



Cross-kingdom mimicry of the receptor signaling and leukocyte recruitment activity of a human cytokine by its plant orthologs

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Human macrophage migration-inhibitory factor (MIF) is an evolutionarily-conserved protein that has both extracellular immune-modulating and intracellular cell-regulatory functions. MIF plays a role in various diseases, including inflammatory diseases, atherosclerosis, autoimmunity, and cancer. It serves as an inflammatory cytokine and chemokine, but also exhibits enzymatic activity. Secreted MIF binds to cell-surface immune receptors such as CD74 and CXCR4. Plants possess MIF orthologs but lack the associated receptors, suggesting functional diversification across kingdoms. Here, we characterized three MIF orthologs (termed MIF/D-dopachrome tautomerase-like proteins or MDLs) of the model plant *Arabidopsis thaliana*. Recombinant *Arabidopsis* MDLs (*AtMDLs*) share similar secondary structure characteristics with human MIF, yet only have minimal residual tautomerase activity using either *p*-hydroxyphenylpyruvate or dopachrome methyl ester as substrate. Site-specific mutagenesis suggests that this is due to a distinct amino acid difference at the catalytic cavity-defining residue Asn-98. Surprisingly, *AtMDLs* bind to the human MIF receptors CD74 and CXCR4. Moreover, they activate CXCR4-dependent signaling in a receptor-specific yeast reporter system and in CXCR4-expressing human HEK293 transfectants. Notably, plant MDLs exert dose-dependent chemotactic activity toward human monocytes and T cells. A small molecule MIF inhibitor and an allosteric CXCR4 inhibitor counteract this function, revealing

its specificity. Our results indicate cross-kingdom conservation of the receptor signaling and leukocyte recruitment capacities of human MIF by its plant orthologs. This may point toward a previously unrecognized interplay between plant proteins and the human innate immune system.

Chemokines are chemotactic cytokines that orchestrate immune cell trafficking in development, homeostasis, and disease. They are small polypeptides characterized by a distinctive chemokine fold and conserved N-terminal cysteine residues. According to the spacing of these cysteines, they are grouped into CC-, CXC-, CX₃C-, and C-type sub-classes, and correspondingly-termed chemokine receptors (CKRs)³ exist. Chemokines interact with G-protein-coupled receptor (GPCR)-type CKRs to constitute a complex network characterized by both specificity and redundancy (1–4). Due to their major role in immune cell migration, chemokines are pivotal players in the host innate and adaptive immune response in infections, but also upon tissue injury and during tumorigenesis. When dysregulated and owing to their role in controlling leukocyte infiltration, chemokines contribute to the pathogenesis of human inflammatory and autoimmune diseases, but also cardiovascular disease and cancer (5, 6). The significance of the chemokine system to host defense against pathogens is additionally highlighted by viral mimicry mechanisms that interfere with host chemokine pathways as an immune evasion strategy (7).

Macrophage migration-inhibitory factor (MIF) is an inflammatory cytokine with chemokine-like characteristics and a regulator of host innate immunity. Dysregulated MIF has been

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This article contains Figs. S1–S6, Tables S1–S4, and supporting Ref. 1.

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³ The abbreviations used are: CKR, chemokine receptor; ACK, atypical chemokine; *AtMDL*, *A. thaliana* MIF/D-DT-like protein; CD74, cluster of differentiation 74; CXCL12, CXC motif chemokine 12; CXCR, CXC motif chemokine receptor; CXCR4, CXC motif chemokine receptor 4; CTX, chemotactic index; D-DT, D-dopachrome tautomerase; *HsMIF*, *Homo sapiens* MIF; HPP, *p*-hydroxyphenylpyruvate; DCME, L-dopachrome methyl ester; MHC, major histocompatibility complex; MDL, MIF/D-DT-like protein; MIF, macrophage migration-inhibitory factor; ANOVA, analysis of variance; MBP, maltose-binding protein; GPCR, G-protein-coupled receptor; HRP, horseradish peroxidase; DMEM, Dulbecco's minimal essential medium; FCS, fetal calf serum; PBMC, peripheral blood mononuclear cell; SOE-PCR, splice overlap-extension PCR; pAb, polyclonal antibody.

identified as a pivotal mediator of human diseases such as acute and chronic inflammatory diseases, autoimmunity, atherosclerosis, and cancer (8–12). MIF is a member of the emerging family of atypical chemokines (ACKs). ACKs lack the chemokine fold and the consensus cysteine residues, but behave as chemoattractants by binding to CKRs (13–15). MIF not only binds to its cognate receptor CD74 to regulate cell proliferation (16), but also engages in binding to the CKRs CXCR2 and CXCR4 (17) as well as CXCR7 (18). MIF/CXCR pathways are drivers in several human diseases. The MIF/CXCR4 signaling axis controls monocyte, T-cell, and B-cell infiltration in atherosclerosis and other inflammatory conditions (17, 19). It also contributes to cancer metastasis, cardiac fibroblast survival, and eosinophil inflammation (20, 21).

MIF is a structurally unique 12.5-kDa protein and the founding member of the MIF protein family that also comprises D-DT/MIF-2 and MIF orthologs in various species (8, 17, 22–25). MIF proteins are structurally distinct from other cytokines and classical chemokines (26), but they share high architectural homology with bacterial tautomerase (8, 13, 27–29). The coding regions of human *MIF* and its paralog *D-DT/MIF-2* are homologous and in close proximity to each other, suggesting an ancestral duplication event (24, 30). D-DT was named for its ability to tautomerize the nonnatural D-stereoisomer of dopachrome, and this catalytic property is shared by MIF (8, 27, 30). Thus, MIF proteins are bifunctional, acting as cytokines/chemokines and enzymes, although the functional significance of the tautomerase activity in mammals has remained elusive. The three-dimensional (3D) structures of MIF and D-DT/MIF-2 are highly similar (26, 31), whereas the amino acid sequence homology is limited to 34 and 27% in humans and mice, respectively. D-DT/MIF-2 shares biological and pathological activities with MIF, but also has distinct characteristics (30).

MIF proteins exhibit a remarkable degree of evolutionary conservation across kingdoms, ranging from mammals to vertebrates, including fish and unicellular parasites (8, 25, 32, 33). Mammalian MIFs are intracellularly expressed and secreted from cytosolic pools via a nonconventional secretion pathway (13). It has been speculated that MIFs are evolutionary ancient cytosolic enzymes that have “acquired” a secondary role as regulatory proteins during evolution from unicellular to multicellular organisms. Consistent with this hypothesis, intracellular MIF has been found to interact with several cytoplasmic proteins to control cell behavior by (co-)regulating cellular redox homeostasis, transcription, and signaling (13, 34). The role as a secreted cytokine/chemokine can be regarded as a further extension of its functional properties in the vertebrate lineage.

Thus, it is not surprising that interactions between MIF/receptor networks from different species/kingdoms have been reported. However, these are so far confined to interactions between a mammalian host and parasitic microbes, with MIF proteins from pathogenic species employing molecular mimicry strategies to contribute to virulence and immune evasion mechanisms (32). For example, *Plasmodium falciparum* produces a MIF ortholog that modulates the host immune response to malaria by suppressing T-cell memory (35). Similar to viral chemokine mimics, parasite MIFs appear to “hijack”

host MIF receptors, albeit so far only interactions with CD74 have been reported (36, 37).

Based on sequence data bank information, the presence of MIF/D-DT-like proteins (MDLs) is also predicted in the plant kingdom, and we recently performed comprehensive *in silico* analyses of MDL genes/proteins across kingdoms and in the model plant *Arabidopsis thaliana* (*AtMDLs*) (33, 38). The *A. thaliana* genome harbors three *MDL* genes, and the predicted proteins exhibit a sequence identity of 28–33% to human MIF (*HsMIF*). A main conclusion of the *in silico* analysis has been that plant MDL proteins share residues reported to be critical for the tautomerase pocket of human MIF/D-DT and may thus have tautomerase activity (38). Interestingly, MIF orthologs from the plant-parasitic aphid *Acyrtosiphon pisum* are secreted in its saliva and mediate aphid survival and feeding on its host plant pea, representing an example of modulating plant immunity by a plant parasite. However, to date no experimental studies have been conducted with plant MDLs and their functions remain completely elusive.

Given the significant degree of sequence homology between MIFs and a predicted structural similarity across kingdoms, including a predicted conserved tautomerase site, we hypothesized that there might exist plant MIF protein-based mimicry mechanisms and that plant MDLs might interact with components of the human MIF network. To test this hypothesis, we cloned and expressed the three known *AtMDL* proteins, studied their structural features by CD spectroscopy, molecular modeling, and site-specific mutagenesis, and explored functional commonalities and potential direct interactions with the human MIF protein/receptor network.

Results

Generation and characterization of His-tagged recombinant *AtMDLs*

To initiate functional characterization of the three predicted *AtMDL* proteins *AtMDL1*, *AtMDL2*, and *AtMDL3* and to test the hypothesis that they might share similarities with mammalian MIFs, we recombinantly expressed these proteins and purified them for functional studies. The nucleotide and amino acid sequences of *AtMDL1*, *AtMDL2*, and *AtMDL3* were retrieved from UniProt and the European Nucleotide Archive (ENA) database. The *AtMDLs* share a sequence identity of 28–33% with *HsMIF*; this value is similar to the homology between human MIF and its paralog D-DT/MIF-2. For purification purposes, the *AtMDLs* as well as *HsMIF* were designed to express a C-terminal hexahistidine (6xHis) epitope tag. Table S1 summarizes key molecular parameters of the studied His-tagged proteins (*AtMDL1*–6xHis, *AtMDL2*–6xHis, *AtMDL3*–6xHis, and *HsMIF*–6xHis), *i.e.* the molecular mass and isoelectric points as predicted by ExPASy M_r /pI point calculator (https://web.expasy.org/compute_pi/).

The protein sequences, including the Leu–Glu linker residues and the C-terminal His-tags, were aligned using ClustalW algorithm (<https://www.genome.jp/tools-bin/clustalw>) (Fig. S1A). Prediction of the 3D structures of the *AtMDL* proteins using the Phyre² Protein Fold Recognition Server (<http://www.sbg.bio.ic.ac.uk/phyre2/html/page.cgi?id=index>) suggests an

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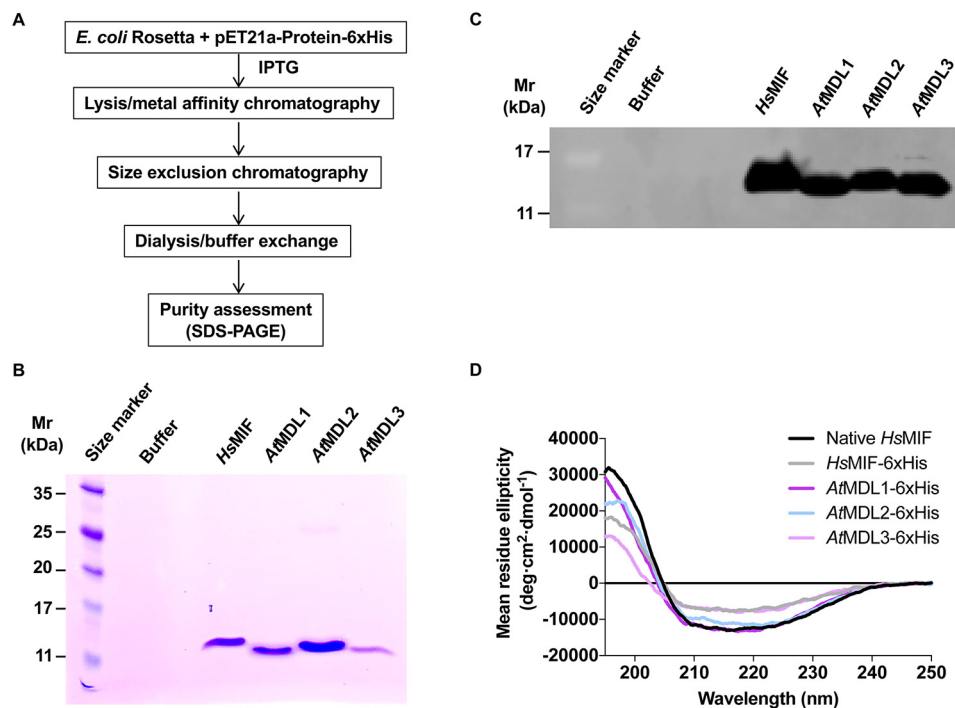


