Cloning, Expression and Characterization of a Peptibody To Deplete Myeloid Derived Suppressor Cells in a Murine Mammary Carcinoma Model

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Abstract

Myeloid derived suppressor cells (MDSCs) are an immature heterogeneous population of myeloid lineage that attenuate the anti-tumor immune responses. Depletion of MDSCs has been shown to improve efficacy of cancer immunotherapeutic approaches. Here, we produced and characterized a recombinant peptibody capable of recognizing and depleting murine MDSCs. Using SOE-PCR, the coding sequence of the MDSC binding peptide and linker were synthesized and then ligated into a home-made expression plasmid containing mouse IgG2a Fc. The peptibody construct was transfected into CHO-K1 cells by lipofectamine 3000 reagent and the resulting fusion protein was purified with protein G column and subsequently characterized by ELISA, SDS-PAGE and immunoblotting. The binding profile of the peptibody to splenic MDSCs and its MDSC depletion ability were then tested by flow cytometry. The purified peptibody appeared as a 70 kDa band in Western blot. It could bind to 98.8% of splenic CD11b^+/Gr-1^+ MDSCs. In addition, the intratumoral MDSCs were significantly depleted after peptibody treatment compared to their PBS-treated negative control counterparts (P <0.05). In this study, a peptibody capable of depleting intratumoral MDSCs, was produced. Our results imply that it could be considered as a potential drug effective for immunotherapy of cancers.

Introduction

Myeloid derived suppressor cells (MDSCs) are known as an immunosuppressive population of myeloid lineage that are enriched in cancer and chronic infections [1, 2]. MDSCs are primarily generated from myeloid precursors of bone marrow in response to tumor-derived factors such as VEGF, TGF-β, IL-6, IL-10, CSF-1 and GM-CSF, and are subsequently recruited to blood, spleen and tumor bed [3]. MDSCs are functionally distinguished from the normal myeloid cells based on their ability to inhibit the immune responses mediated by T cells, B cells and natural killer (NK) cells [4–6]. These immune inhibitory effects of MDSCs mostly contribute to expression of the regulatory mediators including arginase1, S100A8/A9, NO, reactive oxygen species, immunosuppressive cytokines and surface immune checkpoints [7]. Moreover, MDSCs facilitate metastasis of primary tumors to distant organs through formation of premetastatic niches [8]. MDSCs are involved in matrix remodeling by induction of MMP8 and MMP9, angiogenesis process and support the metastatic potency of circulating cancer cells [9, 10]. In a recent study it was reported that epigenetic therapy could disrupt the pulmonary premetastatic niches in murine tumor models by down-regulation of CXCR2 and CCR2 chemokine receptors that mediate the MDSC migration to peripheral organs [11]. Overall, MDSCs take part in the immune escape, invasion and progression of tumors to other organs [12, 13].

MDSC-targeting agents are involved in diverse mechanisms to overcome immunosuppression mediated by MDSCs by interfering in MDSC differentiation, hampering their recruitment to tumor site, depleting MDSCs and by reprogramming MDSCs in tumor microenvironment [14]. In this context, early studies demonstrated that all-trans retinoic acid (ATRA) induce differentiation of MDSCs into mature macrophages and DCs and augment the efficacy of anti-VEGFR2 therapy in a breast cancer animal model [15, 16]. Besides, cytotoxic chemotherapeutic agents including 5-fluorouracil, carboplatin, paclitaxel or gemcitabine have been shown to diminish MDSCs in blood circulation and improve anti-tumor immune responses [17–20]. However, these
drugs reduce other rapidly proliferating cell populations such as T cells and highlight the need for development of more specific agents without off-targeting effects. Francis Mussai et al. discovered depleting effects of an immunotoxin, anti-CD33 monoclonal antibody (gemtuzumab) conjugated to ozogamicin, on human MDSCs that improved the efficacy of CAR-T cell immunotherapy [21]. This immunotoxin was investigated in a phase II clinical trial and could significantly deplete CD33+ MDSCs with promising results [22, 23]. Moreover, anti-GR-1 monoclonal antibody is also able to significantly reduce the MDSC frequency in tumor bearing mice [24, 25]. However it mainly reacted with PMN-MDSCs and thus, M-MDSCs could escape from the elimination by Gr-specific antibodies. Recently, Hong Qin et al. generated a novel therapeutic peptibody that efficiently depleted both PMN-MDSCs and M-MDSCs in tumor-bearing mice [26]. Here, a peptibody containing the MDSC-binding peptide and the Fc domain of mouse IgG2a was produced in a eukaryotic system and was characterized in terms of recognition and depletion of MDSCs in 4T1 mouse tumor model.

