Hematopoiesis

Inhibition of the anti-apoptotic protein MCL-1 severely suppresses human hematopoiesis

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ABSTRACT

Introduction

BH3-mimetics inhibiting anti-apoptotic BCL-2 proteins represent a novel and promising class of antitumor drugs. While the BCL-2 inhibitor venetoclax is already approved by the Food and Drug Administration, BCL-XL and MCL-1 inhibitors are currently in early clinical trials. To predict side effects of therapeutic MCL-1 inhibition on the human hematopoietic system, we used RNA interference and the small molecule inhibitor S63845 on cord blood-derived CD34+ cells. Both approaches resulted in almost complete depletion of human hematopoietic stem and progenitor cells. As a consequence, maturation into the different hematopoietic lineages was severely restricted and CD34+ cells expressing MCL-1 shRNA showed a very limited engraftment potential upon xenotransplantation. In contrast, mature blood cells survived normally in the absence of MCL-1. Combined inhibition of MCL-1 and BCL-XL resulted in synergistic effects with relevant loss of colony-forming hematopoietic stem and progenitor cells already at inhibitor concentrations of 0.1 μM each, indicating “synthetic lethality” of the two BH3-mimetics in the hematopoietic system.
signals, BH3-mimetics are able to bypass this mode of activation. Similar to BH3-only proteins, every BH3-mimetic available so far has specific binding affinities to one or more anti-apoptotic BCL-2 proteins (Online Supplementary Figure S1A). Navitoclax (ABT-263) and its intravenously used precursor drug, ABT-737, bind to BCL-2, BCL-XL and BCL-W. The drug showed good efficacy against non-small lung carcinoma and hematologic malignancies. However, its side effects on the hematopoietic system precluded its full clinical exploration and FDA approval. This indicated that a combined inhibition of more than one pro-survival BCL-2 protein might impede survival of healthy body cells. Later, a BCL-2-specific inhibitor called venetoclax (ABT-199) found its way into clinical trials. Thanks to the much less severe side effects, it was approved by the FDA in 2016 as a second-line treatment for chronic lymphocytic leukemia (CLL) with 17p deletion, and in 2019 for the treatment of all adult CLL and small lymphocytic lymphoma patients. For acute myeloid leukemia (AML), venetoclax was FDA-approved only in combination with hypomethylating agents.

Unfortunately, as for other cytotoxic drugs, tumor cell resistance poses a major problem to the efficacy of venetoclax. Primary resistance is present when tumor cells require anti-apoptotic BCL-2 proteins other than BCL-2 for survival. Naturally, only lymphocytes and melanocytes are dependent on BCL-2 expression, as shown in BCL-2 knockout mice. This might explain why venetoclax is most effective in mature lymphoma while most other tumors show primary resistance. Such primary resistance to venetoclax can also be caused by overexpression of pro-survival proteins other than BCL-2, such as BCL-XL and/or MCL-1.12,13 As shown for CLL, these BCL-2 homologs can be induced by signals from the tumor microenvironment.14 Secondary resistance, in contrast, is acquired by tumor cells to escape previously effective BCL-2 inhibition. Several mechanisms, such as BCL-2 mutations which strongly lower venetoclax affinity, have been implicated in the development of secondary venetoclax resistance.15 Alternatively, BCL-XL and MCL-1 overexpression was noted in relapsed CLL patients who had been previously treated with venetoclax.17,18 Therefore, the development and administration of MCL-1/BCL-XL inhibitors are much needed to overcome primary and secondary venetoclax resistance. MCL-1 inhibitors, in particular, are eagerly awaited by oncologists since this protein plays an essential role in many tumor types (e.g., AML, multiple myeloma, non-small cell lung carcinoma).19

MCL-1 was first identified during the differentiation of monocytes to macrophages in ML-1, a human myeloid leukemia cell line.20 Three isoforms of the gene have been reported; the most abundant anti-apoptotic MCL-1 long (MCL-1L) and two shorter pro-apoptotic isoforms (MCL-1 short, MCL-1 extra short).21,22 In addition, a truncated isoform was shown to localize at the mitochondrial matrix where it facilitates mitochondrial fusion and ATP synthesis.23 Genetic Mcl-1 deletion in mice revealed its essential role in many tissues, both during embryogenesis and in adult mice. Specifically, constitutive MCL-1 deficiency resulted in peri-implantation embryonic lethality,24 while targeted deletion in the fetal hematopoietic system resulted in loss of stem cells.25 When one Mcl-1 allele was deleted in adult mice, hematopoietic stem and progenitor cells (HSPC) were depleted, leading to the death of the animals within 2-3 weeks.26 With regard to human HSPC, there is only indirect evidence for the essential role of MCL-1, given by the BH3 profiling method: Mitochondria isolated from human CD34+ cells were highly sensitive to NOXA BH3 peptides, which typically correlated with MCL-1 dependency.27

