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IMiDs mobilize acute myeloid leukemia blasts to peripheral blood through downregulation of CXCR4 but fail to potentiate AraC/Idarubicin activity in preclinical models of non-del5q/5q-AML

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ABSTRACT
Treatment for acute myeloid leukemia (AML) remains suboptimal and many patients remain refractory or relapse upon standard chemotherapy based on nucleoside analogs plus anthracyclines. The crosstalk between AML cells and the BM stroma is a major mechanism underlying therapy resistance in AML. Lenalidomide and pomalidomide, a new generation immunomodulatory drugs (IMiDs), possess pleiotropic anti-leukemic properties including potent immune-modulating effects and are commonly used in hematological malignancies associated with intrinsic dysfunctional BM such as myelodysplastic syndromes and multiple myeloma. Whether IMiDs may improve the efficacy of current standard treatment in AML remains understudied. Here, we have exploited in vitro and in vivo preclinical AML models to analyze whether IMiDs potentiate the efficacy of AraC/Idarubicin-based standard AML chemotherapy by interfering with the BM stroma-mediated chemoresistance. We report that IMiDs do not exert cytotoxic effects on either non-del5q/5q- AML cells nor BM-MSCs, but they enhance the immunomodulatory properties of BM-MSCs. When combined with AraC/Idarubicin, IMiDs fail to circumvent BM stroma-mediated resistance of non-del5q/5q- AML cells in vitro and in vivo but induce robust extramedullary mobilization of AML cells. When administered as a single agent, lenalidomide specifically mobilizes non-del5q/5q- AML cells, but not healthy CD34+ cells, to peripheral blood (PB) through specific downregulation of CXCR4 in AML blasts. Global gene expression profiling supports a migratory/mobilization gene signature in lenalidomide-treated non-del5q/5q- AML blasts but not in CD34+ cells. Collectively, IMiDs mobilize non-del5q/5q- AML blasts to PB through CXCR4 downregulation, but fail to potentiate AraC/Idarubicin activity in preclinical models of non-del5q/5q- AML.

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AML; BM-MSC; IMiDs; lenalidomide; pomalidomide; AraC; Idarubicin; xenografts

Introduction
Acute myeloid leukemia (AML) comprises a biologically and genetically heterogeneous group of disorders characterized by the rapid expansion of immature myeloid blasts in bone marrow (BM). 1,2 In contrast to acute lymphoblastic leukemia (ALL), treatment of AML has not improved substantially over the last two decades and many patients still fail to respond to standard intensive chemotherapy based on nucleoside analogs such as cytarabine (AraC) or fludarabine and anthracyclines such as idarubicin and daunorubicin. 3,4 Failure of current therapies to eradicate leukemia-initiating/propagating cells (LICs) and chemotherapy refractoriness are the major mechanisms underlying AML progression/relapse. In addition, the BM stroma has been involved in the pathogenesis of a variety of hematologic malignancies including AML. 3 The interaction of AML cells with the BM microenvironment in functional niches is a major mechanism underlying leukemia maintenance and therapy resistance. 5,6 Thus, the high rate of mortality/morbidity and chemo-resistance in AML guides the search for new compounds with higher efficiency and lower toxicity.

New generation of immunomodulatory drugs (IMiDs) such as lenalidomide and pomalidomide are thalidomide analogs, which possess pleiotropic anti-leukemic properties...
including anti-proliferative, anti-angiogenic but also immune-modulating effects.\textsuperscript{7-9} They are approved by the Food and Drug Administration (FDA) and the European Medicines Agency (EMA). Lenalidomide is approved for both 5q- myelodysplastic syndrome (MDS) and multiple myeloma (MM) whereas pomalidomide is approved for MM, which are hematological malignancies associated with intrinsic dysfunctional BM.\textsuperscript{10,11} Some studies reported that lenalidomide has modest (pre)-clinical activity in lymphoma and AML with manageable toxicity.\textsuperscript{9,12,13} Ongoing Phase I clinical trials assaying low- and high-dose lenalidomide for both newly diagnosed and relapsed/refractory AML also support that lenalidomide displays modest activity when given either alone or combined with cytarabine and anthracyclines.\textsuperscript{14}

Whether and how IMiDs may benefit current standard treatment in AML needs further investigation. To bring IMiDs to the forefront of AML therapeutic arsenal, a better understanding of its mechanism of action in AML is required. In fact, it has recently been shown that BM mesenchymal stromal cells (BM-MSCs) derived from diagnostic AML patients are potent immunomodulators and provide similar chemoprotection of AML cells to cytarabine/idarubicin than normal BM-MSCs.\textsuperscript{5} Although IMiDs may have an anti-proliferative effect on leukemic cells, they may likely exert pleiotropic and immunomodulatory effects at the stroma-leukemic cell crosstalk, thus improving standard AML intensive chemotherapy by either potentiating AraC/anthracycline efficacy or reducing BM-MSC-derived chemoresistance.

