Purification and Partial Sequencing of the Major Mitogen for Human Uterine Smooth Muscle–like Cells in Leiomyoma Extracts

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Abstract

We purified the major mitogen for human smooth muscle–like cells in leiomyoma extracts by sequential liquid chromatography on (a) carboxymethyl-Sepharose, (b) heparin-Sepharose columns, (c) cartridges of C4 silica, and (d) linear gradient reverse-phase high performance liquid chromatography. The mitogenic activity of the leiomyoma extract throughout purification was tested by tritiated thymidine incorporation and DNA content in NIH/3T3 fibroblasts and KW human smooth muscle–like cells. Purification of the leiomyoma–derived growth factor (LDGF) for KW smooth muscle–like cells confirmed that its partial NH2-terminal amino acid (aa) sequence (1–20 aa) was identical to 113–132 aa of the human cysteine-rich protein (hCRP).

A synthetic peptide which was engineered based on the purified aa sequence, stimulated the proliferation and growth of KW cells. An oligonucleotide probe constructed by the cDNA of the hcrp gene that encodes this aa sequence depicted the expression of 1.9-kb LDGF mRNA in leiomyomas and myometrium. The expression of the LDGF mRNA was three to sixfold higher in leiomyomas compared with adjacent myometrium of women harboring leiomyomas by in situ hybridization analysis. These data suggest that LDGF may participate in the pathophysiology of uterine leiomyomas. (J. Clin. Invest. 1995. 96:751–758.) Key words: myometrium • leiomyomas • growth substances • cysteine-rich protein • smooth muscle cells

Introduction

Recently, several biologically active peptides, including (a) metastasis-stimulating factor(s) from mouse uterus (1, 2); (b) rat, rabbit, and bovine uterus–derived growth factors (3); (c) 17-kD heparin-binding growth factor (HBGF-8) partly purified from bovine uterus similar to basic fibroblast growth factor (bFGF) (4); (d) mitogens (s) distinct from EGF in porcine uterus extracts (5, 6); and (e) human uterine muscle–derived and leiomyoma–derived growth factors (DGF)1 have been implicated in uterine physiology and pathophysiology (7–11). Growth factors, such as IGFs, EGF, PDGF, TGF/b, hFGF, and their receptors, were postulated to play an important role in human uterine physiopathology (12–27), including leiomyomas (15, 16, 19, 24–27).

Our previous studies documented that uterine tissues contain mitogens and inhibitors for skin fibroblasts, osteoblasts, myoblasts, and rat aorta smooth muscle–like cells (7–9). Moreover, preferential mitogens for rat and mouse muscle–like cells were extracted from leiomyomas that were undetectable in myometrial and endometrial extracts (10, 11).

Herein we report the final purification and the partial amino acid (aa) sequence of an LDGF with preferential activity in KW human uterine smooth muscle–like cells. This material contained activity equivalent to IGF-1, and its NH2-terminal sequence (1–20 aa) encompassed part (113–132 aa) of the LIM motif (zinc finger) of human cysteine-rich protein (hCRP). Using a synthetic peptide comprising this amino acid sequence and an oligonucleotide probe synthetically engineered by the cDNA sequence that encodes this aa sequence, we confirmed that (a) synthetic peptide is mitogenic for uterine smooth muscle cells, and (b) there exists the expression of a discrete signal at 1.9-kb LDGF mRNA in uterine tissues. The expression of the LDGF mRNA was remarkably higher (three–to sixfold) in leiomyomas than in the adjacent myometrium of women harboring leiomyomas as detected by in situ hybridization analysis. These data suggest that this LDGF is possibly linked with the pathophysiology of uterine leiomyomas.

