Oncostatin M (OSM) is a multifunctional cytokine, a member of the interleukin-6/leukemia inhibitory factor (IL-6/LIF) family, that can regulate a number of connective-tissue cell types in vitro including cartilage and synovial tissue-derived fibroblasts, however its role in joint inflammation in vivo is not clear. We have analyzed murine OSM (muOSM) activity in vitro and in vivo in mouse joint tissue, to determine the potential role of this cytokine in local joint inflammation and pathology. The effects of muOSM and other IL-6/LIF cytokines on mouse synovial fibroblast cultures were assessed in vitro and showed induction of monocyte chemotactic protein-1, interleukin-6, and tissue inhibitor metalloproteinase-1, as well as enhancement of colony growth in soft agarose culture. Other IL-6/LIF cytokines including IL-6, LIF, or crdiotrophin-1, did not have such effects when tested at relatively high concentrations (20 ng/ml). To assess effects of muOSM in articular joints in vivo, we used recombinant adenovirus expressing muOSM cDNA (AdmuOSM) and injected purified recombinant virus (10^6 to 10^8 pfu) intra-articularly into the knees of various mouse strains. Histological analysis revealed dramatic alterations in the synovium but not in synovium of knees treated with the control virus Ad-dl70 or knees treated with Ad-muIL-6 encoding biologically active murine IL-6. AdmuOSM effects were characterized by increases in the synovial cell proliferation, infiltration of mononuclear cells, and increases in extracellular matrix deposition that were evident at day 4, but much more marked at days 7, 14, and 21 after administration. The synovium took on characteristics similar to pannus and appeared to contact and invade cartilage. Collectively, these results provide good evidence that OSM regulates synovial fibroblast function differently than other IL-6-type cytokines, and can induce a proliferative invasive phenotype of synovium in vitro in mice on overexpression. We suggest that OSM may contribute to pathology in arthritis. (Am J Pathol 2000, 157:1187–1196)

The process of inflammatory arthritis involves a number of different cell types including resident connective tissue cells of the joint such as synoviocytes, endothelial cells, and chondrocytes, and their interaction with infiltrating cells including macrophages/monocytes, T lymphocytes, B cells, as well as polymorphonuclear cells. Such interactions are mediated in large part by cytokines and particular cytokines secreted by monocyte/macrophages, ie, interleukin-1 (IL-1) and tumor necrosis factor (TNF), are thought to drive effector functions of the local connective tissue cells.1–4 Such effects include activation of synoviocyte proliferation and secretion of matrix metalloproteinases as well as other cytokines and chemokines, and activation of chondrocyte metabolism resulting in cartilage breakdown.1,4 However, other cytokines likely participate in this process and may in fact dominate certain aspects of these complex chronic inflammatory processes.

The 28-kd cytokine oncostatin M (OSM) is another product of macrophages and of activated T cells,5,6 and is found at enhanced levels in synovial fluids of rheumatoid arthritis patients.7–9 OSM can affect a number of cell types including hepatocytes,10 vascular smooth muscle cells,11 Kaposi’s sarcoma cells,12,13 as well as melanoma and carcinoma cell lines.5 Human OSM induces responses by connective tissue cells such as human synovial fibroblasts,7,14–17 chondrocytes,18,19 and osteoblasts,20 in addition to endothelial cells21,22 and epithelial cells.23,24 The regulation of TIMP-1, MMP-1, as well as monocyte chemotactic protein-1 (MCP-1) by OSM in synovial fibroblasts in vitro7 suggest an important role for this cytokine in chronic joint inflammation. Human OSM binds and activates human receptors characterized as type I complex, which is identical to the leukemia inhibi-

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**Murine Oncostatin M Stimulates Mouse Synovial Fibroblasts in Vitro and Induces Inflammation and Destruction in Mouse Joints in Vivo**

Carrie Langdon,* Christine Kerr,* Mohammed Hassen,* Takahiko Hara,† A. Larry Arsenault,* and Carl D. Richards*
tory factor (LIF) receptor, and type II complex (specific for OSM)\textsuperscript{25} and induces intracellular signaling pathways involving JAK-1, JAK-2, and TYK-2 kinases and activation of the STAT transcriptional activator proteins STAT-1 and STAT-3.\textsuperscript{26–28} However, the precise role that OSM plays in chronic inflammation, particularly in comparison to other members of its family of cytokines, and in context of other proinflammatory cytokines, is still not clear.

