Effects of decitabine on allogeneic immune reactions of donor lymphocyte infusion via activation of dendritic cells

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Abstract

Background: Successful prevention of post-transplantation relapse after donor lymphocyte infusion (DLI) depends on its capability to mediate an effective graft-versus-leukemia (GVL) response while minimizing DLI-related toxicity, including graft-versus-host disease (GVHD).

Methods: We assessed the effects of decitabine (DEC), a hypomethylating agent, upon allogeneic immune reaction in a murine model of DLI.

Results: Significantly greater tumor growth retardation and survival prolongation occurred in mice administered with 1.0 mg/kg DEC for 5 days (DEC‑1.0) than in control or DEC‑0.1 mice. Upon prompt DEC and DLI co-administration, dendritic cells (DCs) were activated; DEC‑1.0/DLI induced severe GVHD, and survival was significantly lower than with DLI alone or DEC‑0.1/DLI treatments. IFN‑γ and CD28 levels were higher in splenic DCs of DEC‑1.0 mice than in those of control mice. Assessment of delayed DLI co-administration with DEC, when IFN‑γ levels were normalized to control levels, revealed that DEC‑1.0/DLI successfully facilitated tumor management without causing severe GVHD.

Conclusions: Our results suggest that DEC primes allogeneic immune reactions of DLI via DC activation, and GVHD and GVL effects are separable through optimal DLI timing based on DEC-induced increase in IFN‑γ expression levels.

Keywords: Donor lymphocyte infusion, Decitabine, Graft-versus-host disease, Graft-versus-leukemia effect, Dendritic cells
TKI administration are primarily attributable to its direct antineoplastic cytotoxic effects. HMAs such as 5-azacitidine or 5-aza-2′-deoxycytidine (DEC) can induce direct antitumor toxicities similar to TKIs [3]. In addition, they may also affect tumors by modulating gene expression in key cellular regulatory pathways [4] and exerting their effects on various immune and immune-related cells [5–8]. In an HSCT setting, the administration of 5-azacitidine affected both acute and chronic GVHD [7, 9] and separation of graft-versus-leukemia (GVL) effects from GVHD was also suggested [7]. Overall, these data suggest that HMA administration modulates allogeneic immune reactions; however, further research on its role in cellular immunotherapies including donor lymphocyte infusion (DLI) is needed.

DLI is usually used to prevent imminent relapse or treat early-stage relapse, with the aim of inducing allogeneic immune responses against minimally residual leukemic cells; however, its overall success rate is relatively low due to limited GVL effects or unwanted adverse events, including severe GVHD or marrow aplasia [10, 11]. As limited antitumor effects of conventional DLI are associated with immune tolerance in cancer patients, various strategies have been attempted, including the use of combinations of conventional or novel chemotherapeutic agents [12, 13], induction of tumor antigens or costimulatory molecules [14, 15], or infusion of in vitro-generated leukemia-specific cytotoxic T lymphocytes [16, 17]. Mechanistically, successful tumor targeting by DLI depends on successful tumor antigen presentation and activation of antigen-presenting cells, such as dendritic cells (DCs). The potential of HMAs to disrupt immune tolerance may be due to their up-regulation of HLA, tumor antigens, and costimulatory molecules and induction of T cell responses against tumor antigens [8, 18–21]. More importantly, recent studies have supported the use of HMAs in combination with DLI, since HMAs induce generation or activation of DCs [5, 22], which are primarily involved in the development of immune reactions following allo-HSCT [23].

In the present study, we hypothesized that the administration of DEC activates host DCs and that this DEC priming may enhance the antitumor effects of subsequent DLI. To test this hypothesis, a mouse HSCT model mimicking early-stage relapse, wherein DLI is usually attempted clinically, was selected. Furthermore, to assess the adverse effects of DEC on GVHD, a mouse model wherein DLI typically fails to induce clinical signs of GVHD was used.

Methods

Mice and cell line
Female C57BL/6 (B6, H-2b) and B6D2F1 (F1, H-2b/d) mice were purchased from Orient Bio (Seongnam, Korea) and maintained in our facilities under pathogen-free conditions. Animal care and experiments were conducted in accordance with the guidelines of our institutional animal care and use committee. Murine mastocytoma P815 cells (H-2d) were used to verify GVHD and GVL effects based on previous studies [24] and were obtained from the American Type Culture Collection (ATCC, Manassas, VA, USA) and cultured in accordance with their instructions.

