Posttranslational Modifications of Microfibril Associated Glycoprotein-1 (MAGP-1)†

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ABSTRACT: Microfibril-associated glycoprotein-1 (MAGP-1) is a small molecular weight protein associated with extracellular matrix microfibrils. Biochemical studies have shown that MAGP-1 undergoes several posttranslational modifications that may influence its associations with other microfibrillar components. To identify the sites in the molecule where posttranslational modifications occur, we expressed MAGP-1 constructs containing various point mutations as well as front and back half truncations in CHO cells. Characterization of transiently expressed protein showed that MAGP-1 undergoes O-linked glycosylation and tyrosine sulfation at sites in its amino-terminal half. This region of the protein also served as a major amine acceptor site for transglutaminase and mediated self-assembly into high molecular weight multimers through a glutamine-rich sequence. Fine mapping of the modification sites through mutational analysis demonstrated that Gln20 is a major amine acceptor site for the transglutaminase reaction and confirmed that a canonical tyrosine sulfation consensus sequence is the site of MAGP-1 sulfation. Our results also show that O-glycosylation occurs at more than one site in the molecule.

The formation of a functional elastic fiber requires the processing of tropoelastin monomers into a cross-linked polymer. Ultrastructural studies of elastic fiber assembly in developing tissues have suggested that 10–12 nm beaded filaments called microfibrils provide the alignment scaffold for elastin cross-linking at sites of fiber assembly (1). Much of what is known about microfibrillar composition has been ascertained through experiments using protein extracted from fetal or neonatal elastic tissues. Functional studies with these tissue extracts, however, have yielded little information because the isolation of monomeric microfibrillar proteins requires denaturation with chaotropic agents and the reduction of disulfide bonds. Furthermore, nonreducible cross-links acting to stabilize preexisting interactions form as the tissue ages, making extraction and isolation of microfibrillar proteins particularly difficult from adult tissues. These cross-links have been demonstrated to result, in part, from the action of tissue transglutaminase (2, 3).

It is clear from biochemical studies that fibrillins-1 and -2 play critical structural roles in microfibrillar architecture (4, 5). These large, modular glycoproteins are highly homologous, consisting of repeating calcium-binding epidermal growth factor-like domains interspersed between unique 8-cysteine domains. The fibrillins are thought to create the core structure of microfibrils through a head-to-tail alignment of monomer bundles (6). Ultrastructurally, microfibrils are characterized by a “beads on a string” appearance, with fibrillin molecules contributing to both the bead and interbead regions. The composition of the bead, however, is less clearly defined.

Microfibril-associated glycoprotein-1 (MAGP-1)1 and its structural relative MAGP-2 (7) are small molecular weight proteins that have been localized to fibrillin-containing microfibrils (8). MAGP-1 is associated with all microfibrils with only two exceptions: those directly adjacent to the plasma membrane of aortic endothelial cells and those at the junction of the zonule and the lens capsule of the eye (8–10). Although MAGP-2 is specifically associated with elastin-associated microfibrils and elastin-free microfibrils in a number of tissues, the protein exhibits patterns of tissue localization and developmental expression that are more restricted than those of MAGP-1 (11). The association of MAGP-1 and -2 with other microfibrillar proteins is apparently covalent since reducing agents are required for their extraction.

It is not yet known what role MAGP-1 plays in microfibrillar structure or function. Previous studies have demonstrated that it is capable of binding tropoelastin, an interaction that may facilitate the linking of elastin monomers onto microfibrils for their proper alignment. The amino-terminal half of MAGP-1 is enriched in acidic amino acids resulting in a cluster of negative charge that has been suggested to interact with a positively charged pocket near the C-terminus of the tropoelastin monomer. Anti-peptide

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† Abbreviations: DMEM, Dulbecco’s modified Eagle’s medium; GMEM, Glasgow’s modified Eagle’s medium; FBC, fetal bovine chondroblasts; IPTG, isopropylthio-β-galactoside; MAGP, microfibril associated glycoprotein; MSX, methionine sulfoxamine; SDS, sodium dodecyl sulfate.
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antibodies directed against a domain near the N-terminus of MAGP-1 were able to block the incorporation of tropoelastin into fibers by cultured chondrocytes (12). More recently, Finnis and Gibson (13) have localized tropoelastin binding, as well as binding to type VI collagen, to a tyrosine-rich sequence near the amino-terminus of MAGP-1. This region is immediately upstream of that suggested by the antibody studies of Brown-Augsburger et al. (12).

