Alum increases antigen uptake, reduces antigen degradation and sustains antigen presentation by DCs in vitro

Tirth R. Ghimire a,b, Robert A. Benson b, Paul Garside b, James M. Brewer b,*

a Strathclyde Institute of Pharmacy and Biomedical Sciences, University of Strathclyde, Glasgow, Scotland, United Kingdom
b Institute of Infection, Immunity and Inflammation, 120 Glasgow Biomedical Research Centre, University of Glasgow, Glasgow G12 8TA, Scotland, United Kingdom

ARTICLE INFO

Article history:
Received 30 January 2012
Received in revised form 13 June 2012
Accepted 15 June 2012
Available online 23 June 2012

Keywords:
Alum
Antigen uptake
Processing
Presentation
YAe system

ABSTRACT

Aluminium adjuvants (alum) have been the only widely approved adjuvants for use in human vaccines since the 1920s, however, the mechanism of action of these adjuvants remains elusive. Due to increasing demand for novel adjuvants, a clearer understanding of the mechanisms that allow these important agents to affect adaptive immune responses will make a significant contribution to the rational design of future vaccines.

Using a novel approach to tracking antigen and antigen presentation, we demonstrate that alum induces higher antigen accumulation and increased antigen presentation by dendritic cells (DCs) in vitro. Antigen accumulation was 100-fold higher and antigen presentation 10-fold higher following alum treatment when compared with soluble protein alone. We also observed that alum causes an initial reduction in presentation compared with soluble antigen, but eventually increases the magnitude and duration of antigen presentation. This was associated with reduced protein degradation in DCs following alum treatment. These studies demonstrate the dynamic alterations in antigen processing and presentation induced by alum that underlie enhanced DC function in response to this adjuvant.

© 2012 Elsevier B.V. Open access under CC BY license.

1. Introduction

Despite the tremendous variety of compounds with adjuvant activity, effective adjuvants for use in vaccines against major diseases such as Human Immunodeficiency Virus/Acquired Immunodeficiency Syndrome (HIV/AIDS), tuberculosis and malaria remain elusive [1]. A significant obstacle to the development of new and improved adjuvants is our lack of knowledge of their mechanism of action. This is particularly true of the aluminium adjuvants (alum) that have been applied in many vaccines since the 1920s [2]. It has been proposed that vaccine adjuvants act indirectly via DCs or other antigen presenting cells (APCs) to induce and enhance the activation of antigen-specific T cells and subsequently the adaptive immune response. CD4+ T cells play a central role in cell-mediated immunity and are activated in response to specific vaccine-derived, peptide epitopes bound to MHC class II (MHCII) molecules, displayed on the surface of APCs [3–5]. DCs possess highly controlled antigen processing functions utilising lysosomal proteases and pH changes optimal for the generation of peptides rather than complete protein degradation [6–8]. The duration and magnitude of antigen presentation are key factors in determining the degree and quality of T cell activation [9,10]. Although it has been suggested that an antigen pulse of a few hours is sufficient to support subsequent T cell division [11], the continued engagement of the peptide:MHCII:T cell receptor (p:MHCII:TCR) complex is required for a high degree of T cell expansion [12,13]. Adjuvants such as alum have been proposed to alter the magnitude and duration of antigen presentation through mechanisms such as increased antigen uptake [14,15] and enhanced expression of MHC class II molecules on the surface of APC [16–18]. While the impact of these factors on antigen presentation has previously been read out in terms of T cell expansion [15,19,20], these studies do not directly explain whether alum impacts on the magnitude and duration of antigen presentation that subserves adjuvant function.

