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Insulin-like Growth Factor II Signaling in Neoplastic Proliferation Is Blocked by Transgenic Expression of the Metalloproteinase Inhibitor TIMP-1

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Abstract. Insulin-like growth factor (IGF) II is overexpressed in many human cancers and is reactivated by, and crucial for viral oncogene (SV40 T antigen, [TA g])–induced tumorigenesis in several tumor models. Using a double transgenic murine hepatic tumor model, we demonstrate that tissue inhibitor of metalloproteinase 1 (TIMP-1) blocks liver hyperplasia during tumor development, despite TA g-mediated reactivation of IGF-II. Because the activity of IGFs is controlled by IGF-binding proteins (IGFBPs), we investigated whether TIMP-1 overexpression altered the IGFBP status in the transgenic liver. Ligand blotting showed that IGFBP-3 protein levels were increased in TIMP-1–overexpressing double transgenic littermates, whereas IGFBP-3 mRNA levels were not different, suggesting that TIMP-1 affects IGFBP-3 at a posttranscriptional level. IGFBP-3 proteolysis assays demonstrated that IGFBP-3 degradation was lower in TIMP-1–overexpressing livers, and zymography showed that matrix metalloproteinases (MMPs) were present in the liver homogenates and were capable of degrading IGFBP-3. As a consequence of reduced IGFBP-3 proteolysis and elevated IGFBP-3 protein levels, dissociable IGF-II levels were significantly lower in TIMP-1–overexpressing animals. This decrease in bioavailable IGF-II ultimately resulted in diminished IGF-I receptor signaling in vivo as evidenced by diminished receptor kinase activity and decreased tyrosine phosphorylation of the IGF-I receptor downstream effectors, insulin receptor substrate 1 (IRS-1), extracellular signal regulatory kinase (Erk)-1, and Erk-2. Together, these results provide evidence that TIMP-1 inhibits liver hyperplasia, an early event in TA g-mediated tumorigenesis, by reducing the activity of the tumor-inducing mitogen, IGF-II. These data implicate the control of MMP-mediated degradation of IGFBPs as a novel therapy for controlling IGF bioavailability in cancer.

Key words: TIMP-1 • insulin-like growth factor II • signal transduction • extracellular proteolysis • tumor suppression

 extracellular matrix (ECM) serves as the immediate microenvironment for interactions with the cell surface, besides providing the structural support for all tissues. The ECM is not static. Rather, it is dynamic in nature with a continuous turnover of its protein constituents and growth factor pools. A major determinant of ECM turnover and integrity is the extracellular proteolytic balance between secreted matrix metalloproteinases (MMPs) and their biological inhibitors (TIMPs) (for reviews see Matrisian, 1992; Denhardt et al., 1993; Mignatti and Rifkin, 1993). The function of extracellular proteolysis extends beyond ECM degradation to the processing of cell surface receptors and ligands and release of protein-bound growth factors (for review see Werb, 1997). Therefore, it is conceivable that extracellular proteolytic activity within the cellular microenvironment can directly impact cell proliferation. Despite transgenic studies showing that cellular proliferation is altered by ectopic expression of MMPs or TIMPs (Symson et al., 1994; D’Aramiento et al., 1995; Witty et al., 1995; Martin et al.,
levels in hepatic tissues. Reduced IGF-II bioactivity in vivo inhibition of MMP-mediated IGFBP-3 proteolysis leading to expression attenuates IGF-II bioactivity. This is due to the TAg-induced tumorigenesis, transgenic TIMP-1 overexpression on IGF bioactivity. Here, opportunity to investigate the potential effects of transgenic TIMP-1 overexpression (Christofori et al., 1994), the IGF pathway appears to be crucial for TAg oncoprotein transformation. Indeed, pancreatic tumorigenesis is substantially reduced following IGF-IR ablation (Sell et al., 1993), and TAg-induced pancreatic hyperplasia and liver tumorigenesis (Martin et al., 1996). In this model, we have already shown that free IGF, or IGF-IR signaling. IGFs are considered survival factors, and depending on cell type and situation, demonstrate either a direct proliferative effect (Schoene et al., 1985; Osborne et al., 1989) or an antiapoptotic effect (Bozyczko-Coyne et al., 1993; Resnicoff et al., 1995). In accordance with their diverse biological activities, there is a complex regulation of IGF bioactivity. In the extracellular compartment, IGF activity is controlled by the presence of high affinity IGF-binding proteins (IGFBPs) which bind the IGFs with high or moderate affinity than the type I IGF receptor (IGF-IR; McCuskler et al., 1991; Shimisaki and Ling, 1991). The molar excess of IGFBPs, along with their high affinity, leads to effective sequestration of IGFs by IGFBPs, resulting in little or no free IGFs in most biological systems. Nevertheless, proteolytic cleavage of IGFBPs by aspartic, serine, and metalloproteinases (Cohen et al., 1992; Conover et al., 1994; Fowlkes et al., 1994a) has been shown to release IGFs in vitro, because IGFBP fragments demonstrate significantly lower affinities for IGFs than the intact IGFBPs (Gargosky et al., 1992; Lassarre and Binoux, 1994; Lassarre et al., 1994). Once released from IGFBPs, both IGF-I and IGF-II exert their mitogenic effects through IGF-IR, a protein tyrosine kinase receptor (Osborne et al., 1989; Rubin and Baserga, 1995; Fowlkes, 1997). Therefore, the mitogenic activity of IGFs may be regulated at the level of IGF expression, IGFBP sequestration, proteinase-mediated IGFBP proteolysis to release free IGF, or IGF-IR signaling. We explored whether the TIMP/MMP balance alters cell proliferation by regulating IGF-II bioactivity in vivo by using a double transgenic mouse model that carries the tumor-inducing viral oncprotein, (SV40 T antigen, [TA g]) and the tumor-suppressing TIMP-1 transgene (Martin et al., 1996). In this model, we have already shown that hepatic TIMP-1 overexpression inhibits TA g-induced hepatocyte hyperplasia and liver tumorigenesis (Martin et al., 1996, 1999). Since TA g transformation is unsuccessful following IGF-IR ablation (Sell et al., 1993), and TA g-induced pancreatic tumorigenesis is substantially reduced by IGF-IR ablation (Christofori et al., 1994), the IGF pathway appears to be crucial for TA g oncprotein transformation. Our double transgenic model presented an ideal opportunity to investigate the potential effects of transgenic TIMP-1 overexpression on IGF bioactivity. Here, we demonstrate that despite IGF-II reactivation during TA g-induced tumorigenesis, transgenic TIMP-1 overexpression attenuates IGF-II bioactivity. This is due to the inhibition of MMP-mediated IGFBP-3 proteolysis leading to elevated IGFBP-3 levels and reduced dissociable IGF-II levels in hepatic tissues. Reduced IGF-II bioactivity in vivo was confirmed by decreased signaling through the IGF-IR signal transduction pathway. To our knowledge, this is the first in vivo example of the modulation of growth factor bioactivity by the regulation of extracellular proteolysis.