MAGP-1 has been shown to undergo several posttranslational modifications, including glycosylation, sulfation, and transglutamination (2, 14). To better characterize these modifications with the aim of assessing their contribution to elastic matrix assembly and function, full-length MAGP-1 and two truncated forms of the protein were stably expressed in CHO cells. We show that MAGP-1 undergoes transglutamination, O-linked glycosylation, and tyrosine sulfation at sites in the amino-terminal half of the protein. Precise mapping of these sites using mutational analysis identified Gln20 as the major amine acceptor site for transglutaminase and confirmed that sulfation occurs within a tyrosine sulfation consensus sequence. We also show that at least two sites for glycosylation exist in the amino terminal half of the protein and that a glutamine-rich sequence mediates higher order aggregation.

MATERIALS AND METHODS

All reagents were obtained from Sigma Chemical Company unless otherwise specified. All restriction enzymes were purchased from Boehringer-Mannheim. Fetal bovine chondrocytes (FBCs) were isolated and maintained as previously described (15). All cloning followed protocols described by Sambrook et al. (16).

Mammalian Expression. Full-length bovine MAGP-1 cDNA (17) was ligated into the pEE 14 mammalian expression vector (18) at the EcoR I site. DH-10b bacteria were transformed with ligated DNA, and subclones containing MAGP-1 cDNA were identified by restriction digestion and confirmed by DNA sequencing. In this vector, expression of the inserted protein is driven by the strong CMV promoter/enhancer. A glutamine synthase mini-gene, whose expression is driven by the SV-40 promoter, is employed in selection of stable transfectants with MSX.

The MAGP-1-containing mammalian expression plasmid was transfected into CHO cells using Lipofecttin (Gibco/BRL) according to the manufacturer’s instructions. After 2 days, transfected cells were split 1:3 into GMEM without glutamine (Gibco/BRL) to which 10% dialyzed fetal calf serum (FCS) and 25 mM MSX had been added. After 15 days, colonies were selected and screened for MAGP-1 production by Western blotting of conditioned medium. Positive colonies were cloned by serial dilution, and subclones were screened for MAGP-1 production by Western blotting.

Truncated MAGP-1 constructs encoding the front and back halves of the protein were generated by PCR cloning using full-length cDNA in pBlueScript (pBS, Promega) as a template and oligonucleotide primers containing engineered restriction sites. Amplification with Klenex DNA polymerase (obtained from Dr. Wayne Barnes, Washington University) was done using a hot start followed by 30 cycles of 2 min at 97 °C; 2 min at 55 °C; and 3 min at 72 °C. The amino-terminal half of MAGP-1 was constructed using a 5' primer complementary to the T3 sequence of the cloning vector (5'-ATTAACCTCCTACTAAAG-3'). The 3' primer was reverse complementary to bases 283–306 of the bovine sequence (5'-GGTCCCGGCTCCTCCGGCTCCTGC-3') with bases 293_G and 294_G to generate a Smal restriction site near the 3' end. The 395-bp PCR product was treated with the Klenow fragment of DNA polymerase (Gibco/BRL), gel purified, and digested with Xba I and Smal I. This fragment was ligated into the pEE 14 vector cut with Xba I and Smal I. Ligation into these sites resulted in the construct MAGP-1 (M1-P89) with the addition of six amino acid residues at the carboxyl-terminal end (-Gly-Arg-Ala-Arg-Ile-His) before the vector-derived stop codon was reached.

To generate the carboxy-terminal half of MAGP-1, bases 287–936 (includes some 3' UTR) were PCR amplified using Klenataq DNA polymerase. Full-length bovine cDNA in pBS was used as a template. A 5' primer complementary to bases 286–304 of the bovine sequence with 298ΔT generated a BamH I site near the 5' end (5'-GAGCCCCACGGATCCGCGGA-3'). The 3' primer was used to generate the T7 sequence in pBS. The 725 bp PCR product was treated as above before digestion with BamH I and Hind III and ligation into pBS digested with the same enzymes. The MAGP-1 leader sequence was PCR amplified with Taq DNA polymerase from the full-length cDNA template. The same T3 5' primer described above and a 3' primer reverse complementary to bases 80–103 of the bovine sequence (5'-CAGGGGGTCCTGATCATACTGGCC-3') with base 93ΔA were used to generate a Bcl I restriction site at the 3' end. This 190-bp PCR product was treated as above before digestion with Xba I and Bcl I and was ligated into pBS containing the amino-terminal truncated construct cut with Xba I and BamH I. These steps generated a product consisting of the MAGP-1 leader sequence plus the first four amino acids of the mature protein fused to amino acids 189–183. This 915-bp construct was digested with EcoRI and was ligated into pBS digested with the same enzymes. Both truncated proteins were transfected into CHO cells and stable transfectants were selected as described above for the full-length protein.

