Tumor-Conditioned Macrophages Secrete Migration-Stimulating Factor: A New Marker for M2-Polarization, Influencing Tumor Cell Motility

Graziella Solinas, Silvia Schiarea, Manuela Liguori, Marco Fabbri, Samantha Pesce, Luca Zammataro, Fabio Pasqualini, Manuela Nebuloni, Chiara Chiabrando, Alberto Mantovani and Paola Allavena

*J Immunol* 2010; 185:642-652; Prepublished online 7 June 2010;
doi: 10.4049/jimmunol.1000413
http://www.jimmunol.org/content/185/1/642

References  This article cites 107 articles, 41 of which you can access for free at: http://www.jimmunol.org/content/185/1/642.full#ref-list-1

Subscription  Information about subscribing to *The Journal of Immunology* is online at: http://jimmunol.org/subscription

Permissions  Submit copyright permission requests at: http://www.aai.org/About/Publications/JI/copyright.html

Email Alerts  Receive free email-alerts when new articles cite this article. Sign up at: http://jimmunol.org/alerts
Tumor-Conditioned Macrophages Secrete Migration-Stimulating Factor: A New Marker for M2-Polarization, Influencing Tumor Cell Motility

Graziella Solinas,* Silvia Schiarea,† Manuela Ligouri,* Marco Fabbri,*§ Samantha Pesce,* Luca Zammataro,* Fabio Pasqualini,* Manuela Nebuloni,§ Chiara Chiabrando,† Alberto Mantovani,*§ and Paola Allavena*

Tumor-associated macrophages (TAMs) are key orchestrators of the tumor microenvironment directly affecting neoplastic cell growth, neoangiogenesis, and extracellular matrix remodeling. In turn, the tumor milieu strongly influences maturation of TAMs and shapes several of their features. To address the early macrophage (Mφ) differentiation phase in a malignant context, we mimicked a tumor microenvironment by in vitro coculturing human blood monocytes with conditioned media from different cancer cell lines. Only 2 out of 16 tumor cell lines induced Mφ differentiation due to secreted M-CSF isoforms, including high molecular mass species. A global gene profiling of tumor-conditioned Mφ was performed. Comparison with other datasets (polarized M1-Mφ, M2-Mφ, and TAMs isolated from human tumors) highlighted the upregulation of several genes also shared by TAM and M2-polarized Mφ. The most expressed genes were selenoprotein 1, osteoadivin, osteopontin, and, interestingly, migration-stimulating factor (MSF), a poorly studied oncofoetal isoform of fibronectin. MSF (present in fetal/cancer extracellular matrix; ED, extra domain; FN, fibronectin; Gel-BD, gelatin-binding domain; LC-MS/MS, liquid chromatography tandem mass spectrometry; M, migration-stimulating factor; O/N, overnight; PCA, principal component analysis; rhM-CSF, recombinant human M-CSF; TAM, tumor-associated macrophage; TC-Mφ, tumor-conditioned macrophage.

Abbreviations used in this paper: CM, conditioned media; Ct, cycle threshold; ECM, extracellular matrix; ED, extra domain; FN, fibronectin; Gel-BD, gelatin-binding domain; LC-MS/MS, liquid chromatography tandem mass spectrometry; Mφ, macrophage; MS, tandem mass spectrometry; MSF, migration-stimulating factor; O/N, overnight; PCA, principal component analysis; rhM-CSF, recombinant human M-CSF; TAM, tumor-associated macrophage; TC-Mφ, tumor-conditioned macrophage.

Received for publication February 15, 2010. Accepted for publication April 22, 2010.

This work was supported by Associazione Italiana Ricerca Cancro Italy (to P.A. and A.M.) and grants from the Ministry of Health and Istituto Superiore Sanita Italy (Project Oncology 2006 and Alleanza Contro il Cancro).

Address correspondence and reprint requests to Dr. Paola Allavena or Prof. Alberto Mantovani, Department of Immunology and Inflammation, Istituto Di Ricovero e Cura a Carattere Scientifico Istituto Clinico Humanitas, Rozzano, Milan, Italy. E-mail addresses: paola.allavena@humanitasresearch.it or alberto.mantovani@humanitasresearch.it

Copyright © 2010 by The American Association of Immunologists, Inc. 0022-1767/10/$16.00
suppressor cells, wane the inflammatory/immune response by downregulating M1- and T cell-mediated functions (28, 35–40). On the contrary, inflammatory Mφ (or classically activated or M1) originate upon stimulation with IFN-γ and microbial stimuli, such as LPS, and are characterized by IL-12p40 and IL-23 production and consequent activation of polarized type I T cell response, cytotoxic activity, and good capability as APCs (28, 36–39).

Molecular mechanisms underlying Mφ polarization remain scanty. Tuning of NF-κB activation by p50 homodimers has been associated with M2 differentiation (41, 42). Peroxisome proliferator-activated receptor γ is involved in the M2-like differentiation (43–46), and the phosphatase SHIP was shown to play a key role in balancing Mφ polarization, although recent evidence suggests that it may act indirectly through the regulation of IL-4 production by basophils (47). Recently, Pienta and colleagues (48) reported that CCL2 and IL-6 sustain the survival of myeloid cells at the tumor site and support their differentiation toward tumor-promoting M2-Mφ.

Tumor Mφ are considered attractive targets of anti-cancer strategies. To this end, the first step should be the identification of molecules specifically overexpressed or produced by TAMs but neither by resident Mφ of distant healthy tissues nor by M1 cells, which are important to face pathogens and could take part in anti-cancer actions.

In this study, we focused our work on the initial differentiation phase of monocytes to identify molecules upregulated by Mφ within the tumor context. To mimic a tumor microenvironment, blood monocytes were in vitro exposed to cancer cell-conditioned media. Differentiated Mφ, named tumor-conditioned Mφ (TC-Mφ), were analyzed for phenotype, functional activity, cytokine/chemokine production, and gene expression.

