Hypoxia Selects Bortezomib-Resistant Stem Cells of Chronic Myeloid Leukemia

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Abstract

We previously demonstrated that severe hypoxia inhibits growth of Chronic Myeloid Leukemia (CML) cells and selects stem cells where BCR/Abl protein is suppressed, although mRNA is not, so that hypoxia-selected stem cells, while remaining leukemic, are independent of BCR/Abl signaling and thereby refractory to Imatinib-mesylate. The main target of this study was to address the effects of the proteasome inhibitor Bortezomib (BZ) on the maintenance of stem or progenitor cells in hypoxic primary cultures (LC1), by determining the capacity of LC1 cells to repopulate normoxic secondary cultures (LC2) and the kinetics of this repopulation. Unselected K562 cells from day-2 hypoxic LC1 repopulated LC2 with rapid, progenitor-type kinetics; this repopulation was suppressed by BZ addition to LC1 at time 0, but completely resistant to day-1 BZ, indicating that progenitors require some time to adapt to stand hypoxia. K562 cells selected in hypoxic day-7 LC1 repopulated LC2 with stem-type kinetics, which was largely resistant to BZ added at either time 0 or day 1, indicating that hypoxia-selectable stem cells are BZ-resistant per se, i.e. before their selection. Furthermore, these cells were completely resistant to day-6 BZ, i.e. after selection. On the other hand, hypoxia-selected stem cells from CD34-positive cells of blast-crisis CML patients appeared completely resistant to either time 0 or day-1 BZ. To exploit in vitro the capacity of CML cells to adapt to hypoxia enabled to detect a subset of BZ-resistant leukemia stem cells, a finding of particular relevance in light of the fact that our experimental system mimics the physiologically hypoxic environment of bone marrow niches where leukemia stem cells most likely home and sustain minimal residual disease in vivo. This suggests the use of BZ as an enhanced strategy to control CML, in particular to prevent relapse of disease, to be considered with caution and to need further deepening.

Introduction

We previously demonstrated that severe hypoxia inhibits growth of leukemia cell populations and selects cells exhibiting properties of leukemia stem cells (LSC) [1–3]. Incubation of Chronic Myeloid Leukemia (CML) cells in hypoxia, including K562 as well as the corresponding primary cells (from blast-crisis patients), results in the complete suppression of BCR/Abl protein, but not mRNA, so that hypoxia-selected CML stem cells, while remaining genotypically leukemic, are phenotypically independent of BCR/Abl signaling and thereby refractory to the treatment with Imatinib-mesylate (IM) [1,3]. This is very well in keeping with the notions that IM, despite its impressive efficacy as first-line therapy for patients with chronic phase CML, induces a state of minimal residual disease (MRD), rather than cure, and that LSC are the source of MRD. On this basis, we proposed that LSC responsible for MRD of CML home in vivo within severely hypoxic areas of bone marrow [3] where normal hematopoietic stem cells physiologically reside (the hypoxic stem cell niches) [4–7]. Thus, strategies to target IM-refractory LSC of CML are worth being explored within this context, taking advantage of LSC selection in hypoxia prior to drug administration in vivo to mimic the impact of treatment on cells already adapted to home in hypoxic tissue areas in vivo.

CML treatment with Bortezomib (BZ), a specific and reversible inhibitor of proteasome activity which is licensed for clinical use in mantle cell lymphoma and multiple myeloma, is being proposed to target LSC [8], on the basis of data indicating that BCR/Abl expression results in increased proteasome activity and that proteasome inhibition is cytotoxic against CML cell lines [9]. The study reported here addressed the effects of BZ on K562 as well as primary CML cells in severe hypoxia, to assess the sensitivity to BZ of hypoxia-selected LSC independently of, and in comparison with, that of unselected cells.

The quality of LSC selection, as well as the effects of BZ on hypoxia-selected LSC, were determined by the Culture-Repopulating Ability (CRA) assay, a simple and effective method to test in vitro the effects of drugs on LSC. This assay measures the stem cell potential in growth-inhibitory “selection” primary cultures (LC1) by determining the capacity of cells at the end of LC1 to repopulate growth-permissive “expansion” secondary cultures (LC2) and evaluating the kinetics of this repopulation. Further advantage to our experimental strategy is given by the fact that hypoxia selects LSC from every leukemia cell population tested so...
The hypoxia-dependent protection from the effects of BZ was confirmed by the marked early activation of caspase-3 and induction of apoptosis which followed BZ addition at time 0, but not day 1 (Figure 2), suggesting that hypoxia interfered with BZ by preventing caspase-dependent apoptosis of cell bulk [10]. At later incubation times, as expected [2,3], hypoxia per se induced apoptosis in most cells.