Here, we have exploited in vitro and in vivo AML preclinical models to analyze whether IMiDs potentiate the efficacy of AraC/Idarubicin-based standard chemotherapy by lowering the BM stroma-mediated chemoresistance. Lenalidomide and pomalidomide are not cytotoxic for either non-del5q/5q- AML cells or BM-MSCs, but they enhance the immunomodulatory capacity of BM-MSCs. When administered alone or in combination with AraC and Idarubicin, IMiDs fail to circumvent BM stroma-mediated resistance of non-del5q/5q- AML cells in vitro and in vivo but induce robust mobilization of non-del5q/5q- AML cells, but not healthy CD34\(^+\) cells, to peripheral blood (PB) and spleen likely through specific downregulation of CXCR4 in AML blasts. Global gene expression profiling supports a migratory/mobilization gene signature in lenalidomide-treated AML primary blasts. We conclude that IMiDs mobilize non-del5q/5q-AML blasts to PB through downregulation of CXCR4 but do not improve AraC/Idarubicin activity in a preclinical model of AML.

### Table 1. Biological and cytogenetic-molecular characteristics of blasts from diagnostic AML patients.

<table>
<thead>
<tr>
<th>Patient ID</th>
<th>Diagnostic</th>
<th>Cytogenetics</th>
<th>Molecular</th>
<th>Age (years)</th>
<th>Gender</th>
<th>Blasts (%)</th>
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<tr>
<td>AML#1</td>
<td>AML-M4</td>
<td>46, XX</td>
<td>FL3T-ITD, NPM1(^{MUT})</td>
<td>52</td>
<td>F</td>
<td>81</td>
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<td>46, XX, 20% 47, XX, +8 t(8;21)</td>
<td>AML1-ETO</td>
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<td>F</td>
<td>52</td>
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<td>AML-M0/XY</td>
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<td>86</td>
</tr>
<tr>
<td></td>
<td>M1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AML#4</td>
<td>AML</td>
<td>46, XY, del(7)(q22)</td>
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<td>61</td>
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<td>87</td>
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<td>80</td>
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<td>AML</td>
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<td>BCR-ABL (Ph(^+))</td>
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<td>M</td>
<td>37</td>
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<td>-</td>
<td>43</td>
<td>M</td>
<td>60</td>
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<tr>
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<td>AML-M5</td>
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<td>-</td>
<td>47</td>
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<td>95</td>
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<td>44, XX, del(5)(q13q33), -7, del(13)(q12q14), -15[20]</td>
<td>-</td>
<td>87</td>
<td>F</td>
<td>40</td>
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<td>-</td>
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<td>71</td>
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<td>-</td>
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<td>F</td>
<td>39</td>
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<td>AML-M4</td>
<td>46, XY, del(3)(q21)[26]/46, idem, del(5)(q22)[3]/46, XY[11]</td>
<td>FL3T-ITD, WT-1(^+)</td>
<td>16</td>
<td>M</td>
<td>75</td>
</tr>
</tbody>
</table>

M, male; F, female; dmin, double minute chromosome; -, no mutations found for FLT3, NPM1, cEBP\(^\alpha\), WT and IDH1.
Drugs
AraC (Cytarabine*, Pfizer), idarubicin (Zavedos*, Pfizer) and the CXCR4 antagonist AMD3100 (Sigma) were reconstituted with PBS. Lenalidomide and pomalidomide were kindly provided by Celgene Corp (San Diego, CA), reconstituted in DMSO as per supplier’s guidelines and stored in aliquots at −20°C.

