Macrophage inflammatory protein-3α influences growth of K562 leukemia cells in co-culture with anticancer drug-pretreated HS-5 stromal cells

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Abstract

Stromal cell monolayers have been an important means of studying the regulation of hematopoiesis, because they produce cytokines. Cytosine arabinoside, vincristine, daunorubicin, and doxorubicin are common drugs for hematological cancer therapy, and they may have some effects on bone marrow stroma during chemotherapy. The aim of this study was to elucidate interactions between the bone marrow stroma microenvironment and leukemic cells after drug treatment. We tested the hypothesis that human HS-5 stromal cells, pretreated with anticancer drugs, affected the growth of leukemic K562 cells by changing the cytokines in the culture microenvironment. Thereafter, proliferation of K562 cells increased nearly 2.5-fold compared to the co-cultivation with drugs-pretreated HS-5 stromal cells and drugs-untreated HS-5 stromal cells. The results indicated that co-cultivation with HS-5 stromal cells pretreated with drugs caused significant K562 cell proliferation. Cytokines in the microenvironment were detected via the RayBio® Human Cytokine Antibody Array Membrane. The levels of the cytokines CKβ, IL-12, IL-13, IGFBP-2, MCP-1, MCP-3, MCP-4, MDC, MIP-1β and MIP-1α were decreased, with a particularly marked decrease in MIP-3α. In co-culture medium, there was a 20-fold decrease in MIP-3α in daunorubicin-pretreated HS-5 cells and at least a 3-fold decrease in Ara-C-pretreated cells. This indicated a significant effect of anticancer drugs on the stromal cell line. Using phosphorylated Erk and pRb proteins as cell proliferation markers, we found that phosphorylation of these markers in K562 cells was inhibited during co-cultivation with drug-pretreated stromal cells in MIP-3α-supplemented medium and restored by MIP-3α antibody supplement. In conclusion, anticancer drug pretreatment suppresses the negative control exerted by HS-5 cells on leukemic cell proliferation, via modulation of cytokines in the microenvironment, especially at the level of MIP-3α.

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1. Introduction

The growth and differentiation of most types of hematopoietic cells in vivo require direct contact with stem cells. The bone marrow provides the basic microenvironment for hematopoiesis, and stromal cells are the principal components of this microenvironment. In vitro studies have shown that stromal cells regulate hematopoiesis by secreting cytokines that are important for normal proliferation and differentiation. Recently, some data have suggested that bone marrow stromal cells are involved in the trafficking of malignant B cells within the bone marrow microenvironment by cell contact or production of cytokines. Thus, intercellular cross-talk with activating and suppressive cytokines reflects a common mechanism in the regulation of cell growth in the system (Jiang et al., 1998; Badillo and Flanke, 2006; Wu et al., 2005; Torok-Storb et al., 1999). In treatment with cytostatic drugs, toxicity for hematopoietic progenitor cells is a dose-limiting factor. This may increase the incidence of chemotherapeutic failure in leukemia patients. Stromal cell-based assays have been used to study the growth requirements of hematopoietic cells, since leukemic cells originate from their normal counterparts and also reside within the bone marrow microenvironment, and it is, therefore, likely that stromal cells influence the growth of leukemic cells. Basically, cytostatic drugs damage normal and tumor cells, and therefore, change the bone marrow microenvironment. Cordero et al. (2004) showed that development of the hematopoietic system is influenced by the intrinsic qualities of hematopoietic stem cells and their supportive...
stromal cells. This report also suggests that soluble substance(s) produced from stromal cells play a role in the growth of leukemic cells (Frikin et al., 1993; Aoyagi et al., 1996). In leukemia patients, anticancer drugs may affect bone marrow stromal cells by altering stromal cell secretory cytokines, which influence the leukemic cells. A previous study has shown that stromal elements are not transplantable; the capacity of the recipient’s stroma to support hematopoiesis following chemotherapy and bone marrow transplantation (BMT) is important (Simmons et al., 1987). It implies that the microenvironment of bone marrow is one of the critical factors for success in chemotherapy and BMT.

