Human Pulmonary Endothelial Cells in Culture

ACTIVITIES OF CELLS FROM ARTERIES AND CELLS FROM VEINS

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ABSTRACT Endothelial cells were cultured from various different human vessels, including aortas, pulmonary, ovarian, and umbilical arteries, and pulmonary, ovarian, and umbilical veins. The cultured cells were identified as endothelial cells by the presence of Factor VIII antigen and angiotensin I converting enzyme (kininase II). They retained these markers for at least five passages in culture, and some cells had them for seven passages or more. Endothelial cells from the various vessels were compared with respect to their ability to metabolize angiotensins I and II and bradykinin. Cells from arteries had three to five times the angiotensin I converting enzyme activity as cells from veins. The activity of angiotensinase A (aspartyl aminopeptidase) had a similar distribution, and cells from arteries were consistently more active than cells from veins. Cultures of endothelial cells from pulmonary and umbilical vessels formed prostacyclin in response to mechanical stimulation. Media from cell monolayers that were subjected to a change of medium and gentle agitation inhibited aggregation of human platelets. This inhibitory activity was generated within 2–5 min, and it was not formed by cells that were treated with indomethacin or tranylcypromine. Addition of prostaglandin (PG)H₂ to indomethacin-treated cells restored the ability to form the inhibitor, but cells treated with tranylcypromine were not responsive to PGH₂. In experiments where [¹⁴C]arachidonic acid was added to the cells before stimulation, the major metabolite identified by thin-layer chromatography was 6-keto PGF₁₀. Thus, it appears that pulmonary endothelial cells, as well as umbilical cord cells, can form prostacyclin. In experiments comparing the ability of arterial and venous cells to form prostacyclin, arterial cells were more active than venous cells. These studies of cells from various human vessels suggest that the vascular origin of cultured endothelial cells determines how they metabolize vasoactive peptides and form prostacyclin.

INTRODUCTION

The pulmonary vascular endothelium is a major site for the metabolism of bradykinin and angiotensin I. The angiotensin I converting enzyme (EC 3.4.15.1; kininase II; peptidyl dipeptidase) is localized on plasma membranes of endothelial cells (1, 2) where it can activate angiotensin I and inactivate bradykinin by cleavage of a carboxyl terminal dipeptide from these substrates (3, 4).

Angiotensin I is activated in other vascular beds as well (5–7), and angiotensin I converting enzyme has been identified by fluorescent antibodies in endothelium of aorta (8) and many different organs (2). Although angiotensin II is not inactivated during passage through the pulmonary circulation (9, 10), it is readily inactivated in other vascular beds (11).

The culture of endothelial cells has stimulated studies of their morphologic and functional properties. Endothelial cells have been cultured from veins of human umbilical cords (12–16), vena cavae of rabbits (17), and large arterial vessels of pigs (18–20) and cows (21–24). Most cultured endothelial cells retain the features of endothelium in vivo, including an epithelioid shape and Weibel-Palade bodies in the cytoplasm (25–27). Factor VIII antigen is associated with cells from human veins (12, 15) and with cells from aortas of animals (19, 21, 24). Angiotensin I converting enzyme activity is found in cells from umbilical cord veins (15, 16) and also in aortic endothelial cells from various species (20, 28, 29). Cultured endothelial cells can transport serotonin (22) and adenosine nucleotides (20), and they can form prostacyclin in response to mechanical or enzymatic stimulation (30–33).

The extensive surface provided by pulmonary capillary endothelium could account for the efficiency of the lung in metabolism of vasoactive peptides (11). Alternatively, the pulmonary vascular endothelium may have greater enzymatic activities than endothelium in other vascular beds. This study of endothelial cells cultured from different human vessels was made to determine whether the origin of the cell affects its ability to metabolize vasoactive peptides or to form prostacyclin.

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Some results were reported previously in abstracts (16, 34).

METHODS

**Materials.** Culture media, trypsin, collagenase, and antibiotics (penicillin, fungizone, and streptomycin) were purchased from Grand Island Biological Co. (Grand Island, N.Y.). Bradykinin and angiotensin II were purchased from Bachem Fine Chemicals Inc. (Marina del Rey, Calif.). Assay kits for angiotensin I converting enzyme were from Ventrex, Inc. (Portland, Maine). Antibodies to Factor VIII antigen and fluorescein isothiocyanate (FITC)-labeled goat anti-rabbit gamma globulin were purchased from Behring Diagnostics (Somerville, N.J.). Tranyplycromine was obtained from SmithKline Diagnostics Div., SmithKline Corp., Sunnyvale, Calif. Indomethacin, sodium arachidonate, ADP, α-L-aspartyl β-naphthylamide, and β-naphthylamide were from Sigma Chemical Co. (St. Louis, Mo.). Labeled [1-14C]arachidonic acid (40–60 mCi/mmol) was from New England Nuclear (Boston, Mass.). Prostacyclin, prostaglandin (PG)E₂, PGF₂α, 6-keto PGF₁α, and the stable endoperoxide, U-41669, were generously donated by Dr. J. E. Pike of the Upjohn Co. (Kalamazoo, Mich.). The endoperoxide, PGH₂, was kindly provided by Dr. William Campbell of the University of Texas Health Science Center (Dallas, Tex.).

