The viral anti-inflammatory chemokine-binding protein
M-T7 reduces intimal hyperplasia after vascular injury

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Chemokines and IFN-γ function as central regulators of inflammatory responses to vascular injury. Both classes of cytokines are upregulated during restenosis, a response to vascular injury that leads to recurrent atherosclerotic plaque growth, but the relative impact of each class of cytokines remains undetermined. M-T7 is a secreted myxoma viral immunomodulatory glycoprotein that functions both as a species-specific inhibitor of rabbit IFN-γ and as a chemokine-binding protein, interacting with a wide range of C, C-C, and C-X-C chemokines in a species-nonspecific fashion. We wished to (a) assess the efficacy of purified M-T7 protein in inhibiting intimal hyperplasia after angioplasty injury and (b) exploit unique species-specific functions of M-T7 in order to judge the relative importance of each cytokine class on plaque growth. Anesthetized New Zealand white rabbits and Sprague-Dawley rats received either M-T7 or control at the time of arterial angioplasty injury. Histological analysis at 28 days demonstrated significant reductions in intimal hyperplasia with M-T7 treatment in both models, with an associated early inhibition of inflammatory cell invasion. Purified M-T7 protein inhibits intimal hyperplasia after angioplasty injury in a species-nonspecific fashion, thus implicating the chemokine-binding activity as more critical for prevention of plaque growth after vascular injury.


Introduction

Chemokines and IFN-γ act as regulators of inflammatory reactions during early atherosclerotic plaque development (1–4). Both of these classes of cytokines have been detected in increased amounts at sites of vascular injury and atherosclerotic plaque development (5–12), but the relative roles of these inflammatory mediators in the accelerated atherogenesis after angioplasty (restenosis) remains undetermined.

IFN-γ is released from activated T cells, natural killer (NK) cells, and endothelial cells, possessing not only antiproliferative activities but also functioning as the major activator of macrophages in inflamed tissue (1–4, 13–18). Previous reports have associated administration of IFN with inhibition of smooth muscle cellular proliferation in tissue culture and reduced plaque growth in a hypercholesterolemic rabbit model (19, 20). IFN-γ treatment of human monocytes was also found to reduce acetylated LDL uptake and to reduce intracellular cholesterol ester accumulation, inhibiting the formation of foam cells in tissue culture (16, 17). Other studies have, conversely, found that treatment with an antibody to IFN-γ reduced plaque growth in a mouse heterotopic cardiac transplant model (21) and that the infusion of IFN-γ induced arteriosclerotic arterial changes in pig- and human-to-immunodeficient SCID mouse xenografts (22). The function of IFN-γ in atheroma development remains therefore unclear, with evidence both for proinflammatory and anti-inflammatory actions in animal models.

Chemokines are small 8–10-kDa proinflammatory proteins that act both as chemoattractants and activators of migrating leukocytes. The C-X-C class of chemokines classically is associated with the recruitment and activation of neutrophils whereas the C-C class of chemokines is targeted more to lymphocytes and macrophages, but the cellular specificities of the chemokine classes have subsequently been found to be less distinct (23–27). Mice deficient in monocyte chemoattractant protein-1 (MCP-1), and which also overexpress apo B, were recently reported to have reduced plaque growth indicating that MCP-1 plays a role in plaque development (28). Knockout mice deficient in the CCR2 chemokine receptor (the receptor for MCP-1 and MCP-3) and crossed with atherogenic apo E–null mice were also demonstrated to exhibit reduced atherosclerotic plaque growth, further substantiating a central role for chemokines in early plaque development (29). Monocyte migration into areas of developing plaque has also been found to be closely associated
with chemokine expression (30–32). Finally, an anti-monocyte chemoattractant protein–1 (anti-MCP-1) antibody has also been reported to inhibit neointimal hyperplasia in injured rat carotid arteries (33).

