Sortase-mediated modification of DEC205 affords optimization of antigen presentation and immunization against a set of viral epitopes

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Sortase-mediated modification of αDEC205 affords optimization of antigen presentation and immunization against a set of viral epitopes

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A monoclonal antibody against the C-type lectin DEC205 (αDEC205) is an effective vehicle for delivery of antigens to dendritic cells through creation of covalent αDEC205–antigen adducts. These adducts can induce antigen-specific T-cell immune responses or tolerance. We exploit the transpeptidase activity of sortase to install modified peptides and protein-sized antigens onto the heavy chain of αDEC205, including linkers that contain nonnatural amino acids. We demonstrate stoichiometric site-specific labeling on a scale not easily achievable by genetic fusions (49 distinct fusions in this report). We conjugated a biotinylated version of a class I MHC-restricted epitope to unlabeled αDEC205 and monitored epitope generation upon binding of the adduct to dendritic cells. Our results show transfer of αDEC205 heavy chain to the cytoplasm, followed by proteasomal degradation. Introduction of a labile dipeptide linker at the N terminus of a T-cell epitope improves proteasome-dependent class I MHC-restricted peptide cross-presentation when delivered by αDEC205 in vitro and in vivo. We also conjugated αDEC205 with a linker-optimized peptide library of known CD8 T-cell epitopes from the mouse γ-herpes virus 68. Animals immunized with such conjugates displayed a 10-fold reduction in viral load.

antibody conjugates | vaccines | vaccine development | protein engineering

A monoclonal antibody against the C-type lectin DEC205, αDEC205, recognizes DEC205, a ~250-kDa surface protein on dendritic cells (DCs) (1, 2). Attachment of an antigenic moiety to αDEC205, either through chemical conjugation (3) or as a genetic fusion (4), allows its targeted delivery to DCs in vivo, and elicits potent immune responses even with very small amounts of the αDEC205 adduct. This response includes CD4 and CD8 T cells specific for the antigen attached to the antibody (5). Equipping αDEC205 with a suitable payload, as well as similar modifications of other antibodies directed against yet other surface proteins on professional antigen presenting cells (6), holds promise as a purely protein-based vaccine strategy that does not require replicating or infectious agents.

Most chemical conjugation methods one might consider for modification of αDEC205 lack precision, as they predominantly target exposed lysine or cysteine residues. Even the introduction of a single suitably exposed cysteine by mutagenesis, although it affords good site-specificity of labeling, still limits the range of possibilities for covalent attachment of peptides and proteins. Fusion of αDEC205 and the payload of interest can be attained genetically, but expression and purification of each new individual adduct needs to be explored and optimized. These limitations impelled us to develop an alternative method to install payloads of interest onto αDEC205 in a more flexible manner. An efficient and straightforward chemoenzymatic alternative with the use of sortase A (7) should allow not only the facile installation of a large diversity of antigens, but also an analysis of the mechanisms that underlie the desirable immunogenic properties of αDEC205 adducts, with particular emphasis on the mode of attachment and sequences flanking the antigen. This approach is unique in that it allows the use of nonnatural amino acids as linkers, as well as the selective introduction of chemical modifications on the payload, while leaving the αDEC205 portion otherwise unmodified.

Sortase A from Staphylococcus aureus enables the covalent linkage of proteins that contain a suitably exposed LPXTG motif to probes composed of N-terminal oligoglycines, modified with a cargo of choice: peptide, protein, nucleic acid, (glyco)lipid, or any other entity that can be provided in linkage to an oligoglycine peptide (8, 9). We show that the introduction of an LPXTG motif at the C terminus of the heavy chain of αDEC205 allows the attachment of a T-cell epitope, a fluorescent, or a biotinylated cargo of choice in a stoichiometric manner. This procedure, referred to as sortagging, affords delivery of any T-cell epitope or traceable payload to DEC205+ DCs in vitro and in vivo. Importantly, it allows the installation of a peptide that can be labeled independently from the protein to which it is attached, an essential attribute for investigating T-cell epitope processing.

