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A Novel Member of the Insulin-like Growth Factor Binding Protein Superfamily in Prostate Cancer

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12a. DISTRIBUTION / AVAILABILITY STATEMENT
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13. ABSTRACT (Maximum 200 Words)
The insulin-like growth factors (IGFs) are potent mitogens for normal and cancerous prostatic cells. The IGFs are found complexed to IGF binding proteins (IGFBPs) which modulate IGF bioactivity, but also may themselves act in an IGF-independent manner. We have recently characterized a series of IGFBP-related proteins (IGFBP-rPs) which share homology with the IGFBPs in the amino-terminus, bind IGFs with low affinity, and regulate cell growth through both IGF-dependent and IGF-independent actions. This grant is directed at the study of IGFBP-rP2 (also known as connective tissue growth factor) as a regulator of normal and malignant prostatic growth. The specific aims are to: 1) analyze IGFBP-rP2 mRNA and protein expression and distribution in normal and malignant prostatic tissues; 2) determine the transcriptional, translational and post-translational regulation of IGFBP-rP2; and 3) determine the mechanisms by which IGFBP-rP2 regulates prostatic growth.
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INTRODUCTION:

The insulin-like growth factors (IGFs) are potent mitogens for normal and cancerous prostatic cells. The IGFs are found complexed to IGF binding proteins (IGFBPs), which modulate IGF bioactivity, but may themselves act in an IGF-independent manner. We have characterized recently a series of IGFBP-related proteins (IGFBP-rPs) which share homology with the IGFBPs in the amino-terminus, bind IGFs with low affinity, and regulate cell growth through both IGF-dependent and IGF-independent actions. This grant is directed at the study of IGFBP-rP2 (more commonly known as connective tissue growth factor) as a regulator of normal and malignant prostatic growth. The specific aims are to: 1) analyze IGFBP-rP2 mRNA and protein expression and distribution in normal and malignant prostatic tissues; 2) determine the transcriptional, translational and post-translational regulation of IGFBP-rP2; and 3) determine the mechanisms by which IGFBP-rP2 regulates prostatic growth.

BODY:

Over the course of this grant, we have succeeded in the following accomplishments related to the approved Statement of Work (references indicated in parentheses):

Task 1: Analyze IGFBP-rP2 mRNA and protein expression and distribution in normal and malignant prostatic tissue (months 1-12):

IGFBP-rP2 mRNA and protein expression were analyzed in a panel of prostate cells, including human prostate epithelial cells (HPEC), immortalized prostate epithelial cells (P69), a tumorigenic prostate subline of P69 (M12), as well as the established prostate cancer cell lines, PC-3, DU145 and LNCaP. Cellular localization of IGFBP-rP2 protein expression in these same cells was also performed, although conclusive data were only obtained to permit a determination that IGFBP-rP2 was secreted and/or cell-associated. Thus, Task 1 was completed as indicated in the grant proposal, with the most important results obtained from this series of investigations published in Reference 2 (Lopez-Bermejo, et al, ENDOCRINOLOGY 141:4072-4080, 2000).

In brief, IGFBP-rP2 was detected in most of the normal and malignant prostatic epithelial cells tested. Of the cancer cells, expression at the mRNA level was most pronounced in M12 and PC3 cells, with low expression in DU145 and LNCaP cells. Moderate expression was detected in non-malignant P69 cells and early passage of HPEC (HPEC4), and, somewhat surprisingly, expression at least equivalent to that observed in M12 and PC3 cells was detected in late passage of HPEC (HPEC9). Protein expression, as determined from immunoblot analysis, corresponded with mRNA expression, with protein readily detected in conditioned media (CM) from HPEC9. Additionally, cell-associated IGFBP-rP2 was immundetected in all cell lines. It is of note that a more quantitative method for detecting IGFBP-rP2, employing ELISA, was developed subsequently in our laboratory, employing the antibodies that we generated (Reference 4; Twigg, et al, ENDOCRINOLOGY 142:1760-1769, 2001). We have concluded, on the
basis of this series of investigations, that expression of IGFBP-rP2 could not be correlated with the degree of tumorigenicty, unlike our observations with the structurally related IGFBP-rP1, whose expression decreased in malignant cells, compared to normal cells (Reference 2; Lopez-Bermejo, et al, ENDOCRINOLOGY 141:4072-4080, 2000). On the other hand, it appears that IGFBP-rP2 expression increases with passaging of HPEC (at least form HPEC4 to HPEC9), perhaps representing a marker of senescence in HPEC. Indeed, we characterized HPEC as a “senescing” prostate epithelial cell, both by growth profile and by increases in the senescence marker, p16\(^{ink4a}\). These findings, together with data on the regulation of IGFBP-rP2 (see Task 2, below) suggest that IGFBP-rP2 may be involved in either growth or senescence of normal and prostate cancer cells. These findings represent the first characterization in the literature of IGFBP-rP2/connective tissue growth factor (CTGF) mRNA and protein expression in normal and malignant prostate cells. All aspects of Task 1 have been completed.

A critical goal of our research has been the development of appropriate reagents for investigation of IGFBP-rP protein levels in various cancers and human biological fluids. We had previously generated specific antisera for IGFBP-rP1/mac25, IGFBP-rP2/CTGF and IGFBP-rP3/novH, which were employed in immunoblotting studies. Efforts then turned to development of highly specific antibodies which could be employed to generate sensitive and specific assays and be used for specific immunostaining studies. We have now succeeded in producing both polyclonal and monoclonal antibodies which can differentiate between IGFBP-rP1 and IGFBP-rP2, and have developed an assay which is capable of quantifying concentrations of IGFBP-rP1/mac25 in human biological fluids (Reference 7; Lopez-Bermejo, et al, JOURNAL OF CLINICAL ENDOCRINOLOGY AND METABOLISM 88:3401-3408, 2003). Employing baculovirus-generated IGFBP-rP1, we generated a panel of 11 monoclonal antibodies; 10/11 were of sufficient sensitivity to identify nanomolar amounts of IGFBP-rP1, and none cross-reacted with the six high-affinity IGFBPs nor with IGFBP-rP2 or -3. Evaluation of the newly developed IGFBP-rP1 immunoassay indicated a detection limit of 0.7 ug/L, a dynamic range of 3.13-100 ug/L, and intra- and inter-assay coefficients of variation of 2.5-6.8 and 3.1-6.4% at 24-85 ug/L. In random human adult sera (n=41), the median IGFBP-rP1 concentration was 21.0 ug/L, and values did not correlate with serum levels of IGF-I, IGF-II or IGFBP-3. The monoclonal anti-IGFBP-rP1 antibodies also readily detected IGFBP-rP1 expression in human tissue sections, with preferential expression of IGFBP-rP1 in the microvascular endothelium associated with tumorigenesis. These findings support a role for the IGFBP-rPs in vascular biology and suggest that they may be involved in the process of neoangiogenesis in malignancy. Development of highly specific and sensitive assays for IGFBP-rP2/CTGF should allow further evaluation of this hypothesis.
In further studies on the expression of IGFBP-rP2 in cancer, its expression was evaluated in 12 sporadic hepatoblastomas (Reference 6; Von Horn, et al, INTERNATIONAL JOURNAL OF MOLECULAR MEDICINE, 9:645-649, 2002). The expression profiles for the IGFBP-rPs were found to be disturbed. In the case of IGFBP-rP2, three tumors had increased gene expression, with one sample having greatly enhanced expression.

Limitations of investigations relevant to Task 1: In addition to generating anti-IGFBP-rP2 polyclonal antibodies, for the studies described above, we also produced monoclonal antibodies, in an effort to enhance specificity. The monoclonal antibodies generated, however, proved upon screening to not have the requisite specificity and sensitivity, and proved to be of less use than the polyclonal antibodies. While polyclonal antibodies were highly useful for western immunoblotting and ELISA studies (see above), attempts to employ these reagents in immunohistochemical analysis of tissue sections proved unsuccessful. In parallel with these efforts, we were successful in generating monoclonal antibodies against recombinant IGFBP-rP1 (see above), and to employ both our polyclonal and monoclonal anti-IGFBP-rP1 antibodies in immunohistochemical analysis of normal and cancerous prostatic tissue (Reference 7; Lopez-Bermejo, et al, JOURNAL OF CLINICAL ENDOCRINOLGY AND METABOLISM 88:3401-3408, 2003.

Task 2: Determine the transcriptional, translational and post-translational regulation of IGFBP-rP2 (months 1-24):

We analyzed the regulation of IGFBP-rP2 expression in prostate cells by employing reagents known to inhibit growth in prostate cancer cells, such as transforming growth factor-β (TGFβ) and retinoic acid (RA), as well as IGF-I and related growth factors (Reference 2; Lopez-Bermejo, et al, ENDOCRINOLOGY 141:4072-4080, 2000). Employing our panel of prostate cell lines, we first demonstrated that HPEC was the most sensitive of all the cells tested to TGFβ1 and IGF-I treatment. At TGFβ1 concentrations as low as 5 ng/ml, proliferation of HPEC was inhibited 70%, compared to untreated control cells, and this inhibition correlated with increased IGFBP-rP2 expression at both the mRNA and protein levels. IGF-I, at concentrations of 100 ng/ml, which stimulates HPEC growth in all cell passage numbers, dramatically inhibited IGFBP-rP2 expression, often by 90%. As a comparison, IGFBP-rP1 expression remained unchanged during these manipulations. RA, like TGFβ1, up-regulated IGFBP-rP2 expression. These results strongly suggest that IGFBP-rP2 plays a role in the regulation of prostate cell proliferation in response to a variety of growth factors. The effects of these growth factors on prostate cancer cells (i.e., stimulation by IGF-I and inhibition by TGFβ1) were more modest, than in normal HPEC. Nevertheless, the observed effects upon IGFBP-rP2 followed the same trends as observed in HPEC.

We were also able to demonstrate that TGFβ1 regulated IGFBP-rP2 expression in human dermal fibroblast cells, as do advanced glycosylation end-products (AGE)-BSA
AGE treatment of primary cultures of nonfetal human dermal fibroblasts in confluent monolayers increased IGFBP-rP2 steady-state mRNA levels in a time- and dose-dependent manner. These findings link AGE and IGFBP-rP2 in profibrotic and proangiogenic roles, a finding of potential relevance to both the complications of diabetes and the growth of human cancers. We further demonstrated that the induction of fibronectin by AGE is partially mediated by the AGE-induced up-regulation of cell-derived IGFBP-rP2/CTGF.

The effect of sodium butyrate (NaB), a potent inhibitor of cancer cell growth, on the regulation of IGFBP-rP2 expression was correlated with the inhibitory effects of NaB in a variety of cancer cell lines (Reference 3; Tsubaki, et al, JOURNAL OF ENDOCRINOLOGY 169:97-110, 2001). NaB specifically up-regulated the expression of IGFBP-3 and IGFBP-rP2, with parallel results observed on Northern and protein blots. NaB was found to uniformly suppress DNA synthesis in both cancerous and non-cancerous cells and up-regulate IGFBP-3 and IGFBP-rP2 mRNA and protein levels. In prostate cancer cell lines, a cell growth inhibitory NaB concentration of 10 nM, which up-regulates, IGFBP-3, also up-regulated IGFBP-rP2 expression in PC-3 cells, but not in LNCaP cells. Proteolysis of IGFBP-rP2 was not found to be a significant factor, either under basal conditions, or following treatment with TGFβ, RA, or IGF-I.

Limitations of investigations relevant to Task 2: Given the modest effects observed with TGFβ1 and RA on prostate cancer cells, studies employing IGFBP-rP2 antisense oligo-deoxynucleotides were not completed, and are deemed by us to be, at this point, a relatively low priority. Other than this, all aspects of Task 2 were completed.

Task 3: Determine the mechanism by which IGFBP-rP2 regulates prostatic growth:

A role for IGFBP-rP2 in regulation of cell growth was first ascertained in fibroblasts, where the addition of recombinant IGFBP-rP2 resulted in increased expression of fibronectin mRNA (Reference 5; Twigg, et al, ENDOCRINOLOGY 143:1260-1269, 2002). The similar addition of exogenous C-terminal FLAG-tagged IGFBP-rP2 to prostate cells did not demonstrate consistent inhibition of cell proliferation, suggesting that either our FLAG-tagged IGFBP-rP2 preparation was less active than untagged IGFBP-rP2, or that, in prostate cancer cells, IGFBP-rP2 may need to act in concert with other regulated factors to promote inhibition of cell growth. On the other hand, we have recently demonstrated that at least one of our IGFBP-rP2 polyclonal antibodies possesses neutralizing properties, and, in our fibroblast model, have shown that the addition of the IgG-purified fraction of the anti-IGFBP-rP2 antibody abrogated AGE-BSA induction of fibronectin mRNA. Use of such antibodies may prove to be an alternative method to the use of antisense ODNs for determining the role of IGFBP-rP2 in growth inhibition.

To address the possibility that our FLAG-tagged IGFBP-rP2 preparation may be less biologically active than non-tagged, heparin-column purified IGFBP-rP2, we generated fresh preparations of both forms of IGFBP-rP2 protein in a baculovirus system, and...
purified the protein either via anti-FLAG-agarose column or heparin columns. Further, we elected to test our preparations in the cell model system of NIH3T3 cells with an overexpressed IGF-I receptor (IGFIR) (see figures 1-6 included in appendix). Since we had previously demonstrated that IGFBP-rP2 had the capability to bind IGFs with low affinity in \textit{in vitro} assays (Kim, \textit{et al}, PNAS 94:12981-12986, 1997), we reasoned that the NIH3T3:IGFIR would permit us to determine whether recombinant IGFBP-rP2 is capable of exerting IGF-dependent biological effects (as proposed in Task 3). Our results demonstrated that: a) IGFBP-rP2, whether FLAG-tagged, non-tagged-heparin column-purified, or anti-IGFBP-rP2 antibody column-purified, appeared to potentiate and sustain IGF-stimulated IGFIR signaling; b) potentiation effects of IGFBP-rP2 required preincubation of IGFBP-rP2 and IGF; IGFBP-rP2 alone had no effect on IGFIR signaling, and preincubation of cells with IGFBP-rP2 prior to the addition of IGFs did not enhance IGFIR signaling; and c) potentiation effects appear to be IGF-I-specific, as no potentiation of insulin, IGF-II or EGF signaling pathways was observed.

Studies are currently in progress to determine if prostate cell lines (M12 and DU145) stably transfected with IGFBP-rP2, will demonstrate alteration of cell growth. Otherwise, Task 3 has been completed.

**KEY RESEARCH ACCOMPLISHMENTS OVER LAST 12 MONTHS:**

1. Characterization of IGFBP-rP2 expression in normal and malignant prostatic tissue
2. Characterization of molecular regulation of IGFBP-rP2 by known pro-apoptotic, profibrotic and proangiogenic factors
3. Characterization of IGFBP-rP expression in hepatoblastoma
4. Development of highly sensitive and specific monoclonal and polyclonal antibodies for IGFBP-rP1 and IGFBP-rP2
5. Development of specific immunoassay for IGFBP-rP1; identification of IGFBP-rP1 in human cancer blood vessels
REPORTABLE OUTCOMES:


CONCLUSIONS:

The low-affinity IGFBP-related proteins appear to have important IGF-independent actions in the regulation of normal and malignant cell growth. Studies from our laboratory over the last three years have demonstrated an important role for these factors in senescence, tumor suppression, fibrosis and angiogenesis. Studies of gene expression and protein synthesis have implicated these factors as biologically relevant to cancer. The development of highly specific and sensitive antibodies should permit further delineation of their roles.

“So what”: The entire GH-IGF axis has been the subject of increased interest from the perspective of longevity and cancer. In addition to the actions of IGFs and IGFBPs under in vitro conditions, a variety of epidemiological studies have served to underscore the relevance of these factors to human biology and tumor suppression. The low-affinity IGFBP-related proteins appear to share with several of the high-affinity IGFBPs, the ability to suppress cancer growth through IGF-independent means. Further studies are clearly warranted in delineating the relative contributions of each component of this complex growth axis.

APPENDICES: attached figures and manuscripts.
Figure 1A

<table>
<thead>
<tr>
<th>Treatment</th>
<th>IRS-1</th>
<th>IGFIR β</th>
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<tbody>
<tr>
<td>IGF-I (7nM)</td>
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</tr>
<tr>
<td>Mac25/rP1* (400nM)</td>
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</tr>
<tr>
<td>IGFBP-3 (14nM)</td>
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<td>+</td>
</tr>
<tr>
<td>CTGF/rP2* (400nM)</td>
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<td>+</td>
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</tbody>
</table>

IB: α-phospho-tyrosine (αPY20)
Figure 1B

H = Heparin purified rP2
A = rP2 antibody column purified rP2
F = Flag-tagged M2 column purified rP2
Figure 2

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<th>Preincubation Time (min.)</th>
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<th>0</th>
<th>0</th>
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<th>30</th>
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<tbody>
<tr>
<td>IGFP-I (2.5nM)</td>
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<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>CTGF / rP2 (15nM)</td>
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<td>−</td>
<td>+</td>
<td>+</td>
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IP: IGF-IRβ
IB: αPY20
IB: IGF-IRβ
Figure 3A & B

<table>
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<th>IGF-I (3.5nM)</th>
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<tr>
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<td>-</td>
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<tr>
<td>+</td>
<td>200</td>
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<td>+</td>
<td>100</td>
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<tr>
<td>+</td>
<td>200</td>
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</table>

**IB:** α PY20

**IB:** p-MAPK

<table>
<thead>
<tr>
<th>IGF-I (3.5nM)</th>
<th>CTGF/rP2 (nM)</th>
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<tbody>
<tr>
<td>-</td>
<td>0.5</td>
</tr>
<tr>
<td>+</td>
<td>2.5</td>
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<tr>
<td>+</td>
<td>5</td>
</tr>
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<td>+</td>
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<tr>
<td>+</td>
<td>15</td>
</tr>
<tr>
<td>+</td>
<td>25</td>
</tr>
</tbody>
</table>

**IP:** IGF IRβ

**IB:** α PY20

**IB:** p-MAPK

**ERK1/2**
Figure 4

IGF-1 (nM)  0  3.5  2.0  1.0  0.5
CTGF/ rP2 (25 nM)  -  -  +  -  +  -  +  -  +
IB:α PY20

Densitometric analysis (fold-increase)

IGF-1  3.5  2.0  1.0
CTGF  -  +  -  +  -  +
Figure 5

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<td><strong>IGF-I + rP2</strong></td>
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</table>

**IP: IGF-IRβ**

**IB: αPY20**

**IB: IGF-IRβ**

**WCL**

**IB: p-MAPK**

ERK1/2
### Figure 6

#### A. NIH3T3:IGFIR

<table>
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Binding Properties of Insulin-Like Growth Factor Binding Protein-3 (IGFBP-3), IGFBP-3 N- and C-Terminal Fragments, and Structurally Related Proteins mac25 and Connective Tissue Growth Factor Measured Using a Biosensor

PETER VORWERK, BLANKE HOHMANN, YOUNGMAN OH, RON G. ROSENFELD, AND RONALD M. SHYMKO

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We measured the binding of IGF-I and IGF-II to recombinant human N-terminal [residues 1-97; recombinant human IGF-binding protein-31-97 (rhIGFBP-31-97)] and C-terminal (residues 98-264; rhIGFBP-398-264) IGFBP-3 fragments and compared it with IGF binding to intact IGFBP-3 using biosensor analysis. Experiments were carried out in different configurations, either with binding protein or fragment immobilized or with IGF immobilized. These experiments showed that IGF-I and IGF-II bind to IGFBP-3 with affinities of 4-5 x 10^6 M^-1 and similar binding kinetics. The affinities of both rhIGFBP-31-97 and rhIGFBP-398-264 for IGFBP proteins were approximately 3 orders of magnitude less than that of full-length IGFBP-3. These results further support the concept that high affinity binding of IGF to IGFBP-binding proteins results from a two-site interaction of IGF with both the N- and C-terminal regions of the binding protein.

Binding of insulin to IGFBP-3 and its N- and C-terminal fragments and of IGF-I and IGF-II to the structurally related proteins mac25 and connective tissue growth factor was also investigated. Weak insulin binding to full-length IGFBP-3 could be demonstrated in a few experiments, but we found that binding of IGF-I, IGF-II, and insulin to mac25 or connective tissue growth factor was below the detection limit of the biosensor instrument. (Endocrinology 143: 1677-1685, 2002)

IGF BINDING PROTEINS (IGFBPs) are important regulators of IGF action that act by modulating IGF binding to its receptors. Initially identified as carriers for IGFs in a variety of biological fluids, their presumed function is to protect IGF peptides from degradation and clearance, increase the half-life of IGFs, and deliver them to appropriate tissue receptors (1). A number of IGFBF fragments have been identified in different biological fluids, and a variety of specific and nonspecific proteases for IGFBPs have been described (2, 3). The concept of IGFBPs as simple carrier proteins has been complicated by the discovery of multiple IGF-independent actions of IGFBPs and the identification of a number of CDNAs encoding proteins that bind IGF with substantially lower affinities than IGFBPs (4). The N-terminal regions of these proteins are structurally homologous to the IGFBPs, with conservation of the cysteine residues and a common N-terminal motif, GGGCCXXC (1). This observation initially suggested an IGFBP superfamily consisting of the classical high affinity IGFBPs and a group of low affinity IGFBP-related proteins, later identified as mac25, CTGF, and others (1, 4).

IGFBP-3 is the major serum IGFBP and transports 70-90% of the circulating IGFs (5). In target cell systems it inhibits IGF actions, but also, under specific conditions, potentiates IGF action or exerts IGF-independent effects (5, 6). Proteolysis of IGFBP-3 was initially demonstrated in human pregnancy serum, in which circulating IGFBP-3 was found primarily in low mol wt forms (7, 8). The proteolytic fragments were shown to bind IGF with lower affinities, thereby increasing availability of IGF to target receptors. Subsequent studies in different biological fluids demonstrated that limited proteolysis is not restricted to IGFBP-3, but also occurs in other IGFBP-species, IGFBP-1 through -5 (2, 3). As for IGFBP-3, the resulting fragments have a decreased affinity for IGFs and, therefore, more easily release IGF to the target receptors. Furthermore, various IGFBP fragments are capable of direct stimulatory or inhibitory action at the target cells (9-13).

All members of the proposed IGFBP superfamily preserve the N-terminal cysteine-rich domain, including the IGFBP motif GGGCCXXC, but vary in the intermediate region and C-terminal domain of the protein. Only the high affinity IGFBPs also share homology in the cysteine-rich C-terminal domain, which has led to the hypothesis that the conserved N-terminal domain contains the main IGF-binding activity, forming, together with the C-terminal conserved region, the high affinity IGF-binding activity in IGFBPs (1, 13-15). With the expression and purification of a variety of recombinant IGFBPs, structurally related proteins mac25 and CTGF, and well defined fragments or mutants, our understanding of structure-function relationships of the members of the pro-
posed IGFBP superfamily has improved (12, 13, 15-25). However, exact measurement of the binding properties of these various proteins, and especially their proteolytic fragments, has proved difficult. Different methods have been used to estimate the binding affinities: conventional binding assays using radiolabeled IGFs, affinity cross-linking with radiolabeled ligands, gel filtration or Western blotting using radiolabeled or biotinylated ligands (12-14, 17, 20, 21, 26-32). However, it has been difficult to determine accurate affinities of IGFs for IGFBPs using those assays due to several limiting factors, such as the quality of labeled ligands and proteins, the effect of labeling iodine on affinity, and the limitation of the measurable range of the binding affinity.

In recent years biosensor instruments using surface plasmon resonance technology have been increasingly used to study biomolecular interactions. We (33-35) and others (15, 36-39) have used this technology to study IGFBP interaction with IGFs or their analogs. In the present study our goal was to make a detailed analysis, using a BIACORE biosensor instrument (Biacore AB, Uppsala, Sweden), of the binding of IGF-I and IGF-II to recombinant IGFBP-3 and fragments of this molecule and also to the structurally related proteins mac25 and CTGF.

**Materials and Methods**

**Equipment and reagents**

Experiments were performed on an upgraded BIACORE 1000 instrument (Biacore AB). CM5 certified sensor chips, surfactant P20, and the amine coupling kit (N-hydroxysuccinimide, N-ethyl-N'-[3-dimethylaminopropyl)carbodiimide, and ethanollamine hydrochloride) were also obtained from Biacore AB. Recombinant human nonglycosylated IGFBP-3 (rhIGFBP-3) produced in *Escherichia coli* was a gift from Protegen (Mountain View, CA). N- and C-terminal recombinant fragments, rhIGFBP-3'-97 and rhIGFBP-398-294 as well as structurally related proteins mac25 and CTGF were synthesized by use of a baculovirus expression system in insect cells that synthesize the correct proteins, as previously described in detail (20, 32, 40). The recombinant proteins possess the expected structure based upon purity and size characterized by silver staining and Western immunoblot and appropriate size shifting in Western immunoblot under reducing conditions, indicating the presence of S-S bonds in the proteins. IGF-I and IGF-II were purchased from GroPeP Pty. Ltd. (Adelaide, SA, Australia) and the biotinylation of IGF-I was performed according to the method of Fowkiles (30).

**Immobilization of peptides on the sensor chip**

All immobilizations were carried out at 25°C using an amine coupling procedure (33) with a constant flow rate of 5 µl/min. A more thorough description of the immobilization procedure can be found elsewhere (41). Equal volumes of 0.1 M N-hydroxysuccinimide and 0.1 M N-ethyl-N'-[3-dimethylaminopropyl)carbodiimide were mixed by the BIACORE system's robotics and injected over the surface of the sensor chip to activate the carboxymethylated dextran. For coupling to the sensor chip, peptides were injected over the activated surface in a 10-mM sodium acetate solution. A solution of 1 mM ethanolamine was then passed over the surface to deactivate the remaining active carbonyl groups and to wash out nonspecifically bound protein.

Combinations of peptide concentration, pH during coupling, activation time, and coupling time, respectively, giving a stable surface with satisfactory amounts of the coupled peptide, were as follows: rhIGFBP-3: 10 µg/ml, pH 4.5, 3 min, 7 min; rhIGFBP-3: 2 µg/ml, pH 4.5, 7 min, 7 min; rhIGFBP-3': 20 µg/ml, pH 4.0, 10 min, 10 min; IGF-I: 4 µg/ml, pH 4.5, 7 min, 7 min; and IGF-II: 1 µg/ml, pH 4.0, 7 min, 7 min. All ethanolamine deactivation steps were run for 7 min. Immediately after the immobilization procedure, HBS buffer (10 mM HEPES; 150 mM NaCl; 3.4 mM EDTA; and 0.005% P20, pH 7.4) was flowed over the sensor chip surface for a minimum of 2 h to allow the surface to stabilize.

**Quality control of immobilized surfaces**

In repeated experiments for each peptide, concentration and coupling time were varied to give a range of immobilized peptide coupled to the sensor chip to check for possible mass transport effects, which are expected to be most pronounced at high immobilization levels and high binding affinities (42, 43), and also to control for dependence of fitted binding parameters or of apparent stoichiometry of binding on immobilization level. We calculate the apparent stoichiometry as the ratio of the maximum achievable analyte bound, measured in the instrument's standard resonance units (RU), to the theoretical maximum, where the theoretical maximum is defined as: immobilized RU × molecular weight (analyte)/molecular weight (immobilized peptide), where immobilized RU is the amount of peptide immobilized on the sensor chip, also expressed in RU. The apparent stoichiometry gives a good indication of the degree of inactivation of protein due to immobilization and the steric hinderance of binding due to overcrowding of immobilized molecules on the surface. Immobilization levels in our experiments were kept low both to give as high apparent stoichiometries as possible and to reduce steric hinderance effects. Typical apparent stoichiometries for immobilized rhIGFBP-3 were 0.8-0.9, whereas those for immobilized IGF-I were about 25% and even lower for IGF-II. Note that a low apparent stoichiometry will still give accurate binding results if the immobilized molecules are all either fully active or fully inactive.

Mass transport effects (42, 43) were observed when measuring binding of rhIGFBP-3 to immobilized IGF-I or IGF-II, and these effects could not be fully eliminated even at the high flow rates (50 µl/min) used in our experiments. Mass transport effects were not apparent for any of the other tested molecules. In the case of rhIGFBP-3 binding to immobilized IGF-I, analysis of sensorgram data using models including or lacking mass transport effects showed only minor differences in estimated binding parameters. These differences were especially minimal in global analyses (i.e. simultaneous analysis of multiple sensorgrams at different analyte concentrations) (44, 45). None of the experiments showed a systematic change in binding parameters with immobilization level.

**Kinetic assays on the BIACORE**

All experiments were carried out at 25°C, with a constant flow rate of 50 µl/min HBS buffer. This high flow rate was chosen to minimize shear effects (46). Purified analyte was diluted to various concentrations in HBS buffer using the system robotics, and the solution was injected over the peptides coupled to the chip surface for 5 min (association phase), followed by 10-min flow of HBS buffer alone (dissociation phase). In most experiments the binding phases were preceded by a 10-min wash with HBS buffer alone to allow the surface to equilibrate with the buffer. Bound analyte was removed from the coupled peptide by flowing a solution of 100 mM HCl over the surface for 3 min. This treatment regenerated the surfaces efficiently without any apparent damage. We observed, however, that IGF-I surfaces immobilized at very low levels tended to be somewhat unstable. In these cases the surface was regenerated using 50 mM HCl to minimize any damaging effects of the regeneration procedure.