Post-transplantation relapse mouse model
The early stage of post-transplantation relapse is characterized by a low tumor burden in mixed donor chimerism. Mixed chimeric status was modeled using a haploidentical (B6→F1) transplantation mouse model, receiving nonmyeloablative conditioning [25]. Briefly, an F-1 mouse was conditioned with 400 cGy of total body irradiation, and bone marrow cells at a density of 1 × 10^7 cells were administered intravenously from B6 mice. The early-stage relapse with low tumor burden was modeled via subcutaneous inoculation of P815 cells at a density of 1 × 10^6 cells 1 day before DEC administration to recipient mice.

DEC administration with or without DLI
To investigate the effects of DEC on tumor growth, allogeneic immune reaction, and resultant mouse survival, DEC at variable doses was intraperitoneally administered once daily, and its effects were compared with those in the controls. In the transplantation setting, mice were divided into two groups following DEC administration: DLI and non-DLI. For DLI, spleen cells (2 × 10^7) from B6 donors were resuspended in HBSS (Welgene, Daegu, Korea) supplemented with 2% FBS (GIBCO, Grand Island, NY, USA), washed twice, and injected into the tail vein of recipient mice at specified time points. Tumor growth was assessed every 3–4 days by measuring the largest perpendicular diameter with a caliper; thus, tumor sizes were recorded in terms of area (mm^2). Survival was monitored daily, and the degree of clinical GVHD was assessed every 3–4 days based on weight loss, posture, the occurrence of diarrhea, and skin lesions, such as alopecia and dermatitis [26]. For histological analysis, hematoxylin and eosin staining was performed for the liver, skin, and intestinal tissues. DEC was purchased from Sigma-Aldrich (St. Louis, MO, USA).
T and NK cell and cytokine assays
Frequencies of T cells (CD3+CD4+, CD3+CD8+, CD4+CD25+Foxp3+) and NK cells (CD3-NK1.1+B220+) were analyzed using the following antibodies (BD, Franklin Lakes, NJ, USA): APC-cy7-conjugated anti-CD3 (Clone SK7), PE-conjugated anti-CD4 (Clone RPA-T4), PerCP-conjugated anti-CD8 (Clone RPA-T8), PerCP 5.5-conjugated anti-CD25 (Clone PC61), APC-conjugated-Foxp3 (Clone 259D/C7), PE-conjugated anti-NK1.1 (Clone PK136), and PerCP-conjugated anti-B220 (Clone RA3-6B2). The estimation and analysis of T cells, NK cells, and cytokines were performed using Fortessa (BD Biosciences, San Jose, CA, USA) and FlowJo software (Tree Star, Ashland, OR, USA). T-cell proliferative capacities were assessed via flow cytometric analysis of carboxyfluorescein diacetate succinimidyl ester (Invitrogen, Carlsbad, CA, USA) in the mixed lymphocyte reaction (MLR) with irradiated (13 Gy) splenic DCs (1×10⁵ cells) as stimulators. For cytokine analyses, serial serum samples were collected and analyzed via cytometric Bead Array flex kits (BD Biosciences) containing IFN-γ, TNF-α, IL-2, IL-6, IL-10, and IL-17A.

DC preparation and phenotypical and functional analysis
Bone marrow-derived immature and mature DCs (imDCs and mDCs; im/mDCs) from F1 mice were generated in vitro, as described previously [27]. Briefly, imDCs were generated by culturing bone marrow cells in media containing granulocyte–macrophage colony-stimulating factor (GM-CSF; 10 ng/ml; Peprotech, Rocky Hill, NJ, USA) and IL-4 (10 ng/ml; Peprotech, Rocky Hill, NJ, USA) and major histocompatibility complex (MHC)-I and -II markers (BD). The bone marrow-derived DC function was analyzed via assessment of dextran (Sigma-Aldrich) uptake. Cytochalasin D was inhibited at 1 μg/ml (Sigma-Aldrich) to the media after 6 days. The DC preparation and phenotypical and functional analysis of carboxyfluorescein diacetate succinimidyl ester (Invitrogen, Carlsbad, CA, USA) in the mixed lymphocyte reaction (MLR) with irradiated (13 Gy) splenic DCs (1×10⁵ cells) as stimulators. For cytokine analyses, serial serum samples were collected and analyzed via cytometric Bead Array flex kits (BD Biosciences) containing IFN-γ, TNF-α, IL-2, IL-6, IL-10, and IL-17A.

Statistical analyses
Data are presented as mean±standard error of the mean. The Wilcoxon signed-rank test was performed to compare tumor sizes, body weights, T cell and NK cell numbers, and cytokine levels. Spearman's rank test was performed for correlation analysis. Survival curves were plotted using the Kaplan–Meier method, and differences were evaluated using the log-rank test. P values <0.05 were considered significant throughout.

