Overexpression of Human Aspartyl(Asparaginyl)β-Hydroxylase in Hepatocellular Carcinoma and Cholangiocarcinoma

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Abstract

To characterize genes that become upregulated with malignant transformation of human hepatocytes, a library of monoclonal antibodies was produced against the FOCUS hepatocellular carcinoma cell line. Antibody FB-50 reacted with an antigen that was highly expressed in 4 of 10 primary hepatocellular carcinomas, in all 20 cholangiocarcinomas we studied, and in a variety of transformed cell lines. This antigen was also highly expressed in neoplastic epithelial cells of breast and colon carcinomas in contrast to its low level of expression in normal hepatocytes and in non-neoplastic epithelial cells. Among the normal adult tissues studied, high levels were observed only in proliferating trophoblastic cells of the placenta and in adrenal glands. A 636-bp partial cDNA, isolated from a γGT11 expression library generated with HepG2 human hepatoblastoma cells, and a complete cDNA, generated by reverse transcriptase-PCR, identified the antigen as the human form of aspartyl(asparaginyl)β-hydroxylase. This enzyme catalyzes post-translational hydroxylation of β carbons of specific aspartyl and asparaginyl residues in EGF-like domains of certain proteins. Analyses of extracts prepared from several human tumor cell lines compared to their normal tissue counterparts indicate that the increase in hydroxylase, ~10-fold, is controlled at the level of transcription and the protein is expressed in an enzymatically active form. In similar analyses, comparing hepatocellular carcinomas to adjacent uninvolved liver from five patients, enzymatic activity was much higher in the tumor tissue from the four patients whose immunoblots revealed increased hydroxylase protein in the malignant tissue. EGF repeats in the extracellular domain of Notch or its homologs contain the consensus sequence for hydroxylation. Deletion mutants lacking this domain are gain-of-function mutants, suggesting that the domain modulates signal transduction by the cytoplasmic domain. While the function imparted by β hydroxylation is unknown, our studies raise the possibility that β hydroxylation is regulated in proteins like the mammalian Notch homologs, whose cytoplasmic domains have been shown to be oncogenic. (J. Clin. Invest. 1996. 98:1313–1323.) Key words: transformation • oncogene • EGF domain • β hydroxylation • hepatoma

Introduction

Hepatocellular carcinoma (HCC) is one of the most prevalent tumors in the world today, occurring with especially high frequency in sub-Saharan Africa and the Far East. Despite this, there is actually very little information available regarding the antigens properties of human hepatoma cells (1). Such information could facilitate the development of improved methods for liver tumor detection and could help elucidate the mechanisms underlying malignant transformation of hepatocytes. To study cell surface changes associated with malignant transformation of these tumors, several libraries of mAbs (2–4) have been produced for the FOCUS human hepatocellular carcinoma cell line (5). One of these mAbs, FB-50, has allowed us to characterize an antigen that is highly expressed in some primary HCCs and a variety of transformed cell lines, including HCC-derived cell lines. In addition, the antigen is highly expressed in all cholangiocarcinomas we have studied. We describe the cellular expression of this antigen in transformed hepatic and bile duct epithelial cells, establish its molecular identity as the human aspartyl(asparaginyl)β-hydroxylase (HAAH) enzyme by cDNA cloning and demonstrate that it is expressed in an enzymatically active form.

The bovine form of this enzyme recently has been purified to homogeneity, cloned, and expressed (6–8). It is an α-ketoglutarate–dependent dioxygenase (9–11) that stereospecifically catalyzes the posttranslational hydroxylation of the β carbon of specific aspartyl and asparaginyl residues in one or more EGF-like domains of certain proteins, if the EGF-like domains possess the consensus sequence (12) required by the enzyme. The possible relationship to the malignant phenotype of regulated aspartyl(asparaginyl)β-hydroxylase in EGF-like domains of proteins like the mammalian Notch homologs, known to be involved in cell differentiation and whose cytoplasmic domains have been shown to be oncogenic (13–14) is discussed.

1. Abbreviations used in this paper: HAAH, human aspartyl(asparaginyl)β-hydroxylase; HCC, hepatocellular carcinoma; ORF, open reading frame.
Methods

The following reagents and kits were obtained commercially: all protease inhibitors, catalase, bovine serum albumin, digoxigenin dUTP, antidigoxigenin fluorescent isothiocyanate isomer I, and propidium iodide (Sigma Chemical Co., St. Louis, MO); α-keto[1-14]Cl-glutaric acid (51.8 mCi/mmol), and Solvalbe (DuPont-NEN, Boston, MA); ferrous ammonium sulfate (Fisher Scientific Co., Fairlawn, NJ); α-keto-glutarate (Aldrich Chemical Co., Milwaukee, WI).

