Title: A wave of monocytes is recruited to replenish the long-term Langerhans cell network after immune injury.

Authors: Ivana R. Ferrer-*, Heather C. West-*, Stephen Henderson, Dmitry S. Ushakov, Pedro Santos e Sousa, Jessica Strid, Ronjon Chakraverty, Andrew J. Yates and Clare L. Bennett.

Affiliations:
1 Institute for Immunity and Transplantation, Division of Infection and Immunity, University College London, London NW3 2PF, UK.
2 Cancer Institute Department of Haematology, Division of Cancer Studies, University College London, London WC1E 6DD, UK.
3 Peter Gorer Department of Immunobiology, School of Immunology and Microbial Sciences, King’s College London, New Hunt's House, Newcomen St, London SE1 1UL
4 Division of Immunology and Inflammation, Imperial College London, Hammersmith campus, London W12 0NN, UK.
5 Department of Pathology and Cell Biology, Columbia University Medical Center, New York, NY10032, USA.

*Shared first authors

Correspondence: c.bennett@ucl.ac.uk.

One sentence summary: Immune damage in the epidermis leads to repair of the LC network by monocyte-derived cells that become long-term LC.
Abstract.

A dense population of embryo-derived Langerhans cells (eLC) is maintained within the sealed epidermis without contribution from circulating cells. When this network is perturbed by transient exposure to ultra-violet light, short-term LC are temporarily reconstituted from an initial wave of monocytes, but thought to be superseded by more permanent repopulation with undefined LC precursors. However, the extent to which this process is relevant to immune-pathological processes that damage LC population integrity is not known. Using a model of allogeneic hematopoietic stem cell transplantation, where allo-reactive T cells directly target eLC, we have asked if and how the original LC network is ultimately restored. We find that donor monocytes, but not dendritic cells, are the precursors of the long-term LC in this context. Destruction of eLC leads to recruitment of a single wave of monocytes that engraft in the epidermis and undergo a sequential pathway of differentiation via transcriptionally distinct EpCAM+ precursors. Monocyte-derived LC acquire the capacity of self-renewal, and proliferation in the epidermis matched that of steady state eLC. However, we identified a bottleneck in the differentiation and survival of epidermal monocytes, which together with the slow rate of renewal of mature LC limits repair of the network. Furthermore, replenishment of the LC network leads to constitutive entry of cells into the epidermal compartment. Thus, immune injury triggers functional adaptation of mechanisms used to maintain tissue-resident macrophages at other sites, but this process is highly inefficient in the skin.
Main text:

Introduction

Langerhans cells (LC) are unique mononuclear phagocytes that reside within the epithelial layer of the skin and mucosal tissues (1, 2), where they play a key role in regulating immunity at the barrier surface. Loss of LC-dependent immune surveillance leads to a break-down in skin tolerance, and increased susceptibility to infection (3). Within the skin, embryonic (e)LC differentiate from yolk sac and fetal common tissue macrophage precursors that seed the skin before birth (4, 5)(6)(7)(8). The density of the adult eLC network is established post-birth by a burst of local proliferation of differentiated eLC in response to undefined signals (9). Subsequently, the mature network is maintained by low levels of clonal cell division within adult skin (10-12). By contrast, oral mucosal epithelia are populated by LC-like cells that are continuously seeded from recruited monocytes and dendritic cell (DC) precursors (1). These cells resemble epidermal eLC transcriptionally and phenotypically, but show little evidence of proliferation in situ, and rather depend on recruitment of blood-derived cells to maintain the cellular niche (1, 13).

Ablation of eLC from genetically engineered mice leads to patchy repopulation of the epidermis (14), due to division of surviving eLC with some contribution from bone marrow (BM) cells (12, 15). In this non-inflamed context, the replenishment of the empty niche is characterized by the slow kinetics by which emerging LC expand to fill the epidermis, reflecting the quiescent nature of mature LC within the skin environment (3). By contrast, severe perturbation of eLC in the context of inflammation leads to recruitment of BM-derived cells into the epidermis, and repopulation of the empty niche (10, 16). The cellular mechanisms by which this occurs have largely been defined using models in which transient acute exposure of murine skin to UV irradiation leads to cell death within the epidermis and eLC replacement. Under these conditions, Gr-1- monocytes are recruited to the epidermis (17, 18). While some studies suggest that these monocytes can differentiate into long-term LC (17, 19), others have proposed a two wave model of eLC replacement, in which monocytes can persist for up to 3 weeks as ‘short-term’ LC-like cells,
but are superseded by undefined precursors which become long-lived replacement LC (18). Central to this model is the observation that LC require the transcription factor Id2 for their development and persistence as long-lived quiescent cells (18). But, it remains controversial whether Id2 is required for repopulation of the LC niche after UV-irradiation (20). The nature of the long-term LC precursors remains a key question in the field, with a number of studies suggesting the potential for dendritic cells (DC) or their precursors, to seed epidermal LC in the adult (21-25).

