Laboratories (Mamhead Castle, UK). Rabbit anti-human P-selectin (Coughlan et al., 1994) was kindly donated by Dr M.C. Berndt (Prahran, Australia); rabbit anti-human TFPI was a gift from Dr C. Lupu (Thrombosis Research Institute, London, UK); and rabbit anti-human endothelin-1 was a gift from Dr J. Morton (Royal Postgraduate Medical School, London, UK). Texas red- and FITC-conjugated secondary antibodies, as well as Vectashield fluorescence mounting medium were from Vector Laboratories (Petersborough, UK). Goat anti–mouse and goat anti–rabbit IgG labeled with 5- or 10-nm colloidal gold particles were purchased from Nanoprobes, Inc. (Stony Brook, NY). Lowicryl K4M embedding medium was from Polysciences (Eppelheim, Germany). All other materials used were of analytical grade.

Cells and Tissues

Human umbilical vein endothelial cells (HUVEC) were isolated by collagenase digestion (Jaffe et al., 1973) and cultured in Medium 199 supplemented with 10% (vol/vol) heat-inactivated newborn calf serum, 10% (vol/vol) pooled human serum, 100 μg/ml streptomycin, and 2 mM 1-glutamine, as described (van Hinsbergh et al., 1985). Confluent first-passage cells were used throughout. Regulated secretion (acute release) of tPA and vWF from HUVEC was induced in cells that had been preincubated for 30 min in M199 containing 0.3 mg/ml human serum albumin, 1-glutamine, and antibiotics, but no serum. To induce regulated secretion, 15 μl human α-thrombin (final concentration 1 NIH U/ml) was added, and the medium was collected after 3 min, as described (van den Eijnden-Schrauwen et al., 1995).

The rat endothelial cell line RHE, kindly provided by Dr C.A. Diglio (Wayne State University School of Medicine, Detroit, MI; Diglio et al., 1988) was cultured in DME containing 10% (vol/vol) fetal calf serum, 100 IU/ml penicillin, 100 μg/ml streptomycin, and 2 mM 1-glutamine. For experiments, cells were cultured in 6-well plates and were given fresh medium 24 h before an experiment.

Rat lung, mouse lung, and mouse diaphragm were obtained from Nembutal-anaesthetized (60 mg/kg i.p.) animals. Human lung was obtained as anonymous and nontraceable material from lung cancer surgery. Lung specimens for fractionation were washed free of blood and stored in cold saline.

Cell and Tissue Homogenization

Lung tissue was finely divided by a McIlwain tissue chopper and suspended in homogenization buffer (5 mM Tris-HCl, 220 mM sucrose, pH 7.4, 1 mg/g tissue). Confluent HUVEC (4 cm², ~10³ cells) were washed once and scraped into 2 ml homogenization buffer. For HUVEC homogenates, which contain only low amounts of protein, this buffer also contained 0.001% Tween-80 to improve tPA recovery (at this concentration, Tween-80 did not lyse organelles). For the experiments on thrombin-induced secretion, 300 cm² HUVEC cultures were used and scraped into 5 ml buffer. Cells or tissue were homogenized with ten strokes (1500 rpm) in a glass-teflon Potter-Elvehjem tissue homogenizer, and the homogenates were then centrifuged at 800 g for 5 min at 4°C. An aliquot of the supernatant (“total homogenate”) was retained to calculate recoveries, and the remainder was fractionated.

Nycodenz Density Gradient Centrifugation

A Nycodenz solution containing 35% (wt/vol) of Nycodenz was prepared in 5 mM Tris-HCl buffer (pH 7.4). A Nycodenz stock solution of 30% (wt/vol) was prepared in homogenization buffer. From this stock solution, solutions of Nycodenz of 27.5 to 5% (in increments of 2.5%) were prepared in homogenization buffer. A density gradient was made in 14 ml ultracentrifuge tubes, starting with one ml of 35% Nycodenz, and then in eleven steps of one ml from 30% to 5% Nycodenz, and left to equilibrate overnight at 4°C. Two ml of cell or tissue homogenate was layered on top of the gradient, and the gradient was centrifuged at 250,000 g (202,000 gcv) in a Beckman L-7.57 ultracentrifuge equipped with a SW-40 rotor, at 4°C. Subsequently, one ml fractions were collected, starting at the bottom, and stored frozen. The density of the fractions was determined in a Mettler/PAar Dama 602M density measuring cell, and ranged from 1.03-1.17 g/ml (see Results section). For additional protocol see Wanders et al. (1987).

