Peptides and small molecules blocking the CXCR4/CXCL12 axis overcome bone marrow-induced chemoresistance in acute leukemias

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Abstract. Notable advances in treatment have been made and increases in the cure rates of pediatric leukemia have been achieved. However, the majority of children with relapsed disease are not expected to survive, with chemotherapy resistance acting as the principal cause of treatment failure. Interaction between leukemic cells and the bone marrow microenvironment is the primary cause of relapse. It was identified that a multi-protein membrane complex, formed by potassium voltage-gated channel subfamily H member 2 (hERG1) channels, the β1 integrin subunit and the stromal cell-derived factor 12 (CXCL12) receptor, C-X-C chemokine receptor type 4 (CXCR4), exerts a role in mesenchymal stromal cell (MSC)-mediated chemoresistance in pediatric leukemias. hERG1 blockade was able to overcome chemoresistance in vitro and in vivo. As an alternative strategy to overcome chemoresistance, the present study evaluated the effects of novel tools targeting the CXCR4/CXCL12 axis. The analysis of CXCL12 structural dynamics was used for the selection of a peptide (4-1-17) and a small molecule (8673), which interact with a transient hot spot, identified by a dynamic drug design approach. The present findings indicated that peptide 4-1-17 and small molecule 8673 inhibited leukemia cell proliferation and induced a pro-apoptotic effect, which was not reduced by the presence of MSCs. The combined treatment with 4-1-17 and 8673 had a stronger pro-apoptotic effect, particularly on cells cultured on MSCs in normoxic and hypoxic conditions, and was able to overcome MSC-induced resistance to cytarabine. Overall, the targeting of CXCL12 and the ensuing inhibition of the CXCR4/CXCL12 axis may be proposed as an alternative strategy to overcome chemoresistance in leukemia.

Introduction

Acute leukemia is the most common form of childhood cancer, accounting for ~30% of pediatric cancer cases (1,2). Despite the considerable progress that has been made in its treatment, this disease remains a leading cause of pediatric cancer-associated mortality, and its prognosis is unfavorable for children with relapsed or refractory disease. The principal cause of treatment failure is chemotherapy resistance (2,3).

An emerging concept suggests that leukemia cells and their interactions with the bone marrow (BM) microenvironment are the primary causes of acute leukemia relapse, due to the survival of residual cells following chemotherapy (4-6). A novel mechanism for the protection exerted by BM mesenchymal stromal cells (MSCs) on acute lymphoblastic leukemia (ALL) cells against chemotherapy has been identified. Pivotal to this mechanism is a multiprotein complex expressed on the plasma membrane of leukemic cells, consisting of potassium voltage-gated channel subfamily H member 2 (hERG1) channels, the β1 subunit of integrin receptors and the stromal cell-derived factor 12 (CXCL12) receptor, C-X-C chemokine receptor type 4 (CXCR4) (hERG1/β1/CXCR4) (7). These data gave support to the previously demonstrated functional link between hERG1 K+ channels and CXCL12 in acute leukemic cell migration (8). Overall, by controlling leukemia cell survival and motility, the hERG1/β1/CXCR4 complex has emerged as a target of choice for anti-chemoresistance strategies. Indeed, blocking hERG1 with classical hERG1-specific blockers overcomes MSC-induced chemoresistance, in vitro and in vivo, in ALL mouse models (7).

hERG1 is a voltage-dependent potassium channel, the functional relevance of which in human leukemia has been repeatedly proven (7,9-11). Furthermore, the alternative transcript of the hERG1 gene, hERG1B, is an independent

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prognostic factor of a high risk of relapse in pediatric T-ALL (12). Although a number of hERG1-specific blockers exist on the market, a number of them produce adverse cardiac side effects (13,14). Besides developing hERG1B-specific, non-cardiotoxic blockers (15), an alternative strategy to target the hERG1/B1/CXCR4 complex in leukemia cells may involve targeting CXCR4 and its ligand CXCL12.

