Lenalidomide enhances antitumor functions of chimeric antigen receptor modified T cells

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**ABSTRACT**

Tumor immunotherapy based on the use of chimeric antigen receptor modified T cells (CAR T cells) is a promising approach for the treatment of refractory hematological malignancies. However, a robust response mediated by CAR T cells is observed only in a minority of patients and the expansion and persistence of CAR T cells in vivo is mostly unpredictable. Lenalidomide (LEN) is an immunomodulatory drug currently approved for the treatment of multiple myeloma (MM) and mantle cell lymphoma, while it is clinically tested in the therapy of diffuse large B-cell lymphoma of activated B cell immunophenotype. LEN was shown to increase antitumor immune responses at least partially by modulating the activity of E3 ubiquitin ligase Cereblon, which leads to increased ubiquitylation of Ikaros and Aiolos transcription factors, which in turn results in changed expression of various receptors on the surface of tumor cells. In order to enhance the effectiveness of CAR-based immunotherapy, we assessed the anti-lymphoma efficacy of LEN in combination with CAR19 T cells or CAR20 T cells in vitro and in vivo using various murine models of aggressive B-cell non-Hodgkin lymphomas (B-NHL). Immunodeficient NSG mice were transplanted with various human B-NHL cells followed by treatment with CAR19 or CAR20 T cells with or without LEN. Next, CAR19 T cells were subjected to series of tests in vitro to evaluate their response and signaling capacity following recognition of B cell in the presence or absence of LEN. Our data shows that LEN significantly enhances antitumor functions of CAR19 and CAR20 T cells in vivo. Additionally, it enhances production of interferon gamma by CAR19 T cells and augments cell signaling via CAR19 protein in T cells in vitro. Our data further suggests that LEN works through direct effects on T cells but not on B-NHL cells. The biochemical events underlying this costimulatory effect of LEN are currently being investigated. In summary, our data supports the use of LEN for augmentation of CAR-based immunotherapy in the clinical grounds.

**Introduction**

The adoptive immunotherapy with autologous T lymphocytes genetically modified \textit{ex vivo} to express artificial signaling molecule designated CARs represents a novel and promising treatment modality of cancer. So far, the most successful example of CAR-based immunotherapy achievements came from the treatment of patients with B-cell acute lymphoblastic leukemia and chronic lymphocytic leukemia (B-ALL, CLL).\textsuperscript{1} Successfully targeted antigens include CD19 and CD20 which are major B-cell surface antigens and are strongly expressed by malignant B cells. CARs typically encode an extracellular antibody-derived domain that binds to a surface antigen (CD19, CD20, etc.) linked with an intracellular signaling domain that mediates T-cell activation such as TCRγ chain and co-stimulatory domains from CD28 or 4–1BB intracellular chains. The signaling through CAR substitutes for the signaling through endogenous T-cell receptor and leads to a potent and swift cytotoxicity toward target T cells in non-HLA restricted manner.\textsuperscript{2} In principle, any surface antigen can be targeted with CAR. Up to now, a large number of CARs targeting diverse tumors have been developed and many clinical trials are ongoing.

Despite promising results, resistance to CAR-based immunotherapy is frequently seen.\textsuperscript{3} The most debated reasons for the observed resistance include a loss of the CAR-specific antigen or a limited proliferation of CAR T cells \textit{in vivo} as a result of their inefficient activation or even inhibition due to immunosuppressive microenvironment within the tumor stroma.\textsuperscript{4} Several new approaches that would enhance CAR-based therapy are currently being tested, including an introduction of additional motifs from various co-stimulatory molecules into the intracellular signaling chain of CAR, co-transduction of T cells with genes encoding for essential pro-survival T-cell cytokines, or selective modification of certain T-cell subsets (such as effector memory).\textsuperscript{5} Another strategy to improve clinical efficacy of CAR-based therapy is based on the targeted reversal of tumor stroma immunosuppressive activity by using different immunomodulatory compounds such as monoclonal antibodies (MAbs) that block particular inhibitory receptors (e.g. CTLA-4, PD-1, LAG-3),\textsuperscript{6} or small molecules belonging to the class of immunomodulatory agents (IMiDs), namely LEN.
LEN is an IMiD approved for the treatment of MM, mantle cell lymphoma and 5q-syndrome. It was demonstrated that LEN binds E3 ubiquitin ligase Cereblon and induces degradation of transcription factors Ikaros and Aiolos. It inhibits growth of malignant B cells, inhibits angiogenesis and augments antitumor T-cell responses. It has been reported that LEN triggers tyrosine phosphorylation of CD28 on T cells, followed by activation of nuclear factor kappa B. In addition, LEN modifies T-cell responses in vivo and leads to increased interleukin (IL)-2 production in both CD4+ and CD8+ T cells, induces the shift of T helper (Th) responses from Th2 to Th1, inhibits expansion of regulatory subset of T cells (Treg), and improves functioning of immunological synapses in follicular lymphoma and CLL.