Materials And Methods

Design and cloning of the peptibody coding sequence

The MDSC binding peptibody contains a short specific peptide that is linked to the Fc domain of mouse IgG2a antibody (Gene ID: 380793) by a glycine-serine linker (GS linker) together with an IL-2 signal peptide that is located upstream of the sequence (Fig. 1a) [26]. SmaI and BstbI restriction sites were inserted upstream of the peptide and downstream of the linker sequences, respectively. Two slow codon pairs in GS linker were exchanged with a fast counterpart by only a single nucleotide change of A to T (GGAGGC to GGTCGC). In order to produce codon optimized MDSC-specific peptide and linker sequences (84 bp), three overlapping primers were designed and applied in Splicing by Overlap Extension (SOE) PCR (Bioneer, Daejeon, Korea) (Table 1). The product of each PCR was used as a template for the next PCR. The final PCR product was digested with SmaI and BstbI enzymes (Fermentas, Tokyo, Japan) and subsequently cloned into a home-made expression plasmid containing mouse IgG2a Fc and glutamine synthetase (GS) gene as a selection marker. The recombinant construct was ligated using T4 DNA ligase (Fermentas) and transformed into chemically competent E. coli DH5a host cells through heat shock approach. Positively transformed colonies were screened by colony PCR using primers designed for CMV promoter (forward primer: 5-CAGACATAATAGCTGACAGACTAAC-3) and SV40 poly-A tail (reverse primer: 5-ATACCTACCAGTTCTGC-GC-3) (Bioneer). The PCR reaction was carried out with an initial melting step at 94°C for 5 min, followed by 35 cycles of denaturation at 94°C for 30 Sec, annealing at 60°C for 30 Sec, and extension at 72°C for 1 min with a final extension step at 72°C for 6 min. Moreover, the selected colonies were confirmed using the enzymatic digestion and DNA sequencing (Applied Biosystems 3500, CA, USA).

Table 1. Overlapping primers applied in SOE PCR to extend MDSC-specific peptide and linker sequences. To produce MDSC specific peptide and linker GS, the two overlapping sequences were designed and fused using the complementary part of sequences (underlined in table) in a PCR cycle. Then, the SmaI and BstbI restriction sites were joined in 5′ and 3′ ends of the product, respectively by overhanging primer pairs 1 and
2 carrying special annealing sequence of the template molecule ends. Smal restriction site: CCCGGG, BstBI restriction site: TTCTGAA.

| Primer Name | sequence (5→3) | Size (bp) |
|-------------|----------------|-----------|
| Overlapping primer 1 | TATGGGGCTGGTCTCTGAGCCACGGCTACCAGGTGAAGTCTGGTGGGC | 52 |
| Overlapping primer 2 | TGATCCGCCGCCACCCGACCCACCTCCGCGGCCGACCCGCCACCCAGACTTTCACT | 56 |
| Primer 1, Forward | CGGGTATGGGGCTGGTCTCTGA | 22 |
| Primer 1, Reverse | GAATGATCCGCCGCCACCC | 19 |
| Primer 2, Forward | TTTTTTCCCGGTATGGGGCTGTT | 24 |
| Primer 2, Reverse | CACCATTTCGAATGATCCGCCGCCA | 25 |