Recently, a potent and specific MCL-1 inhibitor, S63845, was developed. This compound can efficiently kill a variety of tumor cell types such as multiple myeloma, lymphomas, leukemias and primary AML cells as well as to some extent solid cancers.28 Treatment of mice with S63845 resulted in only a few side effects in vivo,29 which was rather unexpected considering the many roles of MCL-1 during development and for tissue homeostasis. Here, we extended these studies to human cells and focused on the hematopoietic system. Understanding hematotoxicity of novel anticancer drugs is crucial since suppression of hematopoiesis accounts for most treatment-related morbidity and mortality. By using two different shRNA sequences and the MCL-1 inhibitor S63845, we consistently found that MCL-1 expression is essential for the survival of human stem and progenitor cells, especially during early stages of differentiation. In contrast, mature blood cells are less sensitive to MCL-1 inhibition. Of note, combined inhibition of MCL-1 and BCL-XL was synergistic and already low concentrations of both drugs resulted in profound stem and progenitor cell depletion.

Methods

Lentiviruses

A pLeGO-hU6 lentiviral vector with huU6 promoter and green fluorescent protein (GFP) expression was used to generate shRNA expressing lentiviruses (Online Supplementary Table S1).30 CD34+ cells were transduced with the lentivirus (2x MOI 10, 24 h each) and knockdown efficiencies were determined 24 h later.

Isolation and culture of human CD34+ cells

Umbilical cord blood and bone marrow were obtained immediately after birth or from patients (age: 44-90 years) undergoing orthopedic surgery, respectively. Informed consent was obtained and the ethics committee approved the study. CD34+ cells were isolated (by magnetic activated cell sorting) from mononuclear cells to a purity >90%. Cells were used either immediately or stored in liquid nitrogen (Cryo10 freezing medium, Sigma) for later use. Cells were cultured in serum-free StemPro-34 medium supplemented with embryonic stem cell fetal bovine serum (ES-FBS), penicillin/streptomycin (P/S; Invitrogen), stem cell factor (SCF), FMS-like tyrosine kinase 3 ligand (FLT3L) (200 ng/mL each), thrombopoietin (TPO; 100 ng/mL) and interleukin 3 (IL-3; 20 ng/mL; Immunotools/Peprotech). Where indicated, the BH3-mimetics S63845 (SynMedChem), A-1155463 and ABT-199 (Sellekchem) were added.

Apoptosis assay

CD34+ cells were subjected to cytokine deprivation or treated with etoposide (VP16), tunicamycin, taxol, thapsigargin and brefeldin A (BFA). After 0, 24 and 48 h, cells were stained with annexin V (Biorad) and 7-aminoactinomycin D (7-AAD; Sigma-Aldrich) to detect apoptosis. The percentage of specific apoptosis was calculated as: 100 x (% living cells under control condition - % living cells under treatment)/% living cells under control condition. Control condition was culture with ES-FBS and cytokines.
Colony-forming assays and differentiating culture

One thousand CD34+ cells were seeded in MethoCult SF-H4436 medium. After 10-11 days, colony types (identified by light microscopy) and total cell counts were determined. The percentages of HSPC, erythroid and myeloid cells were determined by flow cytometry (Online Supplementary Table S2).

Proliferating culture

CD34+ cells were cultured for 5-11 days in StemPro-34 medium supplemented with 10% ES-FBS, SCF, FLT3L, TPO and IL-3. The medium was refreshed every 3 days. Cells were analyzed for GFP and immature populations (Online Supplementary Table S2).

Quantitative reverse transcription polymerase chain reaction

RNA was isolated by a Quick RNA Micro Prep kit (Zymo Research) and reversely transcribed to cDNA (Quantitect-Reverse transcription kit, Qiagen). Quantitative reverse transcription polymerase chain reaction (qRT-PCR) was performed by using BIO-RAD (CFX96 Touch) a RT-PCR detection system and SYBR Green master mix (Thermofisher). Expression of the gene of interest was normalized to the expression of either 18S or S6B4.