**Materials and Methods**

**Transgenic Mice**

Transgenic mice expressing the TIMP-1 (Ts') or TA g (TA g') transgenes in liver were generated and bred as described previously (Martin et al., 1996). Single transgenes were crossed to generate four categories of littermates designated as wild-type controls (TA g and TA g'), TIMP-1 controls (Ts'), TA g controls (TA g'), and double transgenic TIMP-1-overexpressing (TA g/Ts') mice. Female littermates were killed at specified ages, and the liver tissue was processed and embedded or flash frozen for analyses.

**Immunoprecipitation and Western Blotting**

Lever tissue was homogenized in lysis buffer (20 mM Tris-HCl, pH 7.4, 1.0% NP-40, 150 mM NaCl, 0.5% PM SF, 1 mM EDTA, 10 μg/ml peptatin, 10 μg/ml leupeptin) at 4°C. Samples were centrifuged for 10 min at 16,000 g, the supernatants collected, and the protein content determined by the Bradford assay. A liquids containing 2.5 μg of protein were adjusted to a volume of 50 μl. TA g was immunoprecipitated by adding 0.5 μg/ml of anti-SV40 40 large small t antibody (clone PA b 108; Pharmingen) and 50 μg of Gamma Bind Plus Sepharose (Amersham Pharmacia Biotech), and rocking for 2 h at 4°C. Immunoprecipitates were collected by centrifugation in a refrigerated Eppendorf centrifuge and washed three times with NET/gel buffer (50 mM Tris-HCl, pH 7.4, 100 mM NaCl, 0.1% NP-40, 1 mM EDTA, 0.25% gelatin, 0.02% sodium azide). The samples were subjected to 10% SDS-PAGE and electrophoresed in formaldehyde agarose gels, subjected to Northern blotting and hybridization with [α-32P]dCTP-labeled and random primed cDNA probes for murine IGF-II (cDNA obtained from Dr. G. Bell, University of Chicago, Chicago, IL), rat IGF-BP-3 (cDNA, obtained from Dr. A. Herington, Queensland University of Technology, Brisbane, Australia), 18S ribosomal RNA, and glyceraldehyde 3-phosphate dehydrogenase (GAPDH) were performed. The latter two probes were used to control for equal loading and transfer of samples.

**Northern Blotting**

Total RNA isolated from individual liver tissue samples (20 μg) was electrophoresed in formaldehyde agarose gels, subjected to Northern blotting and sequential hybridizations with [α-32P]dCTP-labeled and random primed cDNA probes for murine IGF-II (cDNA obtained from Dr. G. Bell, University of Chicago, Chicago, IL), rat IGF-BP-3 (cDNA, obtained from Dr. A. Herington, Queensland University of Technology, Brisbane, Australia), 18S ribosomal RNA, and glyceraldehyde 3-phosphate dehydrogenase (GAPDH) were performed. The latter two probes were used to control for equal loading and transfer of samples.

**Immunohistochemistry and In Situ Hybridization**

Paraffin sections of formalin-fixed liver tissue were generated as described previously (Martin et al., 1999). Clone PA b 108, biotinylated goat anti-mouse IgG, and streptavidin-peroxidase conjugate (Zymed) were used for TA g immunohistochemistry, and peroxidase-conjugated anti–mouse IgG antibodies and ECL chemiluminescence (Amersham Pharmacia Biotech).