Figure 1. Expression, purification, and analysis of recombinant hexahistidine-tagged *HsMIF* and *AtMDLs*. *A*, schematic experimental procedure of protein expression and purification. *B*, Coomassie Blue staining of purified proteins after SDS-PAGE. *C*, protein detection by Western blotting using a monoclonal anti-His antibody. Relative molecular masses (M_r) are indicated on the left in kDa. *D*, CD spectropolarimetry of recombinant *HsMIF* and *AtMDLs*. Representative spectra of His-tagged *AtMDLs* and *HsMIF* as well as native (untagged) *HsMIF* are presented according to the indicated color code. Conformations in the CD spectra were measured as mean residue ellipticity as a function of the wavelength (given in nanometers) in the far-UV range.

apparent high-structural similarity between the authentic (untagged) *HsMIF*, 6xHis-tagged *HsMIF*, and its 6xHis-tagged plant orthologs (Fig. S1B). To assess potential structural similarities also experimentally, *AtMDL1*-6xHis, *AtMDL2*-6xHis, *AtMDL3*-6xHis, and *HsMIF*-6xHis were cloned and expressed in a bacterial expression system (Fig. 1A). All four C-terminally His-tagged recombinant proteins were purified by immobilized metal ion affinity (HisTrap) and subsequent size-exclusion chromatography. All four proteins were obtained in good quantities, without any detectable degradation products, and in high purity of >95% as assessed by SDS-PAGE (Fig. 1B and Table S2), although the yield for *AtMDL3* was lower compared with the other three proteins due to an observed increased aggregation tendency. Endotoxin concentrations were found to be negligible (Table S2). The identity of the enriched proteins was validated by SDS-PAGE/Western blotting detection with an anti-His antibody (Fig. 1C). We additionally conducted Western blot analysis using antibodies raised against human MIF. Both the polyclonal rabbit anti-mouse MIF antibody Ka565 and the monoclonal anti-human MIF antibody MAB289 substantially cross-reacted with *AtMDL1* and also weakly bound to *AtMDL2* (Fig. S2). This result supports the notion that the *AtMDLs* exhibit overlapping antigenic epitopes.

To further compare structural properties of the plant MDLs with those of *HsMIF*, we performed circular dichroism (CD) spectroscopy. This method is a valuable tool to compare the secondary structures of proteins of interest. CD spectra of all *AtMDLs* were recorded at concentrations of 1, 2.5, and 5 μM and compared with those of both His-tagged *HsMIF* and

nontagged (“fully native”) *HsMIF* (Fig. 1D and Fig. S3). This analysis verified that the recombinant *AtMDLs* were folded and that their secondary structure was very similar to that of His-tagged *HsMIF* and native *HsMIF*. It also suggested that the introduced His-tag has a minimal (if any) influence on the structures of the studied proteins, as the spectral features of *HsMIF* and *HsMIF*-6xHis were similar at all concentrations measured (Fig. 1D and Fig. S3). The deconvoluted spectra recorded at a concentration of 5 μM further indicated that the proportions of secondary structure elements of the recombinant *AtMDLs* differ only slightly from those of *HsMIF*. In line with previous CD data for human and mouse MIF (39), the calculated α -helical content of the *AtMDLs* was in the range of 20–37%, the β -strand contribution was between 28 and 33%, and β -turns were predicted to represent 17–22% of the secondary structural contents (Table S3). Overall, *AtMDL1*-6xHis and *AtMDL2*-6xHis showed the highest degree of similarity in experimentally determined secondary structural features to *HsMIF*-6xHis; *AtMDL3*-6xHis exhibited an elevated proportion of unordered secondary structural elements, an observation that is consistent with the reduced solubility of this MDL homolog compared with the other *AtMDLs* and *HsMIF*.

AtMDLs only exhibit minimal residual tautomerase activity

Whereas the functional role of the tautomerase activity of *HsMIF* and its natural substrate in a physiological context has remained elusive, the catalytic pocket residues have also been implicated to contribute to the receptor-binding interface between MIF and its receptors CD74 and CXCR4 (12, 13, 27, 40, 41). As the *AtMDLs* contain the consensus proline residue

in position 2 (Pro-2) known to be essential for tautomerase activity (Fig. S1) (38), we surmised that they may exhibit this catalytic capacity. Purified *AtMDL1*-6xHis, *AtMDL2*-6xHis, and *AtMDL3*-6xHis were subjected to a MIF tautomerase activity assay using *p*-hydroxyphenylpyruvate (HPP) as substrate. Using a MIF concentration of 250 nM, *HsMIF* and *HsMIF*-6xHis exhibited a similar tautomerase activity of 80–90 $\mu\text{mol min}^{-1} \text{mg}^{-1}$ (Fig. 2A). This value is in line with previous reports (42), indicating that the C-terminal His-tag does not interfere with the tautomerase activity of *HsMIF*, and underscoring that the epitope-tagged variant retains structural integrity, which is consistent with previous observations made for MIF orthologs from parasites (43). Surprisingly, all three *AtMDLs* were found to exhibit only minimal HPP tautomerase activity (Fig. 2A). Compared with buffer control conditions, the *AtMDLs* at a concentration of 250 nM exhibited an apparent activity of only 1–3 $\mu\text{mol min}^{-1} \text{mg}^{-1}$, which did not differ from the control at the level of statistical significance. A more detailed comparison of the enzyme kinetics of *AtMDL1*-6xHis with those of *HsMIF*-6xHis confirmed the notion that the *AtMDLs* only have minimal HPP tautomerase activity. Whereas the K_m value of the plant MDL was in the low millimolar range, *i.e.* similar to that of its human ortholog, the k_{cat} of *AtMDL1*-6xHis was substantially lower than that for *HsMIF*-6xHis, translating into a 300-fold higher k_{cat}/K_m value for *HsMIF*-6xHis compared with the plant ortholog (Table 1). To further explore the surprising difference in tautomerase activity between *HsMIF* and the *AtMDLs*, we also compared the tautomerase activities of *AtMDL1*-6xHis and *HsMIF*-6xHis in an assay using L-dopachrome methyl ester (DCME) as substrate. At an enzyme concentration of 100 nM, *HsMIF*-6xHis exhibited an activity of 300 $\mu\text{mol min}^{-1} \text{mg}^{-1}$ toward DCME, whereas *AtMDL1*-6xHis only showed an activity <20 $\mu\text{mol min}^{-1} \text{mg}^{-1}$ (Fig. 2B). The notion that *AtMDL1*-6xHis only has a marginal residual activity was underscored by the kinetic parameters (K_m (*HsMIF*-6xHis) = 0.6 mM; K_m (*AtMDL1*-6xHis) = 4.1 mM; k_{cat} (*HsMIF*-6xHis) = 77.4 s^{-1} ; and k_{cat} (*AtMDL1*-6xHis) = 2.1 s^{-1}) with the k_{cat}/K_m value of *HsMIF*-6xHis 250-fold higher than that of *AtMDL1*-6xHis (Table 1). Thus, although the critical Pro-2 residue of the tautomerase activity site is conserved in all three *AtMDLs*, the catalytic activity itself is drastically reduced in the plant orthologs of human MIF.

To begin to explore the structural basis for this striking difference, we inspected the residues that shape the catalytic tautomerase pocket (42) in 3D space more closely. We noted that in addition to Pro-2 of human MIF, Lys-32, Ile-65, and Tyr-96 are either identical or homologous in the *AtMDLs* (Fig. 2C). By contrast, the basic His-63 residue of *HsMIF* is substituted by a hydrophobic residue (Ile or Val) in the *AtMDLs*, and even more strikingly, Asn-98 of human MIF is replaced by the strongly basic Lys-98 in all *AtMDLs*, introducing a positively-charged residue in a critical position (Fig. 2C). The predicted 3D structures shown in Fig. 2D visualize the positions of residue Asn-98 versus Lys-98 (*top panel*; ribbon structure) and indicate that the Asn \rightarrow Lys substitution may condition a conformational change of the catalytic pocket, paralleled by a different orientation of the Lys-98 side chain (*top panel*; ribbon structure, and

middle panel; surface structure). Furthermore, the electrostatic surface potential model (Fig. 2D, *bottom panel*) highlights the effects of the substitutions on charge distribution around the pocket.

To experimentally test the presumed role of residue 98 for the ablated catalytic activity of the *AtMDLs*, we cloned an *HsMIF* mutant, in which Asn-98 was replaced by Lys-98. The expression and purification characteristics of N98K-*HsMIF*-6xHis were similar to those of WT *HsMIF*-6xHis and the hexahistidine-tagged *AtMDLs*. However, when compared with WT *HsMIF*-6xHis for its HPP- and DCME-dependent tautomerase activity, N98K-*HsMIF*-6xHis exhibited a drastically-reduced catalytic activity (Fig. 2, E and F). Thus, the replacement of asparagine by lysine at position 98 in the *AtMDLs* at least partially explains the essential absence of HPP- and DCME-dependent tautomerase activity noted in the *AtMDLs*.

AtMDLs bind to and signal through human MIF receptors

The Pro-2 residue not only is a central component of the tautomerase cavity of human MIF, but also was identified to contribute to *HsMIF* binding to the MIF receptors CD74 and CXCR4 (40, 41). Furthermore, additional motifs contributing to the binding interface between *HsMIF* and these MIF receptors are also found in the *AtMDLs* (Fig. 3A). Mutations in Pro-2 of *HsMIF* have been found to invoke a conformational change in the MIF structure, resulting in altered binding and activation characteristics of CD74, suggesting that Pro-2 is a critical determinant of the MIF-binding site for CD74 (40). In addition, *HsMIF* residues 80–87 have been identified to contribute to CD74 binding (44), and this site also is well-conserved in all three *AtMDLs* (Fig. 3A). We therefore tested the possibility that *AtMDLs* may bind to CD74, although this receptor is not present in plants. We capitalized on a recently developed MIF/CD74-binding assay that employs an MBP-sCD74 fusion protein, in which maltose-binding protein (MBP) is fused to the MIF-binding CD74 ectodomain (45). Intriguingly, all three hexahistidine-tagged *AtMDLs* were found to bind to MBP-sCD74. *AtMDL3*-6xHis exhibited a binding capacity that was similar to that of *HsMIF*-6xHis (Fig. 3B) and bound to MBP-sCD74 with an apparent K_D of 200 nM (Fig. 3C and Fig. S4A).