The chemokine receptor CXCR4 is overexpressed in leukemia and is associated with poor outcomes in ALL and acute myeloid leukemia (AML) (6,16). The ligand CXCL12 is constitutively produced by MSCs, particularly under hypoxic conditions (17), and contributes to the migration and survival of leukemic blast cells through the activation of phosphatidylinositol 3-kinase/RAC-α serine/threonine-protein kinase (Akt) and mitogen-activated protein kinase (MAPK) pathways (5). Notably, chemotherapeutic treatments have been demonstrated to upregulate CXCR4 expression. Such upregulation represents a mechanism of acquired resistance in pediatric AML (18). A number of tools have been developed to block CXCR4/CXCL12 interactions, and they are currently under different stages of development (5). Research focused on targeting chemokine receptors has also identified promising molecules, which have subsequently been unsuccessful in clinical trials. Certain of these molecules suffered from low oral bioavailability, e.g. the C-C chemokine receptor type 5 (CCR5) inhibitor TAK-779 (19), while others gave rise to severe side effects, e.g. Apalviro, also inhibiting CCR5 and causing hepatotoxicity (20). On the other hand, chemokines, commonly considered ‘undruggable’ due to their small size and shallow surfaces, have re-emerged as drug development targets through novel biochemical approaches (21).

Novel tools targeting CXCL12 have recently been developed by means of combining paramagnetic fragment-based nuclear magnetic resonance (NMR) investigation, molecular dynamics (MD) and docking simulations (22). We here provide in vitro biological data on the effects of some of these molecules on acute leukemia cell survival.

Materials and methods

Peptides and small molecules. Peptides were synthesized by solid-phase synthesis using standard Fmoc chemistry in a Syro multiple peptide synthesizer (MultiSynTech GmbH, Witten, Germany). The final product was cleaved from the solid support, deprotected by treatment with trifluoroacetic acid containing tri-isopropylsilane and water (95/2.5/2.5), and precipitated with diethyl ether. Crude peptides were purified by reverse-phase chromatography. The final peptide purity and identity were confirmed by reverse-phase chromatography on a Phenomenex Jupiter C18 analytical column and by mass spectrometry with a Bruker Daltonics ultraflex matrix assisted laser desorption/ionization-time of flight tandem system (Bruker Corporation, Billerica, MA, USA). Peptide 4-175-185 MS: Calculated for C117H174N26O37S2 was 2,600.91; detected 2,598.29 m/z; HPLC RT (80%A-20%A) 23.33 min. Positive ion mode was used in all cases. All surface plasmon resonance experiments were performed on a BIA T100 system (GE Healthcare, Chicago, IL, USA). The binding of CXCL12 was performed on a streptavidin (SA)-sensor chip previously coated with biotinylated peptides. The immobilization of the biotinylated peptide(s) was achieved by injecting peptide(s) diluted at 50 μg/ml in HBS-EP* (10 mM HEPES, 150 mM NaCl, 3.4 mM EDTA, 0.05% polysorbate 20; pH 7.4) for 60 sec over an SA-coated flow cell at the flow rate of 10 μl/min. The binding of CXCL12 with immobilized peptides was investigated using injections for 180 sec at chemokine concentrations between 1 and 100 nM in HBS-EP*, at a flow rate of 30 μl/min. The regeneration of the matrix was achieved by flushing a short pulse of 1 M NaCl-10 mM NaOH. For small molecule binding, CXCL12 was immobilized on a dextran matrix of a CM4 sensor chip flow cell via a standard amino coupling procedure. A flow cell was used as a reference surface following a blank immobilization. Small molecules were injected over a CXCL12-coated chip for 60 sec at a flow rate of 30 μl/min and diluted at 100 μM in 5% dimethyl sulfoxide-HBS-EP*; which was also used as the running buffer. Molecular docking simulation was performed using the AutoDock Vina package v 1.1.2 (23) using a ZINC-derived library (http://zinc.docking.org) of small molecules probed against a CXCL12 pocket region previously identified by NMR experiments and MD simulations (22), and consisting of residues V23, K24, H25, K27, A40, R41 and K43. Standard Vina parameters were used for the simulation runs. Selected molecules were purchased from ChemBridge Corporation (San Diego, CA, USA).

Cell culture. Leukemia cell lines [B-cell precursor-ALL (BCP-ALL) 697 cells and AML FLG 29.1 cells] and normal Epstein-Barr virus (EBV)-infected B lymphocytes were cultured in RPMI-1640 medium (Sigma-Aldrich; Merck KGaA, Darmstadt, Germany) supplemented with 2 mM L-glutamine (EuroClone SpA, Pero, Italy) and 10% fetal calf serum (FCS; EuroClone). Human BM-derived MSCs immortalized by enforcing the expression of telomerase reverse transcriptase in primary MSCs were established in the laboratory of Dr D. Campana (Department of Pediatrics, National University of Singapore, Singapore) and maintained as described previously (7). MSCs were seeded in 96-well flat-bottomed plates coated with fibronectin (1 μg/well) and grown until confluence prior to undertaking the co-culture experiments.

