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REPORT



Immunotherapeutic targeting of LIGHT/LT β R/HVEM pathway fully recapitulates the reduced cytotoxic phenotype of LIGHT-deficient T cells

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ABSTRACT

Tumor necrosis factor (TNF)/TNF receptor (TNFR) superfamily members play essential roles in the development of the different phases of the immune response. Mouse LIGHT (TNFSF14) is a type II transmembrane protein with a C-terminus extracellular TNF homology domain (THD) that assembles in homotrimers and regulates the course of the immune responses by signaling through 2 receptors, the herpes virus entry mediator (HVEM, TNFSFR14) and the lymphotoxin β receptor (LT β R, TNFSFR3). LIGHT is a membrane-bound protein transiently expressed on activated T cells, natural killer (NK) cells and immature dendritic cells that can be proteolytically cleaved by a metalloprotease and released to the extracellular milieu. The immunotherapeutic potential of LIGHT blockade was evaluated in vivo. Administration of an antagonist of LIGHT interaction with its receptors attenuated the course of graftversus-host reaction and recapitulated the reduced cytotoxic activity of LIGHT-deficient T cells adoptively transferred into non-irradiated semiallogeneic recipients. The lack of LIGHT expression on donor T cells or blockade of LIGHT interaction with its receptors slowed down the rate of T cell proliferation and decreased the frequency of precursor alloreactive T cells, retarding T cell differentiation toward effector T cells. The blockade of LIGHT/LT β R/HVEM pathway was associated with delayed downregulation of interleukin-7R α and delayed upregulation of inducible costimulatory molecule expression on donor alloreactive CD8 T cells that are typical features of impaired T cell differentiation. These results expose the relevance of LIGHT/LT β R/HVEM interaction for the potential therapeutic control of the allogeneic immune responses mediated by alloreactive CD8 T cells that can contribute to prolong allograft survival.

Abbreviations: LIGHT, homologous to lymphotoxin, exhibits inducible expression and competes with HSV glycoprotein D for binding to herpesvirus entry mediator, a receptor expressed on T lymphocytes; mAbM, monoclonal antibody; CD, Cluster of differentiation; HSV, Herpesvirus; TNF, Tumor necrosis factor; TNFR, Tumor necrosis factor receptor; HVEM, Herpesvirus entry mediator; LT β R, Lymphotoxin β receptor; CRD, Cysteine-rich domain; APC, Antigen-presenting cells; CTL, Cytotoxic T lymphocyte; NK, Natural killer; WT, Wild type; KO, Knock-out; Mgfp, monster green fluorescent protein; eGFP, enhanced green fluorescent protein; APC, Allophycocyanin; PE, Phycoerythrin; MHC, Major Histocompatibility Complex; PMA, Phorbol myristate acetate; CFSE, Carboxyfluorescein succinimidyl ester; FF-LIGHT, Flag-Foldon-tagged soluble mouse LIGHT; HVEM-Ig: HVEM.mlgG_{2a}-Fc, Herpesvirus entry mediator bound to mouse IgG_{2a} Fc fragment; LT β R-Ig: LT β R.hulgG₁.Fc, Lymphotoxin beta receptor bound to human IgG1 Fc fragment

Introduction

The tumor necrosis factor (TNF)/TNF receptor (TNFR) superfamily members play essential roles in diverse immunological processes such as T cell activation, costimulation, clonal expansion and T cell differentiation toward effector T cells.^{1,2,3} The therapeutic intervention with biologics antagonizing the TNF/ TNFR interactions can theoretically influence CD4 and CD8 T cell activation, clonal expansion, survival and particularly the process of differentiation toward effector T cells. This converts ARTICLE HISTORY Received 29 September 2015

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these interacting pathways as susceptible targets for the modulation of T cell-mediated immune responses.^{4,5}

Mouse LIGHT (TNFSF14) is a type II transmembrane protein of 239 amino acids with a C-terminal extracellular TNF homology domain (THD) that assembles as homotrimers capable to interact with HVEM ⁶ and LT β R.^{7,8} There are 2 isoforms of mouse LIGHT produced by alternative splicing: an isoform without transmembrane domain that resides in the cytosol, and an isoform with a transmembrane domain that can be proteolytically processed at amino acid 84 to generate a soluble extracellular form of LIGHT.^{6,9} In humans, there is an additional binding partner of LIGHT, a soluble protein named DcR3/TR6 protein (TNFRF6B) that lacks of transmembrane domain.^{10,11}

Mouse LIGHT displays a pattern of expression mainly restricted to activated T cells, NK cells and bone marrow immature dendritic cells.^{6,12} Both LIGHT receptors, HVEM and LT β R, are expressed on haematopoietic and non-haematopoietic stromal cells, although the latter is not expressed on lymphoid cells.^{13,14} Whereas LIGHT/HVEM main functional activity is a cosignaling pathway in T cells, LIGHT/LT β R interaction seems to be more relevant in regulating stromal/antigenpresenting cell (APC)/T cell cross-talk.¹⁵

Preclinical studies in mouse models of disease are required to establish the proof of concept for the function of a target needed to start clinical studies in non-human primates and humans, which could then allow the use of new biologics for the treatment of immune-mediated diseases. However, high quality in vivo reagents to target the molecule of interest and its interactions are needed during the process. Our group has previously shown that antibody-mediated blockade of LIGHT/ LT β R interaction with a partial antagonist antibody of LIGHT/ LT β R interaction reduced short-term cytotoxic allogeneic responses, although it did not fully recapitulate the impaired cytotoxic response observed in LIGHT-deficient T cells.¹²

Based on evidence gained in the field of transplantation using soluble LIGHT receptors as fusion proteins, such as HVEM.Ig and $LT\beta R.Ig.^{16,17,18}$ and from the phenotype of LIGHT-deficient mice,^{19,12,17} we postulated that a complete blockade of LIGHT interaction with its receptors would contribute to achieve a more suitable pharmacological control of the allogeneic immune response. LIGHT blockade on T cells would impede its interaction with $LT\beta R$ or HVEM on dendritic cells, and therefore hinder their maturation, 19,20,21,22 as well as prevent T/T cell collaboration through LIGHT/HVEM interactions that would contribute to maintenance of T cell survival during T cell expansion and differentiation.^{23,24,25,26} To confirm this hypothesis, we characterized a set of anti-LIGHT antibodies raised in LIGHT-deficient mice and chose one that fully blocked the binding of soluble $LT\beta R$ or HVEM to membrane LIGHT.

