Bone marrow stroma cells promote induction of a chemoresistant and prognostic unfavorable S100A8/A9high AML cell subset

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The bone marrow (BM) stroma represents a protective niche for acute myeloid leukemia (AML) cells. However, the complex underlying mechanisms remain to be fully elucidated. We found 2 small, intracellular, calcium-sensing molecules, S100A8 and S100A9, among the top genes being upregulated in primary AML blasts upon stromal contact. As members of the S100 protein family, they can modulate such cellular processes as proliferation, migration, and differentiation. Dysregulation of S100 proteins is described as a predictor of poor survival in different human cancers, including increased S100A8 expression in de novo AML. Thus, we wanted to decipher the underlying pathways of stroma-mediated S100A8/A9 induction, as well as its functional consequences. Upregulation of S100A8/A9 after stromal cross talk was validated in AML cell lines, was contact independent and reversible and resulted in accumulation of S100A8/A9high cells. Accordingly, frequency of S100A8/A9high AML blasts was higher in the patients’ BM than in peripheral blood. The S100A8/A9high AML cell population displayed enhanced utilization of free fatty acids, features of a more mature myeloid phenotype, and increased resilience toward chemotherapeutics and BCL2 inhibition. We identified stromal cell–derived interleukin-6 (IL-6) as the trigger for a Jak/STAT3 signaling-mediated S100A8/A9 induction. Interfering with fatty acid uptake and the IL-6-Jak/STAT3 pathway antagonized formation of S100A8/A9high cells and therapeutic resistance, which could have therapeutic implications as a strategy to interfere with the AML-niche dynamics.

Introduction

Acute myeloid leukemia (AML) is a malignant disorder characterized by clonal expansion of immature myeloblasts that initially accumulate in the bone marrow (BM) and eventually in the peripheral blood (PB).1 The incidence increases with age, with the median age of diagnosis being 67 years.2 Cytogenetic and molecular abnormalities determine disease prognosis. Current therapeutic strategies include chemotherapy (particularly cytarabine in combination with anthracycline); targeted approaches, such as FLT3 or BCL2 inhibitors, epigenetic modifiers, and CD33-directed immunotoxins; and allogeneic
AML blasts

RNA-seq

Figure 1.
hematopoietic stem cell (HSC) transplantation. Despite the improvement in diagnosing and treating AML, the 5-year overall survival rate is only 25% to 30% because of relapse, refractory disease, and disease-related mortality. Therefore, improving treatment outcome remains a high unmet need for most patients with AML.

In line with the “niche” concept, AML blasts closely interact with the stromal component of the BM microenvironment, similar to their nonmalignant counterparts.3 Emerging evidence suggests that BM-derived mesenchymal stromal cells (BM-MSCs) support leukemia cell survival and confer chemoprotection. The underlying mechanisms of stromal-mediated protection of AML blasts is complex. Contact with BM-MSCs promotes proliferation, activates detoxifying ATP-binding cassette transporters, leads to enhanced antioxidative properties, and skews metabolism toward oxidative detoxifying ATP-binding cassette transporters, leads to enhanced antioxidative properties, and skews metabolism toward oxidative phosphorylation.4,5 In this way BM-MSCs protect AML blasts from spontaneous and drug-induced cell death.6 BM-MSC-derived factors that trigger those phenomena include, among others, free fatty acids;7 chemokines, such as CXCL-88 and CXCL-129; mitochondrial transfer10; and the adhesion molecule CD44.11 In fact, cross talk between BM-MSCs and AML blasts is bidirectional, as the fatty acid transporter CD36,17 Toll-like receptor 4 (TLR4),18 S100A8/A9hi population by

