Novel TLR7 agonist stimulates activity of CIK/NK immunological effector cells to enhance antitumor cytotoxicity

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Abstract. Toll-like receptor (TLR) 7/8 agonists have been applied in combination with chemo-, radio- or immunotherapy for lymphoma, and used as topical drugs for the treatment of viral skin lesions and skin tumors. In the present study, the role of an adenine analog, 9-(4-carboxyphenyl)-8-hydroxy-2-(2-methoxyethoxy)-adenine [termed Gao Dong (GD)], a novel TLR7 agonist, in the activation of cytokine-induced killer/natural killer (CIK/NK) cells was determined. The results of the present study indicated that GD was able to activate CIK/NK cells. The proportion of GD-induced CD3+CD56+ CIK and CD3-CD56+ NK cells was ~4% higher respectively compared with the control. Notably, combination therapy with CIK/NK cells stimulated by GD, markedly suppressed the proliferation of the chronic myelogenous leukemia K562 cell line. Following GD treatment, the cytotoxicity improved by ~25 and 21% when the effector/target ratio was 20:1 and 10:1, respectively. The results of the present study suggested a novel protocol for CIK/NK cell proliferation and revealed that GD may serve as a potent innate and adaptive immunomodulator in immunocyte culture.

Introduction

Adoptive cells, including cytokine-induced killer (CIK) and natural killer (NK) cells, aimed at inducing or boosting efficient immune responses have exhibited efficacy and safety in clinical trials (1-6). A typical obstacle to the successful application of these treatments is the difficulty of generating clinically relevant numbers of CIK/NK cells (7). A previous study has demonstrated that immunomodulators may boost the number of immune effector cells and prime their activity efficiently (8). Toll-like receptor (TLR) 7 is expressed in T, CIK and NK cells, in addition to antigen-presenting cells (9). TLR-mediated combination therapies are becoming attractive approaches due to the low toxic side effects and high activity in physiological homeostasis that they exhibit. A phase II clinical study was established in which a TLR9 agonist (CpG ODN1018 ISS) was used with rituximab to treat follicular lymphoma (10). Systemic administration of a synthetic TLR7 agonist (R848) combined with radiation may prime a cytotoxic T cell response against lymphoma cells, and prime a memory immune response that may prevent recurrence of lymphoma (11). Enhanced antitumor efficacy was also identified when the TLR9 agonist (CpG ODN1826) was combined with radio- and chemotherapy (12). These previous studies led to the present study which aimed at determining the role of the TLR7 agonist in the activation of adoptive cells, including CIK and NK cells.

A protocol has been established previously to rapidly and reproducibly expand CIK cells in vitro from human peripheral blood (13-16). In the present study, 9-(4-carboxyphenyl)-8-hydroxy-2-(2-methoxyethoxy)-adenine (termed Gao Dong, GD) (17), a novel TLR7 agonist, was combined with the traditional protocol of culturing CIK cells to determine the role of GD in the activation of CIK/NK cells. The results of the present study demonstrated that GD may activate CIK/NK cells. Notably, the combination therapy with CIK/NK cells, stimulated by GD, markedly suppressed the proliferation of the chronic myelogenous leukemia K562 cell line. The results of the present study suggested a novel protocol for CIK/NK cell proliferation and indicated that GD may serve as a potent innate and adaptive immunomodulator in immunocyte culture. This novel combination therapy concept may be a solution to the difficulty of chemotherapy and adoptive immunotherapy.

Materials and methods

Cell culture media and reagents. RPMI-1640 serum-free medium (Gibco; Thermo Fisher Scientific, Inc., Waltham, MA, USA) was used with 10% fetal bovine serum (Gibco; Thermo
Fisher Scientific, Inc.). CIK/NK cells (1x10⁶ cells/ml, Pierce; Thermo Fisher Scientific, Inc.) were generated and cultured with 80 U/ml gentamycin. Human recombinant interleukin (IL)-2, ≥1,000 U/ml, was purchased from Beijing ShuangLu Pharmaceutical Co., Ltd. (Beijing, China). GD (Fig. 1), synthesized in our laboratory, was added to the cell culture medium at a concentration of 5 µM.

Cell culture and quantification. The control CIK cells were generated as follows: Human peripheral blood mononuclear cells (PBMCs) from three donors were isolated by density gradient centrifugation at 4°C at 700 x g for 20 min using Ficoll (Takeda Pharmaceutical Company, Ltd., Tokyo, Japan). The cells were grown in AIM-V serum-free medium (Gibco; Thermo Fisher Scientific, Inc.) which consisted of 5% autologous plasma and 80 U/ml gentamycin. A total of 1,000 U/ml human recombinant interferon-γ (Gene Tech Co., Ltd., Hong Kong, China) were added on day 0. After 24 h of incubation at 37°C, 50 ng/ml OKT3 antibody (cat. no. 555339) against cluster of differentiation (CD)3 (Ortho Biotech, Inc., Raritan, NJ, USA), 100 U/ml IL-1 (Genzyme, Cambridge, MA, USA) and 300 U/ml IL-2 (Genzyme) were added. Cells were incubated at 37°C in a humidified atmosphere containing 5% CO₂ and were subcultured every 3 days in fresh AIM-V with the addition of 5% auto serum and 1,000 U/ml IL-2 at 1x10⁶ cells/ml.