**Cell cultures.** Endothelial cells were cultured from human umbilical cord veins and arteries as described previously (15). Cells were collected from aortas, pulmonary and ovarian vessels by a modification of this technique (see below). Ovarian vessels were obtained from surgical specimens removed during ovariosalpingectomy. All tissues were from individuals between the age of 23 and 54 yr. Pulmonary vessels and aortas were obtained from cadavers within 2–4 h postmortem. This material was taken only from individuals without obvious vascular or pulmonary disease. The ages of the cadaver donors ranged from 20–45 yr.

Most vessels from surgical specimens or cadavers yielded viable cells, and in 55 separate cultures initiated, only 3 failed to grow. Bacterial contamination was responsible for loss of five others. Cultures from 15 different ovarian arteries, 5 ovarian veins, 18 pulmonary arteries, and 9 pulmonary veins were used in this study. Most cultures were examined for Factor VIII antigen and tested for enzyme activity between the first and third passage, but a few cultures were tested as late as the seventh passage.

Pulmonary arteries and veins were dissected from whole lungs down to the level of the third and fourth branching points. Ovarian vessels were dissected from the ovarian ligament, and all vessels were cleaned free of adhering connective tissues, rinsed in protein-free culture medium, and placed on moist, sterile gauze in a small petri dish. While trypsin (0.125%) was used for some preparations from umbilical (15) or pulmonary (16, 34) vessels, most of the cultures in this study were started from cells detached with 0.25% collagenase. The enzyme was introduced into the vessel lumen with a polyethylene-tipped syringe, and the endothelial surfaces were gently rubbed together, but the branches and ends of the vessel were not clamped. The specimens were covered with moist gauze and placed in a 37°C incubator for 10–15 min. The detached cells were collected by washing the endothelial surfaces with protein-free culture medium. The cells were rinsed into tubes containing 5 ml of medium plus antibiotics and 30% fetal calf serum. They were centrifuged to a pellet, washed once, and resuspended in 2–4 ml of medium that contained 20% fetal calf serum and 10% human serum (type O). The cells were plated in 35 × 10 mm culture dishes and were allowed to remain undisturbed in a 37°C incubator for 1–3 h. After cell attachment was ascertained by examining the dishes with an inverted microscope, the media was removed, and the attached cells were covered with fresh medium. The medium on new cultures was changed daily for the first 3 d and then every 3rd d after that.

Cultures of endothelial cells from pulmonary, ovarian, or aortic vessels initiated from a small number of cells (<10⁴) obtained from enzyme treatment of the vessels. At least one passage was required before adequate material was obtained for enzyme assays, but Factor VIII antigen could be determined from plating a portion of the initial culture on coverslips. Umbilical vessels yielded a larger number of cells, but these cultures were usually transferred once before enzyme assay. The number of cells in each culture was determined by counting detached cells in a Coulter counter (Coulter Electronics Inc., Hialeah, Fla.) or by measurement of DNA (35). All cultures used for enzyme determinations were between 70% and 90% confluent, and all cultures, regardless of origin, were grown in the same media and subjected to the same schedule of media change.

**Enzyme assays.** Enzyme assays were made by disrupted cells. The cell monolayers were washed three times with protein-free culture medium (10 ml) and the cells were scraped with a rubber spatula into 0.5–2 ml of medium. The samples were sonicated for 10 s with a Branson probe sonifier (Branson Sonic Power, Danbury, Conn.). The material was centrifuged to a pellet, washed once, and the wash was pooled with the original supernatant medium. Initial determinations of enzyme activities in the supernate and cell pellets showed that more than 90% of the activity was in the supernatant medium.