Cell lines and human tissues. Most cell lines employed were obtained from the American Type Culture Collection (Rockville, MD) and maintained on Dulbecco's Modified Eagle's medium (M.A. Bio-products, Walkersville, MD) supplemented with 10% fetal calf serum heat inactivated at 56°C, 10 μM nonessential amino acids, 1,000 IU/ml of penicillin, and 100 μg/ml of streptomycin. The human hepatoma FOCUS cell line was developed in the Wands' laboratory and has been characterized previously (5). Samples of normal human tissues, HCCs paired with adjacent uninvolved liver, and human breast and colon carcinomas were obtained by surgical excision or postmortem examination. These specimens were snap-frozen in liquid nitrogen and stored at −80°C. 20 surgical or autopsy cholangiocarcinoma specimens fixed in 2% paraformaldehyde and paraffin embedded were also studied.

Production of the mAbs. The FOCUS human hepatoma cell line was used for immunization. An early passage of this cell line from the original primary HCC tumor was used. Primary immunizations of female Balb/c mice were accomplished intraperitoneally with 4 × 10⁸ intact cells/ml that had been removed from tissue culture plates by EDTA/versene buffer and resuspended in 50% Freund’s complete adjuvant. After 6–10 wk, the secondary immunizations were performed by an intravenous injection of 4 × 10⁹ FOCUS cells in 200 μl PBS, 3 days later, splenocytes from immunized mice were fused with the NS1 myeloma cell line and the resulting hybridomas were maintained and selected in hypoxanthine-aminopterin-thymine–containing medium.

Identification, purification, and specificity testing of mAbs. Antibody-producing hybridoma cells with initial immunoreactivity to FOCUS cells were subsequently screened by binding activity to a panel of human tumor cell lines. Positive hybridomas were identified, and then cloned twice by limiting dilution. Subsequent specificity testing of cloned hybridomas was performed against various cell lines by an indirect binding radioimmunoassay using 125I-labeled sheep F(ab')₂ anti-mouse immunoglobulins (specific activity: 8–12 μCi/μg protein; DuPont-NEN) as described (1). Ascites fluids were prepared by intraperitoneal injection of hybridoma cells into Balb/c mice primed with 2.6,10,14-tetramethylpentadecane (Pristane; Aldrich Chemical Co.). mAbs from double-cloned cell lines of the IgG subclass were purified from the ascites-fluid by using a Sepharose 4B protein-A affinity column (Pharmacia LKB Biotechnology Inc., Piscataway, NJ). The protein concentration was determined by the method of Lowry (15). Purified mAbs were radioiodinated with 125I using the Iodogen method (Pierce Chemical Co., Rockford, IL) (16), to a specific activity of 5–15 μCi/μg protein.

Immunoperoxidase staining. Paraformaldehyde fixed (2%), paraaffin-embedded postmortem and surgical specimens of primary HCC and cholangiocarcinomas, together with adjacent uninvolved liver were studied. In addition, sections of breast and colon carcinomas, normal human liver, and other normal human tissues were also fixed and paraffin embedded. The sections (8-μm thick) were deparaffinized in xylenes, rehydrated in graded alcohol solution, and equilibrated in PBS. After an overnight incubation at 4°C with FB-50 mAb (5 or 10 μg/ml), the sections were immunostained by the avidin–biotin horseradish peroxidase complex method, according to the manufacturer’s protocol (Vector Laboratories, Inc., Burlingame, CA), and with 3,3'–diaminobenzidine (0.5 mg/ml in 0.03% hydrogen peroxide) as the chromagen. Nonspecific binding was defined as binding of a 125I-labeled nonrelevant mAb (antitettanus toxoid B2TT) used under the same conditions.

Tumor cell binding assay. Focus cells (10⁶), removed from plates with EDTA/versene buffer, were resuspended in 100 μl of PBS/20% fetal calf serum and incubated at room temperature for 1 h with 125I-FB-50 mAb (10⁶ cpm). The cells were then washed three times and bound radioactivity was counted in a gamma well counter.