Figure 3 shows that time-0 BZ markedly accelerated BCR/ABL protein suppression occurring in untreated hypoxic cultures [1,3], where BCR/ABL protein was still well expressed at day 2. BZ treatment at day 1 failed to determine such an effect, indicating that a one-day incubation in hypoxia protected BCR/ABL protein from BZ-induced suppression. When compared to time-0 BZ, day-1 BZ was not just ineffective, but actually delayed BCR/ABL protein suppression, indicating that hypoxia suppressed BCR/ABL protein at least in part via proteasoma.

The effects of BZ in hypoxia or normoxia on leukemia progenitor and stem cells were then determined by the CRA assay [1–3,11]. Cells treated or not with BZ in hypoxic LC1 were transferred at different times of incubation therein into BZ-free normoxic LC2 to determine the pattern of their repopulation. Cells replated from day-2 hypoxic LC1, where hypoxia-dependent cell selection had not occurred yet and BCR/ABL protein was still expressed [1,3], rapidly repopulated LC2, to peak at day 10, as an effect of BCR/ABL-dependent growth stimulation (Figure 4A). BZ addition to hypoxic LC1 at day 1 did not alter this kinetics significantly, in keeping with the fact that day-1 BZ did not suppress BCR/ABL protein (see Figure 3). BZ addition to hypoxic or normoxic LC1 at time 0 (or to normoxic LC1 at day 1; data not shown) abolished LC2 repopulation.

On the other hand, cell transfer to LC2 from day-7 LC1, i.e., following a one-log reduction of cell number in LC1 and the relative enrichment of BCR/ABL protein-negative cells (see Figures 1A and 3) [1,3], resulted in a delayed LC2 repopulation, which reached its peak at day 21 (Figure 4B). Such a kinetics typically reflects the content of transplanted LC1 cells with LSC [2,3], as well as normal hematopoietic stem cells [11,12]. LC2 repopulation was significantly reduced, but not abolished, by BZ addition to hypoxic LC1 at either time 0 or day 1. BZ addition to normoxic LC1 at time 0 (or day 1; data not shown) abolished LC2 repopulation. The identical effects of time-0 and day-1 BZ administered in hypoxia indicated that LSC are in part capable per se (i.e., before their adaptation to, and enrichment in, hypoxia) to stand BZ action. To exploit such a resistance, however, LSC are required to be maintained in hypoxia, as BZ addition to normoxic LC1 abolished LC2 repopulation by either day-7 or day-2 LC1 cells. On the other hand, when hypoxic LC1 were treated with BZ at day 6 (i.e., after cell adaptation to, and enrichment in, hypoxia), LC2 repopulation resulted completely drug-insensitive (Figure 5).

This indicated that LSC, once decisively selected in hypoxia, are completely resistant to BZ.

While we were completing our study, the effects of higher BZ doses, although for shorter times than in our experiments, were published [8]. When we tested, on this basis, the effects of the addition of 5–20 nM BZ to hypoxic LC1 at day 1, BZ was completely ineffective in either LC1 or LC2 (data not shown), indicating that a 24-hour pre-incubation in hypoxia confers resistant also to high-dose BZ.
Hypoxia- and Bortezomib-Resistant CML Stem Cells

**Figure 2. Induction of apoptosis by BZ in hypoxia.** (A) Total cell lysates in Laemmli buffer were subjected to immuno-blotting with the indicated antibodies. Anti-H4 was used to verify equalization of protein loading. One representative experiment out of 3 is shown. (B) Percentages of cells undergone “early” (annexin-V+/PI-) or “late” (annexin-V+/PI+) apoptosis, as determined by flow-cytometry. Values are means±S.E.M. of 3 independent experiments.

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**Figure 3. Effects of BZ on BCR/Abl protein expression in hypoxia.** Total cell lysates in Laemmli buffer were subjected to immuno-blotting with an anti-Abl antibody. Anti-vinculin and anti-ERK1/2 antibodies were used to verify equalization of protein loading. One representative experiment out of 3 is shown.