The activity of angiotensin I converting enzyme was measured with two different substrates, bradykinin and [H]-hippuryl glycyglycylcine. When bradykinin (2 μM) was the substrate, inactivation of the peptide was measured by bioassay on the estrus rat uterus in vitro. Cell-free lysates were incubated with bradykinin in 0.2 M Tris-maleate buffer (pH 7.4) with 0.2 M NaCl at 37°C. Aliquots (50 μl) of the reaction mixture were drawn immediately after and at 2.5 min intervals after addition of the substrate. These were diluted with 0.2–0.5 ml of ice-cold saline and tested immediately on the rat uterus. The amount of kinin remaining in each aliquot was estimated by comparison with standard solutions of bradykinin. The specificity of inactivation by angiotensin I converting enzyme (kininase II) was established by incubating selected samples with 0.1 mM SQ 20881 (36). All bioassays were performed in duplicate for a single cell culture. Enzyme activity was expressed as nanomoles of kinin inactivated per hour per 10⁶ cells.

When [H]-hippuryl glycyglycine was the substrate, [H]-hippuric acid was extracted into ethyl acetate, and the radioactivity was measured in a liquid scintillation counter (37). The reaction was linear with time between 0 and 12 h at 37°C. Triplicate samples were incubated both with and without 0.1 mM SQ 20881 for 0–5 h after addition of the substrate. Activity was calculated from the difference in radioactivity in the organic phase between the 0- and 5-h samples and expressed as nanomoles of substrate hydrolyzed per hours per 10⁶ cells.

The activity of angiotensinase A was measured with both angiotensin II and α-L-aspartyl β-naphthylamide as substrates. Angiotensin II (14 μM) was added to cell samples in 0.2 M Tris maleate buffer with 0.2 M NaCl (pH 7.0). Aliquots for bioassay on the rat uterus were drawn immediately after addition of the

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1 **Abbreviations used in this paper:** FITC, fluorescein isothiocyanate; PG, prostaglandin.
substrate and at 2.5 min intervals thereafter. α-Phenanthroline (1 mM) was added to inhibit the reaction in one set of samples (15).

Aspartyl-aminopeptidase activity was measured in cells incubated with 0.7 mM L-aspartyl β-naphthylamide in 0.2 M Tris maleate buffer (pH 7.0) that contained 5 mM CaCl₂ (38). When α-phenanthroline was used to inhibit the reaction, it was added 15 min before addition of the substrate. All samples were incubated at 37°C for 0, 30, and 60 min, and the reaction was stopped by addition of alkaline EDTA (pH 12.0) to a final concentration of 5 mM. Fluorescence of the product, β-naphthylamide, was measured at 410 nm in an Amino-Bowman spectrophotofluorometer (American Instrument Co., Silver Spring, Md.), and the readings were compared with those of standard concentrations of β-naphthylamide.

**Factor VIII antigen.** Factor VIII antigen was identified in initial and passaged cultures of endothelial cells by an indirect fluorescent antibody technique (12, 39). Cultures were grown on glass coverslips from freshly isolated cells or from cells detached with trypsin at the time of transfer. They were fixed with 2.5% glutaraldehyde for 10 min at room temperature, rinsed with 1% sodium borohydride twice, with phosphate-buffered saline once, and rabbit anti-Factor VIII antigen in a dilution of 1:10 was applied to the surface of the coverslip in a volume of 250 μl. The coverslips were incubated at 37°C for 30 min in an atmosphere of 5% CO₂. They were washed well with three rinses of phosphate-buffered saline (5 ml), and 250 μl of a 1:15 dilution of FITC-labeled goat anti-rabbit gamma globulin was applied. After a second incubation of 30 min at 37°C, the preparations were washed three times with phosphate-buffered saline and a final time with distilled water. They were mounted on slides with Permount (Fisher Scientific Co., Pittsburgh, Pa.) and examined with a Leitz orthoplan fluorescence microscope (E. Leitz, Inc., Rockleigh, N. J.). Controls were coverslips treated only with phosphate-buffered saline and FITC-labeled antiserum or coverslips treated first with nonimmun rabbit serum and then with FITC-labeled antiserum.

**Platelet aggregation.** The inhibition of platelet aggregation by media from endothelial cell cultures was used as a bioassay for prostacyclin (30, 31). Venous blood was drawn from volunteers who had not been on any medication for 10 d previously. The blood was diluted with 0.1 volume of 3% trisodium citrate and centrifuged at 150 g for 10 min. The platelet-rich plasma was removed and stored in a capped plastic tube at 25°C. Platelet-poor plasma was prepared by further centrifugation at 600 g for 20 min. The platelet suspensions were adjusted to ~0.8 x 10⁹ platelets/ml with platelet-poor plasma. Aggregation was measured at 37°C in a Chronolog aggregometer (Chrono-log Corp., Haverton, Pa.). Aggregating agents, ADP (15-20 μM), thrombin (0.2-1 u/ml), sodium arachidonate (0.2-0.7 mM), or U-41669 (1 ng/ml) were added in a volume of 10-50 μl to 1 ml of platelet suspension. Endothelial cell media tested for inhibition of aggregation were added in a volume of no more than 100 μl 1 min before addition of an aggregating agent. Prostacyclin was prepared as a stock solution in 100% ethanol and diluted in phosphate-buffered saline just before use. Concentrations of 1-10 ng/ml of prostacyclin blocked all of the aggregating agents used.