The HS-5 cell line is a transformed cell line HPV-16 E6/E7 derived from human bone marrow stromal cells, and it supports growth of hematopoietic progenitor cells (Roeklein and Torok-Storb, 1995). HS-5 has been reported to improve AML cell survival in vitro and attenuate chemotherapy-induced cell killing (Garrido et al., 2001). Observation of stromal cells pretreated with anticancer drugs shows that they sustain cell growth in vitro better than in vivo. We tested the influence of four anticancer drugs on cell growth by co-cultivation of the AML cell line (K562) with HS-5 stromal cells. Based on previous data, we hoped to clarify the influence of anticancer drugs on stromal cells, an influence which should be significant. The purpose of our study was to investigate the effect of cytosine arabinoside (Ara-C), daunorubicin (Dau), doxorubicin (Dox) and vincristine (Vin) on stromal cells (HS-5). The experiment was carried out in vitro by co-cultivation of drug-pretreated HS-5 cells with K562 cells, to assess the cell growth support of stromal cells.

2. Material and methods

2.1. Cells

The human myeloid leukemia K562 cell line which corresponds to AML cell line, derived from a chronic myeloid leukemia patient, was kindly provided by Dr. L.C. Yu of the Institute of Biochemical Sciences of National Taiwan University. The cell line was routinely cultured in RPMI 1640 medium supplemented with 10% newborn calf serum (NCS). The human stromal cell line HS-5 was a gift from Dr. Y.H. Huang at Taipei Medical University. The HS-5 cell line was maintained in RPMI 1640 containing 10% NCS and routinely used for the support of K562 leukemic cell differentiation.

2.2. Chemicals

Ara-C was purchased from David Bull Laboratories (DBL, USA). A stock solution of 100 mg/ml was made and stored at −20 °C. Daunorubicin, doxorubicin, vincristine, and MTT [3-(4,5-dimethylthiazolyl)-2,5-diphenyl-2H-tetrazolium bromide] were purchased from Sigma (MO, USA). All of the other chemicals were reagent grade. The Human MIP-1 alpha ELISA Kit was obtained from RayBiotech. Phosphoprotein 44/42 MAP kinase antibody (for Thr202/Tyr204, p-Erk) was obtained from Cell Signaling Technology (#9101). Phosphorylated Rb antibody (for phospho-S249) was obtained from abcam (ab7488). Anti-Erk 1 antibody was purchased from Santa Cruz Biotechnology (sc-93). Anti-human CCL20/MIP-3α antibody was purchased from R&D Systems (ab9349).

2.3. MTT assay for cell viability

MTT can be reduced to a blue formazan dye and thereby assesses cell viability by measuring mitochondrial function. HS-5 stromal cells were seeded into six-well plates and cells were incubated with various amounts of anticancer drugs. After incubation, cells were washed three times with PBS buffer containing 0.2% NCS and then incubated with 2.5 mg/ml MTT for 40 min at 37 °C. DMSO (0.5 ml) was then added to dissolve the resultant crystals for 30 min at 37 °C. OD570 was measured, and a reduction indicated loss of cell viability and growth.

2.4. Non-contact co-culture

Transwell clear plates (Corning, NY, USA) with 0.4 μm pore size polyester membrane were used for non-contact co-culture. Co-cultures were initiated by seeding K562 cells into the transwell insert on a monolayer of HS-5 cells (1 × 10⁵ cells) grown in six-well plates that contained 5% NCS in RPMI 1640 medium. All experiments used K562 cells at a density of 1 × 10⁴ cells/ml on the first day, and then changed half of the medium, which was finally made up to 5 ml with fresh medium on day 4. K562 cells were harvested after a total of 7 days by pipetting the medium that contained the cells to rule out the presence of stromal cells, and then centrifuging at 2000 g at 4 °C. Daunorubicin, doxorubicin, vincristine, and MTT solutions were used to dissolve the resultant crystals for 30 min at 37 °C. OD570 was measured, and a reduction indicated loss of cell viability and growth.