In most experiments, a standard intermediate concentration of analyte was first injected over the surface to provide a reference sensorgram. Then a series of varying concentrations of analyte was injected, followed by a second reference injection, a repeat of the concentration series, and a final reference injection. With this procedure the immobilized surface could be monitored for loss of activity, and test results checked for reproducibility.

**Solution affinity assays**

Solution affinity assays are designed to measure the equilibrium affinity of two molecules in solution, using the BIACORE instrument as a probe to measure the free concentration of one of the molecules (47). In these experiments rhIGFBP-3 or one of its N-or C-terminal fragments was immobilized on a sensor chip to provide an active surface for measuring the free concentration of IGF-I or IGF-II in a mixture of IGF and binding protein flowing over the surface. The surface was calibrated by flowing IGF at different (free) concentrations over the surface, recording a sensorgram for each concentration. Mixtures of IGF and binding protein or fragment at various concentrations were flowed over the
same surface, and the calibration curve was used to estimate the free IGF remaining in solution for each of the mixed samples. The equilibrium affinity constant, $K_a$, was calculated using this estimate plus the known total concentrations of IGF and binding protein, assuming a one-site binding model.

**Data analysis**

Kinetic analyses were carried out using the BIAEvaluation 3.0 program (Biacore AB), assuming a one-site binding model. If the surface was judged stable by inspection of the repeated series of injected analyte concentrations, all sensorgrams in an experiment were analyzed simultaneously (global analysis). If a surface showed some instability, sensorgrams were analyzed individually, and the results pooled. In all cases, special attention was paid to the fitted $R_{\text{max}}$, which gives the amount of analyte bound at saturation and is therefore a measure of the apparent stoichiometry of binding. We have observed that even a visually good fit of a binding model to sensorgram data sometimes gives an unreasonable value of $R_{\text{max}}$, especially when analyzing single sensorgrams. Binding parameter estimates were rejected in such cases.

Model fitting was first done allowing the BIAEvaluation program to estimate the bulk refractive index effect for each sensorgram. Sensorgrams were then inspected visually to obtain another estimate of these values based on the step up in the sensorgram at the start of the association phase and the step down at the start of the dissociation phase (33). If the fitted estimates did not correspond to the visual estimates, the fitted estimates were rejected, and the fit was re-done with the refractive index values fixed using the visual estimates. In few experiments the surface baseline signal (that is, the signal seen when buffer alone flows over the surface) drifted slowly downward through time (baseline drift), evident in the 10-min preassociation wash and at the end of the dissociation phase. In these cases, the downward drift rate was estimated by fitting a straight line to the 10-min wash, and the sensorgram was corrected for this drift during the fitting.

**Results**

**Binding of IGF-I and IGF-II to rhIGFBP-3 and its N- and C-terminal fragments**

With the BIACORE instrument, a binding reaction can be measured with either of the reacting pair immobilized, and the other injected over the immobilized surface. In this study we attempted to measure all binding reactions in both orientations, which would be expected to give comparable results if the experimental conditions are suitable.

Figure 1 shows the binding of IGF-I to rhIGFBP-3 with either rhIGFBP-3 immobilized (412 RU; Fig. 1A) or IGF-I immobilized (256 RU; Fig. 1B). These sensorgrams show the typical protocol used in these studies: a 10-min wash to allow the surface to equilibrate with the buffer, followed by 5 min of association and 10 min of dissociation phase. The short downward spikes seen in the sensorgrams are disturbances resulting from opening and closing of valves in the BIACORE instrument's flow system. A sharp rise in signal at the beginning of the association phase and a corresponding sharp drop at the beginning of the dissociation phase are due to a small refractive index difference between buffer alone and buffer containing the analyte protein.

In Fig. 1, A and B, the concentration of analyte was 20 nM. The difference in maximum analyte bound (75 RU IGF-I vs. 200 RU rhIGFBP-3) is due to the difference in mol wt. The linearity of the initial portion of the association phase of the sensorgram in Fig. 1B indicates the presence of mass transport effects. However, analyses of these experiments with or without including mass transport in the model gave comparable results, which were also in agreement with results obtained with rhIGFBP-3 immobilized and IGF-I as analyte.

![Figure 1](image-url)  
**Fig. 1**. A, IGF-I (20 nM) binding to immobilized full-length rhIGFBP-3 (412 RU). HBS buffer was flowed over the surface for 12 min, followed by 5 min of association and 10 min of dissociation of IGF-I. B, Full-length rhIGFBP-3 (20 nM) binding to immobilized IGF-I (256 RU).

Figure 2 shows the results of similar experiments using the binding protein fragments rhIGFBP-31-97 and rhIGFBP-398-264. For each of the fragments, the shapes of the sensorgrams were similar whether the fragment or IGF-I was immobilized. The rhIGFBP-31-97 and rhIGFBP-398-264 sensorgrams had different general shapes because both the kinetic association and dissociation constants were slower for rhIGFBP-398-264 than for rhIGFBP-31-97 (see Table 1), although these differences were not statistically significant over the entire set of experiments. Note that there was a downward or upward drift in signal during the initial wash stage in the experiments with binding protein fragments immobilized. This was a general observation during these studies; that is, sensor chip surfaces immobilized with binding protein fragment tend to...
be more unstable than those immobilized with full-length rhIGFBP-3, or with IGF-I.

In Fig. 3 is shown a comparison of the measured binding affinity for IGF-I binding to full-length rhIGFBP-3 or to the binding protein fragments, either with binding protein immobilized or with IGF-I immobilized. The data are plotted with a logarithmic ordinate, and the results show that the affinity of IGF-I for either rhIGFBP-31–97 or rhIGFBP-398–264 is approximately 1000-fold lower than the affinity for full-length rhIGFBP-3. Although the mean affinity for rhIGFBP-31–97 measured in these experiments was higher than that for rhIGFBP-398–264, the differences were not statistically significant. All differences between full-length rhIGFBP-3 and the fragments were significant.

Figure 3 also shows that experiments in either immobilization orientation give similar measured affinities. However, because of apparent self-aggregation of rhIGFBP-3 and the C-terminal fragment (see Discussion), the results with IGF-I immobilized on the sensor chip were considered less reliable.

Whenever possible, binding parameter estimates were obtained from global fitting to multiple sensorgrams at different analyte concentrations. An example of such an experiment is shown in Fig. 4. In this experiment rhIGFBP-3 was immobilized at 726 RU, and IGF-I at concentrations from 0.5–50 nM was flowed over the immobilized surface. Because of the high affinity of IGF-I for rhIGFBP-3, binding is affected by mass transport limitation, which is most pronounced at lower analyte concentrations. The entire set of sensorgram data was fitted using a model that includes mass transport effects, and for comparison was refitted using an ordinary one-site (Langmuir) binding model. For this experiment, the computed binding affinities for the two models were $5.6 \times 10^9$ and $5.8 \times 10^9$ M$^{-1}$, respectively. The respective kinetic association constants were $8.3 \times 10^3$ and $6.4 \times 10^3$ M$^{-1}$s$^{-1}$, and the dissociation constants were $1.4 \times 10^{-4}$ and $1.2 \times 10^{-4}$
Binding of IGF-II or biotinylated IGF-I to rhIGFBP-3 and its fragments

All of the above experiments were repeated using IGF-II instead of IGF-I. In all experiments the sensorgrams for IGF-II were visually very similar to those for IGF-I, and calculated binding parameters were also similar. Figure 5 shows sensorgrams for IGF-I and IGF-II as well as a biotin-labeled IGF-I, all at 50 nm, binding to the same immobilized rhIGFBP-3 surface. These sensorgrams were taken from a larger experiment designed for global analysis. The top two sensorgrams are those for IGF-I and IGF-II. Note that in the association phase the IGF-I signal is higher than that for IGF-II, which is due solely to a slightly higher refractive index of the IGF-I-containing buffer. In the dissociation phase, where both surfaces are exposed to the same buffer, the IGF-II signal is slightly higher, indicating a higher affinity than for IGF-I. The affinities for IGF-I and IGF-II in this experiment, calculated by global analysis, were $2.1 \times 10^9$ and $3.0 \times 10^9$ m$^{-1}$, respectively.

Table 1 gives the summary of the calculated binding parameters for IGF-I and IGF-II binding to immobilized rhIGFBP-3 or its N- or C-terminal fragments. The table only includes results from experiments using immobilized binding protein and fragments, as the experiments with the immobilized IGFs were judged to be less reliable, primarily due to possible self-association of binding proteins (see Discussion). None of the parameter differences between the two binding protein fragments was statistically significant. All differences between full-length rhIGFBP-3 and the fragments were significant, except for those with NS in parentheses.

![Graph](image-url)
rhIGFBP-3. In this experiment, using global analysis, the affinity of biotin-IGF-I was calculated to be $1.1 \times 10^9$ M$^{-1}$.

**Binding of human insulin to rhIGFBP-3 and its N- and C-terminal fragments**

In this study we tested the binding of human insulin to rhIGFBP-3 and its fragments, with either binding protein or insulin immobilized on the sensor chip. In most experiments the observed binding was very weak or absent in either configuration, and reasonable estimates of binding parameters could not be made, nor could comparisons be made of the relative affinities of insulin binding to full-length rhIGFBP-3 or its fragments. Apparent binding was observed in a few experiments, and by comparing this binding with control runs using IGF-I, we could estimate equilibrium $K_a$ values less than $10^6$ M$^{-1}$. Figure 6A shows an example of the weak binding of human insulin (2 μM) to immobilized rhIGFBP-3. For this sensorgram, the $K_a$ was estimated to be less than $10^5$ M$^{-1}$, assuming that insulin binds with the same stoichiometry as IGF-I.

**Binding of IGF-I and IGF-II to mac25 and CTGF**

We also tested the binding of IGF-I, IGF-II, and human insulin to mac25 and CTGF. In no case could we unambiguously demonstrate binding of IGF or insulin molecules to these proteins. Figure 6, B and C, shows examples of tests of mac25 and CTGF binding to immobilized IGF-I.

**Self-association of IGFBP and its fragments**

Most of our experiments using immobilized IGFs gave consistent results, with calculated binding parameters very similar to those for the experiments with IGFBP immobilized (see Fig. 3). However, because of potential inaccuracies in binding parameter determinations due to the two-dimensional geometry of binding, we also attempted to measure equilibrium binding affinities using solution affinity assays on the BIACORE. In an example of one such experiment, IGF-I at 20 nM was incubated with varying concentrations (2–200 nM) of rhIGFBP-3 for 15–20 min at 25 °C, after which the solution was passed over an rhIGFBP-3-immobilized sensor chip to record a sensorgram. Because rhIGFBP-3 in solution should compete for free IGF-I, the binding of IGF-I to the immobilized rhIGFBP-3 was expected to decrease at higher rhIGFBP-3 concentrations. Instead, the signal increased at higher concentrations. The most straightforward interpretation of this was that the rhIGFBP-3 was itself binding to the rhIGFBP-3 immobilized surface, and therefore solution assays could not be performed in this system.

We then investigated directly the binding of rhIGFBP-3 and its fragments to each other. Figure 7 shows an example of binding of rhIGFBP-3 or rhIGFBP-3$^{98-264}$ to a surface immobilized with rhIGFBP-3$^{98-264}$, in which binding is clearly evident. From experiments using all combinations of immobilized and solution proteins, we observed that rhIGFBP-3 and rhIGFBP-3$^{98-264}$ bind weakly to themselves and to each other (apparent $K_a$ $\approx 8 \times 10^6$ M$^{-1}$; that is, more than 100-fold lower affinity than that of IGF-I for rhIGFBP-3), whereas rhIGFBP-3$^{1-97}$ binds very weakly or not at all to itself and to the other proteins ($K_a < 3 \times 10^5$ M$^{-1}$). Because rhIGFBP-3 and rhIGFBP-3$^{98-264}$ would be expected to also aggregate in the flow-through solution in these assays, these results are approximate.

**Discussion**

If the N- and C-terminal regions of IGFBP-3 combine to form a high affinity binding site for IGFBP-3 might individually bind IGF-I or IGF-II, but with lower affinity than the full-length binding protein. The binding regions on the two fragments could combine to form a single larger high affinity site or could interact with two separate regions on the IGF molecule to result in high affinity binding. This last mechanism has been suggested, for example, for insulin binding to the insulin receptor (48, 49). Our results here show that IGF-I and IGF-II bind the N- and C-terminal recombinant fragments rhIGFBP-3$^{1-97}$ and rhIGFBP-3$^{98-264}$ with affinities approximately 1000-fold lower than that of the IGFs for full-length rhIGFBP-3. These results are consistent with the idea that the combination of the two sites results in high affinity binding, but do not distinguish whether there is a single high affinity binding site or two separate sites.

The functional significance of different glycosylation forms of recombinant human IGFBP-3 was studied by Firth et al. (18). It could be shown that different glycosylation patterns (e.g., fully glycosylated vs. nonglycosylated IGFBP-3) do not alter IGF binding to the binding protein in Western ligand blots. Therefore, the difference in binding affinity of nonglycosylated *Escherichia coli*-expressed IGFBP-3 and baculovirus-expressed glycosylated IGFBP-3 fragments is not expected to be due to their glycosylation status.

Recently, Galanis et al. (38), using a BIACORE instrument,
measured equilibrium affinities of $1.03 \times 10^7$ and $6.16 \times 10^7$ M$^{-1}$ for IGF-I and IGF-II binding, respectively, to an N-terminal IGFBP-3$^{1-88}$ fragment, compared with our measurements of about $0.6 \times 10^9$ M$^{-1}$ for both IGF-I and IGF-II. Interestingly, they found that the 1–88 N-terminal fragment could not be immobilized using the amine coupling method, and that a 165–264 C-terminal fragment could be immobilized, but was inactive in BIACORE assays. In contrast, both our 1–97 N-terminal and 98–264 C-terminal fragments could be immobilized and retained activity, and both were also active in binding to immobilized IGF-I.

Our measurements of the binding of IGF-I and IGF-II to full-length rhIGFBP-3 indicate equilibrium affinities in the range of $4–5 \times 10^9$ M$^{-1}$. These estimates are lower than those measured using $^{125}$I-labeled IGFs, where affinities between $8 \times 10^9$ and $152 \times 10^9$ M$^{-1}$ have been reported (quoted and summarized in Ref. 36). We had observed a similar difference in an earlier study of IGF analog binding to IGFBP-3 (33), but it is not clear whether this is due to the different conditions of binding, for example, changes in the ligand due to immobilization or the two-dimensional binding geometry, or to differences in analytical methods.

Wong et al. (36), using a BIACORE instrument, calculated an affinity of $18.4 \times 10^9$ M$^{-1}$ for IGF-I binding to hrIGFBP-3. These results are apparently the average of separate analyses of individual sensorgrams, whereas our results for IGF-I binding to immobilized rhIGFBP-3 are derived from global analyses (44, 45) (see Fig. 4), in which a binding model is fitted simultaneously to multiple sensorgrams at different analyte concentrations. Global analyses should be more accurate, because they assume a common value in all sensorgrams for $R_{\text{max}}$, the maximum analyte bound at saturation, whereas analyses of single sensorgrams from the same experiment can yield different apparent values of $R_{\text{max}}$, which is unrealistic.

Galanis et al. (38) also determined a high affinity ($1 \times 10^{11}$

**Fig. 6.** A, Interaction of human insulin (2000 nM) with full-length rhIGFBP-3 (412 RU). The rising association phase and falling dissociation phase indicate apparent binding, but with very low affinity, and near the detection limit of the instrument. Interaction of mac25 (B) and CTGF (C) with immobilized IGF-I (532 RU). Binding is below the detection limit of the instrument.
assays impossible and raising questions about the accuracy of binding results obtained using binding proteins in the flow solution and immobilized IGFs. With increasing binding protein concentration, self-aggregation should increase both in solution and on the sensor chip surface, introducing errors in the binding analysis. Therefore, results using immobilized binding proteins, for example, as shown in Table 1, should be more reliable than those using immobilized IGFs (36, 38) unless binding protein concentrations are kept well below the $K_d$ for self-aggregation ($\sim 10^{-7}$ M by our estimates).

On the basis of evidence that activation of the insulin receptor is inhibited by IGFBP5, and that mac25 is able to bind insulin (13), we tested the binding of human insulin to rhIGFBP-3 and to recombinant mac25 and CTGF molecules using the BIACORE instrument. We also tested insulin binding to the IGFBP recombinant fragments rhIGFBP3-1-97 and rhIGFBP98-264. Most experiments could not confirm binding, but in a few experiments the binding of insulin to rhIGFBP-3 was detectable, indicating that insulin may bind IGFBP-3 with low affinity, as we had previously observed (33). We could not detect binding of insulin to the IGFBP-related proteins, or to the rhIGFBP-3 fragments, in these biosensor assays.

We tested the binding of IGF-I and IGF-II to the mac25 and CTGF molecules, with either these molecules or the IGF molecules immobilized, at analyte concentrations up to 500 nM. In no case could we detect binding using the biosensor instrument. These results were unexpected in light of the evidence for binding of both IGF-I and insulin to the binding protein molecules from studies using radiolabeled IGF molecules (13, 20). However, as binding affinities less than about $10^6$ or $10^7$ M$^{-1}$ are difficult to detect using the biosensor, we cannot rule out that there is weak binding below the detection limit of the instrument. There is an indication from affinity cross-linking studies that IGF-I binding affinity for mac25 may be higher than these values (13, 40) so it remains to be determined whether our inability to detect binding on the biosensor instrument is because of differences in binding conditions or because cross-linking experiments are, in fact, measuring extremely low binding affinities.

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**Fig. 7.** Interaction of full-length rhIGFBP-3 and rhIGFBP-3<sup>398-264</sup> with immobilized rhIGFBP-3<sup>398-264</sup> (3307 RU). Solid lines, rhIGFBP-3 (bottom to top, 50, 200, and 500 nM); dashed lines, rhIGFBP-3<sup>398-264</sup> (bottom to top, 50, 200, and 500 nM).
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Characterization of Insulin-Like Growth Factor-Binding Protein-Related Proteins (IGFBP-rPs) 1, 2, and 3 in Human Prostate Epithelial Cells: Potential Roles for IGFBP-rP1 and 2 in Senescence of the Prostatic Epithelium*


ABSTRACT

Insulin-like growth factor (IGF)-binding protein (IGFBP)-related proteins (IGFBP-rPs) are newly described cysteine-rich proteins that share significant aminoterminal structural similarity with the conventional IGFBPs and are involved in a diversity of biological functions, including growth regulation. IGFBP-rP1 (MAC25/Angiomedulin/prostacyclin-stimulating factor) is a potential tumor-suppressor gene that is differentially expressed in meningiomas, mammary and prostatic cancers, compared with their malignant counterparts. We have previously shown that IGFBP-rP1 is preferentially produced by primary cultures of human prostate epithelial cells (HPECs) and by poorly tumorigenic P68SV40T cells, compared with the cancerous prostatic LNCaP, DU145, PC-3, and M12 cells. We now show that IGFBP-rP1 increases during senescence of HPEC.

IGFBP-rP2 (also known as connective tissue growth factor), a downstream effector of transforming growth factor (TGF)-β and modulator of growth for both fibroblasts and endothelial cells, was detected in most of the normal and malignant prostatic epithelial cells tested, with a marked up-regulation of IGFBP-rP2 during senescence of HPEC. Moreover, IGFBP-rP2 noticeably increased in response to TGF-β1 and all-trans retinoic acid (atRA) in HPEC and PC-3 cells, and it decreased in response to IGF-I in HPEC.

IGFBP-rP3 (nephroblastoma overexpressed (NOV)), the protein product of the NOV protooncogene, was not detected in HPEC but was expressed in the tumorigenic DU145 and PC-3 cells. It was also synthesized by the SV40-T antigen-transformed P69 and malignant M12 cells, where it was down-regulated by atRA.

These observations suggest biological roles of IGFBP-rPs in the human prostate. IGFBP-rP1 and IGFBP-rP2 are likely to negatively regulate growth, because they seem to increase during senescence of the prostate epithelium and in response to growth inhibitors (TGF-β1 and atRA). Although the data collected on IGFBP-rP3 in prostate are modest, its role as a growth stimulator and/or protooncogene is supported by its preferential expression in cancerous cells and its down-regulation by atRA. (Endocrinology 141: 4072-4080, 2000)

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HE INSULIN-LIKE growth factor (IGF) system is composed of two ligands (IGF-I and IGF-II), six IGF-binding proteins (IGFBPs -1 to -6), and two receptors (type 1 and type 2 IGF receptors) (1). Recently, the IGFBP family of proteins has been expanded to include additional members that share significant structural similarities, the so-called IGFBP-related proteins (IGFBP-rPs) (2), which not only share the conserved aminoterminal domain of the IGFBPs but also show some degree of affinity for IGFs and insulin.

Abnormalities in the IGF system have been identified in prostate disease, such as prostate hyperplasia and cancer. In this respect, prostate cancer growth seems to be poorly dependent on IGFs, because the type I IGF receptor, which mediates most biological functions of IGF-I and IGF-II, is down-regulated during prostate carcinogenesis (3). Additionally, recent epidemiological studies have shown an increased risk of developing prostatic carcinoma in adult males with high-normal serum concentrations of IGF-I (4).

Despite extensive characterization of the IGFBPs in the human prostate, little is known about their roles in prostate physiology. IGFBP-2 and -4 are known to increase during carcinogenesis of the prostatic epithelium (5, 6), whereas IGFBP-3 is a proteolytic substrate for prostate-specific antigen (7, 8). Recent studies have also shown an increase of several IGFBPs in prostate tissue during involution of the gland in castrated rats and in men taking finasteride (9, 10).

These findings suggest that IGFBPs may play a role in apoptosis of prostate cells, either by sequestering IGFs or by direct cellular actions. Our knowledge of the new IGFBP-rPs in the human prostate is only scant at its best.

The IGFBP-rPs are cysteine-rich proteins involved in a diversity of biological functions, including growth regula-
tion. IGFBP-rP1 was originally cloned from leptomeningial cells and was termed meningioma-associated complementary DNA (cDNA) (MAC25) (11). This protein has also been reported as tumor-derived adhesion factor (TAF, recently renamed Angiomodulin) (12, 13), prostacyclin-stimulating factor (14), and T1A12 (15). In the human prostate, we have also recently described preferential expression of IGFBP-rP1 in normal human prostate cells and tissues, compared with their malignant counterparts, and we have shown that this protein is up-regulated by transforming growth factor (TGF-β) and all-trans retinoic acid (atRA) in prostate epithelial cells (16).

IGFBP-rP2 [also known as connective tissue growth factor (CTGF)] was initially isolated from human umbilical endothelial cells and shown to be mitogenic and chemotactic for fibroblasts (17). IGFBP-rP2 belongs to the CCN [for CYR61, CTGF, and nephroblastoma overexpressed (NOV)] family of cysteine-rich proteins involved in a diversity of cellular functions, such as mitogenesis, differentiation, survival, adhesion, migration, and regulation of matrix gene expression (18). IGFBP-rP2 is also a major downstream effector of TGF-β in fibroblasts, where it was evidenced that TGF-β controls IGFBP-rP2 expression via a novel response element in the IGFBP-rP2 gene (19). To date, however, no studies have been reported on the role of IGFBP-rP2 in prostate biology and/or carcinogenesis.

IGFBP-rP3 (also known as NOV) was first recognized as an aberrantly expressed gene in avian nephroblastoma and later shown to be also overexpressed in the human homologue Wilms tumor (20, 21). In the human prostate, it has been reported that NOV is differentially expressed in PC-3 cells, compared with other cancerous and normal cells in culture, but the significance of this finding is unknown (22).

We now report that prostate epithelial cells, in culture, express not only IGFBP-rP1 and 3, but also IGFBP-rP2, and that these proteins are responsive to growth regulators. Interestingly, IGFBP-rP1 and 2 also increase during senescence tissue culture, but the significance of this finding is unknown (22).

We now report that prostate epithelial cells, in culture, express not only IGFBP-rP1 and 3, but also IGFBP-rP2, and that these proteins are responsive to growth regulators. Interestingly, IGFBP-rP1 and 2 also increase during senescence of normal prostate epithelial cells, thus supporting growth-regulatory roles of these proteins in the prostate epithelium.

Materials and Methods

Materials

Epidermal growth factor (EGF), dexamethasone, atRA, and the additive ITS (insulin, transferrin, selenium) were purchased from Sigma (St. Louis, MO). PrEBM human prostate epithelial cell (HPEC) media was obtained from Clonetics (San Diego, CA). RPMI (1640 media) was obtained from Life Technologies (Grand Island, NY). IGF-I and TGF-β1 were both purchased from Austral Biologicals (San Ramon, CA). FBS was obtained from HyClone Laboratories, Inc. (Logan, UT). [3H]methylthymidine was purchased from NEN Life Science Products (Boston, MA), and MTS assay kit (CellTiter 96 AQueous One Solution Cell Proliferation Assay) was obtained from Promega Corp. Corporation (Madison, WI). Nitrocellulose and electrophoresis reagents were purchased from Bio-Rad Laboratories, Inc. (Hercules, CA); nylon membranes (Genescreen) were obtained from NEN Life Science Products. Horseradish peroxidase-linked donkey antirabbit and sheep antimouse IgG antibodies and enhanced chemiluminescence detection reagents were purchased from Amersham Pharmacia Biotech (Arlington Heights, IL). HPEC cells were purchased from Clonetics. LNCaP, DU145, and PC-3 cells were obtained from American Type Culture Collection (Manassas, VA). P69SV40T (P69) and M12 prostate epithelial cells were previously described (23). Polyclonal antibodies against IGFBP-rP1, IGFBP-rP2, and IGFBP-rP3 were generated in rabbits, as previously described (24–26).

Monoclonal IgG antibody against p16INK4a (13251A) was purchased from Phar-Mingen (San Diego, CA).

Cell culture

Figure 1 summarizes the cell lines employed in these studies. HPEC cells were maintained in PrEBM media supplemented with the following: bovine pituitary extract (BPE), insulin, hydrocortisone, CA-1000, retinoic acid, transferrin, levophthyroxine, epinephrine, and human EGF. HPECs were subcultured according to the manufacturer. When they reached 80% confluence, all growth factors were withdrawn, except for BPE, for 12 h. Medium was changed again to PrEBM plus BPE for cell proliferation and growth factor regulation studies. P69 and its M12 subline were grown in RPMI media supplemented with 10 ng/ml EGF, 0.1 µM dexamethasone, 5 µg/ml insulin, 5 µg/ml transferrin, and 5 ng/ml selenium. LNCaP, DU145, and PC-3 cells were maintained in RPMI enriched with 10% FBS. All cultures used were mycoplasma-free, as determined by the Mycoplasma PCR Primer Set (Stratagene, La Jolla, CA) and were grown at 37 C under 5% CO2.

Cellular proliferation assays

Cells were seeded at a density of 1 × 10⁴ cells per well, in 48-well plates (Falcon, Becton Dickinson and Co. Labware, Franklin Lakes, NJ), in 500 µl of either rich defined medium or 10% FBS-enriched medium, depending on the cell line. When 80% confluence was reached, the medium was changed to either basal medium or serum free medium for 12 h and then treated with various doses of TGF-β1 (0–5 ng/ml), atRA (0.001–1 µM), or IGF-I (0–100 ng/ml) in 250 µl of the same serum-free medium for 48 h. [3H]thymidine (0.4 µCi/ml) was added for the last 24 h (HPEC and PC-3 cells) or for the last 6 h (P69 and M12 cells). After labeling, the cell layers were washed twice with PBS and incubated with 10% trichloroacetic acid (TCA) at −20 C for 15 min, followed by another wash with 0.25 N sodium hydroxide, and the precipitated material was read by means of an LS 6500 Scintillation Counter (Beckman Coulter, Inc., Fullerton, CA).

Cell proliferation was also tested by MTS assay kit, according to the manufacturer's instructions. Cells were incubated for the last 2 h of treatment, in the presence of the MTS reagent, and absorbance was measured at λmax nm.

Growth factor regulation studies

Cells were seeded at a density of 10 × 10⁴ cells per well, in 60-mm tissue culture dishes, and grown to 80% confluence. Treatments with TGF-β1 (0–5 ng/ml), atRA (0.001–1 µM), or IGF-I (0–100 ng/ml) were also done under serum-free conditions, for 48 h, after which conditioned media, total cell lysates, and total cytoplasmic RNA were collected for Western immunoblots (see Western Immunoblot analyses) and Northern blots (see RNA analyses) studies.

![Fig. 1. Prostate epithelial cells studied. Primary cultures of normal HPEC were transformed with the SV40-T antigen to obtain the P69 cell line with low tumorigenic potential. M12 cells, originated after several passages of P69 cells in athymic mice, are, however, highly tumorigenic and metastatic. The well-established androgen-dependent (LNCaP) and androgen-independent (DU145 and PC-3) cells lines, derived from clinical samples, were also studied.](image-url)
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Western immunoblot analyses

Conditioned media and total cell lysates, using RIPA buffer [150 mM NaCl, 20 mM HEPES (pH 7.4), 1% (vol/vol) Triton X-100, 1% (wt/vol) sodium deoxycholate, 0.1% (wt/vol) SDS, and Mini EDTA-free protease inhibitors (Roche Molecular Biochemicals, Laval, Québec, Canada)], from both treated and untreated (control) cells, were normalized for protein concentration, using a DC protein assay (Bio-Rad Laboratories, Inc.). Equal amounts of total protein per sample were separated in non-denaturing SDS sample buffer [0.5 M Tris (pH 6.8), 1% SDS, 10% glycerol, and bromophenol blue] and boiled for 5 min. Samples were electrophoresed on 15% SDS-polyacrylamide gels, then electroblotted onto nitrocellulose, and membranes blocked with 4% milk-TBS-T [Tris-buffered saline-Tween-20 (0.1%) for 1 h at 22 C. Western blots were incubated with IGFBP-rPl, -rP2, or -rP3 antisera at a 1:3000 dilution and with p16INK4a IgG antibody at a dilution of 1:500 (1 μg/ml) in TBS-T overnight at 4 C. Blots were washed with TBS-T and then incubated for 1 h at 22 C with a 1:3000 dilution of horseradish peroxidase-linked anti-rabbit or antimouse IgG secondary antibodies. Proteins of interest were detected with ECL chemiluminescence reagents, according to the manufacturer’s protocol.

