Identification of an Endogenously Generated Cryptic Collagen Epitope (XL313) That May Selectively Regulate Angiogenesis by an Integrin Yes-associated Protein (YAP) Mechano-transduction Pathway*

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Extracellular matrix (ECM) remodeling regulates angiogenesis. However, the precise mechanisms by which structural changes in ECM proteins contribute to angiogenesis are not fully understood. Integrins are molecules with the ability to detect compositional and structural changes within the ECM and integrate this information into a network of signaling circuits that coordinate context-dependent cell behavior. The role of integrin αvβ3 in angiogenesis is complex, as evidence exists for both positive and negative functions. The precise downstream signaling events initiated by αvβ3 may depend on the molecular characteristics of its ligands. Here, we identified an RGD-containing cryptic collagen epitope that is generated in vivo. Surprisingly, rather than inhibiting αvβ3 signaling, this collagen epitope promoted αvβ3 activation and stimulated angiogenesis and inflammation. An antibody directed to this RGDKGE epitope but not other RGD collagen epitopes inhibited angiogenesis and inflammation in vivo. The selective ability of this RGD epitope to promote angiogenesis and inflammation depends in part on its flanking KGE motif. Interestingly, a subset of macrophages may represent a physiologically relevant source of this collagen epitope. Here, we define an endothelial cell mechano-signaling pathway in which a cryptic collagen epitope activates αvβ3 leading to an Src and p38 MAPK-dependent cascade that leads to nuclear accumulation of Yes-associated protein (YAP) and stimulation of endothelial cell growth. Collectively, our findings not only provide evidence for a novel mechano-signaling pathway, but also define a possible therapeutic strategy to control αvβ3 signaling by targeting a proangiogenic and inflammatory ligand of αvβ3 rather than the receptor itself.

Angiogenesis, the process by which new blood vessels form from pre-existing vessels, plays a critical role in normal and pathological events. Efforts are underway to more precisely define the interconnected mechanisms that control this crucial biological process to develop more effective strategies to control neovascular diseases. Significant advances have been made in identifying molecular regulators of angiogenesis and their associated signaling pathways. A more precise understanding of angiogenic signaling pathways and the networks of regulatory feedback loops operating within distinct cellular compartments has provided important clues to help explain the modest clinical impact of many current anti-angiogenic strategies. For example, although vascular endothelial growth factor (VEGF) induces pro-angiogenic signaling leading to enhanced endothelial cell migration, growth, and survival, VEGF-induced blood vessels are often characterized as immature, unstable, and leaky and can regress in the absence of additional signaling events. VEGF stimulation under specific circumstances may lead to inhibition of angiogenesis in the context of altered PDGF signaling due to disruption of pericyte recruitment. These unexpected findings provide evidence of a negative role for the VEGF/VEGFR signaling during new vessel development. Similarly, studies have provided evidence for both a positive and a negative role for integrin αvβ3 in angiogenesis. For example, multiple pro-angiogenic roles have been proposed for αvβ3 as cyclic arginine-glycine-aspartic acid (RGD)-containing peptides and antibodies targeting this integrin inhibit angiogenesis in animal models. In contrast, enhanced angiogenesis was detected in tumors growing in αvβ3 null mice. Interestingly, reduced pathological angiogenesis was detected in transgenic mice expressing signaling-deficient β3-integrin, which resulted in part from defective recruitment of bone marrow-derived cells rather than specific endothelial cell defects. Moreover, new evidence suggests that β3-integrin may play a more prominent role in the early stages of angiogenesis when new vessels begin to form, as reduction in endothelial cell expression of β3-integrin impaired early stage pathological angiogenesis, but it had little effect on later maturation stages once vessels had formed. These studies, together with many others, suggest αvβ3-mediated regulation of angiogenesis is complex, temporally regulated, and is not solely dependent on adhesive events but also involves downstream signaling, the consequences of which may depend on the cell type and composition of the local extracellular microenvironment. Given the opposing biological
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responses observed following modulation of some angiogenic regulatory molecules, it is not surprising that anti-angiogenic strategies based on targeting these factors have met with limited clinical success.

Given the importance of integrin-extracellular matrix (ECM) interactions in modulating the intensity and specificity of growth factor signaling (1–5, 17, 18), it is important to define how diverse components within the local vascular microenvironment function cooperatively to regulate angiogenesis. Interestingly, distinct αβ3 ligands may stimulate opposing biological outcomes (7–15). For example, certain NC1 domains of collagen may bind αβ3 and induce pro-apoptotic responses, whereas binding of other αβ3 ligands may promote cell growth and survival (19–22). Given these findings and the complex biological effects observed following direct targeting of αβ3, an alternative therapeutic approach to control signaling from αβ3 might involve specific targeting of the pro-angiogenic ligands of αβ3 rather than directly targeting the receptor itself. Proteolytic remodeling of the ECM can generate integrin binding cryptic epitopes that play functional roles in angiogenesis, including the cryptic epitope present in multiple types of collagen that we have termed HUI177, containing the amino acids, leucine, proline, and glycine (LPGXPG), where X can represent any amino acid and a second cryptic collagen epitope termed HUIV26, which is present in collagen type IV (21–23). Although the HUIV26 epitope is recognized by αβ3, it is not specifically composed of an RGD motif (21). Given the importance of RGD sequences in mediating some integrin-dependent interactions and the roles of amino acids flanking the core RGD motif in establishing integrin-binding specificity and affinity (24–27), we sought to determine whether RGD motifs within collagen differentially regulate angiogenesis. Sequence analysis of collagen type I revealed that five different cryptic RGD motifs are present, each with unique flanking sequences. Surprisingly, the C-terminal KGE flanking sequence of one of these RGD motifs is highly conserved in species as diverse as Xenopus and man. In contrast, significant sequence and positional variation exists within the other flanking sequences among different species.

Here, we provide evidence that an RGDKGGE containing cryptic collagen epitope can be generated by a substrate of macrophages, and this motif but not other RGD peptides may promote rather than inhibit angiogenesis. These novel findings are surprising given the wealth of experimental data indicating the high concentration of RGD peptides inhibit rather than induce angiogenesis (11, 28, 29). A growing body of evidence now suggests that low concentrations of certain RGD peptides may actually enhance angiogenesis and tumor growth (30), which may explain at least in part the minimal impact of cyclic RGD peptide antagonists of αβ3 and αβ5 in human clinical trials (31). Our current studies provide new evidence suggesting that in addition to variations in concentrations that may alter the biological response of certain RGD peptides, the specific composition of the amino acids C-terminal to the RGD motif within naturally occurring epitopes may confer unique pro-angiogenic and inflammatory activity. Taken together, our studies are consistent with a novel mechanism by which the RGDKGGE collagen epitope may induce angiogenesis and inflammation by stimulating mechanical activation of αβ3 leading to Src-dependent phosphorylation of p38 MAPK that promotes nuclear accumulation of the Yes-associated protein (YAP) and enhances endothelial cell growth.

Experimental Procedures

Reagents, Kits, Chemicals, and Antibodies—Ethanol, methanol, acetonitr, bovine serum albumin (BSA), crystal violet, phosphate-buffered saline (PBS), purified human collagen type IV and collagen type I, AMPA, 3,3,5,5-tetramethylbenzidine, phosphatase inhibitor mixture, and CA (cortisone acetate) were from Sigma. MMP2 was from Chemicon/Millipore (Billerica, MA). FBS was from Science Cell (Carlsbad, CA). Fibroblast growth factor-2 (FGF-2) was obtained from R&D Systems (Minneapolis, MN). Nuclear/cytoplasmic fractionation kit was from Thermo Scientific (Waltham, MA). p38 MAPK inhibitor SB202190 was obtained from Calbiochem. RIPA buffer, protease inhibitor, and Src inhibitor (PP2) were from Santa Cruz Biotechnology (Santa Cruz, CA). HRP-labeled anti-biotin was from Southern Biotechnology (Birmingham, AL). Anti-vWF antibody was from Pharmingen. Antibodies directed to p38 MAPK, phospho-p38 MAPK (Thr-180/Tyr-182), Src, and phospho-Src (Tyr-416) were from Cell Signaling Technology (Danvers, MA). Antibodies against tubulin, TATA-binding protein (TBP), YAP, β3, and phospho-β3 (Tyr-747) were from Santa Cruz Biotechnology. Anti-Igfbp4, anti-total, and phosphorylated INK and anti-MMP9 antibodies were obtained from Abcam (Cambridge, MA). Function blocking antibodies P4C10 (anti-β1), LM609 (anti-αvβ3), and P1F6 (anti-αvβ5) were from R&D Systems (Minneapolis, MN). HRP-conjugated secondary antibodies were from Promega (Madison, WI). Anti-collagen type I antibody was from Rockland (Limerick, PA), and anti-collagen type IV was from Millipore (Billerica, MA). Mouse monoclonal antibodies XL313 and XL166 were developed in our laboratory. Alexa-488, Alexa-594, streptavidin Alexa-594, and phalloidin Alexa-594-labeled antibodies were from Invitrogen. Unlabeled and biotin-labeled synthetic collagen RGD-containing peptides (P-1, CKGDRGAPGC; P-2, CQGPRGD-KGEC; P-3, CAGSRGDDGPPC; P-4, CQGIRGDKGE; P-5, CRGPRGDDQGP) and peptide control (P-C, CQGPSAGAP-GECC) were obtained from QED Biosciences (San Diego, CA).