Point mutations for mammalian expression were made using the QuikChange, (Stratagene) site-directed mutagenesis kit per the manufacturer’s specifications with full-length MAGP-1 in pBS as the template. Briefly, this protocol uses Pfu DNA polymerase along with complimentary forward and reverse primers containing the desired mutation(s) to amplify around a circular template. Following amplification, the methylated, wild-type template (plasmid DNA from E. coli) is degraded with Dpn I while the unmethylated PCR product containing the incorporated mutations remains intact. This PCR-derived plasmid containing the mutagenized sequence is then introduced into E. coli and manipulated as desired. Three threonine residues (T54, T79, and T90) were mutated to valine using the following primers: bases 155–156 with 185ΔG and 186ΔT (5'-CCCCAGAGATGGTCCCCCTCGGGCCG-3') for T54ΔVal; bases 251–271 with 260ΔG and 261ΔT (5'-CCTGCCCCCTCGTTAGAACC-3') for T79ΔVal; and bases 285–306 with 293ΔG, 294ΔT, and 295ΔC (5'-CGGAGCCCCGTCGAGCGGGGACC-3') for T90ΔVal. To generate mutation in the tyrosine sulfation consensus sequence, primers complimentary to bases 151–188 of the
bovine sequence with 165ΔC, 168ΔC, and 174ΔC (5’-CGAG-AATCCGA)CTCCTCAGACTCCCCAGATGACC-3’) were used to PCR amplify the MAGP-1 mutant sequence yielding a construct with Y47, Y48, and Y50 all changed to serine residues. In all cases, products were sequenced to confirm incorporation of the desired mutation and to ensure that no additional mutations were introduced. The mutant MAGP-1 sequences were cut out of pBS with Xba I and Hinc II and ligated into the pEE 14 mammalian expression vector cut with Xba I and Sma I. All constructs were transiently transfected into CHO cells along with wild-type MAGP-1 as described for stable transfections above.

**Antibodies.** Bovine MAGP-1 cDNA was digested with Pst I and Apa I, yielding a 270-nt fragment that was filled in with the large fragment of DNA Polymerase and re-ligated into Pst I and EcoR V-cut pBS. This construct was digested with BamH I and Xho I, and the insert was ligated into the pGEX vector (Pharmacia), resulting in a fusion protein of glutathione S-transferase (GST) coupled with amino acids G14–P103 of MAGP-1. E. coli containing this plasmid were grown in an environmental shaker at 37 °C to an O.D. of 0.4 before induction of fusion-protein synthesis with 0.2 mM IPTG (Gibco/BRL) for 4 h. Bacteria were pelleted and subsequently lysed by repeated freeze–thaw cycles followed by addition of 0.1% Triton X-100 and sonication. Bacterial lysate was cleared of debris by centrifugation. The supernatant was then absorbed to a glutathione-Sepharose 4B column (Pharmacia) and bound fusion protein was eluted with PBS containing 10 mM glutathione. The carboxy-terminal construct of MAGP-1 described above for mammalian expression (lacking the leader sequence) was ligated into the bacterial expression vector pQE (Qiangen) at the BamH I and Hind III sites. This expression vector contains an N-terminal 6XHis tag that was used for purification of the recombinant protein. Subclones containing the appropriate recombinant plasmid were identified by restriction digestion. DNA was transformed into M15 bacteria and clones were screened for protein production. One positive clone was selected and was grown for 4 h at 37° C, induced with 1 mM IPTG for 4 h at 37° C, pelleted, and lysed with 6 M urea, pH 8, overnight at 4 °C. Bacterial lysate was cleared by centrifugation, and the supernatant was mixed with nickel agarose (Qiangen). After extensive washing at pH 8 and pH 6, bound protein was eluted with 6 M urea, pH 5.

Antibodies were generated by injection of purified recombinant protein into rabbits. Serum was collected and antibodies were purified over an affinity column containing elastic fiber proteins extracted from bovine nuchal ligament coupled to Sepharose (14). Anti-peptide antibodies generated to peptide 244 (amino acids G19–D31), peptide 243 (amino acids Y38–E52), and peptide 239 (amino acids V74–T87) of bovine MAGP-1 were previously described by Brown-Augsburger et al. (12).