Figure 1. BM stromal cells induce S100A8 and S100A9 in AML cells. (A) FACS-sorted primary CD33+CD34+ AML blasts (n = 6) were cultured for 24 hours in the absence (w/o) or presence (w) of a confluent layer of the human BM-derived stroma cell line HS-5 and subsequently analyzed by microarray analysis (left). The heat map (right) shows the top 30 highest differentially expressed genes in the AML blasts cultured in the presence of HS-5 cells. (B) Publicly available data from TCGA (LAML data set) was divided into S100A8hi/S100A9hi and S100A8lo/S100A9lo groups based on the mean expression, and depicted as survival curves using the Mantel-Cox test for calculation of significance. (C) AML cell lines OCI-AML3 and MOLM-13 were cultured for 48 hours in the presence or absence of HS-5 cells (contact; OCI-AML, n = 11; MOLM-13, n = 10) or HS-5 CM (CM; OCI-AML, n = 5; MOLM-13, n = 5); Gene expression of S100A8 and S100A9 was analyzed by qPCR and is depicted as the fold change of treated/untreated cells (untreated set as 1). (D) Frequency of S100A8/S100A9* cells as shown in the representative pseudocolored flow cytometry plot was determined in AML cell lines (OCI-AML, n = 27; MOLM-13, n = 27) for 48 hours in the absence or presence of HS-5 CM. (E) AML cell lines (OCI-AML, n = 13; MOLM-13, n = 11) were cultured for 48 hours in the absence or presence of CM from MSCs of 7 patients with AML (AML-MSC CM, patient ID 11-17; supplemental Table 1) and analyzed for the frequency of the S100A8/S100A9* population by flow cytometry. (F) Log2-transformed normalized counts of the S100A8 and S100A9 gene expression (from the TCGA LAML data set) were correlated by the Spearman test. (G) The S100A8/S100A9* population among matched-pair PB- and BM-derived AML blasts (n = 10; patient ID 1-10; supplemental Table 1) was analyzed by flow cytometry. Data are expressed as the standard error of mean. *P < .05; **P < .01; ***P < .001. FACS, fluorescence-activated cell sorting; qPCR, quantitative real-time polymerase chain reaction.

Methods

Patient samples and cell lines

PB and BM samples from patients with AML were obtained upon receipt of their informed consent and in accordance with the Declaration of Helsinki (Ethics Committee approval 219_14B). Mononuclear cells from the PB or BM were obtained with Ficoll-Paque (GE Healthcare, Chicago, IL). Patient characteristics can be found in supplemental Table 1. MSCs were isolated from iliac crest BM aspirates taken from patients with newly diagnosed AML (Ethics Committee Approval 4777) by density gradient centrifugation and were expanded as previously detailed, while fulfilling uniformly minimal criteria.24