Sucrose Density Gradient Centrifugation

A 46% (wt/vol) stock solution of sucrose (1.375 M) in 5 mM Tris-HCl, pH 7.4, was prepared (for HUVEC, 0.001% Tween 80 was added). From this stock, sucrose solutions of 43 to 10% (in increments of 3%) were prepared in buffer. A sucrose density gradient (46–10% sucrose; 12 steps of 1 ml) was made in 14-ml tubes and left to equilibrate overnight at 4°C. 2 ml of cell or tissue homogenate was layered on top of the gradient, and the gradient was centrifuged exactly as described above for the Nycodenz gradient. 1-ml fractions were collected, and their densities were measured, as described above. The density of the sucrose gradient fractions ranged from 1.03 to 1.16 g/ml.

Assay of Fractions

In the fractions the following parameters were determined, using minaturized assays in microtiter plates. Where necessary, data were corrected for interference by the Nycodenz or sucrose content of the fractions. Protein concentration was measured spectrophotometrically by the BCA protein assay procedure as prescribed by the manufacturer (Pierce, Rockford, IL). tPA activity (Verheijen et al., 1982) and plasminogen activators inhibitor type-1 (PAI-1) activity (Verheijen et al., 1984) were determined by spectrophotometric assay. Human tPA antigen (Schrauwen et al., 1994), rat tPA antigen (Eimeis et al., 1995), PAI-1 antigen (Niemann-Hilbing et al., 1995), vWF antigen (Tranquille and Eimeis, 1990), and cellular fibronectin antigen (Friedman et al., 1995) were measured by ELISA.

Acid phosphatase (substrate: p-nitrophenylphosphate at pH 5.0), alkaline phosphatase (substrate: p-nitrophenylphosphate at pH 10.5), neutral esterase (substrate: p-nitrophenylacetate at pH 6.6), 5′-nucleotidase (kit 265-3; Sigma Chemical Co.), glutamate dehydrogenase (using α-ketoglutaric acid and NADH at pH 8.0), LDH (kit LD-L 10; Sigma Chemical Co.), and angiotensin-converting enzyme (ACE; substrate N-[3-(2-furyl)acryloyl]-t-phenylalanylglucylglycine by the method of Butterly and Stuart [1993]) were determined spectrophotometrically, using miniaturized procedures adapted to a microtiter plate reader (Titertek Multiscan MCC/360; Flow Laboratories, Irvine, Scotland).
Western Blotting of P-Selectin

The presence of P-selectin in Nycodenz fractions of HUVEC was determined semi-quantitatively by slot blotting, using a polyclonal rabbit anti-human P-selectin (Coughlan et al., 1994). After blotting 25 μl of the fractions on nitrocellulose paper, the paper was blocked with 10 mM Tris-HCl, pH 7.4, containing 150 mM NaCl, 0.05% Tween 80, and 3% BSA. Subsequently, the paper was incubated with polyclonal rabbit anti-P-selectin (2.3 μg/ml), followed by incubation with peroxidase-conjugated swine anti-rabbit Ig, washing, and staining with 2.8 mM 4-chloro-1-naphthol plus 0.01% H2O2 in 20% methanol/80% 200 mM NaCl plus 50 mM Tris-HCl, pH 7.4. Blots were densitometrically scanned, and the background subtracted. Results will be given as relative OD units per fraction.