**Transient expression of peptibody in CHO-K1 cell line**

After cloning of the peptibody sequence, DNA plasmid was gently extracted from the transformed bacterial cells using low endotoxin midiprep kit (Invitrogen, Vilnius, Lithuania). Moreover, CHO-K1 cells were cultured in RPMI-1640 medium (Gibco, Grand Island, NY, USA) supplemented with 10% fetal bovine serum (Gibco), 100 U/mL penicillin (Sigma, Darmstadt, Germany), and 100 µg/mL streptomycin (Sigma). Peptibody construct was utilized to transfect CHO-K1 cells according to the manufacturer's instructions (Invitrogen, Carlsbad, CA, USA). Briefly, the plasmid DNA was mixed with P3000 reagent in serum free OPTI-MEM medium (Gibco) and also the lipofectamine3000 was diluted separately in OPTI-MEM medium. The two solutions were mixed and then added to the seeded cells in 12 well plate. Two days after transfection, the transient expression of recombinant peptibody in cell supernatant was checked by ELISA.

**Detection of secreted peptibody in cell culture supernatant by ELISA**

Briefly, the ELISA 96-well plate (Nunc, Glostrup, Denmark) was coated with 2 µg/ml of affinity purified sheep anti-mouse IgG polyclonal antibody (SinaBiotech, Tehran, Iran) for 2 hours at 37°C. After blocking, 100 µl
transfected cell supernatant were added to the wells and incubated for 1.5 hour at 37°C. The plate was then incubated with horseradish peroxidase-labeled sheep anti-mouse Ig polyclonal antibody (1:1000 dilution in blocking buffer) at 37°C for 1 hour. The absorbance values were measured at 450 nm by an Anthos 2020 Microplate Reader (Biochrom, Cambridge, UK).

**Stable expression of the peptibody by transfected CHO-K1 cells**

The culture medium was replaced with glutamine free DMEM medium (Sigma) supplemented with 10% dialyzed FBS (HyClone™, GE Healthcare, Marlborough, MA, USA) and L-Methionine sulfoximine (MSX, 25µM) (Sigma) 48 hours post-transfection. After two weeks, the production level of the peptibody by each colony in the supernatant was measured by ELISA. Colonies with the highest expression level were selected and subcloned in 96-well plates by limiting dilution method. The final clone was cultured in serum free First CHOice medium (UGA Biopharma, Hennigsdorf, Germany) containing Feed alpha and Feed beta supplements (UGA Biopharma) and maintained for two weeks at 31°C in 5% CO₂.

**Purification of the peptibody from the culture supernatant by protein G column**

The collected supernatant of stable CHO-K1 clone producing the peptibody was centrifuged (4000 ×g for 5 min at 4°C) and then the clear supernatant was concentrated using a stirred ultra-membrane cell with a 10-kDa cut-off (Amicon 8400, Minden, Germany) under N2 gas pressure. The concentrated sample was then passed through a HiTrap Protein G HP column (GE Healthcare). The peptibody was eluted from the protein G column by a glycine-HCl buffer (0.1M, pH 2.7). The concentration of the peptibody was determined using BCA protein assay kit (Thermo scientific, Rockford, IL, USA).

**Characterization of the peptibody by SDS-PAGE and Western blot**

The purified peptibody was mixed with 6X sample buffer with and without β-mercaptoethanol and boiled. The protein bands were visualized by Coomassie brilliant blue staining. Moreover, the extracted peptibody was identified by immunoblotting. After the 10% SDS-PAGE, the protein contents were transferred onto polyvinylidene difluoride (PVDF) membranes in transfer buffer (25 mM Tris; 192 mM glycine; 20% v/v methanol, pH 8.3) at 100 V for 1 hour (BioRad, Hercules, CA, USA). The membranes were blocked with 5% skim milk in PBS with 0.05% Tween20 (PBS-T) and then were individually incubated with HRP-labeled sheep anti-mouse Ig polyclonal antibody (SinaBiotech, Tehran, Iran) in the blocking buffer at 1:1000 dilution under a shaker platform for 1 hour. After repeated washing with PBS-T, the HRP-antibody bound proteins were visualized using DAB substrate kit (BioGenex, Fremont, CA, USA).