In this study, we report that TC-Mφ have a gene expression pattern similar to that of TAMs. Among these genes, we focused on a truncated isoform of fibronectin (FN), migration-stimulating factor (MSF), which is potently chemotactic for tumor cells. We provide evidence that MSF is produced by M2-Mφ and TAMs and therefore may be considered as a novel marker for M2 polarization.

Isolation of human TAMs

Human TAMs were isolated from solid tumors of untreated patients with histologically confirmed epithelial ovarian carcinoma admitted to the Department of Obstetrics and Gynecology, San Gerardo Hospital (Monza, Italy). Briefly, solid tumors were enzymatically digested and centrifuged once as described (41). Tumor Mφ were isolated by plastic adherence (RPMI 1640 without FBS, 1 h, 37°C). The adherent cells were >95% Mφ as assessed by morphology and CD68 positivity. All culture reagents contained <0.125 endotoxin unit/ml as checked by Limulus amebocyte lysate assay (BioWhittaker).

M-CSF blockade

To block M-CSFRs, monocytes were incubated with 2 μg/ml anti-M-CSFR (Santa Cruz Biotechnology, Tuber-bio, Milan, Italy) for 30 min at 37°C or with an anti-human IgG irrelevant Ab 10 μg/ml (Sigma-Aldrich).

Tumor-derived M-CSF neutralization

To neutralize tumor-derived M-CSF, Panc1-CM was incubated for 20 min with different concentrations of Recombinant Human M-CSF R-Fe Chimeric (Symansis, Auckland, New Zealand). As experimental control, a solution of rM-CSF 25 ng/ml was treated following a similar protocol.

Chemotaxis

Cell migration was evaluated using transwell systems with 5 μm (for monocytes) or 8 μm (for Panc1 tumor cells) pore size (Costar, Euroclone, Milan, Italy). CM from TC-Mφ or M1- and M2-Mφ were prepared after 6 to 7 d of differentiation as described above. Medium was replaced with fresh medium and supernatants collected after 24 h. Mφ supernatants or recombinant human MSF (kind gift of Prof. Schor, University of Dundee, Scotland, U.K.) were used as chemoattractants in the lower compartment. To block gelatin-binding domain (Gel-BD), tumor-conditioned supernatants were pretreated for 30 min at room temperature with anti-Gel-BD Ab (Chemicon, Milan, Italy), 10 μg/ml. Results are expressed as the mean number of migrated cells counted in 10 microscope high-power fields (magnification ×1000) after 18 h. Each experiment was performed in triplicate. The p value was calculated by Student t test.

M-CSF and TC-Mφ RT-PCR

Total RNA extraction from monocytes, in vitro-cultured Mφ, and TAMs isolated from human tumors was performed with TRIzol (Invitrogen, Milan, Italy). cDNA was synthesized by random priming from 1 μg total RNA with the GeneAmp RNA PCR kit (Applied Biosystems, Monza, Italy) according to the manufacturer’s instructions. Real-Time PCR was performed using SYBR Green dye and 7900HT Fast Real Time PCR Systems (Applied Biosystems). The sequences of the primer pairs specific for each gene (Invitrogen) were designed with Primer Express Software (Applied Biosystems). All primers were M-CSF isoforms: human CSF-1 isoform A (5′-GCC ATC CCT GAC C-3’ and 5′-TCA AAG GAA CGG AGT TAA AAC GG-3’); human CSF-1 isoform B (5′-GCC AGC AGG AGT ATG ACC G-3’ and 5′-CCC TCA GTT CCC TCA GAG TC-3’); human CSF-1 isoform C (5′-GCT CCT CCA GGA TCT CAT CAC-3’ and 5′-AGG TCT CCA TCT GCC GAC TCT CAC-3’); human CSF-1 isoform D (5′-GTT CCA TCC ATT TAA CAG CAC-3’ and 5′-TTT CTG GTG GGC ATA CTC ACC-3’); extra domain (ED)-A (5′-GGG CAG GAT CCA ACC AATA AAG TGA CAG-3’ and 5′-GAT CCT CAC GGA GGC CAC-3’); ED-B (5′-GGG CAG GAT CCA ACC AATA AAG TGA CAG-3’ and 5′-GAT CCT CAC GGA GGC CAC-3’); ED-C (5′-GGG CAG GAT CCA ACC AATA AAG TGA CAG-3’ and 5′-GAT CCT CAC GGA GGC CAC-3’); ED-D (5′-GGG CAG GAT CCA ACC AATA AAG TGA CAG-3’ and 5′-GAT CCT CAC GGA GGC CAC-3’); ED-E (5′-GGG CAG GAT CCA ACC AATA AAG TGA CAG-3’ and 5′-GAT CCT CAC GGA GGC CAC-3’); ED-F (5′-GGG CAG GAT CCA ACC AATA AAG TGA CAG-3’ and 5′-GAT CCT CAC GGA GGC CAC-3’).

Materials and Methods

Cell line cultures and tumor-conditioned media preparation

Human ovarian cancer cell lines Ovarcar3 and A2780, human colon cancer cell lines HT29, HCT-116, LoVo-N, CaCo-1, and SW620, human pancreatic carcinoma cell lines PaCa44, PaCa3, CFPAC, PC, T3M4, PT45, Panc1, MiaPaCa2, and AsPC1, and the immortalized normal pancreatic epithelial cell lines HPDE6 were cultured in RPMI 1640 (Lonza, Basel, Switzerland) supplemented with 10% FBS. Once grown to 90% of confluence, media were discarded, and flasks were rinsed two times with saline solution. Cells were then incubated with fresh RPMI for 24 h; the conditioned media (CM) was collected and filtered at 0.20 μm, and the supernatant was stored at −20°C. All cell lines were routinely checked for Mycoplasma contamination.

