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Lymph-borne CD8α⁺ dendritic cells are uniquely able to cross-prime CD8⁺ T cells with antigen acquired from intestinal epithelial cells

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Cross-presentation of cellular antigens is crucial for priming CD8⁺ T cells, and generating immunity to intracellular pathogens—particularly viruses. It is unclear which intestinal phagocytes perform this function in vivo. To address this, we examined dendritic cells (DCs) from the intestinal lymph of IFABP-TVA232-4 mice, which express ovalbumin in small intestinal epithelial cells (IECs). Among lymph DCs (LDCs) only CD103⁺ CD11b⁻ CD8α⁺ DCs cross-present IEC-derived ovalbumin to CD8⁺ OT-I T cells. Similarly, in the mesenteric lymph nodes (MLNs), cross-presentation of IEC-ovalbumin was limited to the CD11c⁺ MHCII⁺ CD8α⁺ migratory DCs, but absent from all other subsets, including the resident CD8α⁻ DCs. Crucially, delivery of purified CD8α⁺ LDCs, but not other LDC subsets, into the MLN subcapsular lymphatic sinus induced proliferation of ovalbumin-specific, gut-tropic CD8⁺ T cells in vivo. Finally, in 232-4 mice treated with R848, CD8α⁺ LDCs were uniquely able to cross-prime interferon γ-producing CD8⁺ T cells and drive their migration to the intestine. Our results clearly demonstrate that migrating CD8α⁺ intestinal DCs are indispensable for cross-presentation of cellular antigens and, in conditions of inflammation, for the initial differentiation of effector CD8⁺ T cells. They may therefore represent an important target for the development of antiviral vaccinations.

INTRODUCTION

Cross-presentation is the process by which exogenous antigens are displayed on major histocompatibility complex (MHC) class I molecules to CD8⁺ T cells and is necessary for the generation of cytotoxic T-cell (CTL) responses against tumors and intracellular pathogens, such as viruses. It has long been recognized that the capacity to cross-present is limited to a relatively small population of dendritic cells (DCs). In murine lymphoid tissues these cells are characterized by expression of CD8α.1 Cross-presenting lymphoid tissue CD8α⁺ DCs also have a high capacity for uptake of apoptotic cells,2 and, in conditions of inflammation, initiate the differentiation of naive CD8⁺ T cells into CTLs in a process known as cross-priming.3,4 Cross-presentation by non-lymphoid “peripheral” tissue DCs is less well understood. It was suggested that peripheral DCs are unable to directly cross-present antigen after migrating to the draining lymph nodes (LNs),5 and merely serve to transfer peripheral antigen to CD8α⁺ lymphoid tissue-resident DCs. More recently, a subset of CD103⁺ DCs, which share ontogeny with lymphoid tissue CD8α⁺ DCs, were shown to take up antigen in the lung and skin and, after activation, to cross-prime CD8⁺ T cells in the draining LNs.6,7 The identification of cells capable of cross-priming in the intestinal mucosa remains controversial. Intestinal macrophages are able to capture and cross-present soluble antigen8 but are absent from intestinal lymph9,10 and are therefore unlikely to initiate priming of naive T cells in mesenteric LNs (MLNs). Intestinal CD103⁺ lamina propria (LP) DCs can be separated into CD8α⁺ and CD8α⁻ subsets, both of which are capable of initiating CTL differentiation after incubation with soluble antigen in vitro.11 Although it was suggested that CD8⁺ intestinal DCs derive exclusively from Peyers’ patches and are...
unlikely to migrate to MLNs, we have demonstrated that CD8α+ DCs continuously migrate in intestinal lymph. In fact, they are abundant even in the intestine and lymph of mice lacking all intestinal lymphoid tissues, indicating that the majority of CD8α+ intestinal DCs derive from the LP of the conventional villus mucosa. In addition to the two subsets of CD103+ DCs, a detailed survey of intestinal lymph DCs (LDCs) revealed subsets of bona fide CD103− intestinal DCs. Notably, all CD103+ and CD103− LDC subsets were able to cross-present soluble antigen in vitro. It is not currently clear which of these DC populations is responsible for cross-presentation of intestinal antigen and priming of CTL responses in vivo. Crucially, soluble antigens have been used by the majority of studies to date; this does not represent the physiological source of antigen involved in the cross-priming of CTLs. Cross-priming of virus-specific CD8+ T cells requires presentation of cell-associated antigens. Therefore, it is of vital importance to unambiguously characterize the populations involved in the cross-presentation of cellular antigens and the initiation of CTL responses.