Methods

Source, handling of tissues, and extraction method. Uterine tissues were obtained at hysterectomy of premenopausal women harboring leiomyomas; these women had received no therapy before surgery. Leiomyoma tissues were carefully washed with ice-cold and serum-free HBSS and either extracted or stored at −80°C without preservatives until use. Tissues were minced and extracted in 40 ml/gram of 50 mM of ammonium acetate, pH 5.5, containing 50 mg/liter PMSF, 10 mg/liter soybean trypsin inhibitor, and 1 mg/liter aprotinin for 2 h at 4°C with agitation (10, 11). The extract was centrifuged at 10,000 g for 30 min and filtered with 0.45-μM filters. The extract was dried and redissolved in DME/ F-12, and aliquots were tested for bioactivity and protein content (7, 10, 11). This study was approved by the local ethics committee (Laval University, Ste. foy, Canada) according to the guidelines of Medical

1. Abbreviations used in this paper: aa, amino acid; ACN, acetonitrile; CM–, carboxymethyl-Sepharose chromatography; ELP, embryonal long terminal repeat–binding protein; hCRP, human cysteine-rich protein; HS–, heparin-Sepharose chromatography; LDGF, leiomyoma–derived growth factor; HPLC, reverse phase HPLC; SF–1, steroidogenic factor 1; TFA, trifluoroacetic acid.
Research Council (MRC) of Canada for the use of human tissues in medical research.

Bioassay systems—mitogenic activity. The mouse NIH/3T3 fibroblasts were from the American Type Culture Collection (RL1658; ATCC, Bethesda MD), and the KW human uterine smooth muscle–like cells were spontaneously propagated in a cell line after continuous subcultures of fast-growing cells isolated by enzymatic digestion from normal human uterine muscle (28). In our laboratory, the KW cell line was characterized to express phenotype compatible with human smooth muscle cells, including the expression of smooth muscle cell–specific alpha actin (29).

These cells were used as indicator bioassay systems to assess the mitogenic activity of leiomyoma fractions throughout purification. Cell lines were maintained in DME/F-12 containing 10% calf serum in 75-cm² tissue culture flasks. When the cultures reached confluence, the cells were trypsinized using a mixture of 0.1% trypsin (Sigma Chemical Co.; St. Louis, MO) and 0.01% EDTA (Sigma Chemical Co.) in HBBS and precipitated by low-speed centrifugation. Then the cells were counted in a hemocytometer using trypsin blue exclusion to assess viability and plated at 5 × 10^4 cells/well in 24-well plates using DME/F-12 medium containing 10% calf serum. After 24 h, the cells were changed to serum-free medium for an additional 24 h and then stimulated with aliquots of the fractions under investigation (7–11, 29).

The purity of the tritiated thymidine incorporation was assessed by adding 2 μCi of tritiated thymidine (sp act; 20 Ci-mmol; ICN Radiochemicals, Boston, MA) to each well. The medium was aspirated and the cells washed twice with ice-cold PBS. Then 1 ml of ice-cold trichloroacetic acid (10% TCA) was added. The TCA-insoluble material was dissolved in 0.5 ml of 0.6 N NaOH, and radioactivity (cpm/μg DNA) was measured by liquid scintillation spectrometry (7–11). The DNA content was also evaluated (μg DNA/well) in monolayer cultures as previously described (7–11, 29). Mitogenic activity throughout purification was analyzed as units per microgram of protein; 1 U was defined as the stimulation in tritiated thymidine incorporation and DNA content (percent above control) achieved by 10 ng/ml of IGF-I in the KW smooth muscle–like cells and NIH/3T3 fibroblasts. The cell number was counted by a hemocytometer using the trypsin blue exclusion method in cell cultures treated with mitogens for 4 consecutive days (7–11, 29).

The results were expressed as percent above controls = tritiated thymidine incorporation (cpm/μg DNA), DNA content (μg DNA/well), and number of cells in cell cultures incubated with mitogens minus tritiated thymidine incorporation, DNA content, and number of cells in cell cultures without mitogens (controls), divided by the latter × 100. Results were presented as mean value and standard error (X±SE) in triplicate determinations. Statistical evaluation was by ANOVA (Dunnett’s test).

Carboxymethyl (CM)-Sepharose chromatography (CM-c). CM-c was performed on a 12 × 5.8 cm column preequilibrated with 50 mM ammonium acetate, pH 5.5, at 4°C. The flow rate was 200 ml/h. Leiomysoma extracts were first loaded on the column and then washed with 50 mM ammonium acetate, pH 5.5. CM-c was eluted by 1.2 M NaCl (one-step elution) as previously described (11).