Fig. 1. MTT assay for HS-5 stromal cell viability. HS-5 cells (1 × 10⁵) were cultured in RPMI 1640 medium supplemented with 5% (v/v) NCS and various concentrations of anticancer drugs. After 48 h, cells were washed three times with PBS containing 0.2% NCS and incubated with 2.5 mg/ml MTT for 24 h at 37 °C in the dark. Before reading at OD570, the MTT solution was replaced with 0.5 ml DMSO for dissolving the MTT formazan for 30 min at 37 °C. Results are presented as the mean ± S.D. (n = 10, **p < 0.01, ***p < 0.001). Significantly different from corresponding control group is shown at p-value.
2.5. Cytokines assays in stromal conditioned medium (CM)

The levels of cytokines in CM were measured by a RayBio Human Cytokine Antibody Array V & 5.1 Map. This assay employs a qualitative Western screening technique. The standard array matrix consisted of an 11 × 8 dot grid on a 20 mm × 30 mm nitrocellulose membrane with 79 unique capture antibodies. Array kit included an instruction manual and with the biotinylated-antibodies solution and chemiluminescent substrate. The cytokine array membrane was incubated with 1 ml 7-day-co-culture medium or 7-day control medium for 2 h, and then the membrane was washed three times with washing buffer I for 5 min each, and then washing buffer II for 5 min each. Cytokine was detected using cytokine antibody for 1 h, and then followed by HRP-labeled streapavidin incubation for 1 h. The dilution used for each of these reagents was suggested by the instructions contained in the assay kit. Reactive spots were visualized by enhanced chemiluminescence (ECL) (Amersham Pharmacia Biotech UK Limited) with exposure to X-ray film. Estimation of change in these cytokines was based on the relative ratio to the control, using a non-linear dynamic analysis method with Phoretix™ 2D software.

2.6. ELISA

Cell culture medium was harvested after 7 days and measured by ELISA according to the manufacturer’s instructions (Ray Biotech, USA).

2.7. Western blot analysis

The dissolved proteins in the sample buffer were resolved by SDS-PAGE on a 12% acrylamide gel slab. Proteins were transferred onto a PVDF membrane by electrotransfer (17 V) at room temperature for 18 h. The transferred proteins were detected using HRP-conjugated antibody diluted to 1:1000, 1:3000 and 1:10000 for p-Erk, p-Rb and Erk-1, respectively. The reactive bands were visualized using an ECL kit and exposed on X-ray film.

2.8. Statistical analysis

The statistical analysis was conducted using one-way ANOVA with Dunnett’s post-test using GraphPad InStat version 1.00 for Windows (GraphPad Software, San Diego, CA, USA).

3. Results

3.1. Evaluation of cytotoxic effect of Ara-C on HS-5 cells

Before co-cultivation of K562 cells with drug-pretreated HS-5 cells, we investigated the effect of these drugs on HS-5 stromal cell viability, using the MTT assay. Based on the chemotherapy information, we used a test concentration lower than the level in human blood serum (Seymour et al., 1996; Holstein and Hohl, 2001). HS-5 cells were treated with various concentrations of each drug for 48 h. The data showed that cell viability was inversely proportional to anticancer drug concentration (Fig. 1). When the concentration of drugs reached a certain level, the viability of HS-5 cells was reduced by 50%, which suggested the cytotoxicity of these anticancer drugs on HS-5 cells. The LD_{50} dose of AraC, vincristine, daunorubicin and doxorubicin on HS-5 cells was 1.0, 0.01, 0.1 and 0.2 μM, respectively. Under such circumstances, we suppose that the microenvironment will be changed in half of the HS-5 cells, and secretion of cytokines will be stopped. Based on these data, we used the LD_{50} concentration to pretreat HS-5 cells before co-culturing with K562 cells. We aimed to establish whether the microenvironment of drug-damaged HS-5 cells had an effect on K562 cell growth.

3.2. Effects of microenvironment on cell proliferation

K562 cells undergo differentiation induced by various agents and their optimal microenvironment, including co-culture with stromal cells (Torok-Storb et al., 1999; Barak and Ben-Ishay, 1994). After undergoing cell differentiation, proliferation should be slowed down. Suppression of leukemic cell proliferation by stromal cells is well known. Since K562 cell differentiation is induced by co-culturing with HS-5 cells, we examined the effect of the microenvironment on cell growth. Fig. 2A shows the number of K562 cells cultured alone or co-cultured with drug-pretreated HS-5 cells. Cultured K562 cells alone would keep proliferation constantly. Co-cultivation of K562 cells with untreated HS-5 cells showed fewer cells than for K562 cells cultured alone or co-cultured with drug-pretreated HS-5 cells. In comparison with K562 cell proliferation in RPMI 1640 medium alone (Fig. 2A), co-culture with drug-pretreated HS-5 cells for 7 days maintained K562 cell proliferation. These data
Fig. 3. Conditioned media were detected with a cytokine array and accompanying kit reagents and protocol. The conditional media were collected after 7 days cultivation of K562 or HS-5 alone and with drug-pretreated HS-5 or untreated HS-5 cells. The assay is arranged in the form of an 11×8-lane grid with the configuration shown in (A). The results are shown in (B) and the detail described in the main text. Phoretix™ 2D software (non-linear dynamics) for spots comparison was used for analysis. The arrow indicates the MIP-3β/H9251 spot in array membrane.