Results
Selection of the optimal dose of DEC
We attempted to determine the appropriate dose of DEC for co-administration with DLI by assessing the antineoplastic effects in non-transplanted tumor-bearing mice and systemic toxicities in the transplantation model. First, F1 mice were subcutaneously inoculated with P815 (1×10⁶ cells), and a palpable mass was observed approximately 6 days after inoculation. On day 7, mice were divided into five groups, and daily doses of 0, 0.1, 0.5, 1.0, or 2.0 mg/kg body weight of DEC were administered intraperitoneally for 5 consecutive days, respectively. Tumor size reduction and survival prolongation after DEC treatment were observed at doses of 0.5 mg/kg or higher (Additional file 1: Figure S1a, b). DEC doses other than 2.0 mg/kg were then selected and administered on day 9 after transplantation to assess systemic toxicities in mice undergoing transplantation. No differences in body weight or survival of mice were observed among these three different doses (Additional file 1: Figure S1c, d), and hence, 0.1 and 1.0 mg/kg (DEC-0.1 and DEC-1.0, respectively) were selected to simplify subsequent experiments.

Table 1 Primers used for Real-time PCR

| Primer | Sequence (5’ → 3’) |
|--------|--------------------|
| CD28   | F                  |
| R      | TACTTCTGCAAATTGAGTTCATC |
| INF-gamma | F     |
| R      | GGGGATGTCAGTCCAGTAG |
| β-actin | F     |
| R      | CGACTCCTTTTTGGCTCTCAG |

Differential effects of DEC according to DLI
Figure 1a shows the experimental design of DEC administration with or without DLI in a mouse model of early-stage relapse in a mixed chimeric state. In the absence of DLI, DEC-1.0 group showed significantly greater survival
prolongation and tumor growth retardation than the control and DEC-0.1 groups (Fig. 1b, c). When DLI was combined with DEC, mice in the DEC-1.0/DLI group showed significantly shorter survival but greater tumor reduction compared with the DLI only or DEC-0.1/DLI groups (Fig. 1d, e). Similar to the DLI only group, delayed tumor growth was observed in the DEC-0.1/DLI group, which resulted in prolonged survival. Mice administered DEC-1.0/DLI showed a progressive decrease in body weight (Fig. 2a) and higher clinical GVHD score compared with mice administered DLI only or DEC-0.1/DLI (Fig. 2b).

Histological analyses also revealed that the pathological findings were compatible with those of GVHD and were most prominent in the DEC-1.0/DLI group (Fig. 2c). These data suggest that DEC-1.0 can enhance the allogeneic immune reaction of DLI, thus inducing severe GVHD and concomitant GVL effects.

Changes in immune cells and cytokines following DEC administration

To assess the effect of DEC on immune cells in mice undergoing transplantation, peripheral blood samples
were analyzed 1 day after DEC treatment and 14 days after DEC and DLI co-administration. DEC treatment alone significantly increased the frequency of CD3+CD4+T cells in mice on day 1, while decreasing that of regulatory T-cells; however, differences in the frequency of immune cells between the groups were not statistically significant. Two weeks after DEC and DLI co-administration, when severe GVHD occurred, we observed a significantly high frequency of CD3+CD8+T cells. Nonetheless, frequencies of NK cells were not affected by DEC treatment only or with DLI co-administration (Fig. 3a). Serum cytokine levels decreased 7 days after DEC administration; however, IFN-γ level was significantly increased for approximately 28 days in mice administered DEC-1.0/DLI and exhibiting GVHD (Fig. 3b).

**Phenotypical and functional changes in DCs by DEC administration**

Regarding the role of host DCs in acute GVHD [23], we speculated whether DEC-induced GVHD aggravation was associated with DC activation. First, we investigated the phenotypical changes in marrow-derived DCs, which were induced in vitro with DEC treatment. The expression of CD54, CD80, and CD86 was upregulated in both im/mDCs and that of CD40 and MHC class II was upregulated in imDCs; dextran uptake by imDCs was also enhanced by DEC administration. These findings suggest that DEC treatment functionally modified DCs (Fig. 4a, b). We then selected CD11c+splenic DCs from recipient mice on day 1 after DEC administration to induce an MLR with donor T cells. Carboxyfluorescein diacetate succinimidyl ester assay showed that CD3+ and CD3+CD4+T-cell proliferation induced upon exposure
Fig. 3  

a Changes in CD3+CD4+ , CD3+CD8+ , and CD4+CD25+FOXP3+T cells and NK cells and b various cytokines after decitabine administration with or without donor lymphocyte infusion. The experiment was conducted in 3 times.*P < 0.05, **P < 0.01, ***P < 0.001 for control vs. DEC-treated group
to splenic DCs from mice administered DEC-0.1 and DEC-1.0 was significantly higher than that induced by vehicle (Fig. 5a), suggesting that DEC induced DC activation in a dose-dependent manner.