Immunocytochemistry

First passage HUVEC were cultured to confluency as described above on glass coverslips coated with glutaraldehyde-crosslinked gelatin. Once confluent, cells were fixed with 2% freshly depolymerized p-formaldehyde in PBS containing 5% sucrose (PBS/sucrose) for 1 h at 20°C. Cells were then washed extensively in PBS/sucrose and briefly permeabilized by treatment with a solution of 0.02% saponin in PBS for 10 min at 18°C. Cells were then washed in PBS, quenched with 100 mM glycine in PBS, blocked by treatment with DAKO antibody diluent supplemented with 1% (vol/vol) normal goat or horse serum for 30 min, and incubated for 1 h at 18°C with primary antibody diluted in the same blocking solution. As primary antibodies were used, either a cocktail of monoclonal anti-tPA antibodies (IgG 10 μg/ml) for single labeling or the mixture of these anti-tPA monoclonals with polyclonal rabbit anti-vWf IgG (diluted 1:200) for the double immunostaining approach. To confirm the labeling pattern, in double labeling experiments a rabbit polyclonal anti-tPA IgG in combination with monoclonal anti-vWf antibody was used. All specimens were extensively washed with PBS and incubated for 1 h at 18°C with the secondary antibodies. For single staining we used horse anti-mouse IgG-FITC (diluted 1:100 in PBS) and for double staining a mixture of horse anti–mouse IgG-Texas red and goat anti-rabbit IgG-FITC (both diluted 1:100 in PBS). Cells were again extensively washed in PBS, briefly rinsed in distilled water, and mounted in Vectashield. The fluorescent specimens were examined with a confocal laser scanning unit (MRC600; Bio Rad, Hemel Hempstead, UK) attached to a Nikon Diaphot inverted microscope. The light source was a krypton/argon laser with main lines at 488, 568, and 674 nm. For the visualization of FITC and Texas red staining K1 and K2 filter blocks were used. The recorded images were merged to study colocalization. Some samples were analyzed by serial optical sectioning in the Z-axis of the cells, followed by computer-assisted reconstruction of the images.

Controls to ascertain for the specificity of the binding comprised either replacement of the first antibodies with normal IgG from the same species or incubation of the cells with the conjugates only. Staining for double labeling of tPA and TFPI, endothelin-1, or caveolin, respectively, was performed identically as described for tPA/vWf double labeling, using as antibodies rabbit anti–TFPI, rabbit anti-endothelin-1, and rabbit anti-caveolin, respectively.

Immunogold Electron Microscopy

For immunogold labeling, HUVEC were grown on Thermovan coverslips, washed with serum-free medium, and fixed with 3% (wt/vol) p-formaldehyde (freshly prepared) in PBS for 90 min at room temperature. Mouse lung and diaphragm were fixed by perfusion with 3% glutaraldehyde in PBS for 10 min, followed by immersion fixation at room temperature for a further 2 h. Both the cell monolayers and the mouse tissues were dehydrated in an ascending series of ethanol while the temperature is progressively lowered, following the progressive lowering of temperature protocol (for details see Roth, 1989). Finally, the samples were embedded overnight in 100% Loweryl K4M at −35°C. Polymerization was carried out by irradiating tissue in gelatin capsules with UV light in a low temperature embedding workstation (MS 5000; Microfield Scientific Ltd., Kingston Bagpuize, UK) for 24 h at −35°C, followed by hardening under UV light at room temperature for 1 to 2 d. This sectioning was carried out on a microtome (Ultracut; Reichert-Jung Optische Werke, Wien, Austria), and thin sections were placed on Formvar-coated 200 mesh nickel grids. For immunogold labeling we used the postembedding staining protocol previously described (Lupu et al., 1993). In brief, grids were floated on droplets of PBS containing 20 mM glycine for 5 min to neutralize any free aldehyde groups, transferred to PBS containing 1% (wt/vol) gelatine for 10 min to neutralize “sticky” sites, and washed for 30 min in PBS containing 5% (vol/vol) nonimmune goat serum (PBS-NGS) to quench nonspecific binding. Subsequently, the sections were incubated for 1 h at room temperature with primary antibodies. For HUVEC, we used a cocktail of murine anti–human tPA monoclonals or a polyclonal rabbit anti-human recombinant tPA IgG, both diluted 1:50 in PBS-NGS, and monoclonal mouse anti–human vWf IgG (1:100). For the mouse tissues we have used a polyclonal rabbit anti–mouse tPA diluted 1:50 in PBS containing 1% BSA and then incubated for 1 h at room temperature in the same dilution buffer containing the appropriate gold-labeled secondary antibody. Thereafter, grids were washed with PBS and distilled water, stained with uranyl actate and lead citrate, and observed in a transmission electron microscope (EM201; Philips, Eindhoven, The Netherlands).