Accelerated plaque growth after angioplasty, which can lead to restenosis, is characterized by an early inflammatory cell reaction with associated smooth muscle cellular proliferation and later connective tissue remodeling (34, 35). Restenosis is the major limiting factor in the long-term efficacy of percutaneous transluminal coronary balloon angioplasty (36), stent implantation (37), or other intervention therapies for occlusive coronary atherosclerosis (38). The initial complex cascade of biologic events that produces restenosis at previous angioplasty sites includes thrombosis, macrophage invasion, smooth muscle cellular migration, and proliferation and extracellular matrix production (39, 40).

The large DNA viruses, particularly the poxviruses and herpesviruses, have evolved numerous highly effective mechanisms designed to specifically shelter invading viruses from host immune and inflammatory responses that are intended to eradicate infecting viruses (41–44). Myxoma virus is a member of the poxvirus family that induces a highly virulent and lethal systemic disease in the European rabbit (41, 42). The virulence of myxoma virus infection in the mammalian host is mediated by viral immunomodulating factors that often target host inflammatory responses to infection (41–44). There is a direct interaction between viruses and effector cells that modifies the natural and acquired immune responses mediated through T and B lymphocytes, NK cells, monocytes, and macrophages (43, 44). Poxviruses encode a variety of gene products with significant homology to selected cellular cytokine receptors that are postulated to function by disturbing the cytokine network of infected hosts (41–46). M-T7 is a secreted myxoma viral anti-inflammatory protein discovered to possess the dual functions of inhibiting IFN-γ (45) by mimicking the IFN-γ receptor (IFN-γR) (45) and also acting as a chemokine-binding protein (46). The myxoma virus M-T7 protein bears significant homology to the ligand-binding domain of cellular IFN-γRs (45), but the IFN-γ inhibitory activity is species specific, with binding exclusively limited to rabbit IFN-γ (46). In contrast, M-T7 also binds a wide range of human, rabbit, and mouse C, C-C, and C-X-C chemokines through the conserved COOH-terminal heparin-binding domain (46) and has been postulated to function as a species pan-specific disrupter of chemokine gradients necessary for directed migration of inflammatory leukocytes into damaged or diseased tissues (46). Thus, in rabbits, M-T7 protein interacts with both IFN-γ and chemokines, whereas in rat models, M-T7 possesses only the latter chemokine-binding activity.

We have used this differential species specificity of the myxoma M-T7 viral protein for these two contrasting anticytokine activities in order to assess the relative impact of inhibition of IFN-γ and chemokines in animal models of atherosclerotic plaque growth after angioplasty balloon–mediated vascular injury. Plaque size at 28 days after injury, as well as early (24-hour) inflammatory cell invasion and cytokine expression, were assessed after infusion of purified M-T7 protein or controls in both rabbit and rat models of angioplasty balloon–mediated endothelial denudation.

Methods

Animal models. The effect of M-T7 treatment on cellular invasion and intimal hyperplasia was assessed in both rabbit and rat models of balloon angioplasty injury. M-T7 was infused at the time of balloon-mediated arterial injury in 25 New Zealand white rabbits and in 52 Sprague-Dawley rats. The effects of M-T7 or control infusions were assessed by analysis of plaque growth at a 28-day follow-up. An additional 36 rats had early follow-up at 12 to 24 hours after M-T7 or control infusions given at the time of vascular injury.