Heparin-Sepharose chromatography (HS-c). HS-c was performed on a 1.6 × 18 cm column preequilibrated with 20 mM Tris-HCl containing 0.6 M NaCl and adjusted to pH 7.1 using 1 M Tris-HCl. The flow rate was 50 ml/h. The protein fractions eluted from CM-c were diluted using Tris 20 mM, pH 7.1, to 0.6 M NaCl and adjusted to pH 7.1 using 1 M Tris-HCl. These fractions were then loaded onto HS-c and eluted by 2.0 M NaCl (one-step elution). Eluates were analyzed by SDS-PAGE as previously described (30).

Fractionation over cartridges of C4 silica. Fractions eluted from the HS-c were diluted with 1 vol of 0.1% trifluoroacetic acid (TFA) and then loaded onto cartridges of octadecylsilyle (ODS) silica (C4 Sep-Pak; Waters Associates, Mississauga, Canada) preactivated by acetonitrile (ACN) (80% ACN/0.1% TFA). As a result, three solutions were generated: (a) pass through containing proteins not retained on cartridges of C4 silica; (b) wash containing proteins and salts washed out by the 0.1% TFA; and (c) elute containing the proteins retained on Sep-Pak and eluted by 80% ACN in 0.1% TFA (31). The eluted proteins were analyzed by SDS-PAGE and tested for mitogenic activity.

Reverse-phase HPLC (rHPLC). The material eluted from cartridges of C4 silica was redissolved in 0.1% TFA and further purified using linear gradient elution (linear gradient, 12–60% ACN) (see Fig. 4). The HPLC system was composed of two pumps (model 510, Waters Associates), an automatic gradient controller, and an LKB 2221 recording integrator. Eluates were monitored by ultraviolet absorbance at 214 nm using a wavelength flow-through spectrophotometer (HPLC System model 441; Waters Associates) and HPLC was performed on C4 μBondapak column (3.9 × 300 mm; Waters Associates) at a flow rate of 1.0 ml/min using linear gradients of acetonitrile containing 0.1 TFA as a counter ion as previously described (31, 32). The eluted fractions were collected using a Redirac LKB fraction (Pharmacia, Montreal, Canada) collector (3 min collection = 3 ml), and aliquots were tested for mitogenic activity in indicator bioassay systems. The eluted material was analyzed by SDS-PAGE electrophoresis.

Amino acid sequence analysis and peptide synthesis. Sequence analysis was performed at Clinical Research Institute of Montreal, Montreal, Canada, using a gas-phase sequenator (model 470A; Applied Biosystems, Inc., Foster City, CA) directly coupled to a phenylthiohydantoin (PTH) analyzer (model 120A; Applied Biosystems, Inc.) as previously described (33). The PTH–amino acid yield for each cycle was normalized according to a PTH–Norleucine standard added directly before separation; the initial and repetitive yields were obtained by linear regression from the yields of selected stable PTH derivatives (33). The synthetic peptides were engineered by the solid phase method at Sheldon Biotechnology Center, McGill University, Montreal, Canada. These synthetic peptides were purified by rHPLC and characterized by amino acid analysis and mass spectra analysis.

Extraction of total RNA and Northern analysis. Leiomyomal and myometrial total RNA was extracted from tissues obtained at surgery from premenopausal women harboring leiomyomas; these women had received no therapy before hysterectomy. From each hysterectomy, we extracted leiomyomal and myometrial total RNA using an acid guanidinium thiocyanate, phenol, and chloroform extraction procedure (34). The samples of total RNA (10 μg) were electrophoresed through formaldehyde-agarose slab gels, transferred to GeneScreen Plus, and hybridized overnight with 10⁶ cpm/ml probe at 42°C in a buffer containing 5 × SSPE (750 mM NaCl, 50 mM NaH₂PO₄, pH 7.6, 5.0 mM Na₂EDTA), 50% formamide, 5 × Denhardt’s, and 2% SDS. Membranes were washed in 2 × SSPE, 2% SDS at 65°C for 30 min. Membranes were autoradiographed at −70°C with Fisher Biotech intensifying screens (Fisher, Pittsburgh, PA) for various times, depending on the signal intensities. Quantification of the band intensities was carried out using a PhosphorImager, and data were analyzed using ImageQuant Software v3.0 (Molecular Dynamics, Inc., Sunnyvale, CA). The probes were stripped from the membranes by heating to 95°C in 10 mM Tris, pH 7.5, 1% SDS for 20–30 min after rebhybridization with another probe. The oligonucleotide probe (S–CACCTTCGCGCAGCATAGACTGC–CTGGGTGCACTCGGGGCGACCGCTTGGAAACACCATCAGCTG) was synthetically engineered based on the nucleotide sequence of the hcrp gene, which encodes the 113–132 aa (35). Labeling was by the random primer method using the kit from Boehringer Mannheim Biochemicals (Indianapolis, IN), and GeneScreen Plus membranes from Dupont (Montreal, Canada).