Generation of Hybridomas Producing Antibodies Reactive with RGD-containing Collagen Peptides—C57BL/6 mice were immunized (100 μg/mouse) every 3 weeks with synthetic RGD-containing collagen peptides (described in Table 1) that were conjugated to keyhole limpet hemocyanin carrier protein. Serum samples were analyzed for reactivity with immobilized RGD containing peptides by solid phase ELISA. Spleen cells from mice exhibiting strong immunoreactivity to the RGD containing collagen peptides were fused with myeloma cell line SP-20 by standard techniques. Hybridoma clones expressing antibodies exhibiting differential binding to RGD-containing

2 The abbreviations used are: ECM, extracellular matrix; YAP, Yes-associated protein; TBP, TATA-binding protein; HUVEC, human umbilical vein endothelial cell; HRMVEC, human retinal microvascular endothelial cell; HMVEC, human microvascular endothelial cell; CAM, chick chorioallantoic membrane; CM, conditioned medium; CA, cortisone acetate; MMP, matrix metalloproteinase; P-C, peptide control; vWF, von Willebrand factor.
peptides were isolated following ELISA screening and subcloned by limiting dilution.

Cells and Cell Culture—RAW 264.7 and THP-1 cells were from ATCC (Manassas, VA) and cultured in DMEM and RPMI 1640 medium, respectively, in the presence of 10% FBS, 1.0% penicillin/streptomycin, and 1.0% sodium pyruvate. Immortalized BV-2 cells (32) were a gift from Dr. Ling Cao (University of New England, Biddeford, ME) and were cultured in DMEM with 10% FBS, 1.0% penicillin/streptomycin, and 1.0% sodium pyruvate. Human dermal fibroblasts were obtained from Science Cell (Carlsbad, CA) and cultured in fibroblast growth medium with 2.0% FBS and used between passages 4 and 9. Human retinal microvascular endothelial cells (HRMVECs) were obtained from Applied Cell Biology Institute (Kirkland, WA) and cultured in EBM2 supplemented with EGM-2 growth factors. Human umbilical vein endothelial cells (HUVECS) and human microvascular endothelial cells (HMVECS) were obtained from ATCC (Manassas, VA) and cultured in EBM2 with supplemental growth factors EGM-2 or EGM-2MV, respectively. All endothelial cell growth media contained 2% FBS, 1.0% penicillin/streptomycin, and 1.0% sodium pyruvate and were used experimentally between passages 3 and 9. For collection of conditioned media, cells were cultured in basal media under serum-free conditions for 24 h. Conditioned media were collected and concentrated 10× using an Amicon Ultracell, 3-kDa centrifugal ultrafiltration cartridge.

Cell Adhesion Assays—RGD peptides (P-1, P-2, P-3, P-4, P-5, and P-C) were immobilized (1–100 μg/ml) to wells. HUVEC, HMVEC, HRMVEC, and human dermal fibroblasts cells were suspended in adhesion buffer (RPMI 1640 medium containing 1 mM MgCl₂, 0.2 mM MnCl₂, and 0.5% BSA) and 1 × 10⁶ cells were seeded into the wells in the presence or absence of P4C10, LM609, P1F6, or control antibodies (100 μg/ml). Cells were allowed to attach for 25 min at 37 °C. Media containing non-attached cells were aspirated, and attached cells were washed with PBS and stained with crystal violet (19). Cell adhesion was quantified by measuring the optical density of eluted dye (19). Adhesion assays were completed at least three times with triplicate wells.

Collagen Proteolysis and Solid Phase Binding Assays—Collagen type I and collagen type IV were heat-denatured for 15 min and then incubated with APMA-activated MMP2 for 0.5, 1, 4, 8, and 20 h, at 37 °C, followed by a 5 min boil to deactivate remaining MMP. For solid phase ELISAs, synthetic RGD-containing collagen peptides (P-1, P-2, P-3, P-4, P-5, and P-C) were immobilized (1.0 ng/ml to 100 μg/ml) to wells or wells were coated with 10 μg/ml native or MMP2-proteolysed collagen type I or type IV. Wells were blocked with 1% BSA in PBS for 1 h and then incubated with 1 μg/ml mAbs XL313 or XL166 for 1 h, washed, and incubated with anti-mouse HRP-conjugated antibodies (1:5000). Bound mAbs were detected with a 3,3',5'-tetramethylbenzidine substrate as per the manufacturer’s instructions and quantified via spectrometer measurements. All assays were carried out at least four times in triplicate wells.

Chick Chorioallantoic Membrane (CAM) Inflammation and Angiogenesis Assays—The CAM assays were carried out essentially as described with some modifications (19). Briefly, for all experiments CAMs of 10-day-old chick embryos obtained from Charles River (North Franklin, CT) were separated from the shell membrane. Filter discs were either non-treated (inflammation assays), pretreated (angiogenesis assays) with CA (3.0 mg/ml) containing RPMI 1640 medium only, or FGF-2 (40 ng). CAMs were either non-treated or treated topically with mAb XL313, XL166, or a nonspecific control antibody (10 μg/embryo every 24 h for three consecutive treatments). For peptide induction experiments, CAMs were stimulated with 100 ng/ml of the RGD peptides (P-1, P-2, P-4, or P-C) in the presence or absence of the p38 MAPK inhibitor, SB202190 (10 μM). At the end of the incubation period, the embryos were sacrificed, and the CAM tissues were analyzed. Angiogenesis was quantified by counting the number of angioenic branching blood vessels within the area of the filter disc. The angiogenic index was determined by subtracting the mean number of blood vessel branch points from untreated CAMs from each experimental condition as we described previously (19). Eight to twelve embryos were used per condition, and experiments were repeated at least three times.

Transfections of CAM Tissues and Quantitative PCR Analysis—Chick CAM angiogenesis assays were performed as described previously with slight modification. Briefly, FGF-2-containing filter discs were first placed onto dropped CAM. The siRNA duplex-Lipofectamine RNAiMax complexes were formed by mixing 20 pmol of siRNA (TriFECTa™ dicer-substrate kit (IDT)) and 3 μl of Lipofectamine RNAiMAX (Life Technologies), each in 50 μl of Opti-MEM medium. The three Dicer-Substrate 27-mer siRNAs to chicken Yap designed and purchased from IDT with sequences of 5'-AGCACAGAUGCGGACTUAGCGAUAAG-3', 5'-UGACUCAAAGUAGUUCUGAUCCTTT-3', and 5'-GCCCAAGCGCAUGAAGAG-3' were applied to the filter discs in a total volume of 40 μl. After 72 h, individual CAM tissues were dissected, photographed, and frozen for further analyses. For quantitative PCR analysis, the cDNA templates of individual tissues were generated using iScript™ reverse transcription supermix (Bio-Rad). Quantitative PCR was performed using SsoAdvanced™ Universal SYBR Green Supermix (Bio-Rad) according to the manufacturer’s protocol. Chicken Yap expression was analyzed by The CFX96™ optical reaction module with primers specific for chicken Yap (forward primer, AGCCCAAGTGTCCAGTGA-3', and reverse primer, CTACGGAGAGCCATTCC-TGC-3') and 185 (forward primer, GTTCACCGCACTCCGAGATTGA-3', and reverse primer, CCCATCCAGAATGGGTCAA-3'). Experiments were performed twice.