**Prostacyclin formation.** Replicate cultures of endothelial cells were examined for prostacyclin formation after stimulation by a change of medium and agitation on a rocker platform. Monolayers of cells were treated with fresh culture medium (protein-free) that contained either tranlycypromine (100 μg/ml) or indomethacin (1 μg/ml). The flasks were placed on a rocker platform in the incubator and agitated gently for 2-30 min. Samples of media were taken immediately after the medium change and at 2, 5, 10, 15, and 30 min. Sodium carbonate (0.1 M) was added to each aliquot to bring the pH above 9, and all samples were kept on ice until tested for inhibition of aggregation. For comparison of prostacyclin production by different cultures of cells, replicate monolayers were treated with fresh culture medium or with medium that contained tranlycypromine or indomethacin. After a brief incubation (5 min) this media was removed and 1 ml of protein-free culture medium was added to each flask. The cells were detached immediately by scraping with a rubber spatula and transferred to centrifuge tubes containing 0.1 M Na₂CO₃. The cells were agitated with a vortex mixer, centrifuged to a pellet, and the supernatant medium was tested for inhibition of aggregation. In some experiments PGH₂ (1 μg/ml) was added immediately before scraping.

**Thin-layer radiochromatography.** The production of prostacyclin from arachidonic acid in cultured endothelial cells was established by the formation of 6-keto PGF₁α (30, 32). This metabolite was identified after thin-layer chromatography of radiolabeled samples on silica gel thin-layer plates according to the method of Baenziger et al. (32). Monolayers of endothelial cells were washed with protein-free medium and treated with 5 μM [14C]arachidonic acid in medium adjusted to pH 8.6. The cultures were incubated at 37°C for 30 min with the labeled precursor, then the medium was removed, the cells were scraped as described above, and centrifuged to a pellet. The supernatant medium was acidified to pH 4 with glacial acetic acid and 50 μg of PGE₂, PGF₂α, and 6-keto PGF₁α were added. The medium was extracted twice with 10 volumes of ethyl acetate, dried under N₂, and the residue was dissolved in a small volume of chloroform: methanol (2:1). Each sample was spotted individually on a 5 x 20-cm silica gel plate and chromatographed in a solvent system of ethyl acetate:acetic acid:trimethylpentane (110:20:50). Standards of PGE₂, PGF₂α, and 6-keto PGF₁α were chromatographed on a separate plate. Separation of the prostaglandins was improved by drying each plate and running it a second time in the same solvent. At the end of the second chromatography, the standard plates were sprayed with 3.5% phosphonomylbic acid to develop the spots corresponding to the prostaglandins. The radiolabeled test plates were scanned in a Packard chromatograph scanner (Packard Instrument Co., Inc., Downers Grove, Ill.) and the radioactive peaks were compared with the spots developed on the standard plates.

**RESULTS**

**Cell cultures.** Endothelial cells from all of the vessels examined had a similar appearance under phase microscopy. All cultures started from small clumps of cells and spread into monolayers within ~2-3 wk. Cells from adult vessels grew slowly at first and cells divided only at the edge of each clump. After 10-14 d in culture, however, cell division was more rapid, and transferred cells became confluent within a matter of days instead of weeks. Umbilical cord cells grew at a uniformly slow rate in both initial and transferred cultures. The appearance of endothelial cells from pulmonary artery is shown in Fig. 1.

**Factor VIII antigen.** Cells from all of the vessels examined had Factor VIII antigen as determined by combination with fluorescent antibody. No differences
in the amount of material in cells from different vessels could be detected by this method. All of the cultures treated with antibody to Factor VIII and then with FITC-labeled anti-gamma globulin developed an intense yellow fluorescence. In contrast, cells treated with saline or with normal rabbit serum and FITC-antibody had only pale green background fluorescence. Fig. 2 shows the labeling of pulmonary and umbilical arterial cells with fluorescent antibody. Factor VIII antigen was present in all cultures examined between the first and fifth passages, and in four out of six cultures examined as late as the seventh passage. In two of the older cultures staining was diminished, and the cultures contained large, fibrillar cells that resembled smooth muscle cells. However, all cultures of pulmonary, umbilical, or ovarian cells that retained their epitheliod shape were positive for Factor VIII antigen.