To address this limitation, in the current study we have employed the chimera fluorescent protein, Ealphagreen fluorescent protein (EαGFP), which allows assessment of antigen uptake/degradation and, in combination with the YAe antibody, antigen presentation in situ [21–24]. When EαGFP is internalised by DCs, it is degraded, liberating the Eα peptide for presentation on MHC class II molecules. This p:MHC complex can be detected by staining the cells with the monoclonal YAe antibody which specifically recognises the Eα:A+MHCIId complex [21–24]. Using the EαGFP/YAe system, we have demonstrated that formulation of
antigen in alumin leads to an increase in antigen uptake, a decrease in antigen processing with the eventual result being enhanced magnitude and duration of antigen presentation by DCs in vitro.

2. Materials and methods

2.1. Preparation of DCs from murine bone marrow

Six- to eight-week-old C57BL/6 mice (H-2b) were used to prepare BMDCs, as described previously [25]. Mice were housed in the Central Research Facility, University of Glasgow and procedures were performed according to U.K. Home Office regulations. Bone marrow cells (2 × 10^6/well) were placed in 6-well plates (Corning Incorporated, Corning, NY, USA) and cultured using 10% GMCSF (63 × supernatants) supplemented with RPMI [RPMI 1640 (Sigma, UK), 10% FCS (Gibco, UK), 100 μg/ml penicillin and streptomycin (Invitrogen, UK) and 100 μg/ml l-glutamate (Invitrogen, UK)] at 37 °C in 5% CO₂. At day 3 and 6, cells were fed with each 2 ml/well fresh complete DC media. DCs were used at day 7 for the experiments.

2.2. Antigens and adjuvants

The fluorescent antigen, EoGFP, was prepared in our laboratory using methods described previously [24]. Lipopolysaccharide (LPS; Escherichia coli O111:B4) was bought from Sigma and was used at 1 μg/ml concentration for positive control of MHC class II expression. ALHYDROGEL⃝ was bought from Fremontga Biodscience, Denmark. This adjuvant consists of 3% aluminium hydroxide.

2.3. Analysis of antigen uptake and presentation

To assess the role of alumin in antigen uptake and presentation, we incubated BMDCs with different concentration of EoGFP or EoGFP adsorbed to different concentrations of alumin in a 6-well plate containing 2 × 10^6 cells/5 ml media in each well. Control wells were incubated with media.

To assess the role of alumin in the kinetics of antigen uptake, degradation and presentation, we performed a pulse chase assay. BMDCs (3 × 10^6) were pulsed with pre-determined doses of EoGFP or EoGFP adsorbed to alumin for 1 h. Some of the cells were incubated in media for experimental control. Cells were harvested, washed in HBSS buffer (GIBCO, Invitrogen) (400 × g, 5 min, 4 °C) and BMDC separated from alumin using sterile histopaque (Sigma cat no. H8889; 400 × g, 25 min, 20 °C) followed by washing twice in HBSS buffer (400 × g, 5 min, 4 °C). BMDC (1.5 × 10^6 cells/5 ml) were resuspended in each well of a 6-well plate containing complete DC media and incubated for different chase periods (0 h, 24 h, 48 h and 72 h). After each chase period, cells were analysed by flow cytometry.

2.4. Flow cytometry

The cells were collected in 6 ml fluorescence-activated cell sorting (FACS) tubes (BD FALCON, BD Biosciences Discovery LabWare, USA) and washed (400 × g, 5 min, 4 °C) in FACS buffer (5% FCS, 0.1% sodium azide) and incubated with 100 μL Fc block (2.4G2 hybridoma supernatant) for 30 min. BMDCs were stained with phycoerythrin (PE) anti-mouse CD11c (eBiosciences, clone: N418, cat no. 12-0114), PE armumen hamster IgG (eBiosciences, clone: eBio299Arm, cat no. 12-4888-81), biotinylated anti-mouse E52-68 (eBiosciences, clone: eBioYAe, cat no. 13-5741), bio anti-mouse IgG2b (Southern Biotech, clone: A-1, cat. 0104-08), bio Rat IgG2b, k (BD biosciences, cat no. 553987), alkalicyocyanin (APC) streptavidin (eBiosciences, cat no. 17-4317), peridinin chlorophyll protein (PerCP) anti-mouse I-A/I-E (BioLegends, clone: M5/114.15.2, cat no. 107623), PerCP streptavidin (BD Bioscience, cat no. 554064) for 30 min. Cells were washed twice (400 × g, 5 min, 4 °C) in FACS buffer and analysed by flow cytometry (BD Bioscience, FACS caliber). The results of flow cytometry were analysed by Flowjo software (Flowjo 8.7.1, Stanford University 1995–96). The level of GFP, YAE and MHC class II molecules present on CD11c positive cells were analysed by both mean fluorescence intensity (MFI) and percentage positive cells as determined using isotype controls.