**Western Blotting.** Serum-free GMEM was conditioned overnight by MAGP-1-expressing CHO cells. The conditioned medium was then concentrated and subjected to reducing SDS–PAGE with transfer of protein to nitrocellulose. Filters were blocked in PBS/3% nonfat dry milk for 1 h at room temperature and incubated with primary antibodies, each at a 1:1000 dilution, overnight at 4 °C in PBS/3% dry milk. The filters were then washed three times with PBS/milk and incubated with an HRP-conjugated secondary antibody (Amersham) for 1 h at room temperature in PBS/milk. Filters were washed three times and immunoreactive proteins were visualized by enhanced chemiluminescence.

**Immunoprecipitation.** Confluent cells were incubated for 5 h with 50 μCi/ml [35S]-cysteine (sp. act. >800Ci/mmol; ICN) in cysteine-free DMEM, 42 μCi/ml [3H]-proline (sp. act. >225mCi/mmol; ICN) in DMEM, or 150 μCi/ml [35S]O4 (sp. act. ~43Ci/mg S; ICN) in sulfate-free S-MEM (Gibco/BRL). To assess sugar incorporation, confluent MAGP-1-expressing CHO cells were labeled overnight in 5 mL/p100 tissue culture dish with 50 μCi/ml [3H]-glucosamine (sp. act. 25–40Ci/mmol; ICN) in low-glucose DMEM (Gibco/BRL) containing 2% diazoylc FCS. CHO cells transiently transfected with wild type and point mutation-containing constructs were radiolabeled as above 4 days after transfection. For all immunoprecipitations, radiolabeled medium was pre-cleared with Protein-A immobilized on Trysacryl (Pierce) for 1 h at room temperature. Beads were pelleted, and the supernatant was transferred to a new tube containing antibody and Protein-A/Trysacryl and incubated at 4 °C overnight. Beads were pelleted and washed 4× with detergent buffer containing 0.1% SDS, 0.25% deoxycholate, 2 mM EDTA, 0.5% Triton X-100, 150 mM NaCl, and 50 mM Tris, pH 8. Washed beads were then resuspended in reducing sample buffer before electrophoresis.

**In Vitro Transglutaminase Assay.** To generate a recombinant protein containing the putative N-terminal transglutaminase site, the MAGP-1 (G14–P103)/pGEX construct described above for generation of an MAGP-1 antibody was digested with Eag I. This digestion liberated a fragment consisting of the GST moiety fused to amino acids G14–P27 of MAGP-1 that was then re-ligated into the pGEX vector. Recombinant protein generated from this construct was purified in the same manner as the larger fusion protein described above. Both of these bacterial fusion proteins were incubated in a 50 mM Tris, pH 8.3, buffer containing 100 μM CaCl2 with 1.2 units of guinea pig liver transglutaminase and 0.65 μCi [14C]-spermidine (Amersham) in a final reaction volume of 45 μL. Reactions were allowed to proceed at 37 °C for 1.5 h. Reactions were stopped by the addition of 2 μL of 0.5 M EDTA. Incorporation of radiolabeled spermidine was detected by electrophoresis of the reaction products on 10% SDS–PAGE gels followed by autoradiography.

Point mutations of the glutamine residues within the shorter construct were made using the QuikChange system as described above for mutations in proteins for mammalian expression. The MAGP-1 (G14–P57) construct in pGEX served as a template in generating these mutant fusion proteins. Three of the four glutamine residues contained within this construct, Q20, Q34, and Q41, were mutated independently to Asn by PCR using the following primers: bases 73–98 with 82ΔA and 84ΔT (5′-GGCTCAAGGCAAT- TATGATCTGGACC-3′) for Q20ΔA; bases 112–137 with 125ΔA and 127ΔC (5′-CCGACACGCTGAATACTAC- CCACT-3′) for Q34ΔN; and bases 139–161 with 146ΔA and 148ΔT (5′-CAGCGAATATCGGAATCCGG-3′) for Q41ΔN. These recombinant mutant proteins were isolated by glutathione affinity chromatography, and transglutaminase assays were performed as described above for the wild-type proteins.
Peptide Aggregation and Congo Red Assay. Peptide 127 encoding amino acids 57–80 of MAGP-1 (PPEEQFQFQS-QQQVQEVIPAPTL) was synthesized using FAST-Moc chemistry and dissolved at 10 mg/mL in 7 M guanidine HCl. A total of 10 μL of the peptide stock solution was diluted with 200 μL of PBS and 200 μL of water and subjected to slow rotation overnight at room temperature. Precipitated peptide was pelleted by centrifugation in a microfuge, smeared on a glass slide, fixed in 95% ethanol, and stained with 0.05% congo red in 50% glycerol. Evaluation of congo red staining by polarization microscopy was performed using a Zeiss Axioskop equipped with optimally aligned cross-polarizers.