**Northern Blotting**

Total RNA isolated from individual liver tissue samples (20 μg) was electrophoresed in formaldehyde agarose gels, subjected to Northern blotting and sequential hybridizations with [α-32P]dCTP-labeled and random primed cDNA probes for murine IGF-II (cDNA obtained from Dr. G. Bell, University of Chicago, Chicago, IL), rat IGF-BP-3 (cDNA, obtained from Dr. A. Herington, Queensland University of Technology, Brisbane, Australia), 18S ribosomal RNA, and glyceraldehyde 3-phosphate dehydrogenase (GAPDH) were performed. The latter two probes were used to control for equal loading and transfer of samples.

**Ligand Blotting**

Liver samples were homogenized as for Western blotting. Equivalent amounts of protein from each sample (40 μg) were subjected to 10% SDS-PAGE followed by electroblotting to Hybond N. Membranes were blocked by incubating for 1 h in blocking buffer (10% BSA in TBS, 0.01 M Tris-HCl, pH 7.5, 0.15 M NaCl) followed by overnight hybridization with 50,000 cpm/ml [32P]-IGF-II (Amersham Pharmacia Biotech) in blocking
buffer. Membranes were washed three times for 15 min with TBS, air dried, then subjected to autoradiography.

**IGFBP-3 Substrate Zymography**

Liver samples were prepared as for Western blotting. Protein from each sample (40 μg) was subjected to SDS-PAGE in 10% polyacrylamide gel containing 1.0 μg/ml recombinant human IGF-BP-3 (kindly provided by Dr. C. M.ack, Celtrix Pharmaceuticals, Santa Clara, CA). Following electrophoresis, gels were washed for 30 min with 2.5% Triton X-100. Gels were equilibrated with transfer buffer (50 mM Tris-HCl, pH 8.0, 5 mM CaCl₂), then the proteins were subjected to capillary transfer onto PVDF membranes as described previously (Fowlkes et al., 1997). The transfer was performed either in the absence or presence of 10.0 μg/ml recombinant human TIMP-1 (kindly provided by Dr. H. Nagase, University of Kansas, Kansas City, KS). Transfers took place overnight at 37°C. Under these conditions, only proteolytic fragments of IGF-BP-3 transfer from the membrane. Intact IGF-BP-3 (30 kD) impregnated in the polyacrylamide matrix remains in the gel (Fowlkes et al., 1997). Following transfer, the PVDF membrane was probed with polyclonal antibody against IGF-BP-3 (Fowlkes et al., 1997).

**IGFBP-3 Degradation Assay**

Liver tissue was homogenized as for Western blotting, except that NP-40 and proteinase inhibitors were not added. 50 μg of protein from each sample was brought up to a volume of 28 μl with homogenization buffer containing 50 μM CaCl₂, 125I-IGFBP-3 (50,000 cpm/ml; Diagnostic Systems Laboratories) was added, and the samples were incubated for 5 h at 37°C, followed by SDS-PAGE, electroblotting, and autoradiography (Davenport et al., 1993).

**Affinity Chromatography**

Liver protein homogenates were prepared as above. Total levels of IGF-II were quantified according to a published protocol (D e León and As- merom, 1997). In brief, 10 μl from each sample was dot blotted onto nitrocellulose in triplicate. The membrane was incubated in blocking buffer and then with mAb against IGF-II. The IGF-II concentrations were determined from a standard curve generated with known quantities of purified recombinant IGF-II (kindly provided by Eli Lilly and Co.). To assess the relative levels of dissociable IGF-II, equivalent amounts of total IGF-II from each sample (1.2 μg) were incubated with IGF-BP-4-conjugated Affigel overnight at 4°C with gentle rocking. Samples were centrifuged for 10 min at 1,000 g and the pellets washed three times with TBS. The pellets were then incubated with 0.5 M acetic acid for 5 min to elute IGF-II bound to the immobilized IGF-BP-4. Supernatants were collected and dried under vacuum. Samples were resuspended in TBS and the relative levels of dissociable IGF-II determined using the IGF-II dot blot assay. To validate that increasing the molar ratio of IGF-BP-3 to IGF-II affects IGF-II binding to the IGF-BP-4 Affigel affinity media, increasing amounts of recombinant IGF-BP-3 (10 μl) were mixed with a constant amount of recombinant IGF-II and the mixtures were subjected to the IGF-BP-4 chromatographic separation, as described above.

**Receptor Kinase Assays**

Frozen liver tissue samples were homogenized in TBS containing 1% Triton X-100, 1 mM PM SF, 100 μM Na₃VO₄, 1 mM EDTA, 10 μg/ml pepstatin, 10 μg/ml leupeptin. A aliquots from each supernatant containing 5.0 μg of protein were immunoprecipitated as above, using antibodies against the IGF-IR (clone aR3; Calbiochem). Immunoprecipitates were then incubated with kinase reaction buffer (10 mM Hapes, pH 7.4, 5 mM M-Cl₂, 5 mM MgCl₂, 100 μM Na₃VO₄, 50 mM NaF) containing 2.0 μCi/μl [γ⁻³²P]ATP either with or without 50 μg Poly (Glu, Tyr) 4:1 (Sigma Chemical Co.) for 30 min at 37°C. Samples were electrophoresed on 10% SDS-PAGE Gels, dried onto Whatman 3 M Paper, and autoradiographed.