For CXCR4, Pro-2 supports binding at the site 2 location of the MIF/CXCR4 interface (41, 46). Because *HsMIF* and the *AtMDLs* additionally exhibit an appreciable degree of homology in the extended N-like loop region that fosters site 1 binding of human MIF (Fig. 3A), we also tested the possibility that *AtMDLs* may bind to human CXCR4. As GPCR ectodomains are not amenable to *in vitro* binding studies such as those performed for CD74, we employed a genetically-modified yeast transformant stably expressing human CXCR4. These cells represent a receptor-specific cell system, in which the native yeast GPCR Ste2 was replaced by human CXCR4, which is functionally linked to the yeast Ste/MAPK signaling cascade and enables us to detect receptor-specific binding and signaling responses of CXCR4 following activation with CXCL12 or MIF by β -gal reporter activity. Moreover, the system is devoid of any other mammalian receptors that might be involved in MIF signaling, thus representing a “pure” *in vivo* receptor-binding/signaling system (41, 47). Yeast CXCR4 transformants were incubated

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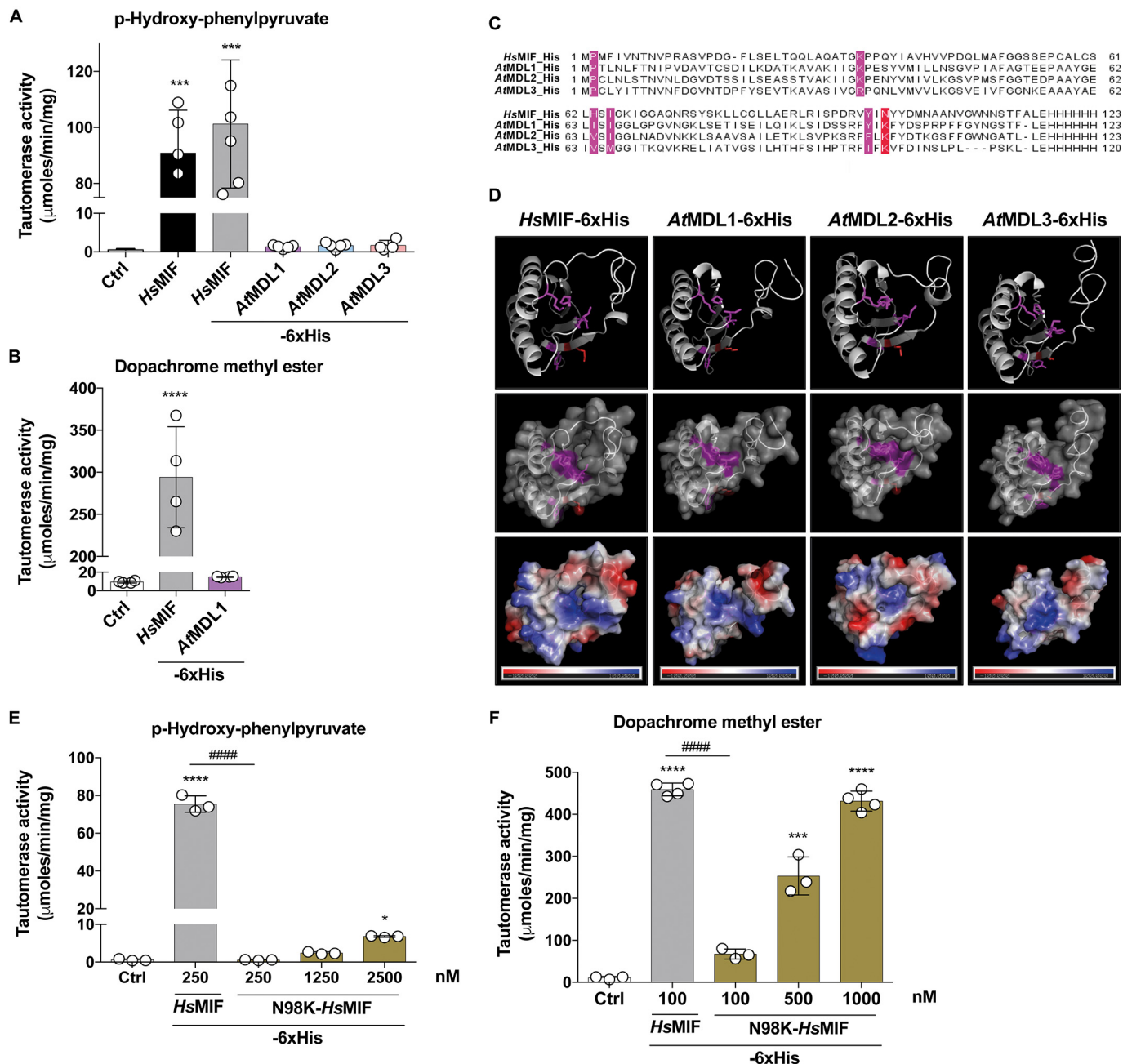


Figure 2. Recombinant *AtMDLs* only have residual tautomerase activity, likely due to a sterically-impeded substrate-binding pocket. *A* and *B*, tautomerase activity of recombinant *HsMIF/AtMDL* protein homologs measured by spectrophotometry. *A*, tautomerase activity of the three 6xHis-tagged *AtMDLs* was compared with that of 6xHis-tagged *HsMIF* using HPP as a substrate. Untagged *HsMIF* was measured for comparison. Data shown are from four to seven independent experiments \pm S.D., each performed in triplicate (scatter plot with *white circles* indicates individual data points). For statistical comparisons, one-way ANOVA between buffer control and the different samples was applied (***, $p < 0.005$). *B*, comparison of the tautomerase activity of *AtMDL1-6xHis* and *HsMIF-6xHis* using DCME as a substrate. Data shown are from four independent experiments \pm S.D., performed in triplicate each (scatter plot with *white circles* indicates individual data points). For statistical comparisons, one-way ANOVA between buffer control and the different samples was applied (***, $p < 0.005$). *C*, multiple sequence alignment of 6xHis-tagged variants of *AtMDLs* and *HsMIF*. Amino acid sequences of *AtMDL1* (identifier Q9LU69), *AtMDL2* (identifier Q9M011), *AtMDL3* (identifier Q8LG92), and *HsMIF* (identifier P14174) were retrieved from the UniProt database and aligned by ClustalW using standard parameters in the Jalview multiple sequence alignment editor desktop application. The amino acid residues forming the tautomerase substrate-binding pocket are highlighted in *magenta* and *red*. *D*, comparative view of predicted 3D structures of 6xHis-tagged *HsMIF* and the three *AtMDLs*. Only the monomers are shown for simplicity reasons. Amino acid sequences of the *AtMDL* proteins were subjected to analysis via the PHYRE² Fold Recognition server and visualized with PyMOL. The predicted 3D structures (ribbon, surface, and electrostatic surface potential models) of *AtMDL1-6xHis*, *AtMDL2-6xHis*, and *AtMDL3-6xHis* were analyzed compared with the known X-ray-resolved 3D structure of *HsMIF-6xHis*. The *upper* and *middle panels* highlight the location of crucial tautomerase pocket residues in the ribbon structure and on the protein surface (*magenta* and *red*). In the *lower panel*, *red* and *blue*, respectively, indicate an excess of negative or positive charges near the surface, and *grayish color* symbolizes neutral regions. *E*, comparison of the tautomerase activity of *N98K-HsMIF-6xHis* and *HsMIF-6xHis* using HPP as a substrate. Data shown are from three to four independent experiments \pm S.D., performed in triplicate each (scatter plot with *white circles* indicates individual data points). For statistical comparisons, one-way ANOVA between buffer control and the different samples was applied (*, $p < 0.05$; ****, $p < 0.001$), as well as comparison between 250 nM *HsMIF* and 250 nM *N98K-HsMIF-6xHis* (####, $p < 0.001$). *F*, same as *E* except that DCME was used as a substrate, and *HsMIF-6xHis* was applied at a concentration of 100 nM (***, $p < 0.005$; ****, $p < 0.001$; ###, $p < 0.01$).

Table 1
Comparison of the kinetic tautomerase activity parameters between recombinant His₆-tagged *HsMIF* and *AtMDL1*

Data represent triplicate measurements \pm S.D.

Enzyme assay	Protein	K_m^a	k_{cat}^b	k_{cat}/K_m
		MM	s^{-1}	$MM^{-1} s^{-1}$
HPP	<i>HsMIF</i> -6xHis ₆	2.31 \pm 0.86	39.78 \pm 0.89	17.22
	<i>AtMDL1</i> -6xHis	7.81 \pm 6.63	0.44 \pm 0.01	0.06
DCME	<i>HsMIF</i> -6xHis	0.59 \pm 0.18	77.39 \pm 1.67	131.17
	<i>AtMDL1</i> -6xHis	4.10 \pm 5.19	2.11 \pm 0.11	0.51

^a K_m is the Michaelis-Menten constant.

^b k_{cat} is turnover number.

with the recombinant *AtMDLs* and their potential CXCR4 binding/signaling activity compared with that of untagged and 6xHis-tagged human MIF. CXCR4 activation by *HsMIF*-6xHis was similar to that of *HsMIF*, confirming that the C-terminal hexahistidine tag neither impairs nor enhances MIF binding to CXCR4 (Fig. 3D). In line with previous data (41), CXCR4 activation by MIF was slightly less potent than that of the cognate ligand CXCL12. Strikingly, all three *AtMDLs* significantly promoted CXCR4 signaling activity. Moreover, CXCR4 activation by the plant MIF orthologs *AtMDL1*-6xHis and *AtMDL3*-6xHis was markedly stronger than that of *HsMIF*-6xHis and *HsMIF* (Fig. 3D).

These data suggested that *AtMDLs* have the capacity to interact with the human MIF receptors CD74 and CXCR4. As the yeast CXCR4-transformant experiments also implied a role in the activation of signal transduction, we focused on CXCR4 for subsequent functional studies and next asked whether *AtMDLs* would also trigger MIF-like CXCR4-facilitated signaling responses in mammalian cells. The MIF/CXCR4-induced PI3K/Akt signaling cascade is a well-studied MIF-mediated response pathway with physiological/pathophysiological relevance in human macrophages and T cells, as well as for cancer cell survival (20, 48, 49). We performed signaling studies in HEK293 cells stably overexpressing CXCR4 (HEK293-CXCR4), fostering appreciable surface expression levels of CXCR4 (Fig. S4B). In accordance with previous results (48), *HsMIF*-6xHis elevated phospho-Akt levels up to 4.5-fold within 15 min of stimulation (Fig. 4, A and B). Of note, all three *AtMDLs* stimulated Akt signaling in these cells. The most pronounced effect was seen for *AtMDL1*-6xHis, which not only shared with human MIF the capacity to trigger Akt phosphorylation, but even exhibited a more pronounced signaling effect, with its maximum shifted to an early peak at 5 min after stimulation (Fig. 4, C and D). *AtMDL2*-6xHis and *AtMDL3*-6xHis had slightly weaker effects than *HsMIF*-6xHis, and their activation maximum was delayed toward 15 min (Fig. 4, E-H). This result suggests that the *AtMDLs* are able to mimic the CXCR4-mediated Akt phosphorylation activity of human MIF in a mammalian cell system.

Plant MIF orthologs engage CXCR4 to act as chemoattractants for human monocytes

Induction of CXCR4-dependent intracellular signaling (Figs. 3 and 4) suggests that *AtMDLs* might modulate or mimic MIF's CXCR4-mediated leukocyte recruitment potential. To investigate this possibility, we tested the effect of the plant orthologs

on monocyte chemotaxis, applying Transwell migration chamber experiments that represent an established setup mirroring chemokine receptor-dependent immune cell migration responses. Recombinant hexahistidine-tagged *AtMDLs* were loaded into the lower compartment of a Transwell device, and their chemoattractant potency for THP-1 monocytes was compared with that of *HsMIF*-6xHis and the classical chemokine CXCL12. In line with prior findings obtained with untagged MIF (17), *HsMIF*-6xHis enhanced monocyte chemotaxis in a concentration-dependent manner with bell-shaped dose-response behavior and a maximal 4-fold chemotactic effect at a concentration of 16–32 nM. This compared well to the effect of the cognate CXCR4 ligand CXCL12 (chemotactic index = 4–7.5-fold compared with the untreated control at a concentration of 8 nM). Notably, all three *AtMDLs* were also able to promote THP-1 cell chemotaxis, featuring dose-dependent bell-shaped behavior, albeit at slightly lower potency compared with *HsMIF*-6xHis (Fig. 5). The maximal chemotactic effect of *AtMDL1*-6xHis and *AtMDL2*-6xHis was 2.5-fold compared with the buffer control and, like for *HsMIF*-6xHis, was observed at a concentration of 32 nM. *AtMDL3*-6xHis was also able to trigger monocyte migration, but its maximum effect was shifted toward a 5-fold higher concentration (chemotactic index = 3.5 at 80 nM) (Fig. 5).

We next sought to further confirm the specificity of this effect and to directly test for the involvement of CXCR4 in this process. ISO-1 is a well-established small molecule inhibitor of MIF that not only inhibits its tautomerase activity but also interferes with proinflammatory activities of MIF and MIF binding to CXCR4 (41, 50, 51). Similarly, AMD3100 is an allosteric inhibitor of CXCR4 that blocks the interaction with its cognate ligand CXCL12 and partially interferes with MIF-mediated CXCR4 activation (17, 41, 52–56). In line with these prior findings, co-application of AMD3100 fully abrogated the chemotactic effect of CXCL12 and also significantly inhibited *HsMIF*-6xHis-mediated monocyte migration (Fig. 6 and Fig. S5). Co-application of ISO-1 blunted the effect of *HsMIF*-6xHis, but did not interfere with CXCL12-triggered migration, confirming the specificity of MIF-driven monocyte chemotactic responses in our experimental system. Strikingly, both inhibitors completely ablated the chemotactic effect of *AtMDL1*-6xHis, which was studied as a representative of the three *AtMDLs*. Although not statistically significant, the drugs lowered the chemotactic effect even below baseline levels, and their effect on *AtMDL1*-6xHis appeared to be even more potent than that on *HsMIF*-6xHis. Together, these data suggested that the *AtMDLs*, in particular *AtMDL1*, are capable of triggering human monocyte recruitment with an efficiency similar to human MIF via interaction with the monocyte-expressed chemokine receptor CXCR4.