Results

Density Gradient Centrifugation of Rat Lung on a Nycodenz Gradient

Rat lung was chosen for the first experiments, since it is, of all rat tissues, richest in tPA (Padró et al., 1990). In pilot experiments, using a Nydodenz gradient with density range 1.05–1.27 g/ml, all tPA and vWF was recovered at densities <1.18 g/ml. In subsequent experiments, therefore, a more shallow gradient (density range 1.03 to 1.17 g/ml) was employed. In such a gradient (Fig. 1, a–d), tPA was found as a single symmetrical peak, with a maximum at density 1.11–1.12 g/ml. Both tPA activity and tPA antigen were found at the same position in the gradient (Fig. 1 a). The correlation between tPA antigen and activity in the 14 fractions was in all four experiments >0.97. In those two fractions that contained maximal amounts of tPA, ~50% of tPA (Fig. 1 a) and 6–7% of protein (Fig. 1, c and d) was recovered. vWF antigen also peaked at density 1.11–1.12 g/ml, but in addition showed a second, smaller peak at density 1.06–1.07 g/ml (Fig. 1 b). Tissue protein, and most of the marker enzymes measured, showed peak values in the fractions with densities of 1.03 to 1.08 g/ml (Fig. 1, c and d). Typically, >70% of the protein was recovered in the these fractions. Maximal alkaline phosphatase and ACE activity (used as plasma membrane marker enzymes) were found around d = 1.05–1.08 g/ml; acid phosphatase (a marker for lysosomes), neutral esterase, and 5’-nucleotidase (markers for microsomes) around d = 1.05–1.07 g/ml, and LDH (a marker for the cytosol) ~d = 1.03 g/ml. Only glutamate dehydrogenase activity, a marker enzyme for mitochondria, was recovered at d = 1.11–1.12 g/ml, the density of maximal tPA and vWF concentrations. Cellular fibronectin, a marker of connective tissue contamination, was found at density 1.06 g/ml.

Centrifugation of the two-peak fractions (d = 1.11–1.12 g/ml; diluted 1:1 with homogenization buffer to reduce density) on an identical Nycodenz gradient resulted in recovery of tPA and vWF at the density of the original fractions (Fig. 2 a). To decide whether the distribution of tPA in the gradient might have been due to binding of particle-free tPA to a particle of d = 1.11–1.12 g/ml, recombinant human tPA (Activase; final concentration 50 ng/ml) was added to a rat lung homogenate before gradient centrifugation. Subsequently, we separately measured the exogenously added human tPA antigen and the endogenous rat tPA antigen by species-specific tPA ELISA assays. Rat tPA was recovered as a single peak at its normal density, while the human tPA was found distributed diffusely...
through the gradient (Fig. 2b). Upon ultracentrifugation of a lung homogenate in homogenization buffer without Nycodenz (d = 1.03 g/ml), >90% of tPA and vWF was recovered at the bottom of the tube. After ultracentrifugation in homogenization buffer without Nycodenz but now containing 1% Triton X-100 to lyse organelles, tPA and vWF were diffusely distributed through the gradient (not shown). From these data it was concluded that almost all tPA was present in rat lung homogenates in sedimentable particles of d = 1.11–1.12 g/ml.

To see whether any of the tPA or vWF could be ascribed to tPA or vWF in a pathway of ongoing protein synthesis, rats were pretreated with cycloheximide (2 mg/kg in vivo) at 3 h before being killed. This procedure inhibits rat protein synthesis fully but does not significantly influence the size of the storage pool in rat lung (Tranquille and Emeis, 1989). Cycloheximide treatment did not significantly influence the distribution of tPA or vWF from rat lung homogenate. Also, the amount of vWF present around d = 1.06 g/ml, which might have been located in the protein synthesis pathway, was unchanged (data not shown). As discussed above, no tPA peak is present at this density.

Fractionation of a cell homogenate from a rat heart endothelial cell line (Diglio et al., 1988) on a Nycodenz gradient showed that the bulk of tPA (antigen and activity) was found distributed similarly as in rat lung, though at a slightly lower density of 1.09–1.10 g/ml (not shown). This cell line does not synthetize vWF.

Density Gradient Centrifugation of Rat Lung on a Sucrose Gradient

Using a sucrose gradient of d = 1.03–1.16 g/ml, tPA (both antigen and activity) was found at the same density (1.11–1.12 g/ml) as in the Nycodenz gradient.
1.12 g/ml (Fig. 3). vWF, however, showed a different distribution than in Nycodenz gradients, as no vWF peak was present in fractions 6 and 7 of sucrose density gradients. This resulted (Fig. 3) in a clear separation of tPA (peak in fractions 6 and 7) and vWF (peak in fractions 10 and 11). Marker enzymes for subcellular fractions, including the mitochondrial marker glutamate dehydrogenase, showed, in sucrose gradients, a similar distribution as in Nycodenz gradients (not shown).