Digestion of Carbohydrate Chains and Lectin Binding. For removal of carbohydrates, Trisacryl beads with immunoprecipitated protein were resuspended in 25 μL of 10 mM calcium acetate/20 mM sodium cacodylate, pH 7.4. Neuraminidase (25 mU) was then added, and samples were incubated for 1 h at 37 °C followed by an overnight incubation with 4 mU O-glycanase (Genzyme). Reactions were stopped by the addition of SDS–PAGE reducing sample buffer. Following electrophoresis, gels were fixed, treated with En3Hance (Amersham), and dried before visualization of immunoprecipitated proteins by autoradiography.

Untransfected CHO cells or CHO cells expressing recombinant protein were incubated overnight in serum-free medium. The conditioned medium was then incubated for 1 h at 37 °C with 1 unit/mL neuraminidase to remove terminal sialic acid residues from glycoproteins. Treated samples were rotated end-over-end at room temperature with uncoupled agarose beads for 1 h to decrease nonspecific protein association with agarose. The beads were pelleted and the supernatants were transferred to new tubes containing washed control agarose or Jacalin-agarose beads (Vector Labs). Samples were allowed to rotate end-over-end at 4 °C overnight. Beads were pelleted by centrifugation and washed four times with serum-free GMEM. Washed beads were resuspended in reducing sample buffer and analyzed by SDS–PAGE followed by Western blotting.

Tropoelastin Binding Assay. MaxiSorp (Nunc) plates were coated with 1 μg/well of recombinant human tropoelastin (kind gift of Dr. Joel Rosenbloom, University of Pennsylvania) in 10 mM bicarbonate buffer, pH 10, in a humidified chamber at 4 °C overnight. Following coating, plates were washed 3× with PBS and blocked with 1% BSA in PBS for 1 h at 37 °C and again washed 3× with PBS. Medium conditioned overnight by MAGP-1-expressing CHO cells was used as a source for MAGP-1 protein. The same cells treated with 10 mM KClO3 were used as a source for MAGP-1 protein. The same cells were centrifuged in a microfuge, smeared on a glass slide, fixed in 95% ethanol, and stained with 0.05% congo red in 50% glycerol. Evaluation of congo red staining by polarization microscopy was performed using a Zeiss Axioskop equipped with optimally aligned cross-polarizers.

RESULTS

Expression of Full-Length and Truncated MAGP-1 Proteins in CHO Cells. To characterize sites of posttranslational modifications in MAGP-1, we expressed full-length, back and front half constructs (Figure 1) of the bovine protein in CHO K1 cells. These cells were chosen because of their ease of transfection and the absence of detectable endogenous MAGP-1 expression. The front and back half constructs were designed to encode the two discreet domains of this protein that are thought to exhibit distinct functions within the elastic matrix (2, 17).

Figure 2 shows that each of the recombinant proteins is expressed by transfected cells and that the antibodies used in this study are specific for their respective antigens and do not react with CHO proteins expressed by untransfected cells. Expression of the amino-terminal half of MAGP-1 results in the secretion of multiple aggregate forms of the protein (Figure 2, panel A). Because this construct contains none of the protein’s cysteine residues, the aggregation is not due to disulfide bonding. It does, however, appear to be an irreversible interaction as we have been unable to either inhibit or reverse aggregate formation by the addition of 8 M urea (data not shown). Like the front half construct, the back half fragment was secreted into the medium at high levels but showed no aggregation (Figure 2, panel B).

Sequences responsible for aggregation of the front half of the molecule were determined by testing the solubility of synthetic peptides that encode most of the front half construct. Of the five peptides tested (G19–D31, Y38–E52, H52–E71, D75–E93, and E93–E118), the latter three peptides demonstrated the aggregation properties of the full-length MAGP-1. 

**FIGURE 1:** Sequence of full-length bovine MAGP-1 and schematic representation of the constructs used in this study. (Top) full-length sequence of bovine MAGP-1 indicating peptide sequences for synthetic peptides (underlined) and sites of posttranslational modification determined in this study. Peptide P57–L80 (dashed underline) contains the glutamine-enriched region of the protein. Threonine residues predicted as sites of glycosylation are indicated by (●). Sites of transglutamination are shaded. Tyrosine residues in the sulfation consensus site are marked by an asterisk. The leader peptide was pelleted by centrifugation in a microfuge, slow rotation overnight at room temperature. Precipitated proteins by autoradiography. Antibodies were diluted in PBS containing 0.1% BSA and incubated with the immobilized tropoelastin for 1 h at 37 °C. The color reaction was quantified using a Bio-Rad microtiter plate reader at 405 nm.