The human BM stroma cell line HS-5 was purchased from ATCC (Manassas, VA). The AML cell lines OCI-AML3 and MOLM-13 were purchased from DSMZ ( Braunschweig, Germany). Mycoplasma contamination was regularly assessed with the Venor GeM Classic Mycoplasma detection kit (Minerva Biologics, Berlin, Germany), according to the manufacturer’s instructions, and authentication of the cell lines was performed.
Figure 2. S100A8 and S100A9 is induced via IL-6/Jak/STAT3 signaling. (A) CM from HS-5 cells (n = 8) was analyzed with a bead-based multiplex assay for 25 human cytokines (IL-9, IL-10, IL-17A, and interferon γ [IFN-γ] not detected). The pie chart displays the individual proportion among the total analytes. (B) Induction of S100A8 and S100A9 on the mRNA (left; OCI, n = 4; MOLM-13, n = 5) and induction of an S100A8/A9hi population (right; OCI-AML, n = 4; MOLM-13, n = 4) were analyzed after treatment with 10 ng/mL of rhIL-6 for 48 hours by qPCR and flow cytometry, respectively. mRNA levels are shown as the fold change with or without rhIL-6 (untreated set as 1). Moreover, the frequency of S100A8/A9hi AML cells in the absence (isotype) or presence of 10 ng/mL anti-IL-6 or anti-IL-6R blocking antibodies. (C) CM from HS-5 cells was prepared after inflammatory priming of the HS-5 cells (with 15 ng/mL tumor necrosis factor-α and 10 ng/mL IFN-γ). Gene expression of IL-6 in HS-5 cells was determined by qPCR and expressed as fold change between nonprimed and primed samples (nonprimed samples set as 1; left, n = 4). The frequency of S100A8/A9hi AML cells (right; OCI-AML, n = 2; MOLM-13, n = 4) cultured in the absence or presence of HS-5 CM for 48 hours was determined by intracellular fluorescence intensity. (D) Gene expression of IL-6 was determined in healthy donor- and AML patient-derived BM-MSCs (both n = 5). (F) Gene expression of PIM1 and SOCS3 (2 STAT3 target genes) was determined in AML cell lines (OCI-AML n = 3, MOLM-13 n = 3) cultured in the absence or presence of HS-5 CM for 48 hours by qPCR and is shown as the fold change (untreated set as 1). (G) Phosphorylation of the STAT proteins (p-STAT) 1, 3, 5, and 6 was determined by PhosFlow in AML cell lines (OCI-AML, n = 3; MOLM-13, n = 3) cultured in the absence or presence of HS-5 CM for 20 minutes and is depicted as a heat map showing the fold change with or without HS-5 CM (left; untreated set as 1). Most prominently, p-STAT3 was induced by HS-5 CM treatment as representatively shown in the histogram (right; scale indicates median fluorescence intensity). (H-J) STATA3 phosphorylation was analyzed by PhosFlow in AML cell lines cultured in the presence of HS-5 CM for 20 minutes after pretreatment for 2 hours with anti-IL-6–blocking antibodies (H; OCI-AML, n = 3; MOLM-13, n = 3), ruxolitinib (I; OCI-AML, n = 2; MOLM-13, n = 2), and Jak1 or Jak2 inhibitor (J; OCI-AML, n = 2; MOLM-13, n = 2). (K-L) Induction of S100A8 and S100A9 gene expression after 48 hours of culture with HS-5 CM was analyzed in the absence (0 μM) or presence (1 μM) of the pan-Jak inhibitor ruxolitinib (K; OCI-AML, n = 3; MOLM-13, n = 3) or the p-STAT3 inhibitor C188-9 (L; OCI-AML, n = 5; MOLM-13, n = 5). Values are depicted as fold change with or without HS-5 CM (untreated set as 1). Data are expressed as the standard error of mean. *P < .05; **P < .01; ***P < .001; ns, not significant. qPCR, quantitative real-time polymerase chain reaction.
Figure 3. S100A8/A9<sup>hi</sup> AML cells display enhanced fatty acid metabolism. (A) Gene expression of the fatty acid translocase CD36 was determined in FACS-sorted S100A8/A9<sup>hi</sup> and S100A8/A9<sup>lo</sup> OCI-AML (n = 3) and MOLM-13 (n = 3) cells cultured for 48 hours in the presence of HS-5 CM by qPCR and is shown as the fold change (S100A8/A9<sup>lo</sup> cells were set as 1). (B) Surface levels of CD36 were determined by flow cytometry on S100A8/A9<sup>hi</sup> and S100A8/A9<sup>lo</sup> cells among the AML cell lines cultured for 48 hours in the presence of CM from either HS-5 cells (left; OCI-AML, n = 4; MOLM-13, n = 4) or AML-MSCs isolated from 6 patients (right; OCI-AML, n = 12; MOLM-13, n = 12, patient ID 11-16; supplemental Table 1). (C) Surface levels of CD36 were analyzed by flow cytometry in matched-pair (n = 10; patient ID 1-10; supplemental Table 1) PB- and BM-derived S100A8/A9<sup>hi</sup> and S100A8/A9<sup>lo</sup> AML blasts, as representatively shown in histograms (left) and summarized in a bar graph (right). (D) Localization of S100A8 and S100A9 was visualized by fluorescence microscopy (left, scale bars = 20 μm). MOLM-13 cells (n = 5) were cultured in the absence (top) or presence (bottom) of HS-5 CM and stained for the cell membrane (WGA, green), S100A8 (left, red) or S100A9 (right, red), and the nucleus (blue, DAPI). Co-localization of S100A8/S100A9 and WGA were calculated with ZEN software (bar graphs; Zeiss). (E) A proximity ligation was performed with a Duolink Flow kit (Sigma-Aldrich) with antibodies against S100A8, S100A9, and CD36. The representative images show the bright-field (left, BF) and the fluorescence signal (right) in MOLM-13 cells. Bars represent 20 μm for 20× magnification.
Cell culture

For coculture experiments, HS-5 cells were seeded 24 hours before the addition of AML cells. Either $5 \times 10^6$ primary, sorted CD45<sup>+</sup>CD33<sup>+</sup>CD34<sup>+</sup> AML blasts were added to $5 \times 10^4$ HS-5 cells, or $2 \times 10^5$ AML cells were added to $1 \times 10^5$ HS-5 cells per well in a final volume of 2 mL in a 24-well plate (Greiner Bioone, Germany). Cultures were maintained in RPMI-1640 (Sigma-Aldrich, St. Louis, MO) supplemented with 10% fetal calf serum (C. C. Pro, Germany), 2 mM L-glutamine (Sigma-Aldrich), and 40 U/mL penicillin-streptomycin (ThermoFisher Scientific, Waltham, MA) at 37°C in 5% CO₂.