In this study, we tested the immunoadjuvant properties of LEN in combination with CAR19 or CAR20 T cells in experimental therapy of aggressive B-cell lymphomas using various mouse xenograft models based on xenotransplantation of both B-NHL cell lines and primary lymphoma cells. Presented data shows that LEN augments activation of CAR19 T cells in vitro and significantly enhances antitumor functions of CAR19 and CAR20 T cells in vivo. Our data additionally demonstrates that LEN works through direct effects on T cells, but not on lymphoma B cell. In general, our data supports the testing of LEN for augmentation of CAR-based immunotherapy in the clinical grounds.

Results

LEN enhances IFNγ secretion and augments signaling in CAR-19 T-cells

The data in Fig. 1 demonstrates that co-incubation of CAR 19 T cells with Ramos cells (Burkitt lymphoma) in the presence of 10 MM LEN results in enhanced production of IFNγ. We also show that the response is specific to CD19 since CAR-negative T cells do not produce IFNγ in the presence of B cells. (right panel).

The activation of CAR19 T cells might be influenced by multiple ligands expressed by Ramos B cells. To minimize such interference we activated CAR19 T cells with antibodies to CAR (or TCR) in the presence of LEN in the absence of Ramos cells. The data in Fig. 2 shows that LEN significantly and dose-dependently enhanced production of IFNγ after stimulation by anti-CAR antibody. Stimulation via endogenous TCR was more efficient than stimulation via CAR19, and in this case LEN did not further augment response of T cells. This finding suggests that stimulation by CAR is suboptimal compared to stimulation via TCR and that such weak stimulation can be markedly augmented by LEN. Further increase of the response in case of strong T cells activation via endogenous TCR cannot be achieved by LEN.

The effect of LEN is mediated by direct costimulation of CAR-19 T cells and more effectively augments the response of weakly stimulated CAR-19 T cells

The data in Fig. 3 shows that LEN pretreatment of CAR19 T cells enhanced the production of IFNγ, while LEN pretreatment of B cell had no effect on IFNγ production. We hypothesized that LEN would enhance the response of weakly or sub-optimally activated T cells, i.e., those expressing low amounts of CAR19. To answer this question we measured by flow-cytometry the upregulation of activatory molecule CD69 on CAR-19 T cells together with the expression of CAR19 receptor to correlate the level of T cell activation with the amount of CAR19. Similarly as in the previous experiment, CAR19 T cells and Ramos B cells were pre-incubated with LEN. The expression of CAR (y-axis, left panel) was separated into three major subgroups: CAR-negative (C-N), CAR-low (C-L), and CAR-high (C-H) subgroups (the gating is shown in the left panel). The level...
of CD69 upregulation was then determined in these three subgroups (right panel), the numbers in histograms indicate the percentage of CD69-positive cells. The data shows that LEN pretreatment markedly enhanced upregulation of CD69 (32% vs. 20%) only in the C-L T cells. No upregulation of CD69 was detected in the C-N T cells, while the difference observed in the C-H T cells was insignificant.