Here we use transgenic mice that express intracellular ovalbumin (OVA) only in mature intestinal epithelial cells (IECs), to assess the capacity of intestinal DC subsets to capture cellular antigens and prime CD8+ T cells. Interestingly, we find that cross-presentation of IEC-expressed OVA in the MLNs is entirely dependent on the CD103+ CD8α+ migratory DC population, and absent from all other intestinal and MLN DC subsets. Furthermore, we demonstrate that the CD103+ CD8α+ subset of migrating intestinal DCs is both necessary and sufficient to induce proliferation of gut-tropic CD8+ T cells in vivo in response to IEC antigen. Finally, we show that under inflammatory conditions the CD103+ CD8α+ subset of intestinal lymph DCs induces cross-priming of naive CD8+ T cells, driving their differentiation into interferon γ (IFN-γ)-producing effector cells. These data demonstrate, for the first time, the unique abilities of migratory intestinal CD8α+ DCs to acquire and transport intracellular IEC-derived antigen from the intestine and to directly drive the initial differentiation of naive T CD8+ T cells into effector cells.

RESULTS

IEC-expressed OVA is efficiently presented to CD8+ T cells

The majority of data on the cross-presenting ability of intestinal DCs concerns their uptake and presentation of soluble antigen. However, following infection with intracellular pathogens that do not infect antigen-presenting cells directly, cross-presentation of cell-associated antigens is essential. Hence we used IFABP-tOVA 232-4 (232-4) mice, which express OVA as an intracellular antigen exclusively in mature IECs. When transferred into steady-state 232-4 mice, transgenic OVA-specific OT-I CD8+ T cells proliferate and migrate to the intestinal LP, but do not mediate killing of OVA-expressing IECs. However, in the presence of an inflammatory stimulus, delivered by viral infection or addition of adjuvant, OT-I cells transferred into 232-4 mice differentiate into CTLs and mediate destruction of the intestinal epithelium. Similarly, we find that this differentiation of OT-I cells into CTLs also occurs after treatment of recipient 232-4 mice with R848, a Toll-like receptor 7 (TLR7) agonist. 232-4 animals that receive OT-I cells and R848 develop small intestinal inflammation characterized by weight loss, prodigious infiltration of CD8+ T cells (Figure 1b), destruction of the epithelial-cell layer (Figure 1c), shortening of the villi (Figure 1d), and high levels of IFN-γ in serum (Figure 1e). Therefore, 232-4 animals can be used to investigate mechanisms involved in the cross-presentation of IEC antigen, and the induction of effector CD8+ T-cell responses in vivo.

Only CD103+ CD11b− CD8α+ migrating DCs present IEC antigen to CD8+ T cells

Since IEC-expressed OVA is efficiently cross-presented to OT-I CD8+ T cells, we used the 232-4 mice to determine which DC populations are able to capture, transport, and present this antigen. In order to identify and purify all migratory DC populations in lymph, unequivocally and without bias, we purified by fluorescence-activated cell sorting the four distinct LDC subsets that are obtained after thoracic duct cannulation of mesenteric lymphadenectomized (MLNx) mice (Figure 2a). This technique allows the collection of all DCs that would normally migrate from the intestine to the MLNs, without contamination by DCs from other tissues. All four subsets of intestinal LDCs from wild-type mice cross-presented soluble OVA to OT-I CD8+ T cells (Figure 2b). In contrast, when the LDC subsets were purified from 232-4 mice and co-cultured with OT-I cells without addition of soluble OVA, only the CD103+ CD11b− DCs could cross-present IEC-associated antigen in vitro (Figure 2c).

