Heterocomplexes between the atypical chemokine MIF and the CXC-motif chemokine CXCL4L1 regulate inflammation and thrombus formation

Markus Brandhofer1, Adrian Hoffmann1,2, Xavier Blanchet3, Elena Siminkovitch1, Anne-Katrin Rohlfing1, Omar El Bounkari1, Jeremy A. Nestele4, Christos Kontos5, Kathleen Hille5, Vanessa Rohde1, Adrian Fröhlich1, Jona Golem6, Özgün Gokce6,7, Christine Krämer1, Patrick K. Otabil1, Nikolaos Tsilimparis8, Nadja Sachs9,10, Wolfgang E. Kempf9, Lars Maegdefessel9,10, Michael K. Otabil1, Remco T. A. Megens3,10,11, Hans Ippel11, Rory R. Koenen11, Junfu Luo12, Benoit Engemann12, Kevin H. Mayo11,13, Meinrad Gawaz4, Aphrodite Kapurniotu5, Christian Weber3,7,11,10, Philipp von Hundelshausen3,10, Jürgen Bernhagen1,7,10

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Abstract
To fulfill its orchestration of immune cell trafficking, a network of chemokines and receptors developed that capitalizes on specificity, redundancy, and functional selectivity. The discovery of heteromeric interactions in the chemokine interactome has expanded the complexity within this network. Moreover, some inflammatory mediators, not structurally linked to classical chemokines, bind to chemokine receptors and behave as atypical chemokines (ACKs). We identified macrophage migration inhibitory factor (MIF) as an ACK that binds to chemokine receptors CXCR2 and CXCR4 to promote atherogenic leukocyte recruitment. Here, we hypothesized that chemokine–chemokine interactions extend to ACKs and that MIF forms heterocomplexes with classical chemokines. We tested this hypothesis by using an unbiased chemokine protein array. Platelet chemokine CXCL4L1 (but not its variant CXCL4 or the CXCR2/CXCR4 ligands CXCL8 or CXCL12) was identified as a candidate interactor. MIF/CXCL4L1 complexation was verified by co-immunoprecipitation, surface plasmon-resonance analysis, and microscale thermophoresis, also establishing high-affinity binding. We next determined whether heterocomplex formation modulates inflammatory/atherogenic activities of MIF. Complex formation was observed to inhibit MIF-elicited T-cell chemotaxis as assessed by transwell migration assay and in a 3D-matrix-based live cell-imaging set-up. Heterocomplexation also blocked MIF-triggered migration of microglia in cortical cultures in situ, as well as MIF-mediated monocyte adhesion on aortic endothelial cell monolayers under flow stress conditions. Of note, CXCL4L1 blocked binding of Alexa-MIF to a soluble surrogate of CXCR4 and co-incubation with CXCL4L1 attenuated MIF responses in HEK293-CXCR4 transfectants, indicating that complex formation interferes with MIF/CXCR4 pathways. Because MIF and CXCL4L1 are platelet-derived products, we finally tested their role in platelet activation. Multi-photon microscopy, FLIM-FRET, and proximity-ligation assay visualized heterocomplexes in platelet aggregates and in clinical human thrombus sections obtained from peripheral artery disease (PAD) in patients undergoing thrombectomy. Moreover, heterocomplexes inhibited MIF-stimulated thrombus formation under flow and skewed the lamellipodia phenotype of adhering platelets. Our study establishes a novel molecular interaction that adds to the complexity of the chemokine interactome and chemokine/receptor-network. MIF/CXCL4L1, or more generally, ACK/CXC-motif chemokine heterocomplexes may be target structures that can be exploited to modulate inflammation and thrombosis.

keywords Heterodimer · protein-protein interaction · platelets · leukocytes · thrombosis · chemotaxis

# M.B. and A.H. contributed equally to the study.

Extended author information available on the last page of the article

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Introduction

Chemokines orchestrate immune cell trafficking in health and disease [1, 2, 3]. Chemokine-directed targeting strategies are pursued in acute and chronic inflammatory conditions, autoimmunity, cancer, and atherosclerosis [2, 4, 5]. The chemokine network encompasses 49 classical chemokines (CKs) and 18 classical chemokine receptors (CKRs), which belong to the class of G protein-coupled receptors (GPCRs) [6, 7, 8]. Depending on the particular chemokine ligand/receptor pair and various disease and microenvironmental factors, chemokine signaling through CKRs overall capitalizes on the principles of specificity, promiscuity, and biased agonism. Accordingly, multiple chemokines can bind to a certain chemokine receptor and vice versa, while ‘biased agonism’ can occur on a ligand, receptor, or tissue basis [9, 10, 11]. Fine-tuning of chemokine responses within this network is further expanded by five atypical chemokine receptors (ACKRs) that serve as decoy receptors and promiscuously bind many chemokines to shape their gradients, but also elicit specific signaling responses [12].

Chemokines are well-known to form homodimers, but the discovery of the chemokine interactome additionally suggested a multitude of heteromeric chemokine-chemokine interactions even across CC- and CXC-chemokine class borders [13, 14]. CC-type heterodimers between CCL5 and CCL17 or CCL5 and CXCL4 (also termed platelet factor 4, PF4) were found to lead to functional synergism by receptor retention or auxiliary proteoglycan binding and enhancement of chemotactic responses, respectively, while CXC-type heterodimers between CXCL12 and CCL5 or CXCL12 and CXCL4 led to signaling inhibition. This has demonstrated yet another level of complexity within the chemokine network and offers novel intervention strategies in inflammatory and cardiovascular diseases [13].

Moreover, some alarmin-like inflammatory mediators such as human β-defensins (HBDs) and secreted fragments of amino acyl tRNA-synthetases (AARSs), which do not belong to one of the four structural classes of CC-, CXC-, CX₃C₄, or C-chemokines, can bind to chemokine receptors by molecular mimicry and exhibit chemokine-like activities [15, 16]. These proteins are also referred to as atypical chemokines (ACKs) [17, 18, 19].

Macrophage migration inhibitory factor (MIF) is an evolutionarily conserved pleotropic inflammatory cytokine [20, 21]. MIF is an upstream regulator of the host innate immune response and, when dysregulated, is a pivotal mediator of inflammatory diseases, autoimmunity, cancer, and cardiovascular diseases [22, 23]. MIF is a structurally unique cytokine [24] and, contrary to its eponymous name, has chemokine-like activities and functions as a prototypical ACK [17, 25]. Accordingly, MIF not only signals through its cognate receptor CD74/invariant chain, but engages in high-affinity interactions with the CXC chemokine receptors CXCR2 and CXCR4 to promote atherogenic monocyte and T-/B-cell recruitment, cancer metastasis, and inflammation [17, 25, 26, 27, 28, 29, 30]. We elucidated the structural determinants of the binding interface between MIF and its CXC-motif chemokine receptors and found that MIF mimics chemokine receptor binding regions such as the ELR motif and the N-loop [31, 32, 33, 35]. Interestingly, CXCL12/SDF-1α (stromal-derived factor-1α), the cognate ligand of CXCR4, was recently found to bind to the non-chemokine proteins galectin-3 [36] and high-mobility group box-1 (HMGB1) [37, 38], but potential interactions between MIF and CXCL12 or CXCL8, the cognate ligand of its chemokine receptors CXCR2, have remained unclear.

Here, we hypothesized that chemokine–chemokine interactions are not only possible between different types of classical chemokines, as demonstrated by chemokine interactome mapping [13], but might extend to ACKs. Choosing MIF as a prototypical ACK, we thus asked whether this mediator would form heterocomplexes with classical chemokines. We tested this hypothesis applying an unbiased chemokine protein array and validated candidate interactors by a battery of biochemical and biophysical methods. We identified the platelet chemokine CXCL4L1 (also termed PF4var1), but not its variant CXCL4, nor the CXCR2 ligand CXCL8 or the CXCR4 ligand CXCL12, as a high affinity interactor of MIF and tested the potential functional role of MIF/CXCL4L1 heterocomplex formation for MIF binding to its receptor CXCR4, and in cell systems that are relevant for the inflammatory, atherogenic, and thrombogenic activities of MIF. Finally, we also asked whether such heterocomplexes can be detected in clinical thrombus specimens from patients with atherosclerosis. Our study extends the chemokine interactome to ACK/CK interactions and demonstrates a functional role for the MIF/CXCL4L1 heterocomplex in disease-relevant activities.

Materials and methods

Proteins and reagents

Biologically active and endotoxin-free recombinant human MIF was prepared as previously described and was obtained at a purity of ~98% as confirmed by SDS-PAGE analysis in combination with silver staining [39, 40]. For the preparation of Alexa Fluor-488- and MST-Red-labeled MIF, a 90–95% pure MIF fraction was used. Alexa Fluor-488-labeled MIF was generated using the Microscale Protein Labeling Kit from Invitrogen-Molecular Probes (Karlsruhe, Germany) and MST-Red-MIF was prepared using the Monolith Protein Labeling Kit RED-NHS second Generation from
NanoTemper (Munich, Germany), following the manufacturers’ instructions. Biotinylated human MIF was produced using d-biotinoyl-e-aminoacproic acid-N-hydroxy-succinimide ester (Biotin-7-NHS) with the Biotin Protein Labe-
ling Kit from Roche (Mannheim, Germany). Alternatively, biotin-amidoheaxanoic acid N-hydroxysuccinimide ester from Sigma-Aldrich (Taufkirchen, Germany) was used.

For the fluorescence polarization assay, a hexahis-
dine-tagged variant of CXCL4L1 was used. Briefly, the coding sequence of human CXCL4L1 with a methionine-
flanked N-terminal His6-tag was cloned into the pET21a vector for recombinant bacterial expression using XhoI and NdeI restriction sites. This construct was then used to transform Rosetta-gami™ 2 (DE3) competent E. coli (Novagen, Merck KGaA, Darmstadt, Germany) for subsequent recombinant protein production following induction with 1 mM IPTG (Carl Roth, Karlsruhe, Germany) according to a standard protocol. For purifi-
cation, bacteria pellets were resuspended in lysis buffer (20 mM Tris–HCl, pH 8.0, 0.5 M NaCl, 5 mM EDTA, 0.1% Triton X-100, with added protease inhibitor tablets according to manufacturer’s instructions) and cells dis-
rupted in an EmulsiFlex-C5 high pressure homogenizer (Avestin Europe GmbH, Mannheim, Germany), the raw extract cleared via centrifugation at 20,000×g and the resulting pellet, containing recombinant His-CXCL4L1 in inclusion bodies, was washed in lysis buffer with and without detergent. Inclusion bodies were solubilized by gentle shaking overnight in 50 mM Tris–HCl, pH 8.0, 6 M guanidine-HCl, 0.5 M NaCl, 10 mM DTT and His-
CXCL4L1 purified from the solubilized pellet via IMAC on a HisTrap HP column on an ÄKTA Pure 25 M FPLC system (Cytiva Europe GmbH, Freiburg, Germany). The obtained protein was subjected to two dialysis steps in refolding buffer (50 mM Tris–HCl, pH 8.0, 0.5 M NaCl, 6 mM methionine, 5 mM cysteine) with and subsequently without 0.9 M guanidine-HCl, followed by a final purifi-
cation step by size exclusion chromatography in 20 mM sodium phosphate buffer, pH 7.4, using a Superdex 75 10/300 GL column (Cytiva Europe GmbH) on an ÄKTA Pure 25 M FPLC system. A purity degree of 90–95% was verified by SDS-PAGE followed by Coomassie staining and Western Blot according to standard protocols.