Enzymatic activity assay. Aspartyl/asparaginylβ-hydroxylase activity was assayed as described previously (6, 7, 10) using as the substrate the first EGF-like domain of bovine protein S with an asparagine replacing the aspartic acid at position 18. Incubations were carried out at 37°C for 30 min in a final volume of 40 μl containing 48 μg of crude cell extract protein and 75 μM EGF substrate.

cDNA library screening and DNA sequencing. A γgt11 expression vector library constructed from the human hepatoblastoma cell line HepG2 (4) was immunoscreened with 125I-FB-50 mAb. About 10⁶ independent recombinant plaque-forming units were screened on blotted filters with 125I-FB-50 mAb as described (4), and one clone with persistent immunoreactivity was plaque purified. The partial cDNA clone was subcloned into the pTI7 blue TA cloning vector (Novagen, Inc., Madison, WI) after PCR amplification using γgt11 screening amplifier primers (Clontech, Palo Alto, CA), and then transformed into Escherichia coli. Miniprep plasmid DNA was prepared from transformed bacteria, and one clone-containing insert was sequenced. DNA sequencing was performed along both strands using the dideoxynucleotide chain termination method (Sequenase version 2.0 T7 DNA polymerase; United States Biochemical Corp., Cleveland, OH) and α-35S-labeled dATP (DuPont-NEN). The DNA sequence was assembled using the Mac Vector Software version 4.1 and analyzed using a Sequence Analysis Software of the Genetics Computer Group version 7.3 as implemented on a MicroVAX II computer (17).

Identification of the complete HAAH cDNA. Total cellular RNA was extracted from the FOCUS cell line cell with a commercially prepared phenol-guanidine isothiocyanate reagent according to the directions of the manufacturer (Ultraspex RNA isolation system; Biotex Laboratories, Houston, TX). RNA was reverse transcribed using RNase H-free reverse transcriptase (Superscript II; Life Technologies, Inc., Grand Island, NY) and random hexamer primers. Two specific oligonucleotide primers, selected from the HAAH open reading frame sequence and modified to include 5'KpnI–BamHI and 3'Xhol–EcoRI

Figure 1. Immunoreactivity of 125I-FB-50 mAb with intact human tumor cell lines as determined by a direct binding assay (see Methods).
restriction enzyme sites, were synthesized in an Applied Biosystem DNA synthesizer: 5'-GGTACCAGGATCCA-3' and 5'-CTCGAGGAATTC-3'. 30 PCR cycles were carried out for 1 min at 94°C, 2 min at 55°C. and 3 min at 72°C, followed by a final 10-min extension at 72°C. The resultant 2,374-bp DNA amplification product was purified from agarose gel, subcloned into the pT7 blue TA cloning vector, and sequenced.

**Expression of bacterial recombinant fusion protein and production of rabbit polyclonal antibodies.** The bacterial expression vector pTrcHis B (Invitrogen Corp., San Diego, CA) was used to subclone the partial FB-50 cDNA into the BamHI and EcoRI sites. This vector was chosen because it encodes a 5'-G-His sequence, which can be used to isolate the resulting fusion protein by chelate chromatography. The open reading frame was confirmed by sequencing both strands at the 5'-end. Recombinant fusion protein induction in transformed E. coli was achieved by the addition of 1 mM isopropyl-β-d-thiogalactopyranoside during the log growth phase. The fusion protein was purified using ProBond resin (Invitrogen Corp.). Two rabbits were immunized by intramuscular injections of 0.5 mg purified recombinant fusion protein emulsified with Freund’s complete adjuvant. Booster injections were performed 2 and 4 wk later with 0.25 mg purified fusion protein emulsified with Freund’s incomplete adjuvant. Sera were obtained 7 d after the final injection.

**Northern blot analysis.** Samples of 15 μg total cellular RNA were electrophoretically fractionated in 1% denaturing agarose gels and transferred to nylon membranes (Hybond N+; Amersham Corp., Arlington Heights, IL). After prehybridization in Rapid Hybrid buffer (Amersham Corp.), the blots were hybridized for 3 h at 65°C with the partial FB-50 cDNA probe labeled with [α-32P]dCTP (DuPont-NEN) using the Megaprime labeling method (Amersham Corp.). Blots were washed at 65°C in 0.1% SSC and 0.5% SDS for 30 min and subjected to film autoradiography at −80°C. To assess RNA integrity and the relative loading in each lane, the blots were stripped and reprobed with [α-32P]ATP labeled synthetic 30 mers corresponding to 18S ribosomal RNA in probe excess.

**Subchromosomal localization of the HAAH gene.** Fluorescent in situ hybridization analysis was performed as described by Pinkel et al. (18) using a human hydroxylase genomic clone as the probe (Genome Systems, Inc., St. Louis, MO).