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The K562 data reported above were confirmed using CD34+ bone marrow cells explanted from blast-crisis CML patients. BZ enhanced the hypoxia-induced time-dependent reduction of the number of viable cells in LC1, irrespective of whether had been added at time 0 or day 1 (Figure 6A), the effect of time-0 BZ being perhaps faster (see day 2). CML cells from day-7 hypoxic LC1 repopulated normoxic LC2 reaching peak after 4 weeks of incubation. LC2 repopulation was not consistently affected by BZ treatment of LC1 at either time 0 or day 1 (Figure 6B). LC2 repopulation was paralleled by re-expression of BCR/Abl protein (data not shown), as expected [1,3]. Thus, primary CML cells were found to contain hypoxia-selectable LSC completely resistant to BZ.

Discussion

This study showed, in primary as well as K562 CML cells, differences with respect to response to BZ between cell bulk (sensitive) and a cell subset containing LSC selected following prolonged incubation in hypoxia (resistant). Two different ways for CML cells to adapt to hypoxia were also evidenced: a “rapid adaptation” (occurring within 1 day of incubation in hypoxia) of cell bulk emerged, capable to prevent BZ-induced caspase activation, apoptosis and BCR/Abl protein suppression, and the consequent destruction of culture. Such a rapid adaptation to hypoxia was not necessary for a minority of cells suitable to survive independently of BCR/Abl protein expression and thereby undergo selection through 7 days of incubation in hypoxia. These cells were in large part BZ-insensitive during their selection irrespective of time (time 0 or day 1) of drug addition (see Figure 4B), and completely insensitive to BZ treatment after cell selection in hypoxia (at day 6; Figure 5). Thus, a leukemia cell subset exists which is predisposed to exhibit a stem cell phenotype and to escape hypoxia-induced apoptosis, so that it can then undergo a relatively slow program of adaptation to hypoxia, which progressively consolidates BZ resistance.

The most relevant result of this study is that hypoxia-adapted LSC of blast-crisis CML are completely resistant to BZ. This resistance, if one assumes hypoxia-adapted LSC to be a subset of total LSC, is likely to explain why BZ was found to determine a marked decline of BCR/Abl-positive chronic-phase LSC, but not their eradication [8]. We believe the hypoxia-adapted LSC subset to be the main source of MRD, as it is adapted to home in the physiologically hypoxic stem cell niches of bone marrow [4–7,13]. Under this perspective, the more interesting results reported here are those obtained following BZ treatment at day 1 (as well as day 6) of incubation in hypoxia, as they mimic more realistically what...
occurs when the drug administered in vivo encounters LSC already placed in hypoxic tissue areas.

The demonstration that hypoxia-selected LSC of CML, which we previously showed to be refractory to IM [1,3], are also resistant to BZ needs to be discussed in view of the possible development of a BZ/IM combination protocol for treatment of CML. Our results indicated that there is no point in using BZ to target hypoxia-selected, BCR/Abl protein–negative LSC. On the other hand, we also showed (see above) that K562 cell bulk can acquire resistance to BZ via a rapid program of adaptation to hypoxia. As this occurs while cells are still IM-sensitive due to the maintained expression of BCR/Abl protein, there is no apparent advantage either for using BZ together with IM to target BCR/Abl protein–expressing cells. Taken together, our results suggest the use of BZ as an enhanced strategy to control CML to be considered with caution and to need further deepening, especially considering the known toxic effects of BZ, including myelosuppression [8].

Materials and Methods

Cells and culture conditions

K562 and primary CML cells were routinely cultured in RPMI 1640 medium supplemented with 10% foetal bovine serum, 50 units/ml penicillin and 50 μg/ml streptomycin (all from Euro-Clone, Paington, UK, E.U.). Primary CML cells were collected after informed consent to the use for basic in vitro research, according to the Helsinki declaration and following approval of the Ethics Committee, at the Division of Hematology of Universita` di Firenze. CD34-positive cells were cultured in the presence of Flt-3 ligand (50 ng/ml) TPO (20 ng/ml) and SCF/KL (50 ng/ml) in LC1 and of SCF (50 ng/ml), G-CSF (100 ng/ml), IL-6 (20 ng/ml) and IL3 (10 ng/ml) in LC2 (all from PeproTech, Rocky Hill, NJ, U.S.A.). Exponentially growing cells were plated at 3×10^5/ml and incubated in hypoxia (~0.1% O2) in a Concept 400 anaerobic incubator (Ruskinn Technology Ltd., Bridgend, UK, E.U.) or normoxia (21% O2), at 37°C in a water-saturated atmosphere containing 0.1% O2, 94.9% N2 and 5% CO2.