Conditioned medium (CM) was produced by culturing either $2 \times 10^6$ HS-5 cells in 20 mL culture medium for 4 days or $5 \times 10^5$ AML-MSCs at passage 3 in 20 mL Dulbecco’s minimal essential medium supplemented with 30% fetal bovine serum and 1% penicillin, streptomycin, and L-glutamine (all from Sigma-Aldrich) for 6 days. Culture supernatant was harvested and made cell free by centrifugation and filtration. AML cells were cultured at a density of $10^3$/mL in fresh culture medium, with or without 50% CM, for 48 hours, unless otherwise stated. In selected experiments CM was depleted from extracellular vesicles by sequential centrifugation at 100 000 g for 18 hours with the supernatant of the last centrifugation step.

All inhibitors, chemotherapeutics, cell culture antibodies, recombinant proteins, and compounds used for in vitro cultures are listed in supplemental Table 2.

Statistical analysis

Outliers were determined with the ROUT test. Differences in means were evaluated with parametric (paired or unpaired t test or 1-way analysis of variance) or nonparametric (unpaired Mann-Whitney, paired Wilcoxon, or unpaired Kruskal-Wallis) tests based on the number of comparisons (2 or $>$2) and distribution levels (as determined by Shapiro-Wilk and Kolmogorov-Smirnov). All statistical analyses were performed with GraphPad Prism, version 7 or 8 (GraphPad Software Inc, San Diego, CA) at a significance level of $P < .05$.

Results

Stromal cells promote S100A8/A9 expression in AML in a cell-to-cell, contact-independent manner

First, we purified primary CD33<sup>+</sup>CD34<sup>+</sup>CD45<sup>+</sup> HSPC blasts by fluorescence-activated cell sorting from the PB of 6 patients with newly diagnosed AML. These cells were either cultured alone or cocultured with HS-5, a human BM-derived MSC line, for 24 hours. Subsequently, AML blasts were repurified (again) based on CD33, CD34, and CD45 coexpression, and microarray analysis was performed. Both S100A8 and S100A9 were among the top 30 significantly upregulated genes upon stromal contact (Figure 1A).

In line with previous studies, our study observed poorer survival in patients with AML with an increased S100A8 or S100A9 mRNA expression when performing Kaplan-Meier survival analyses using data generated by TCGA (The Cancer Genome Atlas) Research Network (www.cancer.gov/tcga) (Figure 1B). Next, we confirmed our findings by coculturing the AML cell lines OCI-AML and MOLM-13 with HS-5 cells in cell-to-cell contact or with HS-5 CM. In fact, cell-to-cell contact was not essential for the upregulation of either S100A8 or S100A9 mRNA and could also be confirmed using primary AML BM-MSC CM (Figure 1C; supplemental Figure 1A). Further analyses revealed the (dose-dependent and reversible) induction of an S100A8/A9<sup>high</sup> population (on the protein and mRNA levels) among OCI-AML and MOLM-13 cells treated with HS-5 or AML-MSC CM (Figure 1D-E; supplemental Figure 1B-D). The simultaneous upregulation of S100A8 and S100A9 is further suggested by the TCGA AML cohort data displaying a strong correlation between S100A8 and S100A9 mRNA expression (Figure 1F). Comparative evaluation of paired AML samples from the PB and BM showed an increased frequency of S100A8/A9<sup>high</sup> AML blasts within the BM, further corroborating the notion that the BM stroma drives S100A8/A9 levels (Figure 1G). Interestingly, primary CD33<sup>+</sup>CD34<sup>+</sup> HSCs did not show such induction of S100A8/A9 gene expression or increased formation of S100A8/A9<sup>high</sup> cells in response to HS-5 CM (supplemental Figure 1E). Because MSCs release bioactive extracellular vesicles (EVs) that shape the BM microenvironment, we depleted EVs in HS-5 CM by ultracentrifugation. However, we did not note any impact of EV depletion on promotion of the S100A8/A9<sup>high</sup> subset by CM, suggesting that alternative soluble factors were the cause (supplemental Figure 1F).

Stroma cell–derived IL-6 promotes the induction of S100A8/A9<sup>high</sup> AML cells via Jak/STAT3 signaling

Next, we used a magnetic bead–based multiplex cytokine assay to analyze the HS-5–derived CM. We identified IL-6 as the most abundant of the 25 cytokines tested (Figure 2A). In fact, previous studies have reported production and secretion of IL-6 by MSCs. Treating the AML cell lines with recombinant human IL-6 (rhIL-6) recapitulated the effects of CM in terms of S100A8/A9 upregulation and enrichment of an S100A8/A9<sup>high</sup> population (Figure 2B). This result is in line with our observation that expression of IL-6 receptor (IL-6R) correlates significantly with S100A8/A9 in the TCGA AML cohort (supplemental Figure 2A). Blocking IL-6 or IL-6R significantly interfered with the CM-mediated induction of S100A8/A9<sup>high</sup> cells (Figure 2C). Inflammatory activation (ie, “priming” of HS-5 cells by interferon-γ and tumor necrosis
Figure 4.