Primary samples. BM samples from children with newly diagnosed AML were analyzed at the Hematology-Oncology Laboratory of the Department of Pediatrics, University of Padua (Padua, Italy). Diagnoses were made according to standard cytomorphology, cytochemistry, an immunophenotypic criteria as described previously (12). Patients studied were enrolled in the AIEOP-BFM ALL 2009 therapy protocol, approved by the local ethical committee (Comitato Etico per la Sperimentazione dell’Azienda Ospedaliera di Padova; no. 0002862-18/01/2012). The parents or legal guardians of
the patients provided written informed consent, following the tenets of the Declaration of Helsinki.

Trypan blue assay. Cell viability was assessed by trypan blue exclusion assay. In brief, 20 µl 0.4% trypan blue solution was added to 20-µl cell suspensions in culture medium. The suspension was gently mixed and transferred to a hemocytometer. Viable and dead cells were identified and counted under a light microscope (x10 magnification) Blue cells failing to exclude the dye were considered nonviable, and transparent cells were considered viable. The percentage of viable cells was calculated on the basis of the total number of cells (viable and non-viable). The median lethal dose (LD50) value was calculated by fitting the data points (following 24 h of incubation) with a sigmoidal curve using OriginPro 2015 (OriginLab, Northampton, MA, USA).

Pharmacology experiments. Leukemic cells were serum-starved for 16 h in RPMI medium and seeded in 96-well flat-bottomed plates (Corning-Costar; Corning Incorporated, Corning, NY, USA) at a cell density of 2x10^5 cells/well in RPMI containing 10% FCS.

Small molecules and peptides were used at the LD50 values indicated in the figures following three different schedules: i) Single treatment (added at timepoint 0); ii) double treatment (added at timepoints 0 and 12 h); and iii) triple treatment (added at timepoints 0, 12 and 24 h).

Following 24, 48 and 72 h of incubation, viable cells (determined by the trypan blue exclusion test) were counted in triplicate using a hemocytometer. Each experimental point represents the mean of four samples from three independent experiments.

Co-culture experiments. Cell suspensions (at a cell density of 2x10^5 cells/well) were placed in a 96-well flat-bottomed plate, with or without bone marrow-derived MSCs and treated with 4-1-17 or 8673 at the LD50 dose alone, or in combination with cytarabine (45 nM) or doxorubicin (0.1 µg/ml). Cultures were maintained for 48 h at 37°C, 5% CO2, 7, 8 and 9. Leukemic cells were treated for 48 h (as discussed above) and subsequently incubated at 37°C, 5% CO2 for 2 h with 2 µl 500X TF2-VAD-FMK. Following PBS washing, cells were resuspended in 0.5 ml assay buffer and analyzed with BD FACSDiva Software 6.1.3 (BD Biosciences).

Hypoxia experiments. Exponentially growing cells were treated as described above and incubated at 37°C in 0.1% O2 (water-saturated atmosphere containing 94.9% N2, and 5% CO2) in a DG250 Anaerobic Workstation (Don Whitley Scientific, Ltd., Bingley, UK) for 48 h.