Isolation of Primary CAM Macrophages and Staining Analysis—CAM tissues were dissected, and minced tissue fragments were incubated at 37 °C for 60 min in 10 ml of dissociated medium containing collagenase/Dispase (Roche Applied Science, at 1 mg/ml) and 20% fetal bovine serum in RPMI 1640 medium. The digested tissues were poured through a 100 μm filter strainer into a 50-ml sterile tube and incubated on ice for 4 min to allow settling of debris. The supernatants were transferred into a new tube and spun at 300 × g for 10 min. Red cells were lysed and removed by ACK lysing buffer (Lonza). The cell pellets were washed twice in 10% RPMI 1640 medium. The final suspension of cells was seeded onto glass slides and incubated at 37 °C for 60 min to allow macrophages to adhere. Non-adher-
ent cells were removed by PBS wash. Adhered cells were fixed in ice-cold methanol/acétone for 5 min.

Fixed adherent cells were washed with PBS and blocked with 1% normal goat serum and 2% BSA in TBS at room temperature for 1 h. The cells were then incubated in 2% BSA/TBS containing KUL anti-chick macrophage/macrophages antibody overnight at 4 °C. The slides were washed with TBS and incubated in 2% BSA/TBS containing Alexa Fluor 488 anti-mouse IgG at room temperature for 1 h. After washing, slides were incubated in avidin/biotin blocking solution according to the manufacturer's recommendation (Vector Laboratories) followed by 1-h incubation of biotinylated XL313 (20 μg/ml). Slides were washed and incubated with Alexa Fluor 594-conjugated streptavidin (Life Technologies, Inc.) for 1 h. Next, slides were rinsed with TBS, counter-stained with DAPI, and mounted with Fluoromount aqueous medium (Sigma).

Immunohistochemistry and Immunofluorescence Analysis—

CAMs examined for accumulation of granulation tissue were harvested, fixed in 4% paraformaldehyde, paraffin-embedded and sectioned (4 μm), and stained with Giemsa. CAM tissues analyzed via immunofluorescence were harvested, embedded with OCT, snap-frozen, and sectioned (4 μm). Frozen sections were fixed in 50% methanol, 50% acetone, air-dried, and blocked with 2.5% BSA for 1 h at 37 °C. For expression of the XL313 and XL166 epitopes as well as monocyte/macrophages, FGF-2-stimulated CAM tissues were stained with mAbs XL313 and XL166, (50 μg/ml), or KUL1 (1:250) and then incubated with Alexa-488-labeled secondary antibodies (1:2000 dilution). Co-staining of monocytes/macrophages and the mAb XL313-reactive epitope in FGF-2 stimulated CAMs was performed by sequential staining by first probing with KUL1 (1:250) and then with a biotin-conjugated mAb XL313 (50 μg/ml) and with secondary antibodies Alexa-488 and streptavidin Alexa-594 (1:2000), respectively. Using a similar strategy, sections of RGD peptide-treated CAM tissues were analyzed for phosphorylated p38 MAPK in angiogenic vessels by co-incubation of anti-phospho-p38 MAPK (1:50) and anti-vWF (1:500) antibodies. Subcellular localization of YAP was observed in HUVECs attached to P-2- and P-1-coated glass coverslips. Cells were allowed to attach for 15, 30, or 60 min in the absence of serum and were fixed with 4% paraformaldehyde. Fixed cells were washed and blocked with 2.5% BSA and stained with anti-YAP (1:200) and phalloidin (1:500). All sections and slides were counter-stained with DAPI. For staining and quantification of actin stress fibers, HUVECs were resuspended in adhesion buffer (RPMI containing 1 mM MgCl₂, 0.2 mM MnCl₂, and 0.05% serum) and allowed to bind for 30 min to peptide-coated coverslips. Cells were stained with phalloidin (1:500) and counter-stained with DAPI. The number of cells with actin stress fibers was counted per ×400 field, from 25 fields per condition. Experiments were complete three times.

Cell Proliferation Assays—HUVECs (WT, shYAP1, or control-transfected) or HRMVECs were plated at 2000 cells per well with complete EGM-2 media containing 2.0% FBS in the absence or presence of P-1, P-2, or P-C (100 μg/ml) and allowed to grow for 24 h. Cell growth was monitored using a BrdU or WST-1 assay kits according to the manufacturer's instructions. All assays were completed at least three times in triplicate wells.

Inhibitor Experiments—Endothelial cells (HUVECs or HRMVECs) were incubated in serum-free media with 1 mM MgCl₂, 0.2 mM MnCl₂ in the presence of an Src inhibitor, PP2 (10 μM), a p38 MAPK inhibitor SB202190 (10 μM), or vehicle only (DMSO) for 10 min at 37 °C. Treated cells were then seeded onto immobilized P-1 and P-2, and lysates were collected at 15 min.

Western Blot—Whole cell and CAM tissue lysates were collected in RIPA buffer supplemented with 1× protease inhibitor and 1× phosphatase inhibitor and were run on polyacrylamide gels using denaturing conditions. Prior to loading the gels, 6× sample buffer was added to each of the lysates (final concentration 1×) and boiled for 5 min. Twenty to 50 μg of total protein were loaded into each lane. For detection of proteins larger than 50 kDa, 10% gels were used; for proteins smaller than 50 kDa, 15% gels were used. Gels were run at 60 V until the dye front passed through the stacking gel, and then the voltage was increased to 100 V for the remainder of the separation. Precision Plus protein standards (Bio-Rad) were used to visualize migration. Protein was transferred to nitrocellulose membranes using a wet tank system and blocked for 1 h using 10% nonfat dried milk in Tris-buffered saline with 0.01% Tween 20 (TBS-T). Membranes were incubated with primary antibodies (anti-coll-1 (1:250), anti-coll-IV (1:250), mAb XL313 (2 μg/ml), mAb XL166 (2 μg/ml), anti-β-actin (1:5000), anti-phospho-β3 (1:7000), anti-β3 (1:1000), anti-phospho-Src (1:500), anti-Src (1:500), anti-phospho-p38 MAPK (1:500), anti-p38 MAPK (1:2000), anti-YAP (1:500), anti-tubulin (1:2000), anti-TBP (1:1000)) in 5% BSA in TBS-T overnight at 4 °C with gentle agitation. Membranes were washed three times in TBS-T for 5 min. Blots were then incubated with HRP-conjugated secondary antibodies (1:15,000) in 1% nonfat milk in TBS-T for 1 h. Membranes were washed a second time as indicated above and exposed to chemiluminescent substrate for 3 min prior to exposure to autoradiography film in a dark room. Western blot bands were quantified using ImageJ software based on pixel intensity.

Transfections and Lentiviral Transductions—Raw cells were transfected with 1 μg of HuSH shRNA plasmids for collagen type and α2 using Amaxa cell line nucleofector kit V (program T024). Constructs expressing 21-nucleotide short hairpin RNAs (shRNA) targeting human YAP (shYAP) or non-targeting control (shNT, Sigma, SHC002) were used. Human-targeting shYAP1 lentiviral shRNA was obtained from the Thermo Scientific RNAi consortium (TRCN0000107625). Constructs were packaged into lentivirus, pseudotyped with the vesicular stomatitis virus glycoprotein. Transduction was performed by incubating cells with lentivirus, and stably transduced cells were subsequently used for studies. All cell lines were verified by morphology and mouse and human YAP-specific PCR. The efficacy of YAP knock down was determined to be between 70 and 80%. Endothelial cells were certified mycoplasma-negative by PCR (Lonza), and primary cell cultures were used within the indicated passage numbers. Cells were transduced and selected using puromycin.

Statistical Analysis—Statistical analysis was performed using the InStat statistical program for Macintosh computers. Data
were analyzed for statistical significance using Student’s t test. p values < 0.05 were considered significant.

Results

Cryptic RGD Containing Peptides from Collagen Type I Support Cell Adhesion—Studies have documented the capacity of ECM proteins containing the short amino acid sequence RGD to support interactions mediated by integrin receptors (33). The ability of cells to interact with RGD sites within the context of larger glycoproteins depends on many factors, some of which include the adjacent flanking sequences surrounding the core RGD tri-peptide as well as the geometrical configuration of the intact molecule and how these molecules are oriented within the context of the interconnected network of other ECM proteins (24, 25, 33). Flanking sequences immediately C-terminal to the RGD site can govern integrin selective binding (24, 25, 33). RGD motifs can be cryptic and inaccessible to cell surface receptors as is illustrated in the case of triple helical collagen (34). In this regard, five different cryptic collagen-containing sites exist within human collagen type I, each with distinct flanking sequences (Table 1).