We demonstrated that efficient blockade of both HVEM/ LIGHT and LT β R/LIGHT interactions attenuated the allogeneic immune response in a mouse model of graft-versus-host reaction and fully recapitulated the reduced cytotoxic phenotype of allogeneic LIGHT-deficient T cells. This study points to LIGHT as a suitable target for a better immunotherapeutic control of cytotoxic responses in transplantation.

Results

Clone 3D11 is a neutralizing anti-LIGHT antibody that efficiently blocks its receptor-binding site

To demonstrate the in vivo effects of LIGHT interaction with its receptors in the course of the allogeneic immune response, an antagonist mouse anti-mouse LIGHT monoclonal antibody (mAb; clone 3D11, mouse IgG_{2b} , k) was raised in LIGHT- deficient mice immunized with LIGHT-transduced cells. The use of LIGHT-deficient mice as recipients for the immunization with LIGHT-transduced cells was critical for obtaining mouse anti-mouse LIGHT antibodies. LIGHT-deficient mice are not tolerant to LIGHT protein and they therefore can recognize LIGHT as a foreign antigen and mount a humoral immune response against multiple antigenic determinants of the protein.

The 3D11 antibody selectively bound NIH 3T3 cells expressing a green fluorescent protein (GFP): LIGHT fusion protein, in which GFP was fused to the intracellular N-terminus of mouse LIGHT, but did not recognize control NIH 3T3 cells expressing GFP only (Fig. 1A, upper panel). The binding specificity was confirmed by pre-incubation of clone 3D11 with recombinant FF-LIGHT, which blocked the ability of 3D11 to bind LIGHTtransduced cells (Fig. 1A, lower panel).

We next evaluated whether the anti-LIGHT mAb would prevent the interaction of mouse LIGHT expressed on cells with its receptors HVEM and LT β R. The 3D11 antibody antagonized the binding of sLT β R.Ig and sHVEM.Ig to mouse LIGHT-transduced cells down to background staining levels, similar to those obtained with isotype-matched control immunoglobulins (Fig. 1B). The binding of anti-LIGHT mAb to LIGHT transduced cells was inhibited by FF-LIGHT at close to stoichiometry ratio assuming a monomeric antibody mass of 150 kDa and 2 recognition sites per antibody and 67.5 kDa for the FF-LIGHT trimer and 3 epitopes per molecule (at EC 50, 0.625 μ g of antibody neutralized 1 μ g FF-LIGHT) (Fig. 1C).

We also assessed the binding avidity of the Fc fragment of anti-mouse LIGHT antibody to the activating and inhibiting mouse Fc γ R by surface plasmon resonance. The avidity constant (K_D = K_d/Ka) for CD64 (Fc γ RI) and CD16.2 (Fc γ RIV) was 84 μ M and 9.9 μ M, whereas for CD16 (Fc γ RIII) and CD32 (Fc γ RIIB) was 0.41 μ M and 0.23 μ M, respectively (Fig. 1C). This indicates that anti-LIGHT antibody (clone 3D11) binds preferentially to activating Fc γ RIII rather than Fc γ RIV or Fc γ RI.

We conclude from these results that 3D11 recognizes specifically mouse LIGHT but also fully inhibits the interaction of LIGHT with its receptors (HVEM and $LT\beta R$).

LIGHT protein expression is transiently detected upon polyclonal activation of T cells

As it occurs for some other members of the TNF superfamily ligands such as CD40L or human LIGHT, its expression is only transient on activated T cells,^{13,27} and visualization of this transient expression in vitro requires the presence of the fluorochrome-labeled antibody throughout the course of activation.¹² In C57BL/6 splenocytes stimulated for 5 h with PMA plus ionomycin, LIGHT expression was specifically detected with anti-LIGHT (3D11) antibody (Fig. 2A), in line with results obtained previously with another anti-mouse LIGHT antibody.¹² Mouse LIGHT was only detected on activated CD4 and CD8 T cells, but not on resting T lymphocytes (Fig. 2A). As expected for a specific staining, pre-incubation of the antibody with a molar excess of Flag-Foldon LIGHT abolished the staining of activated CD4⁺ and CD8⁺ T cells, respectively (Fig. 2B).