Recombinant human peroxiredoxins 1 and 6 (PRX1, PRX6) were purchased from Abcam (Abcam PLC, Cam-
bridge, UK), while recombinant human β-defensin-1 and 2 (HBD-1, HBD-2) were obtained from ProSpec (ProSpec-Tany TechnoGene Ltd., Ness Ziona, Israel). Recombinant human HMGB1 was purchased from Novus (Novus Biologicals Europe, Abingdon, UK). Recombi-
nant human CXCL4L1 (PF4var1) as well as the CXCL4 (PF4) were purchased from ChromaTec (Greifswald, Germany). The other recombinant human chemokines were obtained from Peprotech (Hamburg, Germany). All other reagents and chemicals were purchased from Merck KGaA (Darmstadt, Germany), Carl Roth GmbH (Karslruhe, Germany), or Sigma-Aldrich and were of the highest purity degree available.

**Cell culture and cultivation of mammalian cell lines**

Jurkat T cells were cultured in RPMI1640 medium (Gibco) supplemented with 10% fetal calf serum (FCS), 1% peni-
cillin/streptomycin, and 1× non-essential amino acids (NEAAs, Gibco). The human monocytic cell line Mono-
Mac6 [41] was cultured in RPMI1640 medium + GlutaMAX (1×), supplemented with 1× NEAAs, 10% FCS, and 1% penicillin/streptomycin. Human aortic endothelial cells (HAoECs) were obtained from PromoCell (Heidelberg, Ger-
many) and cultured according to the supplier’s instructions in collagen-coated (collagen type I; EMD Millipore Corp., Darmstadt, Germany) cell culture flasks in Endothelial Cell Basal Medium (EBM®-2, Lonza, Verviers, Belgium), supplemented with 1% penicillin/streptomycin (Gibco, Karlsruhe, Germany) and EGTM™-2 SingleQuots® supple-
ments (Lonza). HEK293 cells stably transfected with human CXCR4 (HEK293-CXCR4) were used at passage 5 and were cultivated in DMEM medium (Gibco), supplemented with 10% FCS and 1% penicillin/streptomycin (Gibco), and used for the experiment between passage 6 and 8.

Unless stated otherwise, cells were cultivated in a tem-
perature- and humidity-controlled incubator at a temperature of 37 °C and 5% CO2. FCS from an EU-approved origin was obtained from Invitrogen-Thermo Fisher Scientific and heat-inactivated prior to usage. Other cell culture rea-
nants, media and supplements were bought from Invitro-
Gen-Thermo Fisher Scientific, unless stated otherwise. Cell lines were originally obtained from the German Society for Microorganisms and Cell Cultures (DSMZ, Braunschweig, Germany) or from the American Type Culture Collections (ATCC).

**Isolation of primary human CD4+ T cells**

Primary human CD4-positive T cells were isolated from enriched peripheral blood mononuclear cell (PBMC) frac-
tions using the human CD4+ T cell isolation kit from Miltenyi Biotec (Bergisch Gladbach, Germany) according to the manufacturer’s instructions. Cells were cultivated in RPMI1640 medium, supplemented with 10% FCS, 1% peni-
cillin/streptomycin, and 1× NEAAs in a cell culture incu-
bator at 37 °C and 5% CO2 and used for functional assays on the next day. PBMC fractions were obtained by apher-
esis from conical chambers of a Leucoreduction System Chamber sourced from anonymous platelet donations at the
Department of Transfusion Medicine, Cell Therapeutics and Hemostaseology of LMU University Hospital. Studies abide by the Declaration of Helsinki principles and were approved by ethics approval 18-104 of the Ethics Committee of LMU Munich, which encompasses the use of anonymized tissue and blood specimens for research purposes.

Isolation of human platelets

For immunofluorescent stainings

Human platelets were isolated from blood, freshly drawn from healthy donors, using a syringe containing 1/10 volume of CTAD–buffer (0.105 M tri-sodium citrate, 10 mM theophylline, 3.7 mM adenosine, 0.198 mM dipyridamole) [42]. To prevent platelet activation, the blood was supplemented with prostaglandine E1 (Merck KGaA), Apyrase (New England Biolabs GmbH, Frankfurt am Main, Germany), and EGTA (Sigma-Aldrich). Briefly, platelets were isolated by sequential centrifugation steps, performed at room temperature (RT) with reduced brake settings. Platelet-rich plasma (PRP) was separated from whole blood by centrifugation for 5 min at 300×g to remove remaining leukocytes. Finally, platelets were sedimented by centrifugation for 10 min at 200×g. Washed human platelets were isolated as previously described [43] and subsequently used for functional flow chamber or platelet spreading assays.

Mice and preparation and cultivation of primary mixed cortical cultures for the microglia motility assay

CX3CR1GFP/+ mice, which were originally obtained from the Jackson Laboratories (strain 005,582; [44]), were established on a pure C57BL/6 background and housed under standardized light–dark cycles in a temperature-controlled air-conditioned environment under specific pathogen-free conditions at the Center for Stroke and Dementia Research (CSD), Munich, Germany, with free access to food and water. Animals were sacrificed under anesthesia with a mixture of midazolam (5 mg/mL), medetomidine and fentanyl (MMF). Mouse maintenance and experiments were reviewed and overseen by the institutional animal use and care committee of the local authorities (Regierung von Oberbayern, ROB, Germany) and performed in accordance with the procedures provided by the animal protection representative of CSD.

Primary mixed cortical cultures containing CX3CR1GFP/+ microglia were prepared in 96-well imaging plates based on a previously established protocol [45] from the cortices of 5 newborn pups of the CX3CR1GFP/+ mouse line (postnatal day 0) in plating medium, consisting of modified Minimum Essential Medium (MEM without glutamine and phenol red) (Gibco), supplemented with 0.5% glucose, 0.02% sodium bicarbonate, 1 × ITS-supplement (Sigma-Aldrich), 2 mM L-glutamine (Gibco), 1% penicillin/streptomycin, and 10% FCS. Cultures were incubated in a humidified atmosphere at 37 °C and 5% CO2 for 10 d. One day after plating, 80% of the plating medium was replaced with growth medium, prepared from MEM (without glutamine and phenol red) supplemented with 0.5% glucose, 0.02% sodium bicarbonate, 5% FCS, 0.5 mM L-glutamine, and serum-free B-27™ supplement (Gibco). On the fourth day after dissection, 50% of the medium was replaced with growth medium additionally supplemented with 4 µM cytosine-1-β-d-arabinofuranoside (Sigma-Aldrich).

Chemokine protein array

Human chemokines and selected atypical chemokines were spotted on a nitrocellulose membrane at 100 ng per spot and left to dry at RT. Membranes were blocked with 1 × ROTI®Block (Carl Roth) for 2 h at RT and then probed overnight with biotinylated human MIF (biotin-MIF) at a concentration of 1 µg/mL in either 10 mM Tris–HCl pH, 8.0 or 10 mM MES, pH 6.0. Subsequently, membranes were washed three times with 0.01% Tween®20 in water and developed with horseradish-peroxidase (HRP)-conjugated streptavidin (Bio-Techne GmbH, Wiesbaden-Nordenstadt, Germany), diluted 1:200 in 1 ×  ROTI®Block, for 2 h. After another washing step, bound biotin-MIF was revealed via chemiluminescence using SuperSignal™ West Pico Chemiluminescent Substrate (Thermo Fisher Scientific) on a LAS-3000 Imaging System (Fuji Photo Film Co., LTD., Japan).

Pull-down of CXCL4L1 from cell lysates and verification of antibody specificity

MonoMac-6 cells were first washed with PBS and then lysed on ice for 30 min with immunoprecipitation (IP) lysis buffer (1 × cell lysis buffer, Cell Signaling), 100 mM PMSF, and 1 × protease and phosphatase inhibitors (ThermoFisher). The purified cell lysates were then incubated with pre-washed streptavidin-conjugated paramagnetic beads (DYNAL™ Dynabeads M-280 Streptavidin; Invitrogen) for 2 h at 4 °C (preclearing step). After centrifugation, the supernatant was incubated with biotinylated human MIF by gentle, constant shaking on a rotary shaker overnight at 4 °C. To capture MIF/CXCL4L1 complexes, prewashed streptavidin-conjugated beads were added to the precleared lysates,
and the mixture was incubated for 2 h at 4 °C on a rotary shaker. Beads were separated from the lysate using a magnetic stand (Dynal™ MCP-S) and washed three times with lysis buffer. The supernatant was removed and the beads were resuspended in LDS sample buffer (Invitrogen), containing 100 mM dithiothreitol (DTT), and boiled at 95 °C for 15 min. Samples were subjected to SDS-PAGE and analyzed by Western blotting. For this purpose, equal amounts of protein were loaded onto 11% SDS–polyacrylamide gels (NuPAGE, ThermoFisher) and transferred to polyvinylidene difluoride (PVDF) membranes (Carl Roth, Karlsruhe, Germany). Membranes were blocked in PBS-Tween-20 containing 5% BSA for 1 h and incubated overnight at 4 °C with rabbit polyclonal anti-MIF antibody Ka565 [25] or rabbit polyclonal anti-PF4V1 IgG PA5-21944 (Invitrogen) diluted in blocking buffer. Proteins were revealed using anti-rabbit HRP as a secondary antibody. Signals were detected by chemiluminescence on an Odyssey® Fc Imager (LI-COR Biosciences GmbH). Imaging of the membrane was performed using a Biacore X100 instrument (GE Healthcare Europe Hamburg, Germany) or CXCL4L1 (ChromaTec, Greifswald, Germany) (final MIF concentrations: 456 nM or 312 nM, respectively). Prior to measurement, the prepared samples were incubated for at least 30 min on ice. MST traces of multiple experiments were analyzed according to the K_D model using the default T-jump settings, focusing on the temperature related intensity change (TRIC) of the fluorescent label (“cold region” from −1 to 0 s, “hot region” from 0.5 to 1.5 s) using the MO. Affinity Analysis V2.3 software (NanoTemper Technologies). Curve fitting for data presentation was performed by GraphPad Prism Version 6.07 ("one site—total binding").