Given the importance of the RGD tri-peptide motif, we sought to establish that the five distinct RGD sites represented sequences may help convey distinct properties, we synthesized the five different cryptic RGD-containing epitopes along with their associated flanking sequences (Table 1).

Five different cryptic RGD-containing sites exist within human collagen type I, each with distinct flanking sequences. Synthetic peptides of these five sequences were generated and designated P-1 through P-5. Additionally, P-C was generating working RGD containing the RGD tri-peptide motif. NA means not applicable.

| Peptide | Amino acid sequence | Location |
|---------|---------------------|----------|
| P-1     | KGDRGDAPG           | Col1a1 (742–750) |
| P-2     | QGPRGDKE            | Col1a1 (1090–1098) |
| P-3     | AGSRGDGGP           | Col1a2 (774–782) |
| P-4     | QGIRGDKE            | Col1a2 (1002–1010) |
| P-5     | RGPRDQGQP           | Col1a2 (819–827) |
| PC      | QGPSGAPGE           | NA       |

mAb XL313 Exhibits Selective Binding for Proteolyzed Collagen Type I—We sought to determine whether mAb XL313 had the capacity to bind its RGD motif within the context of the full-length collagen molecule. To facilitate these studies, we first incubated denatured collagen type I or IV with MMP-2 for 12 h to generate proteolyzed collagen. As expected, MMP-2-mediated proteolysis of collagen type I and IV resulted in the generation of multiple fragments as indicated by Western blot analysis using antibodies specifically directed to either collagen type I (Fig. 2A, left panel) or collagen type IV (Fig. 2A, right panel). mAb XL313 specifically directed to the RGDKGE collagen sequence exhibited weak reactivity with intact collagen type I, and no reactivity with type IV collagen under denaturing and reducing conditions (Fig. 2B). Moreover, mAb XL313 demonstrated little reactivity to intact collagen type I or collagen type IV under non-denatured and non-reducing conditions of the ELISA while readily binding to proteolyzed collagen type I under these identical conditions (Fig. 2C). mAb XL313 failed to recognize other RGD-containing ECM proteins, including vitronectin or fibronectin (data not shown). To further examine the generation of the low molecular weight RGDKGE-containing collagen fragments, we examined a time course of MMP-2-mediated collagen proteolysis. Although mAb XL313 can recognize non-proteolyzed collagen type I that is structurally altered under denaturing and reducing conditions, MMP-2-mediated degradation of collagen type I resulted in a time-dependent generation of mAb XL313 reactive collagen fragments (Fig. 2D). The majority of the collagen was proteolyzed into mAb XL313-reactive fragments of ~14 to 16 kDa by 8 h. These findings are consistent with the notion that XL313 collagen epitopes appear to be cryptic within a structurally intact non-denatured collagen type I molecule and that proteolytic degra-
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Differential Roles of Cryptic RGD Collagen Motifs on Angiogenesis and Inflammation in Vivo—Our previous studies have shown that collagen remodeling within the vascular basement membrane can result in exposure of multiple non-RGD cryptic collagen sites, including the HUIV26 and HU177 epitopes that can play active roles in angiogenesis (21, 23). Given these findings and our current observations, we sought to determine whether distinct RGD-containing epitopes are exposed in vivo. First, to examine whether these RGD epitopes can be generated during angiogenesis, we used the chick chorioallantoic membrane (CAM) model (19). The CAMs of chick embryos were stimulated with FGF-2, and the generation of RGD-containing collagen epitopes was examined using mAbs XL313 and XL166. As shown in Fig. 3A, little mAb XL313 or XL166-reactive epitope could be detected in non-stimulated CAMs. In contrast, RGD-containing collagen epitopes (Fig. 3A, green) were readily detected in FGF-2-stimulated CAM tissues confirming the differential generation of RGD-containing epitopes recognized by these antibodies. Interestingly, mAb XL313- and XL166-reactive RGD epitopes did not exhibit a typical extracellular fibril collagen pattern, but rather they exhibited punctate distribution with both intracellular and scattered extracellular localization. The intracellular distribution is similar to the staining pattern of collagen degradation products previously documented within intracellular vesicles of macrophages (35).

**FIGURE 1. Cryptic RGD-containing peptides from collagen type I support cell adhesion.** Peptides corresponding to the five different RGD-containing motifs of human collagen type I were synthesized with flanking cysteines and immobilized on non-tissue culture plates. The ability of stromal cells to attach to the peptides (A–D) or the ability of monoclonal antibodies to bind the immobilized peptides (E and F) was assessed. A, HUVEC adhesion to immobilized peptides. B, HMVEC adhesion to immobilized peptides. C, HRMVEC adhesion to immobilized peptides. D, human dermal fibroblast (HDF) adhesion to immobilized peptides. E, reactivity of mAb XL313 to immobilized peptides. F, reactivity of mAb XL166 to immobilized peptides. Data bars represent mean binding ± S.D. from at least three experiments each using triplicate wells. *, p < 0.05.
Given the generation of these distinct sets of RGD-containing epitopes, we next examined whether they may play active roles in regulating angiogenesis and inflammation. Angiogenesis was induced within the CAMs of 10-day-old chick embryos with FGF-2 using filter discs coated with CA to reduce growth factor-associated inflammation. As shown in Fig. 3B, treatment with mAb XL313 directed to the cryptic RGDKGE-containing epitope significantly (p < 0.05) inhibited FGF-2-induced angiogenesis by greater than 75% as compared with non-treated or control antibody. Surprisingly, mAb XL166 that specifically binds the remaining three distinct cryptic RGD collagen peptides (P-1, P-3, and P-5) had no effect. These surprising findings are consistent with the possibility that although distinct RGD-containing collagen epitopes are readily generated in vivo, they may not be functionally redundant.

The chick CAM has been routinely used to assess inflammation and granulation tissue formation, which are largely dependent on infiltration of chick heterophils (the avian equivalent of neutrophils), macrophages, and activated fibroblasts (36, 37). To study the potential differential biological impact of RGD epitopes, we examined FGF-2-induced inflammation by carrying out similar experiments in the absence of CA and quantifying CAM thickening. As shown in Fig. 3C, treatment with FGF-2-induced CAM inflammation as indicated by robust tissue thickening (top panels), extensive infiltration of inflammatory infiltrates as indicated by Giemsa stain (middle panel), and increased accumulation of monocytes and macrophages (bottom panel) following staining with an antibody directed to avian-specific monocytes and macrophages. Macrophage infiltration of the CAMs stimulated with FGF-2 was significantly (p < 0.05) enhanced by over 2-fold as compared with control (Fig. 3D) indicating that FGF-2-induced inflammation in this model is associated with the recruitment and accumulation of granulation tissue-associated macrophages as well as other stromal cells, including fibroblasts.

To examine whether the RGD-containing collagen epitopes play a role in the FGF-2-stimulated inflammatory response, we examined this FGF-2-induced inflammatory response in the presence or absence of anti-RGD-specific antibodies XL313 and XL166. Quantification indicated that FGF-2 stimulation in the absence of CA resulted in ~50% of the CAMs showing robust formation of thick granulation tissue (Fig. 3E). Treatment of CAMs with CA, a well documented anti-inflammatory agent, significantly (p < 0.05) reduced the percentage of CAMs exhibiting thickening by greater than 80% as compared with either no treatment or control antibody. Moreover, reduced
levels of inflammatory infiltrates, including macrophages, were also observed (Fig. 3F). Interestingly, treatment of CAMs with mAb XL313 also significantly (p < 0.05) inhibited the number of CAMs exhibiting inflammation by ~75% as well as reducing the relative levels of inflammatory infiltrates as compared with controls (Fig. 3, E and F). In contrast, treatment of CAMs with mAb XL166 showed no significant (p > 0.05) impact. These data are consistent with our previous findings when examining angiogenesis suggesting a differential role for specific cryptic RGD epitopes in vivo.