**Phosphorylation Levels**

Liver proteins were isolated as for the receptor kinase assays. For insulin receptor substrate (IRS)-1, 2.5 mg of protein from each sample was immunoprecipitated with polyclonal antibodies against rat IRS-1 (Upstate Biotechnology) as described above. The immunoprecipitates were then Western blotted and the membranes probed sequentially with antiphosphotyrosine (clone 4G10; Upstate Biotechnology) and anti-IRS-5 antibody. For mitogen-activated protein kinase (MAPK), replicate 10% SDS-PAGE Gels were run with 40 μg of protein from each sample loaded on each gel. Following electroblotting, membranes were probed with antibodies specific for phospho-MAPK or MAPK (New England Biolabs).

**Densitometry and Quantification**

A autoradiographs were scanned using a Molecular Dynamics Densitometer. A biorbance was quantified using ImageQuant® software. Statistical significance was determined using t test. For all gel electrophoresis, each lane corresponds to a tissue sample taken from an individual mouse. A II samples were obtained from 185-d-old female mice unless indicated other- wise, as we determined previously that TIMP-1 modulation significantly affected TA g-induced preneoplastic proliferation at this age (Martin et al., 1999).

**Results**

**TA g Antigen Molecular Interactions Are Maintained in TIMP-1-overexpressing Mice**

We have shown previously that transgenic TIMP-1 expression does not affect TA g oncprotein levels in double transgenic mice that coexpress TA g and TIMP-1 trans- genes (TA g⁺/Ts⁺) (Martin et al., 1996). Equivalent TA g protein levels in both TA g⁺ and TA g⁺/Ts⁺ animals are confirmed here in Fig. 1 a (top panel). Since TA g binds to and inactivates the tumor suppressor gene products p53 and Rb to induce hyperplasia (Tan et al., 1986; DeCaprio et al., 1988), we investigated whether TA g interactions with these tumor suppressor proteins are altered by TIMP-1 overexpression. The amount of p53 and Rb that communoprecipitated with TA g protein was examined by Western blotting and was found to be similar between TA g⁺ and TIMP-1-overexpressing (TA g⁺/Ts⁺) livers, as shown in Fig. 1 a (middle and bottom panels). This indicates that TA g interactions with p53 and Rb proteins are intact, and unaffected by the elevation of TIMP-1.

**Cellular Proliferation Is Inhibited by Transgenic TIMP-1 Despite IGF-II Reaction**

IGF-II is a fetal mitogen in rodents, and its transcription is normally repressed in adult tissues by p53 (Zhang et al., 1996). However, focal reactivation of IGF-II and its local- ization to proliferating cells during TA g-induced tumorigenesis have been reported in two independent transgenic tumor models (Schirmacher et al., 1992; Christofori et al., 1994). Cellular proliferation and tumor development have been shown to be profoundly inhibited in one such model when crossed onto an IGF-II-null background (Christo- fori et al., 1994), revealing the fundamental importance of IGF-II reactivation in TA g-induced tumorigenesis. IGF-II reactivation has been frequently observed during TA g-induced hepatocarcinogenesis (Casola et al., 1995; Haddad and Held, 1997). To investigate whether IGF-II was reactivated in our hepatocellular carcinoma model, we ex- amined its spatiotemporal expression in the livers of experimen- tal and control littermates. IGF-II mRNA was not detected by Northern blot analysis of liver tissue from wild-type mice of all ages, or in the livers of TA g⁺ mice before 165 d of age. However, from 170 d of age, multiple IGF-II transcripts commonly observed in mouse and hu- man tissues (Daughaday and Rotwein, 1989; Casola et al., 1995) were expressed (Fig. 1 b, and data not shown). Fig. 1 c
IGF-II mRNA levels in the livers of TAg transgene–expressing; TAg littermates. Sequential probing of the blot showed that the levels of p53 and pRb that communoprecipitated with TAg were similar in TAg and TAg/Ts littermates. (b) Northern blot analysis of IGF-II mRNA levels in the livers of TAg and TAg/Ts transgenic littermates. IGF-II mRNA was undetectable in livers of non-TAg expressing littermates, but was reactivated between 153 and 191 d of age in TAg-expressing littermates. (c) IGF-II mRNA levels were the same in 185-d-old TAg and TAg/Ts transgenic littermates. RNA was isolated from livers of mice from the indicated age and genotype (wt, wild-type; Ts, TIMP-1 sense transgene-expressing; Ta, TIMP-1 antisense transgene-expressing; Ta*, TAg g transgene-expressing; TAg g/Ts*, double transgenic TIMP-1-overexpressing). 18S ribosomal RNA and GAPDH provided a means of determining RNA loading on the Northern blots.

shows that IGF-II mRNA was expressed in liver tissue from TAg and TAg/Ts littermates at 185 d of age. Densitometric analysis of the major IGF-II transcript (4.2 kb) confirmed that expression was comparable in TAg and TAg/Ts littermates (3.3 ± 1.2 vs. 3.8 ± 0.6, n = 5 per group). The timing of IGF-II reactivation coincided with TAg-induced liver enlargement, and remained unaffected by TIMP-1 overexpression. In contrast to IGF-II mRNA reactivation, IGF-I mRNA expression remained very low and unchanged during all stages of TAg tumorigenesis (data not shown).