To determine the effects of OSM \textit{in vivo}, the use of animal models is necessary. The murine analogue of OSM has been recently identified\textsuperscript{29} and is expressed at high levels in mouse bone marrow and at lower levels in thymus and spleen. Mouse OSM can regulate proliferation of a number of different cell types \textit{in vitro},\textsuperscript{30} supports expansion of multipotential hematopoietic progenitors\textsuperscript{31} and Sertoli cells,\textsuperscript{32} and enhances hepatocyte differentiation.\textsuperscript{33} Murine OSM (muOSM) binds its own specific receptor but, unlike human OSM, does not bind the LIF receptor.\textsuperscript{30,34,35} Furthermore, it is clear that human OSM cannot interact with the mouse OSM receptor to induce responses.\textsuperscript{30,36} We have previously shown that muOSM can stimulate TIMP-1 mRNA expression in mouse and rat fibroblast cell lines, whereas human OSM cannot.\textsuperscript{36} In this investigation of mouse OSM and its regulation of connective tissue cells of joints, we have used recombinant cytokines to study \textit{in vitro} responses of synovial fibroblasts derived from mouse synovium, and have used an adenovirus vector system to overexpress OSM \textit{in vivo} in mouse joint tissues on local intra-articular administration. Our results suggest that muOSM is a potent stimulator of mouse synovial fibroblast responses \textit{in vitro}, and that overexpression of OSM results in dramatic phenotypic changes in the synovium that are consistent with synovial cell proliferation, matrix remodeling, and infiltration of mononuclear cells, and thus results in major physiological alterations to the tissue.

**Materials and Methods**

**Reagents**

Recombinant mouse cytokines OSM, IL-6, LIF, and IL-1β (derived from \textit{Escherichia coli}) were purchased from R&D Systems (Minneapolis, MN). Recombinant mCT-1, expressed in mammalian cells and purified as a fusion protein\textsuperscript{37} was obtained from D. Pennica (Genentech, South San Francisco, CA). Enzyme-linked immunosorbent assay (ELISA) kits for mouse MCP-1 (MCAF/JE) were purchased from Biosource (Camarillo, CA).

**Cells**

Primary mouse synovial fibroblasts were derived from explants of finely minced synovial tissue from normal C57BL/6 mice, 10 to 12 weeks old (Charles River Laboratories, Ottawa, Canada), and passaged in Dulbecco’s modified minimal essential medium (F-15) media. Two separate primary cell cultures of fibroblasts, each derived from pooled mouse synovium (10 animals/preparation) have been analyzed. These were passaged several times before use, show a typical synovial fibroblast morphology, and tested positive for VCAM-1 expression (by immunohistochemistry). Primary rat fibroblasts were derived from explants of Sprague-Dawley rat synovium as above and cultured in Dulbecco’s modified minimal essential medium. All fibroblast cell lines were supplemented with 10% fetal bovine serum, 1% penicillin/streptomycin, and 1.5 µg/ml Fungizone (Bristol-Myers Squibb Canada, Montreal, PQ).

**Assays for IL-6 and mMCP-1 Production**

Cultures of synovial fibroblasts were plated in 24-well Corning plates (Cambridge, MA). Fibroblasts were allowed to recover for 24 hours, and were then refed with medium containing 2% serum and stimulated, in triplicate, with the indicated cytokines. Supernatants were collected after 24 hours and stored at −20°C until analysis. Statistical analysis was completed with one-way analysis of variance using pairwise multiple comparison procedures (Fisher least significant difference method) provided by Sigmastat software.