2.5. Data analysis

Data were analysed using GraphPad Prism version 5.00 for Windows, GraphPad Software, San Diego, CA, USA. Results were expressed as mean ± SEM unless otherwise stated. In the data with one independent variable, we used Tukey post test (one way ANOVA) to test significance between any two different treatment groups. Similarly, in the data having more than one independent variable, we used Bonferroni post test (two way ANOVA) to test the significance between any two different treatment groups either at specific dose or at specific time. A p-value of <0.05 was considered as significant.

3. Results

3.1. The EoGFP/YAE system can be used to study the impact of alumin on antigen uptake and antigen presentation by bone marrow DCs (BMDCs)

To determine the suitability of the EoGFP/YAE system to investigate the impact of alumin on antigen uptake and presentation, BMDCs were incubated with protein, alumin-adsorbed protein or in the presence of media or alumin alone for 24 h. We then analysed the level of GFP and YAE staining within the CD11c positive population (see Fig. 1A). Detection of GFP or YAE staining was dependent on the presence of EoGFP; incubating BMDC with alumin alone did not produce any increase in either of these parameters compared with control cultures (see Fig. 1B and C). Adsorption of EoGFP to alumin produced a significant increase in antigen uptake and presentation (p < 0.0001), with about 5-fold higher MFI of GFP and 2-fold higher MFI of YAE compared with cells incubated in EoGFP alone (see Fig. 1C). Similar results were obtained by analysing the proportion of GFP or YAE positive cells (data not shown). The results suggest that the EoGFP/YAE system is appropriate for the in vitro study of antigen uptake and presentation by DCs following alumin treatment.

3.2. Alumin acts as a delivery vehicle and targets antigen acquisition by DCs in vitro

Adsorption of EoGFP to alumin significantly enhanced antigen uptake and presentation compared with soluble antigen over a range (1–100 μg/ml) of antigen doses tested (see Fig. 2A and B). Comparing the antigen dose response curves demonstrated that compared with soluble antigen, alumin could induce similar antigen uptake by BMDC at a 100-fold lower dose (see Fig. 2B). Similarly, we observed the equivalent levels of YAE expression on cells incubated with 1 μg/ml EoGFP/alumin compared with 10 μg/ml EoGFP alone suggesting that alumin causes 10-fold increase in antigen presenting efficiency of DCs in vitro (see Fig. 2B). We next determined if the enhanced antigen uptake and presentation induced by alumin was dependent on the dose of the adjuvant used. Using a fixed amount of EoGFP (100 μg/ml), doses of alumin between 0.1 and 10 μg/ml produced a small, though significant increase in antigen uptake compared with soluble antigen (see Fig. 2C and D). This was reflected in a corresponding significant increase in presentation of
the Eo peptide. Higher doses of alum (100 and 1000 µg/mL) produced much greater increase in GFP signal in BMDC, however the impact on antigen presentation was more modest, with decreased presentation being observed between 100 and 1000 µg/mL alum (see Fig. 2C and D). The difference in antigen presence and presentation in BMDC implied that the presence of alum may affect the rate of antigen degradation.