**IFN-γ dynamics of DCs and influence on GVHD and GVL effects**

Next, we assessed the DEC-mediated molecular changes in DCs using a commercial kit containing 84 genes associated with DC function. CD11c+ splenic cells from day 1 after drug exposure were used, and upregulation of IFN-γ and CD28 expression was observed in splenic DCs from mice administered DEC-1.0 (Fig. 5b). We also analyzed IFN-γ and CD28 mRNA expression levels using real-time PCR and showed that their levels were significantly higher in splenic cells on day 1 after exposure to DEC-1.0 (Fig. 5c). However, the dynamics of IFN-γ and CD28 expression differed. Upon DEC exposure, IFN-γ expression levels increased and were normalized to control levels over time. Conversely, CD28 levels were persistently higher than the control values. Next, to assess the association between GVHD aggravation and IFN-γ levels in splenic DCs post-administration of DEC-1.0 and DLI, DLI was delayed until 8 days after DEC administration, when IFN-γ levels normalized to control values (Fig. 6a). Greater tumor size reduction and more prolonged survival were observed when DEC-1.0 was administered with DLI than in those groups where mice received only DLI or DEC-0.1 and DLI (Fig. 6b, c). We found that delayed DLI failed to induce severe GVHD, as evidenced by changes in body weight (Fig. 6d) and the clinical GVHD score (Fig. 6e).

**Discussion**

Although HMAs are currently being used to prevent or treat post-transplant relapse either by administering this drug alone [28–30] or in combination with DLI [31–33]; to the best of our knowledge, the mechanism-based roles of HMAs in DLI and their effects on GVHD/GVL have not been analyzed. The purposes of this study were to determine whether pretreatment with DEC enhances allogeneic immune reactions following DLI, whether DCs are associated with altered GVHD/GVL effects, and to evaluate the possibility of modulating immune reaction following DEC and DLI co-administration through analysis of the DEC-induced DC activation mechanism. The mouse bone marrow transplantation model was chosen because GVHD typically does not occur after DLI in this model, resulting in more accurate and reliable information regarding the influence of DEC on GVHD induction. Inadequate antigen presentation is an immune escape mechanism employed by malignant cells [34], and disrupting this immune tolerance is a central step for successful anti-cancer immunotherapy. Accordingly, DCs are crucial for DLI success in terms of their roles in T-cell priming and the development of an immune reaction [23]. Maturation of DCs into effective antigen-presenting cells is a complex process involving epigenetic cell fate control and functional modification of DCs by epigenetic drugs; in fact, decreased antigen-presenting activity of DCs by histone deacetylase inhibitors has been observed [35, 36]. Meanwhile, HMAs have been suggested to stimulate DC activity [22, 36, 37].

In this study, the immune stimulatory influence of DEC upon DCs was first evidenced through the maturation
and activation of DCs following in vitro drug treatment, characterized by higher expression levels of class II MHC molecules and co-stimulatory molecules such as CD80, CD86, and CD54, compared with those in the controls. Functional enhancement of DEC-treated DCs was then confirmed by dextran uptake and MLR; compared with the control, higher proliferation of CD4+ T cells was observed when splenic DCs from mice administered with DEC were used. In line with these observations, the T-cell proliferation pattern was indicative of the effects of DEC on DCs and subsequent immune reactions. High induction of CD4+ T cells soon after DEC administration, especially following DEC-1.0 treatment, was followed by a significant increase in CD8+ T cell levels in DEC/DLI groups at later stages than in the DLI-only group. Regarding the critical importance of CD4+ T cell in the induction and maintenance of tumor antigen-specific CD8+ T cells [38], we assumed that low antigen specificity of conventional polyclonal DLI could be overcome by DEC. Overall, these results suggest that enhanced DC function by DEC (relative to DLI-only group) is associated with significant tumor reduction and severe GVHD in mice administered with DEC-1.0/DLI. Finally, analysis of gene panels related to antigen presentation showed significant upregulation of CD28 and IFN-γ expression in CD11c+ splenic DCs from mice administered with DEC, coinciding with the role of IFN-γ in DC activation [22].