The resident macrophage population in most tissues is maintained by recruitment of Ly6C- classical monocytes from the blood (26). Ly6C- monocytes are short-lived, non-dividing cells once they have left the BM; however they show remarkable plasticity upon differentiation and recent studies have defined a dominant role of the local tissue niche in shaping differentiation of recruited cells (27, 28). One active area of investigation is whether monocyte-derived macrophages can transcriptionally and functionally replace the resident macrophage populations that were originally seeded from intrinsically distinct precursors at birth. In the steady state, genetic ablation of tissue resident cells results in the differentiation of Ly6C- monocytes into Kupffer cells (29) and alveolar macrophages (30) in the liver and lung, respectively, that show few differences from the cells they replace. By contrast, genetic ablation of microglia leads to repopulation by monocyte-derived cells that appear to fulfill the functional roles of their resident counterparts, but remain morphologically distinct, and continue to express monocyte-related genes (31, 32). Whether re-emerging skin LC, driven by immune injury and inflammation in the skin, also retain evidence of their cellular origin, and to what extent repopulating cells can become a long-term quiescent LC network, remain pertinent questions. Not least because monocyte-derived cells in other inflamed tissues remain transcriptionally and functionally distinct from resident cells (33, 34).

We have exploited a murine model of hematopoietic stem cell transplant to define the cellular mechanisms that control re-building of the LC network after immune-mediated pathology. We have previously shown that allo-reactive T cells infiltrate the epidermis in this model, wherein interaction with eLC leads to enhanced cytotoxic function and survival (35). Activated T cells
target recipient keratinocytes and eLC (16), leading to graft-versus-host disease (35). We demonstrate that T cell-mediated destruction of eLC leads to the recruitment of a single wave of monocytes that seed long-term monocyte-derived (m)LC, which are indistinguishable from the embryo-derived cells that have been replaced. We provide evidence for a surge of monocyte recruitment to the epidermis, but differentiation of these cells into mLC is a rate-limiting step, resulting in inefficient rebuilding of the mature LC network. In addition, the epidermal compartment is not re-sealed after entry of T cells, and remains open to circulating cells. Thus, immune injury triggers an adaptive process that converges with mechanisms to regenerate other tissue-resident macrophages, but this is highly inefficient in the skin.
Results

Immune injury leads the gradual replenishment of the epidermis with LC-like cells.

We and others have previously shown that transfer of CD8+ male minor histocompatibility antigen-reactive Matahari (Mh) T cells with BM transplantation (BMT) (35, 36) leads to infiltration of pathogenic T cells into target organs and the development of sub-lethal graft-versus-host disease (GVHD) (37). Intra-vital imaging by our lab demonstrated direct interactions between Mh T cells and host LC within the epidermis (35) and the onset of GVHD pathology 2 weeks post-transplant (35). Once in the epidermis, CD8 T cells kill host LC via Fas ligand-dependent cytotoxicity (16).

We determined the kinetics of LC turn-over following immune injury and eLC destruction in our transplant model (Figure S1). eLC are radio-resistant (16) and persisted in control mice that received BMT alone. A small population of donor BM-derived cells was evident in some mice 10 weeks post-transplant, but we observed significant variability between mice (Figure 1A top panel and B). Loss of eLC was complimented by the emergence of donor BM-derived CD11b-Langerin-LC-like cells in the epidermis 2 weeks post-transplant, which increased sharply in frequency and number between 2-3 weeks (Figure 1A bottom panel and B), concomitant with the peak of Mh T cell numbers in the epidermis (Figure 1C). The development of full donor LC chimerism was gradual and evident by week 10 post-transplant. At this time-point repopulating donor CD11b-Langerin-LC-like cells were phenotypically indistinguishable from host eLC in BMT controls according to markers defined in previous studies (Figure 1D) (18, 38)

DC lineage cells do not become long-term replacement LC.