 Generation of XL313 Cryptic RGDKGE Epitope—During angiogenesis and inflammation, multiple cell types, including...
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endothelial cells, fibroblasts, and macrophages proteolytically remodel extracellular collagen creating a permissive microenvironment that facilitates stromal cell infiltration and new blood vessel growth. A variety of cells, including fibroblasts and endothelial cells, can express collagen, and moreover, partially degraded collagen can be internalized and further processed by activated M2-like macrophages leading to the generation of low molecular weight fragments (35, 38). Given the unique pattern of mAb XL313 immunoreactivity observed in vivo, we examined whether stromal cells associated with angiogenesis and inflammation could generate the RGDKGE-containing epitope. Whole cell lysates were prepared from fibroblasts and endothelial cells, and Western blotting was carried out. Although some mAb XL313-reactive species could be detected in lysates of endothelial cells and fibroblasts, the major immunoreactive species migrated between ~75 kDa and 28 kDa (Fig. 4A, left panel) indicating that these species were unlikely to represent intact collagen. In addition, minimal if any low molecular weight species was detected that might correspond to the major 14- to 16-kDa collagen fragments detected following MMP-2-mediated proteolysis. Interestingly, although a small amount of an ~20-kDa XL313-reactive species was detected in serum-free CM collected from HUVECs, we failed to detect significant levels of immunoreactive fragments in fibroblast CM (Fig. 4A, right panel). Given the minimal reactivity observed in these cell types known to express collagen and the fact that FGF-2 induced a strong angiogenic and inflammatory response in CAMs that was associated with an extensive infiltration of macrophages, we examined FGF-2-treated CAM tissues for the co-distribution of the XL313 epitope and macrophages. As shown in Fig. 4B, the XL313-reactive epitope (red) co-localized with a subset of macrophages (green) in FGF-2-stimulated CAMs. In similar studies, the XL313 reactive epitope localized with primary macrophages isolated from CAM tissues (data not shown), suggesting the possibility that a subset of macrophages may be at least one source of the XL313 collagen epitope. To study this possibility, we examined both whole cell lysates (Fig. 4C, left panel) and serum free CM (Fig. 4C, right panel) from three different macrophage-like cell lines. In contrast to endothelial cells and fibroblasts, strong immunoreactive species were readily detected in cell lysates and CM from multiple macrophage-like cell lines, including Raw 264.7, THP1, and BV2. Importantly, low molecular mass immunoreactive species migrating between 14 and 16 kDa were readily detected in serum-free CM (Fig. 4C, right panel). These 14–16-kDa immunoreactive species corresponded to the size of the low molecular mass mAb XL313-reactive collagen fragments detected following MMP-mediated proteolytic digestion of purified collagen. To further study the XL313-immunoreactive species detected in macrophages, we examined the generation of the 14–16-kDa RGDKGE-containing XL313 fragments in Raw 264.7 macrophages. First, to examine the expression of collagen in these macrophages, we confirmed the expression of mRNA for the α1 and α2 chains of collagen type I in RAW macrophages by PCR (data not shown). Second, although we did not detect any full-length collagen type I in either cell lysates or CM from macrophages using anti-collagen type I-specific antibodies, we did detect low molecular mass fragments migrating at the same molecular mass as the RGDKGE-containing epitope in both cell lysates and CM using either mAb XL313 or an anti-collagen I-specific antibody (Fig. 4D). Importantly, shRNA-mediated knockdown of the α2 chain of collagen type I reduced immune detection of the low molecular mass species (14–16 kDa) in both lysates and CM when using either anti-collagen specific antibodies or mAb XL313 (Fig. 4D). When taken together, these findings are consistent with the possibility that macrophages are at least one cellular source of the RGDKGE collagen epitope.

Induction of Angiogenesis and Inflammation in Vivo by Soluble RGDKGE but Not a Related RGDAPG-containing Collagen Epitope—Given the differential expression and bio-distribution of the RGDKGE epitope in angiogenic CAM tissues and its expression in macrophage-conditioned medium, we examined whether a soluble circulating form of this RGDKGE epitope could be generated. To examine this possibility, chick embryos were either untreated or stimulated with FGF-2, and serum was collected 3 days later. As shown in Fig. 5A, low levels of circulating mAb XL313 immunoreactive epitope could be detected in the serum from non-stimulated control chick embryos. In contrast, a significant (p < 0.05) 2-fold increase was detected in the levels of circulating XL313 epitope following FGF-2 stimulation. Importantly, although the exact nature of the circulating XL313 epitope is not known, mAb XL313 demonstrated the ability to dose-dependently detect increasing levels of P-2 peptide representing the RGDKGE collagen epitope (data not shown), confirming the ability of the ELISA to dose-dependently detect the XL313 epitope. These findings suggest the differential detection of a soluble form of this RGDKGE-containing collagen epitope.

Given that a soluble form of the RGDKGE epitope could be detected in vivo, we next examined whether a soluble peptide containing the XL313 epitope might actively regulate angiogenesis and inflammation. To examine this possibility, we assessed the effects of the RGDKGE-containing collagen peptide on angiogenesis and inflammation in the chick CAM. As shown in Fig. 5B, the XL313 RGDKGE-containing collagen peptide (P-2) dose-dependently enhanced angiogenesis with maximum induction observed at a dose of 100 ng/CAM. Stimulation of CAMs with the RGDKGE-containing collagen peptide P-2, but not the related RGDAPG-containing collagen peptide P-1 or P-C, significantly (p > 0.05) induced angiogenesis as compared with no treatment (Fig. 5C).

Previously published studies implicate inflammation as a contributing factor in regulating angiogenesis (37, 39). Therefore, we also examined the effects of the soluble RGD-containing collagen peptides on inflammation. As shown in Fig. 5D, FGF-2 induced a strong inflammatory response in the chick CAM in the absence of CA as indicated by ~55–60% of the CAMs exhibiting extensive tissue thickening. Interestingly, minimal evidence of tissue inflammation was observed following stimulation with either P-C or the RGDAPG-containing collagen peptide P-1, although the RGDKGE collagen peptide P-2 significantly (p < 0.05) induced inflammation to nearly that of FGF-2 stimulation (Fig. 5D). Given that collagen peptide P-4 contains the identical RGDKGE sequence within its C-terminal region, but is different from that of P-2 in that it contains an isoleucine at the 3rd amino acid position instead of a proline
present in P-2, we compared the ability of these collagen peptides to stimulate inflammation. As shown in Fig. 5, E and F, peptides P-2 and P-4 containing the RGDKGE motif, showed no statistically significant difference in ability to induce CAM inflammation. Interestingly, although FGF-2 stimulated CAM tissues exhibited significantly ($p < 0.05$) higher relative levels of macrophages than either P-2 or P-4, no difference in the relative levels of macrophages was detected between P-2 and P-4 (data not shown). Collectively, these studies suggest that the biological impact of RGD-containing collagen peptides on angiogenesis and inflammation in the chick CAM may depend in part on their associated flanking sequences.
FIGURE 5. Induction of angiogenesis and inflammation in vivo by soluble RGDKGE but not a related RGDAPG-containing collagen epitope. A, serum from chick embryos in which the CAMs had been either not stimulated (NT) or stimulated with FGF-2 was collected. Relative levels of circulating mAb XL313-reactive epitope were quantified by solid phase ELISA. Data bars represent mean mAb XL313 reactivity ± S.E. (n = 12–13). B, quantification of dose-dependent induction of angiogenesis by RGDKGE-containing collagen peptide P-2. Data bars indicate mean number of branch points ± S.E. from 8 to 12 animal per condition. C, quantification of CAM angiogenesis associated with RGD-containing collagen peptides P-1 (KGDRGDAPG), P-2 (QGPRGDKGE), or control peptide P-C (QGPSGSPGE). Data bars indicate mean angiogenic index derived by subtracting the number of branch points from non-stimulated CAMs ± S.E. from 8 to 12 animals per condition. D, quantification of CAM inflammation associated with FGF-2, RGD-containing collagen peptides P-1 (KGDRGDAPG), P-2 (QGPRGDKGE), or control peptide P-C (QGPSGSPGE). E, quantification of CAM inflammation associated with FGF-2, RGD-containing collagen peptides P-2 (QGPRGDKGE), or P-4 (QGIRGDKGE). All experiments were conducted at least three times with similar results. *, p < 0.05. F, representative examples of inflammatory CAM thickening (×40) (top panel), or Giemsa-positive inflammatory infiltrates (×200) (bottom panel) from CAM inflammation associated with FGF-2, peptides P-2 (QGPRGDKGE), or P-4 (QGIRGDKGE). Scale bars, 50 μm.
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RGDKGE-containing Peptide P-2-induced Angiogenesis Depends on p38 MAPK—Studies have indicated that angiogenesis and inflammation in the chick CAM is associated with alterations in MAPK signaling, including p38 MAPK (40, 41). To examine mechanisms that may regulate angiogenesis following stimulation with the RGDKGE-containing collagen peptide P-2, we examined CAM tissues from untreated or RGD peptide-treated animals. As shown in Fig. 6A, elevated levels of phosphorylated p38 MAPK were detected in lysates from CAMs treated with RGDKGE-containing peptide P-2 as compared with either untreated or RGDAPG-containing peptide P-1-treated animals. Quantification of CAM tissues (n = 12)
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indicated a significant ($p < 0.05$) 4-fold increase in phosphorylation of p38 MAPK as compared with controls (Fig. 6B). Consistent with our previous studies (41), although activated p38 MAPK could be detected in multiple cell types in chick CAMs, co-staining analysis indicated expression of phosphorylated p38 MAPK in vWF-positive blood vessels (Fig. 6C). Given these findings we examined the effects of the RGD-containing collagen peptides on the levels of phosphorylated p38 MAPK in endothelial cells. As shown in Fig. 6, D and E, whereas both RGD-containing collagen peptides P-1 and P-2 can support cell binding, interactions with RGDKGE collagen peptide P-2 resulted in enhanced phosphorylation of p38 MAPK as compared with RGDAPG collagen peptide P-1 in HUVECs (left panel) or HRMVECs (right panel). In similar studies, whereas a small increase (1.39-fold) in the phosphorylation of JNK was detected in HUVECs following interactions with P-2 as compared with P-1, this increase did not meet statistical significance (Fig. 6F). Importantly, a functional role for p38 MAPK in mediating RGDKGE peptide P-2-induced angiogenesis in vivo was demonstrated, as P-2-induced angiogenesis was significantly ($p < 0.05$) reduced by a p38 MAPK inhibitor (Fig. 6H). Taken together, these data are consistent with a role for p38 MAPK in the peptide P-2-stimulated pro-angiogenic response observed in vivo.