Figure 1. Effective blockade of LIGHT/LT β **R and LIGHT/HVEM interactions using a mouse anti-mouse LIGHT monoclonal antibody.** 2.5 × 10⁵ NIH-3T3 cells transduced with GFP-tagged murine LIGHT (blue solid lines) or GFP-transduced NIH-3T3 cells (red solid lines) were incubated with a mouse anti-mouse LIGHT mAb 3D11. After an incubation step, antibody binding was revealed with biotinylated rat anti-mouse $|gG_{2b}$ followed by allophycocyanin-coupled streptavidin (A, upper panel). To further demonstrate the specificity of the anti-LIGHT mAb, 1 μ g/well of Hilyte 647-labeled anti-LIGHT mAb alone (blue solid line) or 1 μ g/well Hilyte 647-labeled anti-LIGHT mAb preincubated with 2 μ g/well FF-LIGHT fusion protein (red solid line) were added to LIGHT-GFP transduced NIH 3T3 cells (A, lower panel). (B) 2.5 × 10⁵ LIGHT-transduced NIH-3T3 cells were pre-incubated for 30 min at room temperature with anti-LIGHT mAb (clone 3D11, blue solid lines) or mouse $|gG_{2b}$ isotype control (red dotted lines). Then, cells were stained with $LT\beta$ R-Ig (upper panel) or HVEM-Ig (lower panel). Binding of $LT\beta$ R-Ig was revealed with biotinylated anti-hgG and binding of HVEM-Ig (was revealed with biotinylated anti-hgG and binding of HVEM-Ig (was revealed with biotinylated anti-hgG and binding of HVEM-Ig was revealed with biotinylated anti-hgG and binding of HVEM-Ig (was revealed with biotinylated anti-hgG and binding of HVEM-Ig (was revealed with biotinylated anti-hgG and binding of HVEM-Ig (upper panel). Binding of $LT\beta$ R-Ig was revealed with biotinylated anti-hgG and binding of HVEM-Ig (upper panel), black solid lines) to LIGHT-transduced NIH-3T3 cells (A, lower panel, black solid lines). The case interverse cence intensity (MFI) is indicated in each plot. (C) Serial dilutions of anti-LIGHT mAb (clone 3D11) were preincubated with a fixed amount of 1 μ g/well of soluble recombinant Fr control (red solid line) or FF-LIGHT protein (black solid line). Then, 1 × 10⁵ LIGHT-transduced NIH-3T3 were added to the reaction. After a wash

Antibody mediated blockade of the LIGHT/LT β R/HVEM pathway recapitulates the attenuated cytotoxic allogeneic response of LIGHT-deficient T cells

LIGHT is required for lymph node hypertrophy in response to antigen immunization,²⁸ for T cell differentiation toward effector T cells in the course of an allogeneic immune responses ^{19,29} and for anti-tumor immunity.^{18,30} Due to the lack of well-characterized anti-LIGHT antibodies, the therapeutic potential of modulating LIGHT has remained elusive because selective blockade of LIGHT cannot be properly achieved with soluble HVEM.Ig or LT β R.Ig fusion proteins that both bind several ligands.

With the novel anti-LIGHT antibody, we addressed in mice whether blockade of the LIGHT/LT β R/HVEM pathway could attenuate the course of graft-vs.-host reaction in a semiallogeneic adoptive transfer model of alloreactivity. To that aim, CB6F1 (F1) recipient mice were injected with either F1 splenocytes (control), or semiallogeneic B6 splenocytes (to induce graft-versus-host reaction), or LIGHT-deficient B6 splenocytes (to induce graft-vs.-host reaction in the absence of LIGHT on transferred cells). Mice adoptively transferred with



Figure 2. Mouse LIGHT is rapidly upregulated on activated T cells after polyclonal stimulation. (A) Naïve C57BL/6 splenocytes (2×10^5) were left untreated or were stimulated with PMA (100 ng/ml) and ionomycin (500 ng/ml) for 5 h. 1 μ g/well Hilyte-647-labeled anti mouse LIGHT (3D11) or Hylite-647-labeled isotype control mouse IgG_{2b} were added to cultures during incubation in a final volume of 250 μ l. A lineage cocktail to gate out CD19⁺ CD11c⁺ Ly6G⁺ cells was included in the staining. The expression of mouse LIGHT was then analyzed on live resting and activated CD4 and CD8 T cells. (B) The expression of mouse LIGHT on the surface of resting or PMA plus ionomycin activated CD4⁺ and CD8⁺ T cells was also determined with the anti-LIGHT antibody pre-incubated with Flag-Foldon LIGHT (FF-LIGHT) (2 μ g FF-LIGHT per well). The percentage of each population is represented in each quadrant. PMA: Phorbol Myristate Acetate. Iono: Ionomycin.

semiallogeneic B6 splenocytes were treated with 3D11 (mouse anti-mouse LIGHT, isotype IgG_{2b}) or with an isotype control mouse IgG_{2b} . Nineteen days after the adoptive transfer, at the peak of the acute phase of the disease, the absolute number of host cells residing in the thymus, bone marrow and spleen was evaluated as a readout measurement of graft-versus-host reaction severity.^{31,32} As expected, normal host cell numbers were found in F1 mice receiving F1 splenocytes, but almost all haematopoietic host cells were rejected 19 days after transfer of semiallogeneic B6 splenocytes. An equally efficient rejection of haematopoietic host cells was observed in the bone marrow, thymus (total and double-positive thymocytes) and spleen, i.e.,

in primary and secondary lymphoid tissues (Fig. 3A-D). Transfer of LIGHT-deficient B6 splenocytes still lead to host cell rejection, but with reduced severity compared to wild type (p < 0.005), and the same was true when endogenous LIGHT was blocked with 3D11 upon transfer of wild type B6 splenocytes into F1 recipients (Fig. 3). No significant differences were found between the latter 2 conditions.

These results suggest that LIGHT blockade recapitulates the attenuated cytotoxic phenotype of LIGHT-deficient alloreactive T cells in F1 recipients, pointing to the relevance of LIGHT for the control of cytotoxic responses during the course of the allogeneic immune responses.