**LEN induces Aiolos and Ikaros degradation in T cells**

The molecular mechanisms behind the effects of LEN are not exactly known, but it was recently demonstrated that LEN binds to E3 ubiquitin ligase Cereblon and induces ubiquitinylation and degradation of transcription factors Aiolos and Ikaros. This mechanism was first shown in MM B cells, but was not tested in T cells so far. The data in Fig. 3 shows that LEN indeed similarly induces degradation of Aiolos and Ikaros transcription factors in T cells.

**LEN increases ERK phosphorylation following CAR activation in T cells**

Our data so far demonstrated that LEN augmented CAR signaling resulting in increased production of effector cytokines such as IFNγ, and increased expression of activation markers such as CD69. Additionally, in this experiment we tested the phosphorylation status of ERK kinase following stimulation of T cells with anti-TCR, or anti-CAR antibody. The data in Fig. 3 shows that LEN pretreatment augments the phosphorylation of ERK following stimulation of CAR. LEN did not further increase the level of pERK after TCR activation suggesting that
supra-physiological stimulation such as with anti-TCR antibody is not further enhanced by LEN similarly as in Fig. 2.

**LEN enhances antitumor functions of CAR T cells in vivo**

To translate the *in vitro* data into *in vivo* settings, we tested the immunoadjuvant properties of LEN to antitumor functions of CAR19 or CAR20 T cells using immunodeficient NOD-SCID-gamma chain null mice (NSG mice) transplanted subcutaneously (SC) with human B-NHL cells. In the first series of experiments, we transplanted NSG mice with established B cell lines TMD8 (diffuse large B-cell lymphoma of activated B-cell immunophenotype, ABC-DLBCL) and Ramos (Burkitt lymphoma). Both cell lines grow readily *in vitro* and *in vivo* and form large tumors upon SC injection into NSG mice. In the subsequent series of experiments, we used primary lymphoma cell-based murine xenograft models developed in our laboratory designated NEMO (relapsed mantle cell lymphoma MCL), and KTC (refractory ABC-DLBCL).

It must be emphasized that both NEMO and KTC cells can only be propagated *in vivo* in NSG mice from primary to secondary recipients. 

In the experiment shown in Fig. 4, we transplanted NSG mice on day 1 with 5 million TMD8 cells. These cells are highly CD20 positive and only medium positive for CD19, therefore, we used CAR T cells specific to antigen CD20 in this experiment. On days 2 and 3 mice received intravenous (IV) injection of CAR20 T cells (5 million each) followed by daily intraperitoneal (IP) injection of LEN. Fourteen days later mice were euthanized, and tumors excised and analyzed as in the previous experiment. The data in the left panel shows differences in tumor weight between respective groups and similarly to the previous experiment indicates that the tumor progression was slowest in mice treated with the combination of CAR20 T cells and LEN. Interestingly, tumors obtained from these mice also demonstrated highest levels of infiltration with CD8(+)* T cells. (right panel).

The data in Fig. 5 shows results from the experiment, in which NSG mice received SC injection of 10 million Ramos B-cell followed with one dose of 5 million CAR19 T-cells. In previous experiments (not shown) we found out that Ramos cells are highly sensitive to CAR-mediated therapy and therefore mice received only one dose of CAR T cells. One group of CAR19—treated mice received daily injections of LEN similarly as in the previous experiments. After 3 weeks mice were sacrificed and tumors were analyzed. The results showed that the combination of CAR19 and LEN led to the best anti-lymphoma response, and that the tumors in this group contained significantly more CD8(+)* T cells compared to the tumors obtained from the mice treated with CAR19 T cells only.

The data in Fig. 6 shows results from the experiment, in which NSG mice received SC injection of 5 million NEMO cells followed by injection of 5 million CAR19 T cells 2 and 3 d later, half of these mice received daily LEN. Fourteen days later, mice were euthanized and the tumors were excised and analyzed. Our data showed that LEN markedly augmented CAR-19 mediated suppression of tumor growth *in vivo*. Interestingly, flow cytometry analysis uncovered that tumors in mice treated with CAR19 T cells plus LEN were infiltrated predominantly with CD8(+)CAR19 T cells in contrast to the tumors obtained from the mice treated with CAR19 T cells only, which were infiltrated predominantly with CD4(+) CAR19 T cells.