The DC-like nature of LC has led to the suggestion that DC lineage cells may contribute to adult LC, and/or seed repopulating cells after damage (21, 23). This hypothesis has recently been supported by work demonstrating that circulating human CD1c- DC have the potential to become LC-like cells in vitro, but this has not been directly tested in vivo (24, 25). Therefore, we
investigated the possibility that DC lineage cells may contribute to LC repopulation after immune injury, by transplanting irradiated male recipients with a 1:1 mixture of BM from Vav-Cre.Rosa26LSL<sup>v<sup>−</sup> (Vav<sup>−</sup>) and Clec9a-Cre.Rosa26LSL<sup>v<sup>−</sup> Clec9a<sup>v<sup>−</sup></sup>) reporter lines. The Vav-Cre transgene is expressed by all hematopoietic cells (39) and provided an internal control for the development of BM-derived LC, while Clec9a-dependent YFP marked cells restricted to the DC lineage (40) (Figure 2A). 10 weeks later, an average of 37 ± 14.1% (s.e.m.) of splenic CD11c<sup>+</sup>MHCII<sup>+</sup> cells were derived from Vav<sup>−</sup> BM in the BMT + T group. Of the Clec9a<sup>v<sup>−</sup></sup>-derived cells, 36 ± 2.6% (s.e.m.) expressed YFP. By contrast, while there was a clear contribution of donor Vav<sup>−</sup> cells to repopulating LC after BMT with T cells, we did not detect any YFP LC (Figures 2B and C).

**LC repopulation is preceded by influx of donor CD11b<sup>+</sup> cells.**

Mature LC are identified by the unique concomitant expression of high levels of the cell adhesion molecule EpCAM (CD326) and the C-type lectin receptor, Langerin (CD207), which are simultaneously up-regulated upon differentiation of eLC (9, 19, 41). By contrast, ‘short-term’ LC, do not up-regulate EpCAM after UV irradiation (18). We observed the T cell-dependent accumulation of CD11b<sup>−</sup> cells within the epidermis (Figures 3A and B), that peaked 3 weeks post-transplant and before the shift to donor LC chimerism shown in Figure 1A. Phenotypic analysis of these cells demonstrated the presence of 3 populations sub-divided by expression of EpCAM and Langerin (Figure 3C): donor CD11b<sup>−</sup> cells contained 2 sub-populations that were either negative/low for both markers, suggesting recent arrival in the epidermis, or solely expressed EpCAM; by comparison CD11b<sup>+</sup> cells were EpCAM<sup>−</sup>Langerin<sup>−</sup> and therefore resembled mature LC. For this paper, we will refer to these populations as “CD11b<sup>−</sup>”, “EpCAM<sup>+</sup>” and “donor LC”, respectively. We observed a peak in the frequency and number of EpCAM<sup>+</sup> cells between 2-3 weeks post-transplant, preceding the gradual accumulation of donor LC over time (Figure 3D). These data strongly suggested a developmental trajectory whereby a single wave of epidermal
CD11b- cells repopulated the mature LC network. We reasoned that acquisition of LC-defining proteins would be consistent with the developmental transition of these sub-populations. Indeed, we observed the gradual loss of CD11b, and up-regulation of CD24 and DEC205 with differentiation of donor LC (Figure 3E).