Figure 3. LIGHT blockade attenuated the course of graft-versus-host reaction in a mouse model of alloreactivity. Semiallogeneic splenocytes from C57BL/6 (70×10^6) were intravenously transferred into non irradiated CB6F1 recipients, which were treated with 1 mg of isotype mouse IgG_{2b} control (red squares) or anti-LIGHT mAb (3D11, green triangles) at day 0. In a third experimental group, semiallogeneic splenocytes from LIGHT-deficient mice (70×10^6) were injected into CB6F1 mice (orange diamonds). The fourth group represents the syngeneic control group, in which 70×10^6 F1 splenocytes was injected into F1 recipients (blue circles). The absolute number of host bone marrow (A), thymocytes (B), double positive thymocytes (C) and splenocytes (D) was determined 19 days after the adoptive transfer. Data represent a pool of 3 independent experiments. Statistical significance is indicated as follows: *p < 0.05, **p < 0.005, **p < 0.0005, and ns, non-significant.

Blockade of LIGHT impairs allogeneic T cell proliferation almost as efficiently as its genetic ablation

To understand why LIGHT inhibition reduced the severity of the allogeneic cytotoxic response, we monitored the proliferative capacity of carboxyfluorescein succinimidyl ester (CFSE)labeled donor alloreactive B6 splenocytes transferred into F1 recipient mice, in the presence or absence of genetic deficient LIGHT or immune therapeutic inhibition of LIGHT. Three days after the adoptive transfer of donor splenocytes, precursor frequencies (PF) and proliferation indexes (PI) were measured for CFSE-labeled donor CD4⁺ and CD8⁺ T cells (Fig. 4A). The PF and PtdIns were significantly reduced in the presence of the anti-LIGHT antagonist antibody compared to treatment with the control antibody (Fig. 4B). Moreover, the reduction achieved with the anti-LIGHT antibody was comparable to that obtained using LIGHT-deficient B6 donor cells (Fig. 4B)

These results indicate that the functional blockade of LIGHT retards clonal expansion of alloreactive T cells.

LIGHT inhibition delays differentiation of alloreactive CD8+ T cells toward effector cells

Since LIGHT inhibition reduced proliferation indexes and precursor frequencies of donor alloreactive $CD4^+$ and $CD8^+$ T cells, we investigated whether T cell differentiation toward effector T cells had also been hampered. For this purpose, we monitored expression levels of costimulatory (BTLA, HVEM,



Figure 4. The proliferation index and frequency of donor CD4⁺ and CD8⁺ alloreactive T cells is altered after LIGHT blockade or in LIGHT-deficient T cells. (A-B) 70×10⁶ of CFSE (carboxyfluorescein diacetate succinmidyl ester)-labeled B6 WT or CFSE-labeled B6 LIGHT-deficient splenocytes were adoptively transferred into non-irradiated F1 recipients and treated with 1 mg of isotype-control (mouse $IgG_{2,b}$) or mouse anti-mouse LIGHT (3D11) mAb at day 0. Three days later, the Proliferative Index (PI) and percentage of Precursor Frequency (PF) of donor alloreactive CD4⁺ and CD8⁺ T cells were determined using the ModFit LT software. Black line profile deconvoluted into cells that had divided once (green), twice (light violet), 3x (light blue), 4x (yellow), 5x (red), 6x (dark violet) and 7x or more (light green). X-axis represents CFSE fluorescence on a log scale and Y axis indicates cell counts (number of events). Data are representative of 2 independent experiments with 3 mice per group. Bars indicate mean \pm SEM, and uppaired *t* test was used to compare differences between groups. Statistical significance was indicated as follows: *p < 0.05, **p < 0.005, ***p < 0.0005, and ns, non-significant.

inducible costimulatory molecule (ICOS)) and differentiation (interleukin (IL)-7R α , KLRG-1) molecules on host and donor CD4⁺ and CD8⁺ T cells at 5 and 10 days after the semiallogeneic adoptive transfer of unfractionated parental B6 splenocytes into non-irradiated F1 recipient mice.

During a viral infection, CD8 T cell differentiation toward short-lived effector T cells and memory precursor cells can be tracked with IL-7Rα (CD127) and KLRG-1 surface markers.³³ IL-7R α is the receptor for the homeostatic cytokine IL-7 whereas KLRG-1 is a membrane glycoprotein with a C-type lectin domain and one immunoreceptor tyrosine-based inhibitory motif (ITIM) expressed in NK subsets, effector and memory T cells.^{33,34} KLRG-1 and CD127 (IL-7Ra) distinguish different stages of CD8 T cell differentiation at the peak of the response, just before the contraction phase, in which 95% of responding CD8 T cells are short-lived terminal effector cells (IL-7R α^{low} KLRG-1^{hi}).³³ KLRG-1 is, however, downregulated again during the contraction phase.³⁵ The expression of IL-7R α was down-regulated on donor alloreactive CD4⁺ T cells in all experimental groups at day 5 after the adoptive transfer. This was also true for donor $CD8^+$ T cells in the presence of LIGHT, but not for CD8⁺ T cells in conditions of genetic or pharmacologic impairment of LIGHT: in those cases, a fair percentage of IL-7R α -positive CD8⁺ T cells was still observed at day 5, but these were gone at day 10 post transfer (Fig. 5A-5B, upper panel), suggesting that differentiation toward effector CD8 T cells was delayed in the absence of LIGHT.

KLRG-1 expression was negative in all groups at day 5 postadoptive transfer. At day 10, a KLRG-1-positive population of alloreactive CD8⁺ T cells emerged that was more prominent in experimental groups where LIGHT contribution was genetically or pharmacologically compromised (WT to F1, isotype control, 15.02% \pm 0.16; WT to F1 anti-LIGHT 3D11; 25.96% \pm 2.70; LIGHT knockout (KO) to F1, isotype control; 24.36% \pm 0.96) (Fig. 5A-5B, lower panel). This could be interpreted as a delayed entry of CD8⁺ T cells in the contraction phase in the absence of LIGHT.