Cytotoxicity and apoptosis assays
The cytotoxic effect of the different drugs on HL-60 and MOLM-13 cell was assessed using Cell Counting Kit-8 based on monosodium salt WST-8 according to the manufacturer’s protocol (Sigma-Aldrich). Briefly, 5,000 cells were plated in 96-well plate and incubated with increasing concentrations of the corresponding drug for 48h. After incubation time, absorbance was measured at 450 nm using a microplate reader. The cytotoxic effect of IMiDs on BM-MSCs was assessed using MTT assays as previously described.\textsuperscript{21,22} Briefly, 2 × 10^5 cells were cultured with increasing drug concentrations in 96-well plates for 48 h. To determine the chemoprotective effect of BM-MSCs a total of 2 × 10^6 BM-MSCs were plated in 96-well plates 18 h before addition of AML cells (2 × 10^4 for cell lines and 2 × 10^5 for primary cells). BM-MSC:AML co-cultures were treated with IC\textsubscript{50} concentrations of the AraC (77nM) and Idarubicin (7nM) and 10 µM of lenalidomide/pomalidomide for 48–72 h. Apoptosis of CD33\textsuperscript{+} AML cells was measured using the annexin-V/7-AAD apoptosis detection kit (BD Biosciences) on a FACScanto-II cytometer using FACS-quantified using a FACS Canto II cytometer. The effect of IMiDs on the BM-MSC-mediated immunosuppression and cytokine release (TNF\textalpha, IL1\beta, IL10, IL6 and SDF1) was determined by CSFE and Luminex Multiplex assays as previously described.\textsuperscript{5,24,25}

Immunosuppressive assays and cytokine detection
The effect of IMiDs on the BM-MSC-mediated immunosuppressive properties and cytokine release (TNF\textalpha, IL1\beta, IL10, IL6 and SDF1) was determined by CSFE and Luminex Multiplex assays as previously described.\textsuperscript{5,24,25}

FACS analysis of CXCR4 and SDF1
CXCR4 expression was determined in AML and CD34\textsuperscript{+} cells treated with IMiDs. A total of 0.2 × 10^6 (for cell lines) or 1 × 10^6 (for primary cells) were incubated with 10 µM of IMiDs (the CXCR4 antagonist AMD3100 was used as a positive control). Seven days later (48 h–72 h for primary AML cells) treated cells were stained with anti-human CD45-APC, Cy7, CD33-APC, CD34-PECy7 and CD184-BV421 (BD Biosciences) and the mean fluorescence intensity (MFI) of CD184 within the blast population (CD45\textsuperscript{+}CD33\textsuperscript{-}) was FACS-quantified using a FACS Canto II cytometer. The expression of the CXCR4 ligand SDF1 was analyzed in BM-MSCs (0.5 × 10^6) 48 h after treatment with 10 µM IMiDs. BM-MSCs were trypsinized and stained with anti-human CXCL12/SDF-1 (R&D Systems) using the Cell Permeabilization Kit (FIX&PERM\textsuperscript{+}).\textsuperscript{26}

NSG mice xenotransplantation and analysis of engraftment
All experimental procedures were approved by the Animal Care Committee of The Barcelona Biomedical Research Park (HRH-16–0037). Eight-to-14-week-old NOD.Cg-PrkdcscidIl2rgtm1Wjl/SzJ (NSG) mice housed under pathogen free conditions were used. In order to recapitulate a BM stroma milieu, 1 × 10^5 HL60 (n = 31 mice) or MOLM-13 (n = 33 mice) cells were intra-BM transplanted into non-irradiated NSG mice together with 3 × 10^5 irradiated BM-MSCs.\textsuperscript{25,27} HL60 AML grafts were monitored by BM aspiration and mice were randomized into treatment groups when graft was >1% in the contralateral BM (~2–3 weeks post-transplant). For MOLM-13 AML grafts, mice were randomized for treatment 3 days after intra-BM transplant due to its aggressiveness. At the end of the treatment period, mice were sacrificed and cells from the BM, spleen and PB were stained with anti-human HLA-ABC-FITC, CD33-PE and CD45-APC (BD Biosciences) to analyze human leukemic engraftment by flow cytometry. Similarly, healthy CD34\textsuperscript{+} HSPCs (0.1 × 10^6 cells) were intra-BM-transplanted into sub-lethally irradiated (2.25 Gy) NSG mice (n = 10) together with 3 × 10^5 irradiated BM-MSCs. Six weeks later, when human healthy chimerism was evident, mice were treated with lenalidomide. At the end of lenalidomide treatment, mice were sacrificed and cells from the BM, spleen and PB were stained as above to analyze human normal chimerism by flow cytometry. Multilineage engraftment was assessed using CD33-APC, CD19-PE and CD34-PE.Cy7 (BD Biosciences). Spleen infiltration of AML cells was analyzed by immunohistochemistry using an anti-human-CD45.