The harvested PBMCs from each donor were divided into three equally and treated as follows: One group was cultured following the protocol as aforementioned and termed ‘none’ (the control group); another received the addition of GD with IFN-γ on day 0 and was termed group ‘+GD’; and the third group received GD only and on day 0, was termed group ‘+GD-IFN-γ’.

Cell numbers were counted using the Cell Coulter Technique (Z2 Coulter; Coulter Electronics, Ltd., Luton, UK) on days 0, 3, 5, 7, 9 and 15. Cell viability was analyzed using the trypan blue dye exclusion assay at each time point. Trypan blue dye-exclusion assay was performed by adding 20 µl 0.4% dye solution, and live (unstained) and dead (stained) cells were counted under Leica inverted microscope DMi1 (x10 magnification; Leica Microsystems GmbH, Wetzlar, Germany). Five random microscopic fields were counted in each sample. A total of 300 cells were counted per sample. Non-viable cells were detected as those which took up dye. Written informed consent was obtained from all human donors of PBMCs, and the present study received ethical approval by Shenzhen University Health Science Center Medical Ethics Committee (Shenzhen, China).

Analyses of lymphocyte subsets. Cell cultures were prepared for flow cytometric phenotypic analysis and four-color fluorescence was performed according to standard procedures (16). Briefly, 10⁵ CIK cells derived from PBMCs of donors suspended in 50 µl PBS were stained with 10 µl fluorochrome-conjugated monoclonal antibodies (1:50; BD Biosciences, Franklin Lakes, NJ, USA) against CD3 (cat. no. 561809), CD4 (cat. no. 561841), CD8 (cat. no. 566451), all T-cell antigens and CD56 (cat. no. 557747), NK-cell antigen. Following the addition of the primary antibody, cells were incubated for 20 min at room temperature and subsequently washed with PBS. Phenotype analysis was carried out using an Accuri C6 flow cytometer (BD Biosciences). The data were analyzed using BD CellQuest Pro Software (version 5.1, BD Biosciences).

Cell-mediated cytotoxicity. Cells from the three donors were analyzed on day 15 following culture and the K562 cell line was used as the target. Cell viability of K562 cells was evaluated in vitro using a Cell Counting Kit-8 (CCK-8). K562 cells were seeded in triplicate in 96-well plates (2x10⁴ cells/well) and CIK/NK cells were added at ratios of effector to target of 5:1, 10:1 and 20:1. Following incubation for 18 h at 37°C in a humidified atmosphere containing 5% CO₂, 20 µl CCK-8 was added to each well prior to incubation for 4 h at 37°C. The absorbance of each well was determined with an ELISA reader (iMark; Bio-Rad Laboratories, Inc., Hercules, CA, USA) at a wavelength of 450 nm. Cytotoxicity (%) was calculated using the following equation:

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\text{Cytotoxicity} = \left(1 - \frac{A_{450}}{A_{450}} \right) \times 100
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Statistical analysis. Statistical significance was analyzed using Student's t-test using Prism software version 4.0c (GraphPad Software, Inc., La Jolla, CA, USA). P<0.05 was considered to indicate a statistically significant difference.

Results

Cell proliferation rates. After 15 days, the proliferation rates of cells were increased 127-fold (median; range, 45-178), 46-fold...
Effect of GD on the phenotype of CIK/NK cells. As presented in Figs. 3 and 4, the proportions of CD3^+CD56^+ CIK cells (Fig. 3A) and CD3^+CD8^+ T cells (Fig. 3B) increased over the 15-day period. However, for the CD3^+CD4^+ T cells (Fig. 3C) and CD3 CD56^+ NK cells (Fig. 3D), the proportions decreased over the 15-day period. Furthermore, the proportions of GD-induced CIK cells and NK cells were ~4% higher compared with the control (15.9±5.8 vs. 11.9±2.9 and 7.6±4.2 vs. 3.8±3.7, respectively; Fig. 3A and D; Fig. 4). Additionally, the proportions of GD-induced CD3^+CD4^+ T cells and CD3^+CD8^+ cells were ~9% and 3% lower compared with the control, respectively (Fig. 3B and C; Fig. 4). Notably, there were significant differences between GD treatment and none, or between GD-IFN-γ treatment and none at days 7 and 15, when effector cells (CIK/NK) and other cells were considered (Fig. 3E and F). Treatment with GD may be able to counteract the IFN-γ-stimulated function in CIK culture, which was confirmed by data collected from the ‘+GD-IFN-γ’ group (Figs. 3 and 4). A notable inter-donor difference was that the final proportions of CIK and NK cells yield were linearly dependent on their initial numbers (data not shown).