Plant MIF orthologs act as chemoattractants for human T cells and desensitize T-cell chemotaxis induced by human CXCL12 or MIF

THP-1 cells are monocyte-like human cells but have leukemic properties. To further test the significance of the chemotactic activity of the *AtMDLs*, we wished to study primary cells to ask whether other leukocyte cell types that express CXCR4

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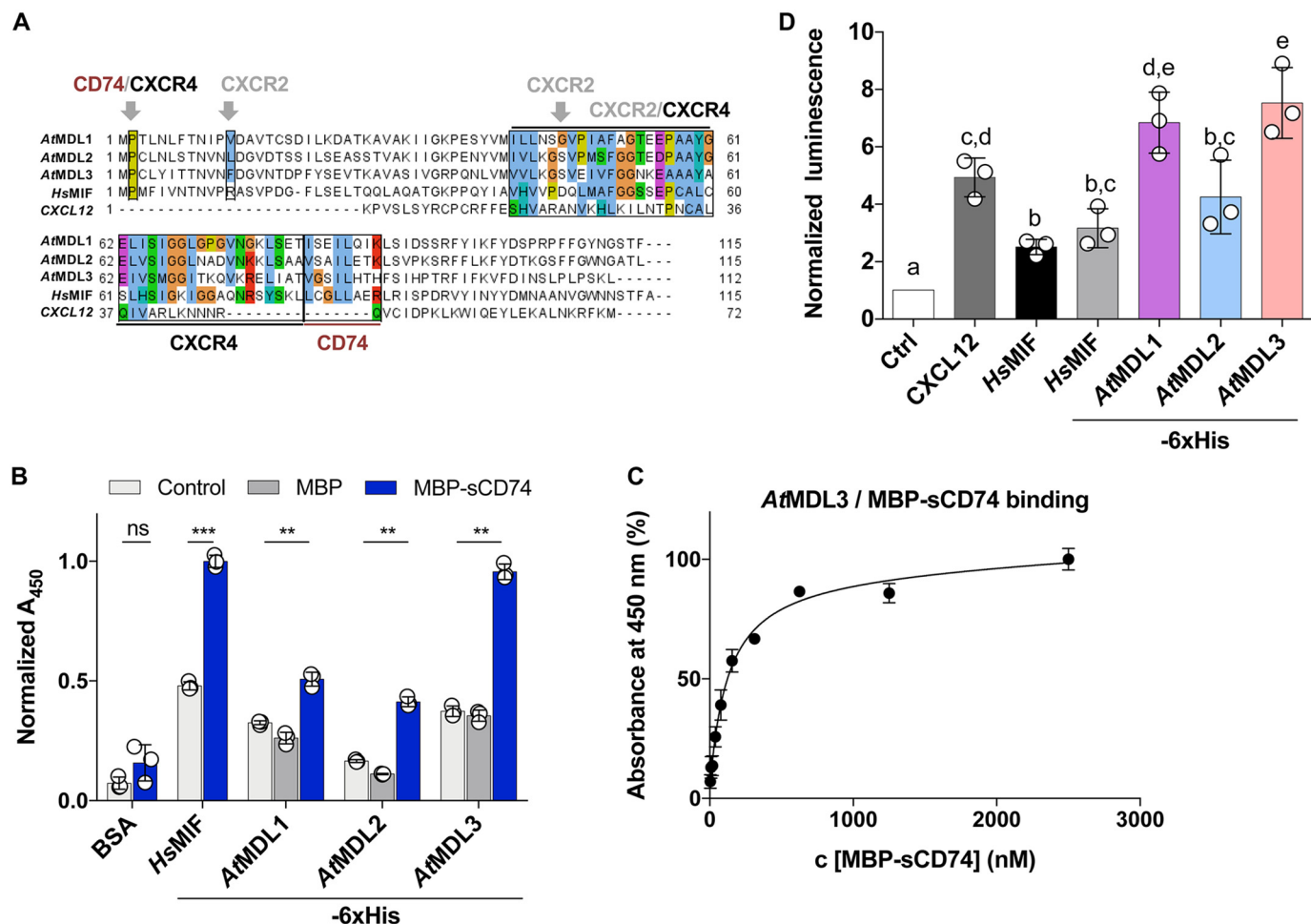


Figure 3. AtMDLs share homology with human MIF in the MIF receptor-binding sites, bind to CD74, and activate CXCR4-mediated signaling in a yeast-based reporter system. *A*, multiple sequence alignment of the AtMDLs, HsMIF, and human CXCL12. Amino acid sequences of AtMDL1 (identifier Q9LU69), AtMDL2 (identifier Q9M011), AtMDL3 (identifier Q8LG92), and HsMIF (identifier P14174) were retrieved from the UniProt database and aligned by ClustalW using standard parameters in the Jalview multiple sequence alignment editor desktop application. The amino acid residues contributing to the site I and II binding interface between HsMIF and CXCR4 (41) or CXCL12 and CXCR4 (90, 91), the binding sites between HsMIF and CD74, and the predicted corresponding regions in the AtMDLs are indicated. Determinants of HsMIF contributing to CXCR2 binding, although not further examined in this study, are indicated for comparison. The degree of homology/identity of the MIF, CXCL12, or AtMDL residues in these regions is highlighted by the following color score: blue, hydrophobic; red, positively charged; magenta, negatively charged; green, polar; pink, cysteine; orange, glycine; yellow, proline; cyan, aromatic; white, unconserved. *B*, comparison of the *in vitro* binding capacity between HsMIF-6xHis and MBP-sCD74 with that of the three His-tagged AtMDLs. Binding was measured by an ELISA-type plate-binding assay. BSA, blank PBS buffer (control), and MBP alone served as negative controls as indicated to account for nonspecific binding effects. Wells were coated with BSA (2% w/v), 500 nM HsMIF, and AtMDL1-6xHis, AtMDL2-6xHis, and AtMDL3-6xHis (500 nM), followed by binding of MBP or MBP-sCD74 (500 nM). After signal development, absorbance at 450 nm was measured, and the signals were normalized by setting the absorbance of HsMIF as 1. *C*, curve for binding of MBP-sCD74 and AtMDL3 using increasing concentrations of MBP-sCD74 as indicated. The data in *B* and *C* are displayed as means \pm S.D. ($n = 3$); (scatter plot with white circles indicates individual data points); ns, not significant; ***, $p < 0.001$; **, $p < 0.01$. *D*, CXCR4-mediated signaling in a yeast-based reporter system. In this assay, the Ste2 GPCR of the pheromone-response pathway of *S. cerevisiae* was substituted by the human CXCR4 receptor. Ligand binding to CXCR4 triggers signaling and expression of the *lacZ* gene, as assessed by β -gal activity. The concentrations of native (untagged) HsMIF, HsMIF-6xHis, and His-tagged AtMDLs were 20 μ M each. The concentration of human CXCL12 was equal to 2 μ M. Reporter activity is given as relative luminescence, normalized to the untreated control (Ctrl). Values shown represent means \pm S.D. from three independent experiments, in which the activity of each was assessed in technical duplicates (scatter plot with white circles indicates individual data points). Statistical analysis was performed using one-way ANOVA and post-hoc Tukey's HSD test with multiple comparisons. Different letters above the bars denote a statistically significant difference between groups ($p < 0.05$), and groups showing the same letters are not statistically significantly different from each other.

also chemotactically respond to the AtMDLs. Peripheral human blood-derived T cells are primary leukocytes that express substantial levels of surface CXCR4 and whose chemotactic response has been shown to be triggered by MIF (17). When examining the chemotactic migration of primary human T cells obtained from healthy donors, the dose optimum for HsMIF-6xHis was found to be between 16 and 32 nM (chemotactic index = 2.3) compared with a chemotactic index of 3.5 for 8 nM CXCL12. Of note, all AtMDLs elicited T-cell chemotaxis, and their pro-chemotactic capacity was compara-

ble with their activity on monocyte migration, except that the optimum dose for AtMDL1-6xHis and AtMDL2-6xHis was at 16 nM (chemotactic index = 2–2.5) and that a significant migratory effect of AtMDL3-6xHis was only seen at a concentration 160 nM (Fig. 7A). Thus, AtMDLs also serve as chemoattractants for primary human T cells.

We employed this physiologically-relevant cell system to mechanistically explore the potential interplay between the AtMDLs and the endogenous human CXCR4 agonists CXCL12 and MIF, hypothesizing that the plant MIF orthologs might act

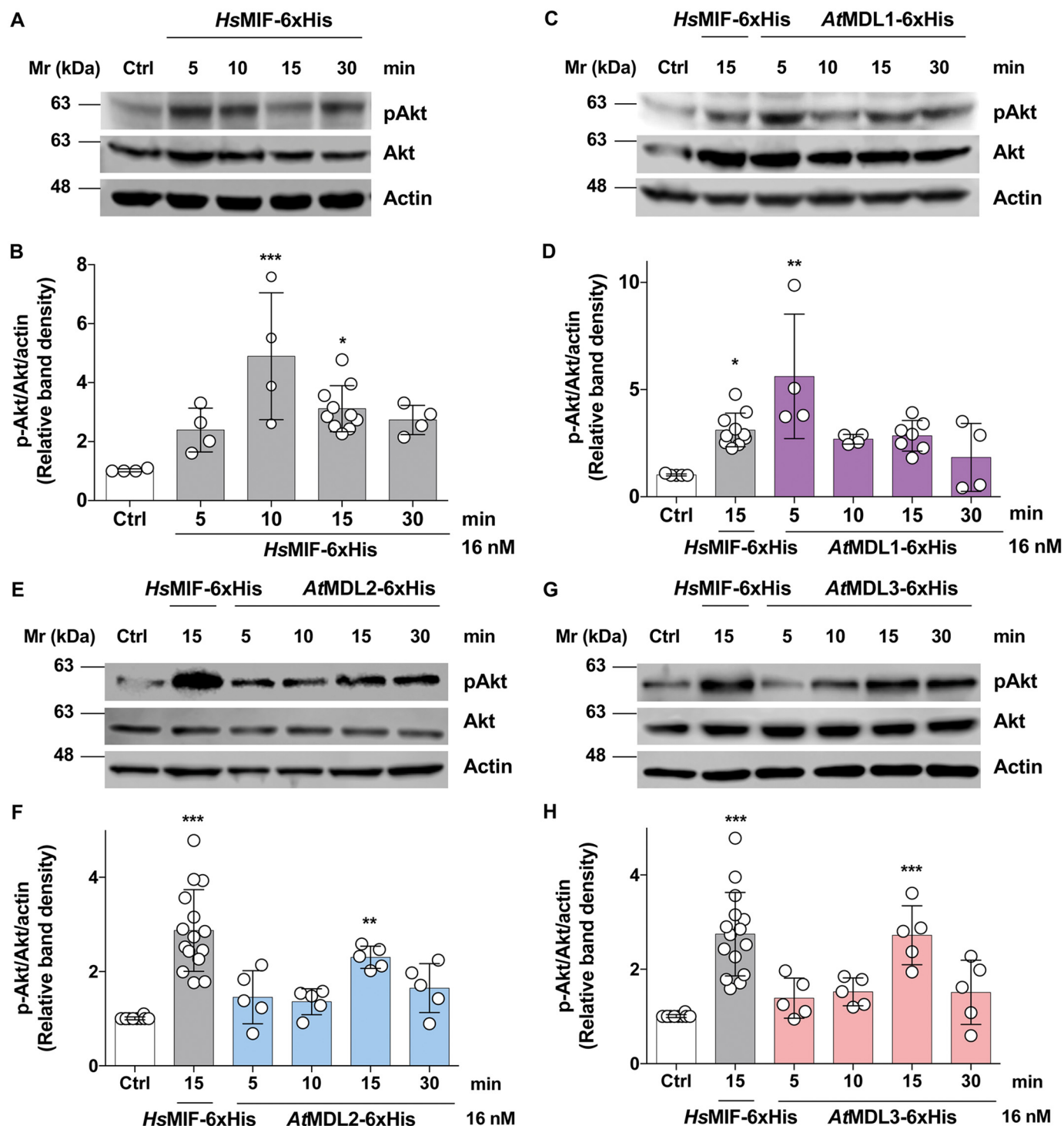


Figure 4. *AtMDLs* activate the CXCR4-PI3K/Akt-signaling pathway in human CXCR4-transfected HEK293 cells. *A*, representative Western blotting indicates Akt phosphorylation (pAkt) at different time intervals as indicated following stimulation with *HsMIF-6xHis* at a concentration of 16 nM. *Ctrl*, untreated control sample. Total Akt and actin were analyzed as loading control and for quantification purposes. *B*, quantification of pAkt band intensities in relation to Akt and actin band intensities according to the Western blot analysis in *A*. *Bar graph* represents means \pm S.D. of 5–15 experiments (scatter plot with white circles indicates individual data points). Statistical analysis was performed using one-way ANOVA between the untreated control (*Ctrl*) and the various time points following treatment (*, $p < 0.05$; ***, $p < 0.005$). *C*, *E*, and *G*, same as *A* except that the time-dependent phosphorylation of Akt following treatment with *AtMDL1-6xHis*, *AtMDL2-6xHis*, and *AtMDL3-6xHis*, respectively, was analyzed and compared with the effect of *HsMIF-6xHis* at 15 min. *D*, *F*, and *H*, same as *B*, except that the quantification refers to the Western blot analysis in *C*, *E*, and *G* and that data are from 5 to 10 experiments (*, $p < 0.05$; **, $p < 0.01$; ***, $p < 0.005$).