**Induction of 4T1 breast cancer model**

Female inbred Balb/c mice (6-8 weeks old) were obtained from Pasteur Institute of Iran (Karaj, Iran) and maintained under pathogen-free condition with a 12/12-hour light/dark cycle at 22±1°C with 50% humidity according to animal welfare guidelines. To induce murine breast tumor, 4T1 cells were cultured in RMPI-
1640 medium supplemented with 7% FBS and were subsequently injected into the mammary fat pad of mice at 5×10^5 cells suspended in 50 µl PBS. All animal experiments were approved by the Ethical Committee of Shahid Beheshti University of Medical Sciences (IR. SBMU. MSP.REC.1396.874).

**Determination of binding specificity of peptibody to MDSCs**

To prepare FITC conjugate, the purified peptibody was dialyzed against bicarbonate buffer (0.1M, pH 8.3) overnight at 4°C. One milligram FITC was dissolved in 1 ml DMSO and subsequently 20 µl FITC was mixed with the peptibody (1 mg) in a final volume of 1 ml at room temperature for one hour. Next, the mixture was dialyzed against PBS overnight at 4°C [27]. Splenocytes were isolated from 4T1 tumor bearing mice 4 weeks after tumor induction using cell strainer (BD Falcon, Bedford, MA, USA). The splenocytes were then simultaneously incubated with FITC-conjugated peptibody, APC-labeled anti-CD11b (Biolegend, San Diego, CA, USA) and PE-labeled anti-Gr1 antibodies (Biolegend) (1 µg/ml) for 1 hour at 4°C. Also, an irrelevant isotope matched FITC-conjugated antibody (2F9G5 clone) was used as a control for peptibody to exclude the nonspecific binding of peptibody.

**In vivo depletion of MDSCs by the peptibody**

The peptibody was administered intraperitoneally at a dose of 50 µg per mouse at 17, 18 and 19 days after 4T1 tumor cell challenge and a control group of mice received PBS alone. On day 20, mice were sacrificed to remove the spleens and tumor tissues. Primary tumors were cut into the 2-4 mm pieces and then dounced mechanically by a tissue grinder (Kimble Chase, Rockwood, TN, USA) to dissociate into the single cell suspensions. The isolated spleen were gently pressed through a 70 µm cell strainer using 1 ml syringe plungers. Then, isolated splenocytes and tumor cells were co-stained with anti-CD11b/Gr1 and FITC conjugated anti-CD45 antibodies (Biolegend). FACS analysis was performed using flow cytometer cell analyzer (BD Biosciences, FACS Lyrics, San Jose, CA, USA) and the data was analyzed by Flow Jo software V10 (Tree Star Inc, Ashland, OR, USA).

**Statistical analysis**

Statistical analyses were conducted by Graph Pad Prism 8 software (Graph Pad Software Inc., San Diego, CA, USA) using two-tailed t-test to compare MDSC percentages in treated group of mice with the control group that received PBS. P value < 0.05 was defined statistically significant and represented as *.

**Results**

**MDSC binding peptide production in eukaryotic CHO-K1 cells**

The sequence of the MDSC binding peptide and the GS linker were assembled by SOE-PCR which resulted in production of a 111 bp fragment (Fig. 1b). The amplified sequence was then inserted into the plasmid that carried the coding sequences of mouse IgG2a Fc protein. Transformed *E. coli* colonies harboring the correct construct had a 1317 bp amplicon in colony PCR using specific primers (Fig. 1c). Two selected colonies (C12 and C13) were confirmed using enzymatic digestion (Fig. 1d). Furthermore, DNA sequencing
analysis of these two colonies indicated that the MDSC binding peptide was successfully cloned into the expression vector in a correct frame shift without any mutation.

It is interesting to mention that before codon pair optimization of GS linker sequence, the maximum transient expression level of the transfected cells was 12 ng/ml despite several attempts to improve the production level. Codon pair optimization of the MDSC-peptide coding and GS linker sequences improved peptibody production level by 20.8 folds (250 ng/ml) after transient transfection. Finally, after multiple rounds of subcloning and selection of stable clones, a stable high producer clone which was producing 8 μg/ml of the peptibody was selected and expanded.