## Results

**Distribution of antigen for the FB-50 mAb.** The FB-50 mAb was initially selected because of its high level of binding to the FOCUS cell line used as the immunogen. Antigen distribution was assessed by determining the binding of 125I-FB-50 mAb to a variety of intact HCC and other transformed cell lines. As shown in Fig. 1, 125I-FB-50 mAb exhibited high level binding to the three human HCC lines (Hep 3B, Hep G2, FOCUS), to

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<table>
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<th>Comments</th>
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<td><strong>Table I. Expression of Proteins in Human Carcinomas and in Normal Human Tissue that React with the FB-50 mAb as Revealed by Immunoperoxidase Staining</strong></td>
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<td>Heterogeneous staining among tumor cells within and between tumors.</td>
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<td>Cholangiocarcinoma</td>
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<td>Diffuse homogeneous staining of all transformed bile duct cells in all tumors studied.</td>
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<td>Breast carcinoma</td>
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<td>Heterogeneous staining among tumor cells.</td>
</tr>
<tr>
<td>Colon carcinoma</td>
<td>6/10</td>
<td>Heterogenous staining among tumor cells within and between tumors.</td>
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<td>Proliferating tissue</td>
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<tr>
<td>Term placenta</td>
<td>4/4</td>
<td>Diffuse homogeneous staining of the placental trophoblastic cells of the chorionic villi.</td>
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<tr>
<td>Normal liver</td>
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<td>Normal bile ducts</td>
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<td>No staining.</td>
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<td>2/2</td>
<td>Intense staining of cells in the medulla and a subpopulation of cells in the zona fasciculata and reticularis.</td>
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<td>Lung</td>
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*All 10 HCCs were reactive with the FB-50 mAb. However, while the level of immunoreactivity in four of the tumors was clearly greater than in normal liver, in the other six HCCs examined, diffuse weak staining similar to that seen in normal liver was observed.
human adenocarcinoma cell lines derived from lung (Calu 6, SK-lu-1), breast (BT 20), and colon (LS180), and to a melanoma cell line (SK-Mel-5). Lower degrees of binding were observed with Chang, Hela, cervical carcinoma (C-33A), and neuroblastoma (IMR-32) cell lines as well as to another lung (A427) and another breast (SK-BR-3) cell line. In contrast, normal human lymphocytes were unreactive.

The cellular localization of antigen in human tissues was confirmed by immunohistochemical staining studies (Table I). Normal hepatocytes exhibited diffuse and very low level expression. Four HCCs exhibited high levels of FB-50 immunoreactivity, while the remaining six had the staining pattern seen in normal liver. Distribution of antigen was heterogeneous among the cells in the tumors with high level immunoreactivity. By contrast, all 20 cholangiocarcinomas exhibited high levels of FB-50 immunoreactivity, while the normal liver and bile ducts (C) have little immunoreactivity. Cholangiocarcinoma stained with a nonrelevant mAb (D) served as a negative control.

Figure 2. Representative examples of immunoperoxidase staining of two cholangiocarcinoma tissues (A and B). There was strong homogeneous cytoplasmic staining of transformed bile ducts in all 20 cholangiocarcinoma specimens studied, while the normal liver and bile ducts (C) have little immunoreactivity. Cholangiocarcinoma stained with a nonrelevant mAb (D) served as a negative control.

Molecular cloning of the antigen: its identification as HAAH. A 636-bp cDNA clone was obtained after screening of a Hep G2 GT11 library with 125I-FB-50 (mAb). The Hep G2–derived cDNA had a single major open reading frame (ORF) capable of encoding for a 212 amino acid protein, as illustrated by the underlined sequences in Fig. 4. The ORF had neither a methionine start codon nor a termination codon, but was in phase with the bacteriophage-derived lac Z gene read-

Figure 3. Representative examples of immunoreactivity. FB-50 immunoreactivity in colon (A–D) and breast (E–G) carcinoma, and in normal placenta (H). Intense FB-50 immunoreactivity in well differentiated glandular secretory neoplastic epithelial cells in both colon (A and B) and breast (E) carcinomas. With loss of glandular architecture, FB-50 immunoreactivity was diminished (C, F, and G). No immunoreactivity was observed with the primary antibody omitted (D).
Elevated Aspartyl(Asparaginyl)/Hydroxylase in Malignancy
ing frame, indicating that the epitope recognized by the FB-50 mAb derived from a linear amino acid sequence. When the FB-50 cDNA and deduced amino acids were compared with the available sequences in the data banks, it was found to be 81% homologous to a region in the enzyme bovine asparaginyl/asparagine \textit{/H9252} hydroxylase (7) and 99% homologous to the same region of the human form of the enzyme cloned from an osteosarcoma cell line MG-63 (19). The partial FB-50 cDNA clone was shown to lack 480 nucleotides in the amino terminal coding region, as well as 1,450 nucleotides spanning the COOH-terminal coding and 3\textit{/H11032} untranslated regions. The full length ORF (Fig. 4) was obtained using specific oligonucleotide-based PCR (see Methods).