A. HS-5 CM

B. AML MSC CM

C. rIL-6

D. PB

BM

E. S100A8/A9 hi S100A8/A9 lo

F. S100A8/A9 hi S100A8/A9 lo

G. S100A8/A9 hi

Figure 4.
factor-α, but not by contact with AML cells, boosted IL-6 expression significantly further, and CM retrieved from primed HS-5 cells displayed a superior ability in driving an S100A8/A9<sup>hi</sup> phenotype (Figure 2D; supplemental Figure 2B). In fact, IL-6 plays a significant role in AML pathogenesis, and IL-6 expression was higher in MSCs derived from patients with AML compared with healthy control–derived MSCs (Figure 2E).

We identified the STAT3 target genes PID1 and SOCS3 among the top 30 significantly upregulated genes after stromal contact (Figure 1A). We validated the microarray findings by quantitative polymerase chain reaction analysis in AML cell lines treated with CM (Figure 2F). The IL-6/JAK/STAT3 signaling pathway is aberrantly hyperactivated in hematopoietic malignancies and solid tumors. Accordingly, phosphospecific flow cytometry (PhosFlow) staining revealed STAT3 activation and phosphorylation in OCI-AML and MOLM-13 cells in the presence of CM, which was most pronounced in S100A8/A9<sup>hi</sup> cells and which could be significantly antagonized by neutralizing IL-6 (Figure 2G-H; supplemental Figure 2C). Simultaneous inhibition of Jak1 and Jak2 (by ruxolitinib) or Jak1 and Jak2 separately also interfered with CM-triggered STAT3 activation (Figure 2H-J). Finally, treatment with both ruxolitinib and C188-9 (a STAT3 inhibitor) attenuated induction of S100A8/A9 by CM (Figure 2K-L; supplemental Figure 2D-E), further corroborating the notion that stroma–derived IL-6 promotes formation of an S100A8/A9<sup>hi</sup> AML subset in a JAK/STAT3-dependent manner.

**S100A8/A9<sup>hi</sup> AML cells are characterized by high CD36 levels and increased uptake of free fatty acids**

The fatty acid transporter CD36 has been linked to adverse prognosis, leukemic growth, and chemoresistance in AML. The gene and protein expression of CD36 was significantly higher in the S100A8/A9<sup>hi</sup> subset of AML cell lines treated with HS-5 or AML MSC–derived CM or IL-6 (Figure 3A-B; supplemental Figure 3A). This finding was confirmed by ex vivo analysis of primary AML blasts isolated from the PB and BM, showing increased CD36 levels in S100A8/A9<sup>hi</sup> cells and in the TCGA AML cohort exhibiting a strong correlation between CD36 and S100A9 determinates the differentiation state of myeloid cells. Moreover, S100A8/A9 gene expression correlated significantly with the myeloid differentiation marker CD11b and CD14 within the TCGA AML cohort (Figure 4A-B). Accordingly, S100A8/A9<sup>hi</sup> AML cells enriched in the presence of HS-5–CM, AML MSC–CM, and rhIL-6 exhibited a significantly increased expression of the myeloid differentiation markers CD11b and CD14 (Figure 4A-C), both of which have been linked to inferior prognosis and higher treatment resistance. Likewise, primary S100A8/A9<sup>hi</sup> PB- and BM-derived AML blasts expressed higher CD11b and CD14 cell surface levels compared with their S100A8/A9<sup>low</sup> counterparts (Figure 4D). Blocking JAK/STAT signaling interferes with the induction of an S100A8/A9<sup>hi</sup> AML population by stromal cells (Figure 2I-L; supplemental Figure 2D-E), and it also prevents upregulation of myeloid differentiation markers (Figure 4E-F). This finding is in accordance with the role of STAT3 as a mediator of myeloid cell differentiation and survival. Blocking fatty acid uptake (by SSO) affected CD14 but not