Collagen Peptide P-2 Binds and Activates $\alpha v \beta 3$ and Stimulates p38 MAPK Activation in an Src-dependent Manner—It is well established that multiple integrins can recognize RGD amino acid motifs within ECM proteins. However, the ability of an integrin to bind distinct RGD epitopes depends in part on its orientation within the parent molecule and the C-terminal amino acid sequences flanking the RGD motif (25–27). Given our observations, we sought to identify potential cell surface receptors that mediate interactions with the RGDKGE-containing collagen peptide P-2. We examined the ability of endothelial cells to bind the RGD collagen peptides P-1 and P-2 in the presence or absence of function-blocking anti-integrin antibodies. As shown in Fig. 7A, anti-$\alpha v$-specific antibody had little effect on HUVEC adhesion to peptide P-2, whereas anti-$\alpha v \beta 3$ antibody significantly ($p < 0.05$) inhibited interactions by 90% as compared with control. Anti-$\alpha v \beta 5$ antibody showed little inhibition. Similar results were observed with P-1, which contains the RGDAPG sequence (Fig. 7B). These data suggest that $\alpha v \beta 3$ can function as a receptor for both RGD-containing peptides in endothelial cells.

Given the ability of both RGD-containing peptides to bind $\alpha v \beta 3$ coupled with the differential effect of these peptides on angiogenesis and inflammation, we sought to determine whether the RGD peptides differentially altered $\alpha v \beta 3$-mediated signaling. Endothelial cells were allowed to attach to P-1 and P-2, and the relative level of $\beta 3$-integrin phosphorylation was examined. As shown in Fig. 7C, in the absence of any growth factors, $\beta 3$ phosphorylation was significantly ($p < 0.05$) increased by over 3-fold following ligation of RGDKGE peptide P-2 as compared with RGDAPG peptide P-1. Ligand binding of $\beta 3$-integrin can induce receptor clustering and initiate downstream signaling events, including activation of protein kinases such as Fak and Src (42). Therefore, we assessed the differential phosphorylation of these protein kinases following the early mechanically-mediated interactions with the RGD-containing collagen peptides. Little if any Fak phosphorylation was detected following ligation of either P-1 or P-2 under these serum-free conditions (data not shown). In contrast, Src phosphorylation was enhanced by nearly 2-fold following interactions with P-2 as compared with P-1 (Fig. 7D). Moreover, activation of $\beta 3$-integrin and p38 MAPK in endothelial cells following binding to P-2 was dependent on Src as incubation of endothelial cells with an Src inhibitor reduced $\beta 3$ and p38 MAPK phosphorylation in cells attached to collagen peptide P-2 (Fig. 7, E and F). Activation of Src and p38 MAPK can lead to enhanced actin polymerization and stress fiber formation (43–45); therefore, we examined the actin cytoskeleton in endothelial cells following binding to the RGD collagen peptides. Interestingly, endothelial cell interactions with the RGDKGE collagen peptide P-2 resulted in accelerated actin stress fiber formation as a statistically significant increase in the percentage of cells with actin stress fibers were observed in endothelial cells attached to P-2 (RGDKGE) as compared with interactions with P-1 (RGDAPG) peptide (Fig. 7, G and H). Given the ability of integrin $\alpha v \beta 3$ to bind both the RGD-containing collagen peptides P-1 and P-2, we examined expression of $\alpha v \beta 3$-integrin in endothelial cells attached to these RGD peptides (data not shown). Although some variation in distribution of $\alpha v \beta 3$ was observed between cells exhibiting a more spread morphology, relatively little difference could be detected in the relative localization of $\alpha v \beta 3$ between cells attached to P-1 and P-2 (data not shown). Taken together, these findings are consistent with the ability of the conserved RGDKGE collagen epitope to stimulate initiation of a $\beta 3$-integrin-dependent mechano-transduction pathway.

Cellular Interactions with RGDKGE-containing Peptide Enhance Nuclear YAP Accumulation and Endothelial Cell Growth—Our data indicate that although both RGD-containing collagen peptides P-1 and P-2 can support an initial $\beta 3$-in-

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integrin-mediated endothelial cell adhesive interaction, collagen peptide P-2 selectively enhanced β3-integrin phosphorylation leading to increased activation of p38 MAPK and accelerated actin stress fiber formation. Actin stress fiber formation and enhanced mechanical tension may contribute to re-localization of the transcriptional co-activator YAP to the nucleus (46). Moreover, YAP has recently been implicated in regulating angiogenesis and endothelial cell growth (47, 48). We examined...
YAP localization following endothelial cell binding to the distinct collagen RGD-containing peptides in the absence of growth factor stimulation. As shown in Fig. 8A, enhanced level of nuclearly localized YAP was detected by 15 min following endothelial cell binding to collagen peptide P-2 as compared with P-1. The differential nuclear localization of YAP was not consistently detected at later times points, likely due to the lack of serum or growth factors during the time course of binding (49). To confirm and quantify the enhanced nuclear accumulation of YAP, Western blot analysis was carried out. As shown in Fig. 8B, the level of nuclear YAP was increased in endothelial cells attached to peptide P-2 as compared with peptide P-1. Quantification of four independent experiments indicated a significant (p < 0.05) 2.7-fold increase in the relative levels of nuclear YAP in cells attached to P-1 set at 1.0. D, Western blot of isolated nuclear fractions from cell lysates of endothelial cells (HUVEC) pre-incubated with Src or p38 MAPK inhibitor and stimulation with soluble RGD-containing collagen peptide P-2 and probed with YAP or TATA-binding protein (TBP). E, quantification of the mean relative levels of nuclear YAP following ImageJ analysis in cells attached to P-2 and pre-incubated with Src or p38 MAPK inhibitor. Data bars represent mean fold change ± S.E. from 2 to 3 independent experiments with the relative levels of nuclear YAP in cells under each condition with control (DMSO-treated) set at 1.0.

**FIGURE 8.** Cellular interactions with RGDKGE collagen-containing peptide facilitates nuclear YAP accumulation. A, representative examples of endothelial cells (HUVECs) attached to RGD-containing collagen peptides P-1 and P-2 stained for YAP (green). Scale bar, 50.0 μm. Photos were taken in a single plane at a magnification of ×630. B, Western blot of nuclear fraction of HUVEC lysates following binding to RGD-containing collagen peptides P-1 or P-2 probed with anti-YAP and anti-TATA-binding protein (TBP). C, quantification of the mean relative levels of nuclear YAP following ImageJ analysis. Data bars represent mean fold change ± S.E. from four independent experiments with the relative levels of nuclear YAP in cells attached to P-1 set at 1.0. D, Western blot of isolated nuclear fractions from cell lysates of endothelial cells (HUVEC) pre-incubated with Src or p38 MAPK inhibitor and stimulation with soluble RGD-containing collagen peptide P-2 and probed with YAP or TATA-binding protein (TBP). E, quantification of the mean relative levels of nuclear YAP following ImageJ analysis in cells attached to P-2 and pre-incubated with Src or p38 MAPK inhibitor. Data bars represent mean fold change ± S.E. from 2 to 3 independent experiments with the relative levels of nuclear YAP in cells under each condition with control (DMSO-treated) set at 1.0.