To further confirm that the FB-50 mAb recognized HAAH and to insure the specificity of the expression cloning technique, the following experiments were performed: first, polyclonal antibodies were produced against a recombinant fusion protein encoded by the FB-50 partial cDNA (B) (A). FOCUS cells were homogenized in 50 mM Tris-HCl buffer, pH 7.4, containing 1% Tween and 0.5% sodium deoxycholate, 0.1% EDTA, 2 mM EGTA, 10 µg/ml phenylmethylsulfonyl fluoride, 0.2 µg/ml leupeptin, 2 µg/ml soybean trypsin inhibitor, 0.2 µg/ml aprotonin, and 2 µg/ml N-tosyl-l-phenylalanine chloromethyl ketone. Following SDS-PAGE of protein samples (60 µg each) and transfer to Immobilon-P (Millipore Corp., Bedford, MA), the membranes were blocked for 1 h at room temperature with PBS, pH 7.5, containing 5% nonfat dry milk, and then probed for 2 h at 20°C with mouse ascites fluid FB-50 diluted to 1:5,000 or 10 µg/ml of purified FB-50 mAb in PBS containing 0.1% Tween 20 and 5% nonfat dry milk. Antibody binding was detected with horseradish peroxidase–conjugated sheep anti–mouse IgG (Amersham Corp.) diluted 1:5,000, using the ECL system (Amersham Corp.). C was blotted with preimmune rabbit serum.

Figure 5. Western immunoblots demonstrating that rabbit polyclonal antibodies raised against a fusion protein encoded by the FB-50 partial cDNA (B) recognize the same cellular proteins (arrows) in FOCUS cell lysates as does the FB-50 mAb (A). FOCUS cells were homogenized in 50 mM Tris-HCl buffer, pH 7.4, containing 1% Tween and 0.5% sodium deoxycholate, 0.1% EDTA, 2 mM EGTA, 10 µg/ml phenylmethylsulfonyl fluoride, 0.2 µg/ml leupeptin, 2 µg/ml soybean trypsin inhibitor, 0.2 µg/ml aprotonin, and 2 µg/ml N-tosyl-l-phenylalanine chloromethyl ketone. Following SDS-PAGE of protein samples (60 µg each) and transfer to Immobilon-P, the membranes were blocked for 1 h at room temperature with PBS, pH 7.5, containing 5% nonfat dry milk, and probed for 2 h at 20°C with mouse ascites fluid FB-50 diluted to 1:5,000 or 10 µg/ml of purified FB-50 mAb in PBS containing 0.1% Tween 20 and 5% nonfat dry milk. Antibody binding was detected with horseradish peroxidase–conjugated sheep anti–mouse IgG (Amersham Corp.) diluted 1:5,000, using the ECL system (Amersham Corp.). C was blotted with preimmune rabbit serum.

Figure 4. Nucleotide and deduced amino acid sequences of cDNA encoding HAAH. The amino acid sequence is denoted under the nucleotide sequence in the standard one-letter code. Amino acids are numbered, starting with the first in-frame methionine as 1, the codon of which is indicated in bold. The translation termination codon is designated by ***, and indicated in bold type. The amino acid residues obtained by immunoscreening and the corresponding FB-50 partial cDNA are underlined. The amino acids and nucleotides found different from the human sequence cloned from the osteosarcoma cell line MG-63 (19) are indicated in relief. A putative transmembrane domain is bracketed. A putative basic dipeptide endoplasmic reticulum retention motif (22) is underlined and indicated in bold type. The His-2 motif (8, 23) is underlined with a dashed line.
protein derived from the FB-50 mAb partial cDNA clone. As shown in Fig. 5, the FB-50 mAb and the polyclonal antibodies raised against the recombinant fusion protein recognized the same cellular proteins in FOCUS cell lysates. Rabbit preimmune serum used as a negative control did not recognize FB-50 protein bands. In addition, immunoprecipitation of a FOCUS cell lysate with both antibodies followed by a Western immunoblot analysis with the reciprocal antibodies also revealed the same protein bands (data not shown). The multiple bands detected (Fig. 5) reflect the proteolysis of the extended, noncatalytic NH₂-terminal region of the hydroxylase that has been described for the bovine enzyme (6).

Characteristics of the HAAH cDNA. The complete ORF (Fig. 4) contains 2,277 bp that encode for a predicted protein having 759 amino acids and a molecular weight of 85 kD. Nearly complete identity (99%) was found between this sequence and that reported by Korioth et al. (19), but there were differences at residues 565 (Tyr to Ile), 575 (Trp-Trp-Thr changed to Cys-Gly), 585 (Asp to Gln) and 709 (Arg to Lys). A nucleotide difference was also found at position 161 (TCG to TCA) that does not modify the corresponding Ser residue. It is noteworthy that the amino acids predicted at residues 565, 575, 585, and 709 from the cDNA derived from FOCUS cells were also found in the bovine sequence (7), suggesting conservation of these residues between the two species.