RNA analyses

Total cytoplasmic RNA was isolated from cells, by use of RNaseasy (QIAGEN, Inc., Chatsworth, CA). Twenty micrograms of each RNA preparation were electrophoresed on a 1.2% agarose-2.2 M formaldehyde gel, transferred overnight onto a nylon membrane (GeneScreen), using 10 X SSC as the transfer solution, and cross-linked to the membrane by UV irradiation in a Stratalinker 1800 (Stratagene). The Northern blots were then probed with an EcoRI/Xho fragment of IGFBP-rPl (27), or a BamHI/Xho fragment of IGFBP-rP2 (28), which were radiolabeled (1 X 10^6 dpm/μg) with [α-32P]deoxyctydine triphosphate (NEN Life Science Products-DuPont; SA, 3000 Ci/mm) using a random priming kit (Prime-a-Gene, Promega Corp.). Northern blots were hybridized overnight at 65 C in hybridization buffer (Rapid-Hyb, Amersham Pharmacia Biotech), according to the manufacturer’s instructions. Blots were then washed for 15 min in 2 X SSC/0.1% SDS at 22 C, followed by two more stringent washes in 0.2 X SSC/0.1% SDS at 65 C for 15 min. Blots were exposed to Kodak Biomax film (Eastman Kodak Co., Rochester, NY) for 12 to 48 h at ~70 C, using one intensifying screen. Membranes were then reprobed with 18S ribosomal RNA, which acted as a loading control for the RNA samples. An image analyzer (GS-700) equipped with MultiAnalyist version 1.02 Software (Bio-Rad Laboratories, Inc.) was used to quantify the resulting bands.

RT-PCR

RT-PCR was performed using 5'-CCCGAATTCTCGGATCCAGTGGTCCAGAAGCAGC-3' and 5'-GGGCGTCCAGGTACATTTCCCTGCGATTGC-3' primers specific for IGFBP-rPl. One microgram of total RNA from each cell line was reversed transcribed in a vol of 20 μl, by use of Reverse Transcription System Kit (Promega Corp.) following the manufacturer’s instructions. The reaction was performed at 42 C for 15 min, denatured at 99 C for 5 min, and placed on ice. One microliter of the mixture and 50 pmol of 5' and 3' primers were employed in PCR amplification reactions using Advantage GC cDNA PCR Kit (CLONTECH Laboratories, Inc., Palo Alto, CA). Amplification of the cDNA was carried out with 25 cycles of denaturing at 94 C for 1 min, annealing at 55 C for 1 min, and extension at 72 C for 2 min. One negative and one positive control were included in all reactions.

Statistical analyses

All experiments were performed at least twice. Statistical analyses were performed by a two-tail Student’s t test, assuming unequal variances, using Excel Data Analysis Software (Microsoft Corp., Redmond, WA). Data are expressed as means ± SE. P < 0.05 was considered significant.

Results

Expression of IGFBP-rPl, 2, and 3 in HPECs

In agreement with previous observations (16), IGFBP-rPl messenger RNA (mRNA) was detected by Northern blot analysis in the P69/M12 lineage and in primary cultures of prostate epithelial cells, with a parallel detection of IGFBP-rP1 in conditioned media from these cultures (Fig. 2). In the malignant LNCaP, DU145, and PC-3 cells, IGFBP-rPl mRNA was undetectable by Northern blot, and an immunoreactive band was only present in cell lysates but not in conditioned media from these cells.

IGFBP-rP2 mRNA was detectable in all but one cell line as a single 2.4-kb band, consistent with the molecular size observed in most studies (29). IGFBP-rP2 protein was present in conditioned media and in cell lysates from these cultures. In contrast to IGFBP-rPl, both normal and malignant prostate epithelial cells express similar amounts of IGFBP-rP2 (Fig. 2).

IGFBP-rP3 message was evaluated by RT-PCR in only a limited number of cell lines but was identifiable in P69, M12, LNCaP, and PC-3 cells. IGFBP-rP3 protein was undetectable in conditioned media from HPEC, compared with readily detectable levels in the immortalized P69 cell line and malignant M12, DU145, and PC-3 cells (Fig. 2).

Expression of IGFBP-rPl and 2 during senescence of HPEC

HPEC cells have a limited life span, with no more than 30 population doublings before they enter replicative senescence (corresponding to our 9th culture passage). Thus, early passages of HPEC (<4th) are highly replicative cells (doubling time for 4th passage is approximately 2 days), whereas senescent cells (>9th passage) are unable to replicate, and they die over time (Fig. 3).

Interestingly, both IGFBP-rPl and 2 were up-regulated on increasing passage of these cells, with concentrations of both mRNA and secreted protein more than 5-fold higher at late passages, compared with early passages (Fig. 4, A and B).

In support of an increase of these proteins during senescence of HPEC, we also investigated the concentrations of cell-cycle inhibitors that are known to be modified during cellular senescence, such as p21WAF1 and p16INK4a (30). In agreement with a previous report in primary cultures of prostate epithelial cells (31), p21WAF1 was barely identifiable in our cell lysate preparations, either at early or late passages (data not shown); whereas a marked up-regulation of p16INK4a at late, low-replicative passages was evidenced (6-fold increase, compared with early passages), paralleling the increases in IGFBP-rPl and -P2 (Fig. 4C).

IGFBP-rP2 is responsive to growth regulators in HPECs

Because TGF-β, atRA, and IGF-I are important regulators of prostate epithelial growth and survival (32-34) and because IGFBP-rP2 is tightly regulated by TGF-β in other cellular systems, we wished to investigate the effects of these growth inhibitors (TGF-β and atRA) and growth stimulator (IGF-I) on cell proliferation and IGFBP-rP2 expression in our normal and malignant prostate cells.

Both TGF-β and atRA caused a dose-dependent inhibition
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Fig. 3. Growth rates of early and late passages of HPEC. Eight hundred cells of an early passage (4th) and of a late passage (9th) were seeded in 500 μl of rich media per well (48-well plate). At the time points indicated, cell proliferation was investigated by MTS assay, in triplicate wells. Early passages of HPEC are highly replicative cells with an approximate duplication time of 2 days, whereas late HPEC passages fail to replicate in vitro, and die over time. Results are expressed as means ± SE of two independent experiments. The lag phase observed at days 3 and 5 is accounted for by a change in the growth medium.

Because IGFBP-rP1 protein and mRNA increase also during senescence of HPEC, we investigated whether IGFBP-rP1 was similarly regulated by these growth regulators. In contrast to IGFBP-rP2, the concentrations of both IGFBP-rP1 protein and steady-state mRNA in HPEC did not change in response to TGF-β treatment. In response to atRA treatment, a 2-fold increase of secreted IGFBP-rP1 protein was detected, although no changes in IGFBP-rP1 mRNA were observed (Figs. 6, A and B).

The effects of IGF-I were also tested in the normal HPEC (late, presenescent passages), where it exerted clear mitogenic actions (Fig. 5A). Importantly, IGFBP-rP2 was also regulated by IGF-I in HPEC, both at the protein and at the mRNA level, with a 90% reduction of both secreted protein and steady-state mRNA, upon treatment with IGF-I at a dose of 100 ng/ml (Fig. 6, C and D). In contrast to IGFBP-rP2, neither IGFBP-rP1 protein nor its mRNA experienced significant changes in response to IGF-I treatment in HPEC (Figs. 6, C and D).

In PC-3 cells, only the effects of TGF-β and atRA on IGFBP-
IGFBP-rP3 is regulated in HPECs

IGFBP-rP3 has been previously shown to be expressed in prostate epithelial cells, but no regulation has been described, as yet, in these cultures (22). To better understand the role of this protein in the human prostate, we also investigated the effects of TGF-β, atRA, and IGF-I on the expression of IGFBP-rP3 in vitro.

IGFBP-rP3 was not produced by normal prostate epithelial cells, nor was it induced by any of the above-mentioned growth regulators in these cells. In the well-established PC-3 (and in DU145) cancer cells, IGFBP-rP3 was readily detected in conditioned media under serum-free incubation, but no significant regulation was observed in these cultures.

In the P69/M12 lineage, both TGF-β and atRA also produced a dose-dependent inhibitory effect on cellular proliferation (Fig. 8). Interestingly, atRA, but not TGF-β, down-

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**FIG. 5.** Effects of growth regulators on cellular proliferation of HPEC and PC-3 cells. A, The growth rates of HPEC cultured with TGF-β, atRA, and IGF-I, for 48 h under serum-free conditions, were investigated by either [3H]thymidine incorporation studies (TGF-β) or by MTS assay (atRA and IGF-I). Both TGF-β and atRA are inhibitory for HPEC, whereas IGF-I is stimulatory for these cells. B, Similarly, the effects of TGF-β and atRA were studied in the malignant PC-3 cells. Only TGF-β was inhibitory for PC-3 cells (atRA did not inhibit the growth of these cells, as judged by either MTS assay or by [3H]thymidine incorporation studies (data not shown)). Results are expressed as means ± SE of three independent studies for HPEC and two independent studies for PC-3 cells. Asterisks indicate statistical significance (compared with control).

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**FIG. 4.** Expression of IGFBP-rP1 and 2 during serial passages of HPEC. A, Western immunoblot studies of conditioned media from HPEC using polyclonal anti-IGFBP-rP1 and 2 antibodies. Three micrograms of total protein were loaded per lane. Both IGFBP-rP1 and 2 concentrations, in conditioned media, increased approximately 6-fold at late, low-replicative passages, compared with early ones. B, Northern blot analyses of same serial passages of HPEC using [32P] radiolabeled IGFBP-rP1 and 2 probes. Twenty micrograms of total RNA were loaded per lane. Concentrations of 18S ribosomal RNA are shown as an internal control for loading. C, Western immunoblot studies of total cell lysates from HPEC using monoclonal anti-p16INK4a antibody. Ten micrograms of total protein were loaded per lane. Note a marked up-regulation (−6-fold) of p16INK4a concentrations at late, low-replicative passages, compared with early ones, as part of the senescence process of HPEC cells.

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rP2 concentrations were studied. TGF-β produced a significant increase (4-fold) in protein levels over nonstimulated values. Similarly, treatment with atRA, under the same experimental conditions, resulted in a 6-fold increase in IGFBP-rP2 concentrations (Fig. 7A). These changes did not, however, follow parallel increases in the concentrations of steady-state mRNA, as the levels of IGFBP-rP2 mRNA did not significantly change on TGF-β treatment and were only increased 2-fold by atRA (Fig. 7B).

In addition, we treated all of the other cell lines with the three above-mentioned growth regulators, but only small responses of IGFBP-rP2 were observed for TGF-β1, atRA, and IGF-I in P69; for TGF-β1 in M12; and for atRA in DU145 cells (data not shown).
regulated the concentrations of secreted IGFBP-rP3 in both P69 and M12 cells (4- and 2-fold, respectively; Fig. 9).

The proliferative actions of IGF-I were also investigated in P69 cells. IGF-I was a potent growth stimulator of these cells (Fig. 8A) but had no observable effects on the concentrations of secreted IGFBP-rP3 protein (Fig. 9).

### Discussion

Recent studies indicate that both IGFBP-rP1 and IGFBP-rP3 are expressed by the prostate epithelium (16, 22). We now report that prostate cells in culture also synthesize another member of the IGFBP superfamily: IGFBP-rP2. Although the biological functions of these proteins in prostate have yet to be defined, they are likely to play a role in the regulation of proliferation of prostate cells, because they are responsive to growth regulators, and, strikingly, IGFBP-rP1 and 2 expression is significantly increased during senescence of the normal prostate epithelium.

IGFBP-rP1 has been shown to be differentially expressed in normal meningeal, breast, and prostate cells, compared with their malignant counterparts (11, 16, 35). In the prostate, it was suggested that IGFBP-rP1 might have an antiproliferative effect, because it was also up-regulated by the epithelial cell growth inhibitors TGF-β1 and atRA (16). Indeed, a more recent study indicates a possible role of IGFBP-rP1 as a tumor suppressor gene for prostate cancer, because over-expression of this gene in the tumorigenic and metastatic P69 and M12 cells caused an antiproliferative effect in vitro and in vivo (36). More difficult to reconcile, however, is a recent report indicating an up-regulation of IGFBP-rP1 during prostate carcinogenesis (37). Because the results by Degeorges et al. are based on immunohistochemistry studies, it is possible that the differences found are attributable to alternative properties or specificity of their antibody; alternatively, different sources of prostatic epithelial cells may vary in these properties.

Further evidence supporting the role of IGFBP-rP1 as a tumor suppressor gene relates to its up-regulation during senescence of normal epithelial cells. Swisshelm et al. (35) have described an enhanced expression of IGFBP-rP1 in senescent human mammary epithelial cells, indicating a possible involvement of this protein in the cell-cycle mechanisms leading to cellular senescence.

HPEC, similarly to human mammary epithelial cells, have a limited life span, undergoing senescence after a limited number of population doublings. Besides a failure to replicate in vitro, senescent HPECs exhibit phenotypic changes, as they become larger and flattened, and induce β-galactosidase activity (data not shown). Consistent with this, a marked up-regulation of the cell-cycle inhibitor p16INK4a was observed at late passages of HPEC, a phenomenon that has been recently reported in these cells (31). Our results are, thus, in
FIG. 7. IGFBP-rP2 is responsive to growth regulators in PC-3. A, Up-regulation of IGFBP-rP2 by the growth inhibitors TGF-β and atRA. Representative Western immunoblot of conditioned media by PC-3 using polyclonal anti-IGFBP-rP2 antibody. Cells were treated with increasing concentrations of TGF-β1 and atRA, for 48 h, in serum-free conditions. A 4-fold increase or greater was seen with both TGF-β1 and atRA in PC-3. B, Northern blot analysis for IGFBP-rP2 indicates only modest increases at the mRNA levels that were statistically different for atRA (graph shows the means ± SE of two independent experiments). Asterisks indicate statistical significance (compared with control).

agreement with observations in human mammary cells, in that the concentrations of both IGFBP-rP1 mRNA and protein were markedly enhanced during replicative senescence of normal prostate epithelial cells. Thus, although the precise role of IGFBP-rP1 in the human prostate has, as yet, to be clarified, our results support the hypothesis that IGFBP-rP1 is a tumor suppressor gene and/or senescence factor.

Several lines of investigation also support roles of IGFBP-rP2 in growth regulation and tumorigenesis. IGFBP-rP2 mediates most of the biological actions of TGF-β in fibroblasts where a novel TGF-β response element in the IGFBP-rP2 promoter has been found (19). Additionally, IGFBP-rP2 is a growth factor for endothelial cells (29, 38). Supporting its role in tumorigenesis, IGFBP-rP2 was expressed by a chondrosarcoma-derived chondrocytic cell line and a fibrosarcoma cell line in vitro (29, 39) and has been found in the stromal component of breast and pancreatic cancer and desmoplastic melanomas (40-42). Additionally, it has been shown that IGFBP-rP2 is up-regulated by TGF-β1 in the breast cancer cell line Hs578T (25) and that IGFBP-rP2 induces apoptosis in the estrogen receptor-positive breast cancer cell line MCF-7 (43). These lines of evidence suggest that IGFBP-rP2 plays a role in modifying the growth of stroma in desmoplastic tumors and in regulating the growth of breast cancer cells.

In this light, our results indicate that IGFBP-rP2 may also regulate the growth of normal and prostate cancer cells, because it is noticeably up-regulated by growth inhibitory factors, such as TGF-β and atRA, in normal and malignant cells, and down-regulated by growth-promoting factors, such as IGF-I, in HPEC. In the prostate, TGF-β is known to limit the proliferation and survival of the normal epithelium, an effect that is lost in malignant cells largely because of a down-regulation of TGF-β receptors during prostate carcinogenesis (44). Retinoids also exert potent inhibitory properties in the normal prostate and have been shown to be present at lower concentration in prostate cancer tissues (33, 45). Thus, both TGF-β and atRA play major roles in controlling the growth and survival of normal prostate cells and probably also in the development of prostate cancer. IGFs regulate also the proliferation of prostate cells (34). Interestingly, the expression of type I IGF-I receptor, which mediates most of the biological activities of IGFs, is markedly down-regulated during carcinogenesis (3). Thus, IGFs seem to play fundamental roles in the regulation of growth and malignant transformation of normal prostate epithelial cells. Because the above-mentioned growth regulators affected the concent-
trations of both IGFBP-rP2 protein and steady-state mRNA in our cultured cells, we speculate that, in the human prostate, IGFBP-rP2 is a downstream effector of growth inhibitors and that IGFBP-rP2 expression must be down-regulated by growth factors to support cell proliferation.

Further observations support the hypothesis that IGFBP-rP2 may act as a growth inhibitor. A striking increase of IGFBP-rP2 was observed during senescence of HPECs, paralleling increases of both IGFBP-rP1 and the cell cycle inhibitor p16INK4A. Thus, IGFBP-rP2 could also be a downstream effector in the mechanisms leading to cellular senescence of normal cells. In summary, IGFBP-rP2 is a potential growth inhibitor that is induced during senescence in prostate epithelial cells and can mediate the effects of growth inhibitors on cell-cycle progression and/or apoptosis in normal and malignant prostate.

IGFBP-rP3 is overexpressed in Wilms tumor, showing an inverse correlation with the concentrations of the tumor-suppressor gene WT1(41), which suggests a potential role of this protein as a protooncogene. Supporting this hypothesis is the observation that overexpression of the aminoterminal truncated NOV molecule was also able to transform chicken embryo fibroblasts (20). More recently, NOV has also shown to induce proliferation of mouse fibroblasts in vitro and to enhance phosphorylation of a 221-kDa protein, suggesting growth-stimulatory properties of this protein through activation of a still unidentified phosphorylated molecule (46).

In the prostate, data on NOV expression are only modest, with one report showing, by RT-PCR, that NOV mRNA is preferentially expressed in PC-3, compared with other normal and malignant prostatic epithelial cells (22). Our results are consistent with this report, and we have extended these findings by demonstrating that IGFBP-rP3 protein was detectable in the condition medium from PC-3 cells. Moreover, by RT-PCR, we were able to detect IGFBP-rP3 mRNA in the immortalized P69 cells and in the malignant M12 and LNCaP cell lines. IGFBP-rP3 protein was also detected in conditioned media from these cells (except LNCaP cells) and in the conditioned medium from the malignant DU145 cells. However, IGFBP-rP3 expression was not detected in normal HPEC cells. Additionally, IGFBP-rP3 expression was suppressed by the growth-inhibitor atRA in P69 and M12 cells. Thus, IGFBP-rP3 expression in prostatic cells is consistent with its potential role as a protooncogene and/or growth factor.

In summary, we report on the expression and regulation of two additional IGFBP-rPs in prostate cells, IGFBP-rP2 and rP3, with results that support the hypothesis that these proteins, like IGFBP-rP1, are involved in the regulation of prostatic cell growth. IGFBP-rP2 may play a role as a growth inhibitor, because its expression is: 1) enhanced during senescence of normal prostate epithelial cells (in a fashion similar to that of IGFBP-rP1); 2) increased by growth inhibitory factors (TGF-β and atRA); and 3) decreased by IGF-I. Conversely, IGFBP-rP3 may act as a growth stimulator for prostate cells, given its preferential expression in malignant cells and its down-regulation by atRA.

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Effects of sodium butyrate on expression of members of the IGF-binding protein superfamily in human mammary epithelial cells

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Abstract

Dietary factors play an important role in both the development and prevention of human cancers, including breast carcinoma. One dietary micronutrient, sodium butyrate (NaB), is a major end product of dietary starch and fiber, produced naturally during digestion by anaerobic bacteria in the cecum and colon. NaB is a potent growth inhibitor and initiates cell differentiation for many cell types in vitro. In this study, we investigated the effects of NaB on three human mammary epithelial cells and regulation of the IGF axis, specifically, IGF-binding protein-3 (IGFBP-3), a known growth regulator in human mammary cells, and IGFBP-related protein 2 (IGFBP-rP2)/connective tissue growth factor.

NaB inhibited DNA synthesis, as measured by [3H]thymidine incorporation, in estrogen-responsive (MCF-7) and estrogen-non-responsive (Hs578T) breast cancer cells, and normal human mammary epithelial cells (HMEC) to a similar degree (up to 90% inhibition at 1–10 mM concentrations). Treatment of cells with NaB induced histone hyperacetylation, suggesting that NaB exerts its biological effects, at least in part, as a histone deacetylase inhibitor in mammary epithelial cells. Treatment of Hs578T cells with NaB caused an induction of apoptotic cell death. NaB treatment resulted in increased levels of p21Waf1/Cip1 mRNA and protein in Hs578T cells and distinct upregulation of p27Kip1 in HMEC, suggesting that NaB activates different genes involved in cell cycle arrest, depending upon the cell type. In the same context, among the IGFBP superfamily members tested, NaB specifically upregulated the expression of IGFBP-3 and IGFBP-rP2. These two proteins are known to be involved in inhibition of mammary epithelial cell replication. Northern blot analysis showed that NaB treatment at 1–10 mM concentrations caused a dose-dependent stimulation of IGFBP-3 mRNA expression in cancerous cells and IGFBP-rP2 mRNA expression in both cancerous and non-cancerous cells. Protein data from Western ligand blot and immunoblot analysis demonstrated parallel results.

In summary, we have demonstrated that NaB (i) uniformly suppresses DNA synthesis in both cancerous and non-cancerous mammary cells, and (ii) upregulates IGFBP-3 and IGFBP-rP2 mRNA and protein levels in cancerous and non-cancerous mammary cells. These results provide the first demonstration that butyrate regulates the IGFBP system in the human mammary system.

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Introduction

Butyric acid is a 4-carbon fatty acid which is the major product from microbial fermentation of dietary fibers in the large intestine (Velásquez et al. 1997). It is a potent growth inhibitor and initiates cell differentiation in several cell types, including breast cancer cells in vitro (Coradini et al. 1997, Velásquez et al. 1997, Gleave et al. 1998, Yamamoto et al. 1998). Although the molecular mechanisms by which butyrate exerts its effects are still unclear, it is known to induce a number of alterations within the nucleus, including histone hyperacetylation (de Haan et al. 1986, Archer & Hodin 1999).

It has been reported that butyrate, as well as trichostatin A (TSA), a specific histone deacetylase inhibitor (Yoshida et al. 1990), modulates specific genes involved in cell cycle regulation and apoptosis. These include the cyclin-dependent kinase (cdk) inhibitor p21Waf1/Cip1 (Archer et al. 1998), p16INK4a (Schwartz et al. 1998), p27Kip1 (Litvak et al. 1998), retinoblastoma protein (Vaziri et al. 1998), cyclin D1 (Lallemand et al. 1996, Slavoshian et al. 1997), Bel-2 and Bax (Mandal & Kumar 1996, Hague et al. 1997). Furthermore, butyrate has also been reported to upregulate the expression of transforming growth factor-β (TGF-β) (Staiano-Coico et al. 1990), which is known to be involved in growth suppression of various...
cancer cells, including breast cancer cells (Knabbe et al. 1987).

Insulin-like growth factors (IGFs) are potent mitogens for several cell types (Macaulay 1992, Resnicoff et al. 1995). The IGF system consists of IGF-I and IGF-II ligands, the transmembrane type I and type II IGF receptors, the IGF-binding proteins (IGFBPs) and IGFBP proteases (Hwa et al. 1999). Recently, the concept of the IGFBP superfamily has been proposed (Baxter et al. 1998); it consists of high affinity IGF binders (IGFBP-1 to -6) and low affinity IGF binders (IGFBP-related proteins (IGFBP-rPs)). The IGFBPs modulate IGF bioactivity, and bind with differential affinities to IGFs in serum and various biological fluids (Kelley et al. 1996, Rajaram et al. 1997). In addition, recent evidence suggests that some IGFBPs may have direct receptor-mediated effects independent of IGFs (Oh et al. 1993). IGFBP-3, for example, has been demonstrated to be an important mediator of other growth inhibitory agents, such as retinoic acid (Gucev et al. 1996), vitamin D (Colston et al. 1998), TGF-β (Oh et al. 1995, Gucev et al. 1996, Rajash et al. 1997), anti-estrogens (Huyhn et al. 1996), tumor necrosis factor-α (Rosen et al. 1998) and p53 (Buckbinder et al. 1995), independently of the IGF signaling system. Furthermore, the importance of IGF-independent biological effects of the IGFBP superfamily, such as IGFBP-3 (Oh et al. 1993), IGFBP-rP1 (Burger et al. 1998) and -rP2 (Hishikawa et al. 1999) on cell replication has been demonstrated in human breast cancer cell systems.

In this study, we have investigated the effects of sodium butyrate (NaB) on members of the IGFBP superfamily in human mammary epithelial cells, using estrogen-responsive (MCF-7) and estrogen-non-responsive (Hs578T) breast cancer cells, and normal human mammary epithelial (HMEC) cells. We report here that NaB upregulates IGFBP-3 and IGFBP-rP2 mRNA and protein in mammary epithelial cells.

Materials and Methods

Materials

NaB, TSA, BSA and 0.4% trypan blue solution were purchased from Sigma Chemical Co. (St Louis, MO, USA). 125I-labeled IGF-I was kindly provided by Diagnostic Systems Laboratories (Webster, TX, USA). Polyclonal anti-IGFBP-3, anti-IGFBP-rP1, anti-IGFBP-rP2 and anti-HEC1 (specific for both human IGFBP-2 and -3) antisera were generated as previously described (Rosenfeld et al. 1990, Oh et al. 1993, Wilson et al. 1997, Yang et al. 1998). Polyclonal anti-IGFBP-5 antibody was purchased from Austral Biologicals (San Ramon, CA, USA). Polyclonal anti-acetyl-lysine, anti-acetylated histone H3 and anti-acetylated histone H4 antibodies were purchased from Upstate Biotechnology (Lake Placid, NY, USA). Polyclonal anti-poly(ADP-ribose)polymerase (PARP) antibody was purchased from Santa Cruz Biotechnology (Santa Cruz, CA, USA). Monoclonal antibodies, anti-p21Waf1/Cip1, anti-p16INK4 and anti-p27Kip1 were purchased from Transduction Laboratories (Lexington, KY, USA), PharMingen (San Diego, CA, USA) and Calbiochem (Cambridge, MA, USA) respectively.

Cell culture

Hs578T estrogen-non-responsive human breast cancer cells and MCF-7 estrogen-responsive human breast cancer cells were purchased from ATCC (Manassas, VA, USA). Both cell lines were maintained in DMEM supplemented with 4.5 g/l glucose, 110 mg/l sodium pyruvate, and 10% fetal bovine serum. HMEC (normal human mammary epithelial cells) were purchased from Clonetics (San Diego, CA, USA), and maintained in mammary epithelium basal medium (MEBM) with growth supplements (bovine pituitary extract (BPE), human epidermal growth factor, insulin, hydrocortisone, gentamicin and amphotericin-B) as directed by the manufacturer.

1H]thymidine incorporation assay

Cells were seeded into 24-well dishes at 37 °C in 5% CO2. At 90% confluence, cells were placed in serum-free media for 12 h, then treated as indicated in the text. After 22 h, 0.1 μCi [1H]thymidine (NEN, Boston, MA, USA) in a volume of 25 μl PBS was added to each well, and the plate was incubated for 4 h at 37 °C. Cells were washed with cold PBS twice, treated with 10% trichloroacetic acid (TCA) for 10 min at −20 °C, washed with 10% TCA followed by 95% ethanol, and lysed with 400 μl 0.25 N NaOH per well. Cell lysates (CL) from each well were transferred to scintillant vials, then 10 ml scintillation fluid with 100 μl 2 N HCl were added to the vials, and the radioactivity was measured in a scintillation counter.

MTS assay

Hs578T cells were seeded into 96-well plates. At 80% confluence, they were incubated for 12 h in serum-free DMEM, then treated with various concentrations of NaB as indicated in the text in serum-free media. After a further incubation of 1–4 days, the MTS reagent (Promega Co., Madison, WI, USA) was added in the ratio recommended by the manufacturer. At 15 min intervals, the absorbance of the formazan product at 490 nm was read with a plate reader (Spectra Shell Reader; SLT Labinstruments GmbH, Austria). On the same plate, cells were dispensed at the differential confluence. These cells were left untreated in serum-free media, and then the MTS reagent was added at the same time as to the butyrate-treated wells. After the reading, cells were trypsinized, then the cell number was counted with a
hemacytometer. Using this method, a linear correlation was obtained between direct cell counts from $1 \times 10^3$ to $1 \times 10^6$ per well and the absorbance ($r=0.85$, $n=70$) (data not shown). The experiment was repeated three times in conditions where the starting untreated control cell number was $8 \times 10^3$ per well, and the absorbance at 490 nm read at 45 min after adding the MTS reagent to the wells was $\sim 0.8$.