Enzymatic activities. Angiotensin I converting enzyme (kininase II) activity was present in all cultures of endothelial cells studied. To establish that this marker was retained over the time of the enzymatic studies, three different cultures (one pulmonary vein and two pulmonary artery cultures) were tested at different passages. Fig. 3 shows the activity of the angiotensin I converting enzyme as measured by the hydrolysis of [3H]hippuryl glycyglycine. Although the activity was higher in the two arterial cultures than in the venous one, it remained constant for at least five passages. In one arterial culture a decrease in the enzymatic activity between the fifth and seventh passages was associated with a change in the morphology of the cells. These cells became larger and assumed an overlapping, interwoven appearance similar to that of smooth muscle cells. This particular culture was also negative for Factor VIII antigen at the seventh passage.

The activity of angiotensin I converting enzyme in cells from arteries and veins was compared with two different substrates. Only cells in the first four passages were used for the enzyme determinations. With either bradykinin or [3H]hippuryl glycyglycine, the activity in cells from arteries was higher than in cells from veins. Fig. 4 shows the activity of angiotensin I converting enzyme measured by hydrolysis of [3H]hippuryl glycyglycine in cells from aorta, pulmonary, ovarian, and umbilical vessels. Table I compares the inactivation of bradykinin by cells from four different pulmonary arteries and two different pulmonary veins. With either substrate the enzyme activity was inhibited more than 90% by SQ 20881.

Angiotensinase activity was also higher in arterial cells than in venous cells. Table II shows the activity in pulmonary and umbilical arterial cells compared to the activity in umbilical vein cells. With either α-L-aspartyl β-naphthylamide or angiotensin II as a substrate, the arterial cells were four to five times more active than venous cells. The inhibition of the enzymatic activity by o-phenanthroline indicates that the enzyme is probably angiotensinase A (15).

Inhibition of platelet aggregation. Media from monolayers of endothelial cells collected after agitation

FIGURE 1  Pulmonary endothelial cells in culture. A shows a monolayer of cells viewed with Nomarski interference phase microscopy. B shows cells viewed with conventional phase microscopy. ×80.
on a rocker platform inhibited the aggregation of platelets caused by thrombin, ADP, sodium arachidonate, and the endoperoxide analog, U-41669. This inhibitory activity was generated within 2–5 min after the cells were stimulated by a change of medium and agitation, but it was not formed by cultures that were treated with either indomethacin or tranylcypromine (Fig. 5). The inhibitory activity formed by control (untreated) cultures was lost if the samples were acidified, or if they were heated to 100°C for 2 min.

Scraping the cells with a rubber spatula provided a more vigorous stimulation for prostacyclin formation. Scraping and vigorous agitation on a vortex mixer increased the amount of prostacyclin to more than 50 times that formed by intact cell monolayers. The amount of prostacyclin formed by scraped cells increased still further if the endoperoxide, PGH₂, was added just before scraping. Indomethacin and tranylcypromine blocked prostacyclin formation in scraped cells, but addition of PGH₂ reversed the inhibition only in indomethacin-treated cells (Table III).

To compare prostacyclin formation by different cultures, duplicate flasks of cells were scraped and tested for inhibition of aggregation under standard conditions. The media were removed by suction, and 1 ml of protein-free culture medium (pH 8.0) was added to each flask. The cells were scraped, centrifuged to a pellet and the supernatant medium was tested immediately in the platelet aggregation bioassay. An estimate of the amount of prostacyclin was made by comparison with standard solutions of prostacyclin. In three separate experiments pulmonary arterial cells formed more prostacyclin than either pulmonary or umbilical venous cells. These data are given in Table IV.

Thin-layer radiochromatography. To establish that the platelet-inhibitory activity from endothelial cells was prostacyclin, samples of media from scraped cells were chromatographed on silica gel plates and compared with several prostaglandin standards. The cell monolayers were treated first with labeled arachidonic acid and the labeled products identified by comigration.
with standard prostaglandins. Fig. 6 shows a scan of media from control cells and cells treated with indomethacin in juxtaposition to a thin-layer plate stained to show the migration of PGE₂, PGF₂α, and 6-keto PGF₁α. In four experiments (two with pulmonary arterial cells and two with ovarian arterial cells) the major peak of radioactivity co-migrated with 6-keto PGF₁α. As shown in the figure, a small peak corresponding to PGF₂α followed the major peak. This product was eliminated by treating the cells with indomethacin before arachidonic acid stimulation, and the major 6-keto PGF₁α peak was greatly diminished.