Several reports suggest that the capacity to cross-present in LNs is limited to a subset of CD8α+ expressing DCs, which are assumed to be blood derived and LN resident. However, our results indicate that migrating lymph-borne CD103+ CD11b− DCs are able to directly present IEC-derived antigen to T cells. In order to compare the relative contributions of migratory and resident MLN DCs to cross-presentation, we purified the migratory (CD11c+ MHCIib) and LN-resident DCs (CD11c+ MHCIib) from the MLNs of 232-4 mice (Figure 3a), using a strategy that has been previously described. To ensure that plasmacytoid DCs did not contaminate these purified populations, B220+ cells were also excluded from the cell sorts. As we have previously observed in lymph, the migratory CD103+ CD11b− DCs express CD8α+ albeit at lower levels than the MLN-resident CD11b− DCs (Figure 3b). The purified migratory and resident MLN DCs were then cultured with OT-I T cells. We found that the ability to cross-present was entirely contained within the CD103+ CD11b− population of the migratory CD11c+ MHCIib MLN DCs, and was absent from the “resident” CD8α+ CD11b− DCs (Figure 3c,d). CD11b+ subsets of both migratory and resident DCs had no cross-presenting activity in these experiments. Consistent with these results, we also observed that when LP DCs from 232-4 mice were co-cultured with OT-I
cells, only the CD103^+ CD11b^- DCs were able to cross-present IEC-derived OVA to the OT-I cells (Supplementary Figure S1 online). In order to eliminate the possibility that 232-4 DCs can express and present endogenous OVA, we used a bone marrow chimera approach. In bone marrow chimeric mice where OVA is only expressed in the non-hematopoietic compartment, donor wild-type CD103^+ CD11b^- MLN DCs efficiently cross-presented OVA to OT-I cells. However, when 232-4 bone marrow was transplanted into wild-type recipients, donor MLN DCs were not able to induce OT-I proliferation (Supplementary Figure S2). Therefore, the induction of OT-I proliferation was entirely due to cross-presentation of IEC-derived OVA, and not due to OVA expression in the DCs themselves. Finally, in order to test whether we can directly observe IEC-associated antigen in any of the MLN DC subsets, we adapted a technique for detecting vesicles containing epithelial-cell cytokeratin in DCs. Discrete cytokeratin^+ inclusions were observed in flow-sorted DC cytospin preparations and were significantly more frequent in the migratory CD103^+ CD11b^- DCs compared to other MLN DC subsets (Figure 3e,f).

These results demonstrate that only a single subset of intestinal DCs—the migratory CD11c^+ MHCI^hi CD103^+ CD11b^- CD8x^+ DCs—is able to cross-present IEC-derived antigen in the MLNs.

Intestinal CD103^+ CD11b^- CD8x^+ migrating DCs are phenotypically and ontogenically similar to other cross-presenting DC populations

Cross-presenting DC populations in the spleen, LNs, and lung share several important characteristics, including the surface expression of DNGR-1^{19,20} and XCR1^{21,22} and a requirement for Bat3 and IRF8 in their development. In order to assess whether the cross-presenting migratory intestinal CD103^+ CD11b^- DCs also display these properties, LDCs were examined for the expression of a range of surface markers by flow cytometry (Figure 4a). All subsets of LDCs lacked the expression of the macrophage markers MerTK, CD64, and CD14 but expressed the DC-specific markers CD26, CD24, and...
CD272. Compared with other subsets of intestinal LDCs, the CD103+ CD11b− LDCs expressed the highest levels of CD272 and were unique in expressing high levels of CD8x, XCR1, and DNGR-1 (Clec-9A) and lacking expression of CD172a (Figure 4a). In order to examine whether migrating intestinal CD103+ CD11b− CD8x+ LDCs share ontogeny with cross-presenting populations in other tissues, we made use of Batf3−/− mice that lack lymphoid tissue-resident CD8x+ DCs and intestinal CD103+ CD11b− DCs.16 As expected, thoracic duct lymph from MLNx Batf3−/− mice is significantly depleted of the CD103+ CD11b− CD8x+ LDCs (Figure 4b). The CD103+ CD11b− CD8x+ subset of LDCs also expressed significantly higher levels of Ifr8 mRNA than the other lymph DC populations (Figure 4c). These results demonstrate that migrating intestinal CD103+ CD11b− CD8x+ LDCs share the ontogeny and many phenotypic characteristics with cross-presenting DCs in other tissues.

**Figure 2** Intestinal epithelial-cell-expressed ovalbumin is cross-presented by migrating CD103+ CD11b− lymph dendritic cells (LDCs). Thoracic duct cannulation was performed on 12-week-old mesenteric lymphadenectomized (MLNx) C57Bl6 or IFABP-tOVA 232-4 (232-4) mice and lymph collected for 16 h, on ice. Lymph cell populations were purified with fluorescence-activated cell sorting as shown in a. (b) 12,500 C57Bl6 LDCs of the indicated subsets were pulsed with ovalbumin, then extensively washed. DCs were then co-cultured with 105 carboxyfluorescein succinimidyl ester (CFSE)-labeled CD8+ T-cell proliferation was then assessed as above. (c) Histograms represent proliferation as assessed by CFSE dilution. (d) 232-4 LDC subsets were plated without the addition of soluble ovalbumin. The DCs’ capacity to drive CD8+ T-cell proliferation was then assessed as above. (e) Histograms represent proliferation as assessed by CFSE dilution. (d) The graph shows the mean percentage of divided T cells; each dot represents an independent experiment. Asterisks denote statistical significance (**P<0.01). MHC II, major histocompatibility complex class II molecules.