Bovine aspartyl(asparaginyl)β-hydroxylase is associated with the endoplasmic reticulum (10). Computer analysis of the deduced HAAH sequence revealed one area of high hydrophobicity as judged by the Kyte and Doolittle model (20). This hydrophobic region (also present in the bovine sequence) is located between residues 54 and 75 and is flanked by positively charged amino acid residues at its amino-terminal side and negatively charged residues at its carboxyl-terminal side. This hydrophobic region is the only putative transmembrane domain in the protein; together with the positions of the adjacent basic and acidic amino acids, it very likely functions as the signal anchor of a type II transmembrane protein (21). The predicted orientation of the hydroxylase would have its NH₂ terminus in the cytoplasm and the COOH terminus, containing the catalytic domain, in the lumen of the endoplasmic reticulum (N₁₅₋₁₇). Furthermore, the Arg-Lys at residues 3 and 4 is an NH₂-terminal basic motif that has been shown to target type II membrane proteins to the endoplasmic reticulum (22).

**Figure 6.** Northern blot analysis of HAAH gene expression in human tumor cell lines. An 18S RNA oligonucleotide has been used as a control for assessment of RNA loading and integrity.

**Figure 7.** Western immunoblot analysis of HAAH extracted from lung and liver carcinoma cells compared to their normal tissue counterparts. Confluent cell cultures were harvested and washed with PBS. Cells (10⁶) were suspended in 3 ml ice-cold lysis buffer (50 mM Tris-HCl, pH 7.4, 200 mM NaCl, 10 μM DTT, 0.9% NP-40, and the following protease inhibitors: 1 μg/ml aprotinin; 1 μg/ml leupeptin; 1 μg/ml pepstatin; 1 mM PMSF; 0.01%, soybean trypsin inhibitor). The cells were then lysed on ice by sonication using three 30-s pulses at 1-min intervals at 50% maximum intensity (Sonifier 450; Branson Ultrasonics Corp., Danbury, CT). The lysate was centrifuged at 105,000 g for 60 min. Proteins in the supernatant (~20 mg/ml) were precipitated by adding 0.7 g solid (NH₄)₂SO₄ to 1 ml. After 60 min at 4°C, the suspension was centrifuged and the pellet was dissolved in 0.5 ml lysis buffer to a protein concentration of 30 mg/ml. 1 g of thawed human lung or liver was cut into smaller pieces that were washed with PBS, resuspended in 10 ml ice-cold lysis buffer, and homogenized on ice three times at top speed (Tissumizer; Tekmar Co., Cincinnati, OH) for 10 s at 1-min intervals. The homogenates were subjected to sonication, and then centrifuged as above. Proteins in the supernatant (~40 mg/ml) were precipitated with (NH₄)₂SO₄ and dissolved in 1 ml of lysis buffer to a final concentration of 30 mg/ml (as described above). Protein (100 μg) from each extract was subjected to SDS-PAGE. After transfer to nitrocellulose, immunoblotting was carried out as follows: the membranes were washed and blocked as described in Fig. 4; blotting was for 18 h at 4°C with rabbit monospecific antisera to the 52 kD bovine hydroxylase (6); an identical membrane was also blotted with a mouse anti-human actin mAb as an internal control; after washing, bound antibodies were detected either by goat anti-mouse IgG antibodies coupled to alkaline phosphatase (Bio Rad Laboratories, Hercules, CA) or by goat anti-rabbit IgG antibodies coupled to horseradish peroxidase using the ECL system. The level of hydroxylase protein in the Hep G2 cells was at least 13-fold greater than its level in normal liver, while in the A549 cells it was approximately sixfold more abundant than in normal lung. Estimates were made by scanning densitometry after normalizing for the internal controls.
Although the bovine enzyme has been shown to be in the endoplasmic reticulum and the cDNA predicts that HAAH would also reside there, FB-50 reacts with intact cells (Fig. 1). This indicates that the epitope recognized by FB-50 is present on the surface of these cells. This situation may be analogous to that of the protein ERGIC-53, which contains an endoplasmic reticulum retrieval signal, but which, when overexpressed, is transported to the cell surface (23).

The overall identity with the bovine enzyme is 68%, but, in regions suggested to be functionally important (7), conservation is considerably higher: for the COOH-terminal catalytic domain, amino acids 310–759 (90%); for the putative transmembrane domain, amino acids 54–75 (100%). In addition, the “His-2 motif” (amino acids 679–697), present in α-ketoglutarate–dependent dioxygenases (8, 24) and containing a histidine essential for Fe²⁺/α-ketoglutarate binding (8), is identical.