Rabbit model of balloon injury. Twenty-five New Zealand white rabbits were fed 0.5% cholesterol in 6% peanut oil for 4 days per week, beginning 2 weeks before balloon injury and continuing for 28 days. A 3.0–3.5-mm angioplasty balloon catheter (1:1 diameter ratio of balloon to aorta) was inserted through a femoral arterial cut down and advanced retrograde to the abdominal aorta under general anesthetic (40 mg/kg ketamine (Veterepharm Canada Inc., London, Ontario, Canada), 8 mg/kg xylazine, and 0.5 mg/kg acepromazine by intramuscular injection). The balloon was inflated to 8 atmospheres (atm) of pressure in the distal abdominal aorta, advanced to the level of the diaphragm, and withdrawn three times, under fluoroscopic guidance (47). Heparin (400 units, Leo Laboratories Canada Ltd., Ax Ontario, Canada) was given immediately after obtaining femoral access. M-T7 was infused via a 3.0–3.5-mm diameter Wolinsky perforated balloon catheter (at an inflation pressure of 6 atm for 2 minutes) positioned in the abdominal aorta approximately 0.5 cm above the iliac bifurcation. M-T7 was infused at doses ranging from 50–5,000 pg as follows: saline (n = 6), M-T7 50 pg (0.017 pg/g; n = 6), M-T7 500 pg (0.17 pg/g; n = 6), M-T7 5,000 pg (1.7 pg/g; n = 7). Each infusate was administered as a diluted infusion in 0.9% saline, total volume of 10 mL, immediately after balloon-mediated injury. To assess the effects of M-T7 on early cellular invasion, four rabbits had infusion of saline (n = 2) or 300 ng M-T7 (n = 2) immediately after angioplasty injury and were sacrificed 24 hours later for immunohistochemical analysis.

Rat model of balloon injury of the iliofemoral artery. To assess the effects of M-T7 on plaque growth after balloon injury and early inflammatory cell invasion, rats were divided into four groups, receiving either purified M-T7, heparin, a heparin/M-T7 mixture, or saline. Fifty-two male Sprague-Dawley rats (Charles River Laboratories, Wilmington, Massachusetts, USA) weighing 250–300 g were anesthetized with ketamine hydrochloride and Rompun 2:1 combination, 1 mL/100 g weight intramuscular injection (Veterepharm Canada Inc.). A 1.5-mm diameter, 20-mm long angioplasty balloon catheter

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(Europass, Cordis Corp., Miami, Florida, USA) was introduced into the femoral artery using a cut-down procedure. All infusions (1.0 mL volume of either M-T7 or controls diluted in 0.9% saline) were given immediately after balloon inflation through the distal lumen of the angioplasty balloon catheter into the arterial circulation, as described previously (48). The balloon was then inflated with saline to the point at which resistance to withdrawal was detected (6 atm). The inflated balloon was then advanced and withdrawn from the iliac bifurcation to the distal femoral artery (4–5 mm) three times. The infusion doses were as follows: saline (0 pg M-T7; \( n = 16 \)), M-T7 5 pg (0.017 pg/g; \( n = 6 \)), M-T7 50 pg (0.17 pg/g; \( n = 6 \)), M-T7 500 pg (1.7 pg/g; \( n = 6 \)), heparin 50 U (\( n = 12 \)), heparin 50 U mixed with M-T7 500 pg (1.7 pg/g; \( n = 6 \)). Heparin and the heparin/M-T7 mixtures were then intravenously injected to assess the effects of heparin alone on plaque development and to assess potential interference by heparin on binding of M-T7 to the heparin-binding domain of chemokines. After the infusion, the catheter was removed and the femoral artery was sealed with local application of n-butyl cyanoacrylate monomer (Nexaband; Veterinary Products Laboratories, Phoenix, Arizona, USA).

To assess the early effects of M-T7 on inflammatory cell invasion, 36 rats had angioplasty injury of the iliofemoral artery (as just described) with immediate infusion of either saline control or 500 pg M-T7 under general anesthesia. Each group of rats was sacrificed at selected time intervals after balloon-mediated vascular injury and M-T7 or control infusion: at 0 hours (\( n = 8 \) for saline, \( n = 6 \) for M-T7), at 4 hours (\( n = 8 \) for saline, \( n = 4 \) for M-T7), and at 24 hours (\( n = 6 \) for saline, \( n = 4 \) for M-T7).

