Original Research

A novel multifunctional anti-PD-L1-CD16a-IL15 induces potent cancer cell killing in PD-L1-positive tumour cells

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ABSTRACT

Cancer is the most acute disease and the leading cause of patient death worldwide. Both chemotherapy and molecular-based therapies play an important role in curing cancer. However, the median and overall survival of patients is poor. To date, immune therapies have changed the treatment methods for cancer patients. Programmed death ligand 1 (PD-L1, also known as B7-H1, CD274) is a well-studied tumor antigen. PD-L1 is overexpressed in colon cancer, lung cancer, and so on and plays a vital role in cancer development. In this study, anti-PD-L1 single-domain antibodies were identified from recombinant human PD-L1 (rhPD-L1)-immunized llamas. Then, we generated a novel multifunctional anti-PD-L1-CD16a-IL15 antibody targeting PD-L1-positive tumor cells. Anti-PD-L1-CD16a-IL15 was constructed by linking the Interleukin-2 (IL-2) signal peptide, anti-PD-L1 single domain antibody (anti-PD-L1-VHH) and anti-cluster of differentiation 16a single domain antibody (anti-CD16a-VHH), and Interleukin-15/Interleukin-15 receptor alpha (IL15/IL-15Ra). This anti-PD-L1-CD16a-IL15 fusion protein can be expressed and purified from HEK-293F cells. In vitro, our data showed that the anti-PD-L1-CD16a-IL15 fusion protein can recruit T cells and drive natural killer cells (NK) with specific killing of PD-L1-overexpressing tumor cells. Furthermore, in the xenograft model, the anti-PD-L1-CD16a-IL15 fusion protein inhibited tumor growth with human peripheral blood mononuclear cells (PBMCs). These data suggested that the anti-PD-L1-CD16a-IL15 fusion protein has a latent function in antitumour activity, with better guidance for future cancer immunotherapy.

Introduction

Cytokines are a broad and loose category of small proteins (~5–20 kDa). Cytokines are essential mediators of communication for the human immune system, and a number of them are crucial for host defense against pathogens [1]. To date, an increasing number of studies have focused on applying many types of cytokines in antiviruses and cancer treatment [2]. Among cytokines, IL-15 has been listed as the most promising antitumor drug candidate [3]. IL-15 and IL-2 belong to the same cytokine family [4], IL-15 and IL-2 have the same receptor β chain and γ chain (IL-2/15γ), but they have different α receptor chains Interleukin-2 receptor alpha/ Interleukin-15 receptor alpha (IL-2Ra/IL-15Ra) [5]. Similar to IL-2, IL-15 can stimulate the proliferation of T cells and natural killer (NK) cells, the expansion of cytotoxic T cells [6], and the activation of NK cells [7]. However, IL-2 but not IL-15 has immunosuppressive effects [5]. IL-15 may have more potent antitumor activity than IL-2. Interestingly, the IL-15/IL-15Ra complex increases the activity of IL-15 by approximately 50 times [8]. Therefore, an IL-15/IL-15Ra fusion protein may perform better than IL-15 alone. Given the multiple functions of IL-15, including increasing the number of activated natural killer (NK) cells, monocytes, and granulocytes, a systemic increase in IL-15 activity may lead to high toxicity [9,10]. Recombinant IL-15 has been used in clinical research for cancer treatment, but its efficacy is limited due to its half-life and toxicity [10]. Therefore, targeting IL-15 to the tumor microenvironment, allowing immune cells to specifically participate in the tumor microenvironment to enhance the antitumor effects of IL-15, will be an attractive strategy. IL-15 activates NK cells, and antibody-dependent cellular cytotoxicity (ADCC) is one of the main cytotoxic mechanisms by which FcγR-expressing effector cells eliminate tumor cells. It has been reported that a bispecific antibody modified the IL-15 crosslinker through genetic insertion to enhance ADCC and activate NK cells to improve anticancer efficacy [11]. In addition, targeting the CLEC12A trispecific killer conjugation molecule can effectively induce NK cells to improve the
efficacy against AML tumors [12]. On the other hand, by increasing IL-15-mediated activation of CD8 T cells, a tumor microenvironment targeting IL-15 using RGD or CEAs was constructed and shown to exhibit strong antitumor activity [13,14].

CD16 (FcγRIII), a receptor for the IgG Fc domain, FcγRIIIa (CD16a) and FcγRIIb (CD16b), facilitates ADCC and plays an essential role in triggering the lysis of target cells by natural killer (NK) cells [15]. It has been shown that monococytes expressing CD16 have a variety of ADCC capabilities in the presence of specific antibodies and can kill cancer cells [16]. Furthermore, CD16a is a potent cytotoxicity receptor on human NK cells, which can be exploited by therapeutic bispecific antibodies [17]. Human NK cells account for ~10% of all lymphocytes and are defined phenotypically by the expression of cluster of differentiation 56/cluster of differentiation 16 (CD56+CD16+) [15]. Two isoforms of human CD16, CD16a and CD16b, share 96% sequence identity in their immunoglobulin-binding regions. CD16a is expressed on NK cells, macrophages, and mast cells and is an activating receptor [18]. On the other hand, CD16b is expressed on granulocytes as a GPI-anchored receptor and does not activate tumor cell killing [18]. Bispecific antibodies are attractive candidates for cancer immunotherapy because they can effectively induce NK cells to kill tumor cells [16]. To activate NK cells, anti-CD16a antibodies have been studied and used to develop bispecific antibodies [16,17].

Programmed death-ligand 1 (PD-L1), also known as cluster of differentiation 274 (CD274) or B7 homologue 1 (B7-H1), is a protein that in humans is encoded by CD274 gene [19,20]. PD-L1 overexpression occurs in different kinds of human cancers, including colon [21], lung [22], pancreas [23], breast [24], ovarian [25], and head and neck cancers [26]. The overexpression of PD-L1 leads to tumor cell progression and tumorigenesis [27]. The tumor microenvironment plays an important role in good prognosis in immunotherapy based on programmed death-ligand 1/programmed cell death protein 1/cluster of differentiation 80 (PD-L1/PD-1/CD80) immune checkpoint inhibitors [28]. Durvalumab is an FDA-approved anti-PD-L1 monoclonal antibody (mAb) for immune checkpoint blockers in patients with lung adenocarcinoma [29]. However, although immune checkpoint inhibitors show great clinical benefits, their efficacy is still modest [30]. To date, there is no benefit against PD-L1-positive colon cancer. Thus, it is better to develop new ways to improve PD-L1-targeting cancer therapy.