IL-6 content of supernatants was assayed using a modified B9 hybridoma proliferation assay.\textsuperscript{38} The IL-6 dependent B9 cells were plated into 96-well microtiter plates at 2.5 × 10\textsuperscript{3} cells/well in 25 µl of RPMI containing 5% fetal calf serum, 1% penicillin/streptomycin, and 2-mercaptoethanol. Seventy-five µl of various dilutions of test sample supernatant was added. Cells were incubated for 3 days at 37°C, 5% CO\textsubscript{2}, and 100% humidity. B9 cellular proliferation was measured using the 3[4,5-dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide (MTT) colorimetric assay as described by Mossman et al.\textsuperscript{38} Briefly, 10 µl of 5 mg/ml MTT (Sigma, St Louis, MO) was added to each of the wells and the cells were incubated for 4 hours at 37°C. Fifty µl of 10% Triton X-100 and 0.5 mol/L of HCl were added to each well to lyse the cells. Plates were incubated overnight in the dark and then absorbances read at 540 nm using a Labsystems Multiskan MCC/340 ELISA plate reader (Labsystems, Helsinki, Finland). IL-6 activity was compared to a standard preparation of human recombinant IL-6 with the limit of detection being 10 pg/ml. Anti-mouse IL-6 (R&D Systems) completely inhibited IL-6 activity of mouse synovial fibroblast (MSF) supernatants whereas a control antibody had no effect. Cell supernatants were analyzed for mMCP-1 content using a commercial ELISA kit (Biosource Cyto-screen, Camarillo, CA) according to the manufacturer’s instructions. The limit of detection was 9 pg/ml.

**Proliferation in Anchorage-Independent Conditions**

Growth in anchorage-independent conditions was assessed by suspending single cell suspensions of synovial fibroblasts in 0.4% agarose in Dulbecco’s modified minimal essential medium containing 20% fetal bovine serum at a concentration of 1 × 10\textsuperscript{3} cells/ml in 24-well plates. The usage of mouse serum in place of fetal bovine serum did not alter results. Cytokines were used at a
concentration of 10 ng/ml, and all treatments were performed in triplicate or quadruplicate. Plates were incubated at 37°C, 100% humidity for 14 to 17 days before being optically graded for colony growth and size. Our studies show that the diameter of our cell/colony is correlated with cell number, in that a colony of >20 μm has a cell number of ~10 or greater. Recently, published work by Imamura et al\textsuperscript{40} defines a human synoviocyte colony as five cells or greater.

**RNA Isolation and Analysis**

Subconfluent fibroblast cultures were stimulated with the indicated cytokines in medium containing 2% serum and incubated for 18 to 24 hours. RNA was isolated from cultures by the method of Chomczynski and Sacchi.\textsuperscript{41} Cells were lysed in guanidium isothiocyanate. RNA was extracted with a 1:1 ratio of phenol-chloroform and then precipitated overnight in isopropanol. RNA was washed in 70% ethanol then dried under vacuum. Northern analysis was performed by running 5 to 15 μg of total RNA on a 1.2% agarose formaldehyde gel. After transfer to nylon membrane (ICN, Biotrans, NY), blots were probed with \textsuperscript{32}P-labeled antisense oligonucleotides corresponding to bp 164 to 191 of the mouse TIMP-1 cDNA sequence (5’-CTTATAACGCTGGTATAAGGTGGTCTCG-3′) (Mobix, McMaster University) and bp 213 to 238 of the mouse MCP-1 sequence\textsuperscript{42} (5’-TGATCCTAATGAGTAGGCTGGAGAGC-3′) (Mobix, McMaster University). Regulation of mTIMP-3, mIL-6 and β-actin mRNA was examined using \textsuperscript{32}P-labeled cDNA fragments to mTIMP-3, rat IL-6, and human β-actin, respectively. RNA levels for NIH 3T3 fibroblasts, mouse synovial fibroblasts, and lung tissue from C57BL/6 mice injected with recombinant muOSM were analyzed by densitometry and normalized to β-actin.