Reagents

Bortezomib (Velcade®, Millennium Pharmaceuticals, Cambridge, MA, U.S.A.) was added to cultures at 0.5 nM (cell lines and primary cells) at the indicated times.

Measurement of cell viability and apoptosis

Viable cells were counted in a hemocytometer by trypan blue exclusion. Apoptotic cells were detected by using a FACSCanto flow-cytometer (Becton & Dickinson, Franklin Lakes, NJ, U.S.A.), after staining with FITC-conjugated annexin V and propidium iodide (PI), using the Annexin V-fluos staining kit (Roche Diagnostics, Basel, Switzerland) and following manufacturer’s instructions. The percentages of annexin-V+ /PI- or annexin-V+/PI+ cells are considered to reflect “early” or “late” apoptosis.

The CRA assay.

This assay estimates the culture-repopulating power of normal [11,12] or leukemic [1–3] hematopoietic cells undergone a selection treatment (e.g. hypoxia) in a primary liquid culture (LC1) by means of their transfer in fresh medium to non-selective conditions (e.g. normoxia) in a growth-permissive secondary culture (LC2) and the measure of its repopulation following a further incubation therein. Cell subsets rescued from LC1 at different times repopulate LC2 with different kinetics, the time necessary to reach the peak of LC2 repopulation reflecting the hierarchical level of stem/progenitor cells enriched in LC1. The CRA assay is a non-clonogenic assay capable to reveal in vitro stem cells endowed with narrow-repopulating ability in vivo [11]. The adaptation of CRA assay to leukemia cell populations enabled to detect different subsets of leukemia stem or progenitor cells [1–3].

Figure 6. Effects of BZ on hypoxia-resistant primary CML cells. Immunomagnetically-selected CD34+ bone marrow cells from 3 blast-crisis CML patients were treated in hypoxic LC1 (established at 3×10^4 viable cells/ml) as indicated (A) and transferred at day 7 into normoxic LC2 (3×10^4 viable cells/ml) (B), where trypan blue-negative cells were counted at the indicated times of incubation. Values represent means±S.E.M. of data from 3 independent experiments.
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Protein separation and detection

Cells (5x10^6) were washed twice with ice-cold PBS containing Na2VO4 100 μM and solubilized by incubating for 10 min at 95°C in Laemmli buffer (Tris/HCl 62.5 mM, pH 6.8, 10% glycerol, 0.005% blue bromophenol, 2% SDS). Lysates were clarified by centrifugation (20000 g, 10 minutes, RT) and protein concentration in supernatants was determined by the BCA method. Aliquots (30 μg/sample) were boiled for 10 min in the presence of 100 mM 2-mercaptoethanol, subjected to SDS-PAGE in 7.6% polyacrylamide minigels and then transferred onto PVDF membranes (Millipore Corporate, Billerica, MA, U.S.A.), by electroblotting. Membranes were blocked in Odyssey Blocking Buffer (OBB)/PBS (1:1) for 1 hour at RT and incubated in a 1:1000 dilution of antibody in PBS 0.1% Tween (T-PBS)/OBB (1:1) overnight at 4°C. Primary (all rabbit) antibodies were: anti-c-Abl and anti-cleaved-caspase 3 (from Cell Signaling Technology, Danvers, MA, U.S.A.), anti-caspase 3 and anti-ERK-1/2 (from Santa Cruz Biotechnology, S. Cruz, CA, U.S.A.), anti-H4 (from Millipore, Billerica, MA, U.S.A.) and anti-vinculin (from Sigma-Aldrich®, St. Louis, MO, U.S.A.). Secondary anti-IgG antibodies were IRDye®800CW- or IRDye®680-conjugated (LI-COR® Biosciences, Lincoln, NE, U.S.A.). Antibody-coated protein bands were visualized by the Odyssey Infrared Imaging System Densitometry (LI-COR®).

Statistical Analysis

All experiments were performed in triplicate or higher numbers of repeats. Statistical significance was evaluated by Student’s t-test for paired samples; p values <.05 (two-sided) were considered statistically significant.

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Author Contributions

Conceived and designed the experiments: MT, SG, PDS. Performed the experiments: MT. Analyzed the data: MT SG VB ER PDS. Wrote the manuscript: PDS. Provided financial support, final editing and approval of manuscript: PDS.

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