Studies have shown that an attractive strategy to improve the antitumor effect of antibodies is to use smaller antibody formats to increase tumor penetration [31]. Many mAb fragments, such as antigen-binding fragment (Fab), single-chain variable fragment (scFv), and nanobodies, have been widely used. However, Fabs and scFv have low yields, poor stability and solubility, and slow half-lives in vivo [9]. Single domain antibodies, also called nanobodies or single domain antibody (VHH), are derived from the variable domains of heavy-chain antibodies in camelids [32]. Although VHH is the smallest fully functional antigen-binding fragment (approximately 15 kDa), its specificity and affinity for recognizing antigens are similar to those of immunoglobulin G (IgG) antibodies [33]. Due to its smaller size, VHH has enhanced tissue penetration and epitope entry [33]. In addition, single-domain antibodies can be expressed in high yields, and VHH also exhibits high thermal stability, conformational stability, and solubility and is resistant to acids and alkalis [34,35]. These excellent characteristics make VHH a valuable and attractive tool for various applications.

Phage display technology has been used to screen and identify ligands for protein, carbohydrate molecules, small peptides and antibodies, improve peptide or antibody affinity, and indicate protein–protein interactions [36,37]. In addition, baculofiber lambda, representing a classical cloning and expression system, can express peptide or protein sequences as fusions to coat proteins [38]. More recently, whole-genome bacteriophage lambda display human libraries are a relatively novel use for identifying new antigens for biomedical applications [39,40].

This study constructed an anti-PD-L1-CD16a construct by fusing an anti-PD-L1 VHH with an anti-CD16a VHH. Another anti-PD-L1-IL15 construct functions by fusing an anti-PD-L1 VHH with IL-15/IL-15Ra. A novel multifunctional anti-PD-L1-CD16a-IL15 antibody was constructed by linking anti-PD-L1-VHH and anti-CD16a-VHH antibodies and IL-15/IL-15Ra. All these fusion proteins can promote immune cell proliferation to drive potent PD-L1-positive tumor cell killing in vitro and in vivo. In addition, compared with anti-PD-L1-CD16a or anti-PD-L1-IL15, anti-PD-L1-CD16a-IL15 showed better activity. These findings hold significant importance for using anti-PD-L1-CD16a-IL15 as a potential cancer immunotherapy.

Materials and methods

Immunized VHH phage display library construction

PD-L1 single domain antibodies were generated as described previously [41]. Second, recombinant human PD-L1-His-protein (Genscript) was used to immunize llamas. Third, the serum titer test was performed using an enzyme-linked immunosorbent assay (ELISA) by coating the antigen rhPD-L1 and following the manufacturer’s instructions (Abcam, cat# ab214565). A high serum titer of a llama was achieved after four immunizations. Then, the peripheral blood of the llama was extracted, and lymphocytes were isolated using gradient centrifugation. Total RNA was isolated from lymphocytes with TRIzol Reagent (Invitrogen, cat# 15596018). The RNA template, via reverse transcription, produces complementary DNA (cDNA), which can be used directly as a template for polymerase chain reaction (PCR); the camelid VHH fragments were amplified using specific primers. Then, the camelid VHH fragments were ligated to the pMECS vector. Finally, a camelid VHH phage library was created by transforming the ligation products into XL1-Blue E. coli cells.

VHH phage display library amplification

To amplify the library, 200 μl of the PD-L1-VHH phage library was inoculated into 40 ml of superbroth medium (10 g of MOPS 3-(N-morpholino)propanesulfonic acid (Sigma, cat# 8899), 30 g of tryptone (BD-Bioscience, cat# 0123–17–3), and 20 g of yeast extract (BD-Bioscience, cat# 0127–17–9); 1 L ddH2O) containing 100 μg/ml ampicillin and 10 μg/ml tetracycline incubated at 37 °C and shaken at 220 rpm until an Optical Density (OD600) of 0.8. Then, 10 μl (2.50×1013 colony-forming units (CFU)) of helper phage VCSM13 was added and incubated for 15 min at 37 °C without agitation. Then, the phage culture was incubated at 37 °C and shaken at 250 rpm for 2 h. After that, the bacteriophages were collected by spinning down at 4000 rpm/min for 10 min at room temperature. Then, the pellet was resuspended in 40 ml fresh superbroth medium containing 100 μg/ml ampicillin, 10 μg/ml tetracycline, and 50 μg/ml kanamycin. They were then incubated at 30 °C with agitation overnight. To transfer overnight culture to an autooclaved 50 ml centrifuge tube, the cells were centrifuged at 4000 rpm/min for 10 min at 4 °C. Next, the phages were precipitated from the bacteriophage supernatant using 5X polyethylene glycol (PEG)/sodium chloride (NaCl) (20% PEG/2.5 M NaCl). In addition, to resuspend the pellet with 1 ml of phosphate buffered saline (PBS), Repeat precipitated steps one more time to remove any bacterial cells. Finally, the phages were resuspended in 200 μl of PBS+1% bovine serum albumin (BSA) buffer.

VHH phage display library screening

To screen the PD-L1-specific VHH binders, as described previously [41], briefly, rhPD-L1-His-antigen was coated on the surface of a 96-well plate. Then, approximately 2.50×103 CFU phages were added to the coated plates and incubated at 37 °C in a 220 rpm shaker for 15 min. To wash plates 6 times with 200 μl/well 0.1% phosphate-buffered saline with Tween 20 (PBST), the weakly bound or nonbinding phages were removed. To adjust the pH, the specific binders were eluted with 100 μl Glycine-BSA buffer ((0.2 M glycine-HCl, 1 mg/ml BSA, pH 2.2) and neutralized with 10 μl 2 M Tris base buffer (pH 9.0). These phage binders
are defined as output phages. The eluted phage-infected competent XL1 blue grew to mid-log phase with OD600=0.6 and was amplified. This phage binder is defined as an input phage and used for the next panning rounds. The panning was repeated three times. After biopanning, we picked 86 clones and amplified them with VCSM13 helper phage in two 2 ml U-bottom 96-well deep blocks. A phage ELISA was performed using 100 µl of the phage supernatant per sample. The positive phage clones precipitated using PEG/NaCl and resuspended in 200 µl PBS. Quantification of bacteriophage by spectrophotometry.