In the last experiment, we tested the efficacy of CAR therapy using a mouse model of primary cell-based ABC-DLBC (i.e., mice xenografted with KTC cells). As the growth of KTC tumors was completely suppressed when treatment with CAR19 T cells was initiated early after tumor inoculation, (Fig. 8A) we decided to modify the approach and delay the treatment initiation. The treatment consisted of three doses of 5 million. CAR19 T cells at days 10, 12 and 17. Two weeks after first injection of CAR19 T cells the mice were sacrificed, tumors excised and weighted. The data in Fig. 8B shows that the tumor growth was significantly suppressed in the group, which received CAR19 T cells plus LEN. In contrast to the previous experiments, we did not find any T cell infiltration in the tumors of any group. The potential explanation for this observation might be the delay in the treatment initiation—the tumors were approximately 0.5 cm in diameter, and the very large tumor mass (compared to numbers of injected CAR T cells) probably led to limited persistence of CAR 19 T cells within tumor stroma.
Discussion

Our data demonstrated that a co-administration of LEN, which is a clinically approved orally available immunomodulatory agent, provides a significant enhancement of CAR-based immunotherapy. LEN leads to stronger signaling via CAR, increased production of IFNγ and upregulation of CD69 by activated CAR T cells. At the subcellular level, we showed that LEN increased phosphorylation of ERK after activation via CAR and the effects of LEN were associated with downregulation of transcription factors Ikaros and Aiolos. We have performed several in vivo experiments using murine models of aggressive human B-NHL (based on xenotransplantation of both cell lines and primary lymphoma cells) and we confirmed that LEN significantly enhanced antitumor functions of CAR T cells in vivo resulting in increased suppression of tumor growth and enhanced infiltration of tumors by cytotoxic CD8+ T cells in most tested models. Our data suggests that that LEN worked through direct effects on CAR T cells by enhancing the cellular response after a weaker stimulus such as after CAR activation.

CAR-based tumor immunotherapy is currently under intensive clinical investigation, treatment induced long-term remission in patients with B cell acute and B cell chronic leukemia who were resistant to standard therapy including allogeneic bone marrow transplantation. Most frequently targeted antigen in these diseases is the CD19 since it is strongly expressed by all malignant B cells, other B cell antigens such as CD20 were also tested. CAR-based therapy is not limited only to B

Figure 5. LEN enhances antitumor response in vivo to TMD8 DLBCL cells. NSG mice were transplanted SC with 5 million DLBCL ABC cells TMD8 and then received two doses of 5 million. CAR20 T cells followed with daily IP injection of 10 Mg of LEN, 14 d later mice were sacrificed and the tumors were excised and weighted (A). In panel B we shown the infiltration of tumors by CD8+ T cells which was determined by flow cytometry in cell suspension prepared from excised tumors. This experiment was performed twice with similar results.

Figure 6. LEN enhances antitumor response in vivo to Ramos Burkitt lymphoma cells. NSG mice were transplanted SC with 5 million Ramos cells and then received one dose of 5 million CAR19 T cells followed with daily IP injection of LEN, 21 d later mice were sacrificed, the tumors were excised and weighted (A). In panel B is shown the infiltration of tumors by CD8+ T cells which was determined by flow cytometry in cell suspension prepared from excised tumors. This experiment was performed once.
cell malignancies, various surface tumor antigens are under pre-clinical and clinical investigation targeting many types of tumors.