We show that conjugation of peptides or a protein, such as GFP, to αDEC205 is achieved with efficiencies approximating 90%. The conjugated antibody was readily separated from the sortase enzyme and unincorporated probes, thus allowing rapid processing of numerous samples in parallel. We used sortagging to install a biotinylated class I MHC-restricted epitope on αDEC205 and unravel the sequence of events that leads to the generation of the epitope upon binding to DEC205. We investigated the factors that influence the presentation by DCs of a peptide conjugated to αDEC205 and show that the introduction of labile dipeptide linkers at the N terminus of a class I MHC-restricted epitope sortaged onto αDEC205 strongly affects the in vivo CD8 immune response upon immunization of mice with conjugated αDEC205 by favoring the generation of the final epitope in a proteasome-dependent manner. We used these findings to design and conjugate to αDEC205 a complex set of peptides corresponding to 19 known epitopes of mouse γ-herpes virus (MHV-68). Immunization with αDEC205 sortaged to the MHV-68 epitope set reduced viral burden upon subsequent infection with live MHV-68. Our study thus addresses the mechanism that underlies antibody-mediated targeting of antigens to DCs and exploits these findings to elicit a CD8 T-cell response that helps curtail a herpesvirus infection.

Results

Sortagging of αDEC205 with (Modified) Peptides or GFP. We modified the DNA construct encoding the heavy chain of αDEC205 (2, 4) to introduce an LPETG motif, required for sortase-mediated installation of payloads of interest, followed by a histidine


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tag (His$_6$, Fig. 1). The modified antibody was expressed in CHO cells and purified from the culture media. The sortase reaction was then performed under native conditions in physiological buffers, with no collateral chemical damage inflicted on αDEC205 or its cargo. We started by coupling peptides (containing an HA or a biotin tag) to αDEC205 and monitored the kinetics of the reaction by immunoblotting against both His$_6$ (input αDEC205 and sortase) and HA or biotin (desired product) tags (Fig. 2A). After 4 h of incubation, 90% of input αDEC205 was conjugated, and, after 20 h, the reaction was nearly complete, as judged from the loss of the His$_6$ tag (Fig. 2A). Gel filtration chromatography or purification by using protein G-coupled beads allowed complete separation of sortagged αDEC205 from sortase A (Fig. S1). αDEC205 could be conjugated to a protein-sized probe with similar efficiency. Fig. 2B shows the result of a sortase reaction in which we used GFP equipped with a five-glycine N-terminal extension as the nucleophile. Incubation of αDEC205 with this modified GFP and sortase allowed its conjugation to the αDEC205 heavy chain in excellent yield (>90%, Fig. 2B, Upper). Sortagging of αDEC205 with the biotinylated peptide or GFP neither inhibited antibody binding nor compromised GFP fluorescence (Fig. S2). The sortase method thus allows conjugation of (biotinylated) peptides or proteins with excellent efficiency and without compromising the binding specificity of αDEC205.

**Insertion of Dipeptide Linkers at N Terminus of T-Cell Epitopes**

Sortagging to αDEC205 Improves Presentation of Class I MHC-Restricted but Not Class II MHC-Restricted Epitopes in Vitro. αDEC205 has been used extensively as a vehicle for the delivery of T-cell epitopes to antigen-presenting cells as a means of evoking a T-cell immune response or to induce tolerance (5, 6, 10). We sortagged a single batch of αDEC205 with a larger set of peptides to explore the importance of the mode of linkage of a T-cell epitope to αDEC205. We measured presentation by DCs as assayd by activation of T cells of the appropriate specificity. We chose the ovalbumin peptide OVA$_{257-264}$ (SIQAVHAAHAEINEAGR) recognized by CD4 T cells from OTII transgenic mice (11, 12) to explore the influence of the flanking residues on presentation via class II MHC. We synthesized a set of sortase-compatible peptides containing three N-terminal glycines followed by the T-cell epitope, flanked by residues contained in intact ovalbumin or preceded by dipeptide linkers that have been used in antibody–drug conjugates (Table S1) (13, 14). We measured the capacity of DCs exposed to αDEC205 sortagged with OVA$_{257-264}$ peptide variants to induce proliferation of carboxyfluorescein succinimidyl ester (CFSE)-labeled CD4 T cells from ovalbumin-specific OTII T-cell receptor (TCR) transgenic mice. Exposure of DCs to αDEC205 sortagged with OVA$_{257-264}$ peptide induced robust proliferation, compared with incubation with αDEC205-HA. All OVA$_{257-264}$ peptide variants sortagged onto αDEC205 showed a similar dose–response curve in this assay (Fig. 3A). We conclude that the flanking residues of this class II MHC-restricted epitope do not affect the capacity of the αDEC205–peptide adducts to activate CD4 OTII T cells.