**EpCAM**-**monocyte-derived cells are distinct from donor LC.**

Our phenotyping data suggested that incoming CD11b- cells differentiated into a unique EpCAM- intermediate before becoming donor LC, but it was possible that EpCAM- cells were already immature LC. To distinguish between these possibilities, we compared the transcriptional profile of EpCAM- cells to donor LC or other CD11b- populations in the skin and blood (Figure 4A and see Figure S2 for sorting strategy). Hierarchical clustering demonstrated that eLC and donor LC were interchangeable, but that EpCAM- cells clustered as a distinct population, and were more closely aligned to blood and dermal monocytes than mature LC (Figure 4B). Analysis of the genes that contributed to differences along the PC1 axis after principle components analysis (Figure S3A) demonstrated that EpCAM- cells were distinguished by the down-regulation, but not loss, of expression of genes associated with monocyte development and function (e.g. *Prr5, Ccl9, Fcgr3* and *Fcgr4, Trem3, Tlr7*) (Figure S3B), including *Treml4*, which is expressed by Ly6C- monocytes with the potential to become moDC (42). However, EpCAM- cells had not up-regulated genes associated with changes to cell structure, adhesion and signaling that defined donor LC (e.g. *Emp2, Kremen2, Nedd4, Ptk7*). Given the distinct clustering of monocytes/EpCAM- cells and eLC/donor LC, we directly tested the contribution of monocytes to emerging LC by transferring mixed congenic Ccr2- and syngeneic Ccr2- BM with T cells (Figure 4C). CCR2 is required for both egress of monocytes from the BM and entry into tissues (43). We found that only Ccr2- cells contributed to CD11b-, EpCAM- and donor LC sub-populations (Figures 4D and E), *demonstrating suggesting* a monocytic origin for these cells.
Ly6C+ monocytes mature into MHCII+ activated monocytes (or monocyte-derived DC) in the dermis (44). Therefore, we directly compared the outcomes of monocyte differentiation within different skin compartments using panels of genes associated with monocyte maturation and function described by Schridde and colleagues (45) (Figure 4F). This analysis demonstrated the clear divergence between Ly6C+ monocytes differentiating within the dermis or epidermis. EpCAM+ cells displayed a unique gene signature associated with tissue homeostasis and modulation of the epidermal niche by matrix metalloproteinases (*mmp12, mmp13, mmp14, mmp25*, but not *mmp2* and *mmp9* which are associated with egress of mature LC out of the epidermis (46)); phagocytosis and uptake of apoptotic cells (*cd9, axl, cd36, itgb5, itgav*); and activation of complement (*c1qa, c1qb, c1qc*). However, EpCAM+ also retained shared patterns of gene expression with moDC that suggested recent extravasation from the blood (*gpr35, itgai, ccr2*), and the potential to activate of T cells (*b2m, tapbp, cd80, cd40, fcgrt*), which was lacking from LC.

**Proliferation of monocytes and LC in situ combine to replenish the LC network.**

Our data implied that monocytes were sufficient to replenish the LC network, but Ly6C+ monocytes are short-lived non-cycling cells, while proliferation of eLC at birth and in adults determines the density LC within the epidermis (9, 12). Therefore, we considered the relative importance of recruitment versus proliferation of epidermal CD11b+ cells for the rebuilding of the LC network. We constructed mathematical models to quantify the flows between CD11b− to EpCAM+ populations and the donor mLC pool using time courses of Ki67 expression (Figure 5A and B). All epidermal populations showed evidence of active or recent cell division after transplant which decreased to homeostatic rates equivalent to eLC by 10 weeks (9.8 ± 1.49\% (s.e.m.) (9). Models of the flow from CD11b− cells to EpCAM+ were fitted simultaneously to the time courses of total cell numbers and Ki67 expression in the two populations. The kinetics of EpCAM+ cell numbers closely tracked that of the CD11b− population, suggesting that EpCAM+cells were short-lived and/or rapidly underwent onward differentiation (Table 1). We first explored whether the
flow of CD11b− and EpCAM+ cells into donor LC were consistent with a linear developmental pathway, as predicted by our experimental data. This was compared to the alternative scenario in which EpCAM+ cells were a ‘dead end’ population and wherein monocytes differentiated directly into LC, which we named the branched pathway (Figure S4A). The fitted predictions of the two models were visually similar (Figure 5C). Nevertheless, we found approximately 10-fold greater statistical support for the linear pathway, as measured by weights calculated using the Akaike Information Criterion (47). Strikingly, however, maturation of EpCAM+ cells was highly inefficient with only 4% becoming donor mLC (Figure 5D and Table 1). Gene set enrichment analysis of EpCAM+ cells compared to donor LC suggests that most EpCAM+ cells underwent apoptosis within the epidermis (Figure S4B).