The mechanisms of antitumor activity of LEN and other IMiDs are broad and may affect many cell types in the organism. Direct effects include anti-proliferative and pro-apoptotic effects especially on malignant plasma cells. Indirect effects are postulated to be mediated by alterations in tumor microenvironment. IMiDs block interaction of MM cells with bone marrow stromal cells through inhibition of expression of surface adhesion molecules and they inhibit formation of bone lesions in MM by directly inhibiting osteoclast maturation. The immunomodulatory effects are associated with non-specific activation of immune system, which then effectively slows tumor progression. Unanswered question, however, remains the true nature of immune system activation by IMiDs. Published data suggests that IMiDs (the most frequently studied compound being LEN) might directly modify phenotype of malignant B cell by reducing the expression of inhibitory ligands such as PD-L1 and thus increase the recognition of tumors cells by T cells. Besides effects on malignant B cell, IMiDs have direct

Figure 7. Lenalidomide enhances antitumor response in vivo to primary mantle cell lymphoma. (A) NSG mice received SC 5 million NEMO cells followed by IV injection of two doses of 5 million CAR19 T cells and daily IP injection of LEN, 14 d later mice were sacrificed and the tumors were excised weighted. (B) The infiltration of tumors by T cells was analyzed by flow cytometry in a cell suspension prepared from excised tumors using antibodies to CD4+, CD8+ and CAR, one representative mouse is shown. The experiment in Fig. 7 was performed once.
NSG mice were SC transplanted with 5 million DLBC cell KTC and left for 10 d to establish cca 0.5 cm tumors, during this period all mice were receiving daily injections LEN. After the tumors were established, mice received three doses of 5 million CAR19 T cells at days 10, 12 and 17 with, or without LEN. Two weeks after treatment with DLBC cell KTC followed with one doses of 5 million CAR19 T cells and daily IP injection LEN, 14 d later mice were sacrificed and weighted. (B) NSG mice were SC transplanted with 5 million DLBC cell KTC and left for 10 d to establish cca 0.5 cm tumors, during this period all mice were receiving daily injections LEN. After the tumors were established, mice received three doses of 5 million CAR19 T cells at days 10, 12 and 17 with, or without LEN. Two weeks after treatment with CAR19 T cells, mice were sacrificed and the size of tumors was determined. The experiment in Fig. 8 was performed once.

Our data shows that LEN increases infiltration of tumors with CD8+ T cells. It is possible that LEN promotes the expansion of CD8+ T cells over CD4+ T cells or reduces the activation-induced cell death (AICD) of activated CD8+ T cells. To investigate this we cultivated CAR T cells in the presence of LEN (up to 7 d) and we did not detect any significant enrichment of CD8+ CAR19 T cells over CD4+ T cells. Similarly, the sensitivity of CAR T cells to AICD in vitro was not affected by LEN regardless if they were CD4+ or CD8+ (data not shown). We have also analyzed the tumors for the presence of T-regulatory cells and we did not find any significant infiltration by Tregs in tumors regardless of LEN treatment. However, the behavior of CAR T cells in vivo following recognition of tumor antigen and subsequent activation might be different from what occurs in vitro. Since LEN provides additional costimulatory signals to T cells for example by mimicking CD28 activation, it is possible that tumor-specific CD8+ CAR T cells will survive longer within tumor stroma in the presence of LEN.

Another evidence showing direct effects of IMiDs on T cells was obtained in patients, who underwent allogeneic bone marrow transplantation for MM and were later on LEN treatment. These patients developed severe graft-versus-host-disease (GvHD) and LEN had to be withdrawn from therapy.19 In mouse models, LEN was shown to enhance the response to antitumor vaccination.20 The conclusion from the available data is therefore quite solid in the sense that IMiDs can stimulate antitumor immune reaction by providing activatory signals to tumor-specific T cells and by enhancing the recognition of malignant B cell by reducing expression of their immunoinhibitory receptors. The overall therapeutic effect of IMiDs thus appears to be a combination of direct anti-proliferative effects on tumor cells, indirect effects on tumor stroma, immunomodulatory effects leading to increased recognition of malignant B cells and enhanced response of antitumor T cells.