**Quantitation of apoptosis**

To quantitate apoptotic cell death, the Cell Death Detection ELISA$^\text{PLUS}$ kit (Roche Molecular Biochemicals, Mannheim, Germany), which measures cytoplasmic histone-bound DNA fragments produced during apoptotic DNA fragmentation, was used (Mandal & Kumar 1996). Hs578T cells were seeded into a 96-well plate at 80% confluence, they were incubated for 12 h in serum-free DMEM, then treated with various concentrations of NaB in serum-free media. After 72 h, cytoplasmic extracts were made from attached cells by adding 100 μl lysis buffer to $5 \times 10^4$ cells per well. Supernatant (20 μl from 100 μl) was analyzed in the ELISA, as directed by the manufacturer’s protocol. Briefly, the samples were placed into streptavidin-coated multi-well plates, a mixture of anti-histone-biotin and anti-DNA-peroxidase was added and incubated. The ELISA was developed with peroxidase substrate, and the absorbance at 405 nm was measured against 490 nm as a reference wavelength. The experiment was performed from duplicate samples for each data point generated, and was repeated twice independently.

**Preparation of conditioned media (CM) and CL**

Cells were seeded in 12-well plates. At 95% confluence, they were incubated for 12 h in serum-free DMEM (Hs578T and MCF-7) or supplement-free MEBM with added BPE (HMEC), then treated as indicated in the text in serum- or supplement-free media. CM samples were collected after 72 h and centrifuged at 1000 g for 10 min CA, USA), and then treated as indicated in the text. Total RNA was isolated from cells per well. Supernatant (20 μl from 100 μl) was analyzed in the ELISA, as directed by the manufacturer’s protocol. Briefly, the samples were placed into streptavidin-coated multi-well plates, a mixture of anti-histone-biotin and anti-DNA-peroxidase was added and incubated. The ELISA was developed with peroxidase substrate, and the absorbance at 405 nm was measured against 490 nm as a reference wavelength. The experiment was performed from duplicate samples for each data point generated, and was repeated twice independently.

**Total RNA isolation and Northern blot analysis**

Cells were grown in 6-well plates until 95% confluent. Cells were then incubated in serum-free media for 12 h, then treated for 18 h in serum- or supplement-free media as indicated in the text. Total RNA was isolated from duplicate wells, using the RNeasy Kit (Qiagen, Valencia, CA, USA), and 5 μg total RNA per sample were separated on a 15% SDS-PAGE gel, transferred onto nitrocellulose membranes and probed with 2'0 x 10$^6$ c.p.m. $^{3}P$-labeled IGF-I. The membranes were washed, dried and exposed to film (Kodak BioMax MS, Eastman Kodak Co., Rochester, NY, USA) for 12-18 h.

**Western immunoblot analysis**

For IGFBP-3, IGFBP-rP1 and IGFBP-rP2 detection, CM samples were separated on non-reducing 12 or 15% SDS-PAGE. For p21$^{\text{Waf1/Cip1}}$ detection, CL samples were separated on reducing 15% SDS-PAGE. Proteins were electrotransferred onto nitrocellulose, and membranes were blocked with 5% non-fat dry milk/TBS-T for 1 h at room temperature, then incubated in 1:3000 dilution of primary antibody at 4 °C overnight. Immunoreactive proteins were detected using enhanced chemiluminescence (NEN, Boston, MA, USA).

**Densitometric analysis**

To quantitate the relative induction after Western blot analyses or Northern blot analyses, densitometric measurement was performed by using a GS-700 imaging densitometer with Multi-Analyst software (Bio-Rad).
Figure 1 Effect of NaB (A) and TSA (B) on DNA synthesis in Hs578T, MCF-7 and HMEC cells. Serum-starved cells (85–90% confluent) were incubated in basal medium for 22 h in the absence or presence of various concentrations of NaB or TSA as indicated. [3H]Thymidine was added, and the incubation was continued for another 4 h. The incorporation of [3H]thymidine was determined relative to control cells incubated without the addition of these reagents. Results represent the average of two independent experiments each performed in triplicate.

Results

NaB treatment inhibits DNA synthesis and causes histone hyperacetylation in human mammary epithelial cells

Since it has been demonstrated that treatment of NaB resulted in growth inhibition in a variety of cell systems in vitro, we first examined the effect of NaB on DNA synthesis in normal (HMEC) and cancerous (Hs578T, MCF-7) human mammary epithelial cells, using the [3H]thymidine incorporation assay. NaB suppressed DNA synthesis in both cancerous and non-cancerous human mammary epithelial cells in a dose-dependent manner, with 90–100% inhibition at NaB concentrations of 5 mM (Fig. 1A). One major function of NaB is inhibition of histone deacetylase activity, resulting in histone hyperacetylation. In order to determine whether NaB-induced inhibition of DNA synthesis might be due to histone hyperacetylation, we treated the cells with TSA, a specific histone deacetylase inhibitor, and compared its effect on DNA synthesis. As shown in Fig. 1B, TSA also suppressed DNA synthesis, with 90–100% inhibition at TSA concentrations of 100 nM in these cells. This suggests that butyrate-induced suppression of DNA synthesis in these human mammary cells may involve histone hyperacetylation, as is shown by Western immunoblot with an anti-acetyl-lysine antibody (Fig. 2). The CL 24 h after treatment with NaB showed an increase of acetylated proteins in a dose-dependent manner, as indicated by the appearance of 11 and 16 kDa bands, which were identified as H4 and H3 histones respectively by Western immunoblot with specific antibodies (data not shown). Histone hyperacetylation was similarly demonstrated by treatment with TSA (Fig. 2). Interestingly, only the 11 kDa band was seen at concentrations over 100 nM.

Effects of NaB on reducing cell number and the induction of apoptosis in human mammary epithelial cells

As NaB inhibits DNA synthesis almost completely by 24 h treatment, cell viability over 4 days after NaB treatment in Hs578T cells was then studied. The MTS assay was used, as described in Materials and Methods, as a marker of relative viable cell number. A progressive reduction in cell number by MTS assay was observed from day 2 onwards using 10 mM NaB, and from day 3 onwards after 5 mM NaB treatment (Fig. 3). In parallel wells, when cell numbers were counted after trypsinizing by direct visualization using a hemacytometer, the attached cell numbers were reduced over the same time course and concentrations of NaB as was detected in the MTS assay above (data not shown). Using trypan blue exclusion during hemacytometer counting, when counted at the same time point, the same number of attached cells were shown to take up trypan blue in the NaB-treated wells compared with the control wells, throughout the full 4 days of the study (data not shown). This trypan blue staining pattern indicates that cell plasma membrane integrity was maintained in the attached cells after NaB treatment compared with the control.

As a reduction in viable cell number by NaB was occurring over time, the possibility that apoptosis was being induced by NaB in the mammary epithelial cells was then addressed. Two independent methods of analysis were used to detect apoptosis: first, nuclear enzyme cleavage, and secondly, DNA fragmentation. The nuclear enzyme PARP is proteolytically cleaved during apoptosis in vitro in many cell types, including breast cancer cells (Kaufmann et al. 1993). Figure 4A shows Western immunoblot with an anti-PARP antibody after treatment of Hs578T and MCF-7 cells with NaB followed by

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Figure 2 Effect of NaB on protein acetylation. Serum-starved cells were treated for 24 h with various concentrations of NaB as indicated. CL harvested from duplicate wells within each experiment were pooled, and 20 μg protein per lane were loaded onto 15% SDS-PAGE under reducing conditions. Gels were immunoblotted with an anti-acetyl-lysine antibody as described in Materials and Methods. The immunoblots of CL treated with 100 nM TSA are shown on the right. Molecular mass markers are also shown. The results are representative of two independent experiments.

Figure 3 Growth inhibition by NaB. Hs578T cells were seeded into 96-well plates, then serum-starved cells were treated with various concentrations of NaB as indicated. On subsequent days, cell proliferation was measured using the MTS assay. A dose-response effect with added NaB is seen over 4 days. Results are expressed in absorbance readings at 490 nm as percent of the untreated controls ± S.E. (n=16).

post-treatment, NaB induced apoptosis in a dose-dependent manner (Fig. 4B). This sensitive assay demonstrated apoptotic effects initially commencing at 1 mM NaB. Taken together, the two methods of measuring apoptosis show that NaB induces apoptosis in the cells studied, in a more delayed time course compared with the earlier effects on [3H]thymidine incorporation. The apoptosis induced by NaB would be expected to reduce the viable cell number, which was observed, as described earlier.

NaB upregulates expression of p21Waf1/Cip1 mRNA and protein levels in human mammary epithelial cells

As the major mechanism for butyrate-induced growth inhibition in various cell systems is known to be through upregulation of cdk inhibitors, in particular p21Waf1/Cip1, the induction of p21Waf1/Cip1 in mammary epithelial cells was next investigated. Figure 5A is a Northern blot of p21Waf1/Cip1 from Hs578T, MCF-7 and HMEC cells treated with or without NaB, p21Waf1/Cip1 mRNA expression was upregulated in all three cell lines, and was most marked in Hs578T cells. As shown in Fig. 5B, an upregulation of p21Waf1/Cip1 protein levels occurred in these cells, with the greatest increase in p21Waf1/Cip1 observed in Hs578T cells, which parallels the mRNA data. As the degree of induction was different between Hs578T cells and the other two cell lines, the NaB effect on cdk inhibitors p27Kip1 and p16INK4 was further investigated. A distinct upregulation of p27Kip1 by NaB treatment (a 2.5-fold increase at 5 mM treatment) was reproducibly seen in HMEC cells, but not in the cancerous cell lines (Fig. 5D). We did not detect p16INK4 in Hs578T and MCF-7 cells, and only a slight induction of this protein was seen in HMEC cells. Taken together, these data suggest that differential cdk inhibitors are induced by NaB treatment, in a cell-type-dependent manner.

Butyrate upregulates IGFBP-3 mRNA and protein levels in cancerous, but not in non-cancerous mammary cells

To investigate any correlation between the effect of NaB and the regulation of IGFBP system, we first examined IGFBP-3, a known growth suppressor in human mammary cells. Northern blotting was firstly performed to measure steady-state mRNA levels. Figure 6A shows the time-course effect of NaB treatment on steady-state levels of IGFBP-3 mRNA in Hs578T cells. NaB induced the expression of IGFBP-3 mRNA in a time-dependent manner, with increases first detectable at 6 h after treatment, and with a 2.5-fold increase after treating cells with 5 mM NaB for 24 h, whereas in the non-cancerous HMEC cells, only a slight induction (<1.2-fold) was observed (Fig. 6B). In MCF-7 cells, IGFBP-3 mRNA was not detected in these analyses.

Figure 4 Effect of NaB on apoptosis. (A) Immunoblot analysis of PARP in Hs578T and MCF-7 CL obtained during treatment with NaB (day 1 and day 3). The 85 kDa fragment characteristic of apoptosis is shown. The data shown are representative of two independent experiments. (B) Induction of apoptosis in Hs578T cells treated with increasing doses (0-10 mM) of NaB for 72 h. Cytoplasmic extracts were prepared from attached cells, and apoptotic cell death was quantitated by ELISA measuring cytoplasmic histone-bound DNA complexes characteristic of apoptosis, as described in Materials and Methods. Results are expressed as mean absorbance ± s.d. of two independent experiments performed in duplicate (n=4).
The CM were then examined for changes in IGFBP-3 protein levels after NaB treatment. In order to ascertain a suitable time point to collect CM samples, the time-course induction of media IGFBP-3 protein in Hs578T cells was studied by Western ligand blot analysis, using $^{125}$I-labeled IGF-I as the ligand, as described in Materials and Methods. The IGFBP-3 level in the CM was detectably increased at 24 h, even after 1 mM NaB, was further increased at 48 h,
these IGFBPs was confirmed by Western immunoblot using IGFBP-3, -2 and -5 specific antibodies, showing the 42–46 kDa doublet bands to be IGFBP-3, the broad 29–36 kDa bands to contain IGFBP-2 and -5, and the 24 kDa band to be IGFBP-4 (not shown). In HMEC cells, the Western ligand blot did not reveal an IGFBP-5 band, whereas the Western immunoblot with IGFBP-5 antibody revealed a low intensity band of the predicted molecular mass for IGFBP-5 (data not shown) (Adamo et al. 1992, Sheikh et al. 1992). IGFBP-3 protein levels were upregulated in both Hs578T (2-1-fold over the control) and MCF-7 (12-6-fold over the control) cells, each after 5 mM NaB treatment (Fig. 7B). Further Western immunoblotting analyses demonstrated no detected IGFBP-3 fragments in all samples tested in these cells (data not shown). As the basal level of IGFBP-3 in MCF-7 cells was nearly undetectable, the induction of IGFBP-3 protein by NaB was more conspicuous in this cell line. In contrast, only slight upregulation of IGFBP-3 (<1-5-fold) was observed in HMEC cells, mirroring the mRNA data (shown earlier in Fig. 6B). Levels of IGFBP-2/-5 and -4 showed no significant change up to 10 mM NaB treatment, after accounting for effects of NaB on cell number (not shown). The effect of TSA treatment on IGFBP-3 protein levels by Western ligand blot and immunoblot was also studied. TSA treatment of both Hs578T and MCF-7 cells caused a dose-dependent increase in IGFBP-3 protein levels (not shown), suggesting that NaB-induced upregulation of IGFBP-3 is, at least in part, through histone hyperacetylation. This effect was not seen in the HMEC cells (data not shown).

NaB upregulates IGFBP-rP2 mRNA and protein expression in both cancerous and non-cancerous mammary cells

The induction of the low affinity IGF binders, especially IGFBP-rP2, was then investigated, as this protein also has recently been shown to have a growth suppressive effect in human mammary cells (Hishikawa et al. 1999). The effect of NaB treatment on IGFBP-rP2 mRNA expression was potent, particularly in Hs578T cells, where effects occurred with 0.5 mM NaB. As seen in Fig. 8A, a 10-fold induction of IGFBP-rP2 by 5 mM NaB treatment was observed in Hs578T cells, whereas a maximal 2- to 3-fold induction occurred in HMEC cells. IGFBP-rP2 mRNA was not detected in MCF-7 cells.

Western immunoblot analysis against IGFBP-rP2 and -rP1 was then performed. As shown in Fig. 8B, NaB highly upregulated IGFBP-rP2 protein levels in all three cell lines in a dose-dependent manner (>10-fold at 10 mM NaB over the control in all three cell lines). Increases in media IGFBP-rP2 were initially detectable within 24 h, even after only 1 mM NaB (data not shown). In contrast to the effects on IGFBP-rP2, the IGFBP-rP1 band intensity was not increased by NaB treatment. The apparent reduction in IGFBP-rP1 by NaB treatment...
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**A**

IGFBP-3 protein secretion

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**B**

**NaB treatment**

- Hs578T
  - IGFBP-3
  - IGFBP-2, 5
  - IGFBP-4

- MCF-7
  - IGFBP-3
  - IGFBP-2, 5
  - IGFBP-4

- HMEC
  - IGFBP-3
  - IGFBP-2
  - IGFBP-4

Figure 7 Effect of NaB on IGFBP production in Hs578T, MCF-7 and HMEC cells analyzed by Western ligand blotting. Serum-starved cells were treated with various concentrations of NaB. CM harvested from duplicate wells within each experiment were pooled, size-fractionated using 12% SDS-PAGE under non-reducing conditions, electroblotted onto nitrocellulose membranes, and treated with \( ^{125}I \)-labeled IGF-I, as indicated in Materials and Methods. (A) Time-course expression of IGFBP-3 protein in Hs578T cells. Serum-starved cells were treated with or without NaB as indicated and Western ligand blot was performed over 4 days. (B) Representative Western ligand blots using 72 h CM in Hs578T, MCF-7 and HMEC cells. The data shown were derived from at least three independent experiments.

compared with control, especially in Hs578T CM, was 52% on average using densitometric analysis at day 3 after 10 mM NaB. This reduction in IGFBP-rP1 could be fully accounted for by considering the effects of NaB on cell number, as shown earlier in Fig. 3, where 40–50% of the cells are non-viable by this time of NaB treatment compared with control. IGFBP-rP1 was not detected in MCF-7 cells. These results show that IGFBP-rP2 mRNA and protein are specifically induced by NaB in both cancerous and non-cancerous breast epithelial cells.

**Discussion**

In this study, NaB effects in the human mammary cell system including cancerous and non-cancerous cells, was
investigated, in order to obtain a greater understanding of the cellular mechanism of action of NaB in this cell type. NaB was found to cause an initial inhibition in new DNA synthesis, followed by apoptotic changes, and a reduction in viable cell number. Subsequently, the cdk inhibitors studied showed some cellular specificity in their

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**Figure 8** Effect of NaB on (A) IGFBP-rP2 mRNA expression, and (B) IGFBP-rP2 and rP1 protein expression in Hs578T, MCF-7 and HMEC cells. (A) Representative Northern blots. Total RNA was harvested at 18 h post-treatment, and 5 µg per lane were electrophoresed. The membrane was subsequently probed with labeled cDNA fragments of IGFBP-rP2. 28S rRNA methylene blue membrane staining is presented as an indicator of equal loading. Densitometric analysis adjusted for 18S and 28S rRNA is also shown. Each result represents two independent experiments. In MCF-7 cells, IGFBP-rP2 mRNA was not detected in 10 µg total RNA. (B) Representative Western immunoblot analysis. Serum-starved cells were treated with various concentrations of NaB for 72 h. CM harvested from duplicate wells within each experiment were pooled, size-fractionated using 12% (for IGFBP-rP2) or 15% (for IGFBP-rP1) SDS-PAGE under non-reducing conditions, electroblotted onto nitrocellulose membranes, and treated with appropriate antibodies, as indicated in Materials and Methods. IGFBP-rP1 was not detected in the CM of MCF-7 cells.
upregulation by NaB. IGFBP superfamily members known to induce growth inhibition and apoptosis in breast epithelial cells were also upregulated by NaB.

A schematic summary of our observations in Hs578T cells is shown in Fig. 9. We first studied the general effects of NaB on cell growth regulation. Butyrate induces diverse and reversible biological effects on cell differentiation, apoptosis and cell growth in vitro (Pouillart 1998). In colon epithelial cells, butyrate is known to exert paradoxical effects, with induction of proliferation in normal cells and growth inhibition in neoplastic phenotypes (Hassig et al. 1997, Archer & Hodin 1999). In the mammary system studied here, butyrate inhibits DNA synthesis in both normal and cancerous cells as measured by [3H]thymidine incorporation at 24 h after treatment. Also, as shown in Fig. 9, after 24 h of NaB treatment at concentrations over 1 mM, cell numbers became reduced in a dose- and time-dependent manner. This reduction in cell number was explained, at least in part, by NaB-induced apoptosis. The observed induction of apoptosis remains consistent with previous reports of NaB-induced apoptosis in other cell systems (Hague et al. 1993, Carducci et al. 1996, Coradini et al. 1997). In summary, NaB was found to induce both inhibition of DNA synthesis and programmed cell death in Hs578T cells.

It is known that butyrate induces a variety of changes within the nucleus (Siavoshian et al. 1997, Schwartz et al. 1998), including histone hyperacetylation, especially of H3 and H4 (Archer & Hodin 1999), and DNA methylation (de Haan et al. 1986). Previous studies indicate that the effect of butyrate and other histone deacetylase inhibitors on cells closely corresponds to the effect of p21\(^{\text{Waf1/Cip1}}\) expression in the regulation of G1 (Hunter & Pines 1994), S (Ogryzko et al. 1997) and G2 (Coradini et al. 1997, Lallemand et al. 1999) phases of the cell cycle (Hassig et al. 1997). The major mechanism for butyrate-induced cell cycle arrest is reported to be through upregulation of p21\(^{\text{Waf1/Cip1}}\) in various cell systems, including hepatocellular carcinoma cells (Yamamoto et al. 1998), colon cancer cells (Nakano et al. 1997, Siavoshian et al. 1997, Archer et al. 1998, Litvak et al. 1998), prostate cancer cells (Huang et al. 1999) and breast cancer cells (Lallemand et al. 1999). In contrast, Vaziri et al. (1998) demonstrated p21\(^{\text{Waf1/Cip1}}\)-independent G1 cell cycle arrest by butyrate in 3T3 fibroblasts. In our present studies, the prominent induction of p21\(^{\text{Waf1/Cip1}}\) was observed in estrogen-non-responsive Hs578T cancer cells, whereas modest induction was observed in estrogen-responsive MCF-7 cancer cells and non-cancerous HMEC cells. Interestingly, butyrate upregulated p27\(^{\text{Kip1}}\) expression only in HMEC cells. The p16\(^{\text{Ink4a}}\) protein was detected only in HMEC cells and no further regulation was observed after butyrate treatment. Taken together, these data imply that the biological function of butyrate is mediated through more than one mechanism, even in butyrate-induced cell cycle arrest, suggesting that butyrate possesses multifunctional mechanisms of action. This is the first demonstration that butyrate upregulates members of the IGFBP superfamily in human mammary cells. One previous report showed that only IGFBP-2 was upregulated by butyrate in colon cancer cells (Nishimura et al. 1998). In contrast, our studies demonstrate that among the six IGFBPs and IGFBP-rP-1 and -2, only IGFBP-3 and IGFBP-rP2 are upregulated by butyrate in cancerous and non-cancerous mammary epithelial cells. Recent evidence has suggested that, in addition to modulating the access of IGFs to their receptors, IGFBP-3 has the ability to suppress proliferation in various cell systems, including...
human breast cancer cells, working through IGF-independent mechanisms (Valentini et al. 1995, Oh 1998). Further studies have demonstrated that IGFBP-3 directly induces apoptosis through an IGF-independent pathway in PC-3 human prostatic adenocarcinoma cells (Rajah et al. 1997) as well as in other cell systems (Nickerson et al. 1997). We, therefore, hypothesized that IGFBP-3 may be a major downstream effector of growth inhibitory and apoptosis-inducing agents.

Indeed, NaB treatment significantly upregulated IGFBP-3 steady-state mRNA and protein levels in a time- and dose-dependent manner in Hs578T and MCF-7 human breast cancer cells, while levels of IGFBP-2 and -4 were unaffected. As summarized in Fig. 9, we observed that the mRNA induction had already started at 6 h after treatment and peaked by 24 h after treatment, while the presence of the protein in CM was barely detectable at the 24 h time point and gradually increased up to day 3. Notably, both the mRNA and protein induction by NaB were detectable earlier than the reduction in cell number and apoptosis, consistent with a specific regulation of these IGFBP superfamily members by NaB rather than effects on these IGFBPs occurring secondary to cell death. In contrast to effects seen in these cancer cell lines, the NaB regulation of IGFBP-3 in normal human mammary epithelial cells was less marked. Additionally, treatment with TSA gave similar results, indicating that histone hyperacetylation may be involved. The differential regulation of IGFBP-3 by NaB in cancerous versus non-cancerous cells may point to an important mechanism for growth inhibition by IGFBP-3 in cancer.

It has been demonstrated that TGF-β, which is a potent growth suppressing factor in human breast cancer cells (Zugmaier et al. 1989, Oh et al. 1995), induces expression of IGFBP-rP2 mRNA and protein levels (Yang et al. 1998). In addition, recent studies have demonstrated that IGFBP-rP2 has a direct apoptotic effect in MCF-7 cells (Hishikawa et al. 1999). Our present data show that IGFBP-rP2 was significantly upregulated by NaB treatment in a time- and dose-dependent manner in both cancerous and non-cancerous human mammary epithelial cell lines, suggesting that IGFBP-rP2 may also play a role in the bioactivity of butyrate on cells. In Hs578T cells, the induction of IGFBP-rP2 mRNA was already marked at 0.5 mM, while the protein levels were significantly increased at 5 mM. One potential explanation for this dose discrepancy is that some post-transcriptional modification by NaB of IGFBP-rP2 may also be occurring.

On the basis of this work, it can be speculated that the IGFBP superfamily members, IGFBP-3 and IGFBP-rP2, participate in butyrate-induced sequential cell growth inhibition, particularly in the later event of apoptosis. Detailed studies involving regulation of IGFBP-3 and IGFBP-rP2 bioactivity following butyrate treatment in breast epithelial cells are now required to formally address this issue. It is hoped that exploration of butyrate-induced biological effects subsequent to the cell cycle arrest in this cell system and the investigation of the interaction between butyrate and the IGF axis will lead to a more complete understanding of the complex mechanisms of cell growth control, and to the development of better therapeutic reagents.

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ABSTRACT

Expansion of extracellular matrix with fibrosis occurs in many tissues as part of the end-organ complications in diabetes, and advanced glycosylation end products (AGE) are implicated as one causative factor in diabetic tissue fibrosis. Connective tissue growth factor (CTGF), also known as insulin-like growth factor-binding protein-related protein-2 (IGFBP-rP2), is a potent inducer of extracellular matrix synthesis and angiogenesis and is increased in tissues from rodent models of diabetes. The aim of this study was to determine whether CTGF is up-regulated by AGE in vitro and to explore the cellular mechanisms involved. AGE treatment of primary cultures of nonfetal human dermal fibroblasts in confluent monolayer increased steady state messenger RNA (mRNA) levels in a time- and dose-dependent manner. In contrast, mRNAs for other IGFBP superfamily members, IGFBP-rP1 (mac 25) and IGFBP-3, were not up-regulated by AGE. The effect of the AGE BSA reagent on CTGF mRNA was due to nonenzymatic glycosylation of BSA and, using neutralizing antisera to AGE and to the receptor for AGE, termed RAGE, was seen to be due to late products of nonenzymatic glycosylation and was partly mediated by RAGE. Reactive oxygen species as well as endogenous transforming growth factor-β1 could not explain the AGE effect on CTGF mRNA. AGE also increased CTGF protein in the conditioned medium and cell-associated CTGF. Thus, AGE up-regulates the profibrotic and proangiogenic protein CTGF (IGFBP-rP2), a finding that may have significance in the development of diabetic complications. (Endocrinology 142: 1760-1769, 2001)

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MULTIPLE MECHANISMS have been described by which chronic hyperglycemia might contribute to the pathological end-organ complications that occur in diabetes mellitus. These include direct effects of elevated glucose on cells, hyperosmolality, oxidant stress, and nonenzymatic glycosylation (1, 2). Advanced glycosylation end products (AGE) are biochemical end products of nonenzymatic glycosylation that are formed irreversibly (3). AGE is elevated in serum (4) and in many tissues in patients with diabetes (5), including skin (6), and has the ability to covalently cross-link and biochemically modify protein structure and affect protein function (7). Additionally, in recent years cell surface receptors for AGE have been identified (8), and postreceptor signaling pathways are being defined (9, 10). Through an AGE receptor-dependent mechanism, AGE induction of cytokines and growth factors has been implicated in contributing to end-organ changes that occur in tissues with diabetes (11–13).

Pathological hallmarks in most tissues where diabetes complications occur include expansion of extracellular matrix (ECM) and angiogenesis (1). The ECM expansion has been proposed to be due to a combination of increased ECM production (14) (15) and biochemically modified matrix, with a reduction in ECM breakdown (16). Connective tissue growth factor (CTGF), also known as insulin-like growth factor (IGF)-binding protein-related protein-2 (IGFBP-rP2) (17), is a potent inducer of ECM in fibroblasts (18, 19) and a potent angiogenic factor (20, 21). A potential role for CTGF in fibrotic disease states is increasingly being described (22–24), suggesting that CTGF may be a mediator of ECM expansion and fibrosis in diabetes. The aim of this study was to determine whether CTGF is up-regulated by AGE and subsequently to explore the cellular mechanism(s) that might be responsible for this effect.

Materials and Methods

Reagents

Polyclonal anti-IGFBP-rP2 (CTGF) antibodies (8799 and 8800) were generated in New Zealand White rabbits, as previously described (25). The anti-AGE antibody, which neutralizes the activity of AGE BSA (26),
was a gift from Dr. H. Miyata, Kissei Pharmaceutical Co. Ltd. (Nagano, Japan). The antihuman polyclonal antibody generated in rabbits against the receptor for AGE (RAGE) was provided as an IgG fraction (gift from Dr. A.M. Schmidt, Columbia University, New York, NY). This antibody binds quantitatively to RAGE, recognized by the antibody (27, 28). The transforming growth factor-β1 (TGFβ1) affinity-purified IgG antibody generated in chickens that neutralizes TGFβ1 bioactivity was purchased from R&D Systems, Inc. (Minneapolis, MN). Nonimmune rabbit IgG, β-glucose, glycolaldehyde, BSA (fraction V, fatty acid and endotoxin free), and aminoguanidine were purchased from Sigma (St. Louis, MO). TGFβ1 was purchased from Austral Biologicals (San Ramon, CA) and Transforming growth factor-β1 (TGFβ1) affinity-purified IgG antibody generated in chickens that neutralizes TGFβ1 bioactivity was purchased from R&D Systems, Inc. (Minneapolis, MN). Nonimmune rabbit IgG, β-glucose, glycolaldehyde, BSA (fraction V, fatty acid and endotoxin free), and aminoguanidine were purchased from Sigma (St. Louis, MO). TGFβ1 was purchased from Austral Biologicals (San Ramon, CA) and from the forearm of a 70-yr-old male) and A305 (newborn foreskin fibroblasts), were gifts from Dr. S. Goldstein, Memorial Veteran's Hospital (Little Rock, AR). These cells were maintained in DMEM with 15% FBS.