Migratory intestinal CD103+ CD11b− CD8x+ DCs are sufficient and necessary for cross-presentation of IEC antigen in the steady-state MLNs

Previous studies have shown that in the absence of an inflammatory stimulus, OT-I CD8+ T cells in 232-4 mice recognize IEC-derived OVA, proliferate, and acquire the capacity to migrate to the intestinal LP. However, they do not differentiate into IFN-γ-producing CTLs.14 To determine whether the CD103+ CD11b− CD8x+ 232-4 LDC subset alone was sufficient to cross-present IEC-derived antigen and drive this proliferation of naive CD8+ OT-I T cells in vivo, we used the approach outlined in Figure 5a. Briefly, carboxyfluorescein succinimidyl ester-labeled OVA-specific OT-I T cells were transferred intravenously into congenic wild-type mice. A day later, purified LDC populations from 232-4 mice were injected directly into the subcapsular lymphatic sinus of the MLNs (subcapsular injection). Three days later, LNs were collected and the transferred OT-I cells examined to determine whether any of the transferred DC subsets had induced their proliferation. Since no exogenous OVA was added in this system, any proliferation of OT-I T cells must result from the cross-presentation of IEC-derived OVA by the LDCs transferred into the MLNs from the 232-4 mice. OT-I cell proliferation was detected in the MLNs of mice that had received CD103+ CD11b− CD8x+ LDC, whereas all the other DC subsets were unable to stimulate OT-I proliferation (Figure 5b). This T-cell priming was localized, as no proliferation of transferred OT-I cells could be observed.

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responses to IEC-derived antigen under inflammatory conditions. Our data is also responsible for cross-priming CTL responses to IEC-derived antigen to OT-I T cells. It is possible that other DC populations may acquire cross-presenting and cross-priming capacities following activation. Indeed it has been proposed that LP-derived intestinal CD8α− DCs are more efficient than CD8α+ DCs at inducing CTL responses to soluble antigen after TLR activation. To examine whether the ability to initiate cross-priming of CD8α− T-cell responses to IEC-derived antigen remained restricted to the CD103+/CD11b− CD8α+ subset of DCs under inflammatory conditions, DCs were isolated from 232-4 mice after a single intraperitoneal dose of R848. CD103+ CD11b+ CD8α+ 232-4 DCs debilitating this subset were then purified and injected subcapsularly into the MLNs of wild-type recipients of OT-I cells. As in steady state, the subcapsular transfer of CD103+ CD11b− CD8α+ DCs induced marked proliferation of OT-I cells in the recipient MLNs, whereas LDCs that had been depleted of the CD103+ CD11b− CD8α+ subset were unable to induce OT-I proliferation (Figure 6A). The CD103+ CD11b+ CD8α+ LDCs from R848-treated 232-4 mice also induced IFN-γ production from the dividing OT-I T cells, although no IFN-γ was detected after transfer of LDCs depleted

in the peripheral inguinal LNs of the same mice (data not shown).

Next, we examined whether the T cells activated by the CD103+ CD11b− CD8α+ LDC in vivo acquired the characteristic gut-homing phenotype induced by intestinal and MLN DCs. Indeed, proliferating CD8α T cells in the MLNs showed increased expression of CCR9, an intestinal homing chemokine receptor (Figure 5B). Furthermore, CCR9+ CFSElim transferred CD8α+ OT-I T cells were readily detectable in the small intestinal LP of mice that had received CD103− CD11b− CD8α+ LPDCs, but not in recipients of other LDC subsets (Figure 5C). These data conclusively demonstrate that the CD103+ CD11b− CD8α+ subset of migrating intestinal DCs is the only migratory DC population capable of cross-presenting IEC-derived antigen to naïve MLN CD8α+ T cells in vivo.