RayBio® Human Cytokine Antibody Array 5.1 (Cat# H0109805)
RayBiotech, Inc.
proteins in the different CMs, which contained compounds secreted from stromal cells. In the array membrane, there were six identical positive control antibodies that contained a biotinylated protein standard (A1, B1, C1, D1, J8 and K8 in Fig. 3A) and three negative controls (E1, F1 and I8 in Fig. 3A). Fig. 3B depicts the results of a typical assay of a 1-ml CM sample using the array kit. Under these assay conditions, a reduced signal was detectable in cell-free culture medium that contained 5% NCS (Fig. 3B); an intense signal was only observed for HGF, with far less intense signals present for the other cytokines. No significant signals appeared in the K562 culture medium (Fig. 3B), which suggests that the membrane array kit was far too insensitive to be used for the K562 cells cultured alone. We detected three intense signals in the media from HS-5 stromal cells cultured alone and co-cultured with K562 cells, including MIP-3α, IL-6 and TIMP-2 (Fig. 3B). The change in cytokines in CM should result from an intercellular reaction between HS-5 stromal cells and K562 cells. When K562 cells were co-cultured with drug-pretreated HS-5 cells, the secreted molecules in the CM were changed in decreasing way (Fig. 3B). Estimation of the change in these cytokines was based on the relative ratio to the control, using the non-linear dynamic analysis method with Phoretix 2D software. In the CM of drug-pretreated HS-5 cells co-cultivated with K562 cells, the level of MIP-3α was significantly reduced. According to Fig. 1, the changes in cytokines in the CM may have resulted from the decrease in HS-5 cell viability. Although the data obtained were non-quantitative, at times we used them to obtain a rough estimate of the changes in some detectable cytokines.

3.4. MIP-3α with cell proliferation

Based on the data in Fig. 3, we attempted to characterize the real changes in MIP-3α levels in CM. Using anti-MIP-3α antibody; we measured MIP-3α in K562 cells cultured alone or in drug-pretreated co-cultured CM, using ELISA. Clearly, MIP-3α levels decreased in drug-treated HS-5 cell medium in a dose-dependent manner (Fig. 4). The amount of MIP-3α in CM of Ara-C-, vincristine-, daunorubicin-, or doxorubicin-pretreated HS-5 cells decreased to 30, 12.7, 5.8 and 12.7%, respectively, at higher concentrations, compared to that from untreated HS-5 cells. The results imply that MIP-3α plays an important role in K562 cell growth. They also indicated that the decrease in MIP-3α levels in CM resulted from HS-5 cell damage induced by the drugs.

The decrease in MIP-3α levels in CM provided a suitable microenvironment for K562 cell proliferation. To confirm whether MIP-3α is an important factor for inhibiting K562 cell proliferation, we used MIP-3α-supplemented medium to analyze the Rb and Erk phosphorylation pattern. Clearly, the Rb and Erk protein phosphorylation decreased after 3 days co-cultivation with HS-5 cells, and in 1 ng/ml MIP-3α-supplemented medium (Fig. 5). The results with MIP-3α-supplemented cultivation indicated that initiation of K562 cell proliferation did result from the presence of MIP-3α in the medium, and not from stromal cell death.

In our study, the doubling time of K562 cells was 24 h (data not shown). During this doubling process, the modification of proliferative molecules should progress; therefore, we chose 3 days cultivation for our phosphorylation assay. Upon adding various concentrations of MIP-3α antibody to K562 cells cultured alone with MIP-3α, or co-cultured with HS-5 cells for 3 days, the phosphoryla-
tion of Rb and Erk proteins was restored to 20–30%. Such a level of antibody supplement may not be sufficient to completely neutralize the effect of MIP-3α on K562 cell proliferation. Or, other minor factors in this microenvironment have synergistic effects on K562 cells. These data indicate that MIP-3α may not be a unique factor, but it is important for inhibition of K562 cell proliferation.