Mφ and TC-Mφ differentiation

Human monocytes were obtained from normal blood donor buffy coats by two-step gradient centrifugation, first by Histopaque-1077 (Sigma-Aldrich, Milan, Italy) and then by Percoll (GE Healthcare, Milan, Italy) (49). Residual T and B cells were removed from monocyte fraction by plastic adherence.

Mφ and TC-Mφ were obtained by culturing 10^5/ml monocytes for 6 d in RPMI 1640 5% FBS supplemented with 25 ng/ml of recombinant human M-CSF (rhM-CSF; PeproTech, Milan, Italy) or in the presence of 30% of tumor cell line supernatants (TC-Mφ). All culture reagents contained <0.125 endotoxin unit/ml as checked by Limulus amebocyte lysate assay (BioWhittaker, Walkersville, MD).

M1- and M2-polarized Mφ were obtained by culturing 10^5/ml monocytes for 6 d in RPMI 1640 5% FBS with 25 ng/ml rhM-CSF. M1 cells were polarized by stimulating overnight (O/N) M-CSF Mφ with LPS (100 ng/ml) (PeproTech) and IFN-γ (500 U/ml) (PeproTech). M2-Mφ were polarized by stimulating O/N M-CSF Mφ with IL-4 (20 ng/ml) (PeproTech).
Tumor-conditioned macrophages influence tumor cell motility

(Affymetrix, Santa Clara, CA), then washed and scanned according to the manufacturer’s guidelines. Expression measures were computed using robust multiarray average. Principal component analysis (PCA) was carried out on all genes analyzed to assign the general variability in the data to a reduced set of variables called principal components. Available were datasets of human TAMs (three different donors) and datasets from M1- and M2-polarized M\(_6\).

PCA (analysis was applied to the complete dataset) is a mathematical algorithm that reduces the dimensionality of the data. It accomplishes this reduction by identifying directions, called principal components, along which the variation in the data is maximal.

In PCA, we obtain a set of orthogonal axes oriented in the directions of largest variance within a set of data points in a high-dimensional space. The first principal component is a vector in the direction of greatest variance, the second principal component is a vector in the direction of greatest variance orthogonal to this, and so on. These vectors are in fact eigenvectors of the empirical data covariance matrix. Values on the \(x\) - and \(y\)-axis express a two-dimensional representation of greatest variance vectors (50).

Technical information requested by Minimum Information about a Microarray Experiment of the latter is available at the Gene Expression Omnibus Web site (www.ncbi.nlm.nih.gov/geo) under the accession number GSE5099. Statistical differences were assessed by a moderated \(t\)-test analysis performed using a limma bioconductor package, and resulting \(p\) values were adjusted using the Benjamini and Hochberg step-up method for controlling the false discovery rate. Genes were defined as regulated when characterized and \(p\) values adjusted using the Benjamini and Hochberg step-up method for controlling the false discovery rate. Genes were defined as regulated when characterized by a fold of induction \(>\)2 and a false discovery rate \(p\) value \(<\)0.05. Computations were conducted using the R statistics programming environment (available at www.r-project.org). Gene trends were organized by K-means clustering using squared Pearson correlation with the TIGR MultiExperiment Viewer (www.tm4.org/). Single-gene comparison was performed by analyzing the relative expression to median value (1) calculated among four Affymetrix datasets (TC-M\(_6\), M1, M2, and TAM).

ELISA
Cytokine production in tumor cell and M\(_6\) supernatants was measured by commercially available ELISA kits (IL-10, IL-12, CCL2, CCL17, IL-6, IL-8, TNF-\(\alpha\)) according to the manufacturer’s instructions (R&D Systems, Space Import, Milan, Italy). M\(_6\) were stimulated or not with LPS (100 ng/ml) (PeproTech) for 24 h.

Bio-Plex Protein Array System
M-CSF in tumor cell line supernatants was measured using the Bio-Plex Protein Array System (Bio-Rad, Milan, Italy) according to the manufacturer’s instructions.

Western blot
Because there are no commercially available reagents for MSF, we kindly received anti-MSF Abs from Prof. Schor (University of Dundee). Rabbit polyclonal anti-MSF Ab (raised against a synthetic peptide containing the MSF-specific C-terminal decapeptide sequence) was used diluted 1:10,000. Protein extracts were processed as described (19).

Flow cytometry
In vitro-differentiated M\(_6\) were analyzed by flow cytometry on FACS-Canto (BD Biosciences, Milan, Italy). Human FcRs were blocked using PBS 1% human serum. For staining, cells were washed and resuspended in FACS buffer (PBS 0.5% BSA, 0.05% NaN\(_3\)). PE-mouse anti-human CD16, APC-mouse anti-human CD14, FITC-mouse anti-human mannose receptor CD206, and PE-mouse anti-human CD68 were obtained from BD Biosciences. For the CD68 staining, we used the BD Fixation/Permeabilization solution kit (BD Biosciences).

Immunohistochemistry
Human surgical samples of pancreatic and ovarian tumors were fixed in 10% buffered formalin for 1–4 h and embedded in paraffin. Sections were cut mounted on Superfrost slides (BioOptica, Milan, Italy), dewaxed in xylene, rehydrated in ethanol, and pretreated in a microwave oven (two cycles for 3 min each at 800 W in 0.25 mM EDTA buffer). Double immunohistochemistry was performed with anti-human CD68 (clone KPI, Dako, Glostrup, Denmark; 1:100 to detect M\(_6\)) and anti-human MSF Ab (mAb clone 7.1, 1:100; Prof. Schor, University of Dundee). Specific secondary Abs conjugated with alkaline phosphatase (Biocare Medical, Concord, CA) for CD68 and ARK peroxidase (Dako) were used. The first reaction (anti-CD68) was developed by using Ferranji blue (blue-stained) and the second (anti-MSF) by using 3,3’-diaminobenzidine–free base as chromogen (brown-stained). For the second staining, double immunohistochemistry was performed with anti-human CD68 (clone KPI, Dako; 1:100 to detect M\(_6\)) and rabbit polyclonal anti-human MSF Ab (home-made, raised against a synthetic peptide containing the MSF-specific C-terminal decapeptide sequence; diluted 1:100). Mach2 double stain 1 (Biocare Medical) was used to detect mouse primary Ab in alkaline phosphatase and rabbit primary Ab in peroxidase. The first reaction (CD68) was developed by using Ferranji blue (blue-stained; Biocare Medical) and the second by using diaminobenzidine as chromogen (brown-stained; Biocare Medical); nuclei were counterstained with Nuclear Fast Red (Bio-Op-tica, Milan, Italy).