Cell treatment

After trypsinization, cells were grown in 12-well plates for 5 days in their respective media with FBS until they were confluent. For experiments requiring the use of blocking antibodies to RAGE and AGE, cells were grown in 24-well plates under the same conditions. Cells were then incubated in their respective serum-free medium for 16 h and then treated with additions on day 0 under serum-free conditions, using fresh media. Unless otherwise indicated in the text, the conditioned media were not changed after adding the treatments. When cells were transiently treated for 8 h with reagents, they were washed with PBS, and fresh serum-free medium with 0.05% BSA was added. Cell lyses and conditioned media were harvested up to 3 days after treatments. For experiments involving the use of blocking antibodies or antioxidants, cells were preincubated with the antibody or reagent for 2 h under serum-free conditions before adding AGE or control BSA directly to the medium.

Total RNA isolation and analysis by quantitative real-time RT-PCR

Total RNA was isolated from duplicate wells using the RNaseasy Mini kit from Qiagen (Valencia, CA) and was then analyzed by quantitative real-time PCR using an ABI Prism 7700 Sequence Detection System (PE Applied Biosystems, Foster City, CA). This system is based on the ability of the 5'-nucleotide activity of Tag polymerase to cleave a nonextendable dual labeled fluorogenic hybridization probe during the extension phase of PCR. The following sequence-specific primers and probes for human CTGF, IGFBP-rP1, and 18S ribosomal RNA were designed using Primer Express Software 1.0 (PE Applied Biosystems): for CTGF: forward, 5'-GCGACGCTCTGATGCA-3'; reverse, 5'-CG-GCACAGCTCTGATGCA-3'; and probe, 5'-FAM-TCTGCACCGAGTTG-TA-3'; for IGFBP-rP1: forward, 5'-GCCACAGCTCTGATGCA-3'; reverse, 5'-GCGACGCTCTGATGCA-3'; and probe, 5'-FAM-TCTGCACCGAGTTG-TA-3'; for 18S: reverse, 5'-GGATTAGTATGATGCG-3'; and probe, 5'-FAM-TCTGCACCGAGTTG-TA-3'. The primer pairs and probes were used in 2-μL reactions using 0.625 μL of cDNA from each treatment. The Ct values were calculated using the comparative threshold cycle number for each sample. The total fluorescence signal was normalized to 18S ribosomal RNA and related to relevant controls, as indicated in the text.

Preparation of conditioned media and cell lyses

Cell lysate samples were harvested after treatment, by washing cells with PBS, then adding 100 μL cold RIPA lysis buffer (20 mm Tris (pH 8.0), 150 mm NaCl, 1% Nonidet P-40, 0.5% sodium deoxycholate, and 0.1% SDS) plus a protease inhibitor cocktail (Roche Molecular Biochemicals, Mannheim, Germany) directly to each well. Plates were rocked for 10 min at 4°C. The supernatants from duplicate wells within each experiment were pooled and stored at −20°C until analysis. The total protein concentration was determined for each sample by use of the DC Protein Assay reagent (Bio-Rad Laboratories, Inc., Hercules, CA). Then 20 μg total protein were loaded per lane for SDSPAGE analysis, and 5 μg total protein were added to each ELISA well for CTGF quantification.
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**Materials and Methods.**

Soaked AGE BSA at 100 μg/ml was added to duplicate wells of confluent human fibroblasts (CRL-2097 cells) under serum-free conditions, and total RNA was collected at the time points shown. Control BSA at the same concentration was added to other wells, also in duplicate. CTGF mRNA was then determined by quantitative RT-PCR in triplicate for each sample, as described in Materials and Methods. The CTGF mRNA level is expressed in arbitrary units normalized to 18S. A, Time course up to 24 h. B, Time course from 1–3 days. C, Treatment for 8 h only with AGE BSA or control BSA at 100 μg/ml, followed by RNA analysis on days 1–3. Data in A–C are the mean ± 1 sd from three independent experiments.

**Western immunoblot analysis.**

Conditioned medium samples were separated on 15% nonreducing SDS-PAGE. Proteins were electrotransferred onto nitrocellulose, and membranes were blocked with 5% nonfat dry milk/TBS with 0.1% (vol/vol) Tween 20 for 1 h at 22 C, then incubated in IGFBP-rP2 (CTGF) antiserum at 1:100 dilution at 4 C overnight. After incubation of membranes with a horseradish peroxidase (HRP)-labeled secondary antibody for 1 h at 22 C, immunoreactive proteins were detected by use of enhanced chemiluminescence (NEN Life Science Products, Boston, MA).

**CTGF (IGFBP-rP2) ELISA.**

The anti-IGFBP-rP2 (CTGF) antibody (8800) (25) was biotinylated by incubating protein A affinity-purified 8800 (0.8 μg) with 150 μg sulfo-NHS-LC biotin (Pierce Chemical Co., Rockford, IL) for 2 h at 22 C, followed by separation from unreacted biotin through a size-fractionation and desalting column with PBS as buffer according to the manufacturer’s instructions. Affinity-purified 8800 antibody (600 ng/well) in 10 mm sodium carbonate, pH 9.6, was adsorbed to 96-well immunoplates (Nalge Nunc International, Rochester, NY) by a 20-h incubation at 4 C. The unbound antibody was removed, and the wells were blocked by incubation with PBS and 0.1% (vol/vol) Triton X-100 (buffer A*) containing 10 g/liter BSA for 2 h at 37 C, then washed four times with buffer A. Purified intact recombinant human (rh) IGFBP-rP2 (CTGF) in buffer A and 1 g/liter BSA was used to generate standard curves. Standard and samples (100 μl/well) were incubated in duplicate at 4 C for 20 h. The plate was washed, then incubated with biotinylated IGFBP-rP2 (CTGF) antibody (80 ng/well) for 20 h at 4 C. After washing, the plate was incubated with streptavidin-HRP (1:50) for 30 min at 22 C, followed by incubation of substrate [0.1 g/liter 3,3',5,5'-tetramethylbenzidine in 0.2 M sodium acetate (pH 6) containing 0.06% (wt/wt) H2O2] for 30 min at 22 C. The reaction was stopped by the addition of 2 M H2SO4, and the absorbance was measured at 450 nm using a microplate reader. The intersample coefficient of variation was 8.1% for the middle concentration (10 ng well) of rhIGFBP-rP2 (CTGF) standard used. No cross-reactivity was detected with 1 μg/well purified rhIGFBP-3, rhIGFBP-rP1 (mac 25), or rhIGFBP-rP3 (Nov H; not shown).

**CTGF (IGFBP-rP2) cell association assay.**

To determine whether increases in rhIGFBP-rP2 (CTGF) in the whole cell lysates after AGE treatment are due to increases in rhIGFBP-rP2 on
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A: Control BSA AGE BSA

CTGF (IGFBP-rP2) mRNA /18S

B: Control BSA AGE BSA TGF-β1

IGFBP-rP1 (mac-25) CTGF (IGFBP-rP2) IGFBP-3

Respective mRNA /18S

Fig. 3. Generalizability of the AGE BSA effect on CTGF (IGFBP-rP2) mRNA to multiple sources and donors of human skin fibroblasts, and specificity of the effect to CTGF (IGFBP-rP2). A, Soluble AGE BSA or control BSA at 100 μg/ml was added to duplicate wells of confluent primary cultures of human skin fibroblasts from multiple donors under serum-free conditions, and total RNA was collected at the time points shown. IGFBP-rP2 (CTGF) mRNA was then determined by quantitative RT-PCR in triplicate for each sample. The IGFBP-rP2 (CTGF) mRNA level is expressed in arbitrary units, normalized to 18S. The donor age and skin site of the fibroblasts studied are: 7-yr-old in human foreskin fibroblast CRL-2097 cells are generalizable to human dermal fibroblasts. When these other cells were treated with AGE BSA, increases in CTGF mRNA were observed 48 h after treatment, compared with the control BSA, in all of the fibroblast cell lines studied whether they were derived from neonatal foreskin (A305), a child’s abdomen (CRL-1474), or the forearm of a mature adult (A35; Fig. 3A). To address whether the changes seen in CTGF mRNA were relatively specific, other members of the IGFBP superfamily were analyzed in the same cell system. In contrast to the observed regulation of CTGF mRNA by AGE, IGFBP-rP1 (mac-25) mRNA was not up-regulated by AGE BSA (Fig. 3B). IGFBP-3 is the predominant IGFBP present in human fibroblasts. Most IGFBP-3 is the predominant IGFBP present in human fibroblasts. Most IGFBP-3

Densitometric analysis

To quantify the relative induction of CTGF after Western immunoblotting, densitometric measurement was performed using GS-700 Imaging Densitometer with Multi-Analyzer Software (Bio-Rad Laboratories, Inc.).

Statistical analysis

Results

To determine whether CTGF mRNA steady state levels are up-regulated by AGE in primary cultures of human dermal fibroblasts, confluent monolayers of CRL-2097 fibroblasts were treated with soluble AGE BSA under serum-free conditions. In response to 100 μg/ml AGE BSA, an increase in CTGF mRNA was initially detectable after 8 h of AGE treatment (Fig. 1A), and a progressive increase occurred over the 3-day time course of the study (Fig. 1B). In contrast, no change in CTGF mRNA over time was seen with the same concentration of control BSA (Fig. 1A and B). These results were confirmed by Northern analysis (not shown). Transient treatment of cells with AGE for 8 h, followed by washing of cells with PBS and replacement with fresh serum-free medium, also caused a progressive increase in CTGF mRNA over subsequent days (Fig. 1C), with a clear persistence of the effect for at least 72 h after AGE addition.

A dose-response study with AGE BSA from 0–200 μg/ml, with continuous AGE treatment and RNA collection at 48 h after initial AGE addition, showed that increases in CTGF mRNA were significant using 10 μg/ml or more AGE BSA, whereas increasing concentrations of control BSA did not produce any change in CTGF mRNA compared with no addition of BSA (Fig. 2).

Primary human skin fibroblasts from other donors were studied to assess whether the changes seen in CTGF mRNA in human foreskin fibroblast CRL-2097 cells are generalizable to human dermal fibroblasts. When these other cells were treated with 100 μg/ml AGE BSA, increases in CTGF mRNA were observed 48 h after treatment, compared with the control BSA, in all of the fibroblast cell lines studied whether they were derived from neonatal foreskin (A305), a child’s abdomen (CRL-1474), or the forearm of a mature adult (A35; Fig. 3A).

To address whether the changes seen in CTGF mRNA were relatively specific, other members of the IGFBP superfamily were analyzed in the same cell system. In contrast to the observed regulation of CTGF mRNA by AGE, IGFBP-rP1 (mac-25) mRNA was not up-regulated by AGE BSA (Fig. 3B). IGFBP-3 is the predominant IGFBP present in human fibroblasts.
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blast conditioned medium (33), and its mRNA also was not increased by AGE (Fig. 3B). In comparison with the lack of effect of AGE treatment, TGFβ1 treatment modestly up-regulated IGFBP-rP1, and possibly IGFBP-3 mRNA, at 48 h (Fig. 3B), as previously described (33, 34). Also consistent with previous observations in cultured human fibroblasts (35), a pronounced up-regulation of CTGF mRNA occurred after TGFβ1 addition (Fig. 3B).

Formation of products of nonenzymatic glycosylation was inhibited by the dihydrazine compound, aminoguanidine (13). When cells were treated with BSA that had been coin-cubated for 10 weeks with both glucose and aminoguanidine as described in Materials and Methods, no increase in CTGF mRNA was observed compared with control BSA treatment alone or with serum-free medium without any addition (Fig. 4A). This result confirms that the active component in the AGE reagent used is a product of nonenzymatic glycosylation.

To determine whether early or advanced glycosylation end products are mediating the effect on CTGF in this cell system, cells were preincubated with anti-AGE IgG before addition of the AGE reagent. Using the anti-AGE antibody, the AGE induction of CTGF mRNA was inhibited, on the average, by 86.2% (Fig. 4B). As this antibody is specific to AGE and it does not bind to amadori products (26), which are early products of nonenzymatic glycosylation, these results show that AGE is the active component in the synthesized reagent responsible for increasing CTGF in these studies.

AGE may bind to and activate one or more of the defined cell surface receptors for AGE (8). The AGE receptor subtype, termed RAGE, has recently been shown to be present on the surface of human fibroblasts (27), and in some cell systems the induction of growth factors by AGE has been shown to be mediated by RAGE (36). When cells were preincubated with a blocking antibody of RAGE activation by AGE ligand, the induction of CTGF mRNA by AGE was attenuated by the RAGE, and is not inhibited by oxygen free radical scavengers. A. Soluble AGE BSA or control BSA at 100 µg/ml, or no treatment, was added to duplicate wells of confluent primary cultures of human fibroblasts (CRL-2097 cells) under serum-free conditions. In other wells BSA was added that had previously been coin-cubated with glucose and aminoguanidine before dialysis, as described in Materials and Methods. Total RNA was collected at 48 h, and CTGF mRNA was determined by quantitative RT-PCR in triplicate for each sample. The CTGF mRNA level is expressed in arbitrary units. Data are the mean ± 1 SD from three independent experiments. **, P < 0.01 vs. all other treatments. B. Wells were preincubated with anti-AGE polyclonal neutralizing IgG, anti-RAGE polyclonal neutralizing IgG (each at 100 µg/ml IgG), or 100 µg/ml normal rabbit serum (NRS) IgG for 2 h. Soluble AGE BSA or control BSA at 100 µg/ml was then added to the wells. Total RNA was collected at 48 h, and IGFBP-rP2 (CTGF) mRNA was determined by quantitative RT-PCR in triplicate for each sample. Data are the mean ± 1 SD from four independent experiments. *, P = 0.05; **, P < 0.01 (vs. AGE BSA added alone). C. Duplicate wells of human fibroblasts under serum-free conditions were incubated with serum-free medium alone, 20 mM N-acetyl cysteine, or 100 mM dimethylsulfoxide (DMSO) for 2 h. Soluble AGE BSA or control BSA at 100 µg/ml was then added to the wells. Total RNA was collected at 48 h, and CTGF mRNA was determined by quantitative RT-PCR in triplicate for each sample. The mRNA levels are expressed in arbitrary units. Data are the mean ± 1 SD from three independent experiments.
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Fig. 5. AGE induction of CTGF (IGFBP-rP2) mRNA is independent of endogenous TGFβ1 activity. Duplicate wells of confluent primary cultures of human fibroblasts (CRL-2097 cells) under serum-free conditions were incubated with AGE BSA or control BSA at 100 µg/ml or with TGFβ1 at 1 ng/ml for 24 h (A) or 48 h (B), and in some wells a chicken antihuman TGFβ1-neutralizing antibody (200 ng/ml) was added simultaneously as indicated. Total RNA was collected at 24 h (A) and 48 h (B), and CTGF mRNA was determined by quantitative RT-PCR in triplicate lysates (not shown). Densitometric analysis of CTGF from whole cell lysates for CTGF protein after AGE treatment was studied. Preincubation of the fibroblasts with the antioxidants dimethylsulfoxide or N-acetyl cysteine, however, did not inhibit the increases in CTGF mRNA (Fig. 4C). These results imply that RO species are unlikely to play a role in the observed AGE effect on CTGF.

A potential role for autocrine TGFβ1 in CTGF mRNA induction by AGE was then examined. TGFβ1 is a potent inducer of CTGF gene expression in this cell system (Fig. 3A), and in addition, AGE may induce TGFβ1 mRNA and protein in some cells (13). Induction of CTGF mRNA by rhTGFβ1 added to the cultured fibroblasts was fully inhibited by a TGFβ1-neutralizing antibody at 24 h (Fig. 5A) and 48 h (Fig. 5B). In contrast, when the same antibody was added under the same conditions in parallel wells, no significant inhibition of the CTGF mRNA increase induced by AGE occurred (Fig. 5, A and B), indicating that the effect of AGE is TGFβ1 independent in this cell system.

The fibroblast cellular protein from CRL-2097 skin fibroblasts was then analyzed to determine whether AGE treatment causes increases in CTGF protein as well as increases in steady state CTGF mRNA levels. By Western immunoblot after SDS-PAGE, using a polyclonal IGFBP-rP2 (CTGF) antiserum (25), CTGF steady state protein levels over days 1–3 in the conditioned medium were increased in response to AGE BSA, compared with control BSA treatment (Fig. 6A). A progressive increase in intact CTGF as well as previously described lower M, immunoreactive forms, at approximately 14 and 20 kDa (38), occurred (Fig. 6A). High Mr immunoreactive material (>80 kDa) was also more prominent in the AGE-treated medium, which may include CTGF covalently cross-linked by AGE. All of these increases were more marked using AGE BSA synthesized from glycolaldehyde compared with AGE BSA synthesized from glucose (not shown). Using densitometric analysis from three independent experiments, the intact CTGF in the medium was increased by AGE compared with control BSA treatment (mean ± SEM) by 3.6 ± 1.1-fold on day 1, 8.0 ± 1.7-fold on day 2, and 14.5 ± 3.1-fold on day 3 (P < 0.05 for all days of AGE treatment compared with control BSA treatment).

As CTGF is an extracellular matrix and cell-associated signaling protein and also exists in cell media (38), analysis of whole cell lysates for CTGF protein after AGE treatment was performed. Western immunoblot analysis of the whole cell lysates after SDS-PAGE showed that intact CTGF was increased by AGE treatment from day 3 compared with control BSA treatment (Fig. 6B). There was no CTGF fragment or high Mr immunoreactive material observed in the lysates (not shown). Densitometric analysis of CTGF from lysates from four independent experiments (mean ± SEM) gave the following results for fold change with AGE treatment compared with control BSA treatment: 0.89 ± 0.33 on day 1, 0.97 ± 0.17 on day 2, and 1.53 ± 0.25 on day 3 (P < 0.05 for day 3 only for AGE compared with control BSA on the respective day).

Considering that the increase in CTGF in whole cell lysates on day 3 after AGE treatment was relatively modest, changes in cell lysate CTGF were further determined by a CTGF ELISA, as described in Materials and Methods. This assay can measure endogenous intact CTGF, which is present in whole cell lysates, but due to a lack of parallelism with the intact
rhCTGF used as the standard, it cannot be used to accurately measure the 14-kDa CTGF fragment (Fig. 7A), which is present in the fibroblast-conditioned media (Fig. 6A). Consistent with the Western immunoblots of cell lysates (Fig. 6B), the ELISA also showed that AGE treatment reproducibly increased CTGF in the fibroblast whole cell lysates on day 3 (Fig. 7B). Thus, in contrast to the increases in CTGF protein observed in the conditioned media, there was no increase in CTGF protein in the first 2 days after AGE treatment in the whole cell lysates compared with control, and there was only a modest and delayed increase in CTGF in the lysates, which was much less striking than the increases in CTGF protein observed in the conditioned media (Fig. 6A).

To determine whether the increase in CTGF in the whole cell lysates seen by day 3 of AGE treatment was accessible to the extracellular environment, a cell association assay for CTGF was performed, as described in Materials and Methods. This assay uses binding of a biotinylated CTGF primary antibody to endogenous CTGF protein, followed by antibody detection using a streptavidin-HRP system. As no plasma membrane-permeabilizing agents were used in the protocol, the specific signal detected by the CTGF primary antibody was due to CTGF present on the cell surface or in the extracellular matrix, rather than CTGF present in an intracellular compartment. As shown in Fig. 7C, at 72 h AGE at 100 µg/ml specifically increased the absorbance signal compared with control (P < 0.05 for analysis of combined data from four independent experiments). In parallel wells, under the same conditions of confluent cell monolayers in serum-free media, cell number determined by hemocytometer counting and trypan blue exclusion was not changed by AGE treatment compared with control BSA (not shown). Thus, these results indicate that at 72 h, AGE treatment increases cell-associated CTGF compared with BSA control treatment alone.

**Discussion**

This study describes the up-regulation of CTGF mRNA and protein by treatment of human skin fibroblasts with advanced glycosylation end products. The effect of AGE on CTGF induction was caused by products of nonenzymatic glycosylation, as coincubation of aminoguanidine, an inhibitor of nonenzymatic glycosylation, with glucose and BSA did not have an effect on CTGF mRNA, nor was an effect seen with increasing control BSA alone. The up-regulation of CTGF in this cell model was mediated by AGE rather than by earlier products of nonenzymatic glycosylation, such as amadori products, as the use of an antibody specific for AGE that does not bind amadori products inhibited the induction of CTGF gene expression. The effect was at least partly mediated through the AGE receptor known as RAGE, as an anti-RAGE antibody significantly attenuated the effect of AGE on CTGF.

The up-regulation of CTGF by AGE appears to be specific for CTGF and is generalizable to skin fibroblasts from dif-
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A: CTGF (IGFBP-rP2) ELISA and parallelism of samples with intact rhCTGF (IGFBP-rP2) standard

B: CTGF (IGFBP-rP2) in whole cell lysates measured by ELISA

C: Cell Associated CTGF (IGFBP-rP2) at day 3

Fig. 7. Whole cell lysate and cell-associated increases in CTGF (IGFBP-rP2) after AGE BSA treatment. A, CTGF (IGFBP-rP2) ELISA as described in Materials and Methods, showing the standard curve generated for intact CTGF (IGFBP-rP2; □) compared with the curves generated with a 14-kDa fragment of CTGF (IGFBP-rP2; △), native intact CTGF present in 2097 cell-conditioned medium (+), and 2097 cell whole cell lysates (+). B, Results of analysis of CTGF in the whole cell lysates by CTGF ELISA after treatment up to 72 h with AGE or control BSA, each at 100 μg/ml. Data are the mean ± 1 SD of four independent experiments. *, P < 0.05 vs. control BSA on day 3. C, Results showing cell-associated CTGF after AGE treatment compared with control BSA (100 μg/ml) at 72 h, using biotinylated anti-CTGF IgG primary antibody followed by HRP-labeled secondary antibody or the secondary antibody alone, as described in Materials and Methods. Data are the mean ± 1 SD of four independent experiments. *, P < 0.05 vs. all other groups.

fering sources and passage number. In each of the four fibroblast cell lines studied, AGE up-regulated CTGF. In the cell line most extensively studied, CRL-2097, CTGF was regulated by AGE in early passages (passage 4) and also at later passages (passage 12). In contrast to effects on CTGF, the two other members of the IGFBP superfamily that were studied, IGFBP-3 and IGFBP-rP1, were not up-regulated by AGE. Further studies will be required to determine whether AGE affects other members of the CCN (CTGF, Cyrbl, Nov) family.

The concentrations of AGE BSA used in these experiments approximate those used in vitro in other studies exploring biological effects of AGE on cells (13, 27). Although there is no universal standard method for measuring specific AGE components at this time, and the AGE antibodies used in assays measuring AGE differ (39), the AGE BSA concentrations studied are in the broad range for AGE concentrations found in diabetic serum (40).

Few AGE components have been defined biochemically to date, and the specific end-product(s) that might be mediating the effect on CTGF was not identified in this work. AGE adducts existing in diabetic tissues that have been shown to signal through AGE receptors include mainly CML (28) and imidazoline-based products (41). Considering that CML adduct is a ligand for RAGE (28), that our AGE reagent contained CML, and that at least part of the AGE effect on CTGF has been shown to be mediated through RAGE, it is plausible that CML adducts are one of the ACE components operative in this study. In the current work, when AGE BSA synthesized from glycolaldehyde was studied in experiments where AGE BSA synthesized from glucose was also used in treatments, each at the same AGE concentration of 100 μg/ml BSA with mRNA measurements over 3 sequential days, the induction of CTGF mRNA by these reagents did not differ (data not shown). As these two AGE reagents contain differing amounts of CML (as described in Materials and Methods), the CML adduct cannot be the only explanation for the observed AGE effect on CTGF mRNA in these AGE preparations. Further experiments with pure CML and other pure AGE adducts, when available, will be required to address this issue.

A number of subtypes of cell surface receptor bind AGE specifically and are responsible for mediating multiple cellular effects of AGE (42). These receptors exist in four main classes: RAGE, AGE-R1, AGE-R2, and AGE-R3 (8). In the diabetic environment, the increased AGE present is hypothesized to bind and activate AGE receptors, and in some studies, the induction of growth factors by AGE has been shown to be mediated by AGE receptors, including RAGE (43). Our studies indicate that RAGE is responsible for mediating at least part of the effect of AGE on CTGF mRNA. The
possibility that other AGE receptors might also contribute to these effects is not excluded by this work.

A role for growth factors in contributing to chronic diabetes-related end-organ complications, particularly vascular endothelial growth factor (VEGF), TGFβ1, IGF-I, and platelet-derived growth factor, is under increasing evaluation, and a potential role for CTGF in chronic diabetic complications is emerging. CTGF is a potent profibrotic agent (18, 44), which is reflected in its ability to induce ECM components and increase fibroblast DNA synthesis (18) and to promote angiogenesis (20, 21). CTGF mRNA levels are up-regulated in many chronic disease states where fibrosis is prominent (22-24). Two separate studies involving renal mesangial cells and differing diabetic rat models recently reported that CTGF gene expression (45) as well as protein (46) are increased in mesangial cells after exposure to high glucose and in vivo in diabetic rat kidneys. Immunohistochemical studies of kidney tissue in human end-stage renal disease showed increased CTGF protein in diabetic kidneys as well as other nephropathies (47), and CTGF mRNA is markedly increased in advanced atheromatous lesions (48).

This is the first report of CTGF induction by advanced glycosylation end products, and it provides a potentially critical linkage among AGE, growth factors, and fibrosis. AGE induction of growth factors and cytokines has been described for VEGF, TGFβ1, IGF-I and platelet-derived growth factor, TNFα, IL-1β, and IL-6 (3) mainly in various endothelial and mesenchymal cultured cells and in some cases by AGE administration in vitro (19). CTGF appears to fit well into this group of proinflammatory and/or profibrotic proteins.

The striking persistent effect over 3 days of AGE on CTGF mRNA even after transient treatment suggests that regulation of CTGF by the AGE reagent tested is complex and may involve multiple interrelated intracellular signals. The cellular mechanism of AGE induction of CTGF mRNA was not defined in this study. ROS species were not implicated, because antioxidants were ineffective in inhibiting AGE induction of CTGF. TGFβ1 is a known potent inducer of CTGF gene expression, and CTGF is implicated as a downstream mediator of TGFβ1 effects (49), particularly in fibrosis (44). We were unable to show, however, that TGFβ1 is a mediator in the AGE induction of CTGF. In the current work, both the early time course of initial induction of CTGF mRNA by AGE at 8 h as well as the inability of TGFβ1-neutralizing antibodies to inhibit AGE induction of CTGF suggest that AGE is operating through mechanisms that are independent of TGFβ1. Although studies involving the use of exogenously added neutralizing antibodies have potential limitations in assessing the role of endogenous protein bioactivity, that total TGFβ1 measurements in conditioned media measured by TGFβ1 ELISA (Promega Corp., Madison, WI) in these cells were not increased by AGE compared with control BSA treatment (data not shown) is also supportive that TGFβ1 is not a mediator of AGE induction of CTGF in this work. These results contrast with studies describing TGFβ-dependent effects of glucose on CTGF up-regulation in human mesangial cells (45, 46), but are consistent with other studies showing that various reagents can potentially up-regulate CTGF mRNA independently of TGFβ1 (50).

In human fibroblast primary cultures, CTGF exists at very low levels in conditioned medium and is often present in low M, fragment forms, which may also have bioactivity (38). That intact CTGF was readily detectable in the medium after AGE treatment may be partly related to posttranslational modification of CTGF, with cross-linking of CTGF protein by AGE into a high Mr, immunoreactive form and redistribution of CTGF from a cell-associated site into the conditioned medium. In addition to the CTGF increases in the conditioned medium and consistent with the progressive increase in CTGF mRNA after AGE treatment, AGE caused increases in intact CTGF in whole cell lysates at 72 h. Further analysis showed that the CTGF increase in the lysates included protein that was cell associated and in a site accessible to the extracellular environment. To what extent the bioactivity of CTGF protein is affected by its presence in the medium compared with a cell-associated site is an important issue for future study of CTGF bioactivity.

There is a rationale to potentially link AGE effects and diabetic complications with the induction of CTGF in skin and, by association, with pathology in other tissues. A feature commonly present in human diabetes, even in late childhood and adolescence, is skin thickening and contracture (51), termed diabetic sclerosis. This process affects mainly the distal extremities and is characterized by expansion of extracellular matrix, fibroblast proliferation, and angiogenesis (52). The presence of overt diabetic sclerosis of skin is correlated with the presence and future development of end-organ complications, particularly diabetic nephropathy and retinopathy (53). AGE products are increased in human diabetic skin (6), and the levels of AGE in skin also correlate positively with the presence of diabetic microvascular kidney and eye disease (5, 6). That the ability of CTGF to induce fibrosis has been well characterized in skin (18, 44) makes skin fibroblasts a relevant cell model for the current study.

Clearly, in vivo and longer term studies are required to substantiate a more definitive role for induction of CTGF by AGE in potentially mediating diabetic fibrotic complications in skin and other organs. Activation of receptors for AGE, particularly RAGE (54), has also been implicated in the pathogenesis of fibrosis that develops in chronic diseases other than diabetes (42, 54, 55). That AGE up-regulates CTGF in nontransformed human fibroblasts suggests that CTGF may be a factor mediating the observed AGE and RAGE effects, which is a hypothesis that requires further testing.