**β-Galactosidase (lac Z) Expression**

Mouse synovial fibroblasts were infected with recombinant adenovirus expressing muOSM (AdmuOSM) and AdmCMVlacZ at multiplicity of infection of 10 and 50. After 24 hours, the cell supernatant was removed and the cells were fixed in 0.5% glutaraldehyde for 10 minutes at room temperature. The fixative was aspirated, and then staining solution containing 50 mmol/L phosphate buffer, pH 7.4, 5 mmol/L K₃Fe(CN)₆, 5 mmol/L K₄Fe₄(CN)₆, 2 mmol/L MgCl₂, 0.05% Triton X-100 (BioRad, Hercules, CA) and 0.5 mg/ml of 5-bromo-4-chloro-3-indoyl-β-D-galactopyranoside (X-gal) (Boehringer Mannheim, Mannheim, Germany) diluted from a 20-mg/ml stock solution of X-gal dissolved in dimethylformamide, was added. Color was allowed to develop for 45 minutes. The stained fibroblasts were visualized under a microscope and photographs were taken using a Zeiss inverted microscope with attached camera (Zeiss, Jena, Germany).

**Results**

**muOSM Regulation of IL-6 and MCP-1 Expression**

Initial experiments sought to establish the effects of mouse OSM on mouse synovial fibroblasts. To assess the regulation of IL-6 and MCP-1 production by muOSM, MSF were stimulated with muOSM, other gp130 cytokines, or mIL-1β for 24 hours. IL-6 production was studied using the B9 hybridoma proliferation assay whereas the production of MCP-1 protein was examined by ELISA. In studies of MSF in cell culture and anchorage-independent growth, two separate cell lines (each from pooled knee synovium of 10 mice) were analyzed. Both cell lines responded with identical trends in each of three separate experiments per cell line. Data shown represents one of three experiments. Stimulation with muOSM resulted in a dose-dependent increase in IL-6 protein detected in supernatants. Marked increases were noted at 1 ng/ml (50 pmol/L) of muOSM and maximal levels were reached at
20 ng/ml (1 nmol/L) of stimulus, resulting in >300-fold increases over control (Figure 1A). Mouse CT-1 (1 nmol/L) or LIF (1 nmol/L) were ineffective when tested at 20 ng/ml (Figure 1B). mIL-1 induced 100-fold increases in IL-6 production whereas co-stimulation with muOSM resulted in a synergistic increase in IL-6 of >800 times that of control levels. OSM had a modest influence on the amount of mMCP-1 protein detected in the same supernatants of mouse synovial fibroblasts, which showed very high constitutive levels of mMCP-1 protein (3 ng/ml) (Figure 1C) and express mMCP-1 mRNA constitutively (Figure 2). Concentrations of muOSM (10 to 50 ng/ml) that were able to stimulate >300-fold increases in IL-6, induced increases in MCP-1 protein of approximately twofold (statistically significant), whereas mCT-1, mLIF, or mL-1β had less effect (Figure 1D). mL-1β also enhanced the levels of mMCP-1 protein by twofold.

To assess levels of IL-6, MCP-1, and TIMP-1 mRNA expression, Northern analysis (Figure 2) was completed after MSF stimulation for 18 hours with muOSM, mIL-1β, and the combination of muOSM and mL-1β (left panel), or with other IL-6/LIF cytokines (right panel). muOSM was found to be a potent inducer of IL-6 mRNA in these cells. Densitometry analysis from both experiments determined this to be ninefold and 20-fold increases in IL-6 transcripts, respectively (Figure 2). None of the other IL-6/LIF cytokines tested induced increases exceeding twofold. The combination of muOSM and mL-1β had an additive effect on IL-6 expression (9.1- and 8.3-fold increases induced by muOSM and mIL-1β, respectively, with the combination of muOSM and mIL-1β resulting in a 13.8-fold increase). Similar trends were seen at the protein level (Figure 1C). Stimulation with muOSM resulted in marked increases in mTIMP-1 mRNA (estimated as 42-fold when analyzed by densitometry and normalized to

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β-actin), whereas mCT-1 and mLIF induced modest increases in TIMP-1 message (approximately fourfold) and mIL-6 gave only a slight increase of twofold (Figure 2, right). mL-1β resulted in a sevenfold increase in TIMP-1 message and the combination of mL-1β and muOSM was similar to OSM alone in ability to regulate TIMP-1 mRNA.