**M-T7 source and purification.** M-T7 was isolated from baby green monkey kidney (BGMK) cells infected with a vaccinia vector expressing M-T7 and purified to apparent homogeneity by a two-step procedure using fast-performance liquid chromatography (FPLC), described in detail elsewhere (45, 49, 50). Briefly, concentrated supernatants harvested from myxoma virus–infected BGMK cells were fractionated by anion-exchange chromatography using a MonoQ HR5/5 column (Amersham Pharmacia Biotech Inc., Baie d’Urfé, Québec, Canada) and NaCl containing 20 mM bis-Tris (pH 6) elution buffer. MonoQ fractions containing M-T7 were pooled, dialyzed against PBS overnight, and further purified by gel-filtration chromatography using a HiLoad Superdex 200 column (Amersham Pharmacia Biotech Inc.). Eluted proteins were subjected to SDS-12% PAGE and visualized by silver staining and immunoblotting to assess the purity and composition of the purified proteins (45, 46, 50). Fractions containing highly purified M-T7, were pooled, concentrated, and filter sterilized before they were injected.

**Histological and morphometric analysis.** At the designated follow-up times, study animals were sacrificed with Euthanyl (Bimda-MTC Animal Health Co., Cambridge, Ontario, Canada), and the arteries with previous angioplasty balloon–mediated injury were isolated for histological and morphometric analysis as described previously (47–49). Arterial sections 2.5 cm long from the rabbit aorta (0.5–3.0 cm) proximal to the iliac bifurcation and from the rat iliac artery bifurcation to the femoral arterial branches were harvested, divided into three sections (0.8–1.0 cm long), and stained with hematoxylin and eosin for light microscopic and morphometric examination. Isolated arterial specimens were fixed in 10% sodium phosphate–buffered formalin, processed, impregnated, embedded in paraffin, and cut into 5-μm sections by microtome, as described previously (47–49). Two sections were cut and stained from each of the three arterial sections from each animal specimen; a total of six sections were thus examined for each arterial specimen.
Total intimal plaque area and the area of invading mononuclear cells were measured by morphometric analysis. Intimal plaque area was quantified using the section with the largest detectable area of atherosclerotic plaque by means of the Empix Northern Eclipse trace application program (Empix Imaging Inc., Mississauga, Ontario, Canada) using a Sony Power HAD 3CCD color video camera attached to the microscope and calibrated to the microscopic objective used. The mean total cross-sectional areas of the intima and the areas of inflammatory cell infiltration were then calculated for arterial specimens taken from the site of angioplasty balloon injury in each animal.

**Immunohistochemical staining.** For immunohistochemical analysis, formalin-fixed samples of arterial sections taken from the rat models (early and late follow-up) were embedded in paraffin blocks. The top and midarterial sections were used for immunohistological staining. After fixation, dehydration, and rehydration steps, the specimens were rinsed in PBS for 5 minutes. Tissue sections (5 μm thickness) were blocked with 20% normal horse serum for 10 minutes, then incubated with primary antibody for 1 hour to overnight, as determined previously (49), after which specimens were again rinsed in buffer. Sections were then incubated with primary antibodies specific to each cell type and selected cytokines. In all immunohistochemical staining, bound antibodies were detected using the indirect peroxidase-labeled antibody system (Vector Laboratories, Burlington, Ontario, Canada) as described previously (49). The primary antibody for arterial smooth muscle cells was mouse monoclonal IgG2 anti-alpha smooth muscle actin diluted 1:400 (Sigma Chemical Co., St. Louis, Missouri, USA). For macrophage staining, mouse monoclonal IgG anti-rat macrophage antibody (specific to ED2-like antigen on rat macrophage, not monocyte) at 1:100 dilution was used as the primary antibody (PharMingen, Mississauga, Ontario, Canada). For NK and maturing T-lymphocyte staining, mouse monoclonal anti-rat CD2 (IgG) thymocyte and T-lymphocyte antibody diluted 1:100 (Cedarlane Laboratories Ltd., Hornby, Ontario, Canada) was used as primary antibody (capable of recognizing maturing T lymphocytes and NK cells). For B-cell staining, mouse IgG anti-rat CD45RA (antibody to a subfraction of 240-kDa rat CD45 specific to B lymphocytes) diluted 1:200 (Cedarlane Laboratories Ltd.) was used for the primary antibody (pan B cell). The primary antibody used for polymorphonuclear leukocytes (granulocytes) staining was mouse anti-rat IgM (1:100 dilution) with biotin-conjugated goat anti-mouse (rat adsorbed) Ig-specific polyclonal antibody (PharMingen). For the rabbit model, mouse RAM-11 anti-rabbit macrophage antibody (Cedarlane Laboratories Ltd.) was used for primary antibody for macrophage recognition. The following primary antibodies were used to assess chemokines present in the tissue specimens: rabbit anti-rat IgG for MCP-1, MIP-1α (macrophage inflammatory protein-1α), and RANTES (Cedarlane Laboratories Ltd.).