Western blotting. Protein extraction and western blotting were performed largely as described in (7). Leukemic cells following treatment were washed with cold PBS and immediately extracted with 1% NP-40 lysis buffer (1% NP-40, 150 mM NaCl, 50 mM Tris-HCl, pH 8, 5 mM EDTA and 10 mM Na2P2O7) supplemented with a tablet of a complete mix of protease inhibitors (Roche Complete Mini; Roche Diagnostics). Cell lysates were centrifuged at 13,000 x g for 10 min (4°C), and the supernatants were collected and assayed for protein concentration with the Bradford protein assay method (Bio-Rad Laboratories, Inc., Hercules, CA, USA). Proteins were eluted by boiling the samples in Laemlli buffer, analyzed by SDS-PAGE (7.5%) under reducing conditions, and transferred to a nitrocellulose membrane (Hybond P; Amersham; GE Healthcare). The membrane was incubated for 4 h at room temperature with 0.1% Tween-20 in PBS (T-PBS) containing 5% bovine serum albumin, Sigma-Aldrich; Merck KGaA) and incubated overnight at 4°C with the appropriate primary antibodies at the concentrations listed below. Membranes were washed three times with T-PBS and incubated with the appropriate secondary antibodies for 45 min at room temperature. Following three washes with T-PBS, the immunoreactivity was determined by an enhanced chemiluminescence reaction (SuperSignal; Pierce; Thermo Fisher Scientific, Inc., Waltham, MA, USA). For the stripping of the membranes, the ReBlot WB recycling kit (Chemicon; EMD Millipore, Billerica, MA, USA) was used, according to manufacturer's protocol.

The following primary antibodies were used: Anti-phospho-p44/42 MAPK (Thr202/Tyr204) (Cell Signaling Technology, Inc., Danvers, MA, USA; cat. no. 9101; dilution, 1:500) and anti-pAkt1/2/3 (Thr308)R (Santa Cruz Biotechnology, Inc., Dallas, TX, USA; cat. no. sc-271966; dilution, 1:500). Anti-α-tubulin mouse monoclonal (Sigma-Aldrich; Merck KGaA; cat. no. T9026; dilution, 1:500), anti-α-tubulin mouse monoclonal (Sigma-Aldrich; Merck KGaA; cat. no. T9026; dilution, 1:500) and anti-IκBα mouse monoclonal ( Sigma-Aldrich; Merck KGaA; cat. no. A0545; dilution, 1:10,000) and anti-IκBα mouse monoclonal (Sigma-Aldrich; Merck KGaA; cat. no. A0545; dilution, 1:5,000).

Western blotting images were acquired with an Epson 3200 scanner, and the relative bands analyzed with Scion Image software version 4.0 (Scion Corporation, Frederick, MD, USA). The intensity of the bands was normalized to the intensity of the bands that corresponded to the total protein. The control cell ratio was set as 1.
Statistical analysis. Graphs and statistical analyses were prepared using Prism 4.00 (GraphPad Software, Inc., La Jolla, CA, USA). Values in all panels are the mean ± standard deviation of three independent experiments. The normality of the data distribution was checked with the Kolmogorov-Smirnov test. In the case of normal distributions, each dataset was first checked for variance homogeneity, using the Brown-Forsythe test for multiple comparisons. For multiple comparisons, one-way analysis of variance followed by Bonferroni’s post hoc test was performed to derive the P-values. P<0.05 was considered to indicate a statistically significant difference.

Results

Selection of peptides from CXCR4. Amino acid (aa) residues involved in, and putatively modulating, the CXCR4/CXCL12 interaction were identified from CXCR4 fragments located in regions in which the interaction with CXCL12 had been inferred either by mutagenesis (24), structural modeling of the ligand receptor/complex (25), NMR evaluation of the binding of CXCR4 N-terminal peptides spanning aa 1-27 (26), or the NMR-derived structure of CXCL12 in complex with a CXCR4 N-terminal fragment spanning aa 1-38 (27). Comparison of the two NMR studies allowed for the accurate selection of a shorter (thus more suitable as a molecular tool) peptide, strongly interacting with CXCL12 when used as a fragment against CXCL12 (26). The left panel of Fig. 1 illustrates the mode of interaction of peptide 4-1-17 with the chemokine as derived from the complex CXCL12/CXCR4_1-38 (27) (Fig. 1; left) and marked binding (in terms of NMR chemical shift perturbation) when used as a fragment against CXCL12 (26). The left panel of Fig. 1 illustrates the mode of interaction of peptide 4-17 with the chemokine as derived from the complex CXCL12/CXCR4_1-38 (Protein Data Bank ID 2K04; http://www.rcsb.org); a strong interaction between the peptide and the crevice surface formed by the N-terminal and the central β-sheet of CXCL12 is apparent.