After activation with a TLR7 agonist, migratory intestinal CD103+ CD11b− CD8α+ DCs induce effector CD8α+ T-cell responses to IEC-derived antigen

We next examined whether the same subset of intestinal DCs is also responsible for cross-priming CTL responses to IEC-derived antigen under inflammatory conditions. Our data clearly indicate that, in the steady state, migratory CD103+ CD11b− CD8α+ LDCs are the only cells able to cross-present IEC-derived antigen to OT-I T cells. It is possible that other DC populations may acquire cross-presenting and cross-priming capacities following activation. Indeed it has been proposed that LP-derived intestinal CD8α− DCs are more efficient than CD8α+ DCs at inducing CTL responses to soluble antigen after TLR activation. To examine whether the ability to initiate cross-priming of CD8α− T-cell responses to IEC-derived antigen remained restricted to the CD103+ CD11b− CD8α+ subset of DCs under inflammatory conditions, DCs were isolated from 232-4 mice after a single intraperitoneal dose of R848. CD103+ CD11b+ CD8α+ 232-4 DCs or DCs depleted of this subset were then purified and injected subcapsularly into the MLNs of wild-type recipients of OT-I cells.

As in steady state, the subcapsular transfer of CD103+ CD11b− CD8α+ DCs induced marked proliferation of OT-I cells in the recipient MLNs, whereas LDCs that had been depleted of the CD103+ CD11b− CD8α+ subset were unable to induce OT-I proliferation (Figure 6A). The CD103+ CD11b+ CD8α+ LDCs from R848-treated 232-4 mice also induced IFN-γ production from the dividing OT-I T cells, although no IFN-γ was detected after transfer of LDCs depleted
of this subset (Figure 6b). Importantly, despite inducing CD8+ T-cell proliferation in vivo, CD103+ CD11b– CD8α+ DCs from control 232-4 mice that did not receive R848 were also unable to drive the production of IFN-γ from transferred OT-I T cells (Figure 6b). These results demonstrate that activation of CD8α+ LDCs by a TLR agonist allows them to prime effector CD8+ T cells. In contrast, despite stimulation with a TLR agonist, the CD8α– LDC subsets were unable to cross-present IEC antigen or induce CD8+ T-cell effector differentiation, indicating that they are not involved in cross-presentation or induction of CTL responses. Finally, as in the steady state, transferred OT-I T cells were only observed in the intestinal LP of mice that had received subcapsular CD8α+ LDCs (Figure 6c).

Taken together, our results demonstrate that, unlike soluble antigen, intestinal cellular antigen is transported and cross-presented to MLN CD8+ T cells in vivo by a single subset of intestinal DCs, defined by their CD103+ CD11b– phenotype, both in the steady state and after TLR7-induced activation.

Figure 4 Characteristics of intestinal lymph CD8α+ dendritic cells (DCs). (a) Thoracic duct lymph cells from mesenteric lymphadenectomized (MLNx) C57Bl6 mice were stained for flow cytometry with phenotypic markers: MerTK, CD64, CD14, CD26, CD24, CD272, CD8α, XCR1, DNGR-1 (Clec-9a), and CD172a. Each plot shows staining of the four lymph DC (LDC) subsets and is representative of at least three independent replicates. The shaded histograms show staining of the total LDC population with the isotype control antibody. (b) Thoracic duct lymph was collected from MLNx Batf3–/– mice, and lymph cells were stained for flow cytometry. Dot plots show staining of LDC subsets. The graph shows the proportion of the CD103+ CD11b– DC subset as a percentage of total LDCs. Each dot represents an independent replicate. (c) LDC subsets were purified with fluorescence-activated cell sorting, the RNA extracted and reverse transcribed into cDNA. Expression of the IRF-8 mRNA was assessed by quantitative PCR. Results presented in arbitrary units (AU), normalized to the housekeeping gene cyclophilin A. In all panels asterisks denote statistical significance (*P<0.05, ***P<0.001).

Figure 5 Only intestinal lymph CD103+ CD11b– CD8α+ dendritic cells (DCs) cross-present intestinal epithelial cell antigen in vivo. (a) Schematic shows the experimental protocol. Five days after subcapsular injections of 50,000 fluorescence-activated cell sorting (FACS) purified lymph DC (LDC) subsets, mesenteric lymph nodes (MLNs) and the small intestine (SI) were harvested from recipient mice. Cells were then stained for flow cytometry. Transferred OT-I T cells were identified as CD45.1+ TCR Vα2+ in the MLNs (b) or SI lamina propria (LP) (c), and assessed for proliferation by carboxyfluorescein succinimidyl ester (CFSE) dilution. (b) For each injected LDC subset, FACS plots show the relative proportion of transferred OT-I T cells (CD45.1+ CD8α+) among TCRVα2+ cells (top left), CFSE dilution (top right), and CCR9 expression (bottom left) in the MLNs. The graph shows mean percentage of divided T cells; each dot represents an independent experiment. Asterisks denote statistical significance (**P<0.01). (c) FACS plots show the relative proportion of transferred OT-I T cells among TCRVα2+ cells (left) or CFSE dilution and CCR9 expression of transferred OT-I cells (right) in the LP, and are representative of two independent experiments. i.v., intravenous.
Figure 5  For caption see page 43.
DISCUSSION