ELISA

For the phage ELISA analysis, a 96-well ELISA plate (Thermo Scientific, Nunc) was coated with 100 µl/well of 2 µg/ml rPhD-L1 antigen in 0.1 M sodium bicarbonate (NaHCO3) (pH 8.6) buffer at 4 °C overnight. The plate was blocked with 200 µl 0.2% BSA blocking buffer and incubated at 37 °C for 2 h. Then, the plate was washed 2 times with 200 µl PBS containing 0.05% Tween-20 (pH 7.2). Add 50 µl of phage culture per well; duplicate wells are used for each clone, incubated at 37 °C for 1 h. The plate was washed 4 times with 200 µl PBS containing 0.05% Tween-20 (pH 7.2). Then, 50 µl of HRP/anti-M13 monoclonal conjugate (GE Healthcare Life Science, cat# 27-9420-01) diluted 1:5000 in PBS containing 1% BSA was added to each well and incubated at room temperature for 1 h. To wash the plate 4 times with 200 µl PBS containing 0.05% Tween-20 (pH 7.2), 50 µl per well of TBMB substrate solution was added and incubated at 37 °C for 10−15 . Stop the reaction with 50 µl stopping buffer. Another uncoated rPhD-L1 antigen was used as a negative control and the same process as described above was performed. The absorbance of the plate was measured on a reader (Tecan) at 450 nm. The positive phage binders Optical Density (OD450) was coated plate OD450-uncoated plate OD450. All of the experiments were repeated at least three times.

For PD-L1 ELISA analysis, recombinant human PD-L1 (Sino Biological, cat# 10084-H02H) or recombinant mouse PD-L1 (Sino Biological, cat# 50010-M02H) (0.1 µg/well) was coated at 4 °C overnight, and the plate was washed with 200 µl/well washing buffer once. Next, 200 µl/well blocking buffer (PBS+0.1% BSA) was added and incubated at 37 °C for 1 h to block nonspecific binding sites. Then, the plate was washed one time. Next, quantification of supernatants with unpurified anti-PD-L1 VHVs by using the BCA Protein Assay Kit (BioRad), then unpurified primary antibodies of anti-PD-L1 VHVs (10 µg/well, HEK-293F medium 100 µl/well) or different purified primary antibodies of PD-L1 and control antibody were added to each well of the plate and incubated at 37 °C for 1 h. Then, wash the plate five times. Next, the HRP anti-6X His-tag antibody (Abcam, cat# ab1187) or HRP-conjugated anti-human IgG (Fab'2) (Abcam, cat# ab87422) (1:2000) was added and incubated at 37 °C for 35 min. After washing five times, the substrate was added, and the absorbance of each well was read at 450 nm. All of the experiments were repeated at least three times.

Antibody design and purification

Diagrams of the anti-PD-L1 VH single antibody are shown in Fig. 2A. Briefly, for anti-PD-L1 VH, the single domain anti-PD-L1 VH was fused with a His-tag. Anti-PD-L1-CD16a antibodies were constructed by fusing anti-PD-L1 and anti-CD16a single domain antibody (GenBank: ABQ52436.1) with a (GGGS)3 linker (Fig. 3A, a). For anti-PD-L1-IL15, anti-PD-L1 VH with IL15/IL15Ra and a (GGGS)3 linker was used (Fig. 3A, b). The IL-15Ra sushi domain (GenBank: CAG33345.1, amino acid from 1 aa to 77 aa in UniProtKB) included the subsequent 12 amino acids from exon 3 as previously described [13]; IL15 (GenBank: CAA62616.1, amino acid from 30 aa to 162 aa in UniProtKB) and IL15/IL15Ra were linked with a SG2SG4SG3SG4SLQ peptide to mimic the physiological trans-presentation of IL-15 as previously described [13,42]. Finally, for anti-PD-L1-CD16a-IL15, anti-PD-L1-CD16a with IL15/IL15Ra with a (GGGS)3 linker (Fig. 3A, c).

Then, the fusion genes were cloned into the pcDNA3.1(+) vector with an IL-2 signal peptide, and a His-tag was added to the C-terminal end for protein detection and purification. The antibodies were finally expressed in HEK-293F cells, as previously described [14]. The fusion proteins were purified by a Ni-NTA affinity (Qiagen, cat# 30210) purification system. After purification, samples of the antibodies were run on 10% sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) gels under reducing conditions, and then the gel was stained with Coomassie brilliant blue R-250 dye (Thermo Fisher Scientific, cat# 20278). All of the experiments were repeated at least three times.

Flow cytometry analysis

For the flow cytometry analysis, LS174T and CHO cells were grown to 80%–90% confluence. First, the cells were digested with 0.25% trypsin, incubated at 37 °C, and collected. Then, 5 × 10⁶ cells/sample were collected by centrifugation at 1000 rpm for 5 min. Then, the cells were washed with 1 ml of ice-cold phosphate-buffered solution (PBS)+0.2% bovine serum albumin (BSA) 2 times. The samples were then incubated with different antibodies (final concentration of 50 µg/ml) for 1 h on ice. The cells were then washed twice, and goat anti-human IgG (H + L) AF488 (Invitrogen, cat# A11013) or anti-6X His-tag FITC (Abcam, cat# ab1206) was used as the secondary antibody for another 1 h. After washing the cells three times, flow cytometry analysis was performed on an FC50 (FC 500, Beckman Coulter). All the experiments were repeated at least three times.