In situ hybridization. We decided to analyze leiomyomal and myometrial tissues from four women who underwent hysterectomy while in the proliferative phase (phase 1 of the menstrual cycle). The phase of menstrual cycle was confirmed by the endometrial histology. The leiomyomal and myometrial tissues were embedded in 10.24% (wt/wt) polyvinyl alcohol, 4.26% (wt/wt) polyethylene glycol, and 85.5% (wt/wt) nonreactive ingredient-embedding medium for frozen tissue specimens (Miles Inc., Elkhart, IN) and mounted on a cryostat. Serial sections were cut (10 mm thick) at −16°C, collected on poly-L-lysine–
coated slides, and stored at −80°C. Tissue sections were fixed for 20 min in 4% paraformaldehyde (wt/vol) at 4°C and then washed in 0.1 M phosphate buffer for 20 min (4 × 5 min each). Before hybridization the sections were washed first in 2 × SSC solution (0.3 M sodium chloride and 0.03 M sodium citrate) and then in a 2 × SSC solution containing 0.1% Triton X-100 for 10 and 20 min, respectively. Prehybridization was performed in a buffer solution containing 50% (vol/vol) formamide, 5 × SSPE (20 × SSPE = 0.18 M NaCl, 10 mM NaH2PO4, 1 mM EDTA, pH 7.4, 0.1% SDS, 0.1% [wt/vol] polyvinylpyrrolidone), 200 μg/ml denatured salmon testis DNA, 2 μg/ml poly(A)+ RNA, 4% [wt/vol] dextran sulfate, and 10 mM DTT) at 39°C for 2 h.

The 32P-labeled oligonucleotide probe was prepared by a typical tailing reaction using terminal deoxynucleotidyltransferase (TdT; 25 U/μl) and [α-32P]-labeled ATP (100 μCi/100 μl; Amersham Corp.) purified on NAP-sephadex G25 DNA grade column (Pharmacia LKB Biotechnology Inc, Piscataway, NJ). Two million copies of this labeled oligonucleotide probe (5′-CACCTTCTCCGACATAGCTGCTGGTGCAATGCGGCGCGGCTCGGAGCCACCAAT-3′, synthetically engineered as described above) was diluted in the prehybridization buffer, and hybridization was performed at 39°C for 16 h. After hybridization the slides were washed at 20°C in (a) 2 × SSC solution for 40 min; (b) 1 × SSC solution for 40 min; and (c) 0.5 × SSC solution × 2 for 40 min each at 20°C and 39°C, respectively. The slides were rinsed with ascending concentrations of ethanol 70, 90, and 100% and air-dried. The slides were exposed to film (Eastman Kodak Co., Rochester, NY) for 7 d, and the intensity of the signals was analyzed as described above (Northern analysis). The sections were coated with a Kodak NTB-2 liquid photographic emulsion diluted 1:1 with distilled water at 45°C and stored in darkness at 4°C. After 5 and 7 d of exposure, the slides were developed and stained with hematoxylin and eosin (H&E). The slides used as negative controls (RNase) were pretreated with ribonuclease A (10 mg/ml) and ribonuclease T (100,000 U/ml) in 2 × SSC for 45 min at 39°C. Signal intensity (OD/mm2) was evaluated by subtracting the RNase signal from each individual signal measurement in leiomyoma (L) and myometrium (M) tissues. The results were expressed as ratio of L/M signal intensity in tissues from each hysterectomy (relative expression of hCRP mRNA).