Cross-presentation of cellular antigens by DCs is crucial for the priming of CTLs capable of killing tumors or virally infected cells. However, the identity of the DCs responsible for cross-presentation in the intestinal mucosa is unclear; several subtypes of intestinal DCs and macrophages are able to cross-present soluble antigen. By isolating DCs directly from lymph, we were able to analyze the function of all the DC populations that migrate from the intestine to the MLNs. All four subsets of migrating intestinal DCs can induce proliferation of antigen-specific OT-I CD8\(^+\) T cells in response to soluble antigen. However, to understand how CD8\(^+\) T-cell...
responses against viral and tumor antigens are initiated, it is important to understand which DCs present peripheral cellular antigens to CD8⁺ T cells. We therefore assessed the cross-presentation capacity of migratory DCs isolated from transgenic 232-4 mice, which express OVA exclusively in small intestinal IECs. In this system, without the addition of any soluble antigen, only the CD103⁺ CD11b− CD8α⁺ LDCs, but not the other LDC subsets, were able to induce proliferation and differentiation of naïve OVA-specific CD8⁺ T cells in vitro.

We show further that cross-presentation does not involve the transfer of antigens to resident CD8α⁺ DCs in the draining LNs, as has been suggested in other systems, as the only DCs from 232-4 MLNs capable of priming CD8⁺ T cells were CD103⁺ CD11b− migratory DCs. All other MLN DC populations, including the CD8α⁻/⁻-resident DCs, were completely unable to cross-present IEC antigen. It is therefore unlikely that antigen transfer from migratory to resident DCs plays a major role in the cross-presentation of IEC-derived cellular antigens during the induction phase of the response. As has been indicated previously, we confirm that migrating CD103⁺ CD11b− also express CD8α, but at a lower level than resident MLN DCs.

The CD103⁺ CD11b− CD8α⁺ population of migrating intestinal DCs shares the Batf3 and IRF-8-dependent ontogeny with cross-presenting populations in both lymphoid and peripheral organs, and expresses a very high level of surface expression of XCR1, a chemokine receptor associated with cross-presentation in mice and human DC subsets. Notably, the cross-presenting population of CD103⁺ CD8α⁺ DCs were the only subset of DCs migrating in mouse lymph that lacked the expression of CD172a (SIRPα). This is consistent with previous evidence that an analogous population of CD103⁺ CD172a⁻ LDCs in rat lymph can take up apoptotic IECs and transport them to MLNs. In line with this, CD103⁺ CD11b− CD8α⁺ LDCs express DNGR-1 (Clec-9a), which is involved in the uptake of dead cells and is expressed on cross-presenting populations in mice and humans. Furthermore, CD103⁺ CD11b− DCs that migrate from the lung selectively sample apoptotic cell antigen. Thus, it is highly likely that intestinal CD103⁺ CD11b− CD8α⁺ DCs and other migratory CD11b− DCs utilize similar mechanisms to sample cellular material, transport it to LNs, and present it to CD8⁺ T cells.

It has been suggested that CD103⁺ CD11b⁻ intestinal DCs mainly derive from the lymphoid tissues of the intestine, such as Peyers patches or isolated lymphoid follicles. However, we have previously shown that these cells are present in normal numbers in the intestine and lymph of mice lacking all lymphoid tissues, suggesting that they mainly derive from the LP. Further characterization of this LDC subset, including the expression of DNGR-1 and XCR1, support this interpretation; DCs expressing DNGR-1 (Clec-9a) can be found throughout human non-lymphoid tissues, including the LP, where they can be observed in the immediate vicinity of the intestinal epithelium. Furthermore, XCR1-expressing cells can be observed throughout the LP of the small intestinal villi (Richard Kroczek, unpublished data) as well as in other non-lymphoid tissues. Although we cannot rule out the possibility that some CD103⁺ CD11b⁻ CD8α⁺ LDCs originate in the lymphoid compartments of the intestine, it is likely that they largely represent LP-derived migratory DCs.