**Characterization of MDSC binding peptide using SDS-PAGE and immunoblotting**

The size and structure of the purified peptibody were investigated by SDS-PAGE and the peptibody migrated as a single band. The molecular weight of the peptibody was around 70 kDa under non-reducing conditions and its reduced form showed a smaller 35 kDa band (Fig. 2a). The recombinant peptibody was also detected by immunoblotting with HRP conjugated sheep anti-mouse IgG as a 70 kDa protein band (Fig. 2b).

We next determined the binding characteristics of the peptibody to MDSCs. When the 4T1 tumor size reached 760 mm\(^3\), the splenocytes were harvested and stained with the FITC-conjugated peptibody and APC-anti-CD11b and PE-anti-Gr1 antibodies as MDSC markers. The peptibody bound to 98.8% of CD11b/Gr1 positive MDSCs, however, the control peptibody failed to recognize this cell population (Fig. 3). The CD45/CD11b/Gr-1 positive MDSCs substituted 66.2% of the splenocytes obtained from the 4T1 tumor bearing mice.

**MDSCs depletion by the peptibody in 4T1 tumor model**

The peptibody was intraperitoneally administrated at 50 μg/mouse/day; 17, 18 and 19 days after induction of 4T1 tumors (tumor volume: 1500 mm\(^3\)). Flow cytometric analysis showed that intratumoral MDSCs were significantly depleted after peptibody treatment compared with the control group that received PBS alone (39.6% compared to 57.3%) (P <0.05) (Fig. 4). However, no significant reduction of MDSCs was observed in splenocytes of tumor bearing mice (Fig. 4).

**Discussion**

It is well known that tumor microenvironment is infiltrated with various types of immune suppressive cells including the immature myeloid cells, MDSCs [28]. MDSCs certainly have critical roles in tumor development and are documented as a major immunosuppressive population against efficient immunotherapy [29, 30]. In this context, MDSC-targeting strategies lead to MDSC depletion or inhibition of their immunosuppressive function in malignancies. ATRA, an isomer of retinoic acid, can mediate differentiation of MDSCs to mature cells such as macrophages, dendritic cells and granulocytes in human and mouse tumors. By induction of MDSC differentiation, ATRA enhances anti-tumor immune responses of CD4\(^+\) and CD8\(^+\) T cells and improves the efficacy of cancer vaccines [31]. In a lymphoma model, 5-
fluorouracil and gemcitabine dramatically decreased MDSC numbers in spleen and tumor tissues. This inhibition of MDSC population was associated with induction of the INFγ production in tumor-specific CD8+ T cells and development of anti-tumor T cell dependent immune responses [32, 33]. In addition, anti-Gr-1 monoclonal antibody eliminates Gr-1+ granulocytic MDSCs and can restore the antitumor T cell responses. However, it causes elimination of mature granulocytes and consequently induces a severe systemic immunosuppression [34]. Thus, the finding of a specific MDSC-depleting agent with poor off-targeting activity is valuable to overcome the robust tumor-induced immunosuppression and leading to enhancement of cancer immunotherapies in clinical practice. In order to discover an efficient MDSC-targeting agent, Hong Qin et al. identified peptibodies by phage display technique that efficiently depleted both PMN-MDSCs and M-MDSCs in lymphoid peripheral organs and intratumoral sites [26].

In this study, we produced an anti-MDSC peptibody containing a MDSC-binding peptide fused to Fc domain of mouse IgG2a by a GS linker in CHO-K1 cells [26]. Purified peptibody showed a 70 kDa dimeric protein in immunoblotting that was dissociated into 35 kDa monomeric subunits under reducing condition. In our first attempt, the MDSC-specific peptide and linker sequences were synthetized (Biomatik) and cloned in a home-made vector harboring FC domain of mouse IgG2a. However, we found that even optimization was not efficient enough for expression in both transient and stable transfected CHO-K1 cells. Then, the sequences of MDSC-binding peptide and GS linker were optimized on the basis of codon pair bias theory to improve the expression level of peptibody [35] (Supplementary Table 1). We exchanged the slow codon pairs with a fast counterpart in GS linker by only a single nucleotide change of A to T (GGAGGC to GGTTGGC). In agreement with Trinh study [35], we observed that a fast codon pair improved peptibody production levels from 12 ng/ml to 250 ng/ml in the CHO-K1 transiently transfected host. Our efforts to produce stable cell line with original sequences also led to increased production to 8 ug/ml. Thus, in both transient and stable transfection plans, codon pair optimization resulted in a significant improvement in peptibody production level.