Table 1. Purification of the Leiomyma-derived Mitogen for Human KW Uterine Smooth Muscle-like Cells

<table>
<thead>
<tr>
<th>Purification step</th>
<th>Protein content (μg)</th>
<th>Activity (U/μg·fold)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude extract</td>
<td>34.39 × 10^6</td>
<td>0.004</td>
</tr>
<tr>
<td>CM-c</td>
<td>4.64 × 10^6</td>
<td>0.02</td>
</tr>
<tr>
<td>HS-c</td>
<td>14.16 × 10^3</td>
<td>0.3</td>
</tr>
<tr>
<td>Sep-PAK</td>
<td>683</td>
<td>6.25</td>
</tr>
<tr>
<td>rHPLC</td>
<td>20</td>
<td>124,000</td>
</tr>
</tbody>
</table>

* 1 U of mitogenic activity = the stimulation of 10 ng/ml of IGF-I in the DNA synthesis of KW smooth muscle-like cells expressed as percent above control (see Methods).

Results

Leiomyoma tissue (wet weight, 2.56 kg) was extracted as described in Methods. Extracts contained 34.39 grams of total protein; aliquots of 50–250 μg/ml stimulated tritiated thymidine incorporation and DNA content in KW smooth muscle cells and NIH/3T3 fibroblasts in a dose-dependent manner, confirming our previous findings (7–11). Leiomyma extracts contained higher activity in fibroblasts than in smooth muscle cells. The leiomyma extract contained 0.004 U/μg of activity in KW smooth muscle–like cells and 0.02 U/μg in NIH/3T3 cells (Fig. 1 A).

The leiomyma extract was further purified onto CM-c as described in Methods. The eluted proteins from the CM-c also contained mitogens for both the fibroblasts and smooth muscle–like cells. The CM-c eluates contained 0.02 U/μg of activity in KW cells and 0.008 U/μg in NIH/3T3 fibroblasts, indicating that CM-c purified by fivefold the leiomyomoral mitogen(s) in KW cells. In contrast, the CM-c eluates contained less activity in NIH/3T3 fibroblasts compared with crude leiomyma extracts (Fig. 1 B). These data suggest that important mitogen(s) for fibroblasts possibly passed through CM-c under these experimental conditions. Indeed, compared with the eluted fractions from CM-c, the pass-through fractions contained twice as much mitogenic activity (0.017 U/μg) in fibroblasts.

The CM-c eluates were diluted and rechromatographed on HS-c as described in Methods. The fractions eluted from heparin-Sepharose contained 0.3 U/μg mitogenic activity in KW cells and 0.2 U/μg in NIH/3T3 fibroblasts (Fig. 1 C), ~75-fold purification in the KW cell-specific activity (Table 1).

The HS-c eluates were fractionated using cartridges of C18 silica as described in Methods. Eluted material contained 6.25 U/μg mitogenic activity in KW smooth muscle–like cells and 0.28 U/μg in NIH/3T3 fibroblasts (Fig. 1 D). This step contributed to 1,562.5-fold purification of the mitogen(s) in smooth muscle–like cells and 14-fold purification in NIH/3T3 cells, suggesting purification of preferential mitogen(s) for human smooth muscle cells (8, 10, 11).

We further purified the smooth muscle cell–specific activity using linear gradient HPLC (Fig. 2). Aliquots containing 75 ng/ml of the peptide(s) eluted by 35% of ACN in a linear HPLC gradient (28–44% ACN in 0.1% TFA within 40 min) equivalently stimulated KW smooth muscle–like cells with 100 ng/ml of IGF-I (Fig. 2). This material was less active in NIH/
laid tritiated thymidine incorporation and DNA content of KW cells in a dose-dependent manner (Fig. 3). Evaluating the mitogenic activity as units per microgram, we calculated overall purification by 31,000-fold (Table I). By SDS-PAGE, we analyzed the proteins eluted by the HS-c, cartridges of C₁₈ silica, and the linear rHPLC gradient; the latter (LDGF) was a 17- to 18-kD protein (Fig. 4).