**FIGURE 2:** Sequence of full-length bovine MAGP-1 and schematic representation of the constructs used in this study. (Top) full-length sequence of bovine MAGP-1 indicating peptide sequences for synthetic peptides (underlined) and sites of posttranslational modification determined in this study. Peptide P57–L80 (dashed underline) contains the glutamine-enriched region of the protein. Threonine residues predicted as sites of glycosylation are indicated by (●). Sites of transglutamination are shaded. Tyrosine residues in the sulfation consensus site are marked by an asterisk. The leader sequence is in italics. (Bottom) Schematic representation of the MAGP-1 constructs stably expressed in CHO cells.
Additional evidence that MAGP-1 contains O-linked carbohydrate(s). Uncertainty as to whether MAGP-1 is a glycoprotein was raised when no N-linked glycosylation sites were found in the protein’s primary sequence (17). To determine whether the protein contains O-linked sugars, we incubated CHO cells expressing full-length MAGP-1 with [3H]-glucosamine. While glucosamine is not incorporated directly into O-linked carbohydrates, most cells will epimerize this carbohydrate to galactosamine, which is incorporated. Figure 4 shows incorporation of the radiolabeled sugar into full-length MAGP-1 immunoprecipitated from culture medium (lane 1). Removal of the glucosamine label with O-glycanase (lane 2) confirms its incorporation into O-linked carbohydrate chains.

Identification of Transglutamination Site in MAGP-1.

Previous studies have suggested that an amine acceptor site for the transglutaminase reaction is located near the amino terminus of MAGP-1 (2). To better localize the site on MAGP-1 that is modified by this enzyme, two overlapping GST bacterial fusion proteins were made. The longer of the two constructs (amino acids G14–P103) contains 13 of the 16 glutamine residues found in the full-length protein. The smaller fusion protein (amino acids G14–P57), which contained the putative acceptor site, was generated by taking advantage of a convenient restriction site in the longer construct. This polypeptide contains the first four glutamine residues in the amino-terminal region of MAGP-1. Figure 3, panel A shows that both the shorter (G14–P57) and longer (G14–P103) proteins incorporate [14C]-spermidine when incubated with transglutaminase. It was necessary to include the GST moiety on peptides containing the glutamine-rich sequence to prevent aggregation. GST alone was not a substrate for transglutaminase (not shown).

To identify the specific residues acting as amine acceptors, three of the four glutamines in the shorter construct were individually mutated to asparagine residues. The glutamine at the amino-terminus of secreted mammalian MAGP-1 (Q18) forms pyroglutamic acid (2) and therefore cannot act as an acceptor residue. Figure 3, panel B shows that two of the mutant recombinant proteins, Q34N and Q41N, continue to serve as substrates for transglutaminase, although the level of labeling of the Q34N mutant is minimal as compared to the wild type or Q41N proteins. Mutation of Q20 completely blocked incorporation of [14C] label, indicating that this residue is the major site for transglutamination in this region of the protein.

MAGP-1 Contains O-Linked Carbohydrate(s). Two constructs encoding amino acids G14–P103 and G14–P57 were expressed as GST fusion proteins and incubated with [14C]-spermidine in the presence of transglutaminase. CB, Coomassie blue stain; 14C, incorporation of [14C]-spermidine. (B) Recombinantly expressed proteins with individual point mutations. WT, unmutated protein; Q20, Glu20 to Asn mutation; Q34, Glu34 to Asn mutation; Q41, Glu41 to Asn mutation. CB, Coomassie blue stain; 14C, incorporation of [14C]-spermidine.
conditioned by CHO cells expressing full-length and truncated MAGP-1 proteins was treated with neuraminidase to remove terminal sialic acid residues and then incubated with either control agarose or Jacalin-coupled agarose. Protein that bound to the lectin was subjected to electrophoresis and MAGP-1 was visualized by Western blotting. Figure 4 shows that the full-length (lane 3) and the amino-terminal half of the protein (lane 5) associate specifically with Jacalin agarose while the carboxy-terminal half (lane 8) does not. This demonstrates that carbohydrate modification of MAGP-1 occurs between residues 1 and 89 of the mature protein.

Submission of the bovine MAGP-1 sequence to the NetOglyc Prediction Server (www.cbs.dtu.dk/service/netOGlyc/) resulted in the identification of three possible threonine glycosylation sites within full-length bovine MAGP-1: T54, T79, and T90, with the first two of these having the highest glycosylation potential. The same three residues are predicted for glycosylation in human MAGP-1. To determine whether these residues are active sites of glycosylation, all three residues were independently mutated in the full-length protein. Figure 5 shows that all three mutant proteins are secreted when transiently expressed in CHO cells. Also shown in Figure 5 is binding of all three mutant proteins to Jacalin agarose, indicating that glycosylation is occurring at multiple sites in the protein and that mutation of only one of these amino acids is not sufficient to prevent glycosylation.