Crucially, CD103⁺ CD11b− CD8α⁺ DCs are also able to cross-present IEC antigen to CD8⁺ T cells in vivo. By adapting a technique described by the Pabst laboratory, we showed that subcapsular injection of CD103⁺ CD11b− CD8α⁺ LDCs, but not other DC subsets, from the lymph of steady-state 232-4 mice into the MLNs led to clonal expansion of OT-I T cells.

The capacity of intestinal DCs to induce gut tropism in a retinoic acid-dependent manner has been well characterized. However, it is not clear which intestinal DCs are primarily responsible for inducing this phenotype in CD8⁺ T cells. In contrast with an earlier report, we have recently shown that CD103⁺ CD11b⁻ CD8α⁺ DCs have high aldehyde dehydrogenase activity and can induce CCR9, a marker of gut tropism on T cells, with a similar efficiency to the other intestinal DC subsets. Here we show that, in vivo, cross-presentation by CD103⁺ CD11b⁻ CD8α⁺ DCs also induces CCR9 expression on responding OT-I T cells, and that, in this experimental system, fully differentiated CCR9⁺ OT-I cells are only detectable in the intestinal LP of mice after subcapsular injection of CD103⁺ CD11b⁻ CD8α⁺ LDCs, but not other LDC subsets.

Subcapsular transfer of steady-state CD103⁺ CD11b⁻ CD8α⁺ LDCs causes T-cell proliferation, but does not induce production of IFN-γ. This replicates the phenotype of OT-I cells transferred to 232-4 mice, which remain tolerant to OVA-expressing IEC under steady-state conditions but, as we show here, develop into CTLs when challenged with the TLR7 agonist R848. Importantly, CD103⁺ CD11b⁻ CD8α⁺ DCs isolated from the lymph of 232-4 mice treated with R848 are sufficient to induce differentiation of CD8⁺ T cells into IFN-γ-producing gut-tropic T cells after subcapsular transfer into MLNs. In addition, despite the suggestion that other DC populations may acquire cross-presenting and cross-priming capacities following activation, LDCs depleted of the CD8α⁺ subset were unable to cross-present IEC antigen.

Taken together, these data definitively demonstrate that the migrating CD103⁺ CD11b⁻ CD8α⁺ DCs play an essential and exclusive role in the initial cross-presentation of cellular antigens to naïve CD8⁺ OT-I T cells in the MLNs and can, in the presence of an inflammatory stimulus, initiate the differentiation of IFN-γ⁺ effector CD8⁺ T cells.

Our results suggest that plasticity of CD103⁺ CD11b⁻ CD8α⁺ intestinal DCs is the crucial mechanism that controls the balance between CD8⁺ T-cell-dependent homeostasis and active immunity. Like other DCs, CD103⁺ CD11b⁻ CD8α⁺ appear to be exquisitely sensitive to their environment. They maintain tolerance under steady-state conditions, but react to stimuli such as TLR ligands, and drive the generation of effector CD8⁺ T cells. Selective targeting of these processes in migratory CD103⁺ CD11b⁻ CD8α⁺ DCs may be a productive strategy for the generation of novel vaccines against mucosal pathogens.
METHODS

Animals. C57/B6e mice were purchased from Harlan and maintained in individually ventilated cages. 232-4 IFABP-1OVA mice (a kind gift from Professor Leo LeFrancois, University of Connecticut) and OT-I mice were bred and maintained under specific pathogen-free conditions at the Central Research Facility, Glasgow. All protocols involving live animals were approved by the local ethical committee and conducted under licenses issued by the UK Home Office. Batf3−/− mice were bred in specific pathogen-free conditions at the Experimental Biomedicine Animal Facility, University of Gothenburg.

Surgical procedures. Mesenteric lymphadenectomy and thoracic duct cannulation procedures were performed according to established protocols. Mesenteric lymphadenectomy was performed on 6-week-old C57/B6e or 232-4 male mice by laparotomy and blunt dissection. Six weeks later, mice were fed 0.2 ml olive oil to visualize the lymphatics and the thoracic duct was cannulated by the insertion of a polyurethane cannula (2Fr, Linton Instrumentation, Diss, UK). Lymph was collected in phosphate-buffered saline with 20 U ml−1 of heparin sodium (Wockhardt UK, Wrexham, UK), on ice, for up to 16 h. Subcapsular MLNs injections of cells were adapted from a technique for intralymphatic injections. Briefly, cells were resuspended in <10 μl of saline and injected into the subcapsular lymph sinus of the MLNs, using a BD Micro-Fine 30G needle (BD Biosciences, Oxford, UK). During surgical procedures the animals were maintained under inhalation anesthesia with isoflurane (Abbott Animal Health, Abbott Park, IL).