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**Figure 4. S100A8/A9<sup>hi</sup> AML cells display signs of myeloid differentiation.** (A-C) AML cell lines were analyzed by flow cytometry for the myeloid maturation markers CD11b and CD14 in S100A8/A9<sup>hi</sup> and S100A8/A9<sup>lo</sup> cells after 48 hours of culture in the presence of HS-5 CM (A; OCI-AML, n = 5; MOLM-13, n = 5), AML–MSC CM from 6 patients (B; OCI-AML, n = 12; MOLM-13, n = 12, patient ID 11-16; supplemental Table 1), and recombinant human IL-6 (C; OCI-AML, n = 3; MOLM-13, n = 3), based on median fluorescence intensity (MdFI). (D) The myeloid maturation markers CD11b and CD14 were analyzed by flow cytometry in matched-pair (n = 10; patient ID 1-10; supplemental Table 1) PB- and BM-derived S100A8/A9<sup>hi</sup> and S100A8/A9<sup>lo</sup> AML blasts, as shown in representative histograms (left) and as summarized by fold change (right; S100A8/A9<sup>lo</sup> set as 1). (E-F) Surface levels of CD11b and CD14 were analyzed in S100A8/A9<sup>hi</sup> and S100A8/A9<sup>lo</sup> (OCI-AML, n = 4; MOLM13, n = 4) populations after culture for 48 hours with HS-5 CM in the absence or presence of the pan-Jak inhibitor ruxolitinib, as indicated. (G) Surface levels of CD11b and CD14 were analyzed in S100A8/A9<sup>hi</sup> cells among the AML cell lines (OCI-AML, n = 5; MOLM-13, n = 5) after 48 hours of culture with HS-5 CM in the absence or presence of the CD36 inhibitor SSO. Data are expressed as the standard error of the mean. *P < 0.05; **P < 0.01; ***P < 0.001.
Figure 5. S100A8/A9 expression is associated with enhanced therapeutic resistance. (A) AML cell lines (OCI-AML, n = 5; MOLM-13, n = 5) were incubated for 48 hours in the absence (−, circles) or presence (+, squares) of HS-5 CM and treated with 50 nM doxorubicin for the last 24 hours of culture. Viability was analyzed by annexin-V/7-AAD staining via flow cytometry, and specific cell death by doxorubicin was calculated. (B-C) Similarly, specific cell death by 50 nM doxorubicin of AML cell lines (OCI-AML, n = 3; MOLM-13, n = 2-5) cultured in the presence of HS-5 CM was determined by flow cytometry after 48 hours of culture in the absence and presence of HS-5 CM and upon treatment with 50 nM doxorubicin for the last 24 hours, as indicated. (E-F) Frequency of S100A8/A9hi OCI-AML (n = 3-4) and MOLM-13 (n = 4) cells after culture for 48 hours in the absence of HS-5 CM and for the last 24 hours in the presence of 50 nM doxorubicin was determined by flow cytometry upon treatment with ruxolitinib (B) or SSO (C). (D) The frequency of S100A8/A9lo OCI-AML (n = 8) and MOLM-13 (n = 11) cells was determined by flow cytometry in untreated (circles) and treated (squares) samples treated with ruxolitinib (B) or SSO (F). (G) Free cytosolic calcium was semiquantified in AML cells (OCI-AML, n = 3; MOLM-13, n = 3) cultured for 48 hours in the absence (medium) or presence (CM) of HS-5 CM with the flow cytometric probe Fluo-8. (H) Specific cell death of AML cells (OCI-AML, n = 3; MOLM-13, n = 2) by 50 nM doxorubicin was estimated via annexin-V/7-AAD staining in untreated samples and upon scavenging extracellular (EGTA) and free cytosolic (BAPTA) calcium. (I) Log2-transformed normalized counts of the S100A8 and S100A9 gene expression (from TCGA LAML data set) correlated with the log2-transformed normalized count of MCL1 by the Spearman test. (J) Cell death of S100A8/A9lo and S100A8/A9hi AML cells (OCI-AML, n = 3; MOLM-13, n = 3) cultured for 24 hours in the presence of HS-5 CM was analyzed upon treatment with increasing concentrations of venetoclax by annexin-V/7-AAD staining in flow cytometry. (K) Frequency of S100A8/A9lo AML cells (OCI-AML, n = 2; MOLM-13, n = 2) cultured for 24 hours in the absence or presence of HS-5 CM was determined by flow cytometry after treatment with increasing concentrations of venetoclax. (L-M) Intracellular protein levels of Mcl-1 (L, OCI-AML, n = 4; MOLM-13, n = 4) and Bcl-2 (M, OCI-AML, n = 3; MOLM-13, n = 3) were measured by flow cytometry in S100A8/A9lo and S100A8/A9hi subsets of AML cells cultured for 48 hours in the presence of HS-5 CM. Data are expressed as the standard error of mean. *P < .05; **P < .01; ***P < .001; ns, not significant.
CD11b expression in S100A8/A9high AML cells, whereas blocking fatty acid oxidation (by etomoxir) had no effect on the expression of either molecule (Figure 4G; supplemental Figure 4C-D).