In vivo chemotherapy and imids treatment
Ara-C and idarubicin were administrated following a well-established 5 + 3 treatment schedule, consisting on intravenous (i.v.) co-delivery of 30 mg/Kg of Ara-C and 0.1 mg/Kg of idarubicin for 3 days followed by intraperitoneal injection (i.p.) of Ara-C for two further days (A + I treatment).\textsuperscript{28} Lenalidomide (25 mg/kg) given as single agent was administrated i.p. for 10 days. When combined with A + I treatment, lenalidomide (25 mg/kg) was administrated alone during days 1–3 and 9–10 and with A + I on days 4–8. 1% DMSO/PBS solution was always used as vehicle treatment. Mice were weighted throughout treatment and drug doses were recalculated as necessary.

Microarray gene expression profiling (GEP)
Non-del5q/Sq- AML primary samples (n = 3) and BM-CD34\textsuperscript{+} HSPCs (n = 2) were FACs purified (CD45\textsuperscript{+}CD33\textsuperscript{-} purity >98%) and cultured with 10 µM lenalidomide for 48h before global gene expression profiling (GEP) as described.\textsuperscript{29} Total RNA was then extracted using a Maxwell\textsuperscript{R} RSC simplyRNA Cells Kit in a Maxwell\textsuperscript{R} RSC Instrument (Promega) and its quality checked in the Agilent 2100 Bioanalyzer. Total RNA samples were labeled with Cy3 using the Quick-Am Labeling Kit and hybridized with the Gene Expression Hybridization Kit to the GeneChip Human Gene 2.0 ST (Affymetrix) following Manufacturer’s instructions.
Hierarchical clustering of genes was performed with the one minus correlation metric and the unweighted average distance. Only genes showing >1.5-fold change expression and p-value<0.05 were considered differentially expressed and were subjected to gene ontology (GO) term analysis using Gorilla\textsuperscript{30,31} publicly available at http://cbl-gorilla.cs.technion.ac.il. Microarray data were deposited in the public Gene Expression Omnibus database, accession number GSE106748.

**Results**

**BM stroma/milieu confers resistance of AML cells to AraC/idarubicin**

Chemotherapy based on the combination of Ara-C with anthracyclines, either idarubicin or daunorubicin, is the first-line induction treatment in diagnostic AML.\textsuperscript{3,4} Here, viability in vitro assays revealed that both cytotoxic drugs, Ara-C and idarubicin, are equally toxic in different AML cell lines with an IC\textsubscript{50} of \(~10^{-7}\)M (Fig 1S). Exposure to Ara-C plus idarubicin (Ara-C/Ida) resulted in 80\%–95\% cell death of AML cell lines, measured by Annexin V staining (Figure 1A). However, when co-cultured with BM-MSCs, AML cell lines were chemo-resistant with only 40–50\% cell death (Figure 1A). To further test whether the BM stroma/milieu protects AML cells from Ara-C/Ida treatment, AML cell lines were co-transplanted via intra BM transplant (IBMT) into NSG mice with human BM-MSCs which largely increase AML engraftment (Fig 2S), as reported in several hematologic malignancies.\textsuperscript{32-35} When engraftment was >1\% in non-injected tibia, mice were treated with Ara-C/Ida administrated following the well-established 5 + 3 treatment schedule.\textsuperscript{28} Similar to the in vitro results, Ara-C/Ida significantly (2-3-fold) reduced AML engraftment/survival in

![Figure 1](image.png)

*Figure 1. BM microenvironment protects HL60 and MOLM-13 cells from AraC+Idarubicin-based chemotherapy in vitro (A) and in vivo (B). (n = 3). *p < 0.05. **p < 0.01.*
extramedular BM hematopoietic tissues and in contralateral BM (CL) whereas the co-transplanted human BM-MSCs conferred chemoresistant to AML cells in injected tibia (IT) (Figure 1B).