to modulate T-cell chemotactic responses induced by the human agonists. We specifically asked whether the *AtMDLs* would be able to enhance or desensitize T-cell chemotaxis responses triggered by the human CXCR4 chemokines. Pre-

treatment with all three hexahistidine-tagged *AtMDLs* added to the upper chamber of the Transwell setup fully ablated the chemoattractant activity of 16 nM *HsMIF-6xHis* in the lower chamber (Fig. 7*B*). Similarly, all three *AtMDLs* also attenuated

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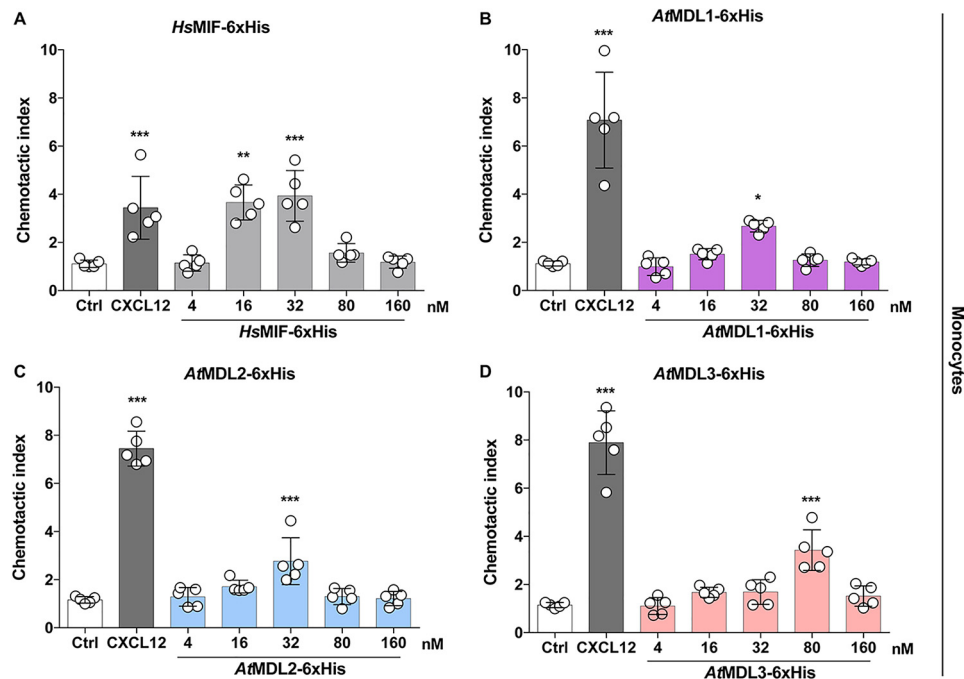


Figure 5. Recombinant 6xHis-tagged *AtMDLs* trigger chemotactic migration of human THP-1 monocytes in a dose-dependent manner. Chemotaxis (referred to here as chemotactic index) of THP-1 monocytes toward *HsMIF*-6xHis (A), *AtMDL1*-6xHis (B), *AtMDL2*-6xHis (C), or *AtMDL3*-6xHis (D) at the different indicated concentrations. The chemotactic potency was compared with human CXCL12 (at a concentration of 8 nM) serving as a positive control and to buffer control (*Ctrl*), which also served to normalize treatments to spontaneous (random) migration events. The bar graphs show means \pm S.D. of five independent experiments (scatter plot with white circles indicates individual data points). Statistical analyses were performed using one-way ANOVA between the buffer control and the treatment groups at the various doses (*, $p < 0.05$; **, $p < 0.01$; ***, $p < 0.005$).

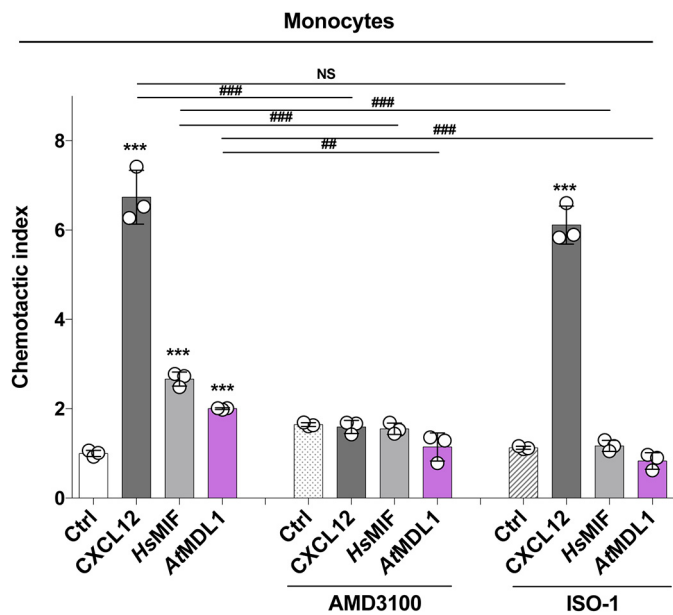


Figure 6. *AtMDL1*-6xHis-triggered monocyte chemotaxis is blocked by the small molecule inhibitors AMD3100 and ISO-1, indicating MIF and CXCR4 specificity. Chemotaxis experiments were performed as shown in Fig. 5 in the absence or presence of the small molecule inhibitors AMD3100 (10 μ M) or ISO-1 (100 μ M) as indicated. Concentrations of the recombinant proteins were equal to 32 nM for *HsMIF*-6xHis and *AtMDL1*-6xHis and 8 nM for CXCL12. Sodium phosphate buffer was used to normalize treatments to spontaneous random migration (control, *Ctrl*). Bar graph shows means \pm S.D. of one of two independently performed experiments carried out as technical triplicates each (for the other experiment see Fig. S5) (scatter plot with white circles indicates individual data points). Statistical analysis was done using one-way ANOVA for comparisons within a group (***, $p < 0.005$) and paired *t* test for comparisons between control and the AMD3100 and ISO-1 treatment groups (##, $p < 0.01$; ###, $p < 0.005$; NS, not significant).

the chemotactic effect of CXCL12, although full blockade was only seen for *AtMDL3*-6xHis (Fig. 7B). As verified by an experiment adding polymyxin B, which neutralizes endotoxin, together with the *AtMDL* pretreatment regimen, the desensitization effect was not due to minute endotoxin contaminants in our *AtMDL*-6xHis preparations (Fig. S6). These data therefore suggested that treatment of primary human T cells with *AtMDLs* desensitizes the cells for subsequent exposure to chemokines signaling through CXCR4. Plant MIF orthologs can thus directly modulate human immune cell behavior elicited by human chemokines.

Discussion

Chemokine-orchestrated immune cell trafficking is a central regulatory mechanism of the host immune response, but pathogens have developed intricate mimicry mechanisms to compromise the host chemokine system at both the ligand and receptor level. This capability is typically limited to viruses (7, 57). Prominent examples include the following: (i) lentiviridae such as HIV with the envelope protein gp120 binding to human CXCR4; (ii) γ -herpesviridae such as HHV8 that encode for the viral CC-chemokine mimic vMIP-II that functions as a chemoattractant and also binds to CXCR4; and (iii) β -herpesviridae such as human cytomegalovirus that encodes for the soluble CKR US28, serving to sequester host chemokines (7, 57, 58). Molecular mimicry of chemokines or their receptors by bacteria and fungi has not been observed, but indirect mechanisms to manipulate the host chemokine system have been reported, e.g. fungal mimicry of a mammalian dipeptidyl-peptidase that cleaves and inactivates CCL2 (59). Upon first view, chemokine mimicry mechanisms are counterintuitive for plants. MIF pro-

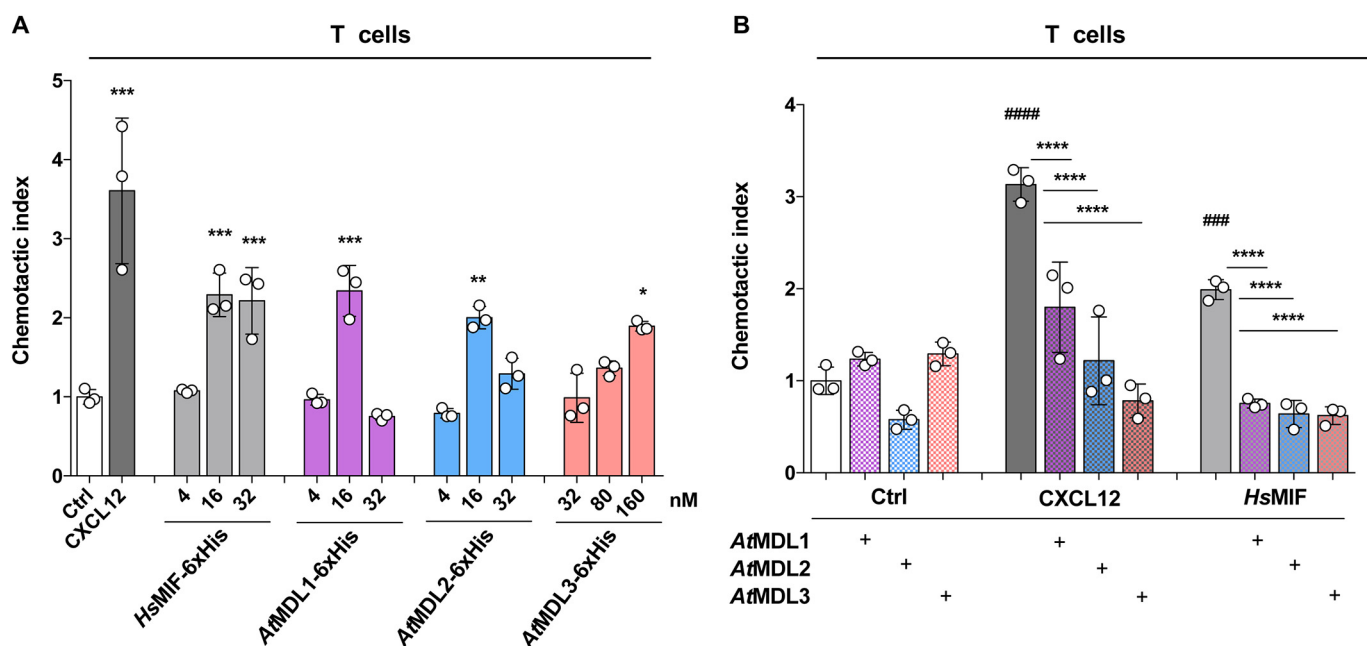


Figure 7. Recombinant 6xHis-tagged AtMDLs dose-dependently trigger chemotactic migration of primary human T cells and desensitize T cells from CXCR4 agonist-triggered chemotaxis. *A*, recombinant 6xHis-tagged AtMDLs trigger chemotactic migration of primary human T cells in a dose-dependent manner. The chemotactic potency was compared with HsMIF-6xHis and human CXCL12 (at a concentration of 8 nM) serving as a positive control and to buffer control (Ctrl), which also served to normalize treatments to spontaneous (random) migration events. Bar graphs show means \pm S.D. of three independent experiments (scatter plot with white circles indicates individual data points). Statistical analyses were performed using one-way ANOVA between the buffer control and the treatment groups at the various doses (*, $p < 0.05$; **, $p < 0.01$; ***, $p < 0.005$). *B*, recombinant 6xHis-tagged AtMDLs desensitize T cells from chemotaxis elicited by CXCL12 or HsMIF-6xHis. Data are the same as in *A*, except that T cells in the upper chamber were preincubated with the His-tagged AtMDLs for 2 h (+), before being subjected to chemoattractant exposure (CXCL12 or HsMIF-6xHis) in the lower chamber. Control (Ctrl) incubations were performed without chemoattractant in the lower chamber (random migration). Bar graphs show means \pm S.D. of three experiments (scatter plot with white circles indicates individual data points). Statistical analyses were performed using one-way ANOVA between control and CXCL12 or HsMIF (###, $p < 0.005$; ####, $p < 0.001$) and between CXCL12 or MIF with and without pre-treatment with AtMDLs (****, $p < 0.001$).