**DISCUSSION**

Endothelial cells cultured from various human vessels have similar characteristics. They are epithelioid in shape, they contain Factor VIII antigen, angiotensin I converting enzyme, and angiotensinase a, and they can form prostacyclin in response to mechanical stimulation.

Cells cultured from aorta, pulmonary, ovarian, or umbilical vessels retain angiotensin I converting enzyme and Factor VIII antigen for five or more passages. Because smooth muscle cells have neither converting enzyme activity (15) nor Factor VIII antigen (12, 40), the retention of these markers by the cultured cells used in this study indicates that they are endothelial cells. In one culture that was followed for multiple passages a decline in converting enzyme activity and Factor VIII antigen reactivity was associated with a change in morphology of the cells consistent with an overgrowth of smooth muscle cells. While a minor contamination with smooth muscle cells in the endothelial cultures cannot be excluded, the cells used in this study retained their endothelial characteristics throughout the time they were studied. Because smooth muscle cells exhibit a distinctive morphology (14, 24, 26), a culture that was prominently contaminated could be easily identified.

Although the collection of the cells and the conditions of culture were the same for all of the cells studied, there was a striking difference in the enzymatic activities of cells derived from different vessels. Both angiotensin I converting enzyme and angiotensinase A were three to five times more active in cells from arteries than in cells from veins. This pattern persisted even though two different substrates were used.

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**TABLE I**

Kininase Activity in Human Pulmonary Endothelial Cells

<table>
<thead>
<tr>
<th>Cell origin</th>
<th>Treatment</th>
<th>Enzyme activity (nmol/h/10⁶ cells)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pulmonary artery</td>
<td>None</td>
<td>64.3±9.1</td>
</tr>
<tr>
<td></td>
<td>SQ 20881</td>
<td>3.2±0.1</td>
</tr>
<tr>
<td>Pulmonary vein</td>
<td>None</td>
<td>14.4, 20.6</td>
</tr>
<tr>
<td></td>
<td>SQ 20881</td>
<td>1.0, 1.2</td>
</tr>
</tbody>
</table>

Inactivation of bradykinin (2 μM) was measured on the rat uterus in vitro. Data for pulmonary arterial cells are mean±SEM of four experiments and data for pulmonary vein cells are individual values from two separate experiments.

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**FIGURE 3** Angiotensin I converting enzyme in human pulmonary endothelial cells at different passages. Two cultures of pulmonary arterial cells and one of venous cells were examined after each passage. Enzyme activity, as measured by the hydrolysis of [³H]hippuryl glycylglycine, is indicated by the height of the bars. The number of each passage is on the abscissa.

**FIGURE 4** Activity of angiotensin I converting enzyme in cells cultured from different human vessels. Stippled bars indicate activity in arterial cells and hatched bars indicate activity in venous cells. All values are mean±SEM of activities from 4 to 12 separate cultures. OV, ovarian cells; UMB, umbilical cord cells; and PULM, pulmonary cells. Enzyme activity was measured by hydrolysis of [³H]hippuryl glycylglycine.
to measure each of the enzymes. Since the activity of angiotensin I converting enzyme remained constant for at least five passages, and the cells were compared within the first three passages, it is unlikely that dedifferentiation can account for the differences in enzymatic activities.

Other investigators found that angiotensin I converting enzyme activity is affected by changes in serum concentration or oxygen tension. Hayes et al. (29) noted that cultures of bovine aortic endothelial cells that were maintained in serum-free medium for several days had twice the activity of cells that were grown in media with 20% fetal calf serum. Leuenberger and associates (41) measured the conversion of angiotensin I to II in the pulmonary vascular beds of dogs and found that hypoxia reduced conversion. Stalcup et al. (42) reported that cultures of human umbilical vein endothelial cells had reduced converting enzyme activity after equilibration with hypoxic gas mixtures. However, since all of the cells in this study were grown under identical conditions, it is unlikely that changes in serum or oxygen can account for differences in enzymatic activities between arterial and venous cells. Probably the differences reflect the specialization of the vessels of origin and indicate that the cultured cells remain highly differentiated.