YAP localization following endothelial cell binding to the distinct collagen RGD-containing peptides in the absence of growth factor stimulation. As shown in Fig. 8A, enhanced level of nuclearly localized YAP was detected by 15 min following endothelial cell binding to collagen peptide P-2 as compared with P-1. The differential nuclear localization of YAP was not consistently detected at later times points, likely due to the lack of serum or growth factors during the time course of binding (49). To confirm and quantify the enhanced nuclear accumulation of YAP, Western blot analysis was carried out. As shown in Fig. 8B, the level of nuclear YAP was increased in endothelial cells attached to peptide P-2 as compared with peptide P-1. Quantification of four independent experiments indicated a significant (p < 0.05) 2.7-fold increase in the relative levels of nuclear YAP in endothelial cells attached to P-2 as compared with P-1 (Fig. 8C). Given the selective ability of P-2 to enhance Src-dependent activation of p38 MAPK, we examined whether the enhanced nuclear accumulation of YAP following RGDKGE peptide P-2 stimulation was Src- and/or p38 MAPK-dependent. As shown in Fig. 8D, nuclear YAP accumulation following P-2-stimulated endothelial cells was reduced by treatment with either Src or p38 MAPK inhibitors as compared with control. Quantification of 2–3 independent experiments indicated a 60–80% reduction in the relative levels of nuclear YAP in endothelial cells attached to P-2 following treatment with either Src or P38 inhibitors (Fig. 8E). These findings are consistent with a role for Src and p38 MAPK in mediating the RGDKGE P-2-stimulated nuclear accumulation of YAP.

**YAP Contributes to P-2-stimulated Endothelial Cell Growth—** Nuclear localization of YAP can contribute to the regulation of endothelial cell growth (47). Moreover, YAP has been shown to play a functional role in regulating angiogenesis (48, 49) and compounds known to inhibit YAP inhibit angiogenesis in the chick CAM model (50, 51). To this end, we identified an siRNA capable of significantly reducing the relative levels of YAP following in vivo transfection of CAM tissues (Fig. 9A). Consistent with these studies suggesting a role for YAP in regulating angiogenesis in the chick CAM, siRNA-mediated knockdown of YAP within the chick CAM significantly reduced FGF-2-induced angiogenesis (Fig. 9B). Next, we examined the effects of RGD-containing collagen peptides on HUVEC and HRMVEC growth. Addition of soluble P-2 significantly (p < 0.05) enhanced endothelial cell growth, although stimulation with RGDAPG peptide P-1 had minimal if any effect (Fig. 9, C and D). YAP is thought to play roles in regulating the expression of multiple angiogenesis and inflammatory factors (47–49, 51); therefore, we examined whether P-2-stimulated endothelial cell growth might depend on YAP.
FIGURE 9. YAP contributes to P-2-stimulated endothelial cell growth. A, siRNAs directed to avian YAP were screened for their ability to knock down expression of YAP following in vivo transfection within FGF-2-induced CAM tissues by quantitative PCR. Data bars represent mean fold change in relative YAP levels ± S.E. from each condition. *, p < 0.05. B, representative examples of angiogenic CAM tissues from each condition (top panel). Photos were taken at a magnification of ×40. Quantification of the number of branching blood vessels from each condition (bottom panel). Data bars represent mean blood vessel branch point counts ± S.E. from 6 to 8 CAMs per conditions. *, p < 0.05. C and D, quantification of HUVEC (C) and HRMVEC (D) growth in the presence or absence of exogenously added RGD-containing peptides P-1 or P-2. Data bars represent mean endothelial cell growth (± S.E.) from 3 to 4 independent experiments expressed as percent of control. E and F, quantification of endothelial cell growth in the presence or absence of exogenously added RGD-containing peptides P-1 or P-2 of cells transduced with nonspecific shRNA (E) or YAP-specific shRNA (F). Data bars represent mean endothelial cell growth (± S.E.) from four independent experiments. G, quantification of endothelial cell growth in the presence or absence of exogenously added VEGF (50 ng/ml) or 10% serum from cells transduced with YAP-specific shRNA. Data bars represent mean endothelial cell growth (± S.E.) from four independent experiments. *, p < 0.05.
To facilitate these studies, we transduced endothelial cells with YAP-specific or nonspecific shRNA. Addition of soluble RGD-KGE-containing collagen peptide P-2 to endothelial cells, but not the related RGDPAG-containing peptide P1, significantly (p < 0.05) enhanced growth in control transduced cells (Fig. 9E), although the RGDKG peptide P-2 failed to induce growth in endothelial cells in which YAP was knocked down (Fig. 9F). Importantly, YAP knockdown cells were capable of proliferating as enhanced growth was observed following VEGF or high serum stimulation (Fig. 9G). Taken together, our findings are consistent with a mechanism by which RGDKG-containing collagen peptide P-2 may initiate a unique mechano-signaling cascade leading to the activation of p38 MAPK in an Src-dependent manner that ultimately leads to nuclear YAP accumulation and enhanced endothelial cell growth.

Discussion

A wide array of alterations in the composition and biomechanical properties of ECM proteins is known to occur during angiogenesis, and studies are beginning to define how these changes contribute to new blood vessel development (18, 21–23, 51, 53–55). Among the key cell surface molecules that play roles in mechano-transduction to facilitate information flow from outside the cell to the inside are integrin receptors. Integrins may act like information hubs by sensing diverse extracellular inputs and relaying this information into a complex network of intracellular circuits that ultimately modulate cellular behavior (4). The precise molecular mechanisms by which cells fine-tune their response to changes within the stromal microenvironment are not completely understood. Further complicating our understanding of new vessel development is the expanding number of cell types that contribute to tissue-specific control of angiogenesis such as distinct subsets of stromal fibroblasts, progenitor cells, and a variety of inflammatory cells such as neutrophils, mast cells, and macrophages. The roles played by these diverse cells during angiogenesis range from secretion of cytokines, chemokines, and proteolytic enzymes to the differential expression of other pro- and antiangiogenic factors. Thus, tight control mechanisms must operate to allow coordination between these diverse compartments to govern tissue-specific vascular responses.

Among the most well studied integrins known to play a role in angiogenesis is αvβ3. The complexity by which αvβ3 regulates angiogenesis is illustrated by the fact that this receptor may exhibit both pro- and antiangiogenic functions (7–15). Studies suggest that the distinct biological responses stimulated by binding to αvβ3 may depend on many factors, including the mechanical and biochemical features of the particular ligands, the cell types within which αvβ3 is expressed, as well as the concentration and manner by which the ligands are presented to the receptor (7–9, 24–27). For example, studies indicate that αvβ3 binding to specific NC1 domains of collagen or selected RGD peptides can induce apoptosis, induce arteriole contraction, and inhibit angiogenesis (19, 29, 56), although other αvβ3 ligands may promote cell survival, induce vascular dilation, and support angiogenesis (21–23, 30, 57). These observations are consistent with the notion that the final outcome of αvβ3-mediated signaling may depend to a large extent on the particular characteristics of the ligand. Although a wealth of data has shown that RGD peptides can inhibit angiogenesis when administered exogenously, the approach of using a cyclic RGD peptide to control tumor growth failed to significantly impact glioblastoma progression and patient survival in late stage clinical testing (31). Interestingly, studies have indicated that specific RGD peptides may activate β3-integrins (58, 59), and under-defined experimental conditions induce angiogenesis and tumor growth (30). These findings and other studies suggesting that amino acids C-terminal to the RGD motif play roles in governing integrin-selective binding prompted us to examine the biological significance of naturally occurring RGD-containing epitopes on angiogenesis.

Sequence analysis of RGD sites within collagen type I indicate that the KGE tri-peptide motif that is C-terminal to the RGD site was highly conserved among diverse species, although considerable variation is observed in the other collagen RGD-flanking sequences. Although all five of the collagen RGD epitopes can support cell binding, the highly conserved RGDKG collagen peptide P-2 may play a functional role in angiogenesis and inflammation given that mAb XL313 directed to this epitope but not an antibody that recognizes the other three RGD collagen sites inhibited angiogenesis and inflammation in vivo. Although we cannot precisely differentiate between the three other naturally occurring non-RGDKG-containing collagen epitopes, one or more of these epitopes were detected in vivo in addition to the RGDKG epitope. Given that these RGD-containing epitopes are thought to be largely cryptic and not readily accessible to cell surface receptors, our findings are consistent with active collagen remodeling resulting in generation of neoepitopes during new vessel formation.