Global proteomic analysis of CM
A proteomic analysis was performed on serum-free CM of tumor cell lines (PT45, Panc-1, MiaPaCa2, ASPC1) (S. Schiara, G. Solinas, P. Allavena, G.M. Scigliuolo, R. Bagnati, R. Fanelli, C. Chiabrando, submitted for publication). CM was concentrated using 5-kDa molecular mass cut-off centrifugal filter devices (Millipore, Bedford, MA). Proteins were separated by one-dimensional SDS-PAGE in reducing conditions on 4–12% NuPAGE Bis-Tris gel (Invitrogen). Each gel lane was cut into 24 bands of equal height. Proteins in each bands were reduced, alkylated, and in-gel digested with bovine trypsin. Digests were analyzed by liquid chromatography tandem mass spectrometry LC-MS/MS with a high-resolution/high-accuracy mass spectrom-eter (LTQ Orbitrap XL, Thermo Fisher Scientific, Waltham, MA). Tandem MS/MS data were analyzed with Mascot version 2.2 (Matrix Science, Boston, MA) against the Swiss-Prot database, version 56.5 (http://us.expasy.org/sprot/download.html).

Results
Effects of tumor cell line conditioned-media on human monocytes
To mimic a tumor microenvironment, we exposed human blood monocytes to tumor CM, in the absence of other exogenous stimuli. We tested several CM from different tumor cell lines (live from colorectal cancer, two from ovarian cancer, and nine from pancreatic carcinoma) and noticed that only those from the pancreatic cancer cell lines PT45 and Panc1 were able to induce evident morphological modifications of monocytes. Therefore, we decided to restrict our investigation to these active cell lines, using as negative controls other pancreatic (cancer or normal) cell lines without differentiating activity (i.e., AsPC1 and MiaPaCa2) and the normal human pancreatic ductal epithelial cell line HPDE6. The CM from PT45 or Panc1 as well as rhM-CSF induced a strong differentiation of monocytes that became larger, with ruffling membrane typical of M\(_6\), whereas the majority of monocytes cultured with AsPC1-, MiaPaCa2-, and -HPDE-CM almost completely died. Representative images of the cultured cells are shown in Fig. 1A. We confirmed by FACS analysis that the differentiation of monocytes (evaluated as percentage of large cells, side scatter) was strongly induced by rhM-CSF (46.77 ± 8.8%) and PT45- (41.62 ± 2.55%) or Panc1-CM (36.24 ± 2.7%). In contrast, only minimal monocyte differentiation was induced by other CM, as well as by RPMI without additions (6.35 ± 1.18%) (Fig. 1B). M\(_6\) differentiated by tumor-CM were defined in this study as TC-M\(_6\).

To study the phenotype of the differentiated cells, we assessed the expression of CD14, CD16, CD68, and mannose receptor. All of these molecules were highly expressed by M\(_6\) cultured with rhM-CSF as well as M\(_6\) differentiated in the presence of PT45- and Panc1-CM (Fig. 1C). The dendritic cell marker CD1a was completely absent from TC-M\(_6\), thus excluding that tumor-CM were inducing a dendritic cell-like differentiation (data not shown). To better characterize the differentiated cells, we measured the constitutive and LPS-stimulated release of cytokines/chemokines by rhM-CSF-M\(_6\), TC-M\(_6\), and TAMs directly isolated from human tumor. In particular, Panc1-CM induced expression of IL-10, IL-6, and TNF-\(\alpha\) and high amounts of CCL2 by M\(_6\). Similarly, PT45-M\(_6\) produced IL-6 and TNF-\(\alpha\) and very high quantities of IL-10 and CCL2. Notably, neither Panc1-M\(_6\) nor PT45-M\(_6\) produced...
IL-12, even after LPS stimulation. The cytokine/chemokine production profiles of TC-Mφ were similar to those of freshly isolated human TAM (Table I).

Identification of monocyte-differentiating bioactivity in tumor-CM

We first established whether the monocyte to Mφ-differentiating activity observed in the two active cell lines was due to the known growth factor, M-CSF. Bio-Plex Protein Array System (Bio-Rad) analysis determined 600 ± 0.01 pg/ml of M-CSF in the active cell line PT45 and 200 ± 0.11 pg/ml in Panc1 and undetectable levels in the inactive cell lines AspCl and MiaPaCa2. As M-CSF is known to exist in separable isoforms with different m.w. (a secreted glycoprotein, a high molecular mass secreted proteoglycan, and a membrane-spanning cell-surface glycoprotein) (51), mRNA levels with primers specific for the three isoforms were tested. The results confirmed a much higher overall expression of M-CSF species in the active than in the inactive cell lines (Fig. 2A). Evidence of the prevalent secretion of high molecular mass M-CSF isoforms was suggested, in an independent study, by a mass spectrometry-based proteomic analysis on the secretome of pancreatic cancer cell lines, in which several M-CSF peptides were identified in the active than in the inactive cell lines (Fig. 2A). Of interest, IL-34, the other only known specific ligand of the M-CSFR, was not identified in the above-cited proteomic analysis of tumor-CM. Taken together, these results demonstrate that the differentiating activity of the active tumor-CM was exclusively due to M-CSF molecular species.