Sequence analysis confirmed final purification revealing a partial NH₂-terminal aa sequence of the putative LDGF (1–20 aa) identical to 113–132 aa of the hCRP, encompassing part of the second zinc-finger “LIM motif”, defined by C₆₋₇₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋₋_-\( \text{Fig. 4. Analysis of purified proteins by CM-c (lane A), HS-c (lane B), Sep-Pak (lane C), and rHPLC (lane D) as detected by SDS-PAGE electrophoresis using Commassie blue staining. Lane D presents similar analysis by silver staining.} \)

To investigate whether the mitogenic activity of purified extracts is linked to the obtained aa sequence, we synthetically engineered three peptides encompassing (a) peptide A, part of the NH₂-terminal of hCRP (1–10 aa); (b) peptide B, the obtained aa sequence identical to 113–137 aa sequence of hCRP, and (c) peptide C, part of the COOH-terminal sequence of hCRP (170–193 aa). These peptides were tested for mitogenic activity by analyzing the [³H]thymidine incorporation, DNA content, and cell number in KW smooth muscle-like cells. Peptide B stimulated only the proliferation and growth of KW cells (Table III). The mitogenicity of the peptide B was comparable to that of purified leiomyoma extracts, thus definitely confirming that this aa sequence was a link to the putative LDGF in leiomyoma extracts.

To assess the corresponding size of the LDGF mRNA in uterine tissues, we extracted the total mRNA of leiomyoma and myometrium, which was further analyzed by Northern blots. For this analysis, we used the radiolabeled oligonucleotide probe synthetically engineered as described in Methods. This probe was engineered based on the cDNA sequence that encodes the NH₂-terminal aa sequence of the LDGF. Northern analysis documented the presence of a discrete signal band of 1.9-kb LDGF mRNA (Fig. 5), which corresponded also to the expected size of hCRP mRNA (35).

Analysis of the relative expression of the LDGF mRNA depicted by this oligonucleotide probe in uterine tissues (n = 4; leiomyoma vs adjacent myometrium of the same uterus) documented a three- to sixfold higher mRNA expression in leiomyoma (L) than in myometrium (M) by in situ hybridization (L/M: A = 6-fold; B = 6-fold; C = 4.5-fold; and D = 3-fold; Figs. 6 and 7).
Table II. The Amino Acid Sequence of the Purified Material (LM) and hCRP;* and the Nucleotide Sequence of the hcrp Gene which Encodes this Peptide Sequence

<table>
<thead>
<tr>
<th>Synthetic peptide</th>
<th>[3H]thymidine incorporation (cpm/μg DNA)</th>
<th>Cell number</th>
<th>hcrp:</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>(ng/ml)</td>
<td></td>
<td>5'------ATT·GGT·GAC·TTG·GAG·GGG·TGC·GGG·GCA·------</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>LM:</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1'-LYS(112)-ILE·GLY·GLY·SER·GLU·ARG·CYS·PRO·ARG·------</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>hCRP:</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1'-LYS(112)-ILE·GLY·GLY·SER·GLU·ARG·CYS·PRO·ARG·------</td>
</tr>
</tbody>
</table>

* The zinc-finger region "LIM motif" (C,H,XX-C,C) of the hCRP amino acid sequence is underlined. Cysteine cannot be detected by sequence analysis.

## Discussion

Although the effects of sex steroid hormones on the development, growth, and evolution of uterine leiomyomas are unquestionable, the mechanism of this action is unclear (36, 37). Recently, the presence of growth substances in normal and neoplastic uterine tissues was documented (1–11, 38), implicating growth substances as mediators of the effects of sex steroid hormones in the physiopathologic aspects of this tissue, including leiomyomas (38 and references therein).