teins are atypical chemokines that control pathogenic cell recruitment in human diseases, such as atherosclerosis and cancer through noncognate interactions with classical CKRs such as CXCR4 (8, 9, 11, 13, 17, 24, 60). CKR engagement by MIF is reminiscent of chemokine mimicry mechanisms (13). Furthermore, MIF proteins are characterized by a unique enzymatic tautomerase activity that is unprecedented in the chemokine/cytokine group of proteins and that is a notable component of the remarkable evolutionary conservation of the MIF sequence (8, 25, 61). MIF has been implicated as a player in various host–parasite interactions, and MIF orthologs have been predicted to exist even in the plant kingdom (32, 33, 38).

Here, we have cloned and experimentally characterized three MIF proteins from the model plant *A. thaliana*, i.e. *A. thaliana* MIF/D-DT-like (MDL) proteins (AtMDLs), and we provide evidence that they are all structurally similar to human MIF (Fig. 1D and Figs. S1B and S3) and are able to mimic biological activities of human MIF (Figs. 3–7). Surprisingly, mimicry of human MIF by AtMDLs is not related to the conserved MIF tautomerase activity, which is almost absent in AtMDLs (Fig. 2 and Table 1), but AtMDLs were found to interact with the human MIF receptors CD74 and CXCR4. Importantly, we discovered that they activate CXCR4-mediated Akt signaling (Fig. 4) and monocyte and T-cell chemotaxis (Figs. 5–7). These activities represent cellular functionalities of MIF typically encountered in MIF-regulated inflammatory and/or host defense responses (8, 9, 12, 25). The AtMDLs desensitized human T cells from responding to gradients of the human

CXCR4-ligating chemokines CXCL12 and MIF (Fig. 7), offering one potential mechanism of how plant MIF orthologs might influence human immunity.

Our findings indicate that AtMDLs preserve a sufficient level of sequence conservation (Fig. 3A) and conformational similarity (Fig. 1D) to enable binding to the human MIF receptors CD74 and CXCR4, thereby allowing them to mimic functions of the endogenous human MIF or to influence human MIF or CXCL12 responses. Notably, although our experiments were only performed with MDLs from the model plant *A. thaliana*, this is to the best of our knowledge the first report demonstrating that plant proteins can phenocopy or “mimic” inflammatory activities of a *bona fide* human cytokine/chemokine.

Moreover, although plant MDLs were identified in data banks and their sequences and structures predicted by *in silico* methods (33, 38), our current study offers the first experimental characterization of plant MDLs. In fact, little is known about plant MDLs. The *in silico* analyses trace back MDLs several hundred million years in evolution and predict a role for them in plant stress-response pathways (33, 38), but an experimental validation of this presumption has been elusive. Also, there has only been a single experimental study in which plant MDL proteins have been “indirectly” implicated. Naessens *et al.* (62) previously showed that a MIF ortholog secreted in the saliva of a plant–parasitic insect modulates the immune response of the host plant. However, whether host plant MDLs are involved in this effect has remained unresolved.

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To study experimentally the three predicted *AtMDLs*, we opted to express these proteins recombinantly in a standard *Escherichia coli* expression system and use a C-terminal hexahistidine tag for purification purposes. Importantly, comparison of the structural and functional properties of C-terminally hexahistidine-tagged *HsMIF* with those of native untagged *HsMIF* in all experiments of this study essentially excludes the possibility that the hexahistidine tag artificially influences the observed mimicry effects of *AtMDLs*. These findings suggest that mimicry of human MIF activities by *AtMDLs* is a true property of these plant proteins.

Unique among cytokines/chemokines, MIF proteins contain a conserved tautomerase cavity that they share with the tautomerase superfamily, an evolutionarily-conserved protein family, the members of which feature an invariant N-terminal proline residue and a characteristic β - α - β -fold (28, 61). In fact, the entire 3D architecture of the MIF structure is remarkably similar to members of this family, such as the bacterial enzymes 4-oxalocrotonate tautomerase (4-OT) or 5-(carboxymethyl)-2-hydroxyisocaproate isomerase (CHMI), and it is similar between MIF-like proteins from different species and kingdoms, as determined by X-ray crystallographic analysis or predicted by *in silico* analyses (61). The tautomerase superfamily is characterized by catalytic promiscuity and diversity and has been suggested to be derived from a common ancestor by divergent evolution (33, 61). The known catalytic tautomerization activity of human MIF is limited to the nonnatural substrate D-dopachrome or DCME and to HPP. Whereas the keto-enol tautomerization of HPP has been generally associated with tyrosine and phenylalanine metabolism, a role for MIF in this process in mammalian cells has not been detected. Moreover, a physiological or pathophysiological relevance of the tautomerase activity of human MIF remains to be demonstrated. Nevertheless, the high degree of sequence similarity among members of the tautomerase superfamily and the conservation of the tautomerase consensus motif in plant MDLs prompted us to speculate initially that the tautomerase activity would be the basis of potential functional similarities, if any, between plant and human MIF proteins. In fact, mammalian MIF proteins have been suggested to exert dual roles with functions both in the extracellular space as cytokines/atypical chemokines and in the intracellular compartment as regulators of cell homeostasis and gene transcription (13, 34, 63). The extracellular cytokine/chemokine activities of MIF proteins are mediated by high-affinity binding to the cell-surface receptors CD74, CXCR2, and CXCR4 (and CXCR7) that are typically expressed in immune cells, and thus they represent the activities of a prototypical innate cytokine/chemokine (9, 16, 17). The molecular basis of the intracellular activities of MIF proteins is much less defined, but protein-protein-binding events and redox processes possibly also involving MIF's catalytic capacity have been implied (13, 25, 34, 63). Functional dichotomy as both the extracellular cytokine/chemokine and intracellular regulator is not unique to MIF proteins but has also been reported for proteinaceous alarmins such as the high-mobility-group box protein 1 (HMGB1) or some ribosomal tRNA transferases (13).

Given these considerations, the observed dramatic reduction in tautomerase activity of the *AtMDLs* as measured in both the

HPP and DCME enzymatic assays is surprising (Fig. 2, A and B). Both assays are widely applied to evaluate tautomerase activities of MIF and its variants, and *HsMIF*-6xHis displayed full catalytic activity comparable with native untagged human MIF (Fig. 2A). It is likely that the almost complete lack of tautomerase activity in the *AtMDLs* is a general property of these orthologs. Neither D-dopachrome nor DCME has been identified in plants, and although L-dopachrome has been implicated as a metabolite that is synthesized in some plant species and expelled via root exudates to compete out other plant species, *A. thaliana* is not a producer of this substance (64), together suggesting that dopachrome or its derivatives are unlikely to be substrates for MDLs in *Arabidopsis*. Similarly, although HPP-metabolizing enzymes have been found in plants, not much is known about the function of this metabolite, let alone its connection with MDL proteins (65). The observed marked reduction of tautomerase activity in *AtMDLs* may be explained by the following mechanism. Although the K_m values of HPP and DCME are higher than that of human MIF, they are in the same order of magnitude, with a 3.5- and 8-fold lower apparent affinity for the plant MDLs, respectively (Table 1), suggesting these substrates may be capable of binding into the tautomerase pocket of *AtMDLs*. However, the measured k_{cat} values indicate that catalytic conversion of both substrate types is highly inefficient in the *AtMDLs*. In fact, although the overall 3D consensus motif of the tautomerase site is conserved in the three *AtMDLs*, we noticed a potentially critical amino acid substitution at position 98, in which a positively-charged lysine residue in all three *AtMDLs* replaces the neutral asparagine residue of *HsMIF* (Fig. 2C). In fact, a model suggested by our *in silico*-modeling analysis predicting that this could trigger a conformational change and an altered charge distribution profile in the vicinity of the pocket (Fig. 2D) was underpinned experimentally by the generation of a site-specific mutant of human MIF, in which Asn-98 was replaced by Lys-98. N98K-*HsMIF* showed a greatly reduced catalytic activity using both HPP and DCME as substrate (Fig. 2, E and F). Thus, the replacement of Asn-98 by Lys-98 in the *AtMDLs* may be the basis for an impeded catalytic turnover in plant MDLs. Furthermore, the observed inhibitory effect of the MIF tautomerase inhibitor ISO-1, which shares structural similarities with HPP and not only fully blocked *HsMIF*- but also *AtMDL*-mediated monocyte migration (Fig. 6), furthers the notion that HPP (and DCME) binding still occurs, but that substrate tautomerization is not efficiently catalyzed by *AtMDLs*. The further resolution of these mechanistic questions will have to await the 3D structural characterization of *Arabidopsis* MIF orthologs by X-ray crystallography and the elucidation of *AtMDL*/inhibitor co-complexes.

In considering alternative molecular mechanisms that could give rise to functional overlaps between plant MDLs and human MIF, we explored the possibility that *AtMDLs* might interact with human MIF receptors. At first glance, this appeared to be an unlikely option. Neither CD74 nor the MIF chemokine receptors CXCR2 or CXCR4 are present in plants. In fact, the existence of *bona fide* GPCRs in the plant kingdom is still controversially discussed, and it seems that only few candidates exist, none of which has been functionally linked to signaling via heterotrimeric G-proteins (66). Plants lack a cir-

cultivation-based immune system and deploy nonproteinaceous phytohormones such as salicylic acid, jasmonic acid, and ethylene as pivotal players for immune signaling (67). However, intriguingly, we found that *AtMDLs* not only share with *HsMIF* some degree of sequence homology regarding Pro-2 and in the site 2 binding region of the MIF/CXCR4 interface (Fig. 3A), but they elicited CXCR4-mediated cell signaling responses in both a yeast-based model cell system (Fig. 3D) and mammalian HEK293 cells (Fig. 4), comparable with the signaling effects triggered by *HsMIF*. Moreover, the data obtained in the yeast transfectant system, which specifically expresses human CXCR4 but none of the other MIF receptors nor other mammalian proteins, argue that *AtMDLs* are able to directly bind to human CXCR4. This appears surprising at first sight, but CXCR4, which was long regarded as a highly-specific chemokine receptor that only binds to one chemokine ligand, namely its cognate ligand CXCL12, has more recently been recognized to be fairly promiscuous, binding to several non-*bona fide* chemokine ligands. These include the atypical chemokine MIF, viral macrophage inflammatory protein II (vMIP-II), viral HIV gp120, human β -defensin-3 (HBD3), and extracellular ubiquitin (9, 13, 17, 56, 57, 68–70). As it seems unlikely that there has been any evolutionary pressure on plant MDLs to develop agonistic properties for a human CKR, we speculate that the CXCR4-binding capacity of plant MDLs may have developed as an “evolutionary side-reaction” of plant interactions with pathogenic animals, such as recently described for plant–parasitic aphids feeding on *Nicotiana benthamiana* leaves (62). The latter scenario is somewhat reminiscent of vertebrate–parasite interactions that rely on cross-species utilization of MIF ligand/receptor pathways (71). Alternatively, the capacity of plant MDLs to activate human CKRs could be the indirect consequence of constraints imposed on MIF evolution to maintain one or several core functions of this protein family.