Profiling of upregulated genes expressed by TC-Mφ

To obtain a comprehensive view of our TC-Mφ, we performed a gene expression analysis. Two time points were considered: 4 and 72 h. After 4 h culture, only 12 genes were upregulated (>2-fold) versus resting monocytes. Among these were three chemokines (CCL2, CCL7, and CXCL1), COX-2, the β form of pro-IL-1, and vascular endothelial growth factor (not shown).

A summary of the results of expressed genes at 72 h culture is shown in Table II. Overall, >500 genes had >2-fold expression and 20 genes had >10-fold expression relative to resting monocytes. Several upregulated genes were characteristic of the M2-Mφ polarization: the mannose receptor (CD206), other members of the C-type lectin R family (e.g., MGL/CLEC10/CD301, CLEC11 and DCL-1/CD302, and CD209), the receptors for the Fc fragment of IgG (CD32 and CD64), and receptors importantly implicated in the uptake of extracellular macromolecules and scavenging of debris, like SR-A, SR-B, CD163, and stabilin-1 (STAB1). The latter is a membrane receptor involved in the binding/scavenging of the matricellular protein secreted acidic and rich in cysteine (52).

The most expressed genes were selenoprotein (SEPP1) with a 95-fold expression relative to resting monocytes. Other upregulated molecules related to the ECM were fibronectin (FN1), F13A1, which catalyses the polymerization of several matrix proteins, and osteopontin (OPN). A set of mRNA coding for adhesion molecules was enhanced; these include integrin β5,

Table I. Protein levels produced by TC-Mφ and TAM (ng/ml)

<table>
<thead>
<tr>
<th>Protein</th>
<th>M-CSF-Mφ LPS -</th>
<th>LPS +</th>
<th>Panc1-Mφ LPS -</th>
<th>LPS +</th>
<th>PT45-Mφ LPS -</th>
<th>LPS +</th>
<th>TAM LPS -</th>
<th>LPS +</th>
</tr>
</thead>
<tbody>
<tr>
<td>IL-10</td>
<td>0.13 ± 0.04</td>
<td>0.93</td>
<td>0.06</td>
<td></td>
<td>&lt;0.01</td>
<td>2.36</td>
<td>0.39</td>
<td></td>
</tr>
<tr>
<td>IL-12</td>
<td>0.03 ± 0.01</td>
<td>0.42</td>
<td>0.11</td>
<td></td>
<td>&lt;0.01</td>
<td>0.01</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>CCL2</td>
<td>6.08 ± 2.75</td>
<td>6.46</td>
<td>1.18</td>
<td></td>
<td>5.36 ± 1.55</td>
<td>8.25</td>
<td>2.30</td>
<td>3.63</td>
</tr>
<tr>
<td>IL-6</td>
<td>&lt;0.01</td>
<td>0.24</td>
<td>0.05</td>
<td></td>
<td>0.01 ± 0.00</td>
<td>0.31</td>
<td>0.05</td>
<td>0.02</td>
</tr>
<tr>
<td>TNF-α</td>
<td>&lt;0.01</td>
<td>1.13</td>
<td>0.34</td>
<td></td>
<td>0.01 ± 0.00</td>
<td>1.05</td>
<td>0.46</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Protein levels of chemokines and cytokines produced by TC-Mφ and TAM with and without LPS (mean ± SD of 2–4 experiments).
the VLA-4, P selectin ligand, CD81 (also involved in cell motility as well as cell activation and signal transduction), two sialic-acid binding proteins (Siglec1 and Siglec7), and SLAMF8. Highly expressed (22-fold) was the gene coding for the myelin-associated protein (PMP22), a binding partner in the integrin/laminin complex and specifically binding α6β4.

Table II. Gene expression analysis of TC-Mφ

<table>
<thead>
<tr>
<th>Family</th>
<th>Symbol</th>
<th>Description</th>
<th>Fold Increase</th>
</tr>
</thead>
<tbody>
<tr>
<td>C-type lectin R</td>
<td>MRC1</td>
<td>Mannose receptor/CD206</td>
<td>44</td>
</tr>
<tr>
<td></td>
<td>CD209</td>
<td>DC-SIGN</td>
<td>3.6</td>
</tr>
<tr>
<td></td>
<td>CD80</td>
<td>DCL-1</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>CLEC10A</td>
<td>MGL/CD301</td>
<td>2.5</td>
</tr>
<tr>
<td>Fcγ receptors</td>
<td>FCGR2B</td>
<td>CD32</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>FCGR3</td>
<td>Fc for IgG transporter</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>FCGR1A</td>
<td>CD64</td>
<td>2.5</td>
</tr>
<tr>
<td>Scavenging receptors</td>
<td>MSR1</td>
<td>Scavenger receptor, SR-A</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>STAB1</td>
<td>Stabilin1/SPARC-R</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>CD163</td>
<td>CD163</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>SCARB1</td>
<td>Scavenger R, class B</td>
<td>3</td>
</tr>
<tr>
<td>ECM molecules</td>
<td>GPNMB</td>
<td>OA</td>
<td>83</td>
</tr>
<tr>
<td></td>
<td>SPP1</td>
<td>OPN</td>
<td>26</td>
</tr>
<tr>
<td></td>
<td>F13A1</td>
<td>Factor XIII A1</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>FN1</td>
<td>Fibronectin</td>
<td>7</td>
</tr>
<tr>
<td>Adhesion molecules and related</td>
<td>PMP22</td>
<td>Myelin-associated protein</td>
<td>22</td>
</tr>
<tr>
<td></td>
<td>SIGLEC1</td>
<td>Sialoadhesin 1</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>SLAMF8</td>
<td>CD2 family member</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>ITGB5</td>
<td>β5 integrin</td>
<td>3.5</td>
</tr>
<tr>
<td></td>
<td>CD81</td>
<td>CD81</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>SIGLEC7</td>
<td>Sialoadhesin 7</td>
<td>2.7</td>
</tr>
<tr>
<td></td>
<td>SELP1</td>
<td>P selectin ligand</td>
<td>2.5</td>
</tr>
<tr>
<td></td>
<td>ITGA4</td>
<td>α4 subunit, VLA4</td>
<td>2.1</td>
</tr>
<tr>
<td>Enzymes</td>
<td>ADAMDEC1</td>
<td>Decysin</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td>CPM</td>
<td>Carboxypeptidase</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>MMP9</td>
<td>MMP9</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>ADAM28</td>
<td>ADAM28</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>PLAU</td>
<td>uPA</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>MMP2</td>
<td>MMP2</td>
<td>3</td>
</tr>
<tr>
<td>MHC molecules and related</td>
<td>HLA-DMA</td>
<td>HLA-DMA</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>CHTA</td>
<td>MHC IL, transactivator</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>CTSC</td>
<td>Cathepsin C</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>CTSD</td>
<td>Cathepsin D</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>CTSB</td>
<td>Cathepsin B</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>CTSK</td>
<td>Cathepsin K</td>
<td>2.01</td>
</tr>
<tr>
<td>Protein trafficking</td>
<td>VPS45</td>
<td>Vacuolar sorting protein</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>VPS13B</td>
<td>Vacuolar sorting protein</td>
<td>3.01</td>
</tr>
<tr>
<td></td>
<td>LAMP1</td>
<td>Lysosomal protein 1</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>LAMP2</td>
<td>Lysosomal protein 2</td>
<td>3</td>
</tr>
<tr>
<td>Miscellaneous</td>
<td>SEPP1</td>
<td>Selenoprotein 1</td>
<td>95</td>
</tr>
</tbody>
</table>