αDEC205 can induce CD8 T-cell stimulation by cross-presentation in vitro and in vivo (3). As cross-presentation involves steps in epitope processing and routing that likely include the cytoplasm, the exact manner of attachment of an epitope sortagged onto αDEC205 potentially affects its presentation via class I MHC. We synthesized a set of sortase-compatible peptides (Table S1) derived from the OVA$_{257-264}$ (SIINFEKL) peptide, an H-2Kb restricted epitope recognized by CD8 T cells from OTI mice (15). Exposure of bone marrow-derived DCs (BMDCs) to αDEC205 sortagged with SIINFEKL activated OTI CD8 T cells, based on the up-regulation of CD69 (Fig. 3B). We observed striking differences in the ability of αDEC205 sortagged with different peptides to activate T cells (Fig. 3B). BMDCs exposed to αDEC205 sortagged with OVA$_{257-264}$ preceded by dipeptide linkers stimulated approximately five times more OTI cells than
These results correlate with an increase in the number of proliferating cells in mice immunized with αDEC205-G3-FR-SIINFEKL compared with mice immunized with αDEC205-G3-SIINFEKL (Fig. 4B). There were two times more IFN-γ-producing OTI T cells in the spleen of mice immunized with αDEC205-G3-FR-SIINFEKL compared with mice immunized with αDEC205-G3-SIINFEKL (Fig. 4D). Proper linker design is therefore important to ensure delivery of T-cell epitopes: even minor modifications may have an outsized impact on the desired immune response.

Presence of Dipeptide Linker Favors Generation of Proteasome-Dependent Class I MHC-Restricted Epitope. Our results show that insertion of an FR motif immediately upstream of the antigen epitope attached to αDEC205 significantly increases the ability of DCs to stimulate antigen-specific CD8+ T cells. Different routes of antigen presentation have been described, involving transfer of the antigen to the cytoplasm or antigen processing in vacuoles with subsequent loading onto class I MHC (19). To

**Fig. 3.** In vitro stimulation of antigen-specific CD4+ and CD8+ T cells by DCs incubated with sortagged αDEC205. (A) DCs were incubated in vitro with αDEC205 sortagged with HA tag (negative) or OVA253-339-containing probes (Table S1) at various concentrations, washed, and incubated together with CFSE-labeled splenocytes from OTI transgenic mice at day 0. The histogram shows the percentage of CFSE-diluted OTI CD4+ T cells at day 4 for each concentration of αDEC205 adduct used. (B) BMDCs were incubated in vitro with medium alone (negative) or αDEC205 sortagged with OVA257-264-containing probes (Table S1) at various concentrations, washed, and incubated together with splenocytes from OTI transgenic mice. The histogram shows the percentage of CFSE-positive CD8+ OTI T cells for each concentration of αDEC205 adduct used after 16 h. Errors bars show SDs of three independent measurements; representative of three independent experiments.

BMDCs exposed to αDEC205 sortagged with most of the other peptides (Fig. 3B). We investigated two additional class I MHC-restricted epitopes: the H-2Ld restricted Rop7161-169 epitope (IPAAAGRRF) from *Toxoplasma gondii* (16), recognized by Rop7-specific transmural cells (17) and the H-2Dk restricted Gp33-41 epitope (KAVYNFATC) from lymphoctic choriomeningitis virus, recognized by P14 TCR transgenic mice (18), to assess the generality of these findings. BMDCs exposed to αDEC205 sortagged with the relevant peptides preceded by a dipeptide linker were significantly more potent in stimulating antigen-specific CD8+ T cell than constructs that lacked them (Fig. S3). The ability of DCs to stimulate antigen-specific CD8 T cells upon incubation with sortagged αDEC205 thus improves upon the introduction of a dipeptide linker preceding the epitope, as shown for three unrelated epitopes, presented by three different class I MHC products.