3.3. Alum adjuvants increase antigen uptake, reduce degradation and sustain antigen presentation by DCs

To understand the role of alum in the kinetics of antigen uptake, processing and presentation by DCs we performed a pulse-chase experiment. We exposed BMDC to a 60 min pulse of antigen and examined antigen degradation and presentation over time. We found a greater proportion of cells were GFP positive (see Fig. 3A and B) and had a greater GFP signal, as determined by assessment of MFI (data not shown), following treatment with EαGFP adsorbed to alum compared with exposure to EαGFP at each chase period tested. Following exposure to soluble antigen, both the GFP signal and proportion of cells that were GFP positive returned to background levels within 24 h. The rate of GFP decay in cells exposed to antigen formulated in alum, was slower and the GFP signal was sustained up to 72 h following exposure (see Fig. 3B), demonstrating that intact antigen was degraded more slowly and persisted for longer in the presence of alum.

This would suggest that alum may also affect the rate of antigen presentation by BMDC. In keeping with this observation, formulation of antigen with alum decreased the fraction of YAE-positive BMDC at 0 h compared with soluble antigen (p < 0.01) which then gradually increased to an equal level as induced by EαGFP treatment at 24 h (p > 0.05) (see Fig. 3A and C). Alum significantly enhanced antigen presentation at 48 h (p < 0.001) which was
Fig. 2. Alum efficiently targets DCs by enhancing both accumulation and presentation of antigen by DCs in vitro. (A) BMDCs (2 × 10^6/5 mL) were incubated with Eα (EαGFP) (0, 1, 10, 100 μg/mL) and EαGFP adsorbed to alum (100 μg/mL) for 24 h and using the gating strategy in Fig. 1A, levels of GFP and/or Yαe on CD11c positive populations were analysed. (B) The levels of GFP (left) and Yαe (right) on CD11c positive populations were analysed and the line graphs were made by the scatter dot plots and joining the mean values. Bonferroni post test (two way ANOVA) were used to compare the level (mean ± SEM of triplicate cultures) of GFP or Yαe between EαGFP- or EαGFP + alum-treated groups at particular dose of EαGFP. ***p < 0.001. Data shown is representative of three independent experiments. (C) BMDCs (2 × 10^6/5 mL) were incubated with EαGFP (0 and 100 μg/mL) and EαGFP adsorbed to alum (0.0, 0.1, 1.0, 10.0, 100.0 and 1000.0 μg/mL) for 24 h and using the gating strategy in Fig. 1A, levels of GFP and/or Yαe on CD11c positive populations were analysed. (D) The levels of GFP (left) and Yαe (right) on CD11c positive populations were analysed and the line graphs were made by the scatter dot plots and joining the mean values. Tukey post test (one way ANOVA) was used to compare the level (mean ± SEM of triplicate cultures) of GFP or Yαe between control (alum; 0.0 μg/mL or EαGFP; 100 μg/mL)-treated group and increasing concentrations of alum. ***p < 0.0001, **p < 0.001, *p < 0.01, ns = not significant. Data shown is representative of three independent experiments. Eα: EαGFP; A: alum.
Fig. 3. Alum enhances antigen uptake, reduces antigen degradation and maintains antigen presentation by DCs in vitro. (A) BMDCs (3 × 10⁶/5 mL) were pulsed with EsGFP (100 µg/mL) or EsGFP adsorbed to alum (100 µg/mL) for 1 h, then incubated for different chase periods (0, 24, 48 and 72 h). Using the gating strategy in Fig. 1A, levels of GFP and/or Yae on CD11c positive populations were analysed. The line graphs show the % of GFP (B), % of Yae (C), % of GFP+Yae+ positive cells (D) and % of GFP+Yae− (E) in different treatment groups. Data have been presented as the mean ± SEM of quadruplicate samples. Bonferroni post test (two way ANOVA) was used to evaluate p-value by comparing the proportion (mean ± SEM) of cells positive for either GFP and/or Yae between EsGFP- and EsGFP + alum-treated groups at specific chase period. ***p < 0.001, ns: not significant. Data shown is representative of two independent experiments.