Reverse transcriptase multiplex ligation dependent probe amplification

RNA samples from CD34+ cells cultured under different conditions were obtained by a Quick RNA Micro Prep kit (Zymo Research). Reverse transcriptase multiplex ligation dependent probe amplification (RT-MLPA) was performed according to the manufacturer’s instructions (MRC Holland, R011-C1). The resultant amplicons were separated by capillary electrophoresis (ABI-3130xl Genetic Analyzer) and Sequence Pilot (JSI Medical Systems) was used for the analysis. The sum of all peaks was taken as 100%, and the values of the single peaks were normalized accordingly.

Xenotransplantation

All experiments were performed after approval from the local ethics committee and in compliance with German law. Rag2−/−, c−/− mice were kept under specific pathogen-free conditions. Newborn mice were sub-lethally irradiated with 2.5 Gy. Animals were sacrificed for analysis after 8 weeks.

Flow cytometry

Single cell suspensions obtained from colony-forming assays or hematopoietic organs from mice were surface stained with monoclonal antibodies: CD34 PE-Cy7(581), CD38 APC (HIT2), CD10 PE/APC (H10a), CD45RA PerCP-Cy5.5 (H100), CD90 APC-Cy7 (SE10), CD117 PE-Cy7 (104D2), CD71 APC (CY14), CD33 PE (WM53), CD14 APC (M5E2), CD115 BV421 (4D2-1E4), CD15 PerCP (W6D3), CD66b PerCP-Cy5.5 (G10F5), CD19 PE-Cy7 (HB19), IgM APC-Cy7 (MHM-88), CD45 Biotin (H130), CD45 PE-Cy7/V500 (30-F11) (Biolegend), and CD235a BV421 (HR2) (BD Biosciences). Streptavidin PerCP-Cy5.5/V450 (Biolegend) was used as a secondary antibody. BD LSRFortessa and FlowJo were used for flow cytometry and analyses, respectively. The gating strategy was published earlier.

Western blot

Purified proteins were size fractioned by 12% sodium dodecylsulfate polyacrylamide gel electrophoresis under reducing conditions and transferred onto polyvinylidene difluoride membranes. The antibodies were used: MCL-1 (D2W9E) rabbit monoclonal antibody, BCL- XL (54H6), BCL-2 (D17C4), BFL1/A1 (D1A1C), α/β-tubulin or β-actin (13E5) (all rabbit, Cell Signaling). A peroxidase-coupled goat anti-rabbit IgG secondary antibody was used (sc-2004, Santa Cruz).

Statistics

Statistical analyses were performed using the unpaired Mann-Whitney test in GraphPad Prism 7 software. P values less than 0.05 were considered statistically significant. Synergy of BCL-XL and MCL-1 inhibitors was calculated using the Bliss synergy score and the program SynergyFinder (https://synergyfinder.fimm.fi).

Results

MCL-1 knockdown sensitizes human CD34+ cells selectively to endoplasmic reticulum stress