The CXCR4_4,27 sequence, on the contrary, exhibited no binding to CXCL12 when used as a fragment. Due to the lack of binding data for the remaining sequence CXCR4_29,35 and the large conformational variability in the CXCL12/CXCR4_1-38 structure, the peptide spanning aa 29-35 (termed 4-29-35) of CXCR4 was selected for the evaluation of binding capability of this region. Although no other experimental structural data are available for CXCR4, mutagenesis and modeling studies (24,25) indicated the second extracellular loop ECD2 as being involved in ligand binding and signaling, due to its interaction with the CXCL12 loop linking β-strands I and II. Thus, the CXR4_4,185 fragment spanning the ECD2 sequence was selected for evaluation (termed 4-175-185). CXCL12 binding to another seven transmembrane span receptor, CXCR7 (28), suggested the selection of an additional peptide from the latter one. Although the structure of CXCR7 remains to be resolved, comparative molecular modelling predicted that interacting regions of CXCR4 and CXCR7 with CXCL12 are similarly located in the N-terminal region (29). Hence, to exploit the CXCR4/CXCR7/CXCL12 chemokine axis, the sequence segment spanning the N-terminal residues 1 to 17 of the CXCR7 was selected to obtain a peptide (termed 7-1-17) homologous to 4-1-17.

Selection of small molecules. Small molecules modulating the CXCR4/CXCL12 interaction were selected from a virtual library [Clean Leads subset from ZINC database; (30)] using experimental data as constraints for a molecular docking simulation. In particular, all molecules were docked to the transient binding pocket opening on Val23/Ala40 (22). Transient pockets, indeed, refer to their rapid appearance and disappearance on flat protein surfaces as a consequence of fluctuations in conformational dynamics [hence the definition of dynamic drug design (DDD)]. The presence of such a pocket was inferred by an NMR paramagnetic perturbation study, while the open conformation was determined by trajectory analysis of molecular dynamics simulation of CXCL12 (22). The trajectory frame representing the pocket open conformation was used for the docking run, and the simulation box was narrowed down to the involved residues V23, K24, H25, K27, A40, R41 and K43. A total of one out of the three chosen small molecules, henceforth termed 8673, had already been reported to interact with CXCL12 in previous investigations (31), and its binding geometry is reported in Fig. 1 (right panel).

Selected peptides and small molecules were tested for chemokine binding by surface plasmonic resonance (see Tables I and II).

Effect of peptides and small molecules targeting CXCL12 on leukemia cell vitality and proliferation. The panel of molecules reported above was tested in a viability assay. For each molecule, the LD50 in AML (FLG 29.1) and ALL cell lines (697) were determined. Dose-dependence curves and LD50 values measured following 24 h of treatment are presented in Fig. 2 and Table I, respectively. Peptide 4-1-17 and the small molecule 8673 had a strong anti-proliferative effect on the two leukemic cell lines at micromolar concentrations. Similar effects were observed for small molecule 9430, although to a lesser extent. On the other hand, 4-175-185 and 9355, which did not bind CXCL12 in the SPR assay (see Table II), did not exert any effects on cell viability at concentrations up to 100 µM.

Fig. 3 illustrates the effects of 4-1-17 (Fig. 3A) and 8673 (Fig. 3B), tested at their LD50 value, on the proliferation of FLG 29.1 and 697 leukemic cells, in addition to that of normal EBV-infected B lymphocytes. The two molecules almost completely inhibited cell proliferation in either leukemic cell lines in the first 24 h of incubation. Subsequently, cells recommenced proliferation, although at a lower rate. However, when the two compounds were re-added to the cells following 12 and 24 h of incubation, leukemia cell proliferation was almost abolished. On the contrary, 4-1-17 and 8673 did not affect the cell viability of normal EBV-infected B lymphocytes (Fig. 3A and B lower panels).

Effect of peptides and small molecules targeting CXCL12 on leukemia cell apoptosis. The effects of 4-1-17 and 8673 on cellular apoptosis were also tested (at their LD50 value) on the two leukemia cell lines (FLG 29.1 and 697) and on primary BCP-ALL samples, in suspension or co-culture with MSCs (as in 7) and in the absence or presence of classical chemotherapeutic drugs (cytarabine in AML, or doxorubicin in ALL). As expected (7,32,33), MSCs significantly protected leukemic cells from either spontaneous, or cytarabine- (in AML) or doxorubicin- (in ALL) induced apoptosis (Fig. 4). On the contrary,
Figure 2. Dose response curves for peptides and small molecules targeting CXCL12 in leukemic cell lines. Acute myeloid leukemia (FLG 29.1) and B-cell precursor-acute lymphoblastic leukemia (697) cells were cultured and exposed to increasing concentrations of peptides and small molecules for 24 h. The percentage of trypan blue-negative cells was measured. LD₅₀ values were evaluated by nonlinear regression analysis using OriginPro 2015. Values in all panels are the mean ± standard deviation of three independent experiments. Statistical analysis was performed with one-way analysis of variance followed by Bonferroni's post hoc test. LD₅₀, median lethal dose.
the pro-apoptotic effects exerted by either 4-1-17 or 8673 were not reduced by MSCs in AML and ALL cells (Fig. 4).