Finally, in a fluorescence in situ hybridization analysis for subchromosomal localization of the gene, all 80 of the metaphase cells examined by in situ hybridization exhibited specific labeling of the proximal long arm of chromosome 8. Measurements of 10 such chromosomes demonstrated that the gene is located 14% of the distance from the centromere to the telomere of chromosome arm 8q, an area corresponding to band 8q12 (data not shown). Band 8q12 is a region known to undergo translocations that are associated with salivary gland pleomorphic adenomas (25, 26), although there is as yet no evidence that these adenomas contain altered levels of HAAH.

### Table II. HAAH Activity of Crude Extract from Human Carcinoma Cells and Normal Tissue Counterparts

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<th>Enzyme source</th>
<th>Activity* (CO₂ released, in picomoles)</th>
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<tr>
<td>A549 cell culture</td>
<td>239±10</td>
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<tr>
<td>Human lung tissue</td>
<td>25±4</td>
</tr>
<tr>
<td>Hep G2 cell culture</td>
<td>133±16</td>
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<tr>
<td>Human liver tissue</td>
<td>14±0.1</td>
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*The average value of two assays, each run in duplicate.

### Table III. HAAH Activity of Crude Extracts from Human Hepatocellular Carcinoma Tissues and from Adjacent Uninvolved Liver

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HAAH gene expression in transformed cell lines. Northern blot analysis of total cellular RNA, using the partial FB-50 cDNA as a probe (Fig. 6), revealed a major transcript of ~2.8 kb and a minor transcript of ~5 kb in the FOCUS, Hep G2, Hep 3B, and HuH 7 human HCC cell lines, the SK-Hep 1 liver adenocarcinoma cell line, and in transformed Chang cells. Transcripts were also detected in a number of other human tumor cell lines. Consistent with the 125I-FB-50 mAb binding studies, little, if any, transcript was detected when total cellular RNA from normal human liver was analyzed. As has been reported (19), when polyA⁺ RNAs were analyzed, transcripts of roughly equivalent abundance were detected in preparations from normal liver, lung, heart, kidney, pancreas, skeletal muscle, and placenta and, at lower abundance, from brain. These transcripts were found to be elevated ~10-fold in several human tumor cell lines relative to their normal tissue counterparts (data not shown). For both the bovine (7) and human (19), the size of the ORFs (~2.3 kb) are such that the smaller of the two transcripts are large enough to contain the entire ORF. While the sizes of the larger transcripts are consistent with that of the full length cDNA, the relative contributions of the two transcripts to cellular levels of the enzyme remain to be determined.

Elevated levels of hydroxylase protein and activity in carcinoma cell lines and in primary HCCs. When Western blotting
was carried out using a monospecific antiserum (6) directed against the carboxyl-terminal catalytic domain of the bovine hydroxylase (as opposed to the FB-50 mAb, which is directed against an epitope in the amino terminal noncatalytic region of the human enzyme), the levels of hydroxylase protein in Hep G2 cells and A549 cells were greater than in normal liver and lung tissue, respectively (Fig. 7). The molecular weight of the major species in both the carcinoma cells and in normal tissues is $\sim 72$ kD. With longer exposure (data not shown), the blots of the carcinoma cell preparations reveal multiple molecular weight species (50–120 kD)$^2$. Previously, multiple species of hydroxylase (52–90 kD) were identified in bovine liver (6).

When equivalent amounts of total protein from preparations of normal lung or liver or from the carcinoma cell lines were assayed for hydroxylase activity, the activities in the carcinoma cells were substantially higher ($\sim$ 10-fold) (Table II). The lower levels of enzyme activity in the extracts of normal tissues did not result from the presence of an endogenous inhibitor(s) since, in incubations of extracts to which were added purified bovine hydroxylase, there was no inhibition of the activity of the bovine enzyme (data not shown). Furthermore, the specific activities of the hydroxylase found in the normal lung and liver are quite comparable to those previously reported for bovine liver and mouse L cells (9, 10); thus, the marked increases seen with the carcinoma cell lines represent true increments over what is observed in normal tissue.

When similar analyses were made comparing HCCs to adjacent uninvolved liver from five patients, enzymatic activity was much higher in the tumor tissue from the four patients whose immunoblots revealed increased hydroxylase protein in the malignant tissue (Fig. 8, Table III). Thus, the increase in hydroxylase enzymatic activity extends to malignant tissue.