It was suggested that overrepresented codon pairs may reduce efficiency of transition process and may thus act as ribosomal pause sites. These slow pairs give the intact polypeptides an opportunity to fold properly by chaperones. So, it seems that elimination of the slow codon pairs to accelerate the translation step may result in increasing the yield of recombinant protein production [36, 37].

When 4T1 murine mammary carcinoma cells were implanted orthotopically into mammary fat pad of female Balb/c mice, they could spontaneously metastasize to bone, liver, lungs and spleen organs which was also affected in late-stage cancer patients [38, 39]. The invasive 4T1 tumor growth causes a leukemoid reaction and splenomegaly accompanied by massive myeloid cell infiltration that was partly mediated via colony-stimulating factor and chemokines secreted by 4T1 tumor cells [40]. The predominant population of these leukocytes has been reported to be CD11b+/Gr-1+ immature myeloid MDSCs that accumulate in primary tumors, spleen, lung and peripheral blood [30, 41]. Transcripts of G-CSF and GM-CSF and myeloid cell chemokines MCP-1, KC, RANTES, MIP-1a and MIP-1b have been detected in cultured 4T1 cells in vitro which suggests that 4T1 tumor cells individually cause a remarkable increase in myeloid infiltrations of primary tumors and metastatic foci. Our findings consistently showed that 4T1 tumor promoted the MDSCs
expansion in spleen and tumor beds [40]. Interestingly, in our experiments 4T1 induced MDSCs to constitute 89% and 57% of CD45+ population in spleen and primary tumor, respectively, four weeks after tumor inoculation (Fig. 4). This suppressive population probably participates in establishment and survival of 4T1 tumors within 3-4 weeks using blocking of innate and adaptive anti-tumor immune responses directed against 4T1 cell antigens and as a result reduction of immune surveillance.

The purified peptibody bound to 98.8% of splenic MDSCs isolated from 4T1 tumor implanted mice and could deplete MDSC population in vivo. Intraperitoneal injection of three doses of peptibody significantly reduced intratumoral MDSC numbers but this reduction was not significant for splenic MDSCs. One explanation for this finding is that four weeks after tumor inoculation the MDSCs constitute 89% of the leukocytes in spleen. Thus, administration of 3 doses of peptibody may not be sufficient to induce significant reduction of MDSCs. However, the frequency of MDSCs in tumor microenvironment is not so high to counteract the peptibody depletion effects. The dose and stability of the peptibody administered into the peritoneum might also affect its functional failure in the spleen. Assessment of the effect of peptibody on MDSCs at early stages of tumor development, during the first 2 weeks of implantation, may help to elucidate the underlying reasons.

**Conclusion**

In this study, we successfully constructed and produced a fusion protein containing MDSC-binding peptide in CHO-K1 cells and improved the expression level of the peptibody by codon pair optimization in GS linker. The peptibody was able to deplete MDSC population resident in 4T1 tumor microenvironment. This peptibody could be applied for combination immunotherapy to overcome the MDSC immune suppressive function.

**Declarations**

**Ethics approval and consent to participate:** Not applicable

**Consent for publication:** Not applicable

**Availability of data and materials:**

All data generated or analyzed during this study are included in the manuscript.

**Competing interests:**

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Authors' contributions:

KRA: performing the tests, data collection, analysis, and writing original draft, VKB: participating in experiments, and data analysis, JM: participating in experiments, and advising, HAO: investigation, methodology, project administration, supervision, FS: methodology, project administration, and supervision, MHF: methodology, data analysis, MJT: data curation, formal analysis, investigation, methodology, project administration, supervision, validation, and editing, MS: conceptualization, data curation, formal analysis, investigation, methodology, project administration, supervision, validation, writing and editing. All authors approved the final manuscript submitted to the journal.