**Stromal cell–induced S100A8/A9high AML cell population displays increased resistance to chemotherapy and Bcl-2 targeting**

The chemoprotective effect of the stromal niches on AML cells is well established. Furthermore, previous studies have suggested that a high expression of the fatty acid translocase CD36, as seen in S100A8/A9high AML cells, is associated with increased chemoresistance. First, we validated that AML cells were significantly more resistant to doxorubicin-induced cell death when treated in the presence of HS-5 CM (Figure 5A). Induction of S100A8/A9high AML cells by stroma was efficiently antagonized by blocking JAK/STAT signaling or fatty acid uptake (Figure 5E-F). S100A8 and S100A9 molecules both contain calcium (Ca2+) binding motifs and can therefore buffer free cytosolic Ca2+, which in turn is essential for chemotherapies to exert their cytotoxic function. Treating AML cells characterized by an overly active CD36-fatty acid oxidation-metabolic switch toward the preferential utilization of endogenous fatty acids, was counteracted by interfering with JAK/STAT signaling or fatty acid uptake (Figure 5E-F). S100A8 and S100A9 molecules both contain calcium (Ca2+) binding motifs and can therefore buffer free cytosolic Ca2+, which in turn is essential for chemotherapies to exert their cytotoxic function.12 Treating AML cell lines with HS-5 CM reduced their intracellular Ca2+ levels, whereas chemically depleting intracellular (not extracellular) free Ca2+ similarly led to a reduced sensitivity toward doxorubicin (Figure 5G-H), which coincided with a stabilized mitochondrial membrane potential (and an according elevation of reactive oxygen species) reflecting a reduced Ca2+ crisis (supplemental Figure 5A). In addition, a recent study demonstrated that elevated S100A8/A9 expression in AML correlates with resistance to Bcl-2 inhibition. However, increased Ca2+ binding by S100A8/A9 was most likely insufficient to cause venetoclax resistance. Mcl-1 overexpression can be found in venetoclax-resistant lymphoma and leukemia cell lines and gene expression of the antiapoptotic Mcl-1 correlated positively with S100A8 and S100A9 in the TCGA AML cohort (Figure 5), whereas Bcl-2 expression showed a negative correlation (supplemental Figure 5B). Next, we validated the earlier report by documenting an increased resilience of S100A8/A9high AML cells in response to venetoclax (Figure 5J). Similar to our observations when treating AML cells with doxorubicin, we found an enrichment of S100A8/A9high cells during treatment with venetoclax, particularly in the presence of HS-5 CM, which at the same time harbored higher Mcl-1 and Bcl-2A1 levels (Figure 5K-L; supplemental Figure 5C-D), as well as lower levels of Bcl-2 (Figure 5M).

**Discussion**

Therapeutic resistance of AML represents one of the main hurdles in clinical practice. BM stroma promotes survival and therapeutic resistance of AML cells. To date, several underlying mechanisms have been described and include, among others, CXCL-12–driven metabolic reprogramming, upregulation of antiapoptotic proteins, and mitochondrial transfer. Obviously, a better understanding of those processes is a prerequisite for developing novel treatment approaches.