Unlike naïve T cells, which are ICOS negative, the expression of ICOS was upregulated at day 5 on most donor alloreactive CD8⁺ T cells in isotype-treated F1 mice. However, in the absence of LIGHT, there was still a fraction of ICOS⁻ donor CD8 T cells at day 5 (14.48% \pm 1.30 in isotype-treated controls vs. 29.63% \pm 1.61 and 30.81% \pm 2.12 in anti-LIGHT-treated and LIGHT-deficient donor cells, respectively), but not anymore at day 10 (Fig. 5C-5D), again pointing at delayed maturation of effector CD8⁺ T cells in the absence of LIGHT.

BTLA and HVEM expression were reciprocally regulated in donor $CD4^+$ and $CD8^+$ alloreactive T cells, after T cell activation and T cell expansion. Expression of BTLA is increased while that of HVEM was decreased. No significant changes in different surface markers were seen on donor alloreactive $CD4^+$ or $CD8^+$ T cells when experimental groups were compared to control (Fig. 5E).

Overall, these results suggest that LIGHT is required on allogeneic $CD8^+$ T cells to accelerate their differentiation.

Discussion

In the years to come, treatment of chronic and devastating diseases to which only palliative treatments are available will improve thanks to the substitution and complementation of chemical-based conventional therapy by biological-based



Figure 5. Altered expression of differentiation (IL-7R α and KLRG-1) and costimulatory (ICOS) surface markers is associated with delayed donor CD8 T cell differentiation in the absence of LIGHT. The pattern of expression of costimulatory and differentiation markers was assessed at days 5 and 10 after adoptive transfer of 70 × 10⁶ of donor syngeneic or allogeneic B6 WT or B6 LIGHT-deficient splenocytes into F1 recipients. Recipients were treated on the day of the adoptive transfer with 1 mg of isotype control mouse IgG_{2b} or anti-LIGHT mAb, clone 3D11. The expression of differentiation markers and costimulatory molecules was assessed by flow cytometry on host (blue lines) and donor (red solid lines) CD4⁺ and CD8⁺ T cells. The percentage of donor alloreactive CD8⁺ B6 WT or B6 LIGHT-deficient splenocytes that express IL7R α or KLRG-1 (A-B) was evaluated 5 and 10 days after the adoptive transfer into F1 recipients. The delayed maturation of effector CD8⁺ T cells (ICOS negative) in the absence of LIGHT was also depicted (C-D). (E) The expression of the costimulatory markers BTLA and HVEM was also assessed at days 5 and 10. Bars indicate mean ± SEM. Statistical significance was used: *p < 0.05, ***p < 0.005, ***p < 0.0005, and ns, non-significant. Data are representative of 2 independent experiments with 3 mice per group.

approaches, including mAbs.^{4,36,37} The understanding of how TNFSF and TNFRSF molecules contribute to the development of the immune response in different models of disease is essential for translation of the research findings into clinical practice. As members of the TNF/TNFRSF family control the absolute number of effector T cells and modulate the speed of the T cell differentiation process, they dictate the frequency of memory T cells that subsequently develop into long-lived memory T cells.^{4,36}

The process of T cell activation, costimulation, clonal expansion and differentiation toward effector T cells offers potential checkpoints for immune intervention. T cell activation with no or inefficient costimulation leads to functional inactivation, unresponsiveness or impaired T cell differentiation.^{38,39,40} Therapeutics targeting the CD28/CTLA4/CD80/CD86 costimulatory pathway, which is dominated by members of the immunoglobulin superfamily, have had a major effect on the control of allogeneic responses. CTLA-4.Ig blockade of CD80 and CD86 interaction with CD28 and antibody-mediated blockade of CD40/CD40L are the 2 most successful therapeutic achievements for targeting costimulation in the field of preclinical transplantation.^{41,42,43,44} CTLA4.Ig (belatacept) has already been introduced in the clinical setting as a maintenance treatment that efficiently prevents rejection while reducing the metabolic side effects of conventional drugs.⁴⁵ Treatment with CTLA4.Ig is particularly effective at low precursor frequency of alloreactive T cells that needs multiple rounds of division to reach a threshold effective to drive rejection.⁴⁶ However, CD8⁺ T cell-mediated rejection that is resistant to costimulation blockade remains an unbridgeable barrier, and thus alternative strategies aimed at targeting CD8 T cell differentiation that is independent of CD4 helper cells are required.^{47,48}

In a previous report with a partial antagonist of the LIGHT/ LT β R interaction, short-term cytotoxic response was attenuated, although to a lesser extent than that seen in LIGHT-deficient mice.¹² Using a murine model of graft-versus-host reaction, in which host hematopoiesis is attacked by donor cytotoxic CD8 T cells, we now provide evidence that a fully antagonist antibody of LIGHT/HVEM and LIGHT/LT β R pathway, can down-modulate T cell responses to the same extent as that seen in LIGHT-deficient T cells. Since in vitro binding of LT β R to LIGHT competitively inhibits HVEM recognition of LIGHT when both receptors are expressed on the same cell, the blockade of LT β R/LIGHT by the anti-LIGHT antibody is likely to be more critical than blockade of HVEM/LIGHT. This accounts for the fact that this novel antibody (clone 3D11) fully recapitulates the phenotype of LIGHT-deficient T cells while a previously reported antibody (rat IgG2a anti-mouse LIGHT, clone 10F12) did not.¹²