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Connective Tissue Growth Factor/IGF-Binding Protein-Related Protein-2 Is a Mediator in the Induction of Fibronectin by Advanced Glycosylation End-Products in Human Dermal Fibroblasts

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Expansion of extracellular matrix with fibrosis occurs in many tissues, including skin, as part of the end-organ complications in diabetes. Advanced glycosylation end-products (AGEs) have been implicated as a pathogenic factor in diabetic tissue fibrosis. Connective tissue growth factor (CTGF), also known as IGF-binding protein-related protein-2, induces extracellular matrix. We have recently shown that CTGF mRNA and protein are up-regulated by AGE treatment of cultured human dermal fibroblasts. The aim of this study was to determine whether CTGF is an autocrine mediator in the induction of fibronectin (FN) by AGE. Primary cultures of non-fetal human dermal fibroblasts in confluent monolayer were treated with synthesized soluble AGE BSA, 0–200 μg/ml. Analysis of mRNA, by quantitative real-time RT-PCR and conditioned media from treated cultures, showed that FN mRNA was increased by approximately 4-fold at 48 h, and FN protein levels by Western immunoblot and FN ELISA were doubled, compared with control. In the same system, added recombinant human CTGF (0–500 ng/ml) induced FN mRNA and protein levels dose dependently and in a rapid time course. To test whether AGE BSA acts through cell-derived CTGF to induce FN, a CTGF neutralizing antibody was shown to significantly attenuate, but not fully inhibit, the AGE induction of FN mRNA. A pan-specific PKC inhibitor, GF109203X, at 0.2 μM, inhibited the induction of FN mRNA by AGE BSA. Although the same inhibitor did not significantly affect the induction of CTGF mRNA by AGE, it blocked the induction of FN mRNA by recombinant human CTGF. In summary, the induction of FN by AGE is partly mediated by the AGE-induced up-regulation of cell-derived CTGF and is dependent on PKC activity. These results have potential implications for the expansion of extracellular matrix in diabetes mellitus by advanced glycosylation end products. (Endocrinology 143: 1260–1269, 2002)

A MECHANISM PROPOSED whereby chronic hyperglycemia contributes to diabetic complications is in the formation of advanced glycosylation end-products (AGEs) (1). AGEs constitute irreversibly formed biochemical end-products of nonenzymatic glycosylation (2) and are elevated in many tissues, including skin (3), in subjects with diabetes (4). One method by which AGEs appear to contribute to pathological end-organ changes that occur in tissues in subjects with diabetes is through the induction of specific cytokines and growth factors, which may act as mediators in causing tissue pathology (5–7).

One hallmark in most tissues in which diabetic complications occur is expansion of extracellular matrix [ECM (8)]. AGEs (7) and increased cellular PKC activity (9) have each been identified as contributors to ECM expansion in diabetes through increased ECM production. An integral component of ECM is the glycoprotein, fibronectin (FN), which acts as a scaffold for collagen, and contributes to an ECM network involved in cell proliferation and migration (10). FN is increased in vivo in the ECM expansion that occurs in diabetes (11, 12).

Recently, we have reported that the cytokine, connective tissue growth factor (CTGF), is up-regulated at the mRNA and protein level by AGEs in confluent monolayers of cultured human dermal fibroblasts (13). Also known as IGF-binding protein-related protein-2 (IGFBP-rP2 (14)), CTGF is a potent inducer of ECM, including FN (15), in fibroblasts (16) as well as an angiogenic factor (17, 18). A potential role for CTGF in fibrotic disease states is increasingly being described (19, 20), suggesting that CTGF is a mediator in the ECM expansion and fibrosis occurring in diabetes. To date, the cellular mechanism of action of CTGF in enhancing ECM has been inadequately studied (21).

Because AGEs and CTGF can both up-regulate fibronectin, and AGEs also up-regulate CTGF in human fibroblasts, the aim of this study was to determine whether the induction of FN by AGEs is mediated through up-regulation of endogenous CTGF and to explore the cellular secondary messenger systems involved in mediating this effect on FN expression. Our work shows that CTGF contributes significantly to AGE up-regulation of FN in human dermal fibroblasts, through a PKC-dependent mechanism.

Abbreviations: AGE, Advanced glycosylation end-products; CML, carboxymethyl lysine; CTGF, connective tissue growth factor; ECM, extracellular matrix; FN, fibronectin; IGFBP-rP, insulin-like growth factor binding protein related protein; NRS, normal rabbit serum; rhCTGF, recombinant human connective tissue growth factor.
Materials and Methods

**Reagents**

The anti-fibronectin mouse monoclonal antibody against human fibronectin was purchased from Neomarkers Inc. (Union City, CA). Polyclonal anti-IGFBP-rP2 (CTGF) antibody (8800), was generated in New Zealand White rabbits, using full-length human CTGF (IGFBP-rP2) as the immunogen, as previously described (22). The anti-AGE polyclonal antiserum generated in New Zealand White rabbits, which neutralizes the activity of AGE BSA (23), was a generous gift from Dr. Miyata (Kissei Pharmaceutical Co. Ltd., Nagano, Japan). The antisera to human CTGF, and to AGE, as well as nonimmune normal rabbit serum (Sigma, St. Louis, MO) were each affinity purified by protein A affinity chromatography, using protein A-Sepharose (Pharmacia Biotech, Uppsala, Sweden). In each case, the eluted IgG protein was dialyzed against PBS, using a low-molecular-mass-cut-off membrane (Spectrapor 1, 6–8 kDa, Spectrum Industries, Los Angeles, CA), and confirmed to be immunglobulin by migration characteristics on nonreducing SDS-PAGE, followed by Coomassie staining. The total amount of IgG protein was quantitated using the DC protein assay reagent (Bio-Rad Laboratories, Hercules, CA) and was further confirmed by spectrophotometer absorbance readings at 280 nm. β-Glucose, BSA (fraction V, fatty acid and endotoxin free), human plasma fibronectin, and aminoguanidine were purchased from Sigma. TGF-β1 was purchased from Austral Biologicals (San Ramon, CA). The pan-specific PKC inhibitor, GF109203X, was purchased from BIOMOL Research Laboratories, Inc. (Plymouth Meeting, PA).

**AGE synthesis**

Advanced glycosylation end-products were synthesized in vitro, following methods previously described (13, 23, 24, 25). BSA (Sigma, Riga grade, fraction VI) at 10 mg/mL, was coincubated in sterile PBS with 0.5 mg/mL-glucose for 10 wk, with 1.5 mg/mmethylsulfonyl fluoride, under aerobic conditions at 37°C. To generate control BSA for comparison with AGE treatments, tubes were prepared with simultaneous incubations under the same conditions without the addition of the glucose. Additionally, in parallel preparations, aminoguanidine at 100 μM, as an inhibitor of formation of products of nonenzymatic glycosylation (7), was added to the BSA and glucose. All preparations were extensively dialyzed in PBS, using a low-molecular-mass-cut-off membrane (Spectrapor 1, 6–8 kDa, Spectrum industries; Ref. 7). The AGE content in the preparations was assessed by means of fluorescence, SDS-PAGE analysis, and ELISA. The fluorescence content, measured with a fluorescence spectrometer at 590 nm emission after a 450-nm excitation, in relative fluorescence units per milligram of BSA, was 11.2 ± 2.5 for control BSA, 52.3 ± 6.3 for AGE BSA, and 9.7 ± 1.2 for aminoguanidine added to BSA and glucose (termed aminoguanidine BSA) (25). By SDS-PAGE analysis under reducing conditions, followed by Coomassie staining, the AGE BSA produced was shown to have high-molecular-mass species, consistent with the intermolecular cross-linking ability of AGE, as described (23). In contrast, the control BSA and aminoguanidine BSA preparations did not have these high-molecular-mass forms (data not shown). By competitive ELISA (as described in Ref. 26) performed by Dr. P. Foles (Alteon Inc., Seattle, WA), using a synthetic Nε-carboxymethyl lysine (CML) analog as the standard, the CML content of the preparations (picomole CML per microgram of BSA ± 95% confidence interval) was: 13 ± 4 for AGE BSA from glucose and was not detectable (<1) for control BSA and also undetectable when aminoguanidine at 100 μM was coincubated with BSA and glucose.

**Synthesis and purification of recombinant human CTGF**

The recombinant human CTGF [rhCTGF (IGFBP-rP2)] protein was produced using a baculovirus expression system (Innositron Corp., Carlsbad, CA). The CTGF 1047-bp cDNA open reading frame was cloned from a recombinant human brain cDNA library and sequenced. The resulting fragment coding for full-length nontagged human CTGF was subcloned into BomHI and XhoI sites in the baculovirus human recombinant baculovirus vector, pFastBac1 (Life Technologies, Inc., Rockville, MD), and insert presence and orientation were verified by DNA sequencing. Recombinant baculovirus stocks were isolated and produced in increasing titer, as recommended by the supplier of the expression system. The rhCTGF protein was then produced by infecting HIGH Five insect cells with recombinant virus under serum-free conditions and collecting the conditioned media. To purify rhCTGF protein from filtered media, heparin-Sepharose affinity chromatography, using HiTrap columns (Pharmacia Biotech) with a step-up salt gradient in the elution, was employed as previously described (15, 27). Peak fractions containing rhCTGF were determined by Western immunoblotting and CTGF protein was quantitated using Coomassie Blue-stained gels with BSA as standard, as previously described (13, 28).

**Cell culture**

Primary cell cultures of nonfetal human dermal fibroblasts, CRL-2097, and CRL-1474 were purchased from ATCC (Manassas, VA). Cells were maintained in MEM supplemented with 10% FBS and were used in these studies between passages 4 and 12. The human primary cultures of dermal fibroblasts, designated A35 (derived from the forearm of a 70-yr-old male) and A305 (newborn foreskin fibroblasts) were generous gifts from Dr. S. Goldstein, Memorial Veteran’s Hospital (Little Rock, AR). These cells were maintained in DMEM containing 450 mg/dl glucose and 15% FBS.

**Cell treatment**

After trypsinization, cells were grown in 12-well plates for 5 d in their respective media with FBS until they were confluent. For experiments requiring the use of blocking antibodies to AGE or CTGF, or using control normal rabbit serum IgG, cells were grown in 24-well plates under the same conditions. Cells were then incubated in their respective serum-free media for 16 h and were then treated with 0 to 10 μM AGE under serum-free conditions using fresh media. Unless otherwise indicated, the conditioned media were not changed after adding the treatments. Cell lysates and conditioned media were harvested up to 3 d after treatments. For experiments involving the use of blocking antibodies or PKC inhibitors, cells were preincubated with the antibody or reagent for 2 h under serum-free conditions before the addition of AGI or control BSA. The recombinant human CTGF protein was then produced by infected insect cells with recombinant virus under serum-free conditions and collecting the conditioned media. To purify rhCTGF protein from filtered media, heparin-Sepharose affinity chromatography, using HiTrap columns (Pharmacia Biotech) with a step-up salt gradient in the elution, was employed as previously described (15, 27). Peak fractions containing rhCTGF were determined by Western immunoblotting and CTGF protein was quantitated using Coomassie Blue-stained gels with BSA as standard, as previously described (13, 28).

**Total RNA isolation and analysis by quantitative real-time RT-PCR**

Total RNA was isolated from duplicate wells, (RNAeasy mini kit, QIAGEN, Valencia, CA) and analyzed by quantitative real-time PCR using an ABI Prism 7700 sequence detection system (PE Applied Biosystems, Foster City, CA) as previously described (13, 29). This system is based on the ability of the 5′ nucleoside activity of Taq polymerase to cleave a nonextendable dye-labeled fluorescent hybrid probe during the extension phase of PCR. The following sequence specific primers and probes for human CTGF, FN, and 18S rRNA were designed, using Primer Express software 1.0 (PE Applied Biosystems): for FN, forward 5′-TCTTCTGCTGTTATAGTGCGAC-3′, reverse 5′-AGACCCCAACCCTCTTACATTGAA-3′, and probe 5′-6FAM-CCACGTGCCAGGATTACCGGCTACAT-TAMRA-3′; for CTGF, forward 5′-GAGGAAAAACATATAGAGAACGCGAAA-3′, reverse 5′-GCCAACAAGCTCTGATGA-3′, and probe 5′-6FAM-TTTGAGCTTCTCCTGCTGCACCACCTTGT-TAMRA-3′; for 18S, forward 5′-CGGCTACCACTCAGAGGAA-3′, reverse 5′-GCTGAAATTCGCCACCCGTCCTCACTCCCTC-TAMRA-3′. Primers were used at a concentration of 200 nm and probes at 100 nm in each reaction. Multiscribe reverse transcriptase and AmpliTaq gold polymerase (PE Applied Biosystems) were used in all RT-PCR. Each RNA sample was analyzed in triplicate. Relative quantitation of 18S rRNA and human FN and CTGF mRNAs were calculated, using the competitive threshold cycle number for each sample fitted to a five-point standard curve (ABI prism 7000 user bulletin no. 2, PE Applied Biosystems). The standard curve was constructed using a serial dilution of total RNA extracted from human cardiac fibroblasts that had been treated with TGF-β1 at 1 ng/ml for 24 h. Expression levels were normalized to 18S rRNA and related to relevant controls as indicated in the text.

**Preparation of conditioned media and cell lysates**

Conditioned media were collected after cell treatment and centrifuged at 10,000 x g for 10 min at 4°C, and then supernatants from duplicate wells within each experiment were pooled and stored at −20°C.
C until analysis. Cell lysate samples were harvested after treatment by washing cells with PBS and then adding 100 µl cold RIPA lysis buffer (20 mM Tris, pH 8.0, 150 mM NaCl, 1% Nonidet P-40, 0.5% NaDOC, 0.1% SDS) plus a protease inhibitor cocktail (Roche Molecular Biochemicals, Mannheim, Germany) directly to each well. Plates were rocked for 15 min at 4°C, and lysates were collected and centrifuged at 10,000 × g for 10 min at 4°C. The supernatants from duplicate wells within each experiment were pooled and stored at −20°C until analysis. Total protein concentration was determined for each sample by use of the DC protein assay reagent (Bio-Rad Laboratories, Inc.). Twenty micrograms of total protein were then loaded per lane for SDS-PAGE analysis.

Western immunoblot analysis

Conditioned media samples were separated on 7.5% nonreducing SDS-PAGE. Proteins were electroblotted onto nitrocellulose, and membranes were blocked with 5% nonfat dry milk/TBS with 0.1% (vol/vol) Tween 20 for 1 h at 22°C, then incubated overnight, 4°C, in FN antiserum (1:800 dilution; NeoMarkers, Inc.). After incubation of membranes with a horseradish peroxidase-labeled secondary antibody for 1 h at 22°C, immunoreactive proteins were detected by use of enhanced chemiluminescence (NEN Life Science Products, Boston, MA).

ELISA for fibronectin

A competitive ELISA was developed based on previously described methods (30). Pure human plasma FN (100 ng/well) in 10 mM sodium carbonate, pH 9.6, was adsorbed to 96-well immunoplates (Nalge Nunc International, Rochester, NY) by a 20-h incubation at 4°C. The wells were then blocked by incubation with PBS, 0.1% (vol/vol) Triton X-100 (buffer A) containing 10 g/liter BSA for 2 h at 37°C, and washed four times with buffer A. Purified human FN in buffer A was used to generate standard curves. Standard and samples (125 µl/tube) were incubated with a limiting amount (1:2500 titer, 0.005 µl/tube) of antifibronectin antibody (NeoMarkers, Inc.) at 22°C for 1 h. The plate was then incubated with the FN standards (0–500 ng) and samples (each at 100 µl/well) for 30 min at 22°C. After washing, the plate was incubated with streptavidin horseradish peroxidase at 1:3000 (Sigma) for 30 min at 22°C and, after four more washes in buffer A, with substrate [0.1 g/liter 3,3',5,5'-tetramethyl benzidine in 0.2 mM sodium acetate, pH 6, containing 0.6% (wt/wt) H2O2] for 10 min at 22°C. The reaction was stopped by the addition of 2 M H2SO4, and the absorbance was measured at 450 nm using a microplate reader. Linearity in the assay was achieved over the range of 15–250 ng/well of FN.

Densitometric analysis

To quantify the relative induction of FN following SDS-PAGE and Western immunoblotting, densitometric measurement was performed using GS-700 imaging densitometer with MultiAnalyst Software (Bio-Rad Laboratories, Inc.).

Statistical analysis

Results are expressed as mean ± SD or mean ± SEM as indicated. Differences between groups were assessed using a two-tailed paired t test in Microsoft Corp. (Redmond, WA) Excel 98, where shown. A level of P < 0.05 was considered statistically significant.

Results

To determine whether FN mRNA steady-state levels are up-regulated by AGE in primary cultures of human dermal fibroblasts, confluent monolayers of CRL-2097 fibroblasts were treated with soluble AGE BSA under serum-free conditions. In response to 100 µg/ml of AGE BSA, an increase in FN mRNA was observed at 24 h of AGE treatment, and a progressive increase occurred over the 3-d time course of the study (Fig. 1A). In contrast, no change in FN mRNA over time was seen after treatment with the same concentration of control BSA (Fig. 1A). A dose-response study with AGE BSA from 0 to 200 µg/ml in 2097 cells, with continuous AGE treatment and RNA collection at 48 h after initial AGE addition, showed that increases in FN mRNA were detectable using 10 µg/ml or greater AGE BSA, but increasing concentrations of control BSA did not show any change in FN mRNA in comparison with no addition (Fig. 1B).

Formation of products of nonenzymatic glycosylation is inhibited by the dihydrazine compound aminoguanidine (7). When cells were treated with BSA that had been coincubated for 10 wk with both glucose and aminoguanidine, as described in Materials and Methods, no increase in FN mRNA was observed, compared with control BSA treatment alone or with serum-free media without any addition (Fig. 2A). This result confirms that the active component in the AGE reagent used is a product of nonenzymatic glycosylation. As a positive control reagent in this system for the induction of FN, TGF-β1 (1 ng/ml) was also seen to induce FN mRNA (Fig. 2A).

To determine whether early or advanced glycosylation end-products mediate the effect of the AGE reagent on FN in this cell system, cells were preincubated with anti-AGE IgG before addition of the AGE reagent. Using the anti-AGE antibody, the AGE induction of FN mRNA was inhibited on average by 81% (Fig. 2B). In contrast, normal rabbit serum
had no effect (Fig. 2B). Because the anti-AGE antibody is specific for AGE and does not bind to Amadori products (23), which are early products of nonenzymatic glycosylation, these results showed that AGE is the active component in the synthesized reagent responsible for increasing FN in these studies.

To test whether the changes seen in FN mRNA in human foreskin fibroblast CRL-2097 cells after AGE treatment can also be observed in human dermal fibroblasts, primary human skin fibroblasts from other donors were studied. When these other cells were treated with 100 μg/ml of AGE BSA, increases in FN mRNA were observed 48 h after treatment, compared with control BSA, in all of the fibroblast cell lines studied, whether they were derived from neonatal foreskin (A305), a child's abdomen (CRL-1474), or the forearm of a mature adult (A35) (Fig. 3).

The 2097 fibroblasts were used in all subsequent experiments. The addition of pure rhCTGF protein (500 ng/ml) caused an induction of FN mRNA. Up-regulation was observed at 24 h and 48 h after reagent addition (Fig. 4A). In a parallel fashion, CTGF mRNA was autoinduced by rhCTGF treatment (Fig. 4A). Such autoinduction of CTGF mRNA, observed with the addition of recombinant CTGF protein, is consistent with a similar effect of added CTGF protein previously described in cultured rodent renal mesangial cells (27). A dose-response study from 0 to 500 ng/ml rhCTGF showed that statistically significant increases in FN mRNA at 24 h after treatment occurred when 100 ng/ml or more rhCTGF protein was added to the conditioned media (Fig. 4B). In parallel with the induction observed for FN, a dose-response induction of CTGF mRNA by added rhCTGF was also seen with a statistically significant increase occurring from 250 ng/ml added protein (Fig. 4B).

To determine whether AGE and rhCTGF treatment up-regulated FN protein levels, conditioned media were collected and analyzed by Western immunoblots and by FN ELISA. Immunoblot analysis indicated that in the presence of no addition, with serum free media alone, basal levels of FN protein accumulated in the conditioned media (Fig. 5A), a finding that has been observed by others in human fibroblasts (28). After treatment with rhCTGF, increased FN protein, compared with the no addition control, was detected in a time- and dose-dependent manner (Fig. 5, A and B). The addition of control BSA resulted in an accumulation of soluble FN protein (Fig. 5C). Over the 3 d of the study, after treatment of cells with AGE BSA, FN protein levels were further increased above that of the control BSA (Fig. 5C). In whole-cell lysates, an increase in FN protein was also observed after rhCTGF and AGE treatment, compared with either no addition or control BSA (data not shown). Quantitation by FN ELISA confirmed the media Western immunoblot results (Fig. 5, D through E). FN protein in conditioned media after 48 h treatment with 500 ng/ml rhCTGF was approximately 1.6-fold higher than in cells in which serum-free media were added alone (Fig. 5D). Increases in FN media
of 72 h (13). GF109203X (data not shown). These results indicate that in treatment and continued to increase for the study duration mRNA were seen with higher concentrations (up to 5 μM) of the conditioned media were detectable within 24 h of AGE further specific inhibitory effects on FN mRNA or CTGF or more of the AGE BSA (13). Increases in CTGF protein in these reagents on FN mRNA induction (Fig. 7, A and C). No after AGE addition to the media and became most marked (unstimulated) FN mRNA were observed, suggesting that

Increases in CTGF mRNA were detectable from 8 h onward isoforms (33). In addition, no significant effects on basal BSA induces up-regulation ofCTGF mRNA and protein (13). (0.2 μM), consistent with specificity of this inhibitor for PKC blast system and under the same cell culture conditions, AGE GF109203X were observed at relatively low concentrations

marked than the observed increases in above the controls, and these increases were somewhat less mRNA to basal levels (Fig. 7C). Autoinduction of CTGF treatment by AGE (Fig. 7B). In a similar manner, preincubation of a PKC inhibitor for 2 h, followed by addition of reagent (rhCTGF or AGE), and total RNA was collected at 48 h. Preincubation of cell monolayers with the pan-specific PKC inhibitor GF109203X at 0.2 μM caused an inhibition of AGE induction of FN mRNA by about 82% at 96 h, compared with the effect of normal rabbit serum IgG (Fig. 6B). These studies showed that anti-CTGF IgG abrogated the induction of FN mRNA by AGE by about 42%, compared with no IgG addition (Fig. 6A). In contrast to the effect of the anti-CTGF IgG, the same amount of normal rabbit serum IgG had no obvious effects (Fig. 6A). A longer-term protocol to 96 h was then performed, with the anti-CTGF IgG or normal rabbit serum IgG also added at 48 h (Fig. 6B).

We have recently reported that in the same dermal fibroblast system and under the same cell culture conditions, AGE BSA induces up-regulation of CTGF mRNA and protein (13). Increases in CTGF mRNA were detectable from 8 h onward after AGE addition to the media and became most marked after 72 h of AGE treatment and were seen using 10 μg/ml or more of the AGE BSA (13). Increases in CTGF protein in the conditioned media were detectable within 24 h of AGE treatment and continued to increase for the study duration of 72 h (13).

Based on our present observation that both AGE and rhCTGF up-regulate FN and that AGE increases endogenous CTGF in fibroblasts, we tested whether cell-derived CTGF, in an autocrine manner, is a contributor to the observed increase in FN mRNA following AGE treatment. Using an IgG affinity purified fraction of polyclonal antibody against rhCTGF, the induction of FN at 48 h by exogenously added rhCTGF (250 ng/ml) was fully inhibited (Fig. 6A). In contrast, no inhibition of FN mRNA up-regulation by rhCTGF occurred when the same amount of normal rabbit serum IgG was added exogenously to cells (Fig. 6A). In other experiments, the anti-CTGF antibody inhibited rhCTGF-induced increases in FN mRNA after 24 h of rhCTGF treatment (data not shown). In addition, the autoinduction of CTGF mRNA by rhCTGF, described in Fig. 4, was fully blocked by the CTGF antibody (data not shown). These results confirm that the anti-CTGF antibody specifically neutralizes the bioactivity of CTGF in this system.

The ability of the CTGF-neutralizing antibody to block AGE induction of FN mRNA was then assessed. When the CTGF antibody was added before AGE treatment of cells, the induction of FN mRNA at 48 h by AGE, was attenuated by about 42%, compared with no IgG addition (Fig. 6A). In contrast to the effect of the anti-CTGF IgG, the same amount of normal rabbit serum IgG had no obvious effects (Fig. 6A). A longer-term protocol to 96 h was then performed, with the anti-CTGF IgG or normal rabbit serum IgG also added at 48 h (Fig. 6B). These studies showed that anti-CTGF IgG abrogated the induction of FN mRNA by AGE by about 82% at 96 h, compared with the effect of normal rabbit serum IgG (Fig. 6B). Higher concentrations of CTGF antibody and more frequent treatments did not produce any greater inhibitory effect on AGE induction of FN (data not shown). These data show that endogenous CTGF contributes to the induction of FN by AGE in this cell system.

Recent studies have implicated PKC in the induction of FN in a diabetic environment (12, 31). Because AGE can regulate PKC isoforms in some systems (32), we tested whether PKC activity might be involved in the induction of FN by AGE and/or by rhCTGF. Cells were preincubated with a PKC inhibitor for 2 h, followed by addition of reagent (rhCTGF or AGE), and total RNA was collected at 48 h. Preincubation of cell monolayers with the pan-specific PKC inhibitor GF109203X at 0.2 μM caused an inhibition of AGE induction of FN mRNA by about 72% (Fig. 7A). In contrast, there was no statistically significant inhibition of CTGF mRNA induction by AGE (Fig. 7B). In a similar manner, preincubation of cells with GF109203X reduced rhCTGF induction of FN mRNA to basal levels (Fig. 7C). Autoinduction of CTGF mRNA by rhCTGF, however, was not significantly inhibited by preincubation with GF109203X (Fig. 7D). The effects of GF109203X were observed at relatively low concentrations (0.2 μM), consistent with specificity of this inhibitor for PKC isoforms (33). In addition, no specificity on basal (unstimulated) FN mRNA were observed, suggesting that the PKC blocker was specifically inhibiting the activity of these reagents on FN mRNA induction (Fig. 7, A and C). No further specific inhibitory effects on FN mRNA or CTGF mRNA were seen with higher concentrations (up to 5 μM) of GF109203X (data not shown). These results indicate that in
The up-regulation of FN by AGE is generalizable to skin expression in the previous work (35), is known to have a role in the demonstration that CTGF contributes to AGE up-regulation of FN, possibly in cooperation with CTGF, is an important topic to address in future studies.

BSA used in these experiments approximate those used in vitro in other studies exploring biological effects of AGE on cells (7, 34). In addition to AGE BSA synthesized from glucose, we have also synthesized AGE BSA from glycoaldehyde as substrate (13) and have confirmed that AGE synthesized from glycoaldehyde also induces FN mRNA (data not shown). BSA was used as the protein for synthesizing AGE adduct because it is highly purified and delipidated and has commonly been used in AGE experiments by others (23–25). We have not yet studied other proteins made from AGE, such as AGE synthesized using extracellular matrix proteins. Few AGE components have to date been defined biochemically, and the specific glycosylation end-product(s) that might be mediating the effect on FN were not identified in this work.

Recent studies using AGE in human dermal fibroblasts have focused on AGE effects on type 1 collagen, rather then FN. These reports have shown that in contrast to the observed up-regulation of FN by AGE BSA in the current work, type 1 collagen mRNA and protein synthesis were inhibited by soluble AGE BSA treatment (35). This prior report and the current work are not inconsistent because regulation of FN and type 1 collagen gene transcription differs. Specifically, epidermal growth factor receptor activation, which was shown to mediate AGE inhibition of type 1 collagen gene expression in the previous work (35), is known to have a role in positively regulating FN transcriptional activity in fibroblasts (36, 37). What role the epidermal growth factor receptor may play in AGE up-regulation of FN, possibly in cooperation with CTGF, is an important topic to address in future studies.

Discussion

This study describes the stimulation of FN mRNA and protein following treatment of human skin fibroblasts with advanced glycosylation end-products and with rhCTGF and the demonstration that CTGF contributes to AGE up-regulation of FN mRNA. The effect of the synthesized AGE reagent on FN mRNA induction was seen to be caused by products of nonenzymatic glycosylation because incubation of aminoguanidine, an inhibitor of nonenzymatic glycosylation, with glucose and BSA did not affect FN mRNA, and no effect was seen with control BSA alone. The up-regulation of FN mRNA in this cell model by the synthesized AGE reagent was mediated by AGE rather than by earlier products of nonenzymatic glycosylation, such as Amadori products, because the use of an antibody specific for AGE, which does not bind Amadori products, inhibited the AGE induction of FN gene expression. The increase in soluble FN protein in this system by AGE and rhCTGF were greater than the accumulation of FN protein over time, which was seen in the presence of either no addition or the addition of control BSA.