**CelluSpot peptide array**

The CelluSpot antibody array method has been described previously [33]. Briefly, 15-meric peptides, positionally frame-shifted by three residues and spanning the entire sequence of CXCL4L1 and CXCL4 (both obtained from ChromaTec) were prepared in 200 µL of PBS and loaded onto a nitrocellulose membrane using a slot blot device (Whatman™ Minifold I, Cytiva). After blocking for 1 h in TBS-Tween-20 containing 1% BSA, the membrane was probed with anti-PF4V1 PA5-21944 diluted in blocking buffer overnight at 4 °C. Bound antibody was detected by incubation for 2 h at RT with a fluorescently labeled donkey anti-rabbit IRDye 800 CW antibody (LI-COR Biosciences GmbH, Bad Homburg, Germany). Imaging of the membrane was performed on an Odyssey® Fc Imager (LI-COR Biosciences GmbH).

**Analysis of protein–protein interactions by surface plasmon resonance (SPR)**

Surface plasmon resonance measurements were performed using a Biacore X100 instrument (GE Healthcare Europe GmbH) and neutravidin-modified C1 sensor chips. Biotin-MIF was immobilized on flow cells to 1064.8 RU. CXCL4 (Peprotech, Hamburg, Germany) and CXCL4L1 (ChromaTec, Greifswald, Germany), used at concentrations in the range of 0.125 to 20 µg/mL in running buffer (HBS-EP + Buffer: 0.01 M HEPES, 0.15 M NaCl, 0.003 M EDTA and 0.05% v/v surfactant P20) were injected at a flow rate of 60 µL/min. The complex was allowed to associate and dissociate for 90 s and 240 s, respectively. Surfaces were regenerated with 2 pulses (60 s) of 30 mM NaOH and 2 M NaCl. Responses from analyte injections were fitted to a 1:1 Langmuir interaction profile using Biacore X100 evaluation 2.0.1 Plus package software.

**Fluorescence spectroscopy**

Fluorescence spectroscopic titrations were carried out using a JASCO FP-6500 fluorescence spectrophotometer...
a 12 h migration interval at 37 °C and 5% CO₂, migrated were added to the lower chamber as a chemoattractant. After 32 nM of CXCL4L1, as well as 32 nM CXCL4L1 alone preincubated (30 min on ice to allow for complex formation) with slatern, Germany). 16 nM MIF, either alone or pre-incubated Transwell insert with 5 μm pore size (Corning, Kaiser- mLeica DMi8 inverted microscope (Leica Microsystems, Wetzlar, Germany) and Leica live cell-imaging software (LAS X version 3.7.4). Images were imported as stacks to ImageJ version 1.51n and analyzed with the manual tracking and Chemotaxis and Migration tool (Ibidi GmbH) plugin for ImageJ.

Motility measurement of primary murine microglia

The motility of mouse microglia was determined using mixed cortical cultures, established and cultivated as stated above. A day prior to imaging, the medium was changed to Hibernate A medium (Gibco) in order to maintain a physiological pH value during imaging. Prior to imaging, different wells of cells from each individual pup were treated with either 8 nM MIF, 1.6 nM CXCL4L1, or both (pre-incubated for 30 min on ice to allow for complex formation). A control group was treated with 20 mM sodium phosphate buffer, pH 7.4. Cell motility was monitored by time-lapse imaging for 15 h at 37 °C with recordings every 5 min, using a Leica DMi8 inverted microscope (Leica Microsystems, Wetzlar, Germany) and Leica live cell-imaging software (LAS X version 3.7.4). Images were imported as stacks to ImageJ software version 1.51n and analyzed with the manual tracking and Chemotaxis and Migration tool Plugin for ImageJ from Ibidi. In order to quantify microglial motility from the time-lapse videos, 20–25 GFP-positive microglia per treatment group were randomly selected and manually tracked throughout all frames. Cells that died or moved out of the frame were excluded from the analysis. Accumulated distance of each tracked microglia was calculated with Chemotaxis and Migration Tool (Ibidi).

Monocyte adhesion under flow conditions

Human aortic endothelial cells (HAoECs) were seeded at a density of approximately 70,000 cells per channel in collagen-coated Ibidi μ-Slides I 0.8 and incubated until monolayers were confluent. The flow adhesion assay was performed essentially as described [40]. Briefly, HAoECs were treated with 16 nM human MIF, 32 nM CXCL4L1, or a mixture of both for 2 h. MonoMac6 cells, fluorescently labeled using the CFSE Cell Division Tracker Kit (BioLegend Europe BV, Amsterdam, Netherlands) according to manufacturer’s instructions, were exposed to human MIF and/or CXCL4L1 at the above concentrations for 2 min, before they were transferred into assay buffer (1 × HBSS, 10 mM HEPES, 0.5% BSA) at a density of 0.5 × 10⁶ cells/mL and kept at 37 °C.

Transwell migration assay

Transwell migration experiments to study the influence of CXCL4L1 on MIF-mediated chemotaxis responses were performed with Jurkat T cells. Briefly, Jurkat cells were diluted in RPMI1640 medium at a density of 1 × 10⁷ cells/mL. Cells were placed in the upper chamber of a 24-well plate protocol for human monocytes, as described previ

3D migration of human CD4⁺ T cells

The migratory behavior of primary human T cells was assessed by three-dimensional (3D) migration methodology using time-lapse microscopy and single cell tracking using the 3D chemotaxis µ-Slide system from Ibidi GmbH (Munich, Germany). The method was performed following a slight modification of the established Ibidi dendritic cell protocol for human monocytes, as described previously [40]. Briefly, isolated CD4⁺ human T cells (3.5 × 10⁶) were seeded in a rat tail collagen type-I gel (Ibidi, Munich, Germany) in DMEM and subjected to a gradient of human MIF, CXCL4L1, or a pre-incubated combination of both. Cell motility was monitored performing time-lapse imaging every 0.5 or 2 min at 37 °C for a period of 120 min to cover either a short or extended migration period, using a Leica DMi8 inverted microscope (Leica Microsystems, Wetzlar, Germany) and Leica live cell-imaging software (LAS X version 3.7.4). Images were imported as stacks to ImageJ version 1.51n and analyzed with the manual tracking and Chemotaxis and Migration tool (Ibidi GmbH) plugin for ImageJ.
Prior to starting the flow adhesion assay, MgSO₄ and CaCl₂ stock solutions were added to MonoMac6 suspensions to a final concentration of 1 mM each. Flow channels containing HAoEC monolayers were then perfused with MonoMac6 cells at a shear rate of 1.5 dyn/cm² for 10 min at 37 °C using the Ibidi pump system and perfusion set. For quantification of adherent monocytes, images from four positions per channel (fields of view, FOV) were acquired using a Leica DMi8 inverted microscope (20 × objective) and quantified with Image J software version 1.51n.

**Fluorescence polarization spectroscopy**

Fluorescence polarization was measured using a JASCO FP-6500 fluorescence spectrophotometer equipped with FDP-223 and FDP-243 manual polarizers (JASCO Deutschland GmbH, Pfungstadt, Germany). Preparation of stock solutions, measurements and analysis were performed essentially following a previously published protocol [40]. For binding/inhibition experiments, mixtures of Alexa 488-labeled MIF (10 nM) in the absence/presence of CXCL4L1 (1.6 µM) (or 20 mM sodium phosphate buffer, pH 7.2), and non-labeled msR4M-L1 peptide (concentration between 1 nM and 10 µM) were prepared in 10 mM sodium phosphate, pH 7.2, containing 2% hexafluoro-isopropanol (HFIP). Where CXCL4L1 was added as a putative inhibitor, Alexa 488-labeled MIF and CXCL4L1 were mixed and incubated for 30 min prior to measurements. Bandwidth for excitation and emission was set at 5 nm and time response at 0.5 s. The excitation wavelength was 492 nm and emission was set at 5 nm and time response at 0.5 s. Fluorescence polarization was measured using a JASCO FP-6500 fluorescence spectrophotometer equipped with FDP-223 and FDP-243 manual polarizers (JASCO Deutschland GmbH, Pfungstadt, Germany). Preparation of stock solutions, measurements and analysis were performed essentially following a previously published protocol [40]. For binding/inhibition experiments, mixtures of Alexa 488-labeled MIF (10 nM) in the absence/presence of CXCL4L1 (1.6 µM) (or 20 mM sodium phosphate buffer, pH 7.2), and non-labeled msR4M-L1 peptide (concentration between 1 nM and 10 µM) were prepared in 10 mM sodium phosphate, pH 7.2, containing 2% hexafluoro-isopropanol (HFIP). Where CXCL4L1 was added as a putative inhibitor, Alexa 488-labeled MIF and CXCL4L1 were mixed and incubated for 30 min prior to measurements. Bandwidth for excitation and emission was set at 5 nm and time response at 0.5 s. The excitation wavelength was 492 nm and emission was recorded at 519 nm. Measurements were taken at RT within 2 to 3 min upon preparation of the solutions. Polarization P was calculated according to the equation $P = (I_\parallel - G \cdot I_\perp)/(I_\parallel + G \cdot I_\perp)$, with $I_\parallel$ as the intensity of emitted light polarized parallel to the excitation light and $I_\perp$ as the intensity of emitted light polarized perpendicular to the excitation light. The $G$ factor was calculated based on the instrumental documentation [47]. Apparent $K_D$ (app. $K_D$) values were calculated assuming a 1:1 binding model [46], using sigmoidal curve fitting with OriginPro 2016 (OriginLab Corporation, Northampton, MA, USA).