Human monocytes were stimulated for 72 h with tumor-CM. Results (mean of three different donors) are fold-increase relative to unstimulated monocytes.
A large number of protease genes were strongly expressed and included: **CPM, MMP2, and MMP9, and ADAM molecules**, such as decysin, **ADAM28**, and **uPA**. Other proteolytic enzymes whose expression was enhanced were the cysteine proteases (**cathepsins type C, D, A, B, and K**). These molecules are extremely important for the maturation of the MHC class II molecules and for the correct loading of antigenic peptides. The genes coding for the **trans-activator of HLA class II CIITA** and genes of the HLA-II family were also increased, especially **HLA-DM**. Other expressed mRNA code for proteins involved in the trafficking and sorting of molecules within specific organelles: **VPS45 and VPS33B** as well as lysosomal-associated membrane proteins (**LAMP1 and LAMP2**).

To broaden our study, we compared the expression of some upregulated genes of **TC-Mφ** with that of TAMs isolated from human tumors and of in vitro **M1- and M2-polarized Mφ** performed in our laboratory with the same Affymetrix technology (53). Some upregulated genes in **TC-Mφ** were similarly expressed in TAM and to a further extent in **M2-Mφ** but not in **M1-polarized cells** (Fig. 3A).

From this restricted gene comparison, we moved to a global gene profiling comparison of **TC-Mφ, TAM, M1, and M2 cells**. To identify sources of variability among these entire databases, a PCA was performed (50, 54, 55).

This PCA showed that **TC-Mφ** are indeed comparable to TAMs, and this observation confirmed the validity of our in vitro tumor-conditioning model of Mφ differentiation. Moreover, their global profiling is closer to that of **M2- than M1-polarized Mφ** (Fig. 3B).

Finally, these data boosted the concept of the M2-like polarization for TAMs.

**Fibronectin isoforms in TC-Mφ**

The mRNA profiling revealed that the **FN1** gene was highly expressed in **TC-Mφ**. Alternative splicing within human FN pre-mRNA results in the generation of ~20 distinct isoforms (56–59).Because specific isoforms of FN are known to contribute to cancer pathogenesis (60, 61), we decided to further investigate their possible expression in **TC-Mφ**.

Using primers specific for distinct domains, we tested in **TC-Mφ** the expression of four FN isoforms: full-length FN, oncofetal ED-A FN, oncofetal ED-B FN, and MSF. We found that the most abundant FN mRNA transcripts expressed by **TC-Mφ** were the full-length isoform and the MSF (Fig. 4A). MSF differs from the other two oncofetal full-length isoforms because it is identical to their 70-kDa N terminus but terminates in a unique 10 aa sequence, and it has been defined as a truncated isoform (62). Importantly, MSF was known to be produced by fibroblasts, fetal skin keratinocytes, tumor cells, and tumor-associated vascular endothelial cells (62) but not by Mφ. We observed an opposite trend over time for the MSF expression compared with full-length FN. The **TC-Mφ** MSF mRNA level increased along with the time of culture and the number of supernatant stimuli, whereas transcripts from the full-length FN decreased (Fig. 4B, 4C).

Next, we investigated whether MSF was preferentially associated with the classically (M1) or alternative (M2) Mφ polarization. By analyzing 6-d in vitro rhM-CSF–differentiated Mφ, further polarized with LPS/IFN-γ or IL-4, it was evident that MSF expression was higher in **M2-Mφ** compared with M1 cells (Fig. 4D). Moreover, MSF expression was specifically associated with M2 cells because its levels were dramatically downregulated in M2 cells reverted into M1 (with LPS/IFN-γ) and, on the contrary, strongly upregulated in M1 cells skewed toward the M2 phenotype (with IL-4) (Fig. 4D). Higher MSF production in **M2-Mφ** was confirmed at protein level by Western blot as shown in Fig. 4E.

To confirm MSF expression by tumor-infiltrating leukocytes in vivo, we performed RT-PCR with purified ex vivo preparations of TAM. Four different preparations isolated from human ovarian carcinoma specimens expressed MSF transcripts (Fig. 5A). In addition, immunohistochemical analysis of tumor sections showed that immunoreactivity for anti-MSF was detected in CD68+ cells infiltrating human tumors (Fig. 5B).