Treatment with the primary antibody was followed by incubation with biotinylated anti-mouse IgG diluted 1:250 (Vector Laboratories) for 20 minutes (for mouse primary antibodies) and goat anti-rabbit IgG (for rabbit primary antibodies) and avidin-biotin-peroxidase complex for 40 minutes. After rinsing, specimens were counterstained with 3’3’-diaminobenzidine tetrahydrochloride (DAB) for 5 minutes. All specimens were then rinsed with distilled water, dehydrated, and clear mounted in cytoseal (Stephens Scientific Division of Cornwell Corp., Riverdale, New Jersey, USA) mounting medium for histologic assessment.
The number of positively staining cells for the cellular analysis or the area staining positive for chemokines or cytokines was measured in three high-power fields, selected as having the largest area of positively stained cells or cytokine staining, from each slide and divided by the area of medial or adventitial layers of the artery wall examined. The final value for each arterial section was determined by the number of positively stained cells, or the area staining positive for chemokines and cytokines, and then divided by the area assessed for each of the three readings. A mean value was then calculated for each study animal. A negative control lacking primary antibody was present on each slide. Control stains lacking the secondary antibodies or using irrelevant primary antibody (IgG1 antibody to the cytoplasmic tail of vesicular stomatitis virus G protein from T. kresi) were performed for each new antibody examined.

Statistical analysis. The mean values for plaque area or positively stained cells and chemokines were measured for all arterial specimens for each study animal. Correlations between the ratio of positively stained cells and measured plaque area with control or M-T7 treatment were determined by ANOVA and unpaired student’s t test. The results are summarized as mean value plus or minus SE, and significance was defined at P less than 0.05.

Results

Effect of M-T7 protein administration on neointima formation after vascular injury. In the control saline-treated rabbits, the 0.5% cholesterol/peanut oil diet, together with aortic injury, induced lipid-rich areas of plaque with increased macrophage invasion at the 28-day follow-up (Figure 1a). In contrast, at 28 days after balloon injury in rats fed a normal diet the plaque that developed in the iliofemoral artery was relatively rich in smooth muscle cells (Figure 1b). In both models, the foam cell rich–plaque in the cholesterol-fed balloon-injured rabbits (Figure 1c; P < 0.0108) and the smooth muscle cell–rich intimal hyperplasia in the rat model (Figure 1d; P < 0.0136), there was a significant reduction in plaque growth with a single injection of M-T7 at the time of vascular balloon injury (Figures 1 and 2).