Discussion

To characterize cellular genes that are upregulated with malignant transformation of human hepatocytes, a library of mAbs was produced against the FOCUS HCC cell line. The antigen recognized by the mAb FB-50 was particularly abundant. The fact that immunoreactivity was maintained in Western blots under reducing conditions suggested that the epitope was linear. Accordingly, it was possible to clone the antigen, which was identified as HAAH, by directly immunoscreening the FOCUS cDNA library. Our studies indicate that high level expression of HAAH is not restricted to the FOCUS cell line; rather, it is found with a wide variety of tumor cell lines. Furthermore, the enzyme is overexpressed in a significant proportion (40%) of primary HCCs and in all 20 cholangiocarcinomas examined as well as in breast and colon carcinomas. The high levels of HAAH seen in syncytiotrophoblasts demonstrate that this increased expression is not restricted to carcinomas. In this regard, in preliminary studies we have also found increased HAAH expression in mouse fetal tissues (data not shown). The increased expression may be characteristic of growth in normal tissues.

Recent results from several independent lines of research highlight a need for determining the relevance of the increase of HAAH in transformed cells. A series of reports have documented that deletion mutants, lacking the EGF-repeat extracellular portions of Drosophila Notch (27–30) or of homologous invertebrate (31, 32) and vertebrate proteins (33–36), which are involved in cell fate determinations and cellular differentiation, are gain-of-function mutants. This suggests that in these transmembrane proteins, the extracellular domain regulates signal transduction by the cytoplasmic domain. Many of the EGF-like repeats in the extracellular domain (more than 20 of the 36 repeats in Drosophila and mammalian Notches) contain the putative consensus sequence (12) for asparyl(asparaginyl)$\beta$-hydroxylation. The low abundance of these proteins has to date precluded a direct demonstration of $\beta$-hydroxyaspartate or -asparagine in them. However, it seems likely that at least some, if not all, of the consensus-containing EGF repeats in Notch homologues are hydroxylated since the enzyme is present in all mammalian (mouse, rat, bovine, human) (6, 9) and invertebrate (Drosophila, Spodoptera) (37) species in which it has been sought and is widely distributed in adult and fetal tissues (19). It is tempting to hypothesize that the extent and/or pattern of $\beta$ hydroxylation modulates the regulation of signal transduction by the extracellular domain.

Two of the aforementioned gain-of-function mutations are particularly noteworthy. First, the mouse mammary tumor virus genome, which contains no known oncogene, induces mammary tumors by acting as an insertional mutagen (14), integrating at one or more specific (int) loci. Integration at the int-3 locus causes transcription of a novel truncated 2.3-kb cellular RNA species, the deduced amino acid sequence of which shows significant homology with the intracellular, ankyrin repeats of Notch (38). The untranslated portion of the gene located 5’ to the integration site encodes for the 36 EGF-repeat rich extracellular domain of Notch. Transgenic mice, containing a DNA fragment composed of the 3’-end of the mouse mammary tumor virus genome and the flanking cellular int-3 sequences, developed focal and, in some instances, poorly differentiated mammary and salivary adenocarcinomas (14). Second, in some human T lymphoblastic neoplasms, the TAN-1 gene, a human homolog of Drosophila Notch that maps to chromosome 9, is broken by a recurrent (7, 9) (q34, q34.3) chromosomal translocation (13). The translocation separates the TAN-1 gene into two nearly equal portions, one consisting of the extracellular domain and the second comprised of the transmembrane and cytoplasmic domains. When a cDNA, composed of this second portion of the gene, was inserted via a retroviral vector into mouse bone marrow cells, which were subsequently transplanted into lethally irradiated syngeneic mice, the mice developed T cell leukemias (14).

In addition to being an integral part of receptors involved in cell growth and differentiation, EGF domains containing the consensus sequence for hydroxylation may also act as important regulation elements in ligands. The vitamin K–dependent gamma glutamyl carboxylation may be important regulation elements in ligands. The vitamin K–dependent gamma glutamyl carboxylation may be
required for full stimulation of receptor autophosphorylation and, presumably, signal transduction (39, 40), a prototype receptor tyrosine kinase is the EGFR receptor; by analogy, the EGFR domains in the growth-arrest–specific gene 6 protein may participate in signal transduction and/or receptor endocytosis and recycling. Here, too, the extent of β hydroxylation could determine the binding affinity of the protein ligands, the extent of autophosphorylation and/or the efficiency of signal transduction by these receptor tyrosine kinases. Receptor endocytosis and recycling could also be affected, as it could for Notch and its homologues.

These experimental systems offer new opportunities to define the function subserved by the β hydroxylation of specific aspartic acid and asparagine residues in proteins. They further provide a starting point for studies designed to determine whether the substantially increased activity of HAAH in carcinomas is merely associative or contributes to the generation and/or maintenance of the malignant phenotype.

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References