sustained up to 72 h (p < 0.001) (see Fig. 3A and C). Similar trends were observed while analysing the data on the basis of MFI of the Yae (data not shown). We also examined the BMDC that were GFP positive for presence or absence of antigen presentation as detected by Yae staining. While the GFP+Yae− population became undetectable within 24 h chase when pulsed with soluble EsGFP, these cells remained detectable for up to 72 h following alum/EsGFP treatment (see Fig. 3E). Very few cells were GFP+Yae+ following a 24 h chase period, this was more apparent in BMDC treated with soluble antigen than EsGFP adsorbed to alum (see Fig. 3D). This suggests that BMDC do not appear to transition through a GFP+/Yae+ population when moving from GFP positivity to Yae presentation and furthermore, that alum slows down antigen processing and presentation.

In summary, as well as targeting antigens to DC, data from these pulse chase experiments suggest that alum decreases the rate of antigen degradation resulting in increased duration and magnitude of antigen presentation by BMDC.
3.4. **Alum increases MHC class II expression in DCs in dose-dependent manner**

While alum may act to make processed peptides available for longer, to mediate increased antigen presentation would presumably require increased cell surface MHC class II expression. We therefore analysed levels of MHC class II expression on BMDC incubated with different concentrations of alum (see Fig. 4A and B). We found that expression of MHC class II was dependent on the dose of alum used, with both the highest proportion of the cells positive for MHC class II molecules and the highest expression level of MHC class II molecules in the cells treated with 100 μg/mL of alum (see Fig. 4B). BMDC treated with 1000 μg/mL had lower MHC class II expression than the cells treated with 100 μg/mL alum, but higher than other doses used in the experiment indicating alum at high doses sustain MHC class II expression on the surface of DCs (see Fig. 4B). A similar dose response was seen for Yae expression in Fig. 2D, suggesting a link between increased antigen presentation and increased availability of MHC class II molecules following alum treatment in vitro.

4. **Discussion**

In the current study we have applied the previously described EoxGFP/Yae system [21,24] to directly track antigen internalisation, degradation and presentation in BMDC and the impact that alum adjuvants have on the magnitude and kinetics of these processes. The fluorescent protein moiety in the chimaeric EoxGFP antigen allowed tracking of antigen uptake and degradation. This approach confirmed previous in vitro studies demonstrating the ability of alum to enhance internalisation of antigens by DCs [14,15], more importantly; we were also able to demonstrate that alum has a significant impact on the rate of degradation of antigen within DCs. While the GFP signal was completely extinguished within 24 h of administration of soluble antigen, formulation in alum allowed intact antigen to persist for up to 72 h. Degradation of antigens by lysosomal proteases is an essential step in liberating peptide antigens from proteins, and agents that interfere with this process, such as protease inhibitors or inhibitors of lysosomal acidificaion have been shown to reduce antigen presentation [26,27]. This would suggest that slowing of antigen degradation by alum may result in poorer peptide loading and antigen presentation on MHC class II molecules. However, by virtue of the ability of the Yae antibody to directly recognise Eox:MHCII complexes, we were able to show that alum actually enhances the duration of antigen presentation by BMDC from less than 24 h, observed with soluble antigen, to at least 72 h. In agreement with our data, previous work has demonstrated that limiting the susceptibility of antigens to lysosomal proteolysis actually acts to increase antigen presentation and immunogenicity [28]. In terms of adjuvant activity in vivo,
slowing down antigen degradation and increasing antigen persistence makes physiological sense. It takes hours for peripheral DCs to migrate to draining lymph nodes where naive cognate T cells are resident. Furthermore, functional interactions between DCs and T cells are thought to occur over the following 48 h or more [21] and blockade or interruption of this interaction is known to block the development of effective T cell responses [12,29,30].

The reduction in the rate of antigen degradation we observed could therefore lead to a temporal increase in availability of peptide for binding to MHCII in peptide loading compartments resulting in increased duration and magnitude of antigen presentation, as we also observed. However, it remains unclear at this point, exactly how this mechanism works. In the current study we also demonstrated that exposure to Alum/EαGFP induces increased cell surface MHC class II expression on BMDC. Similarly, previous studies have shown the high expression of MHCII molecules following alum treatment in human peripheral blood mononuclear cells in vitro [16–18].