The liver-specific C reactive protein promoter, which directs TAg expression (Rüther et al., 1993), resulted in uniform TAg expression in almost all hepatocytes by 185 d of age, as shown by immunohistochemistry (Fig. 2 a). In adjacent sections, in situ hybridization revealed IGF-II mRNA production by most hepatocytes (Fig. 2 b), and extensive hepatocytic hyperplasia was detected by PCNA immunostaining (Fig. 2 c). The liver of a 185-d-old double transgenic TIMP-1-overexpressing (TA g+/Ts−) littermate demonstrated the same ubiquitous expression patterns of both the TAg oncoprotein (Fig. 2 d) and IGF-II mRNA (Fig. 2 e). In contrast, probing the adjacent sections with anti-PCNA antibody revealed far less proliferation in the livers of TAg g+/Ts− mice (Fig. 2 f) compared with TAg g+ mice. In our previous study using morphometric analysis, we found this proliferation to be significantly suppressed in TIMP-1 transgenic liver tissue (Martin et al., 1999). Together, the results demonstrate that the effects of TIMP-1 are exerted downstream of IGF-II reactivation, but before cell proliferation.

For further analyses of the effects of TIMP-1 on IGF-II bioactivity, we chose to use liver specimens from 185-d-old mice. This choice was based on our previous pilot study encompassing 20–250 d, in which we found that hepatocyte proliferation was maximal (>65%) in TAg mice at 185 d of age, and was inhibited 3.3-fold in TAg g+/Ts− littermates at this age (Martin et al., 1999). The difference in proliferation was most accentuated at this age, and therefore we anticipated that in vivo analysis of molecular factors in the IGF-II signaling pathway would be most clearly resolved at this point in time. The above observations that IGF-II is reactivated in both TAg g+ and TAg g+/Ts− livers (Fig. 1 b and Fig. 2, b and e), yet hepatocyte proliferation is only prevalent in TAg g+ tissue at 185 d (Fig. 2, c and f), further supported the use of this age group for IGF-II bioactivity studies.

**IGFBP-3 Levels Are Elevated Due to Reduced Proteolysis**

IGFBPs regulate IGF activity by sequestering free IGFs, thus preventing ligand–receptor interactions. Previous studies from one of our laboratories have demonstrated that MMPs, the primary proteinases inhibited by TIMP-1, can degrade IGFBPs both in vitro and in vivo (Fowlkes et al., 1994a,b; Thrailkill et al., 1995; Fowlkes, 1997). Since the amount of high-affinity, intact IGFBP may be regulated in part by MMP-mediated proteolysis, we compared the levels of intact hepatic IGFBPs using [125]IGF-II Western ligand blotting. Of the hepatic IGFBPs that bound [125]IGF-II on ligand blots, only IGFBP-3 (42–46-kD doublet) levels were strongly affected by TIMP-1 modulation (Fig. 3 a). The levels of this binding protein, which is also the major serum carrier protein for IGFs (Jones and Clemmons, 1995), were increased by more than twofold in the livers of TIMP-1-overexpressing mice (TA g+/Ts−) compared with levels in TAg g− littermates (Fig. 3 d, left panel; P < 0.02). TIMP-1 overexpression did not affect the levels of IGFBP-4 (24–26 kD), although there were notable minor elevations of a doublet of proteins at 28–32 kD that, based on molecular mass, likely represent IGFBP-1, -2, and/or -5 (Fig. 3 a).

In contrast to IGFBP-3 protein levels, Northern blot analysis showed that TIMP-1 overexpression did not affect the levels of IGFBP-3 mRNA (Fig. 3 b), attributing the differences in IGFBP-3 protein levels to posttranscriptional events. MMP-1, -2, -3, and -9 have been shown to proteolytically cleave IGFBP-2, -3, and -5, a process inhibited by TIMP-1 in vitro (Fowlkes et al., 1994a,b; Thrailkill et al., 1995; Fowlkes, 1997). Here, IGFBP-3 substrate zymography (Fowlkes et al., 1997) was used to identify...
IGFBP-3-degrading proteases in liver homogenates. We found two IGFBP-3-degrading activities with molecular masses of \( \sim 62 \) and \( \sim 84 \) kD (Fig. 3 c). Both activities were substantially reduced in the presence of recombinant TIMP-1 (Fig. 3 c), demonstrating that the IGFBP-3-degrading proteinases were MMPs. Next, we determined whether a decreased proteolysis of IGFBP-3 is evident in liver tissue obtained from TIMP-1-overexpressing mice. Liver homogenates were analyzed for their ability to degrade \( ^{125}I \)-IGFBP-3 into smaller molecular weight species, as described in Materials and Methods. There was significantly less degradation of \( ^{125}I \)-IGFBP-3 by liver homogenates from TIMP-1-overexpressing (TA \( g^+ \)/Ts\( s^+ \)) mice compared with TA \( g^+ \)/littermates (Fig. 3 d, right panel; \( P < 0.02 \)). These data suggest that TA \( g^+ \) can induce MMPs that are capable of degrading IGFBP-3, and that coexpression of TIMP-1 can reduce MMP activity, thereby inhibiting IGFBP-3 degradation. Together, these actions allow for a net increase in tissue IGFBP-3 protein levels.

**Dissociable IGF-II Levels Are Reduced by Transgenic TIMP-1**

We determined whether the increased IGFBP-3 levels in the TIMP-1-overexpressing liver tissue affected the amount of dissociable IGF-II. Fig. 4 a shows that using a dot blot procedure developed to quantify total IGF-II (bound and unbound; De Leon and A smer, 1997), recombinant IGF-II can be measured linearly over a range of concentrations (0.4–3.2 \( \mu g \)). Next, dissociable IGF-II was measured by IGFBP-4-conjugated Affigel affinity chromatography. Using recombinant IGF-II and recombinant IGFBP-3 in different molar ratios, we were able to confirm that increasing molar ratios of IGFBP-3 reduced
the amount of IGF-II that bound to the Affigel (Fig. 4 b).