The reduction in intimal hyperplasia detected in both New Zealand white rabbits (Figure 2a) and Sprague-Dawley rats (Figure 2b) at the 28-day follow-up was comparable for both models at equivalent infusion doses. Relative to saline-treated control animals (rabbits or rats), an intra-arterial infusion of 0.017 pg/g or greater per rabbit (50 pg or greater total dose) and 0.017 pg/g or greater per rat (5 pg or greater total dose) given immediately after balloon angioplasty, produced the greatest reductions in neointima formation at the follow-up 28 days after M-T7 infusion. In the rabbit model, a relative increase in plaque area was noted at higher doses at 28 days after balloon injury, but this increase in intimal plaque size at higher infusion doses did not reach significance (P = 0.6504).
M-T7–treated animals, in the rat or the rabbit model, had reduced mononuclear inflammatory cell invasion into both the intimal and the adventitial layers on comparison with saline-treated controls at the 28-day follow-up (Figures 2 and 3). The reduction in macrophage and smooth muscle cellular invasion in the intimal and adventitial layers was dose dependent (Figure 2c), with the greatest reduction seen at the two highest M-T7 infusion doses. Macrophage invasion was significantly reduced \( (P < 0.0011) \) even at the lowest dose of M-T7 (Figure 2c), whereas the smooth muscle cellular invasion was only significantly reduced at the highest M-T7 infusion dose \( (P < 0.0001) \) in the intimal layer (Figure 2d), indicating a dose-dependent effect of M-T7 on arterial cellular responses to vascular injury. Histological analysis and the cross-sectional area of the media did not differ significantly between the two groups after treatment with M-T7.

Immunohistochemical evaluation of cellular invasion. To evaluate the effects of M-T7 on cellular responses to vascular injury, sections of arteries were assessed by immunohistochemical analysis of individual cells, macrophages, smooth muscle cells, and CD-2–positive T lymphocytes known to play a role in early and late atherosclerotic plaque development. The rat model was used initially in this study because of the known rabbit species–specific IFN-\( \gamma \) binding activity of M-T7. The rat model was believed to be capable of reacting only to the species-nonspecific chemokine-binding activity of M-T7, which would allow the rat model to provide information on the specific effects of chemokine binding in early vascular responses to injury. The number of invading cells per high-power field in the intimal, medial, and adventitial layers was measured.

Invading macrophages were seen in arterial sections taken at the 28-day follow-up in the saline-treated rat model in all three arterial layers. The largest areas of macrophage invasion were seen in the intimal and the adventitial layers, with the greatest numbers of invading cells found in the adventitial layers. At 28 days' follow-up, these areas of macrophage invasion into the adventitial layer were markedly reduced after M-T7 infusion \( (P \leq 0.0016) \) (Figure 3a). A corresponding early reduction in macrophage invasion \( (P \leq 0.0016) \) was detected in the adventitial layer of the arterial wall at 24 hours after balloon angioplasty in rats treated with M-T7 (Figure 3d) on comparison with control saline-treated animals (Figure 3c). There was no significant decrease in the numbers of invading macrophages in the adventitial layers at earlier times (at 0–4 hours) after injury with M-T7 treatment (Figure 3e). Very few macrophages were detected in the medial arterial wall layers after angioplasty injury so that any potential effects of M-T7 on macrophage invasion into the media could not be accurately assessed. At 0–24 hours there was no intimal invasion; therefore, only data for cellular invasion from the medial and adventitial layers is discussed.

A trend toward a reduction in smooth muscle cell invasion was seen in the intimal and adventitial layers at the 28-day follow-up after M-T7 infusion (Figure 3f) on comparison with control saline-treated rat arterial
specimens, but this did not reach significance ($P = 0.4659$). Very few smooth muscle cells were detected migrating into the adventitial layer at early time points after balloon injury (0–24 hours after angioplasty injury) in these studies, thus preventing accurate assessment of effects of M-T7 on smooth muscle cellular invasion. Smooth muscle cellular count in the media was, however, unaffected by M-T7 treatment at early follow-up times (0–24 hours; $P = 0.3994$), suggesting that no significant migration of smooth muscle cells out of the media was occurring at these early times after vessel injury.