Putative shRNA sequences specific for MCL-1 were tested in transduced HEK293T cells. For shRNA delivery into human CD34+ cells, a lentiviral system that allowed stable expression of the shRNA and GFP was used. Two different shRNA sequences, both binding to exon 3 of the human MCL-1 gene, were identified to reduce MCL-1 mRNA expression to 25-55% in HEK293T and by 41-55% in human cord blood-derived CD34+ HSPC (Figure 1A). Knockdown of the Luciferase (Luci) gene was used as a negative control. Efficient MCL-1 knockdown was confirmed on a protein level in HEK293T and CD34+ cells and showed, at least in CD34+ cells, no relevant differences between the two shRNA sequences used (Figure 1B, C). Transduction efficiency was similar for the different viruses (Online Supplementary Figure S1B) but CD34+ cells transduced with shRNA specific for MCL-1 showed increased apoptosis rates 24 h after transduction (Figure 1D). The surviving cells were cultured and treated for 24 h and 48 h with different cytotoxic drugs including the DNA damaging agent etoposide, the mitotic spindle inhibitor taxol and different compounds inducing endoplasmic reticulum (ER) stress (i.e., tunicamycin, thapsigargin and brefeldin A) (Figure 1E, F). Interestingly, MCL-1 inhibition selectively increased the sensitivity of CD34+ cells to ER stress (Figure 1E, F). While transduced GFP+ cells expressing MCL-1 shRNA were killed in a dose-dependent manner (Online Supplementary Figure S2A), GFP+ cells expressing Luci shRNA or non-transduced GFP+ cells did not undergo increased cell death upon ER stress (Online Supplementary Figure S2A, B). These findings indicate that MCL-1 inhibition increased cellular sensitivity in a stress-dependent and cell-intrinsic manner. Since knockdown of MCL-1 was not complete in CD34+ cells, we hypothesized that ER stress itself reduced MCL-1 levels, thereby leading to critical depletion of this protein once combined with the gene knockdown. Indeed, we found significant and dose-dependent downregulation of MCL-1 mRNA when CD34+ cells were treated with tunicamycin (Figure 1G). At the same time, tunicamycin led to downregulation of BCL-XL mRNA and upregulation of the BH3-only proteins PUMA and BMF (Online Supplementary Figure S2C). Upregulation of CHOP and PERK mRNA confirmed the presence of ER stress (Online Supplementary Figure S2D). In summary, ER stress shifts the BCL-2 equilibrium towards apoptosis, which is initiated once levels of
Figure 1. MCL-1 inhibition selectively sensitizes human CD34+ cells to endoplasmic reticulum stress. (A) HEK293T cells were transfected with plasmids expressing shRNA specific for Luciferease (shLuci) or human MCL-1 (shM#3 or shM#4). MCL-1 mRNA expression was determined in sorted GFP+ cells and normalized to the 36B4 reference gene. Bars represent mean ± standard error of mean (SEM); n=5 from five independent experiments. Human cord blood-derived CD34+ cells were transduced with the corresponding lentiviruses. GFP+ cells were sorted 24 h after transduction, and knockdown efficiency of MCL-1 was determined by quantitative reverse transcription polymerase chain reaction (qRT-PCR). mRNA expression was normalized to 18S expression. Bars represent mean ± SEM; n=2-4 from four independent experiments. Mann-Whitney test: *P<0.05. (B, C) MCL-1 protein levels were determined in HEK293T (B) and CD34+ (C) GFP+ cells. (D) Measurements of apoptosis in CD34+ cells 24 h after lentiviral transduction revealed that 14-19% of cells undergo apoptosis early after MCL-1 depletion. Bars represent mean ± SEM; n=4 from four independent experiments. (E, F) Transduced CD34+ cells (transduction efficiency 45-65%) were cultured either under optimal conditions (cytokines and 10% serum) or under conditions of stress; in the presence of serum but deprived of cytokines, etoposide (0.5 mg/mL), taxol (0.125 mg/mL), tunicamycin I and II (0.5 and 1 mg/mL, respectively), thapsigargin (3 μM) or brefeldin A (BFA; 0.5 mg/mL). Apoptosis in GFP+ cells was determined by flow cytometry using annexin V and 7-AAD staining 24 h (E) and 48 h (F) later. Bars represent mean ± SEM; n=3-8 from eight independent experiments. Mann-Whitney test: *P<0.05, **P<0.01, ***P<0.001. (G) RNA was isolated from CD34+ cells treated with increasing concentrations of tunicamycin and used for qRT-PCR. MCL-1 mRNA expression was normalized to 18S expression. Bars represent mean ± SEM; n=4 from four independent experiments. Mann-Whitney test: *P<0.05.
either MCL-1 (shown here) or BCL-XL (documented earlier) are further reduced by RNA interference.

Human hematopoietic stem and progenitor cells show impaired colony formation and differentiation upon MCL-1 knockdown

To test the effect of MCL-1 knockdown on colony formation and differentiation of human CD34+ cells, we cultured 1,000 transduced and sorted GFP+ cells for 11 days in MethoCult medium containing the cytokines SCF, IL-3, IL-6, erythropoietin (EPO), granulocyte colony-stimulating factor (G-CSF), granulocyte-macrophage colony-stimulating factor (GM-CSF), insulin and transferrin. The number of colonies arising was significantly lower when MCL-1 expression was inhibited (Figure 2A). All colony types were affected, indicating that all multipotent and lineage-committed progenitor cell types were lost to a similar degree (Figure 2B). In addition, fewer cells could be harvested from plates (Figure 2C). Flow cytometry revealed that all cell types were reduced in number when MCL-1 was depleted (Online Supplementary Figure S3A). Since we noticed a relevant toxicity of the sorting procedure on lentivirally-transduced cells, we repeated the experiment using unsorted cells. With this approach, we could directly compare transduced GFP+ with untransduced GFP cells. Colony numbers and types were similar.