**Label-free dynamic mass redistribution (DMR) assay**

Analysis of dynamic mass redistribution of adherent cells was performed on an EnSpire Multimode plate reader equipped with an Epic® label-free measurement module (PerkinElmer Inc., Waltham MA, USA) according to the manufacturer’s instructions for cell-based label-free DMR measurements [48]. The assay protocol was adapted according to a previous publication to be performed in EnSpire label-free 96-well fibronectin-coated cell assay microplates (Corning Gmbh, Amsterdam, The Netherlands) with HEK293 cells stably expressing human CXCR4 [32]. Briefly, 40,000 cells were seeded into each well and cultivated overnight (37 °C, 5% CO₂) to achieve a confluency of > 70%. Prior to the assay, the medium was exchanged with DMR assay buffer (20 mM HEPES and 1% DMSO in HBSS, pH 7.4) and the assay plate left for 6 h to equilibrate to ambient temperature. Baseline measurements for each well were recorded for 10 min every 30 s prior to treatment of the cells with chemokines, inhibitors, or the corresponding buffers as control. Treatments were applied to each well as a 5 x concentrated stock, prepared in assay buffer. For treatment with MIF/CXCL4L1 complexes, both proteins were mixed in assay buffer and incubated for 5 h at RT. Directly after addition of the stimuli, the DMR response was recorded for the indicated duration. The DMR response resembles the wavelength shift of the light reflected from the sensor integrated in the assay microplates and serves as a cumulative cellular response signal. Measurements were performed on two replicates per treatment and results are presented as their mean value.

**Staining of human thrombus specimens**

Human thrombus tissue specimens, obtained as disposable material from vascular surgery procedures of patients undergoing thrombectomy due to a peripheral artery disease (PAD) diagnosis (ethics allowances LMU Munich 18-104, LMU Munich 21-1187, and TUM-MRI project # 2799/10), were embedded in Tissue Tek O.C.T. Compound (Sakura Finetek Germany GmbH, Staufen, Germany), frozen, and cut into 5 or 8 µm sections using a CM 1950 Cryostat (Leica Biosystems). The cryosections were transferred to microscopy slides and stored at −80 °C until use.

**Histological staining of fibrin and platelets in thrombus sections**

The Carstairs’ method was used to stain fibrin and platelets in sections of arterial thrombus tissue specimens. Briefly, blocks of paraffin-embedded thrombi were cut into continuous cross-sections of 5 or 8 µm thickness. The sections were deparaffinized by immersion in ROTI® Histol (Carl Roth) and rehydrated in gradually decreasing concentrations of ethanol and finally in distilled water. After rehydration, the slides were subsequently placed 5 min into ferric ammonium sulfate (5% solution; Electron Microscopy Sciences, Hatfield, PA, USA), Mayer’s hematoxylin (5 min; Electron Microscopy Sciences) and picric acid-orange G solution ( Electron Microscopy Sciences) and picric acid-orange G solution ( Electron Microscopy Sciences) for 1 h. Then, the sections were stained in Ponceau Fuchsin solution ( Electron Microscopy Sciences) for 5 min. For differentiation, the slides were placed into phosphotungstic acid (1%; Electron Microscopy
Staining of platelets

Freshly isolated platelets were fixed with 4% paraformaldehyde (PFA) in PBS (Morphisto GmbH, Frankfurt a. M., Germany) for 10 min and subsequently permeabilized using 1 × Perm buffer (Invitrogen) for 15 min. After washing, platelets were blocked in ROTI® Block (Carl Roth) for 1 h. Immunofluorescent staining was performed as described above for the human thrombus specimens, except that the anti-human MIF antibody was used at a dilution of 1:20 and the anti-human CXCL4L1 antibody at a dilution of 1:100. Freshly isolated platelets were fixed with 4% paraformaldehyde (PFA) in PBS, pH 7.4, containing calcium. Before perfusion, the blood was incubated with fluorochrome 3,3'-dihexyloxacarbocyanine iodide (DiOC6, 1 mM; Sigma Aldrich) and allocated into separate reaction tubes. 200 µL of each solution were distributed onto separate collagen-coated cover slips (100 µg/mL) and incubated for 2 h. Cover slips were then washed in blocking buffer and mounted on poly-L-ornithine-coated glass slides using Pro-Long™ Glass Antifade mountant (Invitrogen), covered with coverslips and stored at 4 °C until imaging by multiphoton microscopy.

Multiphoton laser-scanning microscopy (MPM) and FLIM-FRET

Imaging was conducted using a multispectral TCS SP8 DIVE LIGHTNING multiphoton microscope (Leica, Germany) equipped with filter-free 4TUNE NDD detection module, an extended IR spectrum tunable laser (New InSight® X3™, Spectra-Physics) (680–1300 nm) and fixed IR laser (1045 nm), advanced Vario Beam Expander (VBE), Ultra-high-speed resonance scanner (8 kHz), HC PL IRAPO 25x/1.0 WATER objective, and FLIM-FRET modality with FALCON fast lifetime contrast imaging. Images were collected in a sequential scanning mode using hybrid diode detectors Reflected Light Hybrid Detectors (HyD-RLD) (Alexa Fluor-488: excitation 965 nm/emission 479–568 nm; Cy3: excitation 1095 nm/emission 538–650 nm) and were handled using the LAS-X software package. Deconvolution microscopy was performed using the Leica LIGHTNING (adaptive deconvolution) application. For fluorescence lifetime imaging (FLIM) and FLIM-FRET measurements, up to 1000 photons per pixel were captured in a time-correlated single photon counting (TCSPC) mode. Fluorescence lifetime decay data were fitted using Leica FALCON (FAstLifetime CONtrast) software applying a multi-exponential model. The fitting was assessed by randomly distributed residuals and by low Chi-square (\(\chi^2\)) values. The number of components used for the fittings was manually fixed to a value (\(n = 2–3\)) to minimize \(\chi^2\) values. The FRET efficiency was calculated by the Leica FALCON software according to the following formula: FRET Eff (\(E\)) = 1 − (\(\tau_{AV,Amp}\)/(\(\tau_D\)) (where \(\tau_{AV,Amp}\): amplitude-weighted average lifetime of the quenched donor (undergoing FRET); \(\tau_D\), amplitude-weighted average lifetime of the unquenched donor). The fluorescence lifetime of the donor was acquired similarly in the absence of the acceptor.

Proximity ligation assay (PLA)

For detection of protein complexes by proximity ligation assay (PLA), the Duolink™ InSitu Orange Starter Kit Mouse/Rabbit (DUO92102) from Sigma Aldrich was used. Following scouting experiments to establish the PLA methodology in thrombus material, cryosections of the thrombi were prepared by treatment with cold acetone for 6 min at 4 °C and for 30 min at RT. Samples were rehydrated in PBS for 10 min and hydrophobic barriers were applied to the microscopy slide using an ImmoEdgeTM Pen (Vector Laboratories Inc. Burlingame, USA).

For PLA detection, the Duolink® PLA Fluorescence protocol provided by the manufacturer was essentially followed, using primary antibodies against human MIF (mouse anti-MIF D2, sc-271631, Santa Cruz Biotechnology Inc., Dallas, USA; 1:100) and against human CXCL4L1 (rabbit anti-CXCL4L1, PA5-21944, Invitrogen; 1:100). Samples were then prepared for microscopy using Duolink® mounting medium with DAPI, and coverslips sealed with commercially available nail polish and stored at 4 °C until imaging by confocal microscopy using a Zeiss LSM880 Airyscan microscope (Carl Zeiss Microscopy Deutschland GmbH) was performed.

Flow chamber assay with platelets

Chemokines were diluted in calcium-free PBS, pH 7.4, at their final concentrations (MIF: 16 nM; CXCL4L1: 32 nM) and allocated into separate reaction tubes. 200 µL of each solution were distributed onto separate collagen-coated cover slips (100 µg/mL) and incubated for 2 h. Cover slips were blocked with PBS, pH 7.4, containing 1% BSA for 1 h. Next, human whole-blood was diluted at a 5:1 ratio with PBS, pH 7.4, containing calcium. Before perfusion, the blood was incubated with fluorochrome 3,3'-dihexyloxacarbocyanine iodide (DiOC6, 1 mM; Sigma Aldrich) for 10 min at RT. Thereafter, the blood was allocated into 1 mL syringes and perfused over the different cover slips, through a transparent flow chamber with high shear rate.
Heterocomplexes between the atypical chemokine MIF and the CXC-motif chemokine CXCL4L1 regulate...

(1000 s⁻¹) for 5 min. Per run, one 2-min video clip was recorded (200 ms/frame, Nikon Eclipse Ti2-A, 20× objective). Afterwards, the chamber was rinsed and pictures were taken of five representative areas using the same objective. The covered area was analyzed using the NIS-Elements AR software (Nikon) and the mean percentage of the covered area, the mean thrombus area as well as the mean thrombus count were determined.

Platelet spreading analysis

Fibrinogen-coated (100 µg/mL, Sigma Aldrich) coverslips were preincubated with MIF (16 nM), CXCL4L1 (32 nM), or MIF (16 nM) and CXCL4L1 (32 nM) together, for 2 h. Afterwards, isolated human platelets were diluted in Tyrodes buffer (pH 7.4) to match a concentration of 15,000 cells/µL. Platelets were supplemented with 1 mM CaCl₂, activated with 1 µg/mL CRP-XL (CambCol, Cambridge, UK), and incubated on the previously prepared fibrinogen-coated coverslips for 30 or 60 min at RT. Thereafter, platelets were fixed with 4% formaldehyde (Sigma Aldrich) for 10 min, and washed three times with PBS, pH 7.4. The coverslips were mounted onto slides and five images from randomly selected areas were taken using a Nikon Eclipse Ti2-A microscope with a 100×DIC objective. Subsequently a quarter of each image with at least 20 cells was analyzed.

Protein structure visualization

Three-dimensional structures as well as the surface charge distribution of human MIF, CXCL4 and CXCL4L1 were visualized using the PyMOL Molecular Graphics System software, version 1.8.2.2 (Schrödinger, LLC). The structures represent the Protein Data Bank (PDB) files for MIF (PDB ID: 3DJH), CXCL4 (PDB ID: 1F9Q), and CXCL4L1 (PDB ID: 4HSV), in our molecular docking results.

Protein–protein docking

To simulate the interaction of monomeric MIF with CXCL4 and CXCL4L1 in their monomeric forms, rigid protein–protein docking, followed by clustering of the 1000 lowest energy structures and removal of steric clashes was performed using the ClusPro 2.0 webserver, with single chains of MIF and CXCL4L1 defined as ‘receptor’ and ‘ligand’, respectively [49, 50].

Statistical analysis

Statistical analysis was performed using GraphPad Prism Version 6.07 software. Unless stated otherwise, data are represented as means ± standard deviation (SD). After testing for normality, data were analyzed either by two-tailed Student’s T test, Mann–Whitney U test, Kruskal–Wallis test or ordinary one-way ANOVA as appropriate. To account for multiple comparisons, either Dunn’s post hoc, Tukey’s or Sidak’s multiple comparisons tests were applied as appropriate. Differences with P < 0.05 were considered to be statistically significant.