Pulmonary endothelial cells appear to be no more active than other endothelial cells in metabolizing bradykinin and angiotensin. Although the lung is assumed to be a major site for conversion of angiotensin I (11), only 20–40% of injected angiotensin I is converted during pulmonary passage in normal human subjects (9, 43). The remainder is converted in extra-pulmonary beds. Angiotensin II is more readily inactivated in peripheral circulation than in the pulmonary circulation (9–11), thus there appear to be regional differences in the metabolism of vasoactive peptides in vivo. Whereas these differences were not apparent in the cultures of cells from different parts of the body, the greater enzymatic activity associated with cells from arteries may reflect one means of controlling

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**TABLE II
Angiotensinase Activity in Endothelial Cells**

<table>
<thead>
<tr>
<th>Cell origin</th>
<th>Substrate</th>
<th>Inhibitor</th>
<th>Enzyme activity (nmol/h/10^6 cells)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Umbilical vein</td>
<td>α-L-aspartyl β-naphthylamide</td>
<td>None</td>
<td>0.5±0.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>o-phenanthroline</td>
<td>0.1±0.0</td>
</tr>
<tr>
<td>Umbilical artery</td>
<td>α-L-aspartyl β-naphthylamide</td>
<td>None</td>
<td>2.7±0.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>o-phenanthroline</td>
<td>0.7±0.0</td>
</tr>
<tr>
<td>Pulmonary artery</td>
<td>α-L-aspartyl β-naphthylamide</td>
<td>None</td>
<td>1.9±0.2</td>
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<tr>
<td></td>
<td></td>
<td>o-phenanthroline</td>
<td>0.3±0.0</td>
</tr>
<tr>
<td>Umbilical vein</td>
<td>Angiotensin II</td>
<td>None</td>
<td>4.5±0.5</td>
</tr>
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<td></td>
<td></td>
<td>o-phenanthroline</td>
<td>0.2±0.0</td>
</tr>
<tr>
<td>Umbilical artery</td>
<td>Angiotensin II</td>
<td>None</td>
<td>23.0±0.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>o-phenanthroline</td>
<td>4.4±0.1</td>
</tr>
<tr>
<td>Pulmonary artery</td>
<td>Angiotensin II</td>
<td>None</td>
<td>18.2±2.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>o-phenanthroline</td>
<td>0.7±0.2</td>
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</tbody>
</table>

The concentration of α-L-aspartyl β-naphthylamide was 0.7 mM, the concentration of angiotensin II was 14 μM, and the concentration of o-phenanthroline was 1 mM.

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**Figure 5** Formation of prostacyclin in endothelial cell monolayers. Pulmonary arterial endothelial cells were stimulated by agitation on a rocker platform. Aliquots were drawn immediately after a change of medium and at 2, 5, 10, 15, and 30 min of agitation. Each aliquot was tested in duplicate for inhibition of platelet aggregation by thrombin (0.75 u/ml). Closed circles are values for control cells (untreated), open triangles are from cells treated with 100 μg/ml of tranylcypromine, and closed triangles are from cells treated with 1 μg/ml of indomethacin.
TABLE III
Inhibition of Platelet Aggregation by Endothelial Cells

<table>
<thead>
<tr>
<th>Sample origin</th>
<th>Treatment</th>
<th>Volume added</th>
<th>Inhibition of aggregation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>µl</td>
<td>%</td>
<td></td>
</tr>
<tr>
<td>Pulmonary artery (M)None</td>
<td>50</td>
<td>55</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Indomethacin</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Tranylcypromine</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>Pulmonary artery (S)None</td>
<td>1</td>
<td>95</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Indomethacin</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Tranylcypromine</td>
<td>100</td>
<td>10</td>
</tr>
<tr>
<td>Pulmonary artery (S) PGH₂2</td>
<td>2</td>
<td>90</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Indomethacin + PGH₂</td>
<td>2</td>
<td>83</td>
</tr>
<tr>
<td></td>
<td>Tranylcypromine + PGH₂</td>
<td>25</td>
<td>5</td>
</tr>
<tr>
<td>Pulmonary vein (M)None</td>
<td>100</td>
<td>28</td>
<td></td>
</tr>
<tr>
<td>Pulmonary vein (S)None</td>
<td>10</td>
<td>40</td>
<td></td>
</tr>
</tbody>
</table>

Samples were media from cell monolayers (M) or from freshly scraped and suspended cells (S). Drugs were added 5 min before samples were drawn. PGH₂ was added just before cells were scraped, after treatment with drugs.

access of vasoactive peptides into specific vascular beds.