Imids exert no toxicity on non-del5q/5q- AML cells but increase immunosuppressive properties of BM-MSCs

The immune-modulating properties of BM stroma/BM-MSCs are a major mechanism underlying BM stroma-mediated chemoresistance in hematological malignances. Lenalidomide and pomalidomide possess pleiotropic anti-tumor properties including also tumor-controlling immune-modulating effects in other hematological BM malignancies such as MDS and MM. We therefore set out to assess the anti-leukemia effects of lenalidomide and pomalidomide in AML. Strikingly, as single agents, lenalidomide and pomalidomide display no cytotoxic effects (assayed at 48h and 72h after drug exposure) on neither AML cell lines (Figure 2A) nor diagnostic primary AMLs (n = 9) covering different cytogenetic/molecular AML subtypes other than del5q/5q- AMLs (Figure 2A, Table 1). Importantly, despite lenalidomide/pomalidomide had neither cytotoxic effects on BM-MSCs (Figure 2A) they did enhance the immunosuppressive capacity of BM-MSCs (Figure 2B). In fact, IMiDs boosted the BM-MSC-mediated inhibition of activated T cells (Figure 2B) through blocking T-cell release of pro-inflammatory cytokines such as TNFα and IL1β while enhancing the production of the master anti-inflammatory cytokine IL10 (Figure 2B). Lenalidomide/pomalidomide also inhibited the release of the pro-inflammatory cytokine IL6 by BM-MSC (Figure 2C). The potent immune-modulatory properties of IMiDs on BM-MSCs led us to combine IMiDs with Ara-C/Ida-based standard AML treatment as a potential therapeutic approach to bypass the BM-MSC-mediated chemoresistance to Ara-C/Ida.

Imids fail to overcome BM-MSC/stroma-mediated chemoresistance of non-del5q/5q- AML cells but induce specific mobilization of AML cells to PB

Neither lenalidomide nor pomalidomide were unable to overcome the BM-MSC-mediated resistance of non-del5q/5q- AML cells to Ara-C/Ida chemotherapy either in vitro (Figure 3A) or in vivo (Figure 3B). Surprisingly, when lenalidomide was combined with chemotherapy we observed a robust (3–4-fold) mobilization of non-del5q/5q- AML cells into extramedular BM hematopoietic sites including PB and spleen (Figure 3B). Next, we performed similar in vivo experiments by treating AML-engrafted NSG mice with lenalidomide as single agent and confirmed that it induces a robust mobilization of non-del5q/5q- AML cells into PB (Figure 4A) and spleen (Figure 4B).

Specific mobilization of leukemic cells from BM to PB is commonly desirable in the clinical setting because circulating blasts are more sensitive to treatment in part due to their detachment from the chemoprotective BM niche. However, mobilization of normal hematopoietic stem/progenitor cells (HSPCs) to PB is not desired because it would accelerate BM aplasia and subsequent hematopoietic defects. Thus, we next analyzed whether lenalidomide also mobilizes normal HSPCs to PB. CB CD34+ cells were IBM-transplanted in NSG mice and total (CD45+HLA-ABC+) and immature (CD45+HLA-ABC-CD34+) engraftment was analyzed 6–7 weeks later. Interestingly, lenalidomide failed to mobilize normal HSPCs indicating that lenalidomide-driven mobilization into PB is specific for AML cells (Figure 4C).

Lenalidomide-mediated mobilization of AML cells to PB is associated to downregulation of CXCR4 and a migratory transcriptomic signature in AML cells

We next sought to gain insights into the mechanisms underlying lenalidomide-mediated mobilization of AML cells. Clearly, the CXCR4/SDF1 axis plays a major role in the interaction of HSPCs and leukemic cells within BM niches. CXCR4 is highly expressed in HSPCs/leukemic cells whereas SDF1 is a cytokine commonly released by BM-MSCs (Figures 5A and 5B). Consequently, we first treated both AML cell lines and primary non-del5q/5q- AML blasts with lenalidomide and pomalidomide and found a significant (25%–50%) reduction in the levels of CXCR4 in AML cells similar to treatment with the specific CXCR4 inhibitor AMD3100 (Figure 5B). In line with the IMiDs-mediated specific mobilization of AML blasts (Figure 4), we confirmed that lenalidomide/pomalidomide do not down-regulate CXCR4 expression in healthy PB- or BM-derived HSPCs (Figure 5B). In contrast, lenalidomide and pomalidomide treatment did not impact the either expression levels or cytokine release of SDF1 by human BM-MSCs (Figure 5C). Together, these results suggest that IMiDs specifically mediate the mobilization of AML cells to PB through down-regulation of CXCR4.