Results

High affinity binding between the atypical chemokine MIF and the platelet CXC chemokine CXCL4L1

To begin to test the hypothesis that chemokine-chemokine interactions may extend to ACKs and that MIF may form heterocomplexes with classical chemokines, we applied unbiased chemokine protein array technology (Fig. 1A, B), as previously successfully used to map formation of heterocomplexes between different classical chemokines [13]. In addition to 47 human chemokines covering all four sub-classes (CXC-, CC-, CX3C- and C-type CKs), we also included structurally related and positively charged protein mediators including ACKs/DAMPs such as HMGB1, HBDs, and peroxiredoxins (Prxs) [51, 52], as well as MIF itself and the MIF homolog D-dopachrome tautomerase (D-DT/MIF-2) as spotted proteins in the protein array. Probing of the array with biotin-conjugated MIF and streptavidin-POD (StrAv-POD) revealed high-intensity spots indicative of a tight interaction of MIF with CXCL4L1 and Prx1 (Fig. 1B, C). Weaker spots were detected for CCL28, CXCL9, Prx6, and MIF itself. No spot intensity whatsoever was observed for any of the other 44 chemokines, indicating that none of the other 44 chemokines interacts with MIF. This also included CXCL8 and CXCL12, which share their receptors CXCR2 and CXCR4, respectively, with MIF [25].

Similarly, no binding signal of biotin-MIF was detected with HMGB1, a DAMP which has been demonstrated to form heterodimers with CXCL12 and for which a functional interaction with MIF has been suggested [37, 53], nor for the human β-defensins HBD1 or HBD2 (Fig. 1B, C). Importantly, when testing a control chemokine array developed with StrAv-POD without a biotin MIF incubation step, only one signal was not fully specific. This was the signal for Prx1, so that its interpretation was not possible (Supplementary Fig. 1A). Biotin-MIF also bound to...
MIF itself, but not to MIF-2 (Fig. 1B). As MIF is known to form homo-oligomers [24] and has been reported to form higher-order hexameric complexes [54], this result further verified the validity of the chemokine array approach for MIF.

A striking observation was that biotin-MIF specifically interacted with the immobilized platelet chemokine CXCL4L1, but not with CXCL4 (Fig. 1C and Supplementary Fig. 1B). CXCL4 and CXCL4L1 are highly homologous chemokines, their sequences only differ by three amino acids, and CXCL4L1 has also been suggested to be a decoy chemokine paralog of CXCL4. Given this remarkable specificity of the interaction with MIF and that the spot corresponding to biotin-MIF and CXCL4L1 was the strongest interaction detected on the array, we focused on CXCL4L1 as a novel candidate interactor of MIF.

We first verified the interaction by co-immunoprecipitation using whole cell lysates of MonoMac6 cells, which we found to express substantial amounts of CXCL4L1. Semi-endogenous pulldown of proteins from MonoMac6 lysates by biotin-MIF and StrAv magnetic beads and Western blot using an anti-human CXCL4L1 antibody revealed a specific band for CXCL4L1, which was absent, when the pulldown was performed without biotin-MIF preincubation (Fig. 2A and Supplementary Fig. 1C). Pulldown specificity was further confirmed by Western blot against MIF. We next applied surface plasmon resonance (‘Biacore’) methodology, which was previously successfully used to characterize interactions within the classical chemokine interactome [13]. To study the MIF/CXCL4L1 interaction, MIF chips were exposed to increasing concentrations of CXCL4L1 in the soluble phase. The obtained surface plasmon resonance response curves indicated that MIF specifically binds to CXCL4L1 (Fig. 2B, C). Quantitative analysis determined a $K_D$ value of $116 \pm 16$ nM (mean ± SD) indicating high-affinity binding between MIF and CXCL4L1. By contrast, no appreciable signal was detectable for the incubation with increasing concentrations of CXCL4 and no $K_D$ could be derived, verifying the specificity of the MIF/CXCL4L1 interaction in this set-up. To further confirm the MIF/CXCL4L1 interaction, we next applied microscale thermophoresis (MST), which relied on the interaction between MST-Red-labeled MIF and its binding partner, with both partners in the soluble phase. This methodology was recently established for MIF [40]. MST titrations of MST-Red-MIF with increasing concentrations of CXCL4L1 revealed a typical sigmoidal binding curve with a derived binding constant ($K_D = 159.8 \pm 16.8$ nM) that was similar to that obtained by surface plasmon resonance (Fig. 2D, E). In contrast, binding was much weaker when CXCL4 was titrated and accordingly a low affinity $K_D$ in the micromolar range was determined ($K_D = 2.0 \pm 0.8$ µM).
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![Image](image.png)

**Heterodimer formation between classical chemokines relies on CC-type or CXC-type interactions.** To determine which residues in CXCL4L1 are critical for the interaction with MIF, we employed peptide array technology. A set of 15-meric peptides derived from the CXCL4L1 sequence, positionally frame-shifted by three amino acids to cover...

**Fig. 2** Validation of MIF/CXCL4L1 complex formation by a variety of protein–protein interaction assays and verification of the specificity of MIF complexation with CXCL4L1 over CXCL4. **A** Semi-endogenous pull-down assay, in which endogenous CXCL4L1 from MonoMac6 lysates was captured with recombinant biotinylated MIF and pulled down by streptavidin-coated paramagnetic beads. Blots, developed against MIF (left) and CXCL4L1 (right), show representative results of three independent experiments. Input corresponds to 5% cell lysate without pull-down and control (Ctrl) refers to pull-downs performed in the absence of biotin-MIF. Molecular weight markers were electrophoresed in the same gel and relevant marker sizes are indicated. **B** Interrogation of MIF/CXCL4L1 complex formation by surface plasmon resonance (SPR) spectroscopy using chip-immobilized biotin-MIF titred against increasing concentrations of CXCL4L1. Measurements indicate an interaction between MIF and CXCL4L1 with an estimated $K_D$ of 116 ± 16 nM. The SPR response signal is given in relative units (RU). **C** Same as **B**, except that titration was performed with CXCL4. Corresponding SPR spectroscopy data for MIF and CXCL4. No detectable binding signal was obtained and no $K_D$ could be derived. **D** Interrogation of MIF/CXCL4L1 complex formation by microscale thermophoresis (MST) utilizing fluorescently labeled MIF and CXCL4L1 in solution. MST analysis revealed a $K_D$ of 159.8 ± 16.8 nM for the interaction of MIF and CXCL4L1. **E** Same as **D**, except that CXCL4 was tested. The derived apparent $K_D$ of 2.0 ± 0.8 µM was ten-fold higher compared to MIF/CXCL4L1.
Microglia motility — trajectories

Control

CXCL4L1

MIF

MIF + CXCL4L1

3D T-cell chemotaxis — trajectories

Forward migration index (FMI)

Control

MIF

MIF + CXCL4L1

CXCL4L1

Monocyte adhesion under flow

Control

MIF

MIF + CXCL4L1

CXCL4L1
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The entire sequence of the processed chemokine, was synthesized and immobilized on glass slides and arrays, and probed with biotin-MIF. The most pronounced binding signal was observed for peptides representing the sequence region, which corresponds to the β2-strand motif IKAGPHCPTAQLIAT of CXCL4L1 (Supplementary Fig. 2A, B). A second peak encompasses the N-terminal sequence QCLCVKTTSQVRPRH. The difference in the 3D structures of CXCL4 and CXCL4L1 is characterized by a significant conformational rearrangement of the α-helix [55], although the sequence of CXCL4 differs from that of CXCL4L1 in only three α-helical residues (L58P, K66E, H67L with the conformational difference being mainly governed by the L67H mutation). In this respect, CXCL4 showed an essentially identical peptide binding profile as that of CXCL4L1 at the N-terminus as expected, but a slightly different pattern at the β2 strand region GPHCPTAQLIATLKN, that is packed onto the C-terminal α-helix (Supplementary Fig. 2A, B). Peptide array-based mapping of the CXCL4L1 residues involved in MIF binding was confirmed by molecular docking simulations. Docking applying the ClusPro software predicted that the β-sheet region including the IKAGPHCPTAQLIAT motif is located near the MIF contact site, facing the 4-stranded β-sheet of a single MIF monomer chain. This interaction could be promoted by an energetically favorable complementary electrostatic interaction between the two surfaces (Supplementary Fig. 2C).

MIF is known to crystallize as a homotrimer. Moreover, solution studies suggest that MIF can form trimeric, dimeric or monomeric species, depending on the conditions and concentrations applied [24, 56, 57]. However, the affinity constant characterizing MIF/MIF associations during its homo-oligomerization has not been systematically determined. Given the above confirmed binding of MIF to CXCL4L1, we next determined the affinity constant between MIF and CXCL4L1 by fluorescence titration spectroscopy and directly compared it to the self-association affinity of MIF under essentially identical experimental conditions. We first titrated 10 nM Alexa-labelled MIF against increasing concentrations of CXCL4L1 (Supplementary Fig. 3A). Analysis of the observed shift in fluorescence emission at the spectral peak of 519 nm (binding curve; Supplementary Fig. 3B) revealed an app. $K_D$ of 86.4 ± 15.4 nM, well in line with the affinities derived by SPR and MST. To determine the $K_D$ for MIF homo-oligomerization/self-association, Alexa-MIF was titrated as a function of increasing concentrations of unlabeled MIF. Quantification of the shift in fluorescence emission yielded an apparent $K_D$ of 18.0 ± 5.1 nM (Supplementary Fig. 3C, D), a value that could reflect either fluorescence changes associated with monomer–monomer interactions and/or formation of higher-order oligomers, e.g. trimer-trimer associations. This experiment suggested that the affinity of MIF/MIF homo-oligomerization is somewhat higher than that of MIF/CXCL4L1 hetero-oligomerization. Because affinities for both hom- and hetero-complex formation are not that far apart and fall in a similar range as physiological concentrations of MIF and CXCL4L1, it is likely that CXCL4L1 indeed competes with MIF to form heterocomplexes, a mass-action effect that should reflect local in situ concentrations dependent on the pathology of the particular vascular or inflammatory event.

Together, the co-immunoprecipitation, Biacore, MST, and fluorescence spectroscopic studies confirmed specific binding between MIF and CXCL4L1 and determined a high-affinity binding constant in the 100–150 nM range for the interaction. Analysis of the binding interface by peptide array-based mapping and molecular docking provides an initial prediction of the residues involved in the CXCL4L1/MIF binding site.