Affinity measurement

The affinity of anti-PDL1 antibodies to the extracellular part of the PD-L1 protein was determined using a ProteOn™ XPR36 Protein Interaction Array System (Bio-RAD). Briefly, human recombinant PD-L1 proteins with an Fc tag (Sino Biological, cat# 10084-H02H-B) in phosphate-buffered saline with Tween 20 (PBST) were loaded onto the surface of a ProteOn XPR36 GLC chip. Immobilization levels between 1.0 and 1.5 nM were reached. The tested anti-PD-L1 antibodies were then applied in concentrations of 0 nM, 0.125 nM, 0.25 nM, 0.5 nM, 1 nM, and 2 nM. Then, the association and dissociation phases were measured for 200 s and 15 min, respectively. ProteOn data were analyzed by ProteOn Manager (version 2.0), and the relation between association and dissociation is the equilibrium dissociation constant which equals the association rate constant divided by the dissociation rate constant (KD=Ka/Kd). All of the experiments were repeated at least three times.

Blood cell fractionation

Human peripheral blood mononuclear cells (PBMCs) were freshly prepared by Ficoll density centrifugation (GE Health) from blood that was collected from 11 health donors as described previously with some modifications [17,43]. According to the manufacturer’s instructions, NK cells or T cells were then isolated from the PBMCs by EasySepTM Human NK Cell Enrichment Kit (Stem cell Co. Ltd, cat# 1905) or EasySepTM Human T Cell Enrichment Kit (Stem cell Co. Ltd, cat#19,661).

Cell proliferation analysis

For cell proliferation, CTLL-2 and T cells were harvested, washed twice with PBS, and incubated with RPMI 1640 with 10% PBS and 1% non-essential amino acid (NEAA) at 37 °C and 5% CO2. After 4 h of incubation, the cell suspension at a density of 2 × 10⁵ cells/ml was seeded into a 96-well plate with duralumab (Creative Biolabs, cat# TAB-417CQ) or rHL-15 or anti-PD-L1-VHH or anti-PD-L1-CD16a or anti-PD-L1-IL5 or anti-PD-L1-CD16a-IL15 at different concentrations (varied from 0.0001 to 1000 nM). After 72 h of incubation at 37 °C and 5% CO2, Cell Counting Kit-8 (Dojindo, cat# CK06–10) was performed. All of the
The experiments were repeated at least three times.

**CFSE labeling of T or NK cells and proliferation assay**

To measure T cells or NK cells proliferation, T cells or NK cells were freshly prepared by Ficoll centrifugation, adjusted to 2 × 10^6 cells/ml, and then stained with 5 μM carboxyfluorescein succinimidyl ester (CFSE) (Biolegend, cat# 423801) according to the manufacturer’s instructions. Stained T cells (0.5 × 10^6 cells/ml in 12-well plates) were incubated with 10 nM anti-PD-L1-IL-5 or anti-PD-L1-CD16a-IL15 for 1 day. Stained NK cells (0.5 × 10^6 cells/ml in 12-well plates) were incubated with 10 nM durvalumab, rhIL-15, anti-PD-L1-VHH, anti-PD-L1-CD16a, anti-PD-L1-IL-5 or anti-PD-L1-CD16a-IL15 for 5 days. Then, T cell (1 day) or NK cell (5 days) proliferation was assessed on an FC50 (FC 500, Beckman Coulter) and analyzed using FlowJo v10 software (BD-Biosciences).

**Western-blot analysis**

For western blotting were performed as described previously with some modifications [44–46]. T cells were harvested, washed twice with PBS, and incubated with RPMI 1640 with 10% FBS and 1% NEAA at 37 °C for 4 h. After 4 h of incubation, the cells suspension at a density of 5 × 10^5 cells/ml were seeded into the 96-wells plate with durvalumab or rhIL-15 or anti-PD-L1-VHH or anti-PD-L1-CD16a or anti-PD-L1-IL-5 or anti-PD-L1-CD16a-IL15 at 100 nM. After 48 h of incubation at 37 °C, 5% CO2, cells were rinsed with ice-cold PBS twice and lysed with RIPA buffer (Sigma-Aldrich, cat# R0278) plus the protease inhibitor cocktail. Protein concentrations were determined using the BCA Protein Assay Kit (BioRad). About 25 μg protein per sample was loaded on a 10% SDS-PAGE gel and transferred to the polyvinylidene difluoride (PVDF) membrane, which was blocked with 5% milk in PBS, and incubated with the anti-STAT5 (Cell Signaling Technology, cat# 94205) or phospho-STAT5 (Tyr694) antibody (Cell Signaling Technology, cat# 9351) or GAPDH antibody (Cell Signaling Technology, cat# 2118) in the cold room overnight. After washing with PBST 3 times for 10 min each, the membranes were incubated with the horseradish peroxidase-conjugated secondary antibody (Abcam, cat# ab6721) at room temperature for one hour. The membranes were washed with PBST three times with 10 min and subjected to chemiluminescent western blot analysis. All of the experiments were repeated at least three times.

**Cytotoxic assays**

Cytotoxic assays were performed as described previously with some modifications [16]. Briefly, CHO and LS174T cells were seeded into 96-well plates at a density of 2500 cells per well and incubated at 37 °C, 5% CO2 for 6 h. A total of 25,000 NK cells, T cells or NK+T cells and different concentrations of antibodies were added to each well. After 72 h incubation, removed the T cells and NK cells, then the CHO or LS174T cells was calculated using the following formula: \[(\text{live tumor cells (sample)} - \text{medium}) / (\text{live tumor cells (control)} - \text{medium})\]. The NK or T cells viability were measured by Cell Counting Kit-8 reagent (Dojindo) following the manufacturer’s instructions. LS174T cells viability were measured by Cell Counting Kit-8 reagent (Dojindo) following the manufacturer’s instructions. After 72 h incubation, removed the T cells and NK cells, then the CHO or LS174T cells were incubated in the cold room overnight. After washing with PBST 3 times for 10 min each, the membranes were incubated with the horseradish peroxidase-conjugated secondary antibody (Abcam, cat# ab6721) at room temperature for one hour. The membranes were washed with PBST three times with 10 min and subjected to chemiluminescent western blot analysis. All of the experiments were repeated at least three times.