**Fig. 4.** OTI T-cell proliferation and activation in mice immunized with αDEC205 sortagged with G3-SIINFEKL or G3-FR-SIINFEKL. CFSE-labeled CD8 T cells from CD45.1 OTI transgenic mice were transferred i.v. into CD45.2 C57BL6 mice. At 24 h after cell transfer, mice were immunized with 0.75 μg of αDEC205 sortagged with G3-SIINFEKL or G3-FR-SIINFEKL together with 25 μg of cDNA and 50 μg of Poly I:C or left untreated. Transferred T-cell expansion and activation was measured 7 d after immunization. (A) CD8 vs. CD45.1 staining of splenocytes from control mice (Left), mice immunized with αDEC205-G3-SIINFEKL or αDEC205-G3-FR-SIINFEKL (Middle and Right, respectively). (B) Dot plots show CFSE vs. IFN-γ staining on donor OTI T cells from the spleen of control mice and mice immunized with αDEC205-G3-SIINFEKL or αDEC205-G3-FR-SIINFEKL in vitro stimulation with SIINFEKL peptide. (C) Histogram shows percentage of transferred OTI T cells in the spleen of control mice and mice immunized with αDEC205-G3-SIINFEKL or αDEC205-G3-FR-SIINFEKL, respectively. (D) Histogram shows percentage of IFN-γ positive cells in vitro stimulation with SIINFEKL peptide among donor OTI T cells from the spleen of control mice or mice immunized with αDEC205-G3-SIINFEKL or αDEC205-G3-FR-SIINFEKL, respectively. Errors bars show SD of control, n = 2; αDEC205-G3-SIINFEKL immunized, n = 4; αDEC205-G3-FR-SIINFEKL immunized, n = 4 (*P < 0.05, Student t test; representative of three independent experiments.

**Insertion of Dipeptide Linkers at N Terminus of T Epitopes Sortagged to αDEC205 Dramatically Increases CD8 Immune Response upon Immunization with αDEC205 Adducts.** Our results suggest that residues preceding a class I MHC-restricted epitope critically influence success of antigen presentation upon αDEC205-mediated delivery. To investigate whether the insertion of dipeptide linker would improve DCs T-cell immune response in vivo, we monitored expansion and IFN-γ production of OTI T cells upon immunization with αDEC205-G3-SIINFEKL vs. αDEC205-G3-FR-SIINFEKL. Immunization of mice with 0.75 μg of αDEC205-G3-SIINFEKL induced a modest increase in the percentage of OTI T cells in the spleen compared with unimmunized mice (Fig. 4A and C). In contrast, immunization of mice with 0.75 μg of αDEC205-G3-FR-SIINFEKL induced a significant increase in the percentage of OTI T cells in the spleen (Fig. 4A and C).

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investigate the role of cathepsins and proteasomes in the generation of the epitopes from dDEC205 adducts, we treated DCs with the cathepsin inhibitor Z-FA-FMK or with the proteasome inhibitor ZL3VS and addressed their ability to stimulate OTI T cells upon incubation with dDEC205-G3-SIINFEKL. Neither Z-FA-FMK nor ZL3VS inhibited the ability of SIINFEKL-loaded BMDCs to stimulate OTI T cells (Fig. 5A). However, the ability of ZL3VS-treated BMDCs to stimulate OTI T cells upon dDEC205-G3-SIINFEKL incubation was fivefold less than seen for control BMDCs (Fig. 5B). In addition, we observed a small but reproducible increase in the ability of Z-FA-FMK–treated BMDCs to stimulate OTI T cells upon incubation with dDEC205-G3-SIINFEKL (Fig. 5B). These data suggest that the proteasome contributes to the generation of DEC205-dependent class I MHC-restricted epitopes. Therefore, dDEC205-G3-SIINFEKL or a fragment derived from it must reach the cytoplasm.