MIF/CXCL4L1 heterocomplex formation attenuates MIF-mediated inflammatory/atherogenic activities

We next wished to determine a potential functional role of MIF/CXCL4L1 heterocomplex formation. CXCL4L1 is an angiostatic chemokine acting through CXCR3 [58], but its role in inflammatory responses and atherogenesis is not well understood. Pro-atherogenic activities of MIF have been extensively characterized and are mainly mediated through non-cognate interaction of MIF with CXCR2 and CXCR4 [25, 26]. Here, we hypothesized that MIF/CXCL4L1 complex formation could influence CXCR4-mediated pathways of MIF.
We first asked whether MIF-elicited T-cell chemotaxis, a well-characterized atherogenic MIF effect mediated via T-cell-expressed CXCR4 [25], is affected by CXCL4L1. Scouting experiments using Jurkat T-cells confirmed that, when added to the lower chamber of a Transwell migration device as a chemotactractant, MIF elicited chemotaxis with a chemotactic index (CTX) of approximately 2. Moreover, when CXCL4L1 was preincubated with MIF to allow for complex formation, no upregulation of Jurkat T-cell chemotaxis was observed in contrast to MIF alone. While CXCL4L1 alone exhibited neither a chemotactic nor inhibitory effect (Supplementary Fig. 4A). In contrast and in line with the observed lack of binding between MIF and CXCL4, CXCL4 had no effect on the chemotactic activity of MIF. CXCL4 itself did not significantly enhance Jurkat T-cell chemotaxis (Supplementary Fig. 4B). To further test the physiological relevance of this finding, we next studied primary CD4+ T-cell chemotaxis and applied a three-dimensional migration setup, following individual cell migration trajectories by live cell imaging. MIF potently triggered T-cell migration as evidenced by a significant increase in forward migration index (FMI) (Fig. 3A, B), confirming previous data showing CXCR4-dependent stimulation of 3D monocyte migration by MIF [40]. This effect was abrogated, when MIF was coinubated with CXCL4L1, while CXCL4L1 alone had no effect on 3D T-cell motility. This suggested that MIF/CXCL4L1 heterocomplex formation interferes with MIF/CXCR4-stimulated chemotaxis of T cells.

To study the potential relevance of these findings for other inflammatory/immune cell types, we next evaluated the effect of MIF/CXCL4L1 complex formation on microglial motility in the physiological setting of cortical brain cultures. MIF promotes the motility of Egfp+ microglia in murine cortical brain cultures ex vivo in a Cxcr4-dependent manner, as read out by live microscopy and as indicated by blockade of the MIF effect by the soluble CXCR4 mimicking peptide msR4M-L1 (Supplementary Fig. 3C). Importantly, MIF-triggered microglia migration in this setting was fully ablated, when CXCL4L1 was added together with MIF following preincubation, while CXCL4L1 alone had no effect on microglia motility (Fig. 3C, D). This indicated that CXCL4L1/MIF heterocomplex formation attenuates MIF’s CXCR4-dependent effect on microglia migration.

Monocyte adhesion on endothelial monolayers under flow stress conditions is another atherogenesis-relevant process and MIF has been previously demonstrated to enhance this process in a CXCR2- and CXCR4-dependent manner [25, 40]. We therefore next interrogated the impact of heterocomplexation on MIF-triggered Monomac6 adhesion on HAoEC monolayers under flow. MIF increased the adhesion of Monomac6 cells on HAoECs by approximately sixfold (Fig. 3E, F). In contrast, when MIF was preincubated with CXCL4L1 to foster heterocomplex formation, the pro-arrest effect of MIF was fully abrogated. CXCL4L1 alone also led to increased monocyte adhesion, albeit less pronounced than MIF (approximately threefold). Thus, CXCL4L1/MIF heterocomplex formation attenuates MIF’s CXCR-dependent effect on atherogenic monocyte arrest.

Together, these experiments indicate that MIF/CXCL4L1 heterocomplexes attenuate MIF-mediated atherogenic/inflammatory activities, such as immune cell migration and endothelial adhesion.

**MIF/CXCL4L1 heterocomplex formation inhibits MIF binding to CXCR4**

The cell migration and flow adhesion experiments implied, but did not directly test, the notion that MIF/CXCL4L1 complex formation affects MIF signaling through the CXCR4 pathway. To test the involvement of CXCR4 directly, we performed a binding competition experiment that capitalized on our recent identification of a MIF-binding CXCR4 ectodomain-mimicking peptide msR4M-L1 [40]. Employing fluorescence polarization spectroscopy (FP), titration of increasing concentrations of msR4M-L1 with Alexa 488-MIF led to a pronounced sigmoidal change in the FP signal (Fig. 4A), in line with previous data showing high affinity binding between MIF and msR4M-L1 [40]. By contrast, when Alexa 488-MIF was preincubated with CXCL4L1 before the titration, the FP signal was ablated (Fig. 4A), suggesting that MIF/CXCL4L1 heterocomplex formation interfered with MIF binding to the CXCR4 mimic.

To further confirm an interference of heterocomplex formation with the MIF/CXCR4 pathway, we next studied dynamic mass redistribution (DMR) responses in HEK293 cells stably transfected with human CXCR4. Incubation of HEK293-CXCR4 transfectants with MIF but not control buffer led to a pronounced time-dependent increase in the DMR signal as a real-time readout of an integrated cellular response of living HEK293 cell activation through the MIF/CXCR4 receptor signaling pathway (Fig. 4C). This signal was markedly attenuated by the small molecule CXCR4 inhibitor AMD3100, whereas the DMR curve of AMD3100 alone was similar to the control buffer curve, confirming CXCR4-dependency of the MIF-induced signal. Of note, preincubation of MIF with CXCL4L1 led to an appreciable reduction in the DMR response curve as well, when compared to cell stimulation with MIF alone, while CXCL4L1 alone and buffer control showed no effect (Fig. 4C). Together, the competition binding study and the DMR experiment confirmed the notion that complexation by CXCL4L1 interferes with binding of MIF to CXCR4 and its ability to activate CXCR4-mediated cell responses.
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MIF and CXCL4L1 colocalize and form complexes in human platelet aggregates and clinical thrombus specimens

CXCL4L1 is an abundant platelet chemokine [59, 60] and we previously found that platelets also are a rich source of MIF [61]. The colocalization of MIF and CXCL4L1 in subcellular platelet compartments has not yet been studied, but a cellular biological characterization of CXCL4 suggested that this paralog may be localized in a different intracellular platelet compartment than MIF [61]. Notwithstanding, we surmised that colocalization and complex formation between MIF and CXCL4L1 may occur extracellularly after secretion from activated platelets.

Initial evidence for a colocalization of CXCL4L1 and MIF following co-secretion from activated platelets came from human platelet preparations that aggregated due to handling stress. Examination of these aggregates by multi-photon microscopy (MPM) using an Alexa 488 signal to label MIF and Cy3 immunofluorescence for CXCL4L1 revealed several areas with an apparent decrease in the cellular response to MIF, when MIF is pre-incubated with CXCL4L1. The DMR response of CXCR4-expressing HEK293 cells to MIF in the presence or absence of the CXCR4-antagonist AMD3100 is also shown, confirming the CXCR4-dependency of the cellular response to MIF.

Fig. 4 MIF/CXCL4L1 complex formation inhibits binding of MIF to CXCR4 and signaling of MIF through the CXCR4 signaling axis. A Fluorescence polarization (FP) spectroscopy shows the interaction of Alexa488-labeled MIF with the soluble CXCR4 receptor mimic msR4M-L1 with an apparent dissociation constant $K_D$ of 237.2 ± 24.2 nM. Data is presented as mean of 3 independent experiments; error bars represent the SD. B Pre-incubation of MIF with CXCL4L1 (160-fold molar excess) prevents the interaction of MIF with msR4M-L1 (app. $K_D$ > 10 µM).

C Dynamic mass redistribution (DMR) measurements with HEK293 cells stably expressing CXCR4 indicate that the cellular response to MIF is reduced, when MIF is pre-incubated with CXCL4L1. The DMR response of CXCR4-expressing HEK293 cells to MIF in the presence or absence of the CXCR4-antagonist AMD3100 is also shown, confirming the CXCR4-dependency of the cellular response to MIF.
significant donor lifetime shortening (from $2.019 \pm 0.069$ ns to $1.496 \pm 0.033$ ns) and FRET events (FRET efficiency peak at 20–25%), when Alexa 488/Cy3 FLIM-FRET was recorded in appropriate regions-of-interest (ROIs) (Fig. 5C, D), an observation that is consistent with the notion that MIF and CXCL4L1 not only colocalize in activated platelet preparations but form true heterocomplexes.
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To further investigate the physiological relevance of these findings, we next examined clinical thrombus specimens derived from vascular surgery procedures from PAD patients undergoing thrombectomy. Thrombi from four PAD patients retrieved from different arterial locations were examined (Supplementary Table 1). To determine whether colocalized MIF and CXCL4L1 formed heterocomplexes in thrombus tissue, a proximity ligation assay (PLA) was performed which detects inter-molecular interactions within a distance of < 10 nm and thrombus specimens were counterstained by Carstair’s staining to further visualize fibrin- and platelet-rich areas (Fig. 6; Supplementary Fig. 5). Specific PLA signals were detected in the atherosclerotic thrombus specimens, while control stainings without primary antibodies, or with anti-CXCL4L1 or anti-MIF alone, were negative (Fig. 6; Supplementary Fig. 5). This suggests that there is an appreciable formation of MIF/CXCL4L1 heterocomplexes in clinical thrombus tissue, which essentially confirms our FLIM-FRET data obtained in platelet preparations from healthy blood samples. Thus, both FLIM-FRET and PLA demonstrated that MIF and CXCL4L1 form heteromeric complexes upon release from activated platelets.

**Heterocomplex formation inhibits MIF-stimulated thrombus formation and alters the effect of MIF on platelet morphology**

Thrombus formation and clot retraction are relevant processes that occur upon vessel injury and in advanced atherosclerotic vessels. MIF was found to modulate these processes [62]. As our data showed that MIF/CXCL4L1 heterocomplexes form in the microenvironment of a thrombus, we...
next determined whether heterocomplex formation affects thrombus characteristics. Thrombus formation under flow perfusing diluted human blood over a collagen-coated surface harboring combinations of MIF and CXCL4L1 was studied as established [63] and was found to double following exposure to MIF when applying a shear rate of 1000 s⁻¹ (Fig. 7). CXCL4L1 alone did not affect thrombus characteristics, but when added together with MIF following pre-incubation, MIF-elicited thrombus formation was blocked. These effects were mainly related to thrombus size/coverage (Fig. 7B, Supplementary Fig. 6) rather than thrombus
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Discussion

Chemokines control numerous pathogenic pathways contributing to inflammation and atherogenesis. The recent systematic characterization of the chemokine interactome revealed that heteromeric interactions between classical CC- and/or CXC-type chemokines represent an important molecular adjustment screw that serves to amplify, inhibit, or modulate chemokine activity [13]. Here, we have identified a heteromeric interaction between MIF, a pleiotropic inflammatory cytokine and ACK, and the classical platelet chemokine CXCL4L1. We also show that CXCL4L1/MIF complex formation affects inflammatory/atherogenic and thrombogenic activities of MIF. The scheme in Fig. 8 summarizes the main findings of this study. This suggests that disease-relevant activities of MIF may be fine-tuned by heterocomplexation with CXCL4L1 and that the chemokine interactome extends to heteromeric interactions between classical and atypical chemokines.