Both CD2-positive T-lymphocyte and B-lymphocyte invasion were detected in very minimal amounts at early and late follow-up after balloon angioplasty injury. Neutrophils were not detected in these sections. In the adventitial layer the B cell and CD2-positive T-cell counts on immunohistochemical analysis were reduced at 28 days after M-T7 infusion (data not shown), but this reduction did not reach significance ($P = 0.0966$). The detected B-cell and CD2-positive T-cell numbers were, however, very small, even in the saline control animals, and analysis of effects of M-T7 on these cells was therefore not considered to be accurate.

For comparison, sections taken from the rabbit model at the 24-hour follow-up were also assessed for the effects of M-T7 infusion on macrophage and smooth muscle cellular invasion. The rabbit aortic histological sections had a similar decrease in the numbers of invading macrophage cells at 24 hours' follow-up after treatment with M-T7. Both the medial and adventitial layers had reductions in cells staining positive for RAM-11 (macrophages). In the abdominal aortic sections the B cell and CD2-positive T-cell counts on immunohistochemical analysis were reduced at 24 hours after M-T7 treatment (data not shown), but this reduction did not reach significance ($P = 0.007$). As noted previously, M-T7 treatment reduced plaque size significantly when compared with saline or heparin infusions (M-T7, $0.006 \pm 0.004$; M-T7 + heparin, $0.018 \pm 0.007$; heparin, $0.025 \pm 0.007$; saline, $0.037 \pm 0.007$). As noted previously, M-T7 treatment reduced plaque growth significantly when compared with saline infusion ($P \leq 0.0209$), but the reduction in plaque size after treatment with M-T7 combined with heparin ($P = 0.1319$) or heparin alone ($P = 0.2813$) was not large enough to reach significance. The presence of heparin thus did interfere with M-T7 antiatherogenic activity in the rat model, although a nonsignificant trend toward a reduction in intimal area was detected on comparison with heparin or saline treatments. M-T7 reduced plaque growth to the same degree as when infused alone. This may be due to an independent activity of heparin or due to competition of heparin with M-T7 for chemokine binding.

**Discussion**

We have demonstrated that the myxoma virus protein M-T7, with the dual functions of species-specific IFN-γ and species-nonspecific chemokine-binding activity, inhibits plaque development with similar efficacy in both rabbit and rat models of angioplasty-mediated vascular injury and with similar ranges of effective dose response. We postulate that the broad-based species-nonspecific chemokine-binding activity of M-T7 is responsible for the reduction in plaque growth in the rat and the rabbit models. The IFN-γ-R homologue activity is rabbit species specific and hence cannot function in the rat model. This IFN-γ-binding activity would also be predicted to increase plaque growth based upon the previous observations of reduced plaque growth seen in cholesterol-fed rabbits treated with interferon (19, 20). The reduction of plaque in both models after a single infusion of M-T7 immediately after vascular injury therefore suggests the following: (a) that chemokines play a critical early role in activation of the inflammatory response in vascular
wound healing and (b) the extent of M-T7 antiathero-
ma activity measured cannot be attributed to the inhi-
bition of IFN-γ in this rat model.