Although interactions between *AtMDLs* and the cognate human MIF receptor CD74 were only studied by *in vitro* binding experiments on a biochemical level (Fig. 3, B and C), the data insinuate that *AtMDLs* might also influence human MIF-triggered CD74 responses. On the one hand, this appears unlikely, as plants do not express a major histocompatibility (MHC) system. On the other hand, CD74 has been found to be an amenable target of parasite MIF orthologs such as leishmania, plasmodium, or hookworm (32, 35, 37, 71, 72), suggesting that it is a receptor molecule prone to be engaged during host–parasite interactions. In fact, it has been suggested that the MIF-binding functionality of CD74, which is best known as the MHC class II chaperone invariant chain Ii, already represents a “secondary” function of this membrane protein (16, 73).

Understanding the multiple functions of mammalian MIF proteins that are mediated by specific interactions with four cell-surface receptors and several intracellular-binding partners has been challenging. We have only begun to decipher the binding determinants that govern the molecular promiscuity of the interactions between MIF proteins and these binding partners (13, 17, 34, 40, 41, 46, 60, 74). Including the herein-described three *AtMDLs* in corresponding comprehensive structure–activity relationship studies will add important information to the sequence and 3D motifs that specify such

interactions. Accordingly, it should assist in gathering further insight for novel site-specific drug discovery approaches against human MIF that could eventually be beneficial in treating human diseases such as atherosclerosis and cancer that are at least partially mediated by MIF pathways (17, 75).

The most notable result of this study is the observation that all three *AtMDLs* were found to promote monocyte and T-cell chemotactic migration (Figs. 5 and 7A). Moreover, the measured chemotactic effect for *AtMDL1* and -2 peaked at the same concentration as that of *HsMIF*, albeit the chemotactic index was lower (2.5–3-fold *versus* 3.5–4-fold, respectively). Importantly, ablation of the effect by the small molecule MIF tautomerase inhibitor ISO-1 (51), which also inhibits the MIF–CXCR4 interaction (41), verified MIF specificity of the observed migration response (Fig. 6). It will be interesting to explore whether other documented small molecule MIF tautomerase inhibitors such as 4-IPP or MIF098 (25, 76) have a similar effect on *AtMDLs*. Monocytes express all four MIF receptors, but pre-incubation of the monocytes with the small molecule inhibitor AMD3100, which is an established specific CXCR4 inhibitor (53) and has already been shown to partially interfere with MIF binding to CXCR4 (41), blocked the *AtMDL*-mediated migration effect, providing evidence that *AtMDL*-triggered monocyte chemotaxis is in fact mediated by CXCR4 (Fig. 6). T cells only express CXCR4 (as well as the relatively poorly-characterized CXCR7), but not CD74 or CXCR2, suggesting that the observed chemotaxis induced by the plant MDLs as well as the desensitization effect toward subsequent human MIF or CXCL12 chemotaxis are also mediated by CXCR4 (Fig. 7, A and B, and Fig. S6).

Thus, *AtMDLs* have the surprising capacity to manipulate human immune cell motility via “hijacking” the chemokine receptor CXCR4. This raises a number of follow-up questions and hypotheses. As we already obtained biochemical-binding evidence, it could be asked whether *AtMDLs* also functionally affect MIF-driven CD74 responses. Moreover, *AtMDLs* might interact with CXCR2, the other chemokine receptor that human MIF engages to modulate immune cell migration and that is a prominent atherogenic arrest (77) and neutrophil recruitment receptor (78). Our results may also justify the hypothesis that MDLs from other plant species have a similar ability to engage human MIF receptors and to modulate mammalian immune cell responses.

Our study, which to the best of our knowledge shows for the first time that a plant protein with homology to a mammalian cytokine/chemokine can interact with two human cytokine receptors, may have broader implications. Following contact with plant MDLs, *e.g.* through the respiratory or the gastrointestinal tract during respiration or dietary ingestion, immune cells in the surrounding mammalian tissue might be modulated in their migratory activity by plant MDLs. Our T-cell migration desensitization data (Fig. 7) argue that this could be one mechanism how MDLs modulate human immunity. Although it is well-known that plant proteins, *e.g.* from pollen, can function as allergens to hyper-activate human adaptive immunity (79), our data may also suggest effects on components of the innate human immune system. Plant MDLs might enhance or suppress MIF-dependent innate immune responses, and this may

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have a role in tissue homeostasis or in MIF-driven diseases such as acute or chronic inflammation, cardiovascular conditions, or cancer. This hypothesis will have to be tested in suitable experimental *in vivo* models in the future.

Materials and methods

Cell lines, reagents, and antibodies

The human monocytic cell line THP-1 and primary human T cells were grown in complete RPMI 1640 medium supplemented with GlutaMAXTM, 10% fetal calf serum (FCS), and 1% penicillin/streptomycin. Primary human T cells were isolated from enriched peripheral blood mononuclear cell (PBMC) fractions using the human Pan T-cell isolation kit from Miltenyi Biotec (Bergisch Gladbach, Germany). PBMC fractions were obtained by apheresis via a Leucoreduction System Chamber (“Kegel”) from anonymous thrombocyte donations at the Department of Transfusion Medicine, Cell Therapeutics, and Hemostaseology of the Klinikum der University Hospital (KUM) of the Ludwig-Maximilians-University (LMU). The studies abide by the Declaration of Helsinki principles and were approved by ethics approval 18-104 of the Ethics Committee of LMU Munich entitled “The MIF Protein/Receptor Network in Atherosclerosis” and encompasses the use of anonymized tissue and blood specimens for research purposes.

The human embryonic kidney cell line HEK293 was cultured in Dulbecco’s minimal essential medium (DMEM) supplemented with GlutaMAXTM, 10% FCS, and 1% penicillin/streptomycin. To obtain HEK293 transfectants stably expressing the human CXC chemokine receptor CXCR4, a high-affinity receptor for both CXCL12 and MIF, WT HEK293 cells were stably transfected with the pcDNA3.1 plasmid containing a 3xHA–*HsCXCR4* insert under the control of a human cytomegalovirus immediate–early promoter by an established procedure as described previously (17, 41). THP-1 monocytes, T cells, and HEK293 cells were incubated at 37 °C in a humidified atmosphere containing 5% CO₂. All cell culture reagents were cell culture grade and were obtained from Thermo Fisher Scientific (Waltham, MA).

Other reagents, *e.g.* salts, chemicals, and miscellaneous reagents, were of the highest research grade possible and were purchased from Merck KGaA (Darmstadt, Germany) and Carl Roth GmbH (Karlsruhe, Germany). Imidazole was from Sigma GmbH (Taufkirchen, Germany), and skimmed milk was from SERVA electrophoresis GmbH (Heidelberg, Germany).

Antibodies used in this study were as follows: polyclonal rabbit anti-actin (A2066; Sigma), polyclonal rabbit anti-Akt (catalog no. 9272; Cell Signaling Technologies, Danvers, MA), and polyclonal rabbit anti-phospho-Akt (Ser-473) (catalog no. 9271; Cell Signaling Technologies).

Multiple sequence alignments

Multiple sequence alignments were performed by the ClustalW algorithm (<http://www.genome.jp/tools-bin/clustalw>) using standard parameters in the Jalview multiple sequence alignment editor desktop application (80–82). Sources of the sequences used in the alignment are given in Table S1.

Cloning of *HsMIF* and the three *AtMDLs* with C-terminal hexahistidine tags

HsMIF and the three *A. thaliana* MIF/DDT-like (*MDL*) genes, *AtMDL1*, *AtMDL2*, and *AtMDL3*, were cloned into the pET21a vector via a classical cloning strategy. The genes were N-terminally fused in-frame to a 6xHis tag present in the vector using the restriction enzymes *NdeI* and *XhoI*. For this purpose, the respective restriction sites were added to the desired cDNAs in a PCR using the corresponding primers *NdeI*–“gene”_Fwd and “gene”–*XhoI*_Rev (Table S4). The reverse primer at the same time served to remove the endogenous stop codon. For *AtMDL1* and *AtMDL3*, an internal *NdeI* restriction site was removed using the splice overlap–extension PCR strategy (SOE-PCR) with the internal primer “gene”–mut_Fwd and “gene”–mut_Rev (Table S4) in combination with the above-mentioned restriction site-adding forward and reverse primers. Through the SOE-PCR, the cDNA sequences of *AtMDL1* and *AtMDL3* were modified at position 177 (GCA to GCG), introducing a silent mutation (A59A). Successfully-fused plasmid products were confirmed by sequencing, propagated in One ShotTM TOP10 chemically-competent *E. coli* (Thermo Fisher Scientific), and then transformed into RosettaTM (DE3)-competent *E. coli* (Novagen/Merck KGaA) for expression of recombinant protein. The N98K–*HsMIF* mutant gene was synthesized by and purchased from BaseClear (Leiden, The Netherlands). After subcloning from the pUC57 plasmid into pET21a and transformation of competent RosettaTM (DE3) *E. coli* cells, all following procedures were identical to those described above.

Expression and purification of proteins

Recombinant nontagged human CXCL12 and *HsMIF* proteins, used as controls in this study, were cloned, expressed, and purified as described before (46). RosettaTM (DE3)-competent *E. coli* cells were used to express the pET21-derived gene constructs to yield *HsMIF*–6xHis-, *AtMDL1*–6xHis-, *AtMDL2*–6xHis-, *AtMDL3*–6xHis-, and N98K–*HsMIF*–6xHis-tagged protein products. Culturing of bacteria and protein expression was carried out essentially as described before (46).

For protein purification, cells were harvested by centrifugation, and cell pellets were resuspended in 2 ml of binding buffer (20 mM sodium phosphate, 0.5 M NaCl, 20 mM imidazole, pH 7.2). After homogenizing the bacteria at 75 megapascals using an Aventin EmulsiFlex C5 high-pressure homogenizer (ATA Scientific Pty. Ltd., Lucas Heights, Australia), recombinant hexahistidine-tagged proteins were initially purified using immobilized metal ion affinity chromatography (HisTrap; GE Healthcare, Freiburg, Germany) with an FPLC system (FPLC; ÄKTA Pure, GE Healthcare). Prior to the run, the FPLC system was equilibrated with binding buffer. His-tagged proteins were eluted using elution buffer (20 mM sodium phosphate, 0.5 M NaCl, 0.5 M imidazole, pH 7.2). Resulting protein fractions were stored at 4 °C until subsequent purification steps or experimental usage. Additional purification of the proteins was performed by size-exclusion chromatography (Superdex 75, GE Healthcare) using 20 mM sodium phosphate buffer, pH 7.2, as elution buffer, *i.e.* conditions previously reported to support stability

and bioactivity of recombinant *HsMIF* (17, 39). Subsequently, purified proteins were sterile-filtered using 0.2- μm pore size filters and then stored at 4 °C until use. Proteins were used for biochemical and biological assays within 4 weeks of purification. Endotoxin content was determined in the final enriched, sterile-filtered protein solution, using the PierceTM LAL Chromogenic Endotoxin Quantitation Kit (Thermo Fisher Scientific).

Western blotting analyses of recombinant *HsMIF*-6xHis and *AtMDL*-6xHis proteins

Assessment of protein purity and integrity was performed by SDS-PAGE, using 15% acrylamide gels under reducing conditions essentially as described (39). Proteins were detected by Coomassie Blue staining and/or Western blot analysis, using nitrocellulose membranes and a Novex[®] Tris-glycine transfer buffer and Tris-buffered saline (TBS: 150 mM NaCl, 20 mM Tris, pH 7.3), supplemented with 0.01% Tween 20 and 1% BSA for blocking/staining. His-tagged proteins were detected by mouse anti-6xHis tag mAb (Ma1-135; Invitrogen) followed by incubation with horseradish peroxidase (HRP)-conjugated secondary goat anti-mouse IgG (ab6789; Abcam, Cambridge, UK). In addition, for testing the relative structural similarity between the mammalian and plant orthologs, anti-*HsMIF* antibodies (anti-human MIF mAb MAB289; R&D Systems, Minneapolis, MN; anti-mouse MIF polyclonal rabbit antibody Ka565 (34)) were used, followed by HRP-conjugated secondary antibodies (catalog no. ab6789, Abcam; catalog no. P0448, DAKO) and imaging with SuperSignalTM West Dura Extended Duration Substrate (Thermo Fisher Scientific) on an Odyssey[®] Fc Imaging System with Image StudioTM software (LICOR Biosciences, Bad Homburg, Germany).