Proliferation in Anchorage-Independent Conditions

Proliferation and colony formation by synovial fibroblasts in anchorage-independent conditions has been considered an indicator of the ability of synoviocytes to contribute to the aggressive and destructive pannus tissue seen in rheumatoid arthritis patients. We examined the response of mouse and rat synovial fibroblast cells to muOSM, mL-6, and mLIF in soft agarose cultures for the ability to induce the formation of cell colonies. Colonies smaller than 20 μm represent single cells or clusters of only a few cells, whereas colonies of 20 μm or larger represent true, thriving colonies. Although mL-6 and mLIF induced minimal increases in colonies 20 μm in diameter or larger, muOSM induced sixfold increases in the formation of colonies by MSF (Figure 3). In standard anchorage-dependent proliferation studies (not shown), we did not observe significant regulation of proliferation by OSM as measured by MTT incorporation. Therefore anchorage-independent growth is unlikely to simply reflect standard proliferation. Cultures of rat synovial fibroblasts, derived in a similar manner to MSF cultures, also showed induction of IL-6 production and an increase (11-fold) in colony growth assays in response to muOSM (not shown). Thus, muOSM is able to induce the anchorage-independent growth of synovial fibroblasts in vitro, under conditions that are suggestive of the behavior of synovial fibroblasts in the inflamed joint in vivo.

Adenovirus Vectors in Vitro and in Vivo

The responses of mouse synovial cells to OSM in vitro suggest they could be a major target of OSM in vivo. To assess the potential of OSM to regulate synovial tissue responses in vivo, we have used adenovirus vectors administered locally. The use of Ad vectors in expressing genes in synovium of rabbits and mice has been reported.44,45 We have constructed and characterized an adenovirus vector that expresses mouse OSM in vitro and in vivo, on administration to mice.46 Intramuscular administration resulted in induction of IL-6 and acute-phase protein levels in serum, and local administration resulted in induction of TIMP-1 in lungs on intratracheal treatment.46 To ensure that mouse synovial cells and tissue can be infected and express adenovirus vector inserts, we first assessed in vitro infection of MSF and in vivo administration of an Ad vector expressing lacZ. Mouse synovial fibroblasts infected in vitro with AdmCMVlacZ (AdlacZ) at a multiplicity of infection of 10 and 50 showed significant expression of the virus as noted by the colorimetric conversion of X-gal to a blue product (Figure 4A) demonstrating that this primary synovial fibroblast cell line is infectable with adenovirus constructs. Both AdmuOSM-
infected and uninfected control cells were negative for X-gal staining. Intra-articular administration of AdlacZ to mice resulted in detection of lacZ activity primarily in the lining cells of the synovium at day 1 after administration (Figure 4B) and was detectable in synovium of animals for up to 21 days after treatment (not shown).