In fact, binding of classical chemokines to non-CC- or CXC-chemokine mediators is not unprecedented. Three examples have been documented: (i) the CXC-chemokine CXCL12 binds to the alarmin HMGB1 and HMGB1/CXCL12 complex formation promotes chemotactic activity through CXCR4 [37, 38]; (ii) the anti-microbial peptide and α-defensin HNP1 binds to CCL5 and enhances monocyte adhesion through CCR5 [67]; (iii) macrophage-expressed galectins such as galectin-3 (Gal-3) bind to CXCL12 and attenuate CXCL12-stimulated signaling via CXCR4 [36]. However, while these studies underscore that classical chemokine activity may be modulated by interaction with various soluble mediators, HMGB1 and Gal-3 have no chemotactic activity on their own; HNP1 has been reported to exhibit chemoattractive properties, but the mediating chemoattractant receptor has remained elusive. In contrast, despite lacking the signature structural elements of classical chemokines such as the chemokine-fold and the N-terminal cysteine motif, MIF is a chemoattractant and depending on the microenvironmental context, can signal through the CXC chemokine receptors CXCR2, CXCR4, and/or ACKR3 to promote atherogenic and inflammatory leukocyte recruitment. Its CXC receptor binding capacity is based on the presence of a pseudo-ELR motif and an extended N-like loop, structurally mimicking the site 1 and 2 receptor binding motifs of the corresponding cognate ligands CXCL1/8 and CXCL12, respectively. Together with the β-defensins HDB1/2 and HBD3, which bind to CCR6 and CXCR4, respectively, and secreted fragments of certain AARSs, which bind to CXCR1 and CXCR2, MIF has therefore been designated an ACK [15, 16, 17, 18, 19, 26]. Our current identification of MIF/CXCL4L1 heterocomplexes thus also shows that the chemokine interactome is not strictly limited...
to interactions between classical CC- and/or CXC-type chemokines, but also encompasses heteromorphic interactions between classical and atypical chemokines, with potential functional modulation of the chemokine receptor pathway of both the classical or atypical chemokine. Although not further validated and pursued in our current study, the detection of additional candidate interactors of MIF in our performed unbiased chemokine array, i.e., CCL28, CXCL9, as well as Prx6 leads us to hypothesize that interactions between classical and atypical chemokines could represent a broader principal of an “expanded ACK/CK interactome”.

The validity of the solid phase chemokine array as an unbiased screening approach for candidate chemokine interactors has been previously established [13]. The general utility and specificity of this methodology was further confirmed in the current study. Out of 47 immobilized classical chemokines, in addition to CXCL4L1, only two other classical chemokines, i.e., CCL28 and CXCL9, were revealed to have positivity. While a functional link between MIF and CCL28 has yet to be unveiled, it is interesting to note that the other detected CXC chemokine was CXCL9, a CXCR3 agonist like CXCL4L1. Intriguingly, biotin-MIF neither bound to CXCL12 nor to CXCL8, indicating that implicated functional interactions between MIF and the cognate CXCR4 and CXCR2 ligands, respectively, are independent of heterocomplex formation.

Furthermore, the specificity of the performed array is underscored by the notion that CXCL4, the highly homologous sister variant of CXCL4L1, did not bind to MIF, both at pH 8 and also when we tested for this interaction at pH 6 (data not shown) to account for pH-dependent charge differences. We hypothesize that the striking difference between CXCL4L1 and CXCL4 in binding to MIF might be due to the suggested different conformation of these two chemokines, e.g. the more exposed and flexible α-helix of monomeric CXCL4L1 [55]. While CXCL4 has been amply characterized by us and others as a pro-atherogenic platelet chemokine, in part also via its intriguing capacity to hetero-oligomerize with CCL5 [14, 59], very little is known about the role of CXCL4L1 in chronic inflammatory diseases and atherosclerosis. Like its sister molecule, CXCL4L1 is also abundantly expressed in platelets; however, it apparently is not localized in α-granules but resides in a different sub-cellular compartment, from where it is constitutively secreted [68]. It is also found in other cell types including mononuclear cells and smooth muscle cells [68]. CXCL4L1 serves as an inhibitor of angiogenesis and has pro-inflammatory effects by inducing the release of CCL2 and CXCL8 from monocytes, while—contrary to CXCL4—it does not
promote monocyte survival [69, 70, 71]. There is only one in vivo study, in which CXCL4L1 was investigated as a prognostic marker in cardiovascular disease. Interestingly, below-median levels of CXCL4L1 were found to correlate with a worse outcome in stable coronary artery disease patients, as indicated by a higher rate of cardiac death, stroke, or myocardial infarction [72]. This finding might argue for a beneficial role of this chemokine in cardiovascular disease, even though the mechanisms behind this remain unclear, but certainly more studies are required. Of note, there is no equivalent of CXCL4L1 in mice [73], limiting functional in vivo studies of this chemokine and its complex with MIF, as predicted from our study.

Importantly, we validated the binding between MIF and CXCL4L1 by semi-endogenous pulldown from monocytes, as well as three different biophysical in vitro methods, i.e. SPR, MST, and fluorescence titration spectroscopy. The combination of these methods also addresses potential disadvantages of having one interaction partner immobilized [74]. The binding affinity constants derived from the SPR, MST, and fluorescence spectroscopy experiments (116, 160, and 86 nM, respectively) are in reasonable agreement with each other. The observed differences could be due to a number of factors, including surface immobilization effects, fluorescence versus biotin labeling, or buffers/solvents employed. Together, the results are suggestive of a relatively high binding affinity between MIF and CXCL4L1. Moreover, the obtained nanomolar $K_D$ is consistent with the reported concentrations of both proteins in inflammatory disease settings [26]. Flanking evidence for MIF/CXCL4L1 complex formation was obtained by our peptide array mapping and molecular docking results. As expected given their high sequence identity, the peptide array predicted identical binding sites for CXCL4 and CXCL4L1. Also, the peptide array methodology interrogates linear binding epitopes but cannot delineate conformational differences. In fact, Kuo et al. suggested that the three-amino acid difference between CXCL4 and CXCL4L1, although marginal, leads to a slight tilting of the C-terminal $\alpha$-helix [55]. We hypothesize that this moderate conformational change could be the basis for the observed preferred binding of MIF to CXCL4L1 compared to CXCL4. Differences in their binding affinity to CCL5 have already been reported for CXCL4 and CXCL4L1 and also the availability of their monomers, regulated by the stability of their tetrameric complexes, differs between these two chemokines [70]. Future structural studies, e.g., by nuclear magnetic resonance (NMR) spectroscopy, may help to further address these and other structural questions.

To investigate the functional consequences of MIF/CXCL4L1 heterocomplex formation, we focused on inflammatory and atherosclerosis-relevant activities of MIF. T-cell migration is one such activity that is regulated by the MIF/CXCR4 pathway [25]. In line with previous results, MIF promoted T-cell migration in a physiologically relevant 3D migration setting. Although T cells generally express the CXCL4L1 receptor CXCR3, CXCL4L1 alone had no effect on the chemotaxis of human PBMC-derived T cells. Lack of CXCL4L1 activity in this assay is likely due to the fact that CXCL4L1 is not a $b$ona $f$ide T-cell chemoattractant [71] and that the preferential CXCL4L1 receptor variant CXCR3B is poorly expressed on T cells [75]. The 3D T-cell migration data are supported by the result that MIF, but not the combination of MIF and CXCL4L1, promoted Jurkat T-cell migration in a 2D Transwell assay. Confirming the remarkable specificity of MIF binding to CXCL4L1 versus CXCL4, coinubation of MIF with CXCL4 did not result in reduced Jurkat T-cell migration. Of note, heterocomplex formation with MIF led to a complete blockade of MIF’s pro-migratory effect on primary T cells in the 3D migration setting. Inhibition of MIF-mediated T-cell migration by CXCL4L1 complexation could potentially be relevant in atherosclerosis, where it might represent a feedback mechanism that could serve to dampen the atherogenic response. Indeed, abundant CXCL4L1 levels may be released by activated platelets in an atherogenic microenvironment, where they could colocalize with endothelial-immobilized or monocyte-secreted MIF and infiltrating T cells. That complexation of MIF by CXCL4L1 can interfere with MIF’s chemoattractant activities was confirmed in a microglia assay, in which the motility of Egfp+ microglia in murine cortical brain cultures ex vivo was studied. In addition to representing an independent cell migration system, the data obtained from the microglia-containing cortical cultures further confirmed that complex formation interferes with MIF signaling through the CXCR4 pathway and underscored that the mechanism could be relevant in in vivo-like physiological tissue settings. Further support for a role of the heterocomplexes in modulating MIF’s atherogenic activities came from experiments testing MIF-mediated monocyte adhesion under flow, an effect that is also dependent on the MIF/CXCR4 (and MIF/CXCR2) axis. That MIF/CXCL4L1 heterocomplex formation interferes with MIF signaling through CXCR4 was independently validated by biochemical experiments using FP spectroscopy that capitalizes on a CXCR4 peptide mimic as receptor surrogate, and DMR analysis of HEK293-CXCR4 transfectants. While our focus was on inflammatory/atherosclerosis-relevant cellular effects of MIF predominantly mediated by the MIF chemokine receptor axis, it is possible that MIF/CXCL4L1 heterocomplexation also affects MIF signaling responses through CD74. This receptor is an abundantly expressed MIF monocyte/macrophage and microglia receptor, although its expression is low or absent in T cells and platelets [76]. Similarly (and not addressed in our current study) MIF effects involving CXCR2 pathways may also be influenced by MIF/CXCL4L1 heterocomplexes.
The inhibitory effect of MIF/CXCL4L1 heterocomplexes also raises interesting questions about affinity differences between MIF homo- versus MIF/CXCL4L1 hetero-oligomerization, as well as about the precise inhibitory mechanism of action. Our side-by-side comparison with fluorescence titration data indicated that the affinity for MIF/MIF homo-oligomers is only somewhat higher than that for heterocomplexes formed between MIF and CXCL4L1. Although the assay buffer contained 1% HFIP to increase conformational flexibility, we did not specifically test for monomer–monomer interactions, and the measured fluorescence shifts may reflect either monomer–monomer or higher-order oligomer interactions. Irrespective of the precise mechanism of homo- versus hetero-oligomerization (MIF monomer/MIF monomer versus CXCL4L1 monomer/CXCL4L1 monomer; or MIF trimer/MIF trimer versus CXCL4L1 monomer or CXCL4L1 tetramer; or other interacting oligomeric species), our data suggest that an excess of CXCL4L1 is necessary to appreciably compete with MIF homo-oligomer formation. Because such concentration differences likely occur in situ in the local microenvironment of an atherosclerotic plaque or thrombus, future experiments are required to clarify the significance and mechanism associated with these stoichiometries. To this end, the oligomeric MIF species supporting the activation of CXCR chemokine receptors by MIF is unknown, whereas trimeric MIF has been found to be the predominant oligomeric species that engages with CD74 [77].