**In vivo antitumour experiments**

In vivo studies were performed as described previously [14,16]. Briefly, PD-L1-positive LS174T cells (1 × 10^6) were mixed with freshly isolated human PBMCs (5 × 10^5) (from two healthy donors). The mixed cells (200 μL/mouse) were then transplanted to the right flank of 5-week-old female nonobese diabetic/severe combined immunodeficiency (NOD/SCID) mice by subcutaneous injection. After transplantation for 1 h, durvalumab (3.5 mg/kg or 0.7 mg/kg), recombinant human Interleukin-15 (rhIL-15) (0.35 mg/kg or 0.07 mg/kg), anti-PD-L1-VHH (0.35 mg/kg or 0.07 mg/kg), anti-PD-L1-CD16a (0.7 mg/kg or 0.14 mg/kg), anti-PD-L1-IL5(0.9 mg/kg or 0.18 mg/kg), and anti-PD-L1-CD16a-IL15 antibodies (1.2 mg/kg or 0.24 mg/kg), and negative control (PBS) were administered intraperitoneally to 200 μL per mouse (i.p.) (n = 6 each group). Treatment was performed every two days. The volume of the tumor was measured every two days. Tumor volume was calculated by the formula (width^2×length)/2.

To further validate the antitumor activities of anti-PD-L1 antibodies in vivo. 1 × 10^6 LS174T cells with 200 μL PBS per mouse, the cell suspensions were then injected subcutaneously into the right flank of NOD scid gamma mouse (NSG) mice (female, 6 weeks old). Then, PBMCs freshly were isolated from 1 healthy donor, and different anti-PD-L1 antibodies or PBS were administered intraperitoneally when the tumor size reached 50–100 mm³. Anti-PD-L1 antibodies or rhIL-15 treatment was performed every two days. In addition, mice were weighed, and tumor growth size was recorded every two days. Tumor volume was calculated by the formula (width^2×length)/2. All animal experiments were carried out with the approval of the Ethics Committee of Guangxi Medical University and were performed in accordance with the Institutional Animal Welfare Guidelines set by Guangxi Medical University.

**Results**

**Phage display library construction and single-domain antibody screening**

We constructed a library of anti-PD-L1 single domain antibodies. First, llamas were immunized seven times with 300 μg of PD-L1 recombinant human protein. Then, we isolated lymphocytes from the llamas immunized and constructed a phage library. Next, we conducted three rounds of panning on phage, and anti-PD-L1 phages were significantly enriched (Table 1). After panning, we randomly picked 86 clones and expanded them by incubating them overnight at 37 °C and 220 rpm. After that, we performed an ELISA using 100 μl phage medium per sample. The results showed that 31 out of 86 clones recognized the rhPD-L1 antigen (data not shown). In addition, 31 positive phage clone supernatants with PEG/NaCl solution are an easy way to obtain higher viral titers by concentrating virus. Finally, the pellet was resuspended in 200 μL PBS. The phage concentration measured the OD value of A280. Furthermore, the quantitative ELISA analysis was performed. The positive phage binders OD_450=coated plate OD_450-uncoated plate OD_450. The data showed that 16 positive clones specifically recognized the rhPD-L1-His-antigen (Figs. 1 and S1).

The anti-PD-L1-VHH antibody can be produced and purified in HEK-293F cells

The anti-PD-L1-VHH antibody was constructed by fusing two parts: the anti-PD-L1-VHH single domain antibody can be produced and purified in HEK-293F cells.

**Table 1**

|                | 1st Round | 2nd Round | 3rd Round |
|----------------|-----------|-----------|-----------|
| Input phage (cfu) | 2.50×10^11 | 2.50×10^12 | 5.00×10^12 |
| Output phage (cfu) | 3.50×10^6  | 5.50×10^6  | 4.00×10^6  |
| Output/Input      | 1.40×10^-5 | 2.20×10^-4 | 8.00×10^-3 |

rhPD-L1 antigen (1 μg) was coated in a 96-well plate. Then, the phage library containing approximately 2.50×10^11 cfu phages was input. The input phages were incubated with the coated plates for 15 min at 37 °C. The weakly bound phages or nonspecific phage binders were washed away. Specific binder phages were eluted and named the Output. The eluted phage binders were used to infect competent E. coli XL1-blue and amplified for the next rounds. To enrich the positive binders, the panning was performed for 3 rounds by coating decreased rhPD-L1 antigen from 1 μg to 200 ng in each cycle, which increased stringency.
anti-PD-L1 single-domain antibodies targeting PD-L1-overexpressing cancer cells. In addition, a His-tag was added at the C-terminus to facilitate protein purification and detection (Fig. 2A). To further analyze the positive clones, an ELISA was performed to investigate the binding of the anti-PD-L1-VHH antibody to the rhPD-L1 antigen. The anti-PD-L1-VHH antibody can be produced in HEK-293F cells. On posttransfection day 6, we collected the supernatant. Again, an ELISA was performed, and the data showed that clone #13 showed a large obvious difference in binding to the rhPD-L1 antigen compared with other clones (Fig. 2B). Based on the results, clone #13 was further analysed.

The anti-PD-L1-VHH single domain antibody can bind PD-L1-overexpressing tumor cells

To study the function of the anti-PD-L1-VHH antibody, clone #13 antibody was expressed in HEK-293F cells. The clone #13 antibody was soluble and purified by Ni-NTA affinity chromatography purification, and the anti-PD-L1-VHH antibody with molecular weights of ~15 kDa was obtained (Fig. 2C).

Furthermore, ELISA was performed to check the binding of the anti-PD-L1-VHH antibody to the rhPD-L1 antigen. Both anti-PD-L1-VHH antibody and durvalumab could bind to rhPD-L1 protein (Fig. 2D). Therefore, these results suggested that the anti-PD-L1-VHH antibody has a strong binding affinity for the rhPD-L1 protein.

In addition, to check whether the anti-PD-L1-VHH antibody can bind to PD-L1-overexpressing tumor cells, flow cytometry analysis was performed using PD-L1-positive cancer cells, LS174T cells, PD-L1-negative CHO cells. For PD-L1-positive cancer cells, LS174T, anti-PD-L1-VHH, or positive antibody (durvalumab) showed positive staining (Fig. 2E, left panel) but very low or no staining on the PD-L1-negative cell line CHO (Fig. 2E, right panel), suggesting that the anti-PD-L1-VHH antibody binds specifically to PD-L1-positive tumor cells. To further evaluate the binding of anti-PD-L1 antibodies to PD-L1, the affinity of anti-PD-L1-VHH to PD-L1 was measured. The data showed that anti-PD-L1-VHH have affinity of 1.04–09 M. In addition, the affinities of the previously mentioned antibodies (1.0–09 M) indicates good candidates for further study [47], suggesting that anti-PD-L1-VHH have good affinity.