To address whether dDEC205 traffics to the cytoplasm with its cargo, we monitored the fate of dDEC205 sortagged to G3-SIINFEKL* (a peptide modified to contain a single biotin on the side chain of its lysine residue) in BMDCs—treated with DMSO or ZL3VS—by immunoblot using streptavidin–horseradish peroxidase (HRP). Arrival of dDEC205-G3-SIINFEKL* in the cytoplasm was assessed by subcellular fractionation using perfringolysin O, a toxin that selectively perforates the plasma membrane and allows retrieval of the cytoplasmic fraction after a simple centrifugation step (20). Inclusion of the proteasome inhibitor ZL3VS does not affect binding of dDEC205-G3-SIINFEKL* to BMDCs (Fig. 5C). After 10 min of incubation, we detected a cytoplasmic fraction of dDEC205-G3-SIINFEKL*. In ZL3VS-treated BMDCs, the level of the cytoplasmic pool of dDEC205-G3-SIINFEKL* was increased twofold compared with control BMDCs (8.1 ± 3.1% and 4.1 ± 3.8%, respectively, as percentage of the total fraction). We monitored the distribution of cathepsin L, protein disulfide isomerase (PDI), and GAPDH (Fig. 5C) to demonstrate the efficacy of this fractionation protocol and to establish that subcellular organellar integrity had been maintained. After 45 min, the cytoplasmic fraction of dDEC205-G3-SIINFEKL* was barely detectable, but the organellar fraction (i.e., pellet) remained stable.

Altogether, our results suggest that a fraction of dDEC205 is internalized by DCs and reaches the cytoplasm, and that the proteasome is key to generate the class I MHC-restricted–dDEC205-conjugated epitope. Thus, the presence of an FR linker must favor proteasome-dependent generation of the (extended) epitope. To address this, we sortagged in silico a total of 235 known class I MHC-restricted T-cell epitopes. DMSO-, ZL3VS-, or Z-FA-FMK–treated BMDCs were incubated in vitro with medium alone (negative), with free SIINFEKL peptide (A), or with dDEC205-G3-SIINFEKL (B) at various concentrations, washed, and incubated together with splenocytes from OTI transgenic mice. Histogram shows percentage of CD8* positive CD8OTI T cells for each concentration of dDEC205 adduct used after 16 h. Errors bars show SDs of four independent measurements; representative of three independent experiments. (C) DMSO- or ZL3VS-treated BMDCs were incubated with dDEC205-G3-SIINFEKL* for 10 min on ice. The cells were washed (total 0′) and incubated for 10 and 45 min, at 37 °C, 5% CO2. Cells were perfringolysin O–treated to separate organelle (i.e., pellet) and cytoplasmic (i.e., supernatant) fractions. All fractions were analyzed by SDS/PAGE under reducing conditions and immunoblotted with streptavidin–HRP. The blot was subsequently probed with anti-PDI (luminal ER-resident protein), anti-cathepsin L (soluble lysosomal enzyme), and anti-GAPDH (soluble cytosolic protein) antibodies as loading and fractionation controls. An anti-mouse HRP (GAPDH) and anti-rabbit HRP (PDI and cathepsin L) were used as secondary antibodies. Five equivalents of supernatant were loaded to facilitate visualization. Lane P, cells not incubated with dDEC205-G3-SIINFEKL*. Background biotinylated protein band is marked with an asterisk. Representative of six independent experiments.

Protection Against MHV-68 Infection upon Immunization with dDEC205 Sortagged with Virus Epitope Library. MHV-68 is a rodent pathogen (22) used as a model for the human gammaherpesviruses Kaposi sarcoma-associated herpesvirus (HHV8) and EBV (HHV4) (23). We individually sortagged 19 known class I MHC-restricted epitopes (Table S1, probe 29–47) covering 14 ORFs of MHV-68 and containing a N-terminal GGGFFR motif onto dDEC205 (dDEC205-G3–FR-MHV-68) (24). C57BL/6 mice were immunized with 20 µg of total dDEC205-G3–FR-MHV-68 (~1 µg of dDEC205 adduct per epitope) in combination with Poly I:C and OAg40, or animals received free MHV-68 peptides together with Poly I:C and OAg40 at 14 and 7 d before intranasal infection with MHV-68. Immunization of mice with dDEC205-G3–FR-MHV-68 resulted in a reduction of lung viral titers by 10-fold upon subsequent challenge with MHV-68 (Fig. 6), whereas administration of free peptides was without discernible effect.
handles or aldehyde tags μα pfu of MHV-68. Histograms show pfu per gram of www.pnas.org/cgi/doi/10.1073/pnas.1214994110 = Sweeney et al. 8) 6 d gens suggests that transfer of the (extended) epitope from the structures such as DEC205, remains largely unknown. The inability of the C-terminal portion of the ER to the cytoplasm to the lumen of the ER or endosomal compartments is required (19), or that class I MHC molecules do not reach endolysosomal compartments in TAP-deficient DCs.