![Fig. 5](image_url)

**Fig. 5** Effects of decitabine (DEC)-treated dendritic cells (DCs) on T cell proliferation and associated molecules. **a** Splenic DCs isolated from recipient mice treated with or without DEC were co-cultured with donor T cells, and proliferation of CD3+, CD3+CD4+, and CD3+CD8+ T cells was assessed via the carboxyfluorescein diacetate succinimidyl ester method. **b** Changes in the expression of molecules following DEC administration assessed using a gene kit of 84 genes known to be involved in DC function, and **c** the dynamics of IFN-γ and CD28 expression over time analyzed by real-time polymerase chain reaction.
We also investigated the mechanisms underlying DEC-induced aggravation of allogeneic immune reactions other than DC activation by assessing the changes in cytokines and immune cells other than CD4+ and CD8+ T cells. Consistent with previous studies using 5-azacitidine [7, 39], our results showed that DEC downregulated the expression of inflammatory cytokines. Contrary to observations in previous studies where 5′-azacitidine was used [40–42], the regulatory T cell frequency was decreased by DEC in this study. Of note, differences in HMA type, immune cell analysis timing, and administration schedules could also affect the differential influences of HMAs on regulatory T cells. Although decrements in regulatory T cells might have contributed, DEC-induced DC activation played a dominant role in enhanced allogeneic immune reactions following combined treatment with DEC and DLI.

Our study showed that DEC induced DC activation and potentiated DLI-induced GVL effects as well as unwanted severe GVHD, emphasizing the need for strategies to separate these effects. Based on our observation that the IFN-γ levels increased by DEC returned to control levels over time (as seen in humans) [43], delayed DLI administration was performed on day 8 post-DEC. 

![Fig. 6](image_url)

Fig. 6  a After injecting C57BL6 mouse-derived bone marrow cells into irradiated BDF1 mice, P815 cells were transplanted subcutaneously 8 days later. Decitabine was injected on days 1–5 after tumor transplantation, and donor lymphocyte infusion (DLI) was performed the day after injection. Changes observed in b survival, c tumor size, d body weight, and e GVHD scoring depending on the delay in DLIs. The experiment was conducted in 3 times. *P < 0.05 for control vs. DEC-treated group.
administration. This resulted in reduced GVHD severity even when combined with a higher dose of DEC. However, we observed better anti-tumor activity and longer survival in mice receiving DEC and DLI than in those administered with DLI alone. These results suggest that bone marrow cells are more sensitive to GVHD than other target organs, and separation of GVHD and GVL effects is possible. Different cell number thresholds exist for GVHD and GVL effects following conventional DLI [44, 45]. Similarly, our results suggest that different thresholds may exist for DC activation status. In addition to assessing the optimum time interval between DEC and DLI, we also tested alternative approaches for the infusion of different cell numbers 1 day after DEC administration. However, infusion of a ten-fold lower number of lymphocytes failed to prevent severe GVHD (data not shown), which further suggested that DEC activation plays a greater role in the induction of DEC-induced DC activation-mediated GVHD than just the number of lymphocytes. Together, these findings suggest the possible use of DEC and DLI combination as a therapeutic tool, warranting further research geared toward clinical application. Based on these observations, we are planning to investigate the effects of IFN-γ or CD28 expression levels of DCs on the occurrence of GVHD or GVL effects following DLI in patients treated with HMA and DLI. Moreover, the optimal DEC dosing schedule for appropriate DC activation, the interval between DEC and DLI, and DC activation markers other than IFN-γ should be investigated.

Conclusions

In conclusion, DEC-induced activation of DCs was shown to facilitate the eradication of remnant tumor cells via GVL effects, and the activation status of DCs could be optimized by the time interval between DEC and DLI to help separate GVL effects from unwanted GVHD.

Supplementary information

Supplementary information accompanies this paper at https://doi.org/10.1186/s40164-020-00178-y.

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Authors' contributions

YJK, YRK, HJK, and WJS designed the study and collected and analyzed the data. YRK, HJK, JYL, KSP, SL, NGC, DCJ, CKM, and YJK conducted the search, collected the data, performed the analyses and wrote the manuscript. All authors critically revised the manuscript. All authors read and approved the final manuscript.

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Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no conflict of interest.

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Abbreviations

DLI: Donor lymphocyte infusion; GVL: Graft-versus-leukemia; DCs: Dendritic cells; DEC: Decitabine; GVHD: Graft-versus-host disease; HSC-T: Hematopoietic stem cell transplantation; TNs: Tyrosine kinase inhibitors; HMAs: Hypomethylating agents.

Additional file 1: Figure S1. Optimal decitabine (DEC) dose selection for co-administration with donor lymphocyte infusion. Effects of DEC on tumor growth and survival in a non-transplant setting. a tumor size and b survival. Systemic toxicities of DEC in the haploidentical (B6F1) transplantation mouse model. c survival and d body weight. The experiment was conducted in 2 times.
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