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Disclosure of potential conflicts of interest: Not applicable

Research involving Human Participants and/or Animals:

All animal experiments were approved by the Ethical Committee of Shahid Beheshti University of Medical Sciences (IR. SBMU. MSP.REC.1396.874). All applicable international, national, and/or institutional guidelines for the care and use of animals were followed. All procedures performed in this study were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Informed consent: Not applicable

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Figures

**Figure 1**

**Design, cloning and confirmatory data on MDSC specific peptibody construct.** a) Peptibody contained MDSC-binding peptide fused to FC domain of mouse IgG2a by a glycine-serine linker (GS linker). His-tag and IL-2 signal peptide coding sequences were added to 3' and 5' ends of peptibody, respectively. b) Electrophoresis of SOE-PCR product contained coding sequence of a MDSC-binding peptide and a GS linker. Target amplicon had a 111 bp size after electrophoresis on 2% agarose gel (marked by white arrow). c) Colony PCR of transformed colonies with peptibody construct. After electrophoresis, the positive colonies had a 1317 bp fragment on 1% agarose gel in comparison to the mock vector (397 bp fragment) (pointed by white arrows). d) Double digestion of DNA plasmids extracted from two selected colonies using HindIII and Sall. The digested constructs having the MDSC binding peptide were cleaved to a 361 bp fragment but digested mock vector had a smaller band (251 bp fragment) on 2% agarose gel (pointed by dark arrows). C12: colony 12; C13: colony 13; V: mock vector contained FC-domain, and L: 100 bp ladder (Invitrogen)
Figure 2

Characterization of the purified peptibody by SDS-PAGE analysis and Immunoblotting. a) Peptibody was purified using protein G column and then separated on 10% SDS-PAGE gel in reducing and non-reducing conditions. **Non-Reduced:** lane 1: mouse IgG (10 µg); lane 2: mouse IgG (5 µg); lane 3: purified peptibody (10 µg); lane 4: purified peptibody (5 µg); M: PageRuler Prestained Protein Ladder (Fermentas). **Reduced:** lane 5: peptibody (10 µg); lane 6: peptibody (5 µg); lane 7: mouse IgG (10 µg); lane 8: mouse IgG (5 µg). b) Immunoblotting of purified peptibody by HRP-labeled sheep anti-mouse IgG. Transferred peptibody was visualized by DAB substrate kit under reducing and non-reducing conditions. **Non-reduced:** lane 1: mouse IgG (0.2 µg/well); lane 2: peptibody (0.4 µg/well); lane 3: peptibody (0.2 µg/well); lane 4: peptibody (0.1 µg/well); M: PageRuler Prestained Protein Ladder (Fermentas). **Reduced:** lane 5: peptibody (0.4 µg/well); lane 6: peptibody (0.2 µg/well)
Figure 3

**Flow cytometric analysis of capability of peptibody binding to splenic MDSCs.** a) Splenic MDSCs were identified as CD45⁺/CD11b⁺/Gr-1⁺ population and accumulated in 4T1 bearing mice (66.2%) in comparison to normal mice (9.24%) (3 mouse pre group). b) To evaluate binding ability of peptibody to MDSCs, FITC-conjugated peptibody, APC-anti-CD11b and PE-anti-Gr-1 antibodies were used to stain the splenocytes isolated from 4T1 bearing and normal mice. Also, a FITC-conjugated control antibody was applied to exclude nonspecific bindings. FITC intensity was determined in the gated MDSCs and represented as the histograms. Peptibody bound to 98.8% of splenic MDSCs in 4T1 mice but recognized only 5.28% of splenocytes in normal mice.
Figure 4

Flow cytometric analysis of MDSCs in the spleen and primary tumor of 4T1 bearing mice after treatment with peptibody. Peptibody was intraperitoneally injected at 50 µg dose per day for three days, and then spleens and tumors were harvested to analyze by flow cytometry. Control group was given PBS and in each group was included 4 female Balb/c mouse. Peptibody treatment could decrease the splenic MDSCs to 70.5% proportion compared to PBS received group and significantly depleted the intratumoral MDSCs to 39.6% (a, b). PEP: peptibody

Supplementary Files

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- SupplementaryTable.1.docx