To identify patterns of gene expression that might provide a molecular explanation for the IMiD-induced mobilization of AML cells, we performed whole-genome GEP in lenalidomide-treated versus DMSO-treated AML blasts (blast purity>95%, n = 3 de novo non-del5q/5q- AMLs). A heatmap representation of hierarchical clustering of genes differentially expressed (1.5-fold regulation, p-value<0.05) between lenalidomide- and DMSO-treated AML cells was performed (Figure 6). A total of 323 genes were differentially expressed between lenalidomide- and DMSO-treated AML cells. Of these, 196 (61%) were upregulated and 127 (39%) downregulated in lenalidomide-treated AML cells, establishing a lenalidomide-specific transcriptomic signature. To get insights into the biological functions affected by differentially expressed genes, we used the Gorilla software and found that many significant biological processes predicted to be activated in the lenalidomide-treated AML cells were associated with "migration/motility/mobilization/chemotaxis" (Figure 6A). In contrast, exposure of CB-CD34+ HSPCs to lenalidomide did not affect such biological functions (Figure 6B). This lenalidomide-induced transcriptomic signature supports a CXCR4-mediated mechanism underlying the mobilization of non-del5q/5q- AML cells to PB in response to IMiDs.
Figure 2. Effects of IMiDs on primary AML blasts and BM-MSC. (A) Lenalidomide and pomalidomide do not have a cytotoxic effect on either AML cell lines (n = 3) (top panel), primary AML blasts (n = 9, bottom left panels) or BM-MSC (n = 3) (right bottom panels). (B) Left panel: IMiDs potentiate the immunosuppressive effect of BM-MSCs, measured as percentage of CFSE+ non-proliferating cells. CSFE-labeled PBMCs were stimulated with PHA and then co-cultured with BM-MSCs in the presence of lenalidomide or pomalidomide for 5 days (n = 3). Middle panels: representative flow-cytometry histograms of cycling (CFSElow) PBMCs. Right panels: Concentration of the indicated cytokines in cell-culture supernatants determined by LumineX Multiplex assays. PBMCs were co-cultured with BM-MSCs in the presence/absence of IMiDs. Error bars indicate the SEM values of the n = 3 biological replicates. (C) IMiDs diminish IL6 production by BM-MSC (n = 3). *p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001.
Figure 3. IMiDs combined with AraC-Idarubicin do not circumvent BM- microenvironment-mediated resistance of AML cells but induce robust extramedullary mobilization of AML cells. (A) Lenalidomide and pomalidomide do not overcome BM-MSC-mediated resistance of AML cell lines to intensive chemotherapy (n = 3). (B) IMiD combination with AraC+Idarubicin highly mobilize AML blasts to PB and spleen in AML xenograft models (n = 3). *p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001.
Lenalidomide overcomes BM-MSC/stroma-mediated chemoresistance of AML cells from del5q/5q- patients

A phase II clinical trial has recently reported promising results for lenalidomide combined with AraC/daunorubicine in both MDS and AML patients with 5q alterations (del5q/5q-). Thus, we next focused our efforts on this cytogenetic subgroup of AML patients as potential candidate patients benefiting from the combination of IMiDs with AraC/idarubicine. Primary cells from del5q/5q- AML patients were treated with lenalidomide, AraC/Ida or their combination for 48-72h, in the presence/absence of BM-MSCs (Figure 7A). Opposite to non-del5q/5q- AML cells (Figure 2A), lenalidomide used as single agent displayed a slight, but consistent, 20% toxicity in del5q/5q- AML cells (Figure 7B). More
importantly, lenalidomide significantly boosted the toxicity of del5q/5q- AML cells when combined with AraC/Ida, overcoming the BM-MSC/stroma-mediated chemoresistance of AML cells from del5q/5q- patients (Figure 7C). Interestingly, lenalidomide did not down-regulate CXCR4 expression on del5q/5q- AML cells (Figure 7D), indicating that...
Figure 6. Global GEP reveals a migratory signature in lenalidomide-treated AML blasts. (A, B) Left panels: Heatmap representation of hierarchical clustering of genes differentially expressed between lenalidomide-treated and untreated AML blasts (n = 3 leukemias) (top panel) and BM-CD34+ cells (n = 2 healthy donors) (bottom panel). Right panel: Statistically significant biological functions identified using IPA on genes differentially expressed in lenalidomide-treated versus untreated AML blasts (top panel) and CD34+ cells (bottom panel). They are ranked by z-score. A z-score >2 indicates a predicted activation of that biological function. Biological functions associated with ‘cell migration/movement/motility’ are shown in black.
lenalidomide partially overcomes BM-MSC/stroma-mediated chemoresistance of AML cells from del5q/5q- patients likely at the expense of blast mobilization.