The effects of the Ad vectors expressing muOSM, lacZ, and mIL-6 on the histology of joint tissue were then examined for up to 21 days after administration. Results shown in Figure 5 display typical tissue histological analysis representative of at least five joints per treatment whereas the contralateral knees of C3H/HeJ mice were untreated. Particle calculation shows that doses of AdmOSM particles administered (as opposed to pfu administered) were similar or less than particle administrations into knees treated with Add170 or AdmIL-6. Therefore the effects of AdmOSM are not because of particle number alone, but because of the specific properties of the mOSM-expressing recombinant virus. Sagittal sections of the contralateral knees (Figure 5A) of any of the treatments showed no differences from untreated animals. Joints treated with AdlacZ, or with empty vector Add170 (Figure 5B) showed little or no evidence of inflammatory changes at day 7, which also reflects earlier time points (not shown) using these doses of Ad vector. Joints from animals treated with AdmuOSM, however, showed a progressive inflammatory response that began to be evident by day 4 (Figure 5C). This included increased numbers of infiltrating cells that were mainly mononuclear with some PMNs, and increased numbers of cells with synovial fibroblast-like morphology that contributed to increased thickness of the synovial lining. At day 7 after AdmuOSM treatment (Figure 5, D and E) the amount of infiltration and hyperplasia of the synovium was enhanced with an apparent decrease in subsynovial adipocytes. Increased matrix deposition was evident throughout the synovial tissue, and hyperplastic synovial lining was found to adhere in places to the cartilage surface, particularly at the margins. Figure 5D shows marked degradation of the outer surface of the meniscal cartilage (arrowhead) again adjacent to the hyperplastic synovial tissue. Furthermore, some sections showed that the hyperplastic synovium invaded cartilage and protruded through to bone marrow (Figure 5G). EVG staining showed a re-organization of collagen deposition toward a more diffuse distribution of collagen fibrils (Figure 5H). Empirical observation during dissection consistently showed swelling evident in the AdmuOSM-treated joints but not in other treatments or contralateral knees. Joints examined at day 14 after treatment showed a further increase in fibroblast-like cells and ongoing infiltration, resulting in a substantial synovial lesion.

The lesions were still evident at day 21, at which time synovial hyperplasia appeared to be no less abundant, but the degree of infiltration was less extensive than days 7 and 14. The results were similar on administration of AdmuOSM to other strains of mice including C57BL/6 and MRL+ +. Joints of MRL/lpr mice treated with AdmuOSM showed a more rapid response to AdmuOSM, such that synovial hyperplasia and tissue remodeling was extensive at day 7 and day 14. Thus, dramatic connective tissue changes were evident in AdOSM treated joints. In our hands the pannus-like tissue in treated mice does not yield sufficient quantities for RNA Northern analysis. We have been able to detect transgene mRNA by Northern blots in lungs of animals treated intratracheally. Development of reagents to measure muOSM protein specifically will enable future studies that determine the expression level of transgene as well as endogenous OSM in mouse joint tissues.

Joints of C3H/HeJ mice treated with AdmIL-6 showed a mild increase in hyperplasia at day 7 (Figure 5F) which was not sustained. This construct was previously shown to induce various effects in lungs of rats treated intratracheally. However, 5 × 10⁷ pfu of AdmIL-6, a dose 10-fold greater than was shown to be effective for AdmuOSM (5 × 10⁶ pfu), did not result in inflammatory responses in C3H/HeJ mice.

To quantify the response of mouse joints to the adenoviral vectors, a rating of inflammatory indices of day 7-treated joints in shown in Table 1. For each treatment, a minimum of three and a maximum of nine joints were assessed. The total pathological score is shown in Table 1. Assessment of contralateral knee joints in AdmOSM treated animals by histopathology indices showed no difference from untreated joints (not shown), indicating intra-articular administration did not result in general effects at other joints. For every measurement of joint pathology, AdmuOSM consistently demonstrated effects not seen by administration of either the control Add170 vector, or AdmIL-6.
Discussion

OSM as well as other IL-6/LIF cytokines are found in elevated levels in synovial fluid of rheumatoid arthritic patients. However, the precise role of OSM in human rheumatoid arthritis is not clear. Its ability to regulate responses in many connective-tissue cell types including synovial fibroblasts and chondrocytes suggests an important role in tissue remodeling processes. The establishment of the actions of OSM in local inflammation in animal models will help to discern the role of OSM in human disease. Cloning of the mouse OSM ho-