![Figure 2. MCL-1 is essential for all hematopoietic progenitor cells.](image)
in all groups (Figure 2E, F) and cell numbers were not consistently reduced when MCL-1 was downregulated (Figure 2G). However, while transduction rates were comparable (Online Supplementary Figure S3B), MCL-1-depleted GFP+ cells were selectively lost during the 11 days of MethoCult culture, indicating their selective disadvantage (Figure 2H, Online Supplementary Figure S3B). Loss of MCL-1 affected all analyzed immature and mature cell types in a similar manner, confirming that all progenitor cell types were dependent on MCL-1 expression (Figure 2H, Online Supplementary Figure S3C). To analyze whether MCL-1-depleted cells were lost immediately or progressively over time, we cultured them for only 5 days in MethoCult medium. At this early time point, only mild loss of GFP+ cells was observed, independently of the lentivirus used (Figure 3A, left). This indicates that progenitor cells became more susceptible to MCL-1 inhibition once they progressed in their differentiation process. Accordingly, immature CD34+ cells and specifically hematopoietic stem cells and multipotent progenitors were enriched in the first days of culture (Figure 3B, Online Supplementary Figure S3D, E). To inhibit differentiation but foster proliferation, CD34+ cells were cultured in the presence of the cytokines SCF (200 ng/mL), FLT3L (200 ng/mL), TPO (100 ng/mL) and IL-3 (20 ng/mL). CD34 and GFP expression was measured after 5 and 11 days. Under this condition, the percentage of CD34+ cells remained high (Figure 3B, right) and GFP+ cells were not depleted in a relevant manner (Figure 3A, right). To exclude that expression of shRNA and, consequently, MCL-1 knockdown was different in the two culture conditions, we measured MCL-1 mRNA after 5 days of culture. While we observed stable MCL-1 knockdown by shRNA #3, the shRNA #4 showed less consistent results with re-expression of MCL-1 mRNA (Online Supplementary Figure S3F). However, no difference in knockdown efficiency was observed between the two culture conditions indicating that the dependence of HSPC on MCL-1 did indeed change during proliferation and differentiation, respectively.

To understand why MCL-1 dependence was so different under the two culture conditions, we determined the composition of all BCL-2 proteins by RT-MLPA. As controls, we used freshly isolated CD34+ cells. Interestingly, both MCL-1 and BCL-XL were expressed at higher levels in cells stimulated to differentiate for 4 days (Figure 3C). Among the pro-apoptotic BCL-2 proteins, PUMA was highly upregulated under both culture conditions (i.e., differentiation and proliferation conditions) while BIM, BID and BAK1 were selectively upregulated under differentiation conditions (Figure 3D, E). Thus, it is possible that differentiation is associated with stronger pro-apoptotic signals that need to be counteracted by higher MCL-1 and BCL-XL levels. Based on its binding affinities, it is conceivable that MCL-1 expression is required to counteract BIM-mediated activation of BAK1.

MCL-1 inhibition severely restricts hematopoietic stem and progenitor cell engraftment in xenografted mice

In order to determine the effects of MCL-1 inhibition on the engraftment potential of human CD34+ HSPC, untransduced and transduced cells were intrahepatically transplanted into sublethally irradiated Rag2-/- mice. To reduce cell stress prior to transplantation, we waived the sorting procedure and transplanted GFP+ cells together with GFP- cells. After 8 weeks, xenografted mice were sacrificed and human engraftment was analyzed. Lentivirally transduced cells had a reduced potential to engraft but
there were no differences between the viruses used (Figure 4A). As known from this model system, most cells differentiated into CD19+ B cells while fewer CD33+ myeloid cells and almost no CD3+ T cells arose (Figure 4B and data not shown). Cells expressing the control shRNA (Luci) engrafted and contributed to all lineages (Figure 4C, D). In contrast, cells expressing shRNA specific for MCL-1 showed only very poor engraftment (Figure 4C). In line with the important role of MCL-1 for survival of immature progenitors with multipotent potential, all cell types found in the xenografted mice were equally affected (Figure 4D).

**MCL-1 inhibition limits the survival of immature but not mature hematopoietic cells**

To determine the effects of MCL-1 inhibition on more mature types of hematologic cells, we used the specific MCL-1 inhibitor S63845. First, we treated freshly isolated immature CD34+ and mature CD34+ cells with increasing doses of the inhibitor. While CD34+ cells were moderately sensitive when compared to cancer cell lines (e.g., IC50 in most multiple myeloma cell lines <0.1 μM),28 no apoptosis was induced after 24 h and 48 h in mature CD34+ blood cells even when very high doses of inhibitor were used (Figure 5A, B).