Density Gradient Centrifugation of Human Lung

Homogenates of human lung were fractionated on a Nycodenz density gradient and on a sucrose density gradient. The distribution of tPA (antigen and activity) and vWF was more complex in human lung, and less tPA and vWF was present in fractions 7 and 8 of the gradients, compared to rat lung. In sucrose gradients, vWF had practically disappeared from fraction 7 and 8. Of the vWF recovered, 14% ± 3% was found in fractions 7 and 8 in Nycodenz gradients, but only 2% ± 2% in sucrose gradients (mean ± SD, \( n = 4 \); \( P < 0.01 \)). No such difference between Nycodenz and sucrose gradients was found for the tPA content of fractions 7 and 8 (19% ± 4% in Nycodenz gradients versus 16% ± 4% in sucrose gradients, \( n = 4 \), \( P = 0.33 \)).

Density Gradient Centrifugation of HUVEC

First passage HUVEC (60 cm²/experiment) were fractionated on Nycodenz gradients. In Fig. 4, the averaged distribution of tPA antigen and of vWF is shown for five such fractionations. Slight differences in density distribution, especially for tPA, were found between cultures, but the overall distribution pattern (Fig. 4) resembled that found in rat lung. For tPA, the major peak was found at \( d = 1.105 \) g/ml (fraction 6), with a (variably pronounced) shoulder at \( d = 1.13 \) g/ml (fraction 4). A second, smaller peak was seen at \( d = 1.06 \) g/ml (fraction 10). vWF had a single major peak (\( d = 1.105–1.115 \); fractions 5 and 6), and, in three out of five cultures, a small shoulder at \( d = 1.06 \) g/ml (fraction 11). No sucrose density gradient centrifugation was performed on HUVEC.

Thrombin Stimulation of tPA Secretion

For experiments involving thrombin stimulation of regulated secretion (acute release) of tPA, 300-cm² HUVEC cultures were used that had been preincubated for 24 h with 1 mM sodium butyrate to increase tPA synthesis and storage (Kooistra et al., 1987). HUVEC were stimulated with 1 NIH U/ml of human α-thrombin for 3 min (van den Eijnden-Schrauwen et al., 1995), scraped into cold buffer, and homogenized. Control cultures, not treated with thrombin, were run in parallel. After thrombin, a loss of tPA antigen from the high density range of the gradient (fractions 5–8) was observed, compared to the parallel control cultures. Two typical cultures (out of five) are shown in Fig. 5, \( a \) and \( b \). In the experiment shown in Fig. 5 \( a \), tPA disappeared mainly from density range 1.11–1.12 g/ml (fractions 6–8), while in the experiment shown in Fig. 5 \( b \), tPA was reduced in fractions 4–6 (density 1.14 g/ml). The differences between cultures possibly reflect greater experimental variation in cultured HUVEC than in experiments involving rat lungs (note the small SDs in Fig. 1, \( a \) and \( b \)).

Although the differences in tPA concentration in the fractions induced by thrombin were relatively small, immunocytochemistry of tPA antigen in these cells showed that, after 3 min of thrombin stimulation, the granular tPA staining had disappeared from most cells (Fig. 6), while in other cells a weak granular staining persisted. This loss of staining is in agreement with biochemical data, which showed that, after 1 NIH U/ml thrombin stimulation, no further acute release of tPA could be obtained by ionomycin (van den Eijnden-Schrauwen et al., 1995). By Western blotting, P-selectin was detected (using the 300-cm² HUVEC cultures) in the density range 1.10–1.15. After throm-
bin stimulation, the distribution of P-selectin shifted (one to two fractions) to higher densities (Fig. 7) and became more intense.

**Immunohistochemical Localization of tPA and vWf in HUVEC**

The cellular localization of tPA in HUVEC was visualized by indirect immunofluorescence labeling and confocal microscopy, as well as at the ultrastructural level by postembedding immunogold labeling and electron microscopy. In formaldehyde-fixed, saponin-permeabilized HUVEC, tPA showed, by immunofluorescence, well defined granular staining in the cytoplasm of the cells (Fig. 8a; compare with Fig. 6). To enable precise localization of the fluorescence, some specimens were analyzed by serial optical sectioning followed by computer-assisted reconstruction of the image. Serial optical sectioning along the Z-axis revealed that granular tPA staining was localized in the perinuclear, organelle-rich area as well as in the attenuated areas at the cell edges (Fig. 9).