To better understand the regulation of MSF in myeloid cells, we stimulated human monocytes with several cytokines and growth factors for 24 h. Interestingly, we noticed unequivocally different effects between inflammatory and anti-inflammatory stimuli. As shown in Fig. 6A, M-CSF, IL-4, and especially TGFβ strongly promoted MSF production, whereas LPS, TNF-α, and IFN-γ did not. As TGFβ was almost undetectable in our tumor-CM, we checked the effect of the natural tumor-derived M-CSF. Pretreatment of monocytes with anti–M-CSF Ab almost completely abrogated MSF expression, indicating that, in our experimental conditions, M-CSF is
the main tumor-derived factor able to induce MSF expression in monocytes (Fig. 6).

**Functional characterization of MSF**

It has been reported that MSF has chemotactic activity for fibroblasts and tumor cells through its Gel-BD domain. We confirmed these results with the tumor cell line Panc1 and human monocytes (Fig. 7A, 7B). Of note, MSF showed its maximal chemotactic activity at picomolar concentrations; whereas monocyte migration peaked at 1 to 2 ng/ml, tumor cells responded at 10-fold less MSF concentration.

We next determined the chemotactic activity of MSF released by TC-Mφ and M2-polarized Mφ. Fig. 7C demonstrates that supernatants from both Mφ populations, but not from M1 cells, did stimulate tumor cell migration. Inhibition of MSF with a specific anti-MSF Ab directed to the Gel-BD (62) reduced the number of migrated tumor cells, thus demonstrating the specific involvement of MSF (Fig. 7D).

**Discussion**

The major focus of this study was to clarify how the malignant context is able to influence the fundamental initial phase of monocyte to Mφ differentiating activity, which was totally due to tumor-derived M-CSF–secreted isoforms.

To identify novel molecules highly expressed in TC-Mφ, we performed a gene expression analysis of these cells by Affymetrix technology, and we next compared these results to three other datasets of M1, M2, and TAM isolated from human tumors. Globally, the comparison of the four datasets confirmed that TC-Mφ are indeed comparable to TAM and resemble more M2-Mφ than M1-polarized cells. Among several common upregulated genes are the mannose receptor, other members of the C-type lectin R family, receptors for the Fc fragment of IgG, and receptors importantly implicated in the uptake and scavenging of debris, as well as several adhesion molecules.

In TC-Mφ, the most expressed gene was **SEPP1**, a plasma transporter of selenium that has been implicated in the protection of tissues and in particular of endothelial cells from oxidative damage (63). In a recent study, SEPP1 was reported to be essential to limit disease severity of African trypanosome infection through its antioxidant activity (64). Within the tumor microenvironment, the presence of Mφ-derived SEPP1 may protect tumor vessels, cancer cells, and Mφ themselves from damage and consequently support tumor survival and growth. Our analysis shows that SEPP1 is highly expressed at the RNA level by M2-polarized Mφ and TAM, but not by M1 cells, indicating that SEPP1 is a putative good marker of M2 cells. SEPP1 expression

**FIGURE 4.** Expression of MSF in TC-Mφ. A, mRNA level expression of full-length FN1, ED-A FN, ED-B FN, and MSF. MSF and full-length FN1 in TC-Mφ at different times of culture (days 0, 3, 6) (B) and upon repeated stimulation (days 1 and 5) with tumor-CM (C). D, MSF mRNA levels in Mφ polarization (white bars). M1 stimuli: LPS + IFN-γ; M2 stimulus: IL-4. Black bars indicate MSF expression in reverted Mφ treated as indicated. E, Western blot analysis of MSF production in TC-Mφ, M-CSF-Mφ, M1-, and M2-polarized Mφ. **p < 0.01.

**FIGURE 5.** Expression of MSF in human TAMs. A, MSF mRNA expression in TAMs. Mean of four different donors. B, Immunohistochemistry of MSF (brown) in CD68+ (blue) TAMs from pancreatic or ovarian tumors; nuclei are green. Arrows indicate CD68+MSF+ cells. C, Immunohistochemistry of MSF (brown) in CD68+ (blue) TAM from pancreatic tumors (original magnification ×100). Nuclei are red. Arrows indicate CD68+MSF+ cells. *MSF-positive tumor cells.
is reduced in IL-10 knockout mice, indicating a role for IL-10 in its regulation (64).

Several upregulated genes were related to the ECM organization and turnover. ECM proteins are continuously produced and degraded by many cell types (tumor, endothelial, and stromal cells), among which TAM are master regulators. The incessant and dysregulated remodeling of the ECM in tumors leads to the aberrant presence of some matricellular components that are not usually found in normal tissues. In TC-Mφ, we identified upregulated transglutaminase F13A1, which catalyzes the polymerization of several matrix proteins, OA, OPN, FN1, and proteolytic enzymes.

OA is a poorly characterized transmembrane molecule having supposed functions in cell adhesion, migration, and differentiation. Its expression has been linked to the upregulation of MMP3 and MMP9 in transformed astrocytes and fibroblasts (65, 66). In OA-transgenic mice, the molecule showed a cytoprotective effect on fibrosis induced by skeletal muscle denervation (67). These findings allow the hypothesis that OA could be involved in the pathophysiological cascade of tissue injury and repair. In addition, uremic Mφ in patients with end-stage renal disease have been recently demonstrated to exhibit increased levels of this protein (68). OA was found overexpressed in various malignant tumors (65, 69, 70); moreover, it has been described that overexpression of OA in glioma and hepatoma cell lines sustains tumor invasiveness (65, 71). OA has an Arg-Gly-Asp integrin-binding domain; in mouse dendritic cells, OA enhances adhesion and transendothelial migration (72). The role of OA in myeloid cells infiltrating the tumor microenvironment is still unknown, but it could be speculatively linked to their efficient tissue remodeling function and MMP activation.