MAGP-1 Is Sulfated on Tyrosine Residues. Bovine MAGP-1 has been shown to be sulfated when expressed in SF9 cells (22). To localize the site of sulfation, CHO cells stably expressing full-length MAGP-1 were incubated in medium containing [35S]-sulfate. Figure 6 (lane 1) shows that MAGP-1 immunoprecipitated from the medium of these cells incorporates the sulfate label as expected. Treatment of the immunoprecipitated protein with O-glycanase to release carbohydrates (lane 2) did not remove the sulfate radiolabel, demonstrating that the sulfate modification is not on O-linked sugars.

A possible site of modification within the MAGP-1 molecule was suggested by the presence or absence of sulfate labeling in two known degradation products of the protein. The lower molecular weight products (bands B and C indicated by arrows in Figure 7) occur in variable amounts and are routinely seen when MAGP-1 is isolated from tissues.
or cells. Western blotting of MAGP-1 with antibodies raised against three different synthetic peptides (Figure 7, left) demonstrates that bands B and C arise as a consequence of amino-terminal deletions. These peptide antibodies also define the missing regions with some accuracy. For example, all three bands react with the antibody raised against amino acids 74–86 recognize all three bands (lane 3), whereas only the two higher molecular weight bands are recognized by the antibody to amino acids 38–52 (lane 2). Since the [35S]-sulfate label is missing from the lower band but is present in the two higher bands (center panel), the sulfate-containing residues must lie between the epitopes recognized by the 38–52 and 74–86 antibodies. The right panel shows an immunoprecipitation from [35S]-sulfate labeled cells demonstrating that the full-length (FL) and the front-half (FH) constructs, both of which contain the putative sulfate site, are able to incorporate sulfate label.

Inhibition of Sulfation Does Not Alter MAGP-1’s Ability to Bind Tropoelastin. Our previous studies have shown that MAGP-1 binds tropoelastin in solid-phase binding assays. It has been suggested that these two proteins associate via an ionic interaction between the negatively charged amino-terminal region of MAGP-1 and the positively charged C-terminus of tropoelastin (12). Sulfation of the amino-terminal region of MAGP-1 would greatly enhance the negative charge of this region, raising the possibility that its interaction with tropoelastin might depend on this modification.

To test this possibility, medium conditioned by untreated and chlorate-treated cells expressing full-length MAGP-1 was collected and used in a solid-phase tropoelastin binding assay. Chlorate is a potent and specific inhibitor of protein and proteoglycan sulfation (24) but does not generally inhibit protein synthesis. Indeed, neither MAGP-1 synthesis nor its ability to interact with antibody was inhibited by the addition of 10 mM chlorate (Figure 9, panel A, lanes 1 and 4). The relative absence of [35S]-sulfate label in MAGP-1 immunoprecipitated from chlorate-treated cells (lane 3) confirms the effectiveness of the treatment. Figure 9, panel B, shows that the absence of tyrosine sulfation does not alter the ability of MAGP-1 to bind tropoelastin, suggesting that sulfation plays little role in the tropoelastin-MAGP-1 interaction.

DISCUSSION

Several studies have shown that MAGP-1 undergoes posttranslational modifications that may alter its biological properties and influence its interactions with other proteins. The present study focuses on a detailed analysis of the sites in the molecule where these modifications occur.

In earlier studies using MAGP-1 derived from a baculoviral expression system (2), the full length, native protein demonstrated a propensity to aggregate into high molecular weight multimers. When the front and back halves of the molecule were separately expressed in CHO cells, it was found that aggregation occurred only with the front half of the protein. Analysis of the front half of the molecule using synthetic peptides identified the glutamine-rich motif located between amino acids 61–72 as the sequence responsible for
aggregation. The ability of the glutamine-containing peptide to form insoluble fibrils that demonstrated green birefringence after Congo red staining indicates that the peptide aggregates have β-pleated sheet structural features in common with amyloid-like fibrils (25, 26). Indeed, Aggeli et al. (27) have shown that synthetic peptides with glutamine-enriched sequences similar to the sequence found in MAGP-1 produce β-sheet structures that self-associate to form insoluble gels. The significance of MAGP-1 self-association through the glutamine-rich region is not known, but it may have some physiological role in the organization of this protein into homo- or heteropolymers within the extracellular matrix. It may also suggest a mechanism whereby MAGP-1 could interact with other molecules capable of forming β-sheet structures.