In our present study, BM stroma cells (including AML patient-derived MSCs) promoted (the reversible) induction of the Ca2+ binding proteins S100A8 and S100A9, leading to formation of an S100A8/S100A9high subpopulation among AML cells. Consequently, S100A8/S100A9high blasts are found at higher frequencies within the BM niche, compared with the circulation. S100A8/A9 are expressed in myeloid but not lymphoid cells. They represent alarmins and are elevated in inflammatory conditions or malignant entities that can mimic inflammation. In fact, S100A8 and S100A9 overexpression has been reported in genomic cohort analyses of patients with AML and may be caused by a nonspecific inflammatory reaction. Furthermore, MSCs, and in particular MSCs, retrieved from patients with AML display enhanced IL-6 production. Signaling through the IL-6R supports proliferation of AML blasts, and using IL-6 blockade has led to prolongation of survival in preclinical AML models. Downstream of IL-6, we detected STAT3 activation, which was pharmacologically targetable with different Jak and/or STAT inhibitors and which prevented S100A8/A9 upregulation. In fact, STAT3 binding sites have been located by chromatin immunoprecipitation in the promoter region of S100A8 and S100A9. Previous studies have identified aberrant STAT3 signaling, which is partly ligand dependent, as an important element in AML cell survival and potentially, chemoresistance. However, as inhibition of the IL-6 signaling by blocking antibodies did not have the same magnitude of effect as Jak/STAT inhibition, it is likely that additional stroma cell-derived factors are involved in the regulation of S100A8 and S100A9 expression in AML. S100A8/A9 molecules can interact with the fatty acid transporter CD36, thereby enhancing fatty acid uptake. In AML, CD36 promotes growth, chemoresistance, and persistence of leukemic precursors and is linked to an adverse outcome. Expression of CD36 and colocalization of CD36 with S100A8/A9 was increased upon stromal cross talk. Consequently, we noted a metabolic switch toward the preferential utilization of endogenous and exogenous fatty acids for fueling mitochondrial respiration. This metabolic phenotype resembles chemoresistant AML blasts that are characterized by an overly active CD36-fatty acid oxidation-phosphorylation axis. Therefore, blocking CD36 with SSO, for example, has attracted considerable attention as a strategy in AML. Formation of S100A8/A9high cells was significantly reduced by SSO, which requires further investigation, as S100A8/A9 heterodimers can bind fatty acids. However, whether S100A8 and S100A9 have a direct effect on fatty acid metabolism remains to be elucidated. In fact, our data indicate that S100A8/A9 induction depends on fatty acids and enhances their usage indirectly via increased CD36 activity.

In terms of myeloid differentiation, S100A8 and S100A9 seem to play contradictory roles. In vitro experiments and preclinical models have revealed that S100A9 promotes differentiation via TLR4, whereas S100A8 blocks differentiation. It has been suggested that the relative ratio between S100A9 and S100A8 finally determines their impact on AML maturation. In this study, S100A8/A9high
AML cells displayed a more mature phenotype, suggesting a dominant role of S100A9. Differentiation, together with S100A8/A9 induction, was triggered by the IL-6-STAT3 pathway. The biological impact of this observation remains unclear, and it could be speculated that pseudomaturation could yield less proliferative, quiescent-like AML cells that are less susceptible to chemotherapeutics that preferentially affect the highly proliferative cell fraction.

As previously published by us and others, stromal cells promote chemoresistance of AML cells, and this effect is successfully antagonized by Jak/STAT and/or CD36 blockade. Furthermore, upon treatment with doxorubicin, we documented an enrichment of S100A8/A9-high cells among the surviving cell fraction, in line with our aforementioned findings (eg, the metabolic phenotype of S100A8/A9-high cells) that imply a superior chemoresistance. Survival of S100A8/A9-high cells was diminished when we combined doxorubicin with Jak/STAT or CD36 inhibition. High levels of S100A8/A9 may exert directly antiapoptotic effects by binding free cytosolic Ca²⁺ (most likely released from the endoplasmic reticulum), which is involved in cell death by apoptosis. In addition to conventional chemotherapeutics, S100A8/A9-high cells were also less susceptible to the Bcl-2 inhibitor venetoclax. Similarly, this finding may be related to less-available cytosolic Ca²⁺ migrating toward the mitochondria to elicit apoptosis. However, we also found a positive correlation between Mcl-1, which, as an antiapoptotic protein, represents one of the main contributors to venetoclax resistance in AML and S100A8/A9 expression. The AML-specific relationship between Mcl-1 and S100A8/A9 requires further investigation, as S100A8/A9 and Mcl-1 do not correlate in acute lymphoblastic leukemia. Based on previous data, STAT3 signaling may represent an integrating hub for both S100A8/A9 and Mcl-1.

In summary, we demonstrated that BM-resident stromal cells promote a therapy-resistant S100A8/A9-high phenotype among AML cells in a contact-independent fashion. Mechanistically, we showed that the induction of S100A8/A9 is mediated by an IL-6/Jak/STAT3 axis and depends on exogenous fatty acids. Therapy resistance is at least partly mediated by the Ca²⁺-binding nature of S100A8/A9 and their positive correlation with antiapoptotic proteins resulting in diminished apoptosis. Thus, our data append to the complex network of bidirectional cross talk between leukemic blasts and their stromal microenvironment and suggest that targeting induction of S100A8/A9 in AML by different means may represent a promising way to improve treatment efficacy by interfering with BM-derived nurturing signals.

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