LIGHT has been proposed by several authors as a target for immunotherapy.^{4,5,26,49,50,51} Nevertheless the development of a specific anti-LIGHT reagent has been complicated until recently due to the difficulty of generating active recombinant mouse LIGHT with productive binding affinity for its receptor and functional biological activity.⁵² The description, functional evaluation and validation of this anti-LIGHT antibody opens up many possibilities to study the role of LIGHT and define its therapeutic potential in preclinical murine models of immunerelated diseases (transplantation, tumor immunity and autoimmune diseases),^{17,18,26,53,54,55} as well as in other pathologies in which LIGHT has been implicated, such as pulmonary fibrosis subsequent to chronic lung inflammation and idiopathic pulmonary fibrosis diseases,^{56,57} skin fibrosis ⁵⁸ and bone destruction through osteoclastogenesis.⁵⁹ This tool offers advantages over the classical approaches of using $LT\beta R.Ig$ recombinant fusion protein because $LT\beta R.Ig$ blocks $LT\alpha\beta/LT\beta R$ in addition to LIGHT/LTβR/HVEM.⁵¹ Administration of anti-LIGHT antibody in the above-mentioned disease models may help to clarify the contribution of LIGHT/LT β R/HVEM to the overall disease protection observed when using $LT\beta R.Ig$ fusion protein. Conclusions drawn from results obtained with $LT\beta R.Ig$ fusion proteins may need to be revisited because most studies utilized LT β R.Ig with a non-mutated Fc fragment of human IgG₁ that binds with relatively high affinity to mouse $Fc\gamma RIV$, the main receptor implicated in antibody-dependent cell-mediated cytotoxicity (ADCC),60,61 conferring this molecule with the potential to deplete cells expressing $LT\beta R$ ligands.

In general, the mouse IgG2a isotype is far more efficient than either mouse IgG1 or IgG2b isotypes at mediating depletion through ADCC. The reason for this difference is because mouse IgG2a binds in vitro to all activating $Fc\gamma Rs$ $(Fc\gamma RI > Fc\gamma RIV > Fc\gamma RIII)$, whereas mouse IgG2b binds mainly to $Fc\gamma RIII$ and to a lesser extent to $Fc\gamma RIV$, but not to the high affinity activating receptor $Fc\gamma RI$. Moreover, the activating-to-inhibitory (A/I) ratio for mouse IgG2b (affinity of individual IgG subclasses for distinct activating FcyR and the inhibitory $Fc\gamma RIIB$) is higher than that of mouse IgG2a subclass, indicating that IgG2b binds with more avidity to inhibitory FcyRIIB than activating FcyR.⁶² Monocytes express both FcyRI and FcyRIV whereas NK cells mainly express FcyRIII. Most in vivo ADCC depleting activity is mediated by monocytes (Fc γ RI and Fc γ RIV) rather than by NK cells (Fc γ RIII).⁶³ We could not demonstrate any in vitro ADCC or antibody dependent cell-mediated phagocytosis on opsonized LIGHT transduced target cells, and therefore it is very unlikely that the mouse IgG2b isotype of 3D11 hybridoma may mediate any in vivo ADCC-depleting activity. Besides, 3D11, an IgG2b subclass, exhibits a high binding avidity for the inhibitory $Fc\gamma RIIB$, and therefore a high A/I ratio. Co-ligation of IgG to inhibitory $Fc\gamma$ RIIB receptor with high avidity reduces cellular responses mediated by activating receptors. The data suggest that 3D11 antibody may mediate in vivo effector function through neutralization of LIGHT interactions with HVEM and LT β R receptors rather than in vivo depletion through ADCC.

The experimental model used to dissect the role of LIGHT interaction with its receptors was the adoptive transfer of 70×10^{6} of semiallogeneic B6 splenocytes into non-irradiated (recipient mice are not conditioned) F1 recipients (B6 \times BALB/c). This large number of allogeneic splenocytes is required to overcome NK cell-mediated host resistance to engraftment of donor cells (hybrid resistance: resistance of host F1 to parental haematopoietic cell engraftment).⁶⁴ In this particular graft-vs.-host reaction model, the donor alloreactive lymphocytes eliminate host hematopoiesis starting at day 10, reaching the peak of rejection at around day 19. After this initial acute phase of rejection, recipient mice recover and parental engraftment starts to decline from week 3-4 post-adoptive transfer and disease very slowly progresses toward chronic inflammatory disease, reaching the peak at about 12 weeks after the semiallogeneic adoptive transfer. Although recipient hematopoiesis is practically abrogated at the acute phase of the disease (around day 19), no signs of gross pathology in the peripheral tissues are observed. Tissue infiltration requires the rupture of the epithelial barrier, and this does not occur in recipient F1 in which conditioning is not required for the implantation of donor T cells.32

The adoptive transfer of parental B6 splenocytes into F1 recipients (BALB/c × B6) induces an allogeneic response mediated by donor CD8 cytotoxic cells that eliminate host hematopoiesis (graft-versus-host reaction). Donor CD4 T cells directly recognize host MHC class II alloantigens on host APC and provide cognate help to host B cells that experience transient expansion and autoantibody production. These activated B cells, as well as the host haematopoietic compartment, are soon attacked by activated B6 donor CD8 T cells that recognize host allogeneic MHC class I alloantigens and differentiate toward cytotoxic T cells with the help provided by donor B6 CD4 T cells.^{32,65}

In this mouse model, a reduced index of proliferation and precursor frequency of CD4 and CD8 T cells was observed after either pharmacological blockade or genetic ablation of LIGHT that correlated with impaired T cell expansion and differentiation toward effector T cells measured in an in vivo short-term assay of T cell proliferation tracked with CFSE. These observations were associated with a reduced cytotoxic activity against host F1 cells in a long-term in vivo assay evaluated at the peak phase of the acute graft-vs.-host reaction, 19 days after the adoptive transfer. This indicated that the cytotoxic activity of alloreactive T cells was reduced in the absence of LIGHT. These results are in line with the first phenotypic description of LIGHT KO T cells that denoted an alteration in allogeneic T cell proliferation in these mutant mice due to a defect on IL-2 secretion by CD4 T cells. This impaired production of IL-2 would perturb clonal expansion and the help provided to CD8 T cells to promote T cell division and differentiation toward effector T cells.12,19