Interestingly, in the current study the dose response of alum induced MHC class II expression was similar to that observed when detecting Eα:MHCI complexes using the YAE antibody. Previous studies have demonstrated that inhibition of lysosomal proteases enhances the stability of p:MHCI complexes and leads to increased accumulation of MHCI complexes on the DC surface [31]. Therefore, if alum was to block lysosomal proteolysis, as suggested by the antigen persistence described above, this would explain the increased cell surface MHCI expression, although further studies would be required to validate this hypothesis. In summary, we have identified how alum modulates the following key steps leading to antigen presentation that could underpin adjuvant function (see Fig. 5). Firstly, formulation of antigen in alum results in increased antigen internalisation by BMDC in vitro, consistent with the hypothesis that alum acts as an antigen delivery system (Fig. 5A). Subsequently, we have demonstrated that alum slows protein degradation, presumably increasing the duration of peptide availability intracellularly (Fig. 5B). Finally, we have demonstrated that alum enhances magnitude and duration of expression of p:MHCI complexes on the DC surface, with an accompanying increase in MHCI expression (Fig. 5C). These consequences may underlie the generation of the long lasting T cell responses in vivo.

Conflict of interest

The authors declare no financial or commercial conflict of interest.

Acknowledgements

This work was funded by grants from the Biotechnology and Biological Sciences Research Council and The Wellcome Trust.
References