The up-regulation of FN by AGE is generalizable to skin fibroblasts from differing sources and passage number. In each of the four fibroblast cell lines studied, AGE up-regulated FN. In the cell line most extensively studied, CRL-2097, FN was regulated by AGE in early passages (passage 4) and also at later passage (passage 12). The concentrations of AGE
CTGF is known to induce FN and to be profibrotic (16). The concentrations of soluble CTGF recombinant protein required in this cell system to induce FN mRNA and protein, at and above 100 ng/ml of rhCTGF, appear higher than others have employed in human fibroblasts (15). It is possible that the purified rhCTGF used in the current study is somewhat less bioactive than endogenous CTGF protein and that used by other groups and/or that quantitation of the purified protein differs between groups. Nonetheless, the current up-regulation of FN mRNA by rhCTGF in human fibroblasts has not been reported before this work. The current work shows that added CTGF induces FN mRNA and protein, and completely blocked what less bioactive than endogenous CTGF protein and that without evidence of regulation by cellular PKC activity (42). It is possible ited extent only. Inhibition of the induction of type I collagen by cAMP in human dermal fibroblasts has been described (31), and inhibition of PKC activity may attenuate chronic diabetes-related events (12). Although effects on PKC activity have been well described for high extracellular D-glucose (9) and early products of nonenzymatic glycosylation (39, 40), only recently has AGE been implicated in up-regulating PKC activity (32) and potentially using PKC pathways in inducing diabetic complications (41), indicating that the pathological effects of AGE and PKC on tissues may be interrelated at the level of induction of PKC by AGE. In addition, PKC pathways have been shown to regulate FN induction in various cell types, including human fibroblasts (30). The PKC inhibitor concentrations of GF109203X used in the current work (0.2 μM) are consistent with the published amounts required for specifically blocking PKC activity of both conventional and novel PKC isoforms (33). Although the isoform(s) of PKC that is mediating the effect of AGE on FN remains to be described, this study demonstrates a link between AGE and PKC in human dermal fibroblasts, two major proposed mechanisms involved in the pathogenesis of diabetic complications.

The intracellular second messenger systems mediating the up-regulation of ECM by CTGF have been studied to a limited extent only. Inhibition of the induction of type 1 collagen by cAMP in human dermal fibroblasts has been described without evidence of regulation by cellular PKC activity (42). Second messenger systems affecting the previously observed up-regulation of FN mRNA by rhCTGF in human fibroblasts (15) have not been reported before this work. The current studies show that the induction of FN mRNA by rhCTGF is fully blocked by a pan-specific PKC inhibitor in this cell system. These data using the PKC inhibitor do not indicate the number of cellular signaling intermediates involved and the sequence of the effect. Further work is needed to better define the second messenger pathways involved, and the specific PKC isoform(s) mediating the effect is also part of an ongoing study. CTGF cellular signaling in fibroblast cells has fibrosis. The CTGF neutralizing antibody studies do not implicate CTGF as the only mediator of AGE induction of FN; that AGE induction of FN was only partially inhibited by CTGF-neutralizing IgG also implicates CTGF-independent pathways in AGE-induced increases in FN. We have previously reported that in the same system, a TGF-β1 neutralizing antibody did not inhibit the induction of AGE by CTGF and that total TGF-β1 levels were not detectably increased over the time course of the study (13). These results suggest that TGF-β1 is not involved in the role played by CTGF in contributing to the AGE induction of fibronectin in this cell system. Up-regulation of CTGF in tissues in rodent models of diabetic nephropathy has recently been reported (27, 38). Although AGE has been shown to up-regulate FN in vivo (12), AGE as a reagent has not yet been reported in vivo to induce CTGF mRNA or protein. In contrast to our experimental model, AGE appears to accumulate slowly in vivo, particularly in long-lived proteins (1, 2), and in vivo studies are now required to further substantiate a role for CTGF in mediating diabetic and, specifically, AGE-related ECM expansion.

Up-regulation of PKC activity and PKC isoforms in a diabetic environment has been shown in vitro and in vivo in cells and in tissues that are susceptible to diabetic complications (31), and inhibition of PKC activity may attenuate chronic diabetes-related events (12). Although effects on PKC activity have been well described for high extracellular D-glucose (9) and early products of nonenzymatic glycosylation (39, 40), only recently has AGE been implicated in up-regulating PKC activity (32) and potentially using PKC pathways in inducing diabetic complications (41), indicating that the pathological effects of AGE and PKC on tissues may be interrelated at the level of induction of PKC by AGE. In addition, PKC pathways have been shown to regulate FN induction in various cell types, including human fibroblasts (30). The PKC inhibitor concentrations of GF109203X used in the current work (0.2 μM) are consistent with the published amounts required for specifically blocking PKC activity of both conventional and novel PKC isoforms (33). Although the isoform(s) of PKC that is mediating the effect of AGE on FN remains to be described, this study demonstrates a link between AGE and PKC in human dermal fibroblasts, two major proposed mechanisms involved in the pathogenesis of diabetic complications.
been linked to CTGF-induced activation of cell surface receptors, such as the platelet-derived growth factor receptor (43) and, when in solid phase, the integrin receptor, α6β1 (44). Which of these receptors, if any, is involved in FN up-regulation by CTGF, through a PKC-dependent mechanism, will require further study.

Recently, PKC activity has been shown to regulate CTGF mRNA in fibroblasts. A previous report showed that blocking conventional and novel PKC isoform activity in combination by using GF109203X and other PKC inhibitors, or PKC depletors, caused an induction of CTGF mRNA under conditions of high FCS in the conditioned media (45). The FCS content likely contributed to prominent basal PKC activity and the detection of the inhibition of CTGF gene expression by basal PKC activity (46). That only a slight and nonsignificant induction of CTGF mRNA was observed by GF109203X in the current study in the basal state (Fig. 7) may reflect low cellular PKC activity in these serum-deprived cells. Low PKC activity under serum-free conditions has been observed previously by other groups (47). As well as having no significant effect in the basal state, GF109203X also did not significantly affect the autoinduction of CTGF mRNA by rhCTGF in the current work (Fig. 7D).

The finding that CTGF is a mediator in AGE induction of FN in vitro may have relevance in diabetic complications. Based on previous work, there is a rationale to potentially link AGE effects and diabetic complications with the induction of CTGF and ECM in skin and, by association, with pathology in other tissues. A feature commonly present in human diabetes is skin thickening and contracture, termed diabetic sclerosis. This affects mainly the distal extremities and is characterized by expansion of extracellular matrix, fibroblast proliferation, and angiogenesis (48). The presence of overt diabetic sclerosis of skin is correlated with the presence and future development of other end-organ complications (49). AGE products are increased in human diabetic skin (3) as is FN (50, 51), and the levels of AGE in skin also correlate positively with the presence of diabetic microvascular kidney and eye disease (3, 4). That the ability of CTGF to induce fibrosis has been well characterized in skin (15, 42) makes skin fibroblasts a relevant cell model for the current study.

Our work provides potential links in diabetic complications characterized by ECM expansion. Interactions between AGE effects and PKC activity have been described, and AGE induced up-regulation of the profibrotic agent CTGF, which itself contributes to AGE-induced ECM expansion through a PKC-dependent mechanism, has been observed. This work contributes toward further understanding mechanisms involved in the development of chronic diabetes complications, particularly those characterized by ECM expansion and fibrosis.

Acknowledgments

We thank Alteon, Inc. (Ramsey, NJ) for measuring the CML adduct concentration in the AGE reagent by their CML ELISA. The generous gift of the anti-AGE antiserum from Dr. Miyata at Kissei Pharmaceutical Co. Ltd. (Hotaka, Japan) is gratefully acknowledged.

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References


Erratum

In the article by Dennis D. Rasmussen et al. (Endocrinology 140:1009-1012, 1999), part of Table 1 appeared incorrectly. The values in the row beginning with T3 actually correspond to T4, and the values in the row beginning with T4 correspond with T3. The correct table appears below. The authors regret the error.

Table 1. Effect of aging and melatonin treatment on hormones involved with energy regulation and body composition

<table>
<thead>
<tr>
<th>Hormone</th>
<th>Units</th>
<th>Young</th>
<th>Middle Age Control</th>
<th>Middle Age Melatonin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leptin</td>
<td>ng/ml</td>
<td>3.48 ± 0.27</td>
<td>8.28 ± 1.48&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.27 ± 0.45</td>
</tr>
<tr>
<td>Insulin</td>
<td>ng/ml</td>
<td>2.75 ± 0.25</td>
<td>3.48 ± 0.83&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.87 ± 0.25</td>
</tr>
<tr>
<td>Testosterone</td>
<td>ng/ml</td>
<td>1.55 ± 0.26</td>
<td>0.67 ± 0.08&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.88 ± 0.23&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Corticosterone</td>
<td>ng/ml</td>
<td>26.6 ± 11.5</td>
<td>12.2 ± 3.8</td>
<td>20.7 ± 7.0</td>
</tr>
<tr>
<td>T4</td>
<td>ng/ml</td>
<td>56.1 ± 2.1</td>
<td>36.0 ± 3.4&lt;sup&gt;a&lt;/sup&gt;</td>
<td>36.3 ± 2.9&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>T3</td>
<td>ng/ml</td>
<td>0.87 ± 0.11</td>
<td>0.93 ± 0.04</td>
<td>0.99 ± 0.03</td>
</tr>
<tr>
<td>IGF-1</td>
<td>μg/ml</td>
<td>1.62 ± 0.05</td>
<td>1.66 ± 0.04&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.65 ± 0.04&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

Data represent the mean ± SEM of 8 rats/group. *P < 0.05 vs. Young. **P < 0.05 vs. Young and vs. Melatonin.
Altered expression of low affinity insulin-like growth factor binding protein related proteins in hepatoblastoma

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Abstract. Hepatoblastoma is a poorly understood rare pediatric liver tumour. We have previously shown that the IGF-axis is seriously disrupted in this tumour type. With the recent discovery that several other proteins also have the potential to bind to IGFs called insulin-like growth factor binding protein related proteins (IGFBP-rPs), we undertook an examination of several such genes in a series of hepatoblastomas with matched normal liver tissue. The expression profiles obtained reveal that the expression of these genes are also disturbed in these tumours, and may have implications for our understanding of the IGF-axis and its importance in this disease.

Introduction

Hepatoblastoma is a rare pediatric liver disease with an incidence of between 0.5-1.5 per million children (1). Most hepatoblastomas are sporadic, but some familial inherited disorders, most notably Beckwith-Wiedemann (BWS), and familial adenomatous polyposis (FAP), are associated with an increased risk for developing hepatoblastomas. Pre-operative chemotherapy regimes have proven to be extremely successful for treating this disease, but an understanding of the molecular processes behind the development of hepatoblastoma has lagged.

Gene expression studies have revealed that several genes show altered expression in hepatoblastomas. These include genes involved with modifying chromatin, cell growth and cell cycle control (2,3). We have previously shown that the insulin-like growth factor axis is greatly altered in hepatoblastoma not only at the level of the growth factors themselves but also to their receptors and binding proteins (4,5). Because the IGF-axis plays an important role in many diverse cellular functions including the promotion of cell growth and cell survival, such alterations may be critical to this tumour type.

Recently an increasing number of proteins have been identified to have the ability to bind to the insulin-like growth factors albeit with low affinity. This has led to them being called insulin-like growth factor binding related proteins or IGFBP-rPs, although some controversy exists as to them being renamed. While the concept of an IGFBP superfamily with shared N-terminal domain is incontrovertible, the role of these proteins in modulating IGF action is still uncertain (6,7). These include the proteins CTGF, NovH and TAF all of which have been shown to have altered expression in cancer (6).

Because these genes may have functional roles with regard to the IGF-axis and have been shown to have altered expression in various cancers we sought to examine their expression in series of hepatoblastoma for which matched normal liver was available and in which the IGF-axis has been shown to be altered, in an attempt to assess whether their expression may also be altered as a consequence of the tumour.

Materials and methods

Samples. Twelve sporadic hepatoblastomas were examined in this study. For eight of these, matched normal liver tissue was available. All of the tumours with the exception of cases 6 and 7 were freeze-sectioned into 1 mm portions interrupted by 5 μm sections. The 1-mm sections were used for RNA isolation, while the interrupted thin sections were prepared for histopathological evaluation. These samples were fixed in formalin, stained and processed with hematoxylin and eosin in the usual manner. The results of this evaluation are presented in Table I, along with particulars for each sample. Human fetal livers (14- and 18-week) were obtained from therapeutic terminations, with the permission of the local ethics committee. Due to the nature of such procedures, limited amounts of tissue were obtained, and where available the mRNA was included in the analyses.

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E-mail: steven.gray@vai.org

Key words: hepatoblastoma, insulin-like growth factor binding protein related protein, gene expression
Table I. Clinical data for the tumour samples used in the study.

<table>
<thead>
<tr>
<th>Case no.</th>
<th>Age at diagnosis (months)</th>
<th>Sex</th>
<th>Histology</th>
<th>Pre-operative chemotherapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6</td>
<td>M</td>
<td>Epithelial</td>
<td>No</td>
</tr>
<tr>
<td>2</td>
<td>19</td>
<td>M</td>
<td>Epithelial</td>
<td>Yes</td>
</tr>
<tr>
<td>3</td>
<td>19</td>
<td>M</td>
<td>Epithelial</td>
<td>Yes</td>
</tr>
<tr>
<td>4</td>
<td>22</td>
<td>M</td>
<td>Epithelial/mesenchymal</td>
<td>Yes</td>
</tr>
<tr>
<td>5</td>
<td>54</td>
<td>M</td>
<td>Epithelial</td>
<td>Yes</td>
</tr>
<tr>
<td>6</td>
<td>2</td>
<td>M</td>
<td>Fetal</td>
<td>No</td>
</tr>
<tr>
<td>7</td>
<td>12</td>
<td>F</td>
<td>Fetal</td>
<td>Yes</td>
</tr>
<tr>
<td>8</td>
<td>36</td>
<td>M</td>
<td>Not available</td>
<td>Yes</td>
</tr>
<tr>
<td>9</td>
<td>11</td>
<td>F</td>
<td>Fetal</td>
<td>No</td>
</tr>
<tr>
<td>10</td>
<td>13</td>
<td>M</td>
<td>Fetal</td>
<td>No</td>
</tr>
<tr>
<td>11</td>
<td>8</td>
<td>F</td>
<td>Epithelial/mesenchymal</td>
<td>No</td>
</tr>
<tr>
<td>12</td>
<td>18</td>
<td>M</td>
<td>Not available</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Nucleic acid isolation. Total RNA was prepared as described previously (8).

Preparation of probe and RNase protection analysis. T3, T7 and Sp6 RNA polymerases (Invitrogen) were used to make antisense RNA probes from the following templates according to the protocol provided in the RPA II kit (Ambion). When incorporating radioactivity into the probe, radioactive ATP-UTP with a specific activity of 800 Ci/mmol was used. Cold UTP was added such that final UTP specific-activity was 80 Ci/mmol for the GAPDH probe and 400 Ci/mmol for the others.

The probes used in this study were prepared as follows: To measure IGFBP-rP1 (Mac25/TAF/PSF) expression, a SmaI/KpnI fragment from the full length cDNA contained in pCDNA3.1 (9), was blunted and subcloned into the EcoRV site of pBluescript II SK (-) (Stratagene). When linearised with HindIII a probe of 320 bases could be generated using T3 RNA polymerase of which 210 bases hybridize to IGFBP-rP1 specific transcripts.

IGFBP-rP2 (CTGF) expression was measured using a 217 bp SmaI fragment from the full length IGFBP-rP2 (pFastBac1 IGFBPrP2) (10) cloned into the EcoRV site of pBluescript II SK (-). Following linearization with EcoRI, a probe of 237 bases could be generated with T7 of which 217 bases hybridize to IGFBP-rP2 mRNA and protect from RNase.

A template for IGFBP-rP4 (Cyr61) was generated by cloning a 214 bp SmaI/PstI fragment of the full length cDNA from a pBK-CMV plasmid containing the full length Cyr61 mRNA (generous gift from Dr P. Berta) (11) into pBluescript II SK (-). Following linearization with EcoRI, a probe of 305 bases could be generated with T7 RNA polymerase of which 214 bases hybridize specifically to IGFBP-rP4 specific mRNA transcripts.

IGFBP-rP5 (L56/HtraA) expression was measured by cloning a 186 bp AvaiII fragment from a pUC19 plasmid containing the full length cDNA (kind gift of Dr B. Treub) (12) into the SmaI site of pGem3ZF (+) (Promega). Following linearization with EcoRI a probe of approximately 255 bases could be generated with Sp6 of which 186 bases hybridize specifically to IGFBP-rP5 mRNA transcripts.

All of the plasmids described above were sequenced, in order to ensure that the fragment cloned corresponded to the gene under examination. Sequencing was carried out with T7 Sequenase according to the manufacturer’s instructions (United States Biochemical). RNase protection was carried out according to the protocol given with the RPA II kit.

Analysis of expression. Quantification of the RPA results was obtained using phosphorimager analysis (BAS-1000, Fuji Photo Film Co., Ltd) with GAPDH mRNA levels utilized as the internal control in each case. The values for the gene under scrutiny were normalised to the internal control.

Results

RNase protection analysis. Following sectioning and histopathological examination (Table I), total RNA was isolated and gene expression was measured using RNase protection analysis. A representative image of the results obtained for each gene examined is shown in Fig. 1. The results of each analysis are described in more detail in the following sections.

Expression of IGFBP-rP1 (Mac25/TAF/PSF) in hepatoblastoma. We examined the mRNA expression levels of IGFBP-rP1 in a series of matched hepatoblastomas and the corresponding normal liver tissue from patients between the ages of 2 and 54 months. Included in the analysis were some hepatoblastomas with no counterpart normal tissues and fetal liver samples. The results of this analysis are shown in Fig. 2. In four of the eight matched tumours expression of this gene was downregulated when compared to their matched normal counterparts. In the unmatched sample tumours, two had expression which was below the average normal liver values.
downregulated in the tumours. Thus, 6 out of 12 (50%) tumours showed downregulated expression of this gene. However, in cases 3, 5 and 7 (25% of the tumours) IGFBP-rP1 was greatly upregulated in these tumours.

Expression of IGFBP-rP2 (CTGF) in hepatoblastoma. Next we examined the expression of IGFBP-rP2 in these samples. One tumour (case 7) showed greatly increased expression over its corresponding normal liver (Fig. 3). Two other samples (cases 3 and 5) showed slightly elevated levels of expression. Including the unmatched tumours, four samples (33%) have upregulated expression of IGFBP-rP2. In the matched tumours, downregulated expression of this gene was observed in three tumours (cases 2, 6 and 8). One of the unmatched tumours also showed decreased expression of this gene and two samples had unaltered expression.

Expression of IGFBP-rP4 (Cyr61) in hepatoblastoma. When IGFBP-rP4 expression was examined most of the tumours were observed to have reduced levels of expression of this gene in comparison with their matched normal liver (Fig. 4). The exceptions to this were cases 3 and 7 which showed upregulated expression. Of these, case 7 had greatly increased expression of IGFBP-rP4. In the unmatched samples two samples showed lower levels of expression than the average.
Table II. Expression patterns of the IGFB-rPs in hepatoblastomas.*

<table>
<thead>
<tr>
<th>Case no.</th>
<th>IGFBP-rP1</th>
<th>IGFBP-rP2</th>
<th>IGFBP-rP4</th>
<th>IGFBP-rP5</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>N/†</td>
<td>N</td>
<td>↓</td>
<td>↓</td>
</tr>
<tr>
<td>2</td>
<td>↓</td>
<td>↓</td>
<td>↓</td>
<td>↓</td>
</tr>
<tr>
<td>3</td>
<td>†</td>
<td>†</td>
<td>†</td>
<td>†</td>
</tr>
<tr>
<td>4</td>
<td>↓</td>
<td>↓/N</td>
<td>↓</td>
<td>↓</td>
</tr>
<tr>
<td>5</td>
<td>†</td>
<td>†/N</td>
<td>↓/N</td>
<td>↑</td>
</tr>
<tr>
<td>6</td>
<td>↓</td>
<td>↓</td>
<td>↓</td>
<td>↓</td>
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<tr>
<td>7</td>
<td>†</td>
<td>†</td>
<td>↑</td>
<td>↓</td>
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<td>↓</td>
<td>↓</td>
</tr>
<tr>
<td>9</td>
<td>N</td>
<td>↓/N</td>
<td>↓</td>
<td>↓</td>
</tr>
<tr>
<td>10</td>
<td>↓</td>
<td>↓</td>
<td>↓</td>
<td>↓</td>
</tr>
<tr>
<td>11</td>
<td>↓</td>
<td>N</td>
<td>†/N</td>
<td>↓</td>
</tr>
<tr>
<td>12</td>
<td>†</td>
<td>†</td>
<td>N</td>
<td>n.d.</td>
</tr>
</tbody>
</table>

*†, increased expression; ǂ, decreased expression; N, normal expression; N/†, normal or slightly increased expression; N/4, normal or slightly decreased expression; n.d., not determined.

Discussion

In the present study we have examined the mRNA expression of four genes whose products have recently been identified as having the ability to bind IGFs at low affinity in a series of hepatoblastomas in relation to their matched normal tissues (summarized in Table II). Previously we have reported that many members of the IGF-axis have altered expression in these samples including the IGFs which may therefore lead to increased mitogenic signalling within their surroundings (2,4). IGFBPs are critical regulators of IGF activity, and by binding to IGFs they form biologically inactive complexes which in addition to increasing the half-life of the growth factor, also modulate the binding of the IGFs to their cognate receptors (6). Thus, the newly identified IGFBP-rPs may also prove to have important roles in regulating IGF signalling in addition to their other roles (6,7).

IGFBP-rP1 has been shown to be a potential tumour suppressor (6) and decreased expression of this gene has been observed in higher stage breast cancer (13). In addition, downregulation of this gene by methylation has been shown to be important for tumorigenesis in a mouse model of liver cancer (14). As such, the decreased expression of this gene in some of our samples may reflect the loss of tumour suppressor capability, or it may indicate that increased methylation of this gene may be occurring in those samples. Cases 3, 5 and 7 however have greatly increased expression of this gene, and this may indicate an attempt to regulate or suppress the tumour.

IGFBP-rP2 or CTGF is a major mitogenic factor for connective tissue cells. It has also been shown to be specifically expressed in the malignant lymphoblasts of patients with acute lymphoblastic leukemia (15). In our samples three tumours have increased expression of this gene, with one sample in particular (case 7), having greatly enhanced expression, and may therefore lead to increased mitogenic signalling in these particular tumours.

IGFBP-rP4 (Cyr61) has also been implicated in cancer where it has been shown to be downregulated in prostate cancer (16). However overexpression of this gene in a xenograft model appears to promote tumour growth (17). In our analysis of hepatoblastomas, one sample (case 7) has a very high overexpression of this gene compared to its matched normal liver, and this may be a reflection of increased growth potential in this tumour. However, five of the samples also showed reduced expression of this gene in a manner similar to that observed for prostate cancer (16).

IGFBP-rP5 (L56/HtrA) is an interesting protein because it has serine protease activity. One feature of the IGF-axis is that the IGFBPs themselves are regulated in part by the actions of serine proteases. The cleavage of such binding proteins could therefore increase the availability of free IGFs, leading to increased mitogenic signalling. IGFBP-rP5 has been shown to be able to cleave IGFBP-5 and as such overexpression of this gene may enhance the mitogenic signalling by IGFs in the vicinity of the tumour (6). In our samples two hepatoblastomas (cases 3 and 5) showed elevated expression of this serine protease. However, in most cases the expression of this gene is reduced, and this may signal an attempt by the tumours to limit the mitogenic signalling in their vicinity. It is interesting to note that the same samples which show elevated IGFBP-rP5 also have a concomitant overexpression of IGFBP-5 indicating that the regulation of IGFBP-5 may be linked to IGFBP-rP5 activity (18).

In conclusion, the data presented indicate that in hepatoblastomas, the IGF-axis is seriously disturbed, extending even to the members of the low affinity insulin-like growth factor binding protein related proteins, confirming the importance of this axis in the pathogenesis of this disease.

Acknowledgements

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Generation of Anti-Insulin-Like Growth Factor-Binding Protein-Related Protein 1 (IGFBP-rP1/MAC25)
Monoclonal Antibodies and Immunoassay: Quantification of IGFBP-rP1 in Human Serum and Distribution in Human Fluids and Tissues

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The IGF-binding protein (IGFBP)-related proteins (rPs) are a group of recently described cysteine-rich proteins that share significant amino-terminal structural similarity with the conventional IGFBPs. IGFBP-rP1 (also known as MAC25/angiomodulin/prostacyclin-stimulating factor and T1A12), regulates cellular proliferation, adhesion, and angiogenesis and stimulates prostacyclin synthesis. We characterized new monoclonal antibodies generated against IGFBP-rP1 and have used them to study the distribution of IGFBP-rP1 in human biological fluids and tissues. Additionally, we have developed a noncompetitive sandwich-type immunoassay to quantify the concentrations of IGFBP-rP1 in human serum. IGFBP-rP1 was readily detectable in serum, urine, amniotic fluid, and cerebrospinal fluid by immunoblot analysis. Evaluation of the newly developed immunoassay demonstrated acceptable analytical performance, with a detection limit of 0.7 ng/liter, a dynamic range of 3.1-100 ng/liter, and intra- and interassay coefficients of variation of 2.5-6.8% and 3.1-6.4% at approximately 24-85 ng/ml IGFBP-rP-1, respectively. No significant cross-reactivity with IGFBP-1-6 was observed. In random normal human adult sera (n = 37), the median IGFBP-rP1 was 21.0 ng/liter, and values did not correlate with levels of IGF-I (r = 0.085, P = 0.61), IGF-II (r = 0.051, P = 0.75), or IGFBP-3 (r = 0.061, P = 0.74). The monoclonal anti-IGFBP-rP1 antibodies also readily detected IGFBP-rP1 expression in human tissue sections, with preferential expression of IGFBP-rP1 in the microvascular endothelium associated with tumorigenesis. In summary, using newly developed IGFBP-rP1 monoclonal antibodies, we confirm the presence of IGFBP-rP1 in the major human body fluids, provide quantitative normative data on the concentrations of IGFBP-rP1 in human serum, and show preferential expression of IGFBP-rP1 in the microvascular endothelium associated with tumorigenesis. The use of these novel IGFBP-rP1 detection tools should prove useful in the elucidation of the biological role(s) of this protein. (J Clin Endocrinol Metab 88: 3401-3408, 2003)

The IGF system comprises two ligands, IGF-I and IGF-II, six IGF-binding proteins (IGFBPs), IGFBP-1 to -6, and two receptors, type 1 and type 2 IGF receptors (1). Recently the IGFBP family has been expanded to include the IGFBP-related proteins (IGFBP-rPs), which share significant structural similarities with the IGFBPs (2, 3). Thus, the IGFBP superfamily includes the six conventional IGFBPs, which have high affinity for IGFs, and at least 10 IGFBP-rPs, which not only share the conserved amino-terminal domain of the IGFBPs but also show some degree of affinity for IGFs and insulin in several, but not all, assay systems (4-7).

The IGFBP-rPs are a group of cysteine-rich proteins that control diverse cellular functions, such as cellular growth, cellular adhesion and migration, and synthesis of extracellular matrix. In addition, these proteins are involved in biological processes that include development and differentiation, reproduction, angiogenesis, wound repair, inflammation, fibrosis, and tumorigenesis (3).

IGFBP-rP1 was initially identified as a gene differentially expressed in normal leptomeningeal and mammary epithelial cells, compared with their counterpart tumor cells, and named meningioma-associated cDNA (MAC25) (8). The expressed protein was independently purified as a tumor-derived adhesion factor (later renamed angiomodulin) (5, 9) and as a prostacyclin-stimulating factor (10). It has additionally been reported as T1A12, a gene down-regulated in breast carcinomas (11).

Although the biological roles of IGFBP-rP1 have not been clearly established, there is a growing body of evidence that suggests that it may act as a tumor suppressor gene. IGFBP-rP1 is preferentially expressed in normal (vs. neoplastic) meningeal, mammary, and prostatic cells (8, 12, 13); it is up-regulated during senescence of mammary and prostatic cells (14, 15); loss of heterozygosity of the IGFBP-rP1 locus has been observed in 50% of cancerous breast tissues in one...
study (11); and IGFBP-rP1 shows growth inhibitory effects when overexpressed in prostate cancer cells (16, 17) or breast cancer cells (18).

In addition, IGFBP-rP1 may have an important role in vascular biology. It has been detected in tube-like structures in vitro (19) and high endothelial cells (20, 21) and is preferentially localized in the basement membrane of neocapillaries, like those seen in tumor tissues (5, 22, 23). Furthermore, analysis of genes differentially expressed in endothelial cells indicates that IGFBP-rP1 expression is up-regulated in tumor-derived endothelium (24). A role for IGFBP-rP1 in vascular biology emerges from demonstrations that IGFBP-rP1 is capable of stimulating the synthesis of the vasodilator prostacyclin in cultured endothelial cells and may, therefore, be involved in maintaining the permeability of newly synthesized capillaries (10).

We recently generated and characterized a polyclonal anti-IGFBP-rP1 antibody and identified IGFBP-rP1 in conditioned media from cultured human mammary and prostatic cells and in human biological fluids (12, 25). In this report, we further characterize the presence of IGFBP-rP1 in human biological fluids using anti-IGFBP-rP1 monoclonal antibodies. In addition, we provide quantitative data on serum IGFBP-rP1 using a newly developed IGFBP-rP1 immunoassay and examine the distribution of IGFBP-rP1 protein in human tissues.

Materials and Methods

HPLC-purified IGFBP-1 from human amniotic fluid was kindly provided by Dr. D. R. Powell (Baylor College of Medicine, Houston, TX); rhIGFBP-2, -4, -5, and -6 were purchased from Austral Biologicals (San Ramon, CA); rhIGFBP-3, a nonglycosylated 29-kDa core protein provided by Dr. D. R. Powell (Baylor College of Medicine, Houston, TX); isolated clones: 1, 1B4A; 2, 1B4B; 3, 1B4C; 4, 2A2B; 8, 2C4A; 9, 2C4B; 10, 2C4C; and 11, 2C5.