Our previous studies established that (a) myometrial and leiomyomatus extracts contain mitogen(s) of a proteaceous nature (7); (b) cartridges of C₁₈ silica effectively retain these mitogens (8, 9); and (c) mitogens are retained and can be eluted from CM-c and HS-c (10, 11). Herein we report the final purification of the LDGF for human smooth muscle–like cells using a purification scheme that used sequential liquid chromatography on CM-c, HS-c, cartridges of C₁₈ silica, and μBondapak C₁₈ hPLC column. The mitogenic activity of LDGF throughout purification was assessed in the KW smooth muscle–like cells and NIH/3T3 fibroblasts, thus selecting for further purification the preferential activity for human uterine smooth muscle–like cells (KW cells). This method of screening for selective mitogens enabled us to identify protein fractions that were otherwise unidentified (10, 11, 29).

Indeed, we purified a LDGF that stimulated the DNA content and tritiated thymidine incorporation of KW human smooth muscle–like cells. The activity of this LDGF was equivalent to IGF-I. Chemical analysis revealed that the NH₂-terminal amino acid sequence of LDGF (1–20 aa), which was identical to 113–132 aa of the hCRP, a protein sequence predicted by the fortuitous isolation of hcrp cDNA (35). The mRNA of hCRP was identified by screening a placental cDNA library at low stringency conditions with a human prolactin cDNA probe. Despite this, its structure and the predicted protein suggested a lack of evolutionary or functional relationship to human prolactin. Therefore, isolation of hcrp cDNA was based on limited sequence homology (35). hCRP is of unknown function, comprises 23.4 kD, and contains two copies of a domain encompassing an unusual double zinc-finger referred to as LIM motif (35). hCRP has a high degree of similarity to cysteine-rich rat and mouse intestinal protein (CRIPT), a 8.5-kD protein of unknown function. In contrast to hCRP, CRIP contains a single copy of this double zinc finger (C₅-X₇-C₅-X₇-H₅-X₇-C₅-X₇-C₅-X₇-C₅ region) (39). There is now evidence that other proteins contain a similar motif (C₅-X₇-C₅-X₇-H₅-X₇-C₅-X₇-C₅-X₇-C₅) (40, 41, and references therein). It is noteworthy that a similar double zinc-finger motif was

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Table III. Comparable Mitogenic Activity of the Synthetic Peptide Containing the NH₂-terminal Sequence of the Purified LDGF to Purified Leiomyoma Extracts in KW Smooth Muscle-like Cells

<table>
<thead>
<tr>
<th>Synthetic peptide (ng/ml)</th>
<th>[3H]thymidine incorporation (cpm/μg DNA)</th>
<th>Cell number (cpm/well)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.1</td>
<td>12.8*</td>
<td>1</td>
</tr>
<tr>
<td>0.5</td>
<td>27.5*</td>
<td>5</td>
</tr>
<tr>
<td>1.0</td>
<td>35.7*</td>
<td>15.5*</td>
</tr>
<tr>
<td>100</td>
<td>32.1*</td>
<td>19.8*</td>
</tr>
<tr>
<td>1,000</td>
<td>10.2*</td>
<td>12.5*</td>
</tr>
<tr>
<td>Purified extracts (ng/ml)</td>
<td>24.5*</td>
<td>21.8*</td>
</tr>
</tbody>
</table>

Mitogenic activity was expressed as percent above controls as described in Methods: [3H]thymidine incorporation, control = 4,845±859 (cpm/μg DNA); DNA content: controls = 1,639±0.03 (μg DNA/well); and cell number: controls = 7,774±547. * P < 0.001.

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Figure 5. An example of the discrete signal produced by the labeled oligonucleotide probe that detected a 1.9-kb mRNA size in leiomyoma (A) and normal myometrium (B) as grossly assessed by Northern blot using as reference the elution position of 185 mRNA.
expression profile parallel to c-myc. The coordinate regulation of hcrp and c-myc suggests that these genes respond to the same regulatory pathways and may share transcription control features during the G₀ to S transition. Although it is not clear that the shared LIM/double-finger domain dictates a commonality of function for this family of proteins, hcrp was the first to be identified as a primary response gene and thus possesses apparent biologic significance (40).