Since 4-1-17 and 8673 bind CXCL12 at different sites, the effects of the combination of the two molecules were tested in FLG 29.1 cells cultured either in suspension or onto MSCs. The combined treatment had a strong pro-apoptotic effect, evidenced by the increase in the percentage of apoptotic cells and by caspase activation (Fig. 5A and B). This effect was more evident in cells cultured on MSCs, and even more when cells were treated with cytarabine. Overall, these data indicated that peptide 4-1-17 and small molecule 8673 overcame MSC-induced resistance to cytarabine (Fig. 5).

Table I. Peptides.

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<th>Code</th>
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<th>CXCR7 sequence span</th>
<th>Sequence</th>
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<th>Kd, 298 K</th>
<th>FLG 29.1</th>
<th>697</th>
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<tr>
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<td>66.7±2.3</td>
<td>28±1.4</td>
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CXCR4, C-X-C chemokine receptor type 4; CXCR7, atypical chemokine receptor 3; Kd, dissociation constant; LD50, median lethal dose; MM, molecular mass. Nd, not determined.
Figure 4. Pro-apoptotic effect of peptide 4-1-17 and small molecule 8673 in AML and ALL cell lines. (A) The AML cell line FLG 29.1 was exposed to the LD_{50} dose of peptide 4-1-17, small molecule 8673 or cytarabine (45 nM) for 48 h. (B) The ALL cell line 697 was exposed to the LD_{50} dose of peptide 4-1-17, small molecule 8673 or doxorubicin (0.1 µg/ml) for 48 h. (C) The AML cell line FLG 29.1 was exposed to the LD_{50} dose of peptide 4-1-17 and small molecule 8673 alone or in combination with cytarabine (45 nM) for 48 h. (D) The ALL cell line 697 was exposed to the LD_{50} dose of peptide 4-1-17 and small molecule 8673 alone or in combination with doxorubicin (0.1 µg/ml) for 48 h. (E) A total of two representative B cell precursor-ALL primary samples were cultured with or without MSCs and treated with LD_{50} doses of 8673 in the presence of doxorubicin. Values in all panels are the mean ± standard deviation of three independent experiments. Statistical analysis was performed with one-way analysis of variance followed by Bonferroni’s post hoc test. *P<0.05; **P<0.01. AML, acute myeloid leukemia; ALL, acute lymphoblastic leukemia; MSC, mesenchymal stem cell; LD_{50}, median lethal dose.

Table II. Small molecules.

<table>
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<td>9430</td>
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<td></td>
<td>309.3</td>
<td>No binding</td>
<td>FLG 29.1: 36.0±2.3, 697: 33±2.4</td>
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K_{d}, dissociation constant; LD_{50}, median lethal dose; MM, molecular mass.
CXCL12/CXCR4 interactions have been demonstrated to trigger Akt and ERK signaling (33); the present study therefore tested the effects of 4-1-17 and 8673, alone or in combination, on the MSC-induced activation of Akt and ERK in FLG 29.1 cells. Fig. 6 demonstrates that 4-1-17 and 8673 downregulated ERK phosphorylation induced by MSCs (left panels), and that the combined treatment abrogated MSC-induced ERK phosphorylation. Similarly, the phosphorylation of Akt was downregulated following treatment with either 4-1-17 or 8673, an effect particularly evident when MSCs were added to the leukemia cells (right panels).

Since hypoxia has a relevant impact on leukemic cell survival (34), the effect of the combined treatment was assessed in FLG 29.1 cells cultured either in suspension or onto MSCs in hypoxic conditions. Similar to what was observed in normoxia, the addition of 4-1-17 and 8673 induced a pro-apoptotic effect and overcame MSC-mediated chemoresistance (Fig. 7).