To measure difference in levels of dissociable IGF-II in liver samples, we first quantified total IGF-II using the dot blot assay. Aliquots containing equivalent amounts of total IGF-II were subjected to the affinity chromatography procedure. TIMP-1 overexpression resulted in a sixfold decrease in dissociable IGF-II levels in the livers of TAg<sup>1</sup>/Ts<sup>1</sup> mice compared with their control TAg<sup>1</sup> littermates (Fig. 4 c; *P*, 0.02). This demonstrates that despite an equivalent extent of IGF-II reactivation, the level of dissociable or bioavailable IGF-II is reduced in TIMP-1–overexpressing animals.

**Reduced Signaling through the IGF-IR**

A reduction in the levels of dissociable or bioavailable IGF-II should result in decreased signaling through the IGF-IR pathway in the livers of TAg<sup>1</sup>/Ts<sup>1</sup> mice. Unlike postnatal IGF-II inactivation that occurs in normal rodent liver, the IGF-IR is expressed at constitutive levels in adult
mouse liver, and IGF-II exerts its mitogenic effect through this receptor (Osborne et al., 1989; Rubin and Baserga, 1995). Therefore, IGF-IR kinase activity, as well as the phosphorylation status of the downstream signaling effectors, IRS-1 and MAPK, were assessed. IGF-IR immunoprecipitated from TA g'/Ts' liver tissue exhibited lower autophosphorylation (data not shown) and showed reduced kinase activity on an exogenous substrate (Fig. 5 a), compared with receptor from TA g' controls. Moreover, tyrosine phosphorylation of IRS-1, which binds to IGF-IR following activation, was also decreased in TIMP-1–overexpressing livers (Fig. 5 b, top panel). Sequential probing of the same blot with an antibody against nonphosphorylated IRS-1 showed that IRS-1 protein levels were not altered in TIMP-1–overexpressing animals (Fig. 5 b, bottom panel). Phosphorylation of the downstream signaling molecules, the MAPKs extracellular signal regulatory kinase (Erk)-1 and Erk-2, was also reduced in TIMP-1–overexpressing liver tissue (Fig. 5 c, top panel), whereas the absolute levels of these proteins were unaffected (Fig. 5 c, bottom panel). These data provide direct evidence that the protein levels of the IGF-IR downstream signaling mediators were not altered, but that signaling from the IGF-IR was attenuated in TIMP-1–overexpressing transgenic tissue.

**Discussion**

**Regulation of the IGF-II Pathway by TIMP-1**

IGFs are critical growth factors involved in growth, transformation, and tumorigenesis and act through the IGF-IR (for reviews see Jones and Clemmons, 1995; Rubin and Baserga, 1995, Werner and Le Roith, 1997). Studies have shown that cells null for the IGF-IR do not display the normal increase in proliferation in response to growth factors or serum as seen in normal cells, and that all phases of the growth cycle are prolonged (for review see Resnicoff and Baserga, 1998). Indeed, in several cell lines, abrogation of the IGF-IR has resulted in enhanced apoptosis. Furthermore, IGF-IR–null cells cannot be transformed by TA g (Sell et al., 1993), activated Ha-ras, a combination of both, or by the overexpression of other growth factor receptors. And finally, a number of studies have shown that ablation of this receptor in tumor cell lines significantly...
decreases their tumorigenic potential in vivo. Together, these data demonstrate that interruption of the ligand-receptor interaction between IGFs and the IGF-IR effectively disrupts several aspects of the tumorigenic process. In nature, under homeostatic circumstances, little or no IGFs are present in the free or bioavailable form due to their sequestration by one or more of the six known high-affinity IGFBPs. Because IGFBPs demonstrate equal or higher affinities for IGFs than does the IGF-IR, little or no IGFs are normally available to interact with receptors. Recent studies have begun to elucidate mechanisms by which IGFs can be released from IGFBPs so they may interact with cell-surface receptors and exert their mitogenic and metabolic effects. These studies have demonstrated that a primary phenomenon invoked to release IGFs from IGFBPs is through decreasing the affinities of IGFBPs for IGFs. The best-characterized mechanism involved in decreasing the affinities of IGFBPs is proteolytic degradation (for review see Fowlkes, 1997).

Proteolytic cleavage has been demonstrated for at least five IGFBPs, IGFBP-2 to IGFBP-6, and occurs in several physiologic as well as pathologic circumstances (Jones and Clemmons, 1995; Fowlkes, 1997), yet little is known of the identity of these IGFBP-degrading proteinases in vivo, or the mechanisms that regulate the proteolytic cleavage. In vitro, we and others have provided evidence that production of proteinases, a common feature of transformed cells (Alexander and Werb, 1989), can result in degradation of IGFBP-IGF complexes, releasing IGFs to interact with cell-surface receptors, thereby triggering proliferation of target cells. We have also shown recently that in vitro, MMPs function as IGFBP-3- and IGFBP-5-degrading proteinases (Fowlkes et al., 1994a,b; Thrailkill et al., 1995).