4. Discussion

Little attention has been paid to the bone marrow stromal damage that results from anticancer drugs during chemotherapy (Schwartz et al., 1998). Previous studies have shown that post-chemotherapy marrow stromal cells are deficient in supporting in vitro hematopoiesis, and have suggested that negative regulators induced by chemotherapy and cytokines may be the cause of this defect. In this study, we investigated the effects of stromal cells on growth of myeloid leukemia cell line K562. As a model for hematopoietic/stromal cell interaction, we used the stromal cell line HS-5, which has been shown to be capable of supporting hematopoiesis, and has recently been used for in vitro support of human progenitors (Torok-Storb et al., 1999; Goerner et al., 2000). Although our result did not confirm that HS-5 cells support myeloid cell differentiation, we found that K562 myeloid leukemia cells grow rapidly in co-cultivation with HS-5 cells pretreated with anticancer drugs. In general, increased differentiation of these cells leads to a halt in cell proliferation and vice versa (Steinman, 2002). This means that ongoing differentiation prevents proliferation. Our results indicated that the changes in diffusible molecules derived from HS-5 cells affected the proliferation of leukemic cells. We have shown above that four anticancer drugs reduced the activity of the HS-5 cell line and altered cytokine secretion. It may be assumed that cytokines secreted from stromal cells can cause defects in hematopoiesis or even prevent the differentiation of myeloid leukemic cells.

The bone marrow microenvironment is characterized by complex interaction between stromal cells, hematopoietic cells and cytokines. Bone marrow stromal cells have been shown to influence differentiation, growth and survival of normal hematopoietic cells through the secretion of a variety of cytokines (Bertrand et al., 2000). Using the cytokines array to analyze cytokine secretions from the HS-5 cell line, we identified cytokines that are potential targets of anticancer drugs. These cytokines might be functionally linked to their roles in K562 cell proliferation. Human hematopoietic cells can secrete numerous regulatory molecules that form the basis of intercellular cross-talk, networks and cell regulation in an autocrine and paracrine manner (Majka et al., 2001); stromal cells might be involved in intercellular cross-talk. Recent studies in mice have indicated that anticancer drugs, particularly at high doses, induce bone marrow stromal damage and/or dysfunction (Ben-Ishay and Barak, 2001). It has been shown that anticancer drugs affect cytokine secretion from stromal cells in vivo and have a suppression effect on hematopoiesis. In fact, our data showed that a large part of the secreted cytokines was decreased in CM from high-dose pretreated HS-5 cells. This suggests that anticancer drugs affect stromal cell activity by reducing cytokine secretion, or gene expression of HS-5 cells. The cytokines should have synergistic effects on cell differentiation and proliferation. It must be a complicated intercellular network, and further studies are needed to clarify this.

Chemokines are a large family of chemotactic cytokines that provide key signals for cell growth. The effect of MIP-3α on cell proliferation and differentiation depends on cell type (Lisignoli et al., 2007; Hausmann et al., 2005). HS-5 cells can secrete MIP-3α alone or in co-cultivation with K562 cells. After anticancer drug treatment, levels of MIP-3α in media were found to decrease to a low level, and K562 cell proliferation increased significantly in medium from co-cultured cells. Inhibition of K562 cell proliferation in MIP-3α-supplemented medium indicates that MIP-3α plays an important role in K562 cell growth. It also implies that the increase in K562 cell proliferation results from an absence of MIP-3α in the medium from co-cultured cells. Our data suggest that MIP-3α is involved in inhibition of K562 cell proliferation.

In summary, we showed that anticancer-drug-pretreated stromal cells have lost their ability to suppress proliferation of K562 cells, via a reduction in MIP-3α or change in microenvironment. The mechanism of the MIP-3α effect on K562 cells needs further study. However, the results imply that leukemia patients undergoing chemotherapy have deficient stromal cells, and a cytokine-deficient host microenvironment should be taken into account in cases of chemotherapeutic failure.

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References