Our results show that, within minutes, a minor fraction of intact αDEC205 adduct is detectable in the cytoplasm. Several observations suggest that degradation of αDEC205 adducts involves the proteasome. First, at no time did we detect other streptavidin-reactive material corresponding to a C-terminal proteolytic fragment of αDEC205-G3-SIINFEKL (Fig. 5C). Second, degradation of the cytoplasmic pool of αDEC205-G3-SIINFEKL was partially inhibited by the proteasomal inhibitor ZL3, even though even a significant amount of αDEC205-G3-SIINFEKL was present in the organellar (i.e., pellet) fraction at 45 min, far less was detected in the cytoplasm (i.e., supernatant; Fig. 5C). This suggests spatial and/or temporal constraints on the transfer of αDEC205-G3-SIINFEKL to the cytoplasm. Whether protein-conducting channels of the type implicated in dislocation (27) or disintegration of a vacuolar compartment are involved in αDEC205 transfer will need further investigation.

We demonstrate that the insertion of a dipeptide linker at the N terminus of a class I MHC-restricted epitope increases its presentation by DCs in vitro and in vivo. The insertion of such motifs favors the generation of the final epitope in a proteasome-dependent fashion. We show here that αDEC205 conjugates bearing a defined dipeptide motif are fivefold more potent in inducing CD8 T-cell response upon incubation with DCs. If a strategy analogous to fusion of antigens to αDEC205 was ever considered for clinical applications, a fivefold difference in administered dose is meaningful, from both a manufacturing and a dosing perspective.

The engineering of antibody-based vaccines requires a platform that considers ease of comparison of potency. Conventional conjugation methods are less well suited to investigate this parameter in a high-throughput manner and could not easily have replaced sortagging. The method described here affords conjugation of large numbers of synthetic probes to an antibody substrate in a straightforward and robust fashion. It not only permits immunization against pathogens by using complex sets of peptides (Fig. 6), but, in an analogous manner, should also allow the induction of tolerance toward comparably large sets of epitopes through αDEC205-mediated differentiation of regulatory T cells (10). This might be of particular interest for autoimmune diseases in which self-antigens are still poorly defined.

The range and number of different synthetic class I or II MHC-restricted peptides that can be conjugated to αDEC205 is essentially limitless, including peptides with nonnatural amino acids, or modified with entities that cannot be encoded genetically, e.g., CpG oligonucleotides, fluorophores, lipids, (oligo) saccharides. The ability to evoke a CD8 T-cell response that reduces viral titers, by using a purely protein-based preparation, demonstrates the feasibility of this approach, as demonstrated here for MHV68.

Discussion

Sortase-mediated modification of antibodies is flexible and robust. It offers considerable advantages vs. genetic fusions and chemical crosslinking methods. First, unlike crosslinking approaches, it allows stoichiometric and site-specific installation of probes at the C terminus of the heavy chain of the antibody in a manner that does not perturb antigen binding. Second, it enables rapid conjugation of a single batch of antibody with numerous different peptides. In this way, without further purification, 49 different αDEC205 adducts were synthesized, each of which would have required a separate genetic construct and transfection if generated by conventional cloning and expression methods. Third, we report here the site-specific conjugation of peptides containing (modified) residues that are not template-encoded (e.g., citrulline, biotinylated lysine) and hence cannot be inserted through genetic fusion. Fourth, sortagging allowed us to install a traceable peptide (i.e., epitope) on αDEC205 and monitor its fate during antibody processing. This procedure can be extended to the installation of isotopically labeled peptide if needed. Fifth, it affords a practical alternative to genetic fusions for the installation of protein-sized fusion partners that may affect the conformation and secretion of one or both fusion partners, as well as heterooligomeric proteins, provided at least one of them has an exposed stretch of glycine residues. The sortase-mediated installation of “click” handles or aldehyde tags further extends the range of possibilities (9, 25).