In addition to degrading IGFBPs, extensive data also support the role of MMPs in tumorigenesis, angiogenesis, and metastasis (Alexander and Werb, 1989; Flaimenhaft and Rifkin, 1991; Liotta et al., 1991; Werb, 1997). In the current studies, we now demonstrate the importance of MMP-mediated IGFBP degradation in neoplastic proliferation in vivo, and a means of controlling this degradation.

We have used a double transgenic, TA g-based tumor model to determine directly whether TIMP-1 inhibits IGF bioactivity in vivo to suppress hepatocyte proliferation that leads to tumorigenesis. We selected this model as it provided important features: (a) IGFs are crucial mitogens for TA g-induced transformation, proliferation, and tumorigenesis (Schirmacher et al., 1992; Sell et al., 1993; Christofori et al., 1994; Casola et al., 1995; Haddad and Held, 1997); and (b) transgenic TIMP-1 overexpression in this model substantially inhibits TA g-induced proliferation and hepatocellular carcinoma (Martin et al., 1996, 1999), making it possible to measure in vivo differences in molecular factors and IGF-IR signal transduction in transgenic tissue. A systematic analysis has now been undertaken to explore connections that might exist between TA g expression and IGF-II bioactivity on cell proliferation to define the molecular mechanisms behind TIMP-1-mediated tumor suppression in vivo. The events explored herein and the results are summarized in Fig. 6, and are discussed below.

First, an examination of p53 and Rb in these mice showed that TIMP did not interfere with the molecular interactions between TA g and these tumor suppressor proteins. Next, we found that IGF-II expression was indeed reactivated at the onset of cell proliferation, similar to the
model of TAg-induced pancreatic tumorigenesis (Christofori et al., 1994), indicating a key mitogenic role for IGF-II in our tumor model. IGF-II reactivation is frequent in TAg-induced liver tumorigenesis (Casola et al., 1995; Haddad and Held, 1997), as well as in liver tumor formation induced by oncogene or growth factors (Liu et al., 1997; Harriss et al., 1998). Furthermore, our data showing that TIMP-1 significantly inhibited hepatocellular proliferation despite IGF-II reactivation suggested that TIMP-1 might directly act at a posttranscriptional level to modulate IGF-II activity. Our investigations reveal the novel finding that hepatic TIMP-1 overexpression specifically inhibits IGFBP-3 proteolysis, leading to a significant elevation of hepatic IGFBP-3 levels. We have determined that MMPs induced during TAg tumorigenesis (Martin et al., 1999) appear to function in degrading IGFBP-3. Furthermore, we demonstrate that dissociable IGF-II levels are decreased in TIMP-1-overexpressing hepatic tissue. The physiologic consequence of reducing dissociable IGF-II levels (e.g., a reduction in IGF-II bioactivity) was confirmed by demonstrating a significantly reduced signal transduction from the IGF-I receptor, as measured by reduced IGF-I receptor kinase activity and tyrosine phosphorylation levels of IRS-1 and MAPK. Thus, despite TAg-induced reactivation of IGF-II in the liver, the transgenic TIMP-1-mediated increase of IGFBP-3 levels blocks TAg-induced hepatocyte proliferation by effectively reducing bioavailable IGF-II levels. Together, our data provide direct evidence that the inhibition of extracellular proteolysis by TIMP-1 attenuates the bioactivity of the tumor-inducing growth factor IGF-II.

Transgenic TIMP or MMP Modulation and Effects on Early Tumor Development

Based on previous studies in transgenic systems, a relationship has begun to emerge between TIMP/MMP expression within a tissue and the tissue’s susceptibility to tumor development. We have demonstrated that overexpression of TIMP-1 in the liver inhibits hepatocellular carcinoma (Martin et al., 1996), and its elevation in the skin compromises the ability of transplanted lymphoma cells to grow as a primary tumor (Küger et al., 1997), whereas a reduction of TIMP-1 in these tissues augments tumor development (Martin et al., 1996; Küger et al., 1997). Consistent with these observations, the overexpression of MMP-3 (stromelysin-1) in mammary tissue leads to spontaneous mammary tumor development (Sympron et al., 1995) and the transgenic expression of MMP-1 (type I collagenase) in skin augments carcinogenesis (D’Armiento et al., 1995). In addition, the ablation of MMP-7 (matrilysin) impairs colorectal tumor development in the min mouse tumor model (Wilson et al., 1997). In many of these studies, ectopic TIMP or MMP expression results in altered cellular proliferation. For example, hepatic TIMP-1 overexpression inhibits hepatocyte hyperplasia (Martin et al., 1999), MMP-3 overexpression leads to mammary epithelial hyperplasia (Sympon et al., 1994; Witty et al., 1995), and MMP-1 overexpression to epidermal hyperproliferation (D’Armiento et al., 1995). Despite the many reports indicating that shifts in the extracellular proteolytic balance have a strong influence on early tumor development and on cell proliferation within the afflicted organ, the molecular mechanisms for these effects have remained elusive. Having previously excluded the effects of TIMP-1 on hepatocyte apoptosis (Martin et al., 1999), our present investigation provides in vivo evidence of a link between the inhibition of extracellular proteolysis, reduced growth factor bioavailability, and reduced cellular proliferation. We have shown previously that this reduction in cellular proliferation precedes suppressed tumor development in this model (Martin et al., 1996).