In the context of dendritic cell activation and maturation for proper costimulation, the majority of members of the TNFSF ligands (CD40L, $LT\alpha\beta$, LIGHT, TL1a, CD40L, OX-40L, and 4-1BBL), are molecules transiently expressed or highly upregulated upon T cell activation that would interact with the corresponding TNFR partner molecule on APC. These activated APCs would costimulate T cells, promote their division, survival and differentiation.² CD40L, along with $LT\alpha\beta$ and LIGHT, are molecules transiently expressed in the early phase of T cell activation that differentially contribute to the licensing of dendritic cells through interaction with CD40 and $LT\beta R$ respectively. $LT\alpha\beta/LT\beta R$ and CD40/CD40L interactions appear to be the most relevant, while LIGHT provides a second layer of regulation.^{20,21,22,66,67,68} Between LIGHT/LT β R and LIGHT/HVEM, the predominant interaction is the former, because $LT\beta R$ outcompetes HVEM due its higher affinity for LIGHT.¹² Indirectly, $LT\beta R$ signaling in stromal cells is also important for maintaining the integrity of lymphoid tissue, which is indispensable for the development of the immune response.^{51,19,67} Indeed, LIGHT/LT β R interaction modulates lymph node hypertrophy by activating stromal cells expressing $LT\beta R$ to release chemokines, upregulate adhesion molecules and thus attract tissue-derived dendritic cells to the draining lymph nodes.²⁸

IL-7R α and KLRG-1 are reciprocally modulated on CD8 T cells as they differentiate toward effector CD8 T cells in viral infection models. Combination of these 2 markers permits the process of T cell differentiation to be followed. 33,69 In our hands, the use of bright fluorochromes attached to anti-mouse CCR7 antibody (clone 4B12) in combination with CD62L gave a weak staining that could not clearly discriminate the different stages of T-cell differentiation. The absence of LIGHT (LIGHT-deficient T cells or antibody blockade of LIGHT) leads to delayed downregulation of IL-7R α and delayed upregulation of ICOS cells at day 5 after the adoptive transfer, suggesting that T cell differentiation was hampered in donor alloreactive CD8 T cells in the late phase of T cell activation and clonal expansion. Moreover, T-cell differentiation toward effector cells is associated with an increased percentage of KLRG-1^{high} shortlived effector cells (SLECs) in the allogeneic adoptively transferred CD8 T cells, whereas the initiation (memory precursor cells) and the contraction phase (long-lived memory cells) is characterized by downregulation of KLRG-1.35 The augmented percentage of KLRG-1^{high} SLECs in the absence of LIGHT suggests a defect in T cell differentiation at day 10, which is linked to a less effective clonal expansion at day 3 after the adoptive transfer.

In summary, the data presented here provide evidence for the therapeutic use of LIGHT inhibitors to dampen immune responses via attenuation of donor allogeneic CD8 T cell responses.

Material and methods

Animals

Eight to 12 weeks-old female C57BL/6 (H-2^b, B6, CD45.2), BALB/c (H-2^d), CB6F1 (BALB/c × C57BL/6 F1) mice (H-2^{d/b}) and LIGHT-deficient mice (LIGHT^{-/-}) backcrossed more than 15 generations onto C57BL/6 background were bred at the animal facility of the University of Leon (Spain).¹⁹ All experiments with rodents were performed in accordance and following animal protocols specifically approved for the Ethical Committee for Animal Research of the School of Veterinary Medicine (University of Leon), the Animal Welfare Committee of University of Alcala de Henares (Madrid) and followed the European Guidelines for Animal Care and Use of Laboratory Animals.

Monoclonal antibodies and surface plasmon resonance

LIGHT-deficient mice were immunized intraperitoneally with 0.25 ml of a 1:1.2 mixture of $5 \cdot 10 \times 10^6$ of GFP-tagged mouse LIGHT transduced into NIH-3T3 cells in Freund's Incomplete Adjuvant (Sigma).¹² Six weeks after the priming immunization, mice received an intravenous booster injection of LIGHT-transduced cells in saline. The immortalization of mouse B cells with myeloma cell line X63 Ag8.653 was previously described ⁷⁰ and supernatants of the heterohybridomas secreting rat mAbs were tested 10 days after for their specificity against LIGHT transduced cells by flow cytometry.

Flag-Foldon-tagged soluble mouse LIGHT (FF-LIGHT),⁷¹ soluble FF-control protein (FF-Ctrl) and LT β R.human IgG1.Fc (LT β R.Ig) were produced and purified as previously reported.⁷² Recombinant mouse HVEM.mouse IgG2a.Fc (HVEM.Ig) produced in insect cells was a gift from Genentech. Commercially available mouse IgG_{2a} and human IgG₁ were used as matched isotype control for the recombinant proteins.

For the screening of anti-LIGHT antibodies displaying antagonist activity and therefore capable to prevent LIGHT interaction with its receptors, 2.5×10^5 LIGHT-transduced NIH-3T3 cells were incubated with a saturating amount of isotype matched control or anti-mouse LIGHT mAbs for 30 min at room temperature (2 µg/ml). In the presence of competitor antibody, cells were then further incubated for 2 h at 37°C with an optimal dilution of either HVEM-Ig (5 µg/well) or LT β R-Ig (1 µg/well) in a final volume of 100 µl. After a washing step, the reaction was developed with the appropriate biotinylated conjugates (biotinylated rat anti-mouse IgG_{2a} isotype specific mAb, clone R19-15, BD Biosciences or mouse anti-human IgG Fc fragment, Jackson ImmunoResearch), followed by allophycocyanin-coupled to streptavidin.