Figure 5. Histopathology of mouse knee joints. Adenovirus vectors were injected into the knee joints of mice, and the animals were sacrificed at various times following administration. Knee joints were fixed, decalcified, paraffin-embedded, and stained using H&E and EVG techniques. Tissues were assessed using pathological criteria as shown in Table 1. Original magnification: \( \times 100 \) (A–D and F), \( \times 200 \) (E, G, and H). A: Contralateral joint of AdmuOSM, day 7. A, Adipocytes; M, meniscus; C, articular cartilage; T, tendon. Note the normal appearance and thickness of the synovial lining (arrow). B: Adil70, day 7. The synovial lining shows little if any thickening (arrow). C: AdmuOSM, day 4. Note the substantial increase in the thickness of the synovial lining (arrow). D: AdmuOSM, day 7. The synovial lining is extensively thickened and the entire synovium is infiltrated with mononuclear and some polymorphonuclear cells (PMN), with a concomitant decrease in adipocyte presence. Note the eroded meniscal outer surface (arrow). E: AdmuOSM, day 7. Higher magnification shows infiltration of the synovium, the PMN being clearly visible (open arrowheads). Note also the erosion of the articular cartilage adjacent to synovial tissue (closed arrowhead). F: AdmIL-6, day 7. The synovial lining shows a minor increase in thickness (arrow). G and H: AdmuOSM, day 7, showing matching fields stained with H&E and EVG, respectively. The synovial tissue has eroded the articular cartilage and underlying bone, and has penetrated the bone marrow (BM) (large arrowhead).
mologue has been followed by studies that show similar, but not identical, effects in the mouse and human systems. Novel observations are described in the present study, suggesting that mouse OSM stimulates a chronic inflammatory response in mouse joints.

Mouse OSM regulates several responses in (MSF) in vitro. MSF produced high levels of IL-6 (30- to 300-fold) in response to muOSM (Figure 1) and this was consistent in several lines of MSF cells as well as fibroblast cultures derived from mouse lung or skin (not shown). In contrast, mLIF, mCT-1, or human OSM were not able to induce IL-6 expression. Marked induction of TIMP-1 mRNA by muOSM was evident at day 1 (Figure 2) and at day 3 (not shown) after treatment in vitro. This is consistent with other studies on mouse fibroblast NIH 3T3 cells and in human systems using the human OSM protein. Similar to expression of IL-6, TIMP-1 did not respond to LIF, CT-1, or IL-6 stimulation in MSF. This could be because of low receptor number on fibroblasts for these cytokines, however mLIF and mCT-1 can induce STAT-1/3 activation in MSF indicating receptor presence on these cells (Carl D. Richards, unpublished observations). Alternatively, additional signaling pathways induced by the specific OSMR complex may participate in the regulation of IL-6. The stimulation of IL-6 by OSM may contribute to the levels of IL-6 that are elevated in rheumatoid arthritis synovial fluid and tissue as established in the literature.

MCP-1 was also induced in vitro by muOSM, although to a much lesser extent (twofold induction) than IL-6. Basal mRNA levels and protein content in cell culture supernatants were high (3 ng/ml) possibly making it more difficult to detect induction. Analysis in human synovial fibroblasts showed lower basal levels of MCP-1 production and more marked MCP-1 responses to human OSM. This may reflect species differences in responses of MSF to these cytokines. Mouse MCP-5 has recently been characterized and shows greater identity to human MCP-1 (66% amino acid identity) than mouse MCP-1 does (55% homologous). Whether OSM does indeed regulate other MCPs has yet to be determined. The stimulation of MCP-1 will contribute to the pool of chemotactic factors that control the infiltration of inflammatory cells. MCP-1 is primarily chemotactic for monocytes/macrophages and T cells and because MCP-1 is found in elevated levels in rheumatoid arthritis patient synovial fluid and tissue, the infiltration of such mononuclear cells in rheumatoid arthritis may thus be mediated in part by the OSM-mediated regulation of MCP-1.

The ability of synovial fibroblast-like cells to grow in soft agarose has been described previously, and this has been noted as a reflection of the aggressive nature of the fibroblasts from synovium or pannus of rheumatoid arthritis patients. This can be regulated in culture by a number of growth factors, including platelet-derived growth factor, transforming growth factor-β, and fibroblast growth factor. Our data also shows that OSM can stimulate the formation of colonies in soft agarose, and therefore act as a stimulus of such phenotypic change. Collectively, this data suggests that muOSM is a potent stimulus of synovial cell responses in vitro, supporting a potentially important role in inflamed joints in vivo, unique from other IL-6/LIF cytokines.