Enhancement of cytotoxic capacity of CIK/NK cells stimulated by GD. The killing rates were notably increased in the ‘+GD’ and ‘+GD-IFN-γ’ groups compared with the control group (P<0.05; Fig. 5). The killing rates for group ‘+GD’ vs. group ‘none’ were 98.9±2.4 vs. 74.4±12.0, 54.0±15.3 vs. 32.8±21.5 and 28.0±27.2 vs. 26.3±26.3, respectively. Killing rates of group ‘+GD-IFN-γ’ vs. group ‘none’ were 89.0±4.2 vs. 74.4±12.0, 48.7±16.6 vs. 32.8±21.5 and 34.6±26.8 vs. 26.3±26.3, at effector/target ratios of 20:1, 10:1 and 5:1, respectively. The results of the present study indicate that GD induced a marked cytotoxic effect of CIK/NK cells on K562 cells.

Discussion

CIK cells are a heterogeneous subset of lymphocytes generated by incubating PBMCs with various cytokines in vivo. CIK and NK cells in CIK culture systems are important antitumor
effectors in immunotherapy (18). Culture systems used to provide adequate numbers of immune cells have focused primarily on stimulation with more effective cytokine and monoclonal antibodies (19,20). To the best of our knowledge, existing methods used for expanding CIK cells were almost negative with regard to expanding NK cells, but provided a high ratio of CD3⁺CD8⁺ T cells (21). Excessive CD3⁺CD8⁺ T cell numbers are not beneficial for eliminating tumor cells; however, they increase the risk of graft-versus-host disease (22,23). Thus, a method in which CIK/NK cells expand synchronously and CD3⁺CD8⁺ T cells proliferation decreases, may be valuable, economical and practical for clinical application.

TLR7 agonists are effective immunomodulators as they directly promote the activation of T, CIK and NK cells, induce tumor cell apoptosis or sensitize tumor cells to be killed by cytotoxic T lymphocytes and NK cells (24). CIK/NK cells alone and in combination with other therapies, including TLR7/8 agonists and imidazoquinoline analogs, are proven to be useful and safe (10-12). In the present study, the activity of the TLR7 agonist GD for the proliferation of CIK was determined for the first time, to the best of
our knowledge. The results indicated that GD significantly promoted the proliferative capacity of CIK/NK and decreased the proliferative capacity of no-CIK/NK cells, including CD3+CD4+ and CD3+CD8+ T cells, compared with the control. Additionally, GD improved the cytotoxic effect of CIK/NK cells on K562 cells. Thus, positive T cell co-stimulation by TLR7/8 ligands is dependent on the cellular environment (25). TLR7/8 agonists have been demonstrated previously to markedly upregulate IFN-γ and IL-12 (26), thus it may be possible to decrease the amount of exogenous IFN-γ used in the cell culture, and consequently decrease the toxicity exhibited from exogenous IFN-γ. It is hypothesized that local expression of IFN-γ and IL-12 may be less harmful to the patient than systemic administration of high amounts of IFN-γ and IL-12. The results of the ‘4-GD-IFN-γ’ group indicated that the local IFN-γ is a component of exogenous IFN-γ and, notably, the amount of IFN-γ secreted by the cells appeared to be sufficient to increase proliferation of CIK/NK cells. IL-12 enhances efficacy and decreases the enrichment time in CIK immunotherapy (27). Imidazoquinoline analogs (TLR7/8 agonists) promotes the proliferative capacity of NK cells to enhance antitumor effects (8). The results of the present study indicated a marked increase in the cytotoxic activity at a low ratio of effector to target cells. This appears to be of major importance, as it is difficult to inject patient effector cells at a ratio of 40:1 of effector to target cells in a clinical situation.

The results of the present study have demonstrated for the first time, to the best of our knowledge, that a TLR7 purine agonist promoted the proliferative capacity of CIK/NK cells to enhance antitumor effects. Since GD is involved in immune responses in immunological cell culture in vitro, GD was separated in culture medium while cells were harvested, therefore GD-free cells were reinfused in vivo. This strategy may reduce the risk of toxic cytokine syndrome induced by TLR7 ligands if injected directly in vivo. Furthermore, the heterogeneous cultured cell subset rich in CIK/NK cells and a low proportion of CD3+CD8+ T cells may be valuable in attenuating side effects and may be economical and practical for clinical application. This novel combination therapy concept may be a solution to the problem of chemotherapy and adoptive immunotherapy.

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