In a second approach, we let untreated CD34+ cells differentiate for 11 days in MethoCult medium. Cells were then isolated and put into stem cell medium containing 10% ES-FBS and cytokines (SCF, TPO, FLT3L, IL-3). Different concentrations of the MCL-1 inhibitor were added, and cell numbers were determined after 24 h. Again, differentiated CD34+ cells were much less sensitive than immature CD34+ cells (Figure 5C, D). As a consequence, only a mild and non-significant reduction in cell numbers was noted (Figure 5C) and both myeloid and erythroid cells were depleted only to a minor and non-significant degree (Figure 5D).

**Stem and progenitor cells from neonates and adults are equally sensitive to MCL-1 or BCL-XL inhibition**

Our experiments indicate a strong dependence of human HSPC on MCL-1 expression, which is not unexpected considering the high relevance of MCL-1 for survival of murine HSPC.25,26 However, other authors described an overall good tolerability of human HSPC to MCL-1 inhibitors.33-35 One reason for this discrepancy could be that we used CD34+ cells derived from cord blood while CD34+ cells derived from the bone marrow of aged persons were used in other studies.33 We therefore compared these two cell types with regards to protein levels of MCL-1 and other anti-apoptotic BCL-2 family members. Bone marrow CD34+ cells were obtained from patients with orthopedic problems (age range: 44 to 90 years). While MCL-1 and

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Figure 4. Human CD34+ cells lacking MCL-1 show poor engraftment in xenografted mice. (A-D) Lentivirally transduced or untransduced human hematopoietic stem and progenitor cells (HSPC) were transplanted intraparenchymally into newborn Rag2−/− mice after sub-lethal irradiation. Mice were sacrificed 8 weeks after transplantation and bone marrow (BM) and spleen populations were analyzed. By using antibodies specific for human or murine CD45, percentage human engraftment was determined (A). The various human hematopoietic populations were determined within the huCD45+ cells using flow cytometry (B). GFP expression was determined in huCD45+ cells (C) and in each of the subpopulations (D). Bars represent mean ± standard error of mean, n = 9-13 from eight independent experiments. Mann–Whitney test was performed; *P<0.05, **P<0.01.
BCL-XL levels were identical in both cell types and A1 was not expressed, BCL-2 levels were higher in adult HSPC than in HSPC from neonates (Figure 6A).

Next, we performed comparative functional studies using the MCL-1 inhibitor S63845, the BCL-XL inhibitor A-1155463 and the BCL-2 inhibitor ABT-199. By treating bone marrow immature CD34+ and differentiated CD34+ cells with the MCL-1 inhibitor for 24 h and 48 h, we obtained similar results as those with cord blood cells (compare Figure 6B with Figure 5A, B). Chemical inhibition of MCL-1 also confirmed our RNA interference experiments: when we added the inhibitor S63845 to cord blood CD34+ cells cultured in MethoCult medium, colony formation was impeded in a dose-dependent manner (compare Figure 6C with Figure 2). The numbers of immature cell types, including hematopoietic stem cells, multipotent progenitors and mixed lymphoid progenitors, were significantly reduced (Figure 6C, right panel). Similarly, all emerging erythroid and myeloid cells (Figure 6C, right panel) were depleted in a dose-dependent manner. Importantly, there was no difference in MCL-1 inhibitor sensitivity in CD34+ cells derived from cord blood (Figure 6C) or bone marrow (Figure 6D).

We have shown earlier that BCL-XL, too, is important for keeping human cord blood CD34+ cells alive, a finding that was unexpected considering its dispensable role for mouse HSPC. We now extended our studies to bone marrow-derived cells and showed that CD34+ cells were more sensitive than CD34 cells to the BCL-XL inhibitor A-1155463 (Figure 7A). We compared colony-forming potential CD34+ cells derived from cord blood or bone marrow and observed no difference between the two cell types: BCL-XL inhibition resulted in significant reduction of colony formation (Figure 7B) and cell numbers (Figure 7C), independently of the source of the human HSPC. Notably, effects on colony formation were less pronounced than those caused by MCL-1 inhibition. As documented earlier, immature CD34+ cells (Figure 7D) as well as mature erythroid cells (Figure 7E) were more severely affected than myeloid cells (Figure 7F).

Finally, the BCL-2 inhibitor ABT-199 did not negatively affect survival or colony formation of cord blood- or bone marrow-derived CD34+ cells, even when used at the high concentration of 1 μM (Online Supplementary Figure S4). This finding is consistent with the relatively mild myelosuppressive effects of venetoclax observed in clinical trials.