OPN has long attracted the interest of immunologists because of its many functions in inflammation, immunopathology, and hematopoiesis (73–78). Its expression is increased in response to cell injury or infections and is induced by a variety of proinflammatory mediators and growth factors such as IL-1, TNF-α, and platelet-derived growth factor (79). Elevated levels of OPN are associated with a remarkable variety of pathological processes ranging from atherosclerosis, autoimmune diseases, and cancer (74, 76, 79–81). A correlation between elevated OPN expression and malignant invasion (82–89) was suspected because OPN controls tumor cell motility and invasion through the engagement of CD44 receptors (90–92). Stromal cells and fibroblasts in particular are known to secrete OPN around tumor cell nests. Our study indicates that OPN was also highly expressed by TC-Mφ.

We found FN upregulation in TC-Mφ of special interest. FN is the most well-studied matrix protein, and it is known that in...
transformed cells and in malignancies, the splicing pattern of FN pre-mRNA becomes altered, leading to an increased expression of oncofetal isoforms (93–95). The ED-A (59) and ED-B (96–98) are well-known oncofetal isoforms of FN. They are highly expressed in many tumors and promote cell spreading and angiogenesis (61, 93, 99). In particular, ED-B (96) is produced during active tissue remodeling like wound healing and, in tumor angiogenesis processes, gives rise to a prominent perivascular expression pattern (58).

A third oncofetal FN isoform, known as MSF, has been cloned in 2003 by Schor and colleagues. MSF differs from the other two oncofetal full-length isoforms because it is identical to their 70-kDa N terminus but terminates in a unique 10 aa sequence. Thus, MSF has been defined as truncated isoform (62). Unlike the angiogenic ED-A and ED-B, MSF was first identified by its motogenic activity in the CM of cultured fetal fibroblasts (100, 101). In this study, we show that TC-Mφ produce MSF, whereas the ED-A and ED-B isoforms are not significantly expressed. MSF was known to be produced by fetal skin keratinocytes, neoplastic cells, and tumor-associated vascular endothelial cells (62), but has never been reported in Mφ. It is also expressed during wound healing and strongly overexpressed in a wide range of common human cancers (62, 102). rMSF displays a number of potent bioactivities relevant to cancer development processes, including stimulation of cell migration, hyaluronan synthesis (103, 104), proteolytic activity (105), and angiogenesis (62, 102).

Of note, we observed an opposite trend over time for the MSF expression compared with full-length FN. TC-Mφ–derived MSF increased along with either the time of culture or the number of supernatant stimuli, whereas transcripts from the full-length FN decreased. These data suggested a strict influence by the tumor microenvironment on FN isoform production. Importantly, MSF expression is upregulated in M2 compared with M1-Mφ and MSF RNA and protein (immunohistochemistry) were found also in human TAMs. Thus, MSF may represent a good candidate marker of M2-Mφ polarization and TAM.

In regards to its functions, due to its Gel-BD motogenic site, MSF had strong chemotactic activity for monocytes and tumor cells. Interestingly, this bioactivity is not expressed by full-length FN. It is known in fact that some biological activities of FN are acquired by denaturation and/or proteolytic cleavage. Schor et al. have reported that the Gel-BD domain is cryptic in full-length FN and acquires potent motogenic activity postproteolysis (62). Thus, two independent mechanisms for locally generating the same migratory activity are available in tissues: one by degradation of the extracellular matrix (Gel-BD) and the other by genetic control of FN truncation during gene transcription (MSF). The generation of MSF via a genetic mechanism is likely more efficient and does not rely on the activation of proteolytic enzymes; in addition, some cleaved FN fragments (e.g., Cell-BD) also have potent inhibitory effects on Gel-BD and MSF motogenic activity. We have shown in this study that MSF-containing supernatants of TC-Mφ induced migration of both monocytes and tumor cells, underlining autocrine regulatory loops of leukocyte recruitment as well as paracrine effects on cancer cells. The observation that Mφ secrete a chemotactic molecule that induces the migration of tumor cells highlights a new protumoral mechanism of these cells, already implicated in the process of distant metastasis and more recently in the preparation of the metastatic niche (23, 24, 106).

Another relevant consideration is MSF as therapeutic target. It was recently reported that the in vivo biodistribution of a labeled anti-MSF Ab was selectively concentrated in tumor vessels in a mouse model of xenograft esophageal carcinoma (107). Interestingly, administration of the anti-MSF Ab to tumor-bearing mice significantly reduced tumor growth, thus indicating that targeting of MSF in tumors may have therapeutic potential (107). Up to now, Mφ markers with prognostic utility detectable in biological fluids of patients with cancer have been almost ignored. Of interest, MSF has been measured in a pilot study and detected in the sera of the majority of breast cancer patients but not of healthy subjects (101, 108).

In conclusion, we have described in this study that, in terms of gene profiling, TC-Mφ are strongly comparable to TAMs and resemble M2-like polarized cells, thus confirming and boosting the concept of an M2-like polarization of tumor Mφ. Among some interesting upregulated molecules, we focused on MSF, a truncated FN with motogenic activity. We suggest it as a new specific marker for M2-polarized Mφ and TAM that might be considered in the design of anti-Mφ approaches for novel anti-cancer therapies. Further studies are needed to elucidate the involvement of MSF in the protumoral role of Mφ and as a biomarker in patients with cancer.

Acknowledgments

We thank Profs. Ana and Seth Schor (University of Dundee, Scotland, U.K.) for kindly providing recombinant human MSF and anti-MSF Abs. We also thank Dr. Giuseppe Peri for support and assistance during our anti-MSF Ab production and Dr. Federica Marchesi (Istituto Clinico Humanitas, Milan, Italy) and Dr. Fernando O. Martinez (University of Oxford, Oxford, U.K.) for help and suggestions.

Disclosures

The authors have no financial conflicts of interest.

References