Along the same lines, our MIF/CXCL4L1 binding studies and cellular experiments demonstrating inhibition of MIF effect by MIF/CXCL4L1 heterocomplexes do not directly address the mechanism of inhibition. Overall, our results allow us to propose three principal mechanistic possibilities: (i) heterocomplex formation reduces the concentration of bioactive MIF monomers or homo-oligomers and/or shields the ‘receptor-active’ interface on MIF; (ii) signaling-incompetent heterocomplexes directly interact with CXCR4 or another MIF activity-mediating receptor, thus competing with free MIF; (iii) heterocomplex formation modifies the MIF conformation resulting in a MIF species with attenuated receptor activity.

The interaction between MIF and CXCL4L1, likely resulting in local inhibition of MIF’s pro-inflammatory effects, was especially interesting to us in the context of previous studies, in which we identified human and mouse platelets as an abundant source of MIF [61, 62]. Here, we used confocal (CLSM) and multiphoton microscopy (MPM) to verify expression and localization of MIF in human platelets, as well as in platelet-rich clinical thrombus tissue. As expected, these experiments also showed the abundant presence of CXCL4L1 in platelets and thrombi, and suggested the colocalization and/or complex formation of MIF and CXCL4L1 in the vicinity of platelets. Due to the optical resolution limits of the CLSM and MPM methods, true colocalization and the specific subcellular compartment could not be determined. Evidence for the presence of MIF/CXCL4L1 heteromers is suggested by PLA performed on cryosections of human thrombi obtained from four PAD patients undergoing thrombectomy. In fact, PLA is an established method to detect CK heteromers as shown previously for HNP1/CCL5 complexes [67]. While specific PLA-positive signals were detected in all four patient specimens, the abundance of heterocomplexes appeared to substantially vary between specimens as expected for surgical tissue preparations. Due to the small sample size, this study may only be hypothesis-generating, and no associations with sex, risk factors, or comorbidities can be drawn. Further studies will be required to address this issue in larger cohorts.

Having confirmed the formation of this novel heterocomplex in platelet preparations and thrombus tissue, we assessed the effect of MIF, CXCL4L1 and their heterocomplex on platelet function and thrombus formation. Confirming our previous results, these results showed that MIF promoted thrombus formation leading to a larger thrombus-covered area in an in vitro setting under flow conditions. This effect was abrogated upon co-incubation with CXCL4L1. It is interesting to note that in the settings used in our experiment using a shear rate of 1000 s\(^{-1}\) for 5 min, MIF enhanced thrombus formation. On the other hand, we showed in a previous study that employed a shear rate of 1700 s\(^{-1}\), MIF reduced thrombus size, thus confirming that MIF is a modulator of thrombus formation, but also indicating that the directionality of the effect depends on the specific microenvironmental context.

Investigation of morphologic changes in isolated platelets during adhesion and activation on a fibrinogen-coated surface revealed that both MIF and CXCL4L1 favored a switch from large to small lamellipodia at an early time point. Interestingly, in this setting no inhibition by the heterocomplex on MIF-mediated effects was observed, but a synergistic behavior of MIF and CXCL4L1 was observed, suggesting that this effect may occur independently of CXCR4.

In addition to their classical role in wound closure and haemostasis, thrombus formation and platelet activation are processes that are closely linked to inflammatory processes that drive atherosclerosis [78, 79, 80, 81]. MIF has been amply linked to atherosclerotic pathogenesis both clinically and experimentally, with evidence for a number of contributing mechanisms including leukocyte recruitment and platelet activation [25, 26, 63, 82, 83]. In this present study, we observed heteromerization of MIF and CXCL4L1 and the effect of the MIF/CXCL4L1 heterocomplex on immune cell migration and adhesion. The heterocomplex impacts on thrombus size and platelet morphology implying that CXCL4L1 could have a protective role in atherosclerosis by mitigating the pro-atherosclerotic effects of MIF via
Heterocomplexes between the atypical chemokine MIF and the CXC-motif chemokine CXCL4L1 regulate the existing chemokine interactome by including ACKs. Identification of other potential interactors could broaden our understanding of classical chemokine receptors, but also by direct binding to classical chemokines. Interestingly, the identified chemokine interactor of MIF is not one of the cognate ligands of MIF receptors CXCR2 or CXCR4, but CXCL4L1, a prominent platelet chemokine not previously implicated in MIF biology or MIF-mediated pathologies. While evidence from experimental in vivo disease models will have to be obtained in future studies, our present data do suggest that MIF/CXCL4L1 complex formation could serve to attenuate inflammatory/atherogenic activities of MIF through the CXCR4 receptor axis. However, the influence of heterocomplexes on other MIF-driven receptor pathways, e.g., via CD74, have not been tested and deserve further scrutiny.

Our study also provides insight into the growing "chemokine interactome" with a particular focus on ACKs. While modulatory effects on the interactome by mediators not belonging to the class of chemokines have already been exemplified by intriguing studies involving HMGB1, HNPs, as well as galectins [36, 37, 67], the current study is first in demonstrating a role for MIF family proteins, and bona fide ACKs in general, as defined by their chemotactic activity mediated by engagement with classical chemokine receptors. Future identification of other potential interactors could broaden our existing chemokine interactome by including ACKs.

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Author contributions J.B., P.v.H., M.B., A.H., and X.B. conceived and designed the study with help from C.W., A.K., M.G., O.G., R.T.A.M., H.I., R.R.K., and K.M; M.B., A.H., X.B., E.S., A.-K.R., O.E.B., N.S., M.O., J.A.N., A.B., C.K., K.H., V.R., A.F., J.G., J.L., and C.K. performed research and analyzed data. M.B., A.H., X.B., E.S., A.-K.R., C.W., M.G., A.K., O.E.B., P.v.H., and J.B. contributed to the interpretation of the data. The first draft of the manuscript was written by M.B. and J.B. with help from A.H. and P.v.H. All authors revised and commented on the manuscript drafts. N.S., L.M., P.S., W.E.K., B.E., and R.K. contributed to critical materials, and J.B., P.v.H., C.W., A.K., R.T.A.M., A.H., M.G., B.E., and L.M. provided funding for the study.

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Declarations

Conflict of interest J.B., P.v.H., C.K., O.E.B., C.W., and A.K. are inventors on patent applications related to anti-MIF and anti-chemokine strategies in inflammatory and cardiovascular diseases. The other authors declare that they have no competing interests.

Ethics approval and consent to participate Mouse maintenance and experiments were reviewed and overseen by the institutional animal use and care committee of the local authorities (Regierung von Oberbayern, ROB, Germany) and performed in accordance with the procedures provided by the animal protection representative of CSD. Human thrombus tissue specimens were obtained as disposable material from vascular surgery procedures under ethics allowance LMU Munich 18-104 and TUM-MRI project # 2799/10.

Consent for publication N/A.

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Authors and Affiliations

Markus Brandhofer1 · Adrian Hoffmann1,2 · Xavier Blanchet3,4 · Elena Siminkovitch1 · Anne-Katrin Rohlfing4 · Omar El Bounkari1 · Jeremy A. Nestele4 · Alexander Bild5 · Christos Kontos5 · Kathleen Hille5 · Vanessa Rohde1 · Adrian Fröhlich1 · Jona Golemi6 · Ozgun Gokce6,7 · Christine Krammer1 · Patrick Scheiermann2 · Nikolaos Tsilimparis8 · Nadja Sachs9,10 · Wolfgang E. Kempf9 · Lars Maegdefesse9,10 · Michael K. Otabil1 · Remco T. A. Megens3,10 · Hans Ippel11 · Rory R. Koenen11 · Junfu Luo12 · Bernd Engelmann12 · Kevin H. Mayo13,14 · Meinrad Gawaz4 · Aphrodite Kapurniotu5 · Christian Weber3,7,11,10 · Philipp von Hundelshausen3,10 · Jürgen Bernhagen1,7,10

1 Division of Vascular Biology, Institute for Stroke and Dementia Research (ISD), LMU University Hospital, Ludwig-Maximilians-Universität (LMU) München, Feodor-Lynen-Straße 17, 81377 Munich, Germany
2 Department of Anesthesiology, LMU University Hospital, Ludwig-Maximilians-Universität (LMU) München, 81377 Munich, Germany
3 Institute for Cardiovascular Prevention (IPEK), LMU University Hospital (LMU Klinikum), Ludwig-Maximilians-Universität (LMU) München, Pettenkofer Straße 8a/9, 80336 Munich, Germany
4 Department of Cardiology and Angiology, University Hospital Tübingen, Eberhard Karls University Tübingen, 72076 Tübingen, Germany
5 Division of Peptide Biochemistry, TUM School of Life Sciences, Technische Universität München (TUM), 85354 Freising, Germany
6 Systems Neuroscience Group, Institute for Stroke and Dementia Research (ISD), LMU University Hospital, Ludwig-Maximilians-Universität (LMU) München, 81377 Munich, Germany
7 Munich Cluster for Systems Neurology (SyNergy), 81377 Munich, Germany
8 Department of Vascular Surgery, LMU University Hospital, Ludwig-Maximilians-Universität (LMU) München, 81377 Munich, Germany
9 Philadelphia, PA, USA
10 Nuremberg, Germany
11 Munich, Germany
12 Freiburg, Germany
13 Utrecht, The Netherlands
14 Munich, Germany

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