To test whether anti-PD-L1-VHH cross-reacts with murine PD-L1, ELISA was performed. The results showed that neither durvalumab nor anti-PD-L1-VHH were cross-reactive with murine PD-L1 (Fig. 2F). Furthermore, ELISA competition assays were performed to examine whether durvalumab and anti-PD-L1-VHH bound to the same epitope. The results showed that both durvalumab and anti-PD-L1-VHH can recognize the rhPD-L1 protein. However, the effect of anti-PD-L1-VHH on rhPD-L1 protein was not abolished by durvalumab. These data showed that neither durvalumab nor anti-PD-L1-VHH bound to the same epitope (Fig. 2G).

Anti-PD-L1-CD16a-IL15 stimulates immune cell proliferation in vitro

Anti-PD-L1-CD16a-IL15 was constructed to contain three functional modules (Fig. 3A). The anti-PD-L1 VHH targets PD-L1-overexpressing tumor cells; anti-CD16a VHH engage NK cells; and the IL-15/IL-15Ra complex activates and increases in the NK cells and T cells population. In addition, in the binding of to detection and purification. The fusion gene of anti-PD-L1-CD16a-IL15 was cloned into the pCDNA3.1(+) vector and then transiently transfected into HEK-293F cells. Anti-PD-L1-CD16a as a control antibody was constructed to contain anti-PD-L1-VHH, anti-CD16a VHH, and His-tag. Anti-PD-L1-IL15, another control antibody, contained anti-PD-L1-VHH and IL-15/IL-15Ra complexes and His-tag. These fusion proteins were purified from the HEK-293F cell culture medium by Ni-NTA affinity chromatography. Bands of ~28 kDa (anti-PD-L1-CD16a), 38 kDa (anti-PD-L1-IL15), and 53 kDa (anti-PD-L1-CD16a-IL15) were observed under reducing conditions (Fig. 3B).

The cytokine activity of anti-PD-L1-CD16a, anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15 was then assessed using a cytokine-dependent cell proliferation assay with the CTLL-2 cell line and human T cells. The purpose is to assay IL-15/IL-15Ra complex activity. The data showed that all the rhIL-15, anti-PD-L1-IL15 and anti-PD-L1-CD16a-IL15 dramatically stimulated the proliferation of the CTLL-2 cell line and human T cells at a similar level, not anti-PD-L1-VHH, anti-PD-L1-CD16a and durvalumab, demonstrating that the IL-15/IL-15Ra complex of anti-PD-L1-IL15 and anti-PD-L1-CD16a-IL15 maintains IL-15 cytokine activity (Fig. 3C).

To measure the activity of anti-PD-L1-CD16a, anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15 on primary immune cells, human T cells or NK cells were prepared, T cells stained with CFSE and incubated with anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15 for 5 days. After 1 day, anti-PD-L1-IL15 and anti-PD-L1-CD16a-IL15 significantly stimulated proliferation of human T cells compared to cells in the control group (Fig. 3D). After 5 days, rhIL-15, anti-PD-L1-IL15 and anti-PD-L1-CD16a-IL15 significantly stimulated proliferation of human NK cells, and anti-PD-L1-CD16a and durvalumab but not anti-PD-L1-VHH slightly stimulated the proliferation of human NK cells (Fig. 3E).

The effects of anti-PD-L1-CD16a-IL15 treatment on the activation of STAT-signaling pathways were then evaluated using human T cells. Following receptor engagement, IL-15 signaling activates signal transducer and activator of transcription 5 (STAT5) transcription factors, to exert its regulatory functions in immune cells [48]. Indeed, in human T cells, compared to vehicle, rhIL-15 or anti-PD-L1-IL15 or anti-PD-L1-CD16a-IL15 induced an increase in STAT5 phosphorylation (Fig. 3F). Different from anti-PD-L1-CD16a-IL15, both anti-PD-L1-VHH and anti-PD-L1-CD16a showed no effects on the phosphorylation of STAT5 in T cells (Fig. 3F). These results suggest that...
anti-PD-L1-CD16a-IL15 have significant effects on the STAT5-signaling pathway.