[1] Brewer JM. (How) do aluminium adjuvants work? Immunol Lett 2006;102:10–5.
[2] Clements CJ, Griffiths E. The global impact of vaccines containing aluminium adjuvants. Vaccine 2002;20(Suppl. 3):S24–33.
[3] Villadangos JA, Ploegh HL. Proteolysis in MHC class II antigen presentation: who’s in charge? Immunity 2000;12:233–9.
[4] Watts C. Capture and processing of exogenous antigens for presentation on MHC molecules. Annu Rev Immunol 1997;15:821–50.
[5] Nakagawa TY, Rudensky AY. The role of lysosomal proteinases in MHC class II-mediated antigen processing and presentation. Immunol Rev 1999;172:121–9.
[6] Nelson CA, Vidavsky I, Viner NJ, Gross ML, Unanue ER. Amino-terminal trimming of peptides for presentation on major histocompatibility complex class II molecules. Proc Natl Acad Sci USA 1997;94:628–33.
[7] Trombetta ES, Ebersold M, Garrett W, Pyapaer M, Mellman I. Activation of lysosomal function during dendritic cell maturation. Science 2003;299:1400–3.
[8] Trombetta ES, Mellman I. Cell biology of antigen processing in vitro and in vivo. Annu Rev Immunol 2005;23:975–1028.
[9] Iezzi G, Karjalainen K, Lanzavecchia A. The duration of antigenic stimulation determines the fate of naive and effector T cells. Immunity 1998;8:89–95.
[10] Zinkernagel RM. Localization dose and time of antigens determine immune reactivity. Semin Immunol 2000;12:163–71, discussion 257–344.
[11] Lee WT, Pasos G, Cecchini L, Mittler JN. Continued antigen stimulation is not required during CD4(+) T cell clonal expansion. J Immunol 2002;168:1682–9.
[12] Obst r, van Santen HM, Mathis D, Benoist C. Antigen persistence is required throughout the expansion phase of a CD4(+) T cell response. J Exp Med 2005;201:1555–65.
[13] Schrum AG, Turka LA. The proliferative capacity of individual naive CD4(+) T cells is amplified by prolonged T cell antigen receptor triggering. J Exp Med 2002;196:793–803.
[14] Moorefield GL, Sokolovska A, Jiang D, HogenEsch H, Robinson JP, Hem SL. Role of aluminium-containing adjuvants in antigen internalization by dendritic cells in vitro. Vaccine 2005;23:1588–95.
[15] Mannhalter JW, Neychov HO, Zlabinger JG, Ahmed R, Eibl MM. Modulation of the human immune response by the non-toxic and non-pyrogenic adjuvant aluminium hydroxide: effect on antigen uptake and antigen presentation. Clin Exp Immunol 1985;61:143–51.
[16] Rimaniol AC, Gras G, Verdier F, Capel F, Grigoriev VB, Porcheray F, et al. Aluminum hydroxide adjuvant induces macrophage differentiation towards a specialized antigen-presenting cell type. Vaccine 2004;22:3127–35.
[17] Seubert A, Monaci E, Pissa M, O’Hagan DT, Wack A. The adjuvants aluminium hydroxide and MF59 induce monocyte and granulocyte chemoattractants and enhance monocyte differentiation toward dendritic cells. J Immunol 2006;178:5402–12.
[18] Ulanova M, Tarkowski A, Hahn-Zoric M, Hanson LA. The common vaccine adjuvant aluminium hydroxide up-regulates accessory properties of human monocytes via an interleukin-4-dependent mechanism. Infect Immun 2001;69:1151–9.
[19] Sokolovska A, Hem SL, HogenEsch H. Activation of dendritic cells and induction of CD4(+) T cell differentiation by aluminium-containing adjuvants. Vaccine 2007;25:4575–85.
[20] Rimaniol AC, Gras G, Clayette P. In vitro interactions between macrophages and aluminium-containing adjuvants. Vaccine 2007;25:6784–92.
[21] Itano AA, McSorley SJ, Reinhardt RL, Ehl BD, Ingulli E, Rudensky AY, et al. Distinct dendritic cell populations sequentially present antigen to CD4 T cells and stimulate different aspects of cell-mediated immunity. Immunology 2003;19:47–57.
[22] Rudensky A, Rath S, Preston-Hurlburt P, Murphy DB, Janeway Jr CA. On the complexity of self. Nature 1991;353:660–2.
[23] Rudensky A, Preston-Hurlburt P, Hong SC, Barlow A, Janeway Jr CA. Sequence analysis of peptides bound to MHC class II molecules. Nature 1991;353:622–7.
[24] Rush CM, Brewer JM. Tracking dendritic cells in vivo. Methods Mol Biol 2010;626:169–85.
[25] Lutz MB, Kukutsch N, Ogilvie AL, Rossner S, Koch F, Romani N, et al. An advanced culture method for generating large quantities of highly pure dendritic cells from mouse bone marrow. J Immunol Methods 1999;223:77–92.
[26] Vidard L, Rock KL, Benacerraf B. The generation of immunogenetic peptides can be selectively increased or decreased by proteolytic enzyme inhibitors. J Immunol 1991;147:1786–91.
[27] Watts C, Moss C, Mazzeo D, West MA, Matthews SP, Li DN, et al. Creation versus destruction of T cell epitopes in the class II MHC pathway. Ann NY Acad Sci 2003;987:9–14.
[28] Delamarre L, Couture R, Mellman I, Trombetta ES. Enhancing immunogenicity by limiting susceptibility to lysosomal proteolysis. J Exp Med 2006;203:2049–55.
[29] Celi S, Lemaitre F, Bousso P. Real-time manipulation of T cell-dendritic cell interactions in vivo reveals the importance of prolonged contacts for CD4+ T cell activation. Immunity 2007;27:625–34.
[30] Bajenoff M, Granjeaud S, Guerdier S. The strategy of T cell antigen-presenting cell encounter in antigen-draining lymph nodes revealed by imaging of initial T cell activation. J Exp Med 2003;198:715–24.
[31] Shiz JS, Ebersold M, Pyapaer M, Delamarre L, Hartley A, Mellman I. Surface expression of MHC class II in dendritic cells is controlled by ubiquitination. Nature 2006;444:115–8.