The granular localization of tPA was confirmed by postembedding immunogold labeling of ultrathin sections of HUVEC (Fig. 8a and b) and of vascular endothelial cells in mouse lung and diaphragm capillaries (Fig. 8d and e). The gold particles delineating tPA were present throughout the cells, often in clusters over small, fairly electron-dense vesicles, which in favorable sections could be seen to be surrounded by a membrane (Fig. 8b, d, and e). The location of the gold particles suggested that they were superimposed on the small vesicles that are abundantly present in endothelial cells after routine fixation and embedding and that are much smaller than Weibel-Palade bodies. The gold particles delineating vWf were detected on much larger, elongated particles, which contained densely packed fibrillar material and had the morphological characteristics compatible with Weibel-Palade bodies (Fig. 8c, W-Pb).

**No Colocalization of tPA with vWf in HUVEC**

To further differentiate between tPA- and vWf-containing granules in HUVEC, we performed double immunolabeling for tPA and vWf with Texas red- and FITC-labeled secondary antibodies and simultaneous visualization of the two proteins by confocal microscopy. To detect whether the two proteins were colocalized, the two images were merged; any superimposed green and red areas indicate colocalization. The images show that tPA and vWf are not colocalized in HUVEC, as evidenced by the distinct red and green fluorescence patterns (Fig. 6a and b). The gold particles delineating tPA were present throughout the cells, often in clusters over small, fairly electron-dense vesicles, which in favorable sections could be seen to be surrounded by a membrane (Fig. 8b, d, and e). The location of the gold particles suggested that they were superimposed on the small vesicles that are abundantly present in endothelial cells after routine fixation and embedding and that are much smaller than Weibel-Palade bodies. The gold particles delineating vWf were detected on much larger, elongated particles, which contained densely packed fibrillar material and had the morphological characteristics compatible with Weibel-Palade bodies (Fig. 8c, W-Pb).
would then appear in yellow. As illustrated by Fig. 8, f-h (f, tPA; panel g, vWF), no colocalization of the two antigens was observed (h, in which e and f are superimposed, so that colocalization of tPA and vWF would show as yellow). Control experiments gave no labeling in all conditions tested (not shown).

No Colocalization of tPA with endothelin-1, caveolin, or TFPI in HUVEC

Using identical techniques of indirect immunofluorescence and confocal microscopy as described above for vWF, we looked for colocalization of tPA with three other endothelial proteins: endothelin-1 and tissue factor pathway inhibitor, which are both secreted by HUVEC, and caveolin, a component of caveolae (Lisanti et al., 1994). As shown in Fig. 10, no evidence for colocalization of these proteins with tPA in HUVEC was obtained. We show elsewhere (Lupu et al., 1997) that caveolin and TFPI, but not tPA, colocalize in HUVEC in caveolae.

Discussion

Remarkably, the first studies on the subcellular fractionation of plasminogen activators (at that time called “fibrinolyso kinase activators,” now known to be tPA) from tissue date 1950 (Tagnon and Palade, 1950; Lewis and Ferguson, 1950) and describe that the enzyme activity resides in the “large granule and microsome” fractions of tissue homogenates (Lewis and Ferguson, 1950). Apart from a few similar studies in the mid-sixties (Lack and Ali, 1964; Ali and Lack, 1965; Sugiyama, 1965; Beard et al., 1968), which located tissue PA activity to lysosomes, no major advances in the localization of tPA in tissues were made. Recently, it was shown that, in cells transfected with tPA, tPA is sorted to a granular compartment in the regulated pathway (Harrison et al., 1996).

 Meanwhile, the Weibel-Palade body has been firmly established to be the storage particle for vWF. This particle, first described as an endothelial organelle characterized by specific morphological features (Weibel and Palade, 1964), was subsequently shown to be the storage organelle for vWF in endothelial cells (for review see Reinders et al., 1988; Wagner 1990, 1993; Mayadas and Wagner, 1991; Hop and Pannenkoek, 1996).

In clinical and in experimental (in vivo and in vitro) studies, the regulated secretion (acute release) of tPA and vWF generally occurs simultaneously (Tranquille and Emeis, 1989, Emeis, 1995, van den Eijnden-Schrauwen et al., 1995). This might be due to the fact that the cellular mechanisms involved in tPA and vWF release from endothelial cells are very similar but also due to colocalization of the proteins in a single particle. The question whether tPA and vWF are colocalized is of practical importance, because colocalization would make it impossible to induce selectively the release of either the procoagulant protein vWF or the anticoagulant protein tPA.