Our finding that LEN works via direct effects on T cells but not B cells is the result of in vitro observation and can be very hardly proved in vivo. However, this finding does not contradict any so far published data and we indeed agree that LEN has direct antitumor effects, which is demonstrated by inhibition of tumor growth by single-agent LEN (in the absence of CAR treatment, see Results). Our in vivo data suggests that CAR T cell specific and tumor cell specific effects of LEN are synergistic and complementary. The literature shows that direct immunomodulatory effects of LEN on T cells results in enhanced proliferation and stronger production of IL2, IFNγ and TNF-α in vitro.21 Also the exposure of primary CLL cells to LEN in vitro led to induction of costimulatory molecules including CD80, CD86 and FASL on tumor cells, restoring immunological synapse formation and improving autologous tumor cell recognition by T cells.17 We have analyzed the phenotype of B-NHL cell lines Ramos and TMD8 after in vitro treatment with LEN and we did not detect any reproducible changes in the expression of CD19 or CD20, and both cell lines were negative for the expression of PD-L1 or PD-L2 regardless of LEN treatment. In concordance with these findings, the viability and proliferation of B-NHL cell lines in vitro not affected by LEN.

The molecular mechanisms behind LEN actions are connected to the degradation of transcription factors Aiolos and Ikaros via E3 ubiquitin ligase Cereblon.7 Degradation of these substrates, which are negative regulators of IL2 expression in T cells results in enhanced production of IL2 and other cytokines known to regulate T cell function. This mechanism was first shown in malignant plasma cells and later in other B cell types and T cells as well.8 We have confirmed that LEN induced degradation of Ikaros and Aiolos in CAR T cells and besides...
already discussed effects, we have also found out that LEN leads to increased activation of ERK following CAR stimulation. This finding is another manifestation of increased T cell response after LEN since ERK pathway activity reflects the magnitude of T cell activation.22

Presented experiments were performed with CAR19 T-cells except experiment in Fig. 5, however, we have done several key experiments such as tests for IFNg production also with CAR20 T cells, the results were similar to the results obtained with CAR19 T cells and therefore are not shown. Next, our results suggest that the costimulatory effects of LEN might be related to the strength of receptor-mediated activation and become significant in sub-optimally stimulated CAR T cells. This is supported by in vitro experiments, which have shown that LEN more effectively enhanced upregulation of CD69 in T cells expressing low amounts of CAR-19 compared to T cells expressing high levels of CAR-19. The lack of enhancement of endogenous TCR signaling by LEN can be the result of the fact that LEN might not further enhance supra-physiological activation of T cells triggered by anti-CD3 antibody. This hypothesis fits with the current model of T cell activation, which integrates signals originating form TCR, costimulatory and cytokine receptors.23

In conclusion, the key message of our paper is that the antilymphoma efficacy of CAR-based immunotherapy can be robustly enhanced by concomitant application of immunomodulatory agent LEN. Such combined treatment might lead to a highly effective cancer immunotherapy.

Material and methods

Construction of recombinant scFv fragments

The coding sequences for VL and VH variable domains of anti-CD20 mAb MEM-97 and anti-CD19 mAbs B-D3 were obtained by RT-PCRs using total RNA from respective hybridoma cells. PCR products were sequenced and used as templates for reamplification with modified primers allowing assembly of the VL (EcoRV-Xhol) and VH (PstI-BstEII) coding DNA fragments into scFv molecule in the format (VH)-(Gly4Ser)4-(VL). The scFv protein was produced in bacteria and tested for binding to the surface of CD19 and CD20 positive cells by flow-cytometry (data not shown). Subsequently, the scFv segments were in silico assembled with the CAR sequence, codon optimized and the whole sequence was chemically synthesized (Origene) and cloned into pWPXLd vector (Addgene) via BamHI-EcoRI using standard molecular biology techniques.