Far-UV CD spectroscopy

Far-UV CD spectroscopy was performed with a JASCO J-715 spectropolarimeter (JASCO, Tokyo, Japan). Measurements were carried out at room temperature between 195 and 250 nm at 0.1-nm intervals with a response time of 1 s in a buffer containing 10 mM sodium phosphate, pH 7.4. CD spectra were recorded at protein concentrations of 1, 2.5, and 5 μM in a 1-, 0.5-, or 0.2-cm quartz cuvette, respectively. The background spectrum of buffer alone was subtracted from the CD spectra of the protein solutions. The final data result from an average of three CD spectra. The CD spectra of native *HsMIF* and their deconvolution were published previously (46) and were represented in this study for comparison after verifying similarity/identity with a representative current native *HsMIF* preparation. Dynode voltage values were below 850 and did not interfere with CD measurements. Deconvolutions were performed using Dichroweb online software (<http://dichroweb.cryst.bbk.ac.uk>), and estimation of the secondary protein structure was carried out with the analysis program Contin LL, using the reference spectra Set 7 (83–86).

Tautomerase activity assays

The HPP tautomerase activity assay was performed as described before (87) with the following adjustments: 500 mM boric acid, pH 6.2, was used instead of 435 mM. Enzymatic mea-

surements were conducted in a solution of 8 mM HPP and 250 nM of the respective recombinant protein. An increase in absorbance due to complex formation was recorded at 306 nm every 5 s for a duration of 300 s.

The DCME tautomerase activity assay was performed essentially as described before (50, 88). Briefly, DCME was prepared at a final concentration of 1 mM by oxidizing *L*-3,4-dihydroxyphenylalanine methyl ester with 1 mM sodium *m*-periodate. DCME was then dissolved in 25 mM potassium phosphate buffer, pH 6.0, containing 0.5 mM EDTA, and enzyme (recombinant *HsMIF*-6xHis or *AtMDL1*-6xHis) was added at a final concentration of 100 nM. The decrease in absorbance at 475 nm was measured for 240 s in 10-s intervals.

All tautomerase activity experiments were performed using a JASCO version 650 spectrophotometer (JASCO).

MBP-sCD74/MIF-binding assay

The MBP-sCD74/MIF binding assay was performed essentially as described previously (45). Briefly, freshly thawed *HsMIF*-6xHis or *AtMDL*-6xHis aliquots were diluted into phosphate-buffered saline (PBS) to prepare a solution of 500 nM from which 100 μl was used for coating the wells of a medium-binding 96-well plate overnight at 4 °C. Wells were washed three times with 220 μl of washing buffer (PBS + Tween 0.05%) and subsequently blocked for nonspecific binding with 210 μl of a commercial blocker solution (Rockland Immunochemicals Inc., Limerick, PA) at room temperature for 30 min. During all incubation steps, the plate was shaken slowly. The blocker solution was removed, and the plate was washed three times with washing buffer. Subsequently, the wells were incubated with 100 μl of a 500 nM maltose-binding protein-soluble CD74 fusion protein (MBP-sCD74) solution in PBS for 30 min at room temperature. 100 μl of PBS were used as control at this step to exclude nonspecific binding of the anti-CD74 polyclonal antibody (pAb). After washing, wells were incubated with 100 μl of a rabbit anti-CD74 pAb solution (1:2500 dilution in PBS) (Sinobiological, Vienna, Austria) for 30 min at room temperature. After removing the anti-CD74 solution and washing, a solution of 100 μl of goat anti-rabbit horseradish peroxidase conjugate (1:2000 dilution in PBS) (Life Technologies, Inc., The Netherlands) was added and incubated for 30 min at room temperature. After washing, binding was visualized by conversion of 100 μl of aqueous tetramethylbenzidine solution (Sigma, Zwijndrecht, The Netherlands), which was quenched with an aqueous 1 N H₂SO₄ solution (100 μl). Absorbance was detected at 450 nm. A freshly-prepared solution of 2% (w/v) BSA in PBS was used as negative control for coating. An additional control was done to confirm that binding between the *AtMDL*s and MBP-sCD74 is not caused by the fused protein MBP. Toward this aim, the incubation with MBP-sCD74 was replaced by 500 nM MBP (ProSPEC Inc., Fullerton, CA) in PBS. MBP binding was detected using mouse anti-MBP mAb (1: 2500 dilution in PBS) (Sigma) as primary antibody and goat anti-mouse horseradish peroxidase conjugate (1:1500 dilution in PBS) (Thermo Fisher Scientific, Landsmeer, The Netherlands). The binding curve between *AtMDL3* with MBP-sCD74 was determined by using titrations of different concentrations of MBP-sCD74.

Mimicry of human cytokine activity by *Arabidopsis* orthologs

Yeast-signaling assay

The functional human CXCR4-expressing transformant of *S. cerevisiae* strain CY12946 was described before (46, 47). Briefly, agonist binding to the CXCR4 cell-surface receptor leads to the activation of a MAPK kinase-type signaling cascade initiating the transcription of the β -gal (*lacZ*) reporter gene. CXCR4 engagement and activation can therefore be measured and quantified by a β -gal enzymatic assay.

S. cerevisiae CY12946 cells were diluted to an OD₆₀₀ of 0.3–0.8 and incubated with the respective test proteins (*Hs*MIF–6xHis, *At*MDL1–6xHis, *At*MDL2–6xHis, and *At*MDL3–6xHis) or the established control agonists human CXCL12 and *Hs*MIF (41, 46). All test proteins were applied at a final concentration of 20 μ M, a concentration that had previously been shown to mediate MIF-based activation of CXCR4 in this system (41). CXCL12 was used at a final concentration of 2 μ M. It should be noted that in this system, elevated MIF and CXCL12 ligand concentrations are needed for appreciable receptor activation due to the barrier properties of the yeast cell wall. β -Gal activity was detected using the Beta-Glo[®] assay system (Promega Corp., Madison, WI). Luminescence signals were recorded in a multimode plate reader (Enspire[®] 2300, PerkinElmer Life Sciences).

Monocyte and T-cell chemotaxis assay

The chemotactic potential of hexahistidine-tagged *At*MDL proteins was assessed using the Transwell migration device essentially as described previously (18) using THP-1 monocytes or primary human T cells and an overnight transmigration interval. THP-1 and T cells were cultured in RPMI 1640 medium supplemented with 10% FCS and 1% penicillin/streptomycin. The evening before the assay, cells were transferred into standard RPMI medium without FCS. For the assay, the upper chambers of a 24-well format Transwell device (Sigma-Corning; pore size 5 μ m) were loaded with 1×10^6 THP-1 cells. Tested proteins (*Hs*MIF–6xHis, *At*MDL1–6xHis, *At*MDL2–6xHis, or *At*MDL3–6xHis) were studied at a concentration range of 4–160 nM, representing the chemoattractant in the lower chamber. As a positive control, 8 nM human CXCL12 was used; 20 mM sodium phosphate buffer, pH 7.2, served as negative control. For the desensitization experiment, 16 nM recombinant hexahistidine-tagged *At*MDLs were added to the upper T-cell-containing chamber 2 h before exposure to the chemoattractants in the lower chamber. In the endotoxin control experiment, 20 μ g/ml polymyxin B was added to the upper chamber together with the *At*MDLs.

Cells that migrated into lower chambers were quantitated and obtained values normalized to buffer control (“chemotactic index”) (17). The CXCR4 and MIF inhibitors AMD3100 (Sigma) and ISO-1 (Abcam, Cambridge, UK) were used at final concentrations of 10 and 100 μ M, respectively, and were added to the cell suspensions 30 min before the addition of the chemotactic stimulus. Following a 16-h migration interval, cells were removed from the lower chambers, mixed with Count-Bright[™] Absolute Counting Beads (Invitrogen/Thermo Fisher Scientific), and enumerated using a BD FACSVers[™] flow cytometer (BD Biosciences, Heidelberg, Germany).

Akt cell-signaling assay

HEK293 transfectants stably-expressing CXCR4 or non-transfected HEK293 control cells were subjected to treatment with *Hs*MIF–6xHis or *At*MDL1–6xHis, *At*MDL2–6xHis, or *At*MDL3–6xHis at a concentration of 16 nM. After different stimulation intervals, treated cells were lysed in NuPAGE[®] lithium dodecyl sulfate/dithiothreitol lysis buffer containing PhosSTOP[™] reagent (Roche Applied Science, Mannheim, Germany). Lysates were boiled at 95 °C for 5 min, sonicated for 5 min, and electrophoresed in 11% SDS-polyacrylamide gels. For Western blotting detection of phosphorylated Akt, proteins were transferred to nitrocellulose membrane using Novex[®] Tris-glycine transfer buffer. Human MIF, which has been previously shown to activate Akt signaling (48), was used for comparison. Phospho-Akt band densities were determined by usage of an anti-phospho-Akt(Ser-473) antibody and an HRP-conjugated secondary antibody. Total Akt and actin were used for normalization applying polyclonal rabbit anti-Akt and -actin antibodies. Band densitometry was performed with the Odyssey[®] Fc imaging system (LICOR) using Image Studio[™] software (LICOR Biosciences).

Fluorescence-activated cell sorting (FACS)

The cell-surface expression of the human CXCR4 receptor in HEK293 cells was verified by flow cytometry. HEK293 transfectants stably-overexpressing CXCR4 were grown in standard DMEM, supplemented with GlutaMAX[™], 10% FCS, and 1% penicillin/streptomycin, until the cells reached confluence. After washing in cold phosphate-buffered saline (PBS) buffer, cells were incubated with fluorescein-isothiocyanate (FITC)-labeled mouse anti-*Hs*CXCR4 or FITC-labeled isotype control antibody (IgG2a) (both R&D Systems) at 4 °C for 2 h in the dark. Cells were washed in cold PBS before subjection to FACS analysis using a FACSVers[™] flow cytometer (BD Biosciences).

Structure prediction

Prediction of the 3D protein structures of *At*MDL1–6xHis, *At*MDL2–6xHis, and *At*MDL3–6xHis was performed using the Phyre² Protein Fold Recognition Server (www.sbg.bio.ic.ac.uk/phyre2) (89). Modeling was carried out in intensive mode, using the amino acid sequences obtained from public repositories (Table S1) and accounting for the His-tag–modified variants used in this study. For structure prediction and modeling, Phyre² performs template-based modeling together with *ab initio* folding simulations for sequence segments for which no appropriate model was found.

Structure visualization

The predicted 3D protein structures of monomeric *Hs*MIF–6xHis and *At*MDL1–6xHis, *At*MDL2–6xHis, and *At*MDL3–6xHis were visualized using the PyMOL Molecular Graphics System Version 1.8.2.2 (Schrödinger Ltd. Liability Co.). The structures shown in this work correspond to the Protein Data Bank (PDB) file for human MIF (PDB identifier 3DJH) or our structure prediction results. The 3D structures were rendered both as a cartoon model showing the secondary protein structure and as a space-filling model depicting the protein surface.

This surface model was also used to visualize the electrostatic surface potential, as calculated by PyMOL. For visualization of the three 6xHis-tagged *AtMDL* proteins and their comparison with *HsMIF*, the initial methionine residue was removed, as it has been found to be processed in all cell systems studied so far (8, 12).

Statistics

Unless otherwise indicated, statistical analyses were performed using one-way analysis of variance (ANOVA) followed by post hoc comparison with the Bonferroni test using GraphPad Prism 6 (GraphPad Prism Software Inc., San Diego, CA) with multiple comparisons. Data are presented as means \pm S.D. Considered as significant: $p < 0.05$. Asterisks indicate statistically significant differences as follows: *, $p < 0.05$; **, $p < 0.01$; ***, $p < 0.005$; ****, $p < 0.001$. Letter symbols above bars in Fig. 3 indicate statistically significant different groups (at least $p < 0.05$) according to one-way ANOVA using multiple comparisons and Student's t-test.

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