Our data show that tPA and vWF are not located in a single storage granule but in granules that can be separated by sucrose density gradient centrifugation and that can be visualized as separate by light- and electron-microscopic immunocytochemistry. Ultracentrifugation of rat lung homogenate in buffer showed that tPA (and vWF) were present in a sedimentable particle, from which the proteins could be released by Triton treatment. By density gradient ultracentrifugation on a Nycodenz gradient, the density of the particle was determined to be 1.11–1.12 g/ml. The distribution of tPA in the Nycodenz gradient was not due to nonspecific binding of soluble tPA (Fig. 2 b), while the particle was stable upon recentrifugation (Fig. 2 a). In previous experiments, we have shown that tissue stores of tPA, including tPA stores in rat lung, are metabolically highly stable, hardly showing any decrease 3 h after protein synthesis has been inhibited by cycloheximide (Tranquille and Emeis, 1989). Cycloheximide treatment did not cause any change in the density distribution of tPA and vWF either. The large peak of tPA (activity and antigen) found after density gradient fractionation of lung must thus be either active tPA stored in a storage pool, or tPA present in a metabolically inert pool, e.g., extracellular tPA bound to connective tissue. A contribution from extracellular tPA is, however, most unlikely, because connective tissue components (e.g., cellular fibronectin, see Fig. 1 d) equilibrate at a much lower density. The tPA present at $d = 1.11–1.12$ was, moreover, enzymatically fully active, while extracellular tPA would have been present largely as an inactive complex with its inhibitor PAI-1 (Kooistra et al., 1986; Fears et al., 1996).

In Nycodenz gradients, the distribution of vWF and tPA was nearly identical, suggesting that the two proteins could be present in a single storage granule (Figs. 1, a and b, and 2 a). The equilibrium density of this putative storage granule was, moreover, similar to the equilibrium density of Weibel-Palade bodies in other types of density gradient, e.g., Percoll gradients (Reinders et al., 1984; Ewenstein et al., 1987).

In sucrose gradients, however, the distribution of vWF differed from that of tPA. tPA showed essentially the same density distribution as in Nycodenz gradients, but vWF had disappeared from the $d = 1.11–1.12$ g/ml fractions (compare Fig. 3 with Figs. 1, a and b, and 2 a). This observation showed that the two proteins were not stored
in the same granule in rat lung. An explanation for the density shift of vWF might be destabilization of the Weibel-Palade body by sucrose, leading to collapse, or dehydration by the hypertonic sucrose solutions, leading to a change in density. The tPA storage particle would then be immune to these forces. Similar to rat lung, sucrose density centrifugation resulted in human lung in a significant loss of vWF antigen from the $d = 1.11–1.12$ g/ml fractions (from 14% in Nycodenz to 2% in sucrose), without causing a loss of tPA from these fractions (from 19 to 16%).

The distribution of tPA from rat heart endothelial cells was very similar to that from rat lung. In human lung and HUVEC (Figs. 4 and 5) the distribution pattern of tPA and vWF was more complex, although peaks of tPA and vWF could still clearly be identified at $d = 1.11–1.12$ g/ml. Presumably, the presence of other cells (including blood platelets), more connective tissue in the lung, and more subendothelial matrix in HUVEC contributed to the more complex patterns observed in homogenates from human lung and from HUVEC.
Two other small particles have been isolated from endothelial cells. Caveolae (Lisanti et al., 1994; Schnitzer et al., 1995) are plasmalemmal vesicles characterized by the presence of, among others, GPI-anchored proteins and caveolin. Recently, a storage particle for endothelin-1 has been isolated from bovine aortic endothelial cells (Harrison et al., 1995). It is not likely that tPA storage granules are identical to caveolae, since the former are much denser than caveolae and do not contain plasma membrane markers such as alkaline phosphatase (Fig. 1c), as do caveolae (Lisanti et al., 1994). In addition, by double immunofluorescent staining, tPA- and caveolin-containing structures were different (Fig. 10). Tissue factor pathay inhibitor (TFPI) has, in HUVEC, also been localized (Lupu et al., 1995a) by immunocytochemistry in a particle, which contains, in addition, caveolin but not tPA (Lupu et al., 1997), in agreement with Fig. 10. Together, these data suggest that the tPA-containing particle does not contain caveolin, ET-1, and TFPI. We also found no evidence for a colocalization of tPA with ET-1 (Fig. 10i). The data thus suggest, though they certainly do not prove, that the proteins mentioned above, those four that are secreted from HUVEC (vWF, tPA, ET-1, and TFPI) may be located in different structures, which opens the possibility that endothelial secretory proteins are packaged in separate secretory pathways.