Cells, media and antibodies

HEK and Ramos cell lines were obtained from ATCC,12 TMD8 was a kind gift of dr. Hernandez-Ilizariturry- its origin was authenticated in our laboratory by mutational analysis showing the presence of the specific mutation of CD79B gene (Y196H). The cell lines were grown in IMDM media supplemented with 10% FCS, antibiotics (penicillin, streptomycin) and were mycoplasma negative. Human blood was obtained from volunteers under approval of local ethical committee, samples were processed by Ficoll gradient centrifugation and isolated leukocytes were further used for lentiviral infections. Following fluorescein labeled antibodies were used: Alexa 647 goat antimouse (Jackson Immuno), pacific Blue anti-CD4+, PerCP anti-CD8+, APC anti-CD69, PE anti-CD19 and PE anti-CD20 (all Exbio), for stimulation and western blotting were used unla-

Cell activation, lenalidomide treatment

Goat anti-mouse polyclonal serum (Jackson Immuno), or anti-

Virus production and cell transduction

Lentiviruses (2nd generation vectors) were prepared by transfection the HEK cells with packaging plasmids pSPax and pVSVG (Addgene) and lentiviral vectors pWPXLd containing CAR insert using Metaphetene Easy reagent, supernatants were then used directly to infect primary human T cells. For infections, first, tissue culture plastic was coated with antibodies to CD3 and CD28, then ficoll-purified leukocytes were resuspended in LV sup. and spin-infected (90 min, 1500 g, RT) in the presence of 5 Mg/ul polybrene, the infection was repeated next day, after 3 d the cells were replated in new flasks without antibodies and were further grown in RPMI media in the presence of 100 U/mL of IL2 (Proleukin).

Interferon gamma assay

The concentration of IFNg was determined by ELISA assay kit (Bilegend), the absorbances were read with ELISA plate reader at wavelengths specified in the kit. For the assay, we usually incubated 10,000 T cells with 3,000 B cells, in 100 ul of media for 12 h without the presence of IL2, before analysis cells were freeze-thawed. The concentration of IFNg was calculated in Prism software according to the dilution of IFNg standard.

Flow cytometry

Cells (1 million) in 50 ML PBS with 1% FCS were incubated with fluorescently labeled antibodies at dilution specified by the manufacturer, usually for 30 min at 4°C, washed twice and analyzed with flow-cytometer (Canto, BD). Single cell suspension was prepared from tumors by cutting them into small pieces with a blade and gentle dissociation by pipetting.

Cytotoxic assay

First, Ramos B cells were labeled with CFSE (5 um for 5 min at RT in PBS), washed and co-incubated with T cells, usually 3 ×
10^5 T cells plus 1 × 10^4 Ramos cells in 200 μL media for 24 h, the viability of CFSE^+ cells was analyzed by flow cytometry after adding the DNA dye Hoechst.

**Western blotting**

Cells were lysed at concentration 20 million/mL in 1% SDS loading buffer, reduced with 1% DTT (Sigma) at 95°C for 3 min, sonicated and resolved by SDS-PAGE. Proteins were visualized after transfer onto PVDF membrane by immunoblotting.

**In vivo experiments and statistical analysis**

**In vivo** studies were approved by the institutional Animal Care and Use Committee. Immunodeﬁcient NOD.Cg-Prkdcscid Il2rgtm1Wjl/SzJ mice (referred to as NSG mice, Jackson Laboratory) were maintained in individually ventilated cages. Primary DLBCL based mouse model designated KTC (treatment refractory DLBCL of activated B-cell immunophenotype) was established in the same way as previously described for a primary MCL based mouse model designated NEMO. KTC or NEMO cells were chosen for **in vivo** experiments, because they represent lymphoma subtypes, for the therapy of which LEN is currently approved (MCL) or under advanced clinical testing (ABC-DLBCL).

NSG mice were kept in individually ventilated cages under SPF conditions, mice 6–12 weeks old of the same sex in a group of 5 to 6 animals were injected SC with indicated number of tumor cells in 0.2 mL of PBS. After specified period, mice received intravenously two doses of 5 × 10^6 CAR T cells, followed with daily LEN treatment (10 mg/kg in 0.1 mL PBS intraperitoneally). After indicated time, mice were sacrificed, tumors were excised and weighted, then the tumor tissue was minced and dissociated by pipetting and analyzed by flow cytometry. The differences between LEN-treated groups were analyzed by one-tailed unpaired t-test using the Graph Pad software. All animal work was performed under approved ethical guidelines.

**Disclosure of potential conflicts of interest**

No potential conflicts of interest were disclosed.

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