MMP Substrates of Importance other than ECM during Tumorigenesis

TIMPs have traditionally been considered regulators of cell invasion and motility by virtue of their ability to inhibit MMP-mediated ECM degradation. Now it is recognized that in addition to the effects on ECM structural proteins, extracellular proteolysis has the potential to control the release of growth factors tethered to the ECM (Dallas et al., 1995; Whitelock et al., 1996), the processing of soluble growth factor binding proteins, i.e., IGFBPs (Fowlkes, 1997), and the processing of cell surface molecules such as membrane-bound TNF-α, the Notch receptor in Drosophila (Bielob, 1997), and the FGFR receptor 1 (Levi et al., 1996). Although these and other studies (Kimura et al., 1998) support the overall concept that extracellular proteolysis can broadly influence cell proliferation, behavior, and fate, the observations were made in vitro systems. To our knowledge, our data are the first in vivo evidence to substantiate the importance of proteolytic degradation of a tethering or binding/carrying protein for a growth factor in the pathological development of cancer, and of its regulation by the natural inhibitor, TIMP-1. Our data demonstrate that the dynamics of IGF-II bioactivity can be altered in vivo by proteolytic modulation, and emphasize that factors other than ECM proteins constitute important physiological targets of MMP-mediated cleavage within the extracellular microenvironment.

Role of IGF-II in Tumorigenesis and Hepatocarcinogenesis

The importance of IGFs in cellular transformation has been demonstrated by findings that cells which do not express IGF-I receptors cannot be transformed by any of a number of dominant oncogenes (for review see Werner and Le Roith, 1997). Indeed, TAg was unable to transform IGF-IR-null fibroblasts, despite the presence of various growth factors present in serum (Sell et al., 1993). Thus, IGF-IR is necessary for TAg-mediated transformation to take place. Consistent with these findings, TAg expression has been shown to reactivate IGF-II gene transcription in vivo. IGF-II reactivation occurs in pancreatic beta cells transgenically expressing TAg (Christofori et al., 1994), and here we show that IGF-II reactivation occurs in hepatocytes expressing TAg transgene coincident with the onset of hepatocyte hyperplasia. Similar to our finding, other investigators have reported that TAg expression in the liver tissue coincides with reactivation of IGF-II gene transcription and this reactivation has been associated with neoplastic transformation (Casola et al., 1995; Haddad and Held, 1997). Such reactivation of IGF-II has also been re-
ported during c-myc- and TGF-α-induced hepatocellular carcinomas in transgenic mice (Liu et al., 1997; Harris et al., 1998). The consequence of IGF-I or II reactivation as it relates to neoplastic transformation has only been addressed recently in TAg models of pancreatic cancer and hepatocellular carcinoma. In the former study, the investigators bred TAg transgenics into an IGF-II–null background (Christofori et al., 1994), whereas in the latter TAg transgenics were bred into an Igf2 (+/−) background (H addad and Held, 1997). In both instances, tumor size and incidence were decreased, strongly suggesting a role for IGF-II in the pathogenesis of these two tumor types. Furthermore, several transgenic lines overexpressing IGF-II demonstrate that IGF-II is associated with tumor formation, including mammary tumors, lymphomas, and hepatocellular carcinomas (Rogler et al., 1994; Bates et al., 1995; for review see Wolf et al., 1998). Although these studies provide compelling support for IGF-II as a causal mitogen in the tumorigenesis evidenced in the current animal model, final resolution of this hypothesis must await studies examining our findings in models that are either null or up-regulated for IGF-II. Such studies are currently being pursued in our laboratories.

Significance of Regulated IGF Bioavailability in Clinical Medicine

Both IGF-I and IGF-II are widely implicated in promoting several human cancers, including liver, prostate, and breast cancer, and their expression correlates with poorer prognosis (Osborne et al., 1989; Tennant et al., 1996; Werner and L ER oth, 1996; Sohda et al., 1997; Chan et al., 1998; Hankinson et al., 1998). Furthermore, in transgenic models, IGF-I and IGF-II are causally implicated in tumorigenesis (Rogler et al., 1994; Bates et al., 1995; Haddell et al., 1996; for review see Wolf et al., 1998). These studies point to IGF bioactivity as a target for cancer therapeutics. Our results indicate that TIMP-1–like biomolecules or synthetic MMP inhibitors may be promising candidates for the therapeutic modulation of IGF dosage in novel clinical strategies. A Laternatively, strategies to alter specific IGFBP levels or the production of proteinase-resistant IGF BPs (Chernausek et al., 1995; Conover et al., 1995; Imai et al., 1998; Rees et al., 1998) may prove to be effective therapeutic interventions. A distinct feature of all of these approaches will be to target the bioavailability rather than the production of a growth factor.

The results presented here provide compelling evidence for a novel mechanism by which endogenous TIMPs contribute to the cellular microenvironment. We demonstrate that the inhibition of extracellular proteolysis in vivo impairs the activity of a specific growth factor responsible for hyperplasia during TAg-induced tumorigenesis. Because TIMPs are also capable of inhibiting invasion, metastasis, and angiogenesis (Hokha et al., 1989; DeClerck et al., 1992; Hokha, 1994; A nand-A pte et al., 1997; Krüger et al., 1997, 1998; Wang et al., 1997; Martin et al., 1999), all of which are promoted by IGF action (Bae et al., 1998; D unn et al., 1998), the combined outcome of TIMP-1 elevation may be to suppress multiple stages of tumor development, maintenance, and progression.

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