To definitely link the obtained amino acid sequence with the putative LDGF in leiomyoma extracts and to test the possible relationship with hcrp, we synthetically engineered three peptides comprising (a) part of the NH₂-terminal of hcrp; (b) the purified NH₂-terminal of LDGF; and (c) part of the COOH-terminal of hcrp. Analysis of mitogenic activity documented that the synthetic peptide that corresponded to the NH₂-terminal sequence of purified material stimulated only the proliferation ([³H]thymidine incorporation) and cell growth (DNA content and cell number) of KW smooth muscle–like cells. Therefore, sequence analysis was on the putative LDGF in leiomyoma extracts. Because this synthetic peptide and aliquots of the purified leiomyoma extracts were mitogenic in KW cells grown in serum-free medium or culture medium containing limited calf serum (0.5%), it is conceivable that LDGF is a growth factor enabling smooth muscle cells to enter from the G₀/G₁ phase to the S phase of the cell cycle. This is compatible with the proposed role of products of various early response genes such as c-myc and hcrp (40).

Because the oligonucleotide probe used in our study detected threefold up to sixfold higher LDGF mRNA expression in leiomyomas compared with adjacent myometrium in the proliferative phase of the menstrual cycle, we believe that LDGF is probably the major mitogen for smooth muscle cells in uterine leiomyomas. Moreover, this oligonucleotide probe hybridized with a 1.9-kb mRNA similar to the expected hcrp mRNA size, suggesting that limited proteolysis of hCRP possibly generates the LDGF in leiomyoma tissues. It is noteworthy that the posi-

**Figure 6.** Relative expression of mRNA in leiomyomas versus myometrium as depicted by the labeled oligonucleotide probe using in situ hybridization. Signal density was analyzed as described in the Methods section. Results were expressed as the ratio of leiomyoma/myometrium (L/M) expression of mRNA (relative expression). Leiomyoma tissues contained three to sixfold higher expression than normal myometrium in each individual uterus. The uterine tissues (4) were all from women at the same phase of the menstrual cycle (phase I).

also predicted in the protein sequence of an estrogen-stimulated cDNA isolated from rat C6 glioma cells (rESP1) (42).

The biologic importance of the double zinc-finger motif (hCRP-like) or single-finger (cysteine-rich rat and mouse intestinal protein–like) proteins is still unknown. The zinc finger is suggestive of an ability to bind DNA and RNA, possibly acting as transcriptional factors and is also suggestive of an ability to bind zinc ion and iron-sulfur clusters (35). It is interesting that hCRP is the product of an immediate response gene with an
tion 112–113 (Lys-Ile) of hCRP corresponding to the start of the NH$_2$-terminal sequence of purified LDGF is a potential site for trypsin-like and endopeptidase lysine C--like enzyme action. In this context, the assessment of possible overexpression of hCRP (product of an early response gene leading cells from $G_0$ to $S$ phase of cell cycle) and intracellular processing by endopeptidases generating this LDGF is an attractive field for further investigation.

Against this hypothesis stands the fact that at least theoretically limited proteolysis in position 112–113 aa of hCRP should generate a hCRP fragment no bigger than 9–10 kD (hCRP is 23.4 kD; 113–193 is an 80-aa long sequence). Our data documented that the purified LDGF was 17–18 kD on SDS-PAGE. Therefore, the LDGF may contain only partial homology to hCRP. This homology possibly exists in its NH$_2$-terminal part that contains a LIM motif identical to the second zinc-finger of hCRP. Alternatively, LDGF and hCRP could be generated by the same gene using different promoters or alternative splicing as exemplified by the steroidogenic factor 1 (SF-1), an orphan nuclear receptor and the embryonal long term repeat-binding protein (ELP) (43). ELP possesses 78 additional aa in the NH$_2$-terminal region and 57 different aa in the COOH-terminal region, whereas SF-1 contains 131 additional aa in the COOH-terminal region. There also exists 330 common aa between SF-1 and ELP. It is noteworthy that SF-1 possesses a zinc-finger and stimulates the expression of steroidogenic enzymes (43) as well as is also essential for adrenal and gonadal development (43) and sexual differentiation (44). In contrast, the function and role of the ELP is still unclear (43, 44).

In conclusion, our data partially identified the LDGF for human uterine smooth muscle cells and documented a three- to sixfold higher expression of LDGF mRNA in leiomyoma compared with adjacent myometrium. We conclude that LDGF is probably the major mitogen for smooth muscle cells in uterine leiomyomas.

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