Generation of bone marrow chimeras. Eight-week-old mice received a lethal dose of radiation (10 Gy) and were reconstituted with 5 × 106 bone marrow cells by intravenous injection. Six weeks later, MLNs were harvested for DC isolation.

Reagents. Cells were cultured in RPMI 1640, supplemented with 100 U ml−1 penicillin, 100 μg ml−1 streptomycin, 2 mm 1-glutamine, 5% fetal calf serum (all from Invitrogen, Paisley, UK), and 50 μg/ml of R848 (Resiquimod; Invivogen, Billerica, MA) according to the manufacturer’s instructions. Minimum detectable concentration was <5 pg ml−1. RNA extraction. RNA was extracted using the MicroRNA kit (Qiagen, Hilden, Germany) or MACSQuant (Miltenyi Biotec)flow cytometers, or sorted and analyzed by the FACSAria cell sorter (BD Biosciences). For intracellular cytokine staining, cells were incubated for 4 h with phorbol 12-myristate 13-acetate, ionomycin, brefeldin A, and monensin (all from eBioscience). Cells were stained for surface marker expression and washed. Cells were then fixed in 4% paraformaldehyde, permeabilized, and stained for intracellular cytokine. Acquired data were analyzed using FlowJo software (version 9.3.1; Tree Star, Ashland, OR).

Microscopy. Dissected tissues were immediately fixed in neutral buffered formalin. They were embedded in paraffin, sectioned (5 μm), and stained with hematoxylin and eosin. Images were obtained using a light microscope at ×10 magnification and were analyzed, and archived using cell^B software (Olympus, Tokyo, Japan).

Cell isolation. Thoracic duct leukocytes were collected on ice in phosphate-buffered saline with 20 U ml−1 heparin, passed through a 40-μm cell strainer (BD Biosciences), and RBCs lysed with ACK (ammonium–chloride–potassium) lysis buffer (Sigma-Aldrich). Cells were stained and analyzed by flow cytometry or sorted by fluorescence-activated cell sorting using the FACSaria cell sorter (BD Biosciences).

Small intestines were flushed with Hank’s balanced salt solution with 2% fetal calf serum and the Peyers’s patches excised. The intestines were opened longitudinally and cut into 0.5-cm segments, which were incubated twice in Hank’s balanced salt solution with 2 μg/ml ethylenediaminetetraacetic acid at 37 °C while shaking for 20 min. Supernatants were discarded and the tissue digested with 1 mg ml−1 of collagenase VIII (Sigma-Aldrich) at 37 °C with shaking for 15 min. MLNs were macerated and incubated with 0.4 Wunsch units per milliliter of Blendzyme and 50 μg ml−1 of DNAse (both from Roche, Penzberg, Germany) for 45 min at 37 °C while shaking. Single-cell suspensions were passed through a 40-μm cell strainer and stained for flow cytometry.

Flow cytometry. Cell surface staining was performed in phosphate-buffered saline with 2% fetal calf serum and 10 mM ethylenediaminetetraacetic acid for 30 min on ice. Where biotin-conjugated antibody was used, cells were further stained with a streptavidin-fluorochrome conjugate for 15 min. Samples were acquired on LSRII (BD Biosciences) or MACSQuant (Miltenyi Biotec) flow cytometers, or sorted and analyzed by the FACSaria cell sorter (BD Biosciences). For intracellular cytokine staining, cells were incubated for 4 h with phorbol 12-myristate 13-acetate, ionomycin, brefeldin A, and monensin (all from eBioscience). Cells were stained for surface marker expression and washed. Cells were then fixed in 4% paraformaldehyde, permeabilized, and stained for intracellular cytokine. Acquired data were analyzed using FlowJo software (version 9.3.1; Tree Star, Ashland, OR).

Statistical analysis. For comparison of means between two groups, the data were analyzed by Student’s t-test. For comparisons involving more than two data sets, analysis of variance was used. P values < 0.05 were considered significant and Bonferroni post-test was performed on the data sets. Unless otherwise stated, asterisks denote significant difference of the indicated group from all other groups. All statistical analysis was performed using GraphPad Prism and Microsoft Excel.
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