Throughout this and other studies (2), we have noticed that aggregation of full-length MAGP-1 occurs much more slowly and is much less complete than what is observed for the front half construct. One explanation for this difference in aggregation is that the Gln-rich region in the full-length protein may not be as accessible for protein–protein interactions as it is in the front half construct alone. This raises the interesting possibility that conformational changes induced by proteolysis or by binding of MAGP-1 to other proteins may convert the sequence into an active conformation.

MAGP-1 was initially described as a glycoprotein based on its reactivity with PAS reagent and the detection by sugar analysis of associated N-acetylgalactosamine, galactose, mannose, and glucose carbohydrates (14). When the full-length cDNA for MAGP-1 was characterized and no N-linked glycosylation consensus sequences were found, it was suggested that carbohydrates must be O-linked (17). Our finding that MAGP-1 secreted by CHO cells incorporates radiolabeled carbohydrate and interacts specifically with lectins confirms that the protein undergoes O-linked glycosylation. It is important to note that the extent of MAGP-1 glycosylation is not extensive since the apparent molecular weight of the fully glycosylated protein (either the full-length or front half construct) does not differ significantly from the nonglycosylated form. This suggests that the carbohydrates linked to MAGP-1 are the small tri- and tetrasaccharide varieties and not the larger mucin-like chains (28).

The precise sites of glycosylation within the molecule remain elusive. Because the specificity of the transferase that links the carbohydrate GalNAc to the serine or threonine side chain is presently unknown, the prediction of O-linked glycosylation sequences is still inaccurate. Our studies have localized carbohydrate incorporation to the front half of the protein. However, the capacity of single glycosylation site mutants to bind lectins indicates that MAGP-1 contains more than one site of glycosylation. The biological function of MAGP-1 carbohydrate modification is still unknown, although glycosylation of other proteins has been shown to regulate protein–protein and protein–cell interactions (29).

The presence of sulfate moieties within the MAGP-1 molecule were first demonstrated by Tomasini-Johansson et al. (30) who noted that a 30-kDa sulfated protein, identified as MAGP-1 by antibody reactivity, interacted with a monoclonal antibody raised against human vitronectin. The antibody did not react with protein isolated from cells treated with chlorate to inhibit sulfation, suggesting that the antibody recognized a sulfate moiety on both molecules. In addition, sulfation of MAGP-1 was suggested to be on tyrosine residue(s) since digestion with chondroitinase did not affect the level of sulfation, confirming that MAGP-1 is not a proteoglycan. Our studies confirm that sulfation of MAGP-1 is occurring within a consensus sequence for tyrosine sulfation (Y 47, 48, and 50) that is homologous to that present in vitronectin (30).

The role of MAGP-1 sulfation in elastic matrix assembly or function is unknown, but the importance of the modification was suggested by our studies showing that inhibition of sulfation prevents microfibrillar protein incorporation into the ECM of cultured cells (31). Our direct binding studies show that sulfation of MAGP-1 is not required for its
interaction with tropoelastin. We cannot exclude the possibility, however, that sulfation might alter the affinity of the interaction.

In addition to containing sites for sulfation and glycosylation, the amino-terminal region of MAGP-1 contains an amine acceptor site for tissue transglutaminase. Our previous studies with full-length MAGP-1 suggested that a transglutaminase site was located near the amino-terminus (2), but the precise residues involved in the reaction were not determined. Results in Figure 3 show that a small amino-terminal fragment of MAGP-1 containing the first four glutamine residues (Q18, Q20, Q34, and Q41) serves as an amine acceptor. On the basis of the preference of transglutaminase for modifying the second glutamine residue in a Q-X-Q pair, Q20 was the most likely target residue. Indeed, the mutation of Q20 prevents modification by this cross-linking enzyme, verifying that this residue can serve as a substrate for transglutaminase. Mutation of Q34 resulted in a significant decrease in [14C]-spermidine incorporation, suggesting that this site could influence the transglutaminase reaction at residue Q20, or that it might be a second transglutaminase site. We cannot exclude the possibility, however, that sulfation might alter the affinity of interaction with tropoelastin. We cannot exclude the possibility, however, that sulfation might alter the affinity of the interaction.

As with most proteins of the extracellular matrix, post-translational modifications add diversity and specificity to biological function. Characterizing the modifications that occur on MAGP-1 has been particularly difficult because of the protein’s unusual physical properties. The expression of the recombinant MAGP-1 proteins described in this report has allowed us to more finely map many of the modifications that occur. These results are important in understanding the macromolecular interactions, and therefore the function, of MAGP-1.

REFERENCES