Prostacyclin production by endothelial cells may also depend upon the vessel of origin. Cultured pulmonary endothelial cells, like those from umbilical veins (30, 31) can inhibit aggregation of human platelets by a variety of agents. Several observations indicate that this inhibition is due to formation of prostacyclin. First, there is no inhibition by media from cells treated with either indomethacin or tranylcypromine. Tranylcypromine inhibits prostacyclin synthetase, which forms prostacyclin from the endoperoxide, PGH₂, and indo-

TABLE IV
Prostacyclin Formation by Endothelial Cells

| Sample origin       | Culture No. | Prostacyclin
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>nmol/10⁶ cells</td>
</tr>
<tr>
<td>Pulmonary artery</td>
<td>1</td>
<td>0.55</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>4.30</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>2.68</td>
</tr>
<tr>
<td>Pulmonary vein</td>
<td>1</td>
<td>0.10</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>0.28</td>
</tr>
<tr>
<td>Umbilical cord vein</td>
<td>1</td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>0.08</td>
</tr>
</tbody>
</table>

Samples were supernatant media from scraped and suspended cells. Medium was removed from monolayers and replaced with 1 ml of protein-free M-199 (pH 8.0). The cells were immediately scraped, mixed on a vortex and centrifuged to a pellet. The cell number was determined by assay of DNA in cell pellets. Prostacyclin in the media was estimated by comparison with standard prostacyclin for inhibition of platelet aggregation.

FIGURE 6 Identification of 6-keto PGF₁₀ in media from endothelial cells. Cultures of pulmonary arterial cells were stimulated by scraping with a rubber spatula. The transformation of [¹⁴C]arachidonic acid into prostaglandins was determined by thin-layer chromatography in a solvent system of ethyl acetate:acetic acid:trimethylpentane. Labeled prostaglandins were detected by a radiochromatograph scanner and compared with standards. The arrow indicates the origin on the silica gel plate and the dotted line is the solvent front. Spot 1, PGE₂; spot 2, PGF₂₀; and spot 3, 6-keto PGF₁₀.
methacin inhibits cyclooxygenase, which forms endo-
peroxides from arachidonic acid (44). Pulmonary
endothelial cells treated with indomethacin could still
generate inhibitory activity if they were supplied with
PGH₂, but cells treated with tranylcypromine could
not. Since Marcus and associates (45) showed that cul-
tured endothelial cells can convert PGH₂ to prostacy-
clin via prostacyclin synthetase, it seems likely that the
inhibitory material formed by pulmonary endothelial
cells is prostacyclin.

The inhibitory material is formed within 2–5 min
after endothelial cell monolayers are stimulated by
agitation or scraping. This time-course is similar to that
reported for prostacyclin production of fibroblasts and
smooth muscle cells (46) and umbilical vein endo-
thelial cells (31–33). The ability to inhibit platelet aggrega-
tion is lost, however, if the samples are acidified or
heated to 100°C. Prostacyclin is similarly labile under
these conditions (44).

Finally, radiolabeled 6-keto PGF₁α was identified by
thin-layer chromatography in acidified extracts of endo-
thelial cell media. When cell monolayers were incu-
bated with [³H]arachidonic acid before stimulation by
scraping, the major radioactive product co-chromato-
graphed with 6-keto PGF₁α. Formation of this metabo-
lite was inhibited by indomethacin.

Comparison of media from different endothelial cell
cultures on platelet aggregation suggested that pulmo-

nary arterial endothelial cells form more prostacyclin
than either pulmonary or umbilical venous cells. Weksler et al. (30) noted that endothelial cells cultured
from bovine aorta had approximately 100 times more
inhibitory activity than umbilical cord cells on platelet
aggregation. From the platelet inhibition bioassay, we
estimate that pulmonary arterial cells form between 0.55
and 4.30 nmol of prostacyclin per 10⁹ cells, and pulmo-
nary or umbilical venous cells form between 0.1 and
0.28 nmol per 10⁹ cells. Czervionke and associates (33)
reported that endothelial cells from umbilical veins formed ~0.1 nmol per 10⁹ cells when the monolayers
were treated with thrombin. Thrombin, like scraping,
is a strong stimulus for prostacyclin formation (31).
Although Czervionke et al. (33) used a radioimmuno-
assay for 6-keto PGF₁α, their data for prostacyclin in
umbilical vein cells is similar to that obtained with the
platelet bioassay.

Others found differences in arterial and venous ves-
sels with respect to their ability to form prostacyclin
and 6-keto PGF₁α. Skidgel and Pintz (47) used a radio-
labeled endoperoxide to measure prostacyclin forma-
tion in homogenates of vessels from rats. They found
that whereas all of the vessels that they examined
could metabolize PGH₂ to 6-keto PGF₁α, arteries had
consistently more activity than veins.

Several conclusions can be drawn from this study of
cultured endothelial cells. First, endothelial cells from
all the different human vessels studied contain two
markers associated with endothelium in vivo, Factor
VIII antigen and angiotensin I converting enzyme.
Second, retention of these markers by passaged cells
indicates that the cells remain differentiated for at
least five passages in culture. Finally, cultured endo-
thelial cells, like endothelium in vivo, can metabolize
vasoactive peptides and form prostacyclin. These func-
tions appear to be influenced by the vascular origin of
the cultured cells.

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