**Anti-PD-L1-CD16a-IL15 mediates potent cytotoxic activities against PD-L1-positive tumor cells**

To check whether anti-PD-L1-CD16a, anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15 can mediate specific tumor cell killing, cytotoxic assays were performed using the PD-L1-positive cell lines LS174T and the PD-L1-negative cell line CHO. Both LS174T and CHO cell lines were incubated with durvalumab, rhIL-15, anti-PD-L1-VHH, anti-PD-L1-CD16a, anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15 in the presence of T cells, NK cells, or mixed T cells and NK cells, respectively (Figs. 4 and S2). For PD-L1-negative CHO cells, durvalumab, rhIL-15, anti-PD-L1-CD16a-IL15 had no cell killing activity either alone or in the presence of NK cells.
Fig. 3. Biochemical characterization of the anti-PD-L1-CD16a-IL15 fusion protein
(A) Schematic representation of anti-PD-L1-CD16a and anti-PD-L1-IL15 and anti-PD-L1-CD16a-IL15 fusion proteins. (B) Coomassie blue staining of anti-PD-L1-CD16a (28 kDa), anti-PD-L1-IL15 (38 kDa), and anti-PD-L1-CD16a-IL15 (53 kDa) fusion proteins.
(C) In the left panel, CTLL-2 proliferation stimulated by durvalumab, rhIL-15, anti-PD-L1-VHH, anti-PD-L1-CD16a and anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15.
In right panel, human T cell proliferation was stimulated by durvalumab, rhIL-15, anti-PD-L1-VHH, anti-PD-L1-CD16a and anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15.
(D) T-cell proliferation stimulation in vitro by anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15. CFSE-labeled T cells were incubated with 10 nM of different proteins for 1 day. The proliferation of T cells was assessed by flow cytometry.
(E) NK-cell proliferation stimulation in vitro by durvalumab, rhIL-15, anti-PD-L1-VHH, anti-PD-L1-CD16a, anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15. CFSE-labeled NK cells were incubated with 10 nM of different proteins for 5 days. The proliferation of NK cells was assessed by flow cytometry.
(F) Human T cell proliferation was stimulated by durvalumab, rhIL-15, anti-PD-L1-VHH, anti-PD-L1-CD16a and anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15, then total cell lysates were analyzed by immunoblot for the STAT5 protein or STAT5 phosphorylated protein or GAPDH protein. The protein expression level, quantified by band intensity and normalized to GAPDH, is displayed in the right panel. The results are the averages of duplicates from three independent experiments. Data are the means ± SEM, N.S., not significant, **P < 0.01; ***P < 0.001 compared with the control.
cells (Fig. S2A), T cells (Fig. S2B), or mixed NK cells and T cells (Fig. S2C). For PD-L1-positive tumor cells, LS174T, durvalumab, rhIL-15, anti-PD-L1-VHH, anti-PD-L1-CD16a, anti-PD-L1-IL15 or anti-PD-L1-CD16a-IL15 antibodies and freshly isolated NK cells. The effector (NK cells) (25,000 cells/well) and target LS174T cells (2500 cells/well) at a ratio of 10:1. (B) Different cell lines were treated with durvalumab, rhIL-15, anti-PD-L1-VHH, anti-PD-L1-CD16a, anti-PD-L1-IL15, or anti-PD-L1-CD16a-IL15 with freshly isolated T cells. The effector (T cells) (25,000 cells/well) and target LS174T cells (2500 cells/well) at a ratio of 10:1. (C) Different cell lines were treated with durvalumab, rhIL-15, anti-PD-L1-VHH, anti-PD-L1-CD16a, anti-PD-L1-IL15, or anti-PD-L1-CD16a-IL15 with freshly isolated T+NK cells. The effector (T+NK cells at a ratio of 1:1) (25,000 cells/well) and target LS174T cells (2500 cells/well) at a ratio of 10:1. (D) Different cell lines were treated with durvalumab, rhIL-15, anti-PD-L1-VHH, anti-PD-L1-CD16a, anti-PD-L1-IL15, or anti-PD-L1-CD16a-IL15 fusion proteins with freshly isolated T+NK cells. The effector (T+NK cells at a ratio of 1:1) (25,000 cells/well) and target LS174T cells (2500 cells/well) at a ratio of 10:1. All fusion proteins induce T+NK cell-mediated cytotoxicity in a dose-dependent manner. The results are the averages of duplicates from three independent experiments. Data are the means ± SEM, N.S., not significant, *P < 0.05; ** P < 0.01; *** P < 0.001 compared with the control. See also Fig. S2.

Dose-dependent cell killing by anti-PD-L1-CD16a, anti-PD-L1-IL15,
and anti-PD-L1-CD16a-IL15 was further analysed using different concentrations of anti-PD-L1-CD16a, anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15. No cytotoxic activity against CHO cells was observed regardless of the presence of NK+T cells or the concentrations of anti-PD-L1-CD16a, anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15 (Fig. S2D). However, for PD-L1-positive cells, LS174T (Fig. 4D), increased cytotoxic activities were observed with increased concentrations of anti-PD-L1-CD16a, anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15 in the presence of NK+T cells. Interestingly, anti-PD-L1-CD16a-IL15 shows much stronger activities than anti-PD-L1-CD16a and anti-PD-L1-IL15. These data suggested that the cytotoxic activity of anti-PD-L1-CD16a, anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15 depends on the expression of PD-L1 and the presence of NK+T cells. In addition, the anti-PD-L1-CD16a-IL15 fusion protein is considered a novel therapy for cancer.

Anti-PD-L1-CD16a-IL15 targets tumor tissues and has potent antitumour activities in vivo

To evaluate whether anti-PD-L1-CD16a, anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15 can suppress tumor growth in vivo. First, LS174T PD-L1-positive cancer cells were grafted into NOD/SCID mice. Then, LS174T cells were transplanted together with freshly prepared human PBMCs into NOD/SCID mice. When mice were treated with durvalumab, rhIL-15, anti-PD-L1-VHH, anti-PD-L1-CD16a, anti-PD-L1-IL15, and anti-PD-L1-CD16a-IL15 in the presence of PBMCs, dramatic tumor growth inhibition was observed (Fig. S5A). However, anti-PD-L1-CD16a-IL15
When the tumor volume reached 50 to 100 mm
alone, no significant tumor growth inhibition was observed. Further
more, the mice were also treated with the same molar amount of the
control antibody anti-PD-L1 VHH, which recognizes PD-L1-positive cells
in the presence of PBMCs. As anti-PD-L1 VHH cannot activate immune
cells, no significant tumor growth inhibition was observed for anti-PD-
L1 VHH treatment (Fig. 5A). In addition, no weight loss or apparent
toxicity was observed in any mouse (Fig. 5B). These data confirmed that
anti-PD-L1-CD16a-IL15 can inhibit PD-L1-positive tumor growth in vivo.

To further evaluate anti-PD-L1-CD16a-IL15 antitumor activity, a
synergetic model using the NSG mice was also evaluated. After trans-
planting LS174T cells in NSG mice, rapid tumor growth was observed.
When the tumor volume reached 50 to 100 mm3, the mice were
administered freshly isolated human PBMCs and treated with PBS,
durvalumab, anti-PD-L1-CD16a, anti-PD-L1-IL15, and anti-PD-
L1-CD16a-IL15 every 2 days. Tumor growth inhibition was then
measured (Fig. 5C). Interestingly, anti-PD-L1-CD16a-IL15 showed
stronger antitumor effects than durvalumab, rhIL-15, anti-PD-L1-
CD16a, and anti-PD-L1-IL15. However, rhIL-15 treatment resulted in
severe systemic toxicity, as indicated by dramatic body weight loss,
which was consistent with previous report [49] (Fig. 5D). Interestingly,
no weight loss or apparent toxicity was observed in any other group
(Fig. 5D). We hypothesize that anti-PD-L1 fusion proteins improve
tumor tissue penetration, reduce the off-tumor expansion of NK cells or
T cells, diminish systemic toxicity, and efficiently inhibit tumor growth.
These data demonstrated that anti-PD-L1-CD16a-IL15 has potent in vivo
antitumor effects without gross toxicity in NSG mice.