**Synthetic lethality of MCL-1 and BCL-XL inhibitors in human hematopoietic stem and progenitor cells**

We noted a striking functional homology of MCL-1 and BCL-XL for survival of human HSPC and concluded that combined inhibition of both anti-apoptotic proteins could result in complete depletion of colony-forming stem and progenitor cells. To test this, we treated cord blood-derived CD34+ cells with increasing doses of the

![Figure 5. Survival of mature hematopoietic cells is independent of MCL-1. (A-B) Freshly isolated cord blood was subjected to density gradient centrifugation and mononuclear cells were divided into CD34+ and CD34- cells using magnetic activated cell sorting technology. Both cell fractions were treated with the indicated concentrations of the MCL-1 inhibitor S63845. After 24 h (A) and 48 h (B), percentages of living cells were determined by flow cytometry using annexin V/7-AAD. Bars represent mean ± standard error mean (SEM); n=3-6 from six independent experiments. (C-D) CD34+ cells were differentiated in MethoCult culture. After 11 days, differentiated cells were isolated and treated with S63845 for 24 h. Total cell numbers (C) were determined and erythroid and myeloid cell populations (D) were analyzed by flow cytometry (n=5 from 5 independent experiments). Bars represent mean ± SEM. Mann-Whitney test: *P<0.05, **P<0.01, ***P<0.001.](insert figure here)
Figure 6. MCL-1 is essential for survival of both cord blood- and bone marrow-derived human CD34+ cells. (A) Protein levels of the anti-apoptotic BCL-2 family members MCL-1, BCL-XL, BCL-2 and BFL1/A1 were determined in CD34+ cells isolated from both cord blood and bone marrow. β-actin served as a loading control. (B) Freshly isolated bone marrow was subjected to density gradient centrifugation and mononuclear cells were divided into CD34+ and CD34- cells using magnetic activated cell sorting technology. Both cell fractions were treated with the indicated concentrations of the MCL-1 inhibitor S63845. After 24 h (upper panel) and 48 h (lower panel), percentages of living cells were determined by flow cytometry using annexin V/7-AAD. Bars represent mean ± standard error of mean (SEM); n=6-7 from seven independent experiments. (C-D) Human CD34+ cells isolated from either cord blood (C) or bone marrow (D) were differentiated in MethoCult medium containing 0.1 or 1 μM S63845. As controls, untreated and dimethylsulfoxide (DMSO)-treated cells were used (n=7 from seven independent experiments). After 11 days, total colony numbers (left) and total cell numbers (middle) were determined using light microscopy and hemocytometry, respectively. Different immature and differentiated cell types were determined by flow cytometry, and their absolute cell numbers were calculated (right). The following cell types were studied; HSC: hematopoietic stem cells (CD34+CD38CD45RACD90+), MPP: multipotent progenitors (CD34+38–CD45RA–CD90–), MLP: mixed lymphoid progenitors (CD34+CD38–CD45RA+CD10+), GM: granulocytic– monocytic progenitors (CD34+CD33CD115+); M: monocytes (CD34+CD33CD14+CD115–), immE: immature erythrocytes (CD71HI CD235a–), matE: mature erythrocytes (CD71 CD235a+). Bars represent mean ± SEM. Mann-Whitney test: *P<0.05, **P<0.01, ***P<0.001, ****P<0.0001.
MCL-1 inhibitor S63845, together with increasing doses of the BCL-XL inhibitor A-1155463. Apoptosis induction was determined 24 h later (Figure 8A). Using SynergyFinder (https://synergyfinder.fimm.fi) a dose-response matrix was calculated (Figure 8B). The resulting Bliss score of 21.26 indicates strong synergy between the two inhibitors. Synthetic lethality was confirmed in colony-forming assays, both with cord blood- and bone marrow-derived CD34+ cells. Already at concentrations of 0.1 µM each, the drug combination resulted in a substantial loss of colony-forming cells (compare Figure 8C, D with Figures 6B, C and 7).

To determine the number of immature cells with self-renewal potential that survived BCL-XL and/or MCL-1 inhibition, we used 10,000 cells isolated from primary colonies for serial colony-forming assays. Interestingly, only BCL-XL inhibition in the first plating resulted in depletion of progenitor cells able to form colonies in the second plating. However, there was a synergistic effect when this BCL-XL inhibition was combined with MCL-1 inhibition (Online Supplementary Figure S5).