Discussion

The present study provided evidence that two different molecular tools targeting the CXCL12/CXCR4 complex, peptide 4-1-17 and the small molecule 8673, inhibited the proliferation of AML and ALL cell lines by inducing apoptosis. In contrast to what occurs with chemotherapeutic drugs, this effect was not reduced by the presence of MSCs. Furthermore, the two tools significantly increased the sensitivity of leukemic cells to cytarabine in the presence of MSCs.

The principal cause of treatment failure in acute leukemia, particularly in the pediatric setting, which is generally associated with better outcomes, is chemotherapy resistance (3,35). Leukemic cells have been reported to take refuge within the BM niche (36), which thus leads to survival of residual leukemic cells following chemotherapy, resulting in disease relapse. In other words, leukemic cells that adhere to MSCs...
Figure 5. Continued. (B) Caspase-activity in FLG 29.1 cells, cultured with or without MSCs and exposed under normoxic conditions for 48 h. Representative histograms for pan-caspase analysis are presented on the right. The right side of the vertical line indicates the pan-caspase-positive region in each panel. The position of this line was the same for all treatments, as the control setting. Values in all panels are the mean ± standard deviation of three independent experiments. Statistical analysis was performed with one-way analysis of variance followed by Bonferroni's post hoc test. *P<0.05; **P<0.01. FITC, fluorescein isothiocyanate; PI, propidium iodide; MSC, mesenchymal stem cell.

Figure 6. Combined treatment with 4-1-17 and 8673 inhibits MSC-induced activation of Akt and ERK. Acute leukemia cells (FLG 29.1) were cultured with or without MSCs (suspension) and exposed to the LD50 dose of peptide 4-1-17 and small molecule 8673, alone or in combination, for 48 h. Phosphorylation of p-ERK1/2 Thr202/Tyr204 (42/44 kDa) and p-Akt Thr308 (60 kDa) was detected by western blot analysis. Membranes were also reprobed with an anti-ERK1/2, anti-Akt or anti-tubulin antibody. The corresponding densitometric analyses are presented in the bar graphs, displayed as the ratios of p-proteins to total proteins. Akt, RAC-α serine/threonine-protein kinase; ERK, extracellular signal-regulated kinase; MSC, mesenchymal stem cell; p, phosphorylated.
through CXCR4 are protected from the effects of cytotoxic chemotherapy and represent a reservoir for minimal residual disease and relapse (37). CXCL12 and its cognate receptor, CXCR4, are two key mediators in the cross-talk between leukemia cells and their microenvironment.

A novel mechanism of chemoresistance in ALL was previously elucidated, centered on a plasma membrane macromolecular complex consisting of the hERG1 potassium channel, the CXCR4 chemokine receptor and the β1 integrin subunit (7). The role of hERG1 was critical, as hERG1 inhibitors abrogated the protective effect of MSCs and enhanced the cytotoxicity of drugs commonly used to treat leukemia. However, the targeting of hERG1 channels with classical hERG1-specific blockers is difficult to propose in the clinical setting, due to the potential cardiotoxicity displayed by classical hERG1 blockers (13,14,38). For this reason, aside from using non-cardiotoxic hERG1 blockers (15,38), an alternative therapeutic strategy was proposed to specifically target the hERG1/β1/CXCR4 complex, and in turn overcome chemoresistance in leukemia. Such a strategy consisted of inhibition of the interaction between CXCL12 and its receptor CXCR4 with either peptides or small molecules developed following a previously described procedure (22), and targeting the chemokine CXCL12.

Previous studies provided evidence that targeting CXCR4 with different agents (AMD3100, AMD3465, BKT140, RCP168 and TN140) increased the sensitivity of leukemia cells to chemotherapy (33,39,40). AMD3100 was approved by the FDA in 2008 for the mobilization of hematopoietic stem cells as an injectable agent for short-term treatments (in patients with non-Hodgkin's lymphoma and multiple myeloma) and is currently in phase I clinical trials for AML and chronic myeloid leukemia. The safety and feasibility of combining AMD3100 with chemotherapy in patients with myeloma and lymphoma has already been demonstrated. On the contrary, concerns remain regarding the administration of AMD3100 to patients with acute leukemia.