Discussion

Programmed death-ligand 1 (PD-L1) is also known as cluster of dif-
ferentiation 274 (CD274) or B7 homologue 1 (B7-H1) [50]. When PD-L1
binds to its receptor, PD-1, the PD-1 signal activates T cells, B cells, and
myeloid cells to modulate strong activation or inhibition [51]. There
is also emerging evidence that PD-L1 recognizes its receptor PD-1 on
T cells to deliver a signal that suppresses TCR-mediated activation of IL-2
and T cell proliferation [52]. However, it appears that overexpression of
PD-L1 may help cancers escape host immune responses [21,50]. Thus,
PD-L1 is a cancer immunotherapy target, as it is upregulated in various
cancers, including non-small-cell lung cancer (NSCLC) [53], anal car-
cinoma [54], cervical carcinoma [55], head and neck cancers [56], and
hepatocellular carcinoma (HCC) [57]. This study screened anti-PD-L1
VHH single-domain antibodies from the phage display library and pu-
rified anti-PD-L1-VHH antibodies from HEK-293F cells. In addition, we
tested the anti-PD-L1 antibodies function in PD-L1-overexpressing can-
cer in vitro and in vivo. In summary, our study confirmed that
anti-PD-L1-CD16a and anti-PD-L1-IL15 and anti-PD-L1-CD16a-IL15
fusion proteins have potent antitumour activity against PD-L1-positive
cancer cells.

IL-15 is a potential immunotherapy candidate for cancer, and it can
stimulate the proliferation of T cells and NK cells [6]. However, the
defect in IL-15 is its nonspecific systemic distribution, which leads to
toxicity in various tissues [10]. Thus, their efficacy as immunity therapy
is limited. To reduce the toxicity of IL-15, researchers selectively local-
ized IL-15 in the tumor to minimize toxicity. Therefore, a tumor
microenvironment that uses RGD peptides to target IL-15 was con-
structed, and it has been previously shown to have strong antitumor
activity [13]. In addition, anti-CEA-IL-15 induced strong immune cell
proliferation and antitumor activity in vitro and in vivo [14]. In this
study, to increase the local concentration of IL-15 in tumor tissues,
anti-PD-L1 VHH was fused with anti-CD16a VHH and IL-15/IL-15Ra
(Fig. 3A). First, anti-PD-L1 VHH showed a strong binding affinity,
enhanced tumor targeting and reduced toxicity. In addition, IL-15/IL-15Ra
could amplify T cells and NK cells, and the anti-CD16a VHH antibody could engage NK cells to enhance their ability to kill
tumor cells. Furthermore, the use of anti-PD-L1-CD16a-IL15 (53 kDa)
instead of regular IgG (150 kDa) reduced the molecular weight of
anti-PD-L1-CD16a-IL15 by approximately 3 times, which may improve
tumor tissue penetration. Interestingly, the function of anti-PD-L1-CD16a-IL15 is similar to that of anti-PD-L1 bispecific anti-
odies. However, compared to those of mAbs, the lower molecular
weights of anti-PD-L1-CD16a, anti-PD-L1-IL15 and anti-PD-L1-CD16a-IL15 may have the disadvantage of more rapid
clearance in vivo. To enhance its potential in the clinic, conjugation of
anti-PD-L1-CD16a-IL15 with PEG or other means to increase the in vivo
half-life may be considered.

The ELISA results suggested that anti-PD-L1-VHH did not share the
same binding epitope as durvalumab on the rhPD-L1 protein (Fig. 2G).
This provides great potential for using anti-PD-L1-CD16a-IL15, alone or
in combination with durvalumab, to treat PD-L1-positive cancer. The
combination therapy of trastuzumab with pertuzumab, another anti-
Her2 antibody that binds to a different epitope than trastuzumab, is
more effective in the clinic [58,59]. In addition, we are assessing
whether anti-PD-L1-CD16a-IL15 has synergistic effects in inhibiting
PD-L1-positive tumors when in combination with durvalumab, avel-
ubah, or atezolizumab. This study will provide potentially more effective
combinations than the current treatments in PD-L1-positive cancer.

In summary, we provide a novel format for a trifunctional antibody-
fusion protein, anti-PD-L1 VHH, with anti-CD16a VHH and IL-15/IL-
15Ra. This format consisted of a tumor-directed anti-PD-L1 single
domain antibody, anti-CD16a VHH, and IL-15/IL-15Ra. In addition,
anti-PD-L1-CD16a-IL15 fusion proteins can be expressed and purified
from HEK-293F cells with good solubility and stability. Furthermore,
the flow cytometry assay and ELISA showed that the anti-PD-L1 VHH anti-
body could bind to PD-L1-positive cells. Interestingly, anti-PD-L1-
CD16a-IL15 fusion proteins also potently and specifically stimulate
immune cell proliferation in vitro. In vivo studies demonstrated that
anti-PD-L1-CD16a-IL15 fusion proteins could suppress tumor growth in the
presence of PBMCs in NOD/SCID and NSG mice.

In conclusion, considering its potential anticancer activities, the lower
molecular weight of anti-PD-L1-CD16a-IL15 may allow for
improved tumor tissue penetration, reduce the off-tumor expansion of
NK cells or T cells, diminish systemic toxicity, and efficiently inhibit
tumor growth. The potential to use anti-PD-L1-CD16a-IL15 to treat PD-
L1-overexpressing cancer combined with current anti-PD-L1 therapeu-
tics. Therefore, trifunctional antibody-fusion proteins provide a new
promising immunity therapeutic.

Author contributions

Conception and design of the study: Y.L., X.F., H.W.; experimental
performance: Y.L., L.W., Y.L., S.M., B.H; data collection and analysis and
explanation of experiments: Y.L., H.W.; drafting and critical revision of
the manuscript: Y.L., X.F., H.W. All authors approved the final version of
the manuscript.

RB/Ethical review board approval or patient consent/approval
as a statement

Procedures for the collection and use of blood samples were
approved by the Human Research Ethics Committee of the First Affili-
ated Hospital of Guangxi Medical University, and all subjects provided
informed consent. All animal experiments were carried out with the
approval of the Ethical Committee of Guangxi Medical University.

Data available on request from the authors

The data that support the findings of this study are available from the
corresponding author, [X.F] and [H.W], upon reasonable request.