**Discussion**

AML represents a clinically and genetically very heterogeneous group of diseases characterized by an accumulation of proliferative, abnormally differentiated hematopoietic cells in the BM and other tissues, leading to interference of normal hematopoiesis and BM failure. AML is one of the most common hematopoietic malignancies with ~25,000 new cases per year in United States and Europe. Overall, AML has an unfavorable clinical outcome, largely determined by age, cytogenetic/genetic lesions as well as patient's initial response to standard treatment. The large genetic/cytogenetic and phenotypic heterogeneity together with a median age at diagnosis of ~65–67 years old have long represented a limitation towards the development of better treatments. Despite many clinical trials have been conducted over the last two decades, the standard current treatment in AML remains unchanged, and relies on intensive induction chemotherapy based on nucleoside analogs such as Ara-C or fludarabine and anthracyclines such as idarubicin and daunorubicin followed by high doses Ara-C consolidation. Although complete remission rates (CRR) of ~80% are commonly achieved with conventional chemotherapy, the median survival time is short and the outcome of relapsed patients is dismal. Novel agents or better treatment combinations/schedules with higher efficacy and less toxicity are therefore in high demand for de novo and relapsed AML.

Failure of current therapies to eradicate LICs and chemotherapy refractoriness are the major mechanisms underlying AML progression/relapse, being in part because the BM provides a protective microenvironment to support the survival of LICs/bulk leukemic cells, hence promoting drug resistance and influencing clinical outcome. In AML, the high rate of relapse may in part be a result of the inability of current treatments to effectively overcome the protective influence of the BM niche. In fact, it has been recently shown that BM mesenchymal stromal cells (BM-MSCs) derived from diagnostic AML patients are strong immunomodulators and provide similar or even higher chemoprotection of AML cells to cytarabine/idarubicin than normal BM-MSCs. Therefore, new compounds or improved combinatory regimens targeting the interaction between the BM microenvironment and AML leukemic cells are very attractive candidates.

The new generation of immunomodulatory drugs (IMiDs) such as lenalidomide and pomalidomide are thalidomide analogs, which possess pleiotropic anti-leukemic properties including anti-proliferative, anti-angiogenic but also immune-modulating effects. They are approved by the FDA and EMA for 5q- MDS and/or MM which are hematological malignances associated with intrinsic dysfunctional BM with remarkable clinical responses. However, whether IMiDs may potentiate current standard treatment in AML remains an issue of debate. To bring IMiDs to the forefront of AML therapeutic arsenal, a better understanding of its mechanism of action in AML cells and in the interphase

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**Figure 7.** Effects of Lenalidomide on primary blasts from patients with del5q/5q- AML. (A) Scheme depicting the experimental design. (B) Lenalidomide toxicity on primary del5q/5q- AML cells (n = 4). (C) When combined with standard chemotherapy lenalidomide largely overcomes BM-MSC-mediated resistance of del5q/5q- AMLs to AraC+Idarubicin (n = 4). (D) Effect of lenalidomide on the expression levels of CXCR4 in del5q/5q- AML cells (n = 4). AMD3100 was used as positive control. *p < 0.05; n.s, no statistical differences.
tumor-stroma is required. In this study, we hypothesized that the testing of IMiDs may be facilitated by in vitro and in vivo assays that model the tumor-BM stroma interactions.\(^{47-49}\) Hence, we used human primary BM-MSCs and intra BM co-transplantation with AML cells into NSG mice to mimic the interaction of AML cells with the BM stroma as a platform to assess whether IMiDs add efficiency to current standard AML chemotherapy. We report that lenalidomide and pomalidomide, when administered as single agents, have cytotoxic effects on neither non-del5q/5q AML cells nor BM-MSCs, even at doses as high as 100 µM. This contrasts the anti-proliferative effects reported for IMiDs in other hematopoietic malignancies, evidencing the immature nature of AML cells (progenitor-like cells) and the complex biology of AML.\(^{9-11}\)

However, IMiDs enhanced the immunomodulatory properties of BM-MSCs as demonstrated by the suppression activated T cells proliferation, inhibition of the production of pro-inflammatory cytokines including TNF\(\alpha\), IL1\(\beta\) and IL6 and increased production of the anti-inflammatory cytokine IL10, suggesting that IMiDs may represent safe candidate drugs interfering with the protective BM stroma for combination with current AML chemotherapy.