Figure 7. Addition of 4-1-17 and 8673 in hypoxic conditions induces a pro-apoptotic effect and an overcoming of MSC-associated chemoresistance. (A) The acute leukemia cell line (FLG 29.1; acute myeloid leukemia) was cultured with or without MSCs (suspension) and exposed to combined treatment with peptide 4-1-17 and small molecule 8673 (the LD50 dose) alone or in the presence of cytarabine (45 nM) for 48 h, under hypoxic conditions (0.1% O2). The percentage of Annexin V+ cells was measured. Representative dot plots of Annexin V and PI staining in FLG 29.1 cells cultured under hypoxia are presented in the right-hand panels.
Concerns are primarily associated with BM aplasia, delayed hematopoietic recovery following chemotherapy and CXCR4 upregulation (41). In addition, such treatment may result in a preferential mobilization of leukemic blasts over normal cells. Another compound, the orally active AMD11070, which binds overlapping non-identical residues in the binding pocket of the receptor, is currently in phase I clinical trials for cancer (42,43). Furthermore, a number of novel tools to block CXCR4/CXCL12 interactions are under development, being either peptides or small molecules. Among the peptide class, T140 and its stable derivative BKT140 have recently been reported to target AML anchorage in the BM, in addition to the differentiation and survival of leukemic cells (44-46). The same molecule is undergoing further development via substitution of uncharged and negatively-charged side chains with positively-charged ones. Still in the ‘biological’ domain, the fully human anti-CXCR4 antibody BMS-936564 has exhibited exerting effects similar to those of the AMD3100 small molecule drug, in addition to pro-apoptotic activity (47). Regarding small molecules, the isothiourea derivative IT1t has been proven to be an antagonist for CXCR4, and the structure of IT1t/CXR4 complex has been resolved by X-ray crystallography (25), opening the way to rational design of molecules with improved efficacy (48).

All of the tools reported above target the CXCR4 receptor, since chemokines are generally viewed as ‘undruggable’ proteins. The two molecular tools here proposed target the CXCL12/CXCR4 complex, although they bind to the ligand, CXCL12, rather than the receptor. Notably, targeting a specific ligand may facilitate regulation rather than the elimination of receptor activity. The two compounds, 4-1-17 and 8673 are very different in nature, the former being a 17-mer peptide derived from the N-terminal domain of CXCR4, and the latter, a small drug-like molecule selected by virtual screening. The two compounds target CXCL12, although they
use distinct binding sites. Standard screening for potential drugs is usually performed by examining the static features of target protein surfaces. On the other hand, in the present study, CXCL12 structural dynamics were used as a rational framework for drug selection. Accordingly, disruptors of the CXCL12/CXCR4 interaction were predicted and tested for their activity. Thus, one peptide, which was shorter than those previously suggested, and one small molecule, which interacted with a transient hot spot, were highlighted by the present DDD approach. The fact that 4-1-17 and 8673 bind CXCL12 at different sites suggests that the conjugation of the two molecules may exhibit stronger biological activity. Taken together, the present findings demonstrated the efficacy of the CXCR4/ hERG1/ β1 integrin, and hERG K+ channel for a macromolecular signaling complex in acute myeloid leukemia: Role in cell migration and clinical outcome. Blood 110: 1238-1250, 2007.


Ethics approval and consent to participate

Patients studied were enrolled in the AIEOP-BFM ALL 2009 therapy protocol, approved by the local ethical committee (Comitato Etico per la Sperimentazione dell’Azienda Ospedaliera di Padova; no. 0002862/18/01/2012). The parents or legal guardians of the patients provided written informed consent, following the tenets of the Declaration of Helsinki.

Patient consent for publication

The parents or legal guardians of the patients provided written informed consent, following the tenets of the Declaration of Helsinki.

Competing interests

The authors declare that they have no competing interests.

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Availability of data and materials

The datasets used during the present study are available from the corresponding author on reasonable request.

Authors’ contributions

BL performed the peptide synthesis. AB conceived the peptide and small molecules approach, performed modelling and docking work and contributed to writing the manuscript. OS performed the bioinformatics analysis. LB and NN contributed to the analysis of the results. SP performed the cell viability assay, pharmacology experiments, western blotting and hypoxia experiments. GP performed the caspase activity assay. AA, NN and SP designed the study. AA supervised the study and wrote the paper. SP and AB revised the paper. All authors read and approved the final manuscript.