ECM remodeling occurs as an early event during angiogenesis and multiple proteolytic enzymes, including MMPs as well as serine and cysteine proteases that have the capacity to degrade intact or structurally altered forms of collagen (18, 60). Although our in vitro studies indicate that MMP-2-mediated degradation of collagen can lead to the generation of low molecular weight fragments recognized by mAb XL313, the precise mechanism by which the RGDKG collagen epitope is generated in vivo is not completely understood. Analysis of angiogenic CAM tissues suggests that a subset of macrophages may be an important source of the RGDKG epitope. Activated macrophages with M2-like characteristics can express multiple enzymes capable of degrading collagen and in turn can internalize and further degrade collagen into small low molecular weight fragments (35, 38). Although little evidence exists that macrophages generate and deposit intact triple helical collagen type I, studies have indicated that certain isoforms of collagen may be expressed (61). Consistent with previous reports, we did not detect intact collagen; however, we did detect low molecular weight RGDKG-containing collagen fragments in both whole cell lysates and serum-free conditioned medium from macrophage-like cell lines. Although our studies do not rule out macrophage-mediated collagen internalization as a contributing factor to the in vivo generation of the RGDKG collagen epitope, our in vitro studies were carried out in the absence of serum or exogenously added collagen, and thus are consistent
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with the active generation of the RGDKGE collagen fragment by macrophages.

Activated macrophages, including M2-polarized macrophages, have been implicated in supporting angiogenesis and inflammation as multiple factors secreted by these cells exhibit pro-angiogenic activities (35, 62). Although many studies indicate that synthetic RGD-containing peptides inhibit angiogenesis and tumor growth, our findings provide the first evidence that macrophages may generate and release an RGDKGE-containing collagen epitope that may exhibit pro-angiogenic activity. Importantly, previous studies have suggested that certain RGD peptides can activate αvβ3 (58, 59) and may enhance vascular permeability (45, 57), which might lead to release of inflammatory factors, which in turn may contribute to the formation of new blood vessels.

To examine possible mechanisms by which the RGDKGE collagen peptide might regulate angiogenesis, we first sought to identify endothelial cell receptors for this motif. Although we do not rule out the possibility that additional non-integrin receptors may bind this collagen epitope, our data suggest that αvβ3 can function as an endothelial cell receptor for the RGDKGE motif. Interestingly, αvβ3 bound both the RGDKGE and RGDAPG collagen peptides, yet only RGDKGE peptide significantly induced angiogenesis and inflammation in vivo. These findings are consistent with the notion that distinct RGD-containing αvβ3 ligands may promote different biological responses. Signaling downstream from αvβ3 is complex, and studies have indicated that simple binding of β3-integrin does not necessarily lead to productive outside-in integrin signaling (63). In fact, the capacity of β3-integrins to promote outside-in signaling depends on multiple factors, including the extent of receptor clustering and subsequent generation of mechanical tension within the actin cytoskeleton, recruitment of adaptor and accessory proteins such as Gax13 and Kindlin-2, and the association of the integrin with protein-tyrosine phosphatases and certain growth factor receptors (64–67). Although the exact mechanisms leading to RGDKGE-mediated αvβ3 signaling is not completely understood, endothelial cell interactions with the RGDKGE peptide in the absence of serum led to enhanced phosphorylation of β3-integrin on tyrosine 747 and of Src phosphorylation at tyrosine 416. These data and others are consistent with an early mechanically mediated activation of β3-integrin that depends on Src given that blocking Src activity reduced β3 phosphorylation following binding to the RGDKGE motif.

Integrin signaling and Src activation are known to regulate the architecture of the actin cytoskeleton (66–68). Moreover, Src family kinases regulate p38 MAPK, and activation of p38 MAPK is thought to enhance actin stress fiber formation in endothelial cells and regulate angiogenesis in vivo (39–45). Our findings provide new insight into the coordinated roles of p38 MAPK and Src in regulating RGD-dependent endothelial cell signaling through αvβ3 as interactions with the RGDKGE cryptic collagen epitope leads to enhanced p38 MAPK phosphorylation in an Src-dependent manner. Moreover, RGDKGE-induced angiogenesis in vivo was associated with enhanced levels of phosphorylated p38 MAPK, and the angiogenic response was reduced by inhibiting p38 MAPK. These findings are consistent with the notion that RGDKGE-stimulated angiogenesis depends on p38 MAPK.

New studies have suggested a role for actin stress fibers and mechanical tension in promoting nuclear accumulation of YAP, where it is thought to function in conjunction with TEAD transcription factors in regulating gene expression (46–49, 52). Given data suggesting a role for YAP in regulating endothelial cell growth and angiogenesis, we examined the subcellular distribution of YAP in endothelial cells following interaction with the RGDKGE collagen peptide. Our data indicate that endothelial cell interaction with the RGDKGE epitope was associated with enhanced nuclear accumulation of YAP. Integrin signaling may play a role in the regulation of YAP as studies have implicated a role for β1-integrins expressed in skeletal stem cells and αv-integrins expressed in osteoblasts in governing YAP subcellular localization (69, 70). Our findings are consistent with a mechanism by which αvβ3-mediated binding to the RGDKGE epitope, but not the related RGDAPG epitope, stimulates a signaling cascade leading to enhanced nuclear accumulation of YAP that depends on Src and/or p38 MAPK. This possibility is supported by our findings that reduced level of nuclear YAP was detected following αvβ3-mediated interaction with RGDKGE peptide in endothelial cells in which Src or p38 MAPK was inhibited. Given the documented role of YAP in governing cell growth coupled with the ability of the RGDKGE collagen peptide to stimulate nuclear accumulation of YAP and enhance endothelial cell growth, it is possible that the RGDKGE collagen peptide-induced endothelial cell growth is YAP-dependent. Consistent with this possibility, no enhancement of endothelial cell growth was detected following knockdown of YAP in endothelial cells stimulated with the RGDKGE collagen peptide, even though these cells are capable of proliferating as stimulation with VEGF or high levels of serum enhanced their growth. Given our studies, it would be interesting to speculate that part of the FGF-2-induced angiogenic response observed in the chick CAM model might involve the recruitment of macrophages that generate a previously uncharacterized RGDKGE-containing cryptic collagen epitope that binds to αvβ3 leading to Src and p38 MAPK activation and nuclear accumulation of YAP. Given that YAP is known to regulate a wide array of genes that may impact angiogenesis and inflammation, including CTGF and Cyr61, it is likely that the RGDKGE collagen epitope may initiate a complex pro-angiogenic program in vivo involving YAP-associated regulation of multiple pro-angiogenic molecules and not simply be restricted to only enhancing endothelial cell growth.

Collectively, our studies provide new evidence that a highly conserved RGDKGE-containing collagen epitope can be generated by a subset of macrophages, and the RGDKGE collagen epitope can stimulate pro-inflammatory and angiogenic activity. Binding of the RGDKGE collagen epitope to β3-integrin can initiate a signaling pathway in endothelial cells leading to activation of Src and p38 MAPK ultimately leading to nuclear accumulation of YAP and enhance cell growth. These novel studies provide new cellular and molecular insight into how an endogenously generated RGD-containing cryptic collagen epitope may promote rather than inhibit angiogenesis. Given the complexity of αvβ3 functions and the growing body of evidence that
the final outcome of αvβ3 binding may depend on the nature of the particular ligand, our findings provide support for an alternative strategy to help control the biological activity of β3-integrin by specific targeting of endogenous pro-angiogenic ligands of αvβ3 rather than direct targeting of the receptor itself.

Author Contributions—J. A. conceived and coordinated studies, designed, performed, and analyzed experiments, and wrote the paper. L. C. coordinated studies and designed, performed, and analyzed experiments. J. M. C. coordinated the studies and designed, performed, and analyzed experiments. E. T. coordinated studies and performed and analyzed experiments. X. Y. coordinated studies, designed, and analyzed experiments. P. C. B. conceived and coordinated the studies, designed, performed, and analyzed the experiments, and wrote the paper. All authors approved the final version of the manuscript.

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