Whether there is an upper limit to the size of incoming nucleophiles that can be used in a sortase-mediated transpeptidation reaction remains to be determined, but we have successfully used polypeptides of as much as ~100 kDa in reactions that effectuate head-to-tail protein–protein fusions (26).

The development of a sortase-compatible version of αDEC205 allowed us to optimize αDEC205-mediated antigen presentation in an iterative process, as well as to dissect the route of αDEC205-dependent antigen cross-presentation. Although it has been reported that αDEC205-mediated antigen cross-presentation is transporter of antigenic peptides (TAP)-dependent (3), the sequence of steps that lead to class I MHC-restricted epitope generation, especially when the antigen is delivered via specific surface structures such as DEC205, remains largely unknown. The inability of TAP-deficient DCs to cross-present αDEC205-conjugated antigens suggests that transfer of the (extended) epitope from the
incubated at 1 × 10^6 cells per milliliter of supplemented RPMI with sortagged αDEC205 or agonist peptide for 45 min at 37 °C and washed three times with PBS solution or supplemented RPMI. A total of 0.3 to 1 × 10^6 DCs were then seeded in supplemented RPMI in round-bottom 96-well plates (Corning), together with 1 to 2 × 10^5 CFSE-labeled (Invitrogen) T cells per milliliter of supplemented RPMI with sortagged αDEC205 or free peptide. When indicated, BMDCs were incubated with 20 μM of proteasome inhibitor Z-LEV-S for 1.5 h before and during incubation with sortagged αDEC205 or free peptide and throughout the assay.

In Vivo Experiments. Splenic CD8 T cells from C57/BL6 OTI mice were enriched by negative selection using magnetic beads (cat. 130-985-236; Miltenyi Biotech), labeled with CFSE (cat. 21888; Sigma), and transferred i.v. into 7- to 12-week-old C57BL6/6 mice. One day after T cell transfer, mice were immunized with sortagged αDEC205 together with 50 μg of Poly I:C (cat. P0913; Sigma) and 25 μg of αCD40 (FGK1.5). At 7 d after immunization, mice were killed, and T-cell proliferation was measured by monitoring CFSE dilution by flow cytometry. For measurements of IFN-γ production, 2 × 10^5 splenocytes were incubated for 4 h with OVA peptide at 1 μg/mL in 96-well U-bottom plates in complete RPMI medium and in the presence of brefeldin A (Golgi-Plug; cat. 555029; BD Biosciences). Cells were then stained by using a fixation/permeabilization kit (cat. 554722; BD Biosciences) according to the manufacturer's instructions.


Dissection Assay. For in vitro dissection analysis of αDEC205-G2/3-SIFK*+L in semicellants, 13 × 10^6 BMDCs were treated with 20 μM Z-LEV-S dispersed from a DMSO stock solution or with the corresponding volume of DMSO for 45 min at 37 °C, 5% CO2. The cells were then incubated with 35 μg/mL αDEC205-G2/3-SIFK*+L on ice for 10 min, washed with ice-cold PBS solution, and resuspended in RPMI media. An aliquot was taken, and the cells were immediately transferred to 37 °C, 5% CO2, for 10 and 45 min. Subcellular fractionation using perfringsolin O was performed as described previously (20).

Immunization and Infection of Mice. MHV-68 peptide probes were sortagged to αDEC205 in separate reactions as described earlier. Reactions were pooled after 4 h, and the final product was purified by gel filtration chromatography as described in SI Materials and Methods. C57/BL6 mice (age 7–12 wk) were immunized with 20 μg of sortagged αDEC205 or an equimolar quantity of free MHV-68 peptides together with 50 μg of Poly I:C and 25 μg of αCD40 (FGK1.5) 14 and 7 d before intranasal challenge with 10^6 pfu of MHV-68. Viral titers were measured in lungs 6 d after infection. Infections and viral titer measurements were performed as described elsewhere (28). All described procedures were approved by the Massachusetts Institute of Technology Committee on Animal Care.

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