Analysis of mouse joints treated with the AdnuOSM vector showed extensive synovial phenotypic change including hyperplasia, mononuclear cell infiltration, extracellular matrix remodeling, and cartilage damage. These effects are consistent with the activities of muOSM on MSF in vitro. Anchorage-independent growth stimulation may reflect the hyperplasia and formation of aggressive cells that overlay cartilage and penetrate the tissue, as has been suggested by others. The presence of mononuclear cells may reflect in part the regulation of MCP-1 or other chemotactic factors by OSM. The development of inflammation is evident by day 4, but hyperplasia and extracellular matrix remodeling is most evident at day 7 and thereafter in this model. The expression of cytokines by this and other Ad vectors are typically for 3 to 7 days in other tissues, but it is not known at this time whether muOSM is expressed for a similar period in this model. Certainly the effects of a single administration are evident for 14 to 21 days, suggesting sustained phenotypic change has been induced characteristic of chronic inflammation. Neither the control vector (Add170) nor the AdmIL-6 vector produced such change, even though 10-fold greater doses of each have been tested. The lack of effects of AdmIL-6 (previously published to express high levels of IL-6 and induce lung and liver responses) would suggest that the effects of AdnuOSM are not mediated through IL-6, which is also consistent with the differences in responses of MSF to these cytokines. Whether the sustained effects of AdnuOSM depend on the recruitment of certain cell populations, and activation of certain cytokines, is the subject of further examination.

### Table 1. Inflammatory Indices of Day 7-Treated Joints

<table>
<thead>
<tr>
<th>Pathological index</th>
<th>Add170</th>
<th>AddOSM</th>
<th>AdIL-6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Synovial inflammation</td>
<td>0.0 (0.0)</td>
<td>1.7 (0.3)</td>
<td>0.3 (0.6)</td>
</tr>
<tr>
<td>Synovial hyperplasia</td>
<td>0.2 (0.4)</td>
<td>2.0 (0.0)*</td>
<td>0.3 (0.6)</td>
</tr>
<tr>
<td>Pannus formation cartilage erosion</td>
<td>0.0 (0.0)</td>
<td>1.6 (0.5)*</td>
<td>0.3 (0.6)</td>
</tr>
<tr>
<td>Bone destruction</td>
<td>0.0 (0.0)</td>
<td>1.3 (0.5)*</td>
<td>0.3 (0.6)</td>
</tr>
<tr>
<td>(\text{Synovitis} )</td>
<td>0.0 (0.0)</td>
<td>0.48 (0.5)</td>
<td>0.0 (0.0)</td>
</tr>
<tr>
<td>Pathological score</td>
<td>0.2 (0.4)</td>
<td>7.0 (1.8)*</td>
<td>1.2 (2.3)</td>
</tr>
</tbody>
</table>

*Significant difference from Add170 (P < 0.001).
Recent data has shown that muOSM binds to gp130 and signaling requires the specific OSM receptor subunit. In contrast to human OSM, muOSM interacts very weakly with the mouse LIF receptor complex and does not mediate responses in cells that express the LIF receptor only. Furthermore, in vitro binding assay data clearly shows that human OSM does not act on the muOSM receptor complex, suggesting species specificity of this interaction. This suggests that recently published work examining the effect of human OSM administered to mice reflects interaction of human OSM with receptors other than the specific muOSM receptor complex. This is likely, but possibly not exclusively, the mouse LIF receptor. Since the work of Wallace et al. showed an inhibitory action by systemic administration of human OSM in models of chronic inflammation, this implies that such LIF receptor activation may result in anti-inflammatory effects in mice. We believe that OSM receptor activation induces chronic inflammation on local administration, and therefore suggest that elevated levels of OSM found in human rheumatoid arthritis tissue contribute to the pathology of the disease through the specific OSM receptor.

Acknowledgments

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References

development requires a paracrine action of oncostatin M through the gp 130 signal transducer. EMBO J 1999, 18(suppl 8):2127–2136