The attraction of inflammatory cells to damaged or infected tissues is essential for the host responses to injury and infection. The inflammatory cell response process is controlled by chemokines, chemotactic cytokines (22–26), peptides that are also believed to have central roles in early events leading to plaque growth at sites of vascular injury (27–29). The importance of chemokines in the pathophysiologic process of disease is now understood, but the role of chemokines in recurrent plaque growth after angioplasty is still being defined. Further, the relative effects of chemokine and IFN-γ inhibition on later intimal hyperplasia produced by angioplasty injury remains to be determined. We have tested for local early arterial expression of three C-C chemokines, specifically MIP-1α, MCP-1, and RANTES, as well as rat IFN-γ, in the rat iliofemoral arterial injury model. The rat model alone was tested to assess the effects of the isolated chemokine-binding activity of M-T7 on early expression of chemokines after vascular balloon-mediated injury. IFN-γ expression, although not predicted to be altered by M-T7 treatment in the rat model of balloon injury, was also tested as a control. We detected a significant reduction in chemokine antigens (Figure 4, c–e), but not IFN-γ (Figure 4f), using immuno-
histochemical analysis of arterial specimens taken at 0, 4, and 24 hours after angioplasty. This reduction in chemokine staining may be the result of a primary alteration in local tissue levels of chemokines by M-T7. The reduction in chemokine staining at early times after balloon injury in the presence of M-T7 may also be the product of an inhibition of chemokine mRNA or protein expression, or due to interference with antibody binding to chemokines already present in the vascular wall during immunohistochemical staining, or poten-
tially even due to a downstream effect of an as yet unknown function of M-T7 on the inflammatory sys-
tem response. The fact that IFN-γ antigen expression was not altered after M-T7 treatment in rats, however, would suggest that this observed reduction in plaque growth is not due to a generalized downregulation of inflammatory cytokine responses. It must be noted, however, that the amount of IFN-γ detected was small as seen by immunohistochemical analysis and thus accurate detection of a reduction of IFN-γ by this method is difficult and should be investigated further. Future experiments, perhaps with M-T7 variants lacking chemokine-binding activity, will need to be done to assess the definitive mechanism(s) of action of M-T7 in the prevention of injury-induced plaque growth. This study does, however, demonstrate that the protective antiatherogenic state after arterial trauma in these two models of angioplasty injury by M-T7 cannot be explained by the species-specific inhibition of IFN-γ.

C-X-C chemokines induce neutrophil, but not mono-
cyte, migration, whereas C-C chemokines act predomin-
antley on monocytes and macrophages with no activity on neutrophils (22, 23, 25, 26). Both the C-X-C and C-C family members act on a number of cell types, including basophils, eosinophils, lymphocytes, endothelial cells, epithelial cells, and SMCs. They have also been shown to induce human leukocyte migration both in vitro and in vivo (25, 26, 29–32). Lalani et al. (46) have tested the ability of M-T7 to bind to a variety of human and mouse chemokines using a gel-shift assay. It was found that M-T7 binds to all members of the C-C subfamily (RANTES, MIP-1α, MCP-1, MCP-3), the C-X-C subfamily (IL-8, platelet factor 4), and the C subfamily (mouse lympho-
taxin) (45). In sections taken at early times after angio-
plasty in the rat angioplasty injury study (0 hour, 4 hours, 24 hours) M-T7 inhibited invasion of macrophages, but not T lymphocytes nor smooth muscle cells. This early inhibitory activity is consistent with an inhibition of chemokine-directed cellular migration, but may be due to either direct chemokine binding or to altered regula-
tion of chemokine expression as noted above.

The rat model of vascular balloon injury is charac-
terized by a predominant smooth muscle cellular response with less-pronounced fatty plaque areas or macrophage/foam cell invasion at sites of damage. The cholesterol-fed rabbit model of balloon injury has a more pronounced inflammatory cell component with areas of lipid-filled macrophage invasion. In both cases, however, significant reductions in plaque area were detected with M-T7 infusion. Based upon previous work demonstrating a reduction in plaque growth with IFN treatment in cholesterol-fed rabbits (19), the fact that M-T7 protein known to bind IFN-γ and chemokines reduced plaque also suggests a pre-
dominant effect from chemokine binding in the inhibi-
tion of plaque in this rabbit model of balloon injury. This also suggests that M-T7 is acting at an early initiating step in atherogenesis and that this step is common to both smooth muscle and macrophage cellular responses and invasion and later proliferation at sites of vessel wound healing. These findings also support chemokine activity as a central mediator in the initiation of plaque growth after angioplasty.

In conclusion, we have demonstrated that M-T7, an IFNγ-R homologue and chemokine-binding protein, inhibits plaque growth and inflammatory cell invasion into the vascular wall after balloon injury in both rat and rabbit models. Our study suggests that chemokine activity plays a greater role than IFN-γ activity in early atherogenesis after vascular injury.

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