Unfortunately, when combined with AraC and Idarubicin, both lenalidomide and pomalidomide fail to circumvent BM stroma-mediated resistance of non-del5q/5q AML cells in in vitro and in vivo AML models. The clinical efficacy of lenalidomide has been explored as either low-dose or high-dose lenalidomide as monotherapy\(^{50-54}\) and in combination with azacitidine\(^{55,56}\) for both newly diagnosed and relapsed/refractory AML. Overall, these clinical trials were somewhat disappointing with CRR as low as 11%–20% and remission duration below 5–7 months.\(^{14}\) Our pre-clinical studies mechanistically support previous modest clinical activity reported for IMiDs in AML, and discourages future clinical trials using either lenalidomide or pomalidomide for non-del5q/5q AML either as monotherapy or polytherapy. Of note, a recent Phase II clinical study of lenalidomide combined with intensive chemotherapy in AML has reported that only del5q/5q- AML patients show promising CRR.\(^{39}\) Here, our pre-clinical studies confirm that lenalademediated as single agent displays ~20% toxicity only in del5q/5q AML cells and it overcomes BM-MSC/stroma-mediated chemoresistance of AML cells exclusively from del5q/5q- patients. A potential mechanism of action of lenalidomide involves the degradation of casein kinase 1A1 (CK1\(\alpha\)).\(^{57}\) In fact, del5q MDS patients possess a heterozygous deletion CK1\(\alpha\).\(^{58}\) Collectively, our data is in line with Ades and colleagues\(^{39}\) and suggest that the mechanisms of action of lenalidomide on AML may depend on cell intrinsic/autonomous genetic makeup, further pinpointing genetic and biological heterogeneity in AML. Despite the overall limited activity of lenalidomide during induction/consolidation therapy its efficiency during AML maintenance therapy has yet to be tested.

Intriguingly, we found that lenalidomide, administered either as monotherapy or combined with conventional chemotherapy, induces robust mobilization of non-del5q/5q AML cells to extramedullary hematopoietic tissues including PB and spleen, suggesting that IMiDs might function as plerixafor, a small molecule inhibitor of the CXCR4/SDF1 axis, extensively used for chemosensitization of AML blasts by disrupting the interaction of leukemic blasts with the environment.\(^{43,36,37,59}\) In fact, lenalidomide mediated the mobilization of AML cells to PB through downregulation of CXCR4 in non-del5q/5q AML cells but does not affect either the expression or the production of its ligand SDF1 by BM-MSCs. In contrast to plerixafor, lenalidomide does not mobilize normal CD34\(^+\) HSPCs to PB. Whole-genome GEP revealed a transcriptomic migratory signature of lenalidomide-treated non-del5q/5q AML primary blasts but not lenalidomide-treated CD34\(^+\) cells, confirming that the IMiD-mediated mobilization is specific of AML blasts. We cannot rule out that other signaling pathways/mechanisms contribute to the IMiD-induced mobilization phenotype. Importantly, in contrast to plerixafor, IMiDs do not chemosensitize AML blasts to chemotherapy, further supporting that the addition of IMiDs to cytotoxic chemotherapy seems not feasible in AML.\(^{36,59}\) Lenalidomide-attenuated pro-inflammatory cytokines including TNF\(\alpha\), IFN\(\gamma\), IL1\(\beta\) and macrophage inhibitory factor (MIF) have been previously described to interact with master pathways/molecules such as CXCR4/SDF1 axis, collagenases, integrins and eicosanoids in controlling cell migration and mobilization.\(^{60-63}\) Thus, the immunomodulatory effects exerted by IMiDs seem to influence the BM niche in a way non-del5q/5q AML cells physically detach from BM stroma. Future in vivo imaging studies will precisely define the mechanisms underlying the lenalidomide-mediated AML chemotaxis.

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**Disclosure statement**

The authors have nothing to disclose.

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**Author contributions**

BGL: designed and performed experiments, analyzed data and wrote the manuscript. HRH, RDG, DRM, CP, FG, PC, AI, CB: designed and performed experiments, and analyzed data. EA, JAPS, JABG, FAA, MR, PM, AUI: provided leukemic samples. PM: conceived the study, designed experiments, analyzed data and wrote the manuscript.