Discussion

Because of the narrow spectrum of cancer entities susceptible to venetoclax, specific MCL-1 and BCL-XL inhibitors are eagerly awaited by oncologists. However, observations made in genetically modified mice indicate that inhibition of MCL-1 or BCL-XL could have more severe side effects than BCL-2 inhibition. Mice deficient for either MCL-1 or BCL-XL have severe developmental phenotypes while BCL-2-deficient mice lack lymphocytes and melanocytes but are otherwise normal. Hematopoietic toxicity of anticancer drugs is responsible for most therapy-related morbidity and mortality and a
A common reason for treatment delays or dose reductions. We, therefore, consider it important to generate preclinical data to evaluate the hematotoxicity profile of such novel drugs. Recently, we described the detrimental effects of BCL-XL inhibition on human HSPC and erythroid progenitors.\textsuperscript{31}

Here, we dissected the effects of MCL-1 inhibition on different immature and mature types of hematologic cells. By using an RNA interference approach and the specific small molecule inhibitor S63845, we consistently found that MCL-1 expression is crucial for multipotent stem and progenitor cells, as well as for myeloid progenitors, while erythroid progenitors are less susceptible to MCL-1 inhibition. During later stages of blood cell differentiation, MCL-1 becomes dispensable for cell survival. Interestingly, we noted re-expression of MCL-1 mRNA after some days of culture, when shRNA \#4 was used, while the shRNA \#3 resulted in stable knockdown. Nevertheless, the resulting phenotypes were strikingly similar indicating that the loss of stem and multipotent progenitors occurs early after MCL-1 depletion and cannot be compensated by later MCL-1 re-expression.

Looking more closely at the stem and progenitor cell compartment, we noted that cells that specifically enter the differentiation process are highly dependent on MCL-1 expression, while proliferating CD34+ cells remain fairly resistant. What is the reason for this difference? It is possible that the differentiation process is associated with increased stress levels reflected by accumulation of activated BH3-only proteins. Alternatively, it is possible that the cytokines TPO, FLT3L, SCF and IL3, which induce cell proliferation and are used for CD34+ cell culture, not only induce proliferation but also confer CD34+ cells with survival signals, thereby rendering them independent of MCL-1 expression. Indeed, we have shown earlier that these cytokines induce BCL-XL mRNA upregulation and at the same time repress expression of the pro-apoptotic BCL-2 proteins BIM and Bmf.\textsuperscript{30} Also in this study, BIM mRNA levels were lower and BCL-XL mRNA levels higher when cells were cultured in the presence of TPO, FLT3L, SCF and IL-3. Delbridge et al.\textsuperscript{26} recently showed that MCL-1 expression in murine HSPC is critically required to counteract PUMA-induced apoptosis. While mice lacking only one Mcl-1 allele in the hematopoietic system rapidly succumbed to bone marrow failure, additional deletion of both Puma alleles rescued all animals.\textsuperscript{26} However, our in vitro studies showed strong upregulation of PUMA mRNA in human CD34+ cells irrespective of the
BCL-XL and MCL-1 inhibitors were successfully combined in multiple myeloma, melanoma, prostate cancer and multiple pediatric tumors. These observations make the combination of different BH3-mimetics very attractive. However, our results point strongly towards synthetic lethality also in healthy tissues. Specifically, combined BCL-XL and MCL-1 inhibition might be detrimental to healthy hematopoietic tissue. This fact should be kept in mind when new clinical trials are designed.

Most data available so far, including our own, were acquired either in vitro or in artificial mouse models. Clinical data gathered from already initiated phase I trials (e.g., NCT03218683, NCT02675452, NCT03465540, NCT02979366) will provide better insight into both the anticancer efficacy of MCL-1 inhibitors and their frequent side effects. Importantly, some trials using small molecule MCL-1 inhibitors were placed on hold by the FDA in September 2019 because of cardiac toxicity. No data on hematologic toxicities have been published yet. Based on the very promising preclinical data one could speculate that the overall benefit-to-risk profile of MCL-1 inhibitors will be favorable, especially for tumors otherwise refractory to chemotherapy. In case of severe irreversible hematopoietic damage created by MCL-1 inhibitors, these compounds could still be used within high-dose chemotherapy regimens given prior to autologous or allogeneic hematopoietic stem cell transplantation.

Disclosures
No conflicts of interest to disclose.

Contributions
SB, SA, JFO, EMD, CM, YW, JMW and VRM designed and performed experiments; SB, SA, JFO, EMD, YW and ME analyzed and interpreted data; SB, SA and ME wrote the manuscript; LK, HS and MK provided human samples.

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