Stimulation, by thrombin, of acute secretion of tPA and vWF from HUVEC (van den Eijnden-Schrauwen et al., 1995, 1997) resulted in a loss of tPA from the higher-density fractions (d = 1.10–1.14). This loss coincided with the appearance of tPA and vWF in the culture medium and thus likely reflects the secretion of tPA (vWF) from its storage pool. The loss was accompanied by a slight shift to higher densities. Stimulation with 1 NIH U of human α-thrombin for 3 min induces maximal release of tPA from HUVEC, since neither an increase in thrombin concentration above 1 U/ml nor subsequent stimulation with the calcium ionophore ionomycin induced any additional tPA secretion (van den Eijnden-Schrauwen et al., 1995). Stimulation with 1 U/ml of thrombin was also accompanied by a complete loss of granular tPA staining in some cells and a severe reduction in tPA staining in the others (Fig. 6). The quantitatively, only moderate disappearance of tPA from homogenates of thrombin-stimulated cells was, therefore surprising and may reflect the presence, in endothelial cell cultures, of relatively large amounts of tPA outside the secretory pathway.

Thrombin-induced release of tPA was accompanied (Fig. 7) by a shift to slightly higher densities and an increase in immunoreactivity of P-selectin, a component of Weibel-Palade bodies (Wagner, 1993). Whether the tPA storage granule also contained P-selectin could not be decided from the data and is still under investigation. The cell biological substrate of this density shift of P-selectin remains unexplained. It could reflect the fusion of multiple particles, as described by Richardson et al. (1994) in rat endothelial cells after thrombin stimulation in vivo. The remarkable increase in P-selectin immunoreactivity after thrombin (Fig. 8) is also not explained but might be due to an increased accessibility of P-selectin to antibody after thrombin treatment.

The morphological observations on HUVEC, obtained using immunocytochemical labeling procedures with confocal microscopy and immunoemnicroscopy on thin sections, showed that tPA was localized throughout the cytoplasm of HUVEC in small vesicular structures, which differed from Weibel-Palade bodies in size and morphology. Particles staining for vWF were larger and contained a more electron-dense and fibrillar core. No significant colocalization of tPA- and vWF-containing particles was observed in double immunocytochemical labeling experiments. In summary, the morphological data support and extend the biochemical data that tPA-containing vesicles and vWF-containing Weibel-Palade bodies are distinct entities.

Recent observations (van den Eijnden-Schrauwen, 1996; van den Eijnden-Schrauwen et al., 1997) on regulated secretion of tPA and vWF from HUVEC have shown that several mechanisms are involved in (thrombin-induced)

Figure 9. Sequential display of 10 optical sections in the Z-axis through HUVEC. The cells were double stained for tPA and vWF, and the two proteins were simultaneously visualized in serial optical sections from the apical (section 1) to the basal (section 10) side of the cells. tPA is detected throughout the cell, while vWF is distributed mainly in the perinuclear area. Bar, 25 μm.
Figure 10. Double staining of first passage HUVEC for tPA (left column, a, d, and g) and either TFPI (b), caveolin (e), or endothelin-1 (h). The panels in the right column represent the superposition of staining for tPA (green) with staining (red) for either TFPI (c), caveolin (f), or endothelin-1 (i). In case of colocalization of two proteins, the resultant superimposed image in the right hand column is yellow. Note that only in the case of tPA and endothelin-1 (i) in some cells is superimposition observed. For technical details, see Materials and Methods. Bar, 25 μm.
regulated secretion of tPA and vWF. Regulated secretion of vWF is mainly dependent upon an increase in intracellular calcium and calcium influx, while regulated secretion of tPA requires, in addition, the activation of one or more G proteins (van den Eijnden-Schrauwens et al., 1997). The two secretory pathways also show different sensitivities to actin depolymerization (van den Eijnden-Schrauwens, 1996). Such differences in secretory mechanisms presuppose the existence of two separate storage particles. The present data provide evidence that two such storage particles indeed exist: the Weibel-Palade body for vWF, and a separate, newly described, particle for tPA.

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