The BIACORE 3000 system, sensor chip CM5, surfactant P20, amine coupling kit containing N-hydroxysuccinimide (NHS) and N-Ethyl-N'-dimethylaminopropyl carbodiimide (EDC), were from BIACORE (Upsala, Sweden). All biosensor assays were performed with HEPES-buffered saline (HBS-EP) as running buffer (10 mM HEPES, 150 mM sodium acetate, 3 mM magnesium acetate, 3.4 mM EDTA and 0.005% surfactant P20, pH 7.4). The different compounds were dissolved into running buffer.⁷³

Fc γ RI (CD64), Fc γ RIIB (CD32), Fc γ RIIIA (CD16) and Fc γ RIV (CD16.2) were immobilized at 50 μ g/ml in formate buffer, pH 4.3 by injection onto the EDC/NHS-activated surface of a CM5-type sensor chip until a signal of approximately 6000 RU was obtained. Free activated sites of the matrix were saturated by injection of 20 μ L ethanolamine hydrochloride, pH 8.5. All the binding experiments were carried out at 25°C with a constant flow rate of 30 μ l/min. Different concentrations of mouse anti-LIGHT (clone 3D11) were injected for 3 min followed by a dissociation phase of 3 min. The sensor chip surface was regenerated after each experiment by injection of 20 μ l of 10 mM NaOH.

The kinetic parameters were calculated using the BIAeval 4.1 software on a personal computer. Global analysis was performed using the simple Langmuir binding model. The specific binding profiles were obtained after subtracting the response signal from the channel control (ethanolamine) and from blank buffer injection. The fitting to each model was judged by the reduced chi square and randomness of residue distribution.

Flow cytometry

Mouse anti-mouse LIGHT (3D11, mouse IgG_{2b}) and mouse IgG_{2b} isotype control (MPC-11, Biolegend) mAbs were labeled with Hilyte-Fluor 647 according to the manufacturer's protocol (Anaspec Inc.). CD4 (GK1.5), CD8 α (53-6.7), Ly6G (1A8), CD19 (6D5), ICOS (7E.17G9), CD127 (IL-7R α , A7R34), and killer cell lectin-like receptor subfamily G, member 1 (KLRG-1, MAFA) mAbs were all purchased from Biolegend (USA). To distinguish haematopoietic cells of donor parental B6 and host F1, cells from distinct haematopoietic compartments were stained with FITC-conjugated anti-H-2^d (SF1-1.1) and Alexa 647-conjugated anti-H-2^b (AF6-88.5). Rat mAb anti-mouse BTLA (CD272, clone 4G12b)⁷⁴ and anti-mouse HVEM (CD250, clone 10F3)²⁹ were produced and labeled in-house and used in this study.

Fc gamma receptors were blocked by incubating cell suspensions with 2 μ g/ml of blocking anti-Fc γ R (rat IgG_{2b} anti-Fc γ RII/III mAb, clone 2.4G2) before staining with fluorochrome-labeled antibodies.⁷⁵ Dead cells were excluded from the acquisition gate by staining with propidium iodide. Samples were acquired on a Cyan 9 cytometer (Beckman Coulter, Miami, FL, USA) and data analysis was performed using Win-List version 7.0 (Verity Software House, Topsham, ME, USA).

In vitro polyclonal T cell activation to induce LIGHT expression

Naïve C57BL/6 splenocytes were polyclonally activated in vitro with Phorbol Myristate Acetate (PMA, 100 ng/ml) plus ionomycin (500 ng/ml) or were left untreated for 5 h at 37°C. The transient expression of LIGHT was analyzed on resting and polyclonally activated T cells (2×10^5 cells / well) stained with 1 µg/well of Hylite 647-labeled anti-LIGHT mAb (clone 3D11) or Hylite 647-labeled isotype-matched mouse IgG_{2b} (MPC-11) during the time of incubation, using a lineage cocktail to gate out all cells with phagocytic phenotype (CD19⁺ Ly6G⁺ CD11c⁺) that take up the antibody unspecifically.

In vivo proliferative assay of CFSE-labeled donor alloreactive T cells

 70×10^6 of B6 WT or LIGHT-deficient unfractionated splenocytes were labeled with 5 μ M carboxyfluorescein diacetate succinmidyl ester (CFSE) and adoptively transferred into nonirradiated F1 recipients, according to Lyons et al.⁷⁶ The day of the adoptive transfer, recipient mice were treated with 1 mg (40 mg/kg) of isotype-control (mouse IgG_{2b}) or mouse antimouse LIGHT (3D11) mAb. Three days later, the PI and PF of alloreactive CD4⁺ and CD8⁺ T cells were determined by deconvoluting and analyzing the reduced CFSE fluorescence using the ModFit LT 4.1 version (Verity software, ME). PtdIns, is a measure of the increase in cell number over the course of the assay. PF returns the fraction of cells in the initial population that responded to the stimulus by proliferating. The location of each generation of cells is represented by a unique peak color.

In vivo murine model of Graft-versus-Host Reaction

In a third experimental group, F1 recipient mice were inoculated with 70×10^6 splenocytes from LIGHT-deficient mice in B6 background and treated with 1 mg isotype control. Finally, a syngeneic control group was included in which 70×10^6 syngeneic F1 splenocytes were injected into non-irradiated F1 recipients. The absolute number of haematopoietic cells in primary and secondary lymphoid organs was recorded 19 days after the adoptive transfer.

Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.

Statistical analysis

Collected data from experimental and control groups were analyzed using Graph Pad prism Version 5. Statistical significance was assessed using the non-parametric Mann-Whitney t test or Student's t test with Welch's correction to compare the means among groups. A p value less than 0.05 was considered statistically significant.

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