Materials

HPLC-purified IGFBP-1 from human amniotic fluid was kindly provided by Dr. D. R. Powell (Baylor College of Medicine, Houston, TX); rhIGFBP-2, -4, -5, and -6 were purchased from Austral Biologicals (San Ramon, CA); rhIGFBP-3, a nonglycosylated 29-kDa core protein expressed in Escherichia coli was a generous gift from Celltrix, Inc. (Santa Clara, CA). C-terminally FLAG-tagged rhIGFBP-rP1, CTGF, and NovH were expressed in a baculovirus system as previously reported (2, 4, 26). Baculovirus-generated nontagged rhIGFBP-rP1 protein was purified over SP Sepharose (Sigma Chemical Co., St. Louis, MO) column equilibrated in MES buffer [50 mM N-morpholino-ethanesulfonic acid (pH 6.0), 2 mM EDTA, and 1 mM phenylmethylsulfonyl fluoride]. IGFBP-rP1 protein retained on the column was eluted with an NaCl gradient [0.2-1.0 M in 2-(N-morpholino) ethane sulfonic acid buffer] Fractions containing recombinant human rhIGFBP-rP1 were pooled and dialyzed against PBS. Analysis for protein purity and quantitation was as previously described (4). 125I-IGF-I and 125I-IGF-II were gifts from Diagnostic Systems Laboratories (Diagnostic Systems Laboratories, Webster, TX).

Nitrocellulose and electrophoresis reagents were purchased from Bio-Rad Laboratories, Inc. (Hercules, CA). Polyclonal antibodies against IGFBP-rP1FLAG, CTGFFLAG, and NovHFLAG were generated in rabbits, as previously described (2, 25, 26). The IgG fractions were purified by a protein A affinity column (Amersham, Arlington Heights, IL). Monoclonal antibodies against IGFBP-rP1 (designated no. 1 through 11) were produced against the baculovirus-generated rhIGFBP-rP1FLAG described above. Horseradish peroxidase (HRP)-linked donkey antirabbit and sheep antiguinea IgG antibodies and enhanced chemiluminescence detection reagents were purchased from Amersham.

Human biological fluids were obtained as anonymous samples from the Oregon Health Sciences University Central Laboratory and were residuals from routine clinical test samples. They were from apparently healthy adult subjects (i.e., with no known acute or chronic diseases); for cerebrospinal fluid (CSF), samples from both apparently healthy adult and pediatric subjects were analyzed. Additional serum and urine samples were collected from apparently healthy adult volunteers. The protocol was approved by the hospital Institutional Review Board, and informed consent was obtained from the healthy adult volunteers.

Adult serum samples included in the analysis were from 18 women, aged 25-76 yr and 19 men, aged 25-67 yr. On collection, blood samples were allowed to clot, then separated; after clinical testing, the residuals were used for these studies within 48 h of collection. Spot urine samples were from 50 women, aged 18-78 yr, and 10 men, aged 26-71 yr, collected after the first morning void. Urinalysis abnormalities excluded samples to be further tested for IGFBP-rP1. Samples were stored at -80 °C after centrifugation to discard the cellular pellet. Residual samples of amniotic fluid were from normal pregnancies between 15 and 20 wk gestation and were stored, after centrifugation, at -80 °C until use. Residual samples from normal CSF (two women, four men, two boys, age range for the whole group: 4-88 yr) were also stored at -80 °C until use. Urine, amniotic fluid (AF), and CSF were screened for IGFBP-rP1 within 2 months of collection.

Generation of monoclonal anti-IGFBP-rP1 antibodies

Five 10-wk-old Balb/c mice (Charles River, NC) were immunized with 50 μg/ml baculovirus-generated rhIGFBP-rP1FLAG protein with Freund's complete adjuvant (ICN Biomedicals Inc., Aurora, OH). Three boosts (100 μg rhIGFBP-rP1FLAG with Freund's incomplete adjuvant) were given at 3-d intervals. Two of the mice with high titers of antibodies against rhIGFBP-rP1FLAG were identified and were subsequently used for hybridoma generation. A third booster (75 μg rhIGFBP-rP1FLAG) was administered to the selected mice, and fusion was performed according to Lane (27).

The resulting hybridomas were screened as follows: IgG from hybridoma cell culture supernatants was captured on goat antimouse IgG-coated plates (DSL). After 2 h, the plates were washed three times with PBS-Tween 20 (0.01 m), and further incubated with 100 μl biotinylated IGFBP-rP1FLAG protein (200 ng/ml in PBS-Tween with 1% BSA and 1% goat serum). After washing with PBS-Tween, the plates were incubated with avidin-HRPO (Zymed Laboratories Inc., South San Francisco, CA) for 30 min, and the signal was developed with TMB (Life Technologies Inc., Grand Island, NY). To obtain pure clones, limiting dilution to 1 cell/well was performed for all positive hybridoma colonies and the resultant monoclonals confirmed by plate assays. The 11 clones are designated no. 1 to 11 in this report and represent the following isolated clones: 1, 1B4A; 2, 1B4B; 3, 1B4C; 4, 1D1A; 5, 1D1B; 6, 2A2A; 7, 2A2B; 8, 2C4A; 9, 2C4B; 10, 2C4C; and 11, 2C5.

Immunoadsorption of IGFBP-rP1 polyclonal antibody

Aliquots of the IGFBP-rP1 polyclonal antiserum were incubated overnight at 4 °C with either rhIGFBP-rP1FLAG or a ratio of 15 μg/μl of anti-FLAG M2 agarose beads (Sigma) were added to the aliquots above, and samples were incubated for another hour at 4 °C. Samples were pelleted (5 min, at 12,000 g) and supernatants collected and frozen for subsequent Western blot (WB) studies.

WIB studies

For the initial characterization of the IGFBP-rP1 monoclonal antibodies, conditioned medium from normal human prostate epithelial cells, which are known to secrete large amounts of IGFBP-rP1 protein (12), was used as the source of endogenous IGFBP-rP1. Equal amounts of total protein per lane were dissolved in nondenaturing SDS sample buffer [0.5 mol/liter Tris (pH 6.8), 1% SDS, 10% glycerol, and bromophenol blue] and boiled for 5 min. Samples were electrophoresed in 15% SDS-polyacrylamide gels, electroblotted onto nitrocellulose, and membranes blocked with 4% milk-TBS-T [Tris-buffered saline-Tween-20 (0.1%)] for 1 h at 22 °C. Western blots were incubated with rhIGFBP-rP1 monoclonal antibody (ιg/ml) at a 1:3000 dilution or with IGFBP-rP1 monoclonal antibodies (ιg/ml) at a dilution of 1:2000 in TBS-T overnight at 4 °C. Membranes were washed with TBS-T and incubated for 1 h at 22 °C with a 1:3000 dilution of HRP-linked anti-rabbit or anti-mouse IgG secondary antibodies. Proteins of interest were detected with enhanced chemiluminescence reagents, according to the manufacturer's protocol.

For the characterization of IGFBP-rP1 in human biological fluids, representative samples from healthy human subjects were prepared.
similarly and resolved on 15% SDS-polyacrylamide gels. Normal human serum was concentrated 10-fold using a heparin affinity column (Amersham) before these studies. Immunoblotting of nitrocellulose membranes involved: 1) both the immune and the preadsorbed fractions of the IGFBP-rP1 antiserum, and 2) anti-IGFBP-rP1 monoclonal antibody no. 5 and/or 10.

**Western Ligand blotting**

Equimolar amounts of IGFBPs and IGFBP-rPs were resuspended in SDS sample buffer and resolved on 15% SDS-polyacrylamide gels. Separated proteins were electroblotted onto nitrocellulose membranes. Membranes were rinsed in 3% IGEPAL (Sigma) in TBS-T for 30 min at 22 C, blocked with 1% BSA IGEPAL (Sigma) in TBS-T for 1 h at 22 C and incubated with 2 x 10^4 cpm of a mixture of [125I]-IGF-I and [125I]-IGF-II in 1% BSA/TBS-T overnight at 4 C. Membranes were washed, dried, and exposed to Biomax film (Eastman Kodak Co., Rochester, NY).

**IGFBP-rP1 ELISA development**

Anti-IGFBP-rP1 monoclonal antibodies were employed to construct a noncompetitive sandwich-type immunoassay (see below). The antibody selection was based on extensive pairwise-wise evaluations in both one-step (equilibrium) and two-step (sequential) immunoreaction formats. Using this protocol, combinations of the 11 different anti-IGFBP-rP1 monoclonal antibodies were analyzed. The protocol optimization was based on the initial evaluation of a number of factors that could potentially affect detection limit, dynamic range, precision, and delayed sample addition (28). Antibody combinations demonstrating favorable analytical performances were further assessed for accuracy and comparative IGFBP-rP1 determinations. The sources of the raw materials and composition of the various buffers employed have been previously described (29, 30).

IGFBP-rP1 antibody coating to microtiter wells was performed at a concentration of 0.25-20 mg/liter by using previously published methods (29, 30). The IGFBP-rP1 detection antibodies were coupled to HRP as previously described (29). IGFBP-rP1 calibrators were prepared by appropriately diluting the recombinant IGFBP-rP1 in a protein-based buffer matrix (0.05 mol/liter sodium phosphate (pH 7.4), 9 g/liter NaCl, 6 g/liter BSA, 0.5% Proclin 300). The preparation was stable for at least 5 d at 4 C and more than 6 months at -70 C.

**IGFBP-rP1 ELISA protocol**

Calibrators or samples (0.020 ml) were added in duplicate to the precoated wells, followed by addition (0.1 ml) of the detection antibody-HRP conjugate (diluted in the assay buffer to approximately 0.1-0.25 mg/liter) and 4 h of incubation at room temperature with continuous shaking. The wells were washed five times and incubated with 0.1 ml/well of the TMB/H2O2 substrate solution for 10 min. Stopping solution (0.1 ml) was then added and absorbance measured by dual-wavelength measurement at 450 nm with background wavelength correction set at 620 nm. Absorbance measurements and ELISA data analysis were performed with the Labsystems Multiskan Multisoft microplate reader (Labsystems, Helsinki, Finland).

The best performances were obtained with a coating antibody concentration of 10 mg/liter (1000 ng/0.1 ml/well), a detection antibody concentration of approximately 0.1-0.25 mg/liter (10-25 ng/0.1 ml/well), a sample size of 0.02 ml, and a 4 h one-step (equilibrium) ELISA configuration. With this protocol, the differences in assay results caused by 1- to 20-min delay between addition of the same samples into the coated wells was less than 10%.

**IGFBP-rP1 ELISA validation procedures**

The lower limit of detection (sensitivity) was determined by interpolating the mean plus 2 SD of 12 replicate measurements of the zero calibrator. The intraassay coefficients of variation were determined by repeat analysis (n = 12) of four samples at IGFBP-rP1 concentrations of approximately 10-50 µg/liter in one run and interassay coefficients of variation by duplicate measurement of the samples in 12 separate assays. Recovery was assessed by adding 25 µl recombinant IGFBP-rP1 diluted in the standard matrix to 225 µl of three sera and analyzing the spiked and unspiked samples. Percent recovery was determined by comparison of the amount of added IGFBP-rP1 with the amount measured after subtracting the endogenous IGFBP-rP1 levels. Linearity was tested by analyzing three serum samples diluted serially (2- to 8-fold) in the zero calibrator of the assay.

The standard range and performance characteristics of IGFBP-rP1 are summarized in Table 1. Analysis of IGFBP-1, IGFBP-2, IGFBP-4, and IGFBP-3 (up to 500 µg/liter) did not show any cross-reactivity or interference. There was no cross-reactivity with IGF-I or IGFBP-11 (up to 600 µg/liter) added to the assay zero standard followed by IGFBP-rP1 analysis (data not shown).

**Immunohistochemistry**

Anomalous samples of normal and neoplastic human tissues were provided by the Cancer Pathology Shared Resource of the Oregon Cancer Center. These samples had been collected in the fresh state shortly after surgical resection and either snap frozen in optimal cutting temperature embedding compound, using a dry-ice/pentane slurry, or fixed in 10% buffered formalin. The fixed tissues were processed and embedded in paraffin using standard techniques.

Five-micrometer sections of the frozen samples were prepared in a cryostat, placed on Fisherbrand Plus slides (Fisher Scientific, Pittsburgh, PA) and allowed to air dry for 15 min at room temperature. Dried slides were wrapped in cellophane and stored at -80 C. The cryostat sections were allowed to warm to room temperature and then fixed for 10 min in freshly prepared 1% paraformaldehyde/PBS. Five-micrometer sections of the paraffin-embedded tissues were cut on a microtome and placed on Fisherbrand Plus slides. The slides were deparaffinized through xylene and alcohol and then placed in TBS buffer for use in immunohistochemistry.

**Results**

**Characterization of IGFBP-rP1 monoclonal antibodies**

Anti-IGFBP-rP1 monoclonal antibodies (IgG fractions) were initially screened by WB for their ability to recognize rhIGFBP-rP1FLAG protein. As shown in Fig. 1A, baculovirus-generated rhIGFBP-rP1FLAG protein was detected by both the panel of monoclonal antibodies (no. 1 through 10) and the polyclonal anti-IGFBP-rP1 antibody. Anti-IGFBP-rP1 monoclonal antibody no. 11 was significantly less potent in rec-

**TABLE 1. IGFBP-rP1 ELISA validation data**

<table>
<thead>
<tr>
<th>Assay parameters</th>
<th>Performance characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Detection limit (µg/liter)</td>
<td>0.7 µg/liter</td>
</tr>
<tr>
<td>Standard range (µg/liter)</td>
<td>3.1-100</td>
</tr>
<tr>
<td>Intra-assay CV (%)</td>
<td>2.6-6.8</td>
</tr>
<tr>
<td>Inter-assay CV (%)</td>
<td>3.1-6.4</td>
</tr>
<tr>
<td>Recovery of added IGFBP-rP1 (%)</td>
<td>106 ± 10%</td>
</tr>
<tr>
<td>Recovery after dilution (%)</td>
<td>108 ± 5.6%</td>
</tr>
</tbody>
</table>

CV, Coefficient of variation.
Characterization of IGFBP-rP1 in normal human biological fluids by WIB

Once the sensitivity and specificity of the anti-IGFBP-rP1 monoclonal antibodies were demonstrated, we investigated the presence of IGFBP-rP1 in the major human body fluids, such as serum, urine, AF, and CSF using these antibodies. In pooled normal human serum from healthy adults, both the polyclonal anti-IGFBP-rP1 antibody and the panel of monoclonal antibodies recognized an approximately 31-kDa protein that ran at slightly higher molecular mass than the baculovirus-generated rhIGFBP-rP1FLAG. Furthermore, the specificity of this band was demonstrated by immunoblotting with an rhIGFBP-rP1FLAG preadsorbed fraction of the polyclonal antibody (Fig. 3A). Similarly, distinct and specific IGFBP-rP1 bands were detected by both the polyclonal anti-IGFBP-rP1 antibody and monoclonal anti-IGFBP-rP1 antibodies in pooled normal human urine, AF, and CSF from healthy adults (Fig. 3B). The lower sensitivity of the monoclonals can be attributed to the fact that nonreducing conditions were employed for these WIB analyses because, as demonstrated above (see Fig. 1), reducing conditions en-

![Fig. 1. Characterization of anti-IGFBP-rP1 monoclonal antibodies. Immunoreactivity of 11 anti-IGFBP-rP1 monoclonal antibodies was compared with our polyclonal anti-IGFBP-rP1 antibody (25).](image)

![Fig. 2. Specificity of anti-IGFBP-rP1 monoclonal antibodies. Monoclonal anti-IGFBP-rP1 antibody (no. 5) was tested against equimolar amounts of IGFBPs (IGFBP-1 through 6) and rhIGFBP-rP1FLAG, CTGFFLAG (2), NovHFLAG (26). Note that rhIGFBP-3 is Escherichia coli expressed and nonglycosylated, with a molecular mass of 29 kDa. Identical results were obtained with monoclonal anti-IGFBP-rP1 no. 10 (data not shown).](image)

![Fig. 3. WIB studies of IGFBP-rP1 in human body fluids. A. Two-microliter aliquots of 10-fold heparin affinity column-concentrated normal human serum were electrophoresed in triplicate and immunoblotted with polyclonal anti-IGFBP-rP1 antibody, rhIGFBP-rP1FLAG-preadsorbed fraction of anti-IGFBP-rP1 polyclonal antibody, or a representative monoclonal antibody (no. 5 and/or 10). Baculovirus-generated rhIGFBP-rP1 (100 ng) was run as control ("C"). B. Similar WIB studies were carried out with human urine (50 µl), AF (20 µl), CSF (50 µl), and rhIGFBP-rP1 (100 ng) as control ("C").](image)
hance the affinity of the monoclonals for endogenous IGFBP-rP1.

Quantification of IGFBP-rP1 in normal human serum by ELISA

An IGFBP-rP1 noncompetitive sandwich-type immuno-asset (see Materials and Methods) was developed to analyze individual human serum samples. The IGFBP-rP1 ELISA standard curve for the assay is shown in Fig. 4.

In normal human adult sera (n = 37), the median IGFBP-rP1 was 21.0 µg/liter (Table 2). The values did not correlate with IGF-I (r = 0.085, P = 0.61), IGF-II (r = 0.051, P = 0.75), or IGFBP-3 (r = 0.061, P = 0.74) levels. A sexual dimorphism in circulating IGFBP-rP1 was evident, with higher IGFBP-rP1 concentrations in male subjects (26.3 ± 6.8, n = 19 vs. 19.8 ± 9.2, n = 18; P = 0.01). In matched samples from normal adult males and females (n = 13), median IGFBP-rP1 levels were similar in serum, plasma-EDTA, and plasma-heparin (ANOVA P = 0.768) and were 20.8, 19.9, and 22.0 ng/ml, respectively. The anti-IGFBP-rP1 antibodies also immuno-detected circulating IGFBP-rP1 in other mammals (e.g. pigs and cows) but not rodents (data not shown). IGFBP-rP1 in the matched samples, in fetal bovine serum, as well as the recombinant (nontagged) form, was stable at 4°C for at least 4 d.

Distribution of IGFBP-rP1 in human tissues

Immunostaining of human tissues with IGFBP-rP1 antibodies generated against a decapptide in the C terminus of IGFBP-rP1 has been reported by Akaogi et al. (5) and more recently by Degeorges et al. (33). With our panel of characterized antibodies, the tissue distribution of IGFBP-rP1 in cryostat sections of normal and malignant human tissues was analyzed.

To ascertain the specificity of the immunostaining, parallel tissue sections (Fig. 5, A and B, lung squamous carcinoma tissue) were initially prepared with both the polyclonal anti-IGFBP-rP1 antibody and antibody preadsorbed with rhIGFBP-rP1FLAG. Figure 5A shows typical positive immunostaining (arrows), which was reduced to background with the preadsorbed fraction of the polyclonal antibody (Fig. 5B, arrows). Identical patterns of staining were observed in these sections with the monoclonal anti-IGFBP-rP1 antibodies (Fig. 5C, lung squamous carcinoma tissue).

In normal tissues (prostate, breast, and colon), immunoreactivity was weak and limited to small blood vessels (Fig. 5D, normal prostate). Diffuse staining of the stroma was also observed. The immunostaining was, however, notably enhanced in the endothelial cells of the microvasculature of most of the human cancer tissues examined (lung, prostate, colon), using either the polyclonal or the monoclonal anti-IGFBP-rP1 antibodies (Figs. 5, A, C, E, and F).

Immunohistochemical analysis of IGFBP-rP1 expression was extended to include paraffin-embedded tissue sections (data not shown). Unlike frozen tissue sections, only the monoclonal anti-IGFBP-rP1 antibodies were capable of detecting IGFBP-rP1 expression, but the overall signal was less robust than that detected in frozen tissue sections. The pattern of staining, however, remained the same, with immunostaining predominantly identified in vascular endothelium, particularly in the microvasculature of tumor tissues.

Discussion

The biological roles of IGFBP-rP1 have been evaluated in numerous studies and include diverse actions, such as tumor suppression (8, 11, 12, 14, 16, 34), stimulation of prostacyclin synthesis (10), and involvement in angiogenesis (5, 19, 24).

We previously characterized a polyclonal anti-IGFBP-rP1 antibody and identified IGFBP-rP1 in conditioned media from cultured human cells as well as in human biological fluids (25). Here we extend these studies with the characterization of new anti-IGFBP-rP1 monoclonal antibodies to study the distribution of IGFBP-rP1 in human biological fluids and tissues. These monoclonal antibodies, like the polyclonal antibody, do not cross-react with the six conventional IGFBP proteins or proteins of the CCN family. All 11 of the monoclonal antibodies specifically recognize the approximately 31-kDa secreted IGFBP-rP1 protein in conditioned medium from cultured human cells.

Here we also show for the first time quantitative data on serum IGFBP-rP1 in humans and indicate a dimorphic distribution of circulating IGFBP-rP1, with higher concentrations of IGFBP-rP1 in males. Because IGFBP-rP1 could be down-regulated by estrogens, according to one report (8), a plausible mechanism for the lower concentrations of circulating IGFBP-rP1 in females could be its tonic inhibition by estrogens. Although the assay was not developed to quantitate IGFBP-rP1 in other body fluids, the assay readily detected IGFBP-rP1 in human urine, AF, and CSF (data not shown), consistent with the findings obtained by immuno-

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**TABLE 2. IGFBP-rP1 levels in human body fluids**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Mean</th>
<th>Median</th>
<th>sd</th>
<th>Range</th>
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<tr>
<td>Serum</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All</td>
<td>23.1</td>
<td>21.0</td>
<td>8.6</td>
<td>9.9-40.0</td>
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</tr>
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<td>Male</td>
<td>26.3</td>
<td>27.3</td>
<td>6.8</td>
<td>15.6-37.9</td>
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</tr>
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<td>Female</td>
<td>19.8</td>
<td>17.5</td>
<td>9.2</td>
<td>9.9-40.0</td>
<td>18</td>
</tr>
</tbody>
</table>
FIG. 5. Immunohistochemical analyses of IGFBP-rP1 in normal and malignant human tissues. Cryostat sections of various human tissues were fixed on slides and immunohistochemistry analysis performed using either the polyclonal or representative monoclonal antibodies (no. 5 and/or 10). Representative sections are as shown. Arrows indicate regions of specific immunostaining. A, Lung squamous carcinoma immunostained with the IgG fraction of anti-IGFBP-rP1 polyclonal antibody (aIGFBP-rP1 polyclonal). B, Parallel section of A immunostained with the preadsorbed fraction of aIGFBP-rP1 polyclonal antibody. C, Lung squamous carcinoma immunostained with the IgG fraction of anti-IGFBP-rP1 monoclonal antibody no. 10. Identical results were obtained with monoclonal antibody no. 5 (data not shown). D, Normal prostatic tissue immunostained with aIGFBP-rP1 polyclonal. E, Prostate cancer stained with aIGFBP-rP1 monoclonal no. 10. F, Colon cancer stained with polyclonal aIGFBP-rP1 antibody. Magnification, ×200 (A–D) and ×400 (E and F).

Both the polyclonal and the panel of monoclonal anti-IGFBP-rP1 antibodies were capable of detecting IGFBP-rP1 in major human body fluids, such as serum, urine, AF, and CSF. Interestingly, the IGFBP-rP1 protein detected in most of these samples (with the exception of amniotic fluid) appeared to be of a slightly higher molecular mass than the rhIGFBP-rP1FLAG protein (Fig. 3). This observation may be accounted for by differences in glycosylation or other posttranslational modifications between the baculovirus-generated recombinant protein and human IGFBP-rP1 detected in body fluids. These observations are consistent with the molecular weight of IGFBP-rP1 protein detected in the conditioned media of mammalian cells (12, 15, 25).

To further characterize IGFBP-rP1 protein expression in vivo, we used the panel of anti-IGFBP-rP1 antibodies to evaluate IGFBP-rP1 distribution in tissues. For both the polyclonal and panel of monoclonal antibodies, frozen tissue sections were superior to paraffin-embedded sections for immunodetection. Identical patterns of staining in the frozen tissues were observed with polyclonal and monoclonal antibodies, and the staining appears to be specific because preclearing of the relevant IgG fraction with rhIGFBP-rP1FLAG protein abrogated the signals. Most striking was that the distribution of IGFBP-rP1 was predominantly among endothelial cells of tumor tissues, with reduced, but detectable, signals in endothelial cells of normal tissues and in the stroma of all tissues examined. Similar observations were made with the paraffin-embedded tissue sections.

Our findings concur, in part, with other reports of the IGFBP-rP1 distribution in human tissues (5, 11, 33, 35). In these previous studies, the tissue sections examined were all paraffin embedded, and immunohistochemical analysis employed an incompletely characterized monoclonal antibody (5) or a polyclonal antibody (11, 33, 35) generated against a decapeptide in the C terminus of IGFBP-rP1 and, therefore, of uncertain specificity. The latter polyclonal antibody (11) recognized only the reduced approximately 37-kDa form of IGFBP-rP1 protein (33). Because our polyclonal and monoclonal antibodies were generated against intact IGFBP-rP1 protein, the patterns and sensitivity of staining might be expected to differ from that of other laboratories. Nevertheless, Akaogi et al. (5) detected tumor-derived adhesion factor (IGFBP-rP1) immunoreactivity in the vascular membrane of small blood vessels as well as capillaries of diverse cancer tissues but not those associated with normal tissues. Degorges et al. (33), in contrast, did observe immunostaining of normal endothelial cells, but intense immunoreactivity was associated predominantly with supporting cells of peripheral nerves and stromal cells of numerous tissues.

The implication from this present study (and others) is that not only do endothelial cells express IGFBP-rP1 protein but also expression is considerably higher in endothelial cells.
associated with cancers. Intriguingly, a recent comparative study profiling gene expression in endothelium derived from normal and tumor tissues supports these observations (24). Analysis by serial analysis of gene expression determined that IGFBP-rP1 was the pan endothelial marker most abundantly expressed in endothelial cells and there was a 2-fold increase in gene expression in malignant tissues (24). These observations support an important role of IGFBP-rP1 in vascular biology and, as first proposed by Akaogi et al. (5), suggest that IGFBP-rP1 may be involved in the process of neoangiogenesis in malignancy. Furthermore, because IGFBP-rP1 appears to induce prostacyclin synthesis in endothelial cells (10), IGFBP-rP1 may also have roles in non-cancerous human vascular diseases, such as atherosclerosis and hypertension. The new tools presented here should, therefore, prove useful in dissecting out the involvement of this protein in vascular function of normal and malignant tissues.

Other than endothelial cells, IGFBP-rP1 was detected rarely in epithelial cells in our immunohistochemical analysis of IGFBP-rP1 distribution. This was somewhat unexpected because we previously demonstrated that epithelial cells, at least in vitro, do express IGFBP-rP1 protein, and in situ hybridization studies of normal prostate tissues indicate IGFBP-rP1 mRNA in glandular epithelium that surrounds the lumen (12, 15). The apparent contradiction could be due to the sensitivity of antibodies. Akaogi et al. (5), similarly, did not detect immunostaining of epithelial cells with their antibody. However, the polyclonal antibody raised against the C-terminal region of IGFBP-rP1 was reactive to the luminal epithelial cell of normal lobules and ducts of breast tissues (11); the tumor epithelial cells of prostate tissue (35); and ciliated cells of bronchial, epididymal, and fallopian epithelium (33). Thus, the variations in the immunoreactivity of tissues and cell types in each study clearly depend on the source of antibodies and, most likely, also on the method of tissue preparation and processing.

Existing data on IGFBP-rP1 indicate that its role in cancer remains to be defined. Based on recent serial analysis of gene expression analysis (24) and immunohistochemical studies, including the present report, IGFBP-rP1 is implicated in the neovascularization process so critical for tumor growth. This contrasts with a growing body of evidence that suggests IGFBP-rP1 is a potential tumor suppressor gene. In a number of cancers, such as breast (8, 11, 13, 18), meningiomas (8), prostate cancer (12), and liver tumorigenesis (36), IGFBP-rP1 expression was down-regulated, although expression appeared to be up-regulated in carcinogenesis of colon mucosa (23, 37). Interestingly, IGFBP-rP1 is associated with a 50% loss of heterozygosity (LOH) in breast cancer (11). Furthermore, overexpression of IGFBP-rP1 in a prostate cancer cell line was shown to dramatically reduce tumorigenic potential of the cell line (16, 17), and exogenous addition of the protein to cancer cell lines appeared to inhibit cell growth (38). Taken altogether, these observations suggest that although IGFBP-rP1 is clearly important in tumorigenesis, its specific role(s) is still unclear, and most likely depends on a number of factors, including the tissue and cell type under study, the sensitivity, and specificity of reagents used as well as the techniques employed.

In summary, using a panel of newly developed IGFBP-rP1 monoclonal antibodies and an immunoassay, we demonstrate that IGFBP-rP1 is detectable in serum and other biological fluids and provide quantitative data on the concentrations of IGFBP-rP1 in human serum. We also have shown distribution of IGFBP-rP1 in human normal and cancerous tissues. IGFBP-rP1 appears to be involved in vascular biology, possibly in the process of neoangiogenesis that occurs in tumor tissues. The use of these novel IGFBP-rP1 detection tools should prove useful in the elucidation of the biological role of this IGFBP related protein.

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