Production of donor LC was initially dominated by recruitment from CD11b− and EpCAM+ cells but proliferative self-renewal replaced recruitment by week 10 when the mature LC pool had reached steady state (Figure 5E; Table 1). At this point, donor LC resided in the skin for approximately 10 weeks on average and divided once every 78 days (Table 1). These estimates were consistent with other observations of the rates of turnover and division of eLC in the steady state (10), and matched eLC doubling time within the unperturbed eLC network (12, 48). Fits were based on the assumption that the CD11b− and EpCAM+ cells died or differentiated at a constant per cell rate. However, we had to include density-dependent proliferation of donor LC in order to explain the waning of Ki67+ cells over time. Thus, division occurred more frequently at low cell densities (Figure 5F; Table 1).

Expression of Ki67 by CD11b− cells suggested that accumulation of LC precursors required local proliferation of undifferentiated cells. This hypothesis was supported by the over-representation of cell cycle pathways in EpCAM+ cells (Figure S4C). This gene signature was in contrast to dermal monocytes, which up-regulated pathways associated with innate receptors and T cell activation (Figure S4D). To pinpoint active cell division within epidermal cells in vivo, we injected EdU into mice 3 weeks after BMT with T cells, and analyzed the frequency of EdU+ cells 4
hours later. Within this window, we detected incorporation of EdU by cycling CD11b\textsuperscript{−}EpCAM\textsuperscript{−} cells in the epidermis, and less so in donor LC (Figures 5G and H).

**Long-term mLC are homologous eLC and up-regulate Id2.**

To determine how much of the eLC transcriptional profile was determined by origin we compared the transcriptional profiles of long-term (10 weeks) mLC and eLC from age-matched untreated mice in more detail. Correlation analysis and direct comparison of gene expression as a heat map demonstrated that donor mLC were virtually indistinguishable from eLC (Figure 6A and S5A), and the few up-regulated genes were dominated by functions associated with cell adhesion and motility, suggesting the positioning and establishment of mLC within the epidermis (Table S2). To understand whether emergence of long-term mLC depended on restoration of a steady state environment in the epidermis, we also compared eLC and donor mLC 3 weeks post-transplant, at which point chimerism was incomplete and both populations shared the same inflammatory environment. We again found that donor mLC were homologous to eLC (Figure S5B), but that eLC showed evidence of prolonged exposure to the inflammatory environment due to conditioning and T cell damage (Figure S5C). Consistent with this, eLC isolated from the epidermis 3 weeks post-transplant primed CD8 T cells more efficiently than donor mLC from the same environment (Figure S5D).

Given that monocyte-derived cells rapidly differentiated into quiescent, long-lived LC, we reasoned that this must require programming by lineage-defining transcription factors (LDTF), namely Runx3 and Id2 in LC. Thus, we first assess expression these genes in donor mLC and their precursors. Figure 6B shows the sequential up-regulation of *Runx3*, *Id2* and *Cbfβ2* (41), and down-regulation of monocyte-associated *Irf8*, as the cells became mLC. In addition, EpCAM\textsuperscript{+} cells showed evidence of early responsiveness to the dominant epidermal cytokine TGFβ (Figure S5E), and matured into LC-like cells upon culture with TGFβ *ex vivo* (Figure S5F). Therefore, these data suggested that LDTF, and responsiveness to TGFβ, are switched on in EpCAM\textsuperscript{+} cells within the
epidermal environment, before differentiation into mature LC. We next used an in vitro screen to identify the growth factors, in addition to TGFβ, that controlled Id2 expression and LC identity after differentiation from BM cells. We selected BMP7, CSF-1 and IL-34 based on expression of their cognate receptors (BMPR1a and CSF1R respectively) by EpCAM+ cells in vivo (Figure S5G) and their requirement for LC repopulation after UV-irradiation (43, 49, 50), and tested the impact of each factor on LC development in BM cultures (Figure S6) (20, 41, 51). IL-34, but not CSF-1 or BMP7 enhanced LC numbers (Figure 6C), and this was associated with the specific up-regulation of Id2 by LC in IL-34 cultures (Figure 6D). BM-LC were derived from cells that proliferated before maturation in these cultures (Figure 6E), mirroring cycling of CD11b+ cells in the epidermis. Cell division was not affected by addition of IL-34, which instead increased the survival of LC (Figure 6F).

Immune damage and loss of eLC opens the epidermal compartment.

Having considered LC repopulation at the cellular level, and demonstrated that monocytes differentiate into bona fide LC within the epidermis, we now considered the impact of immune pathology on the LC network and integrity of the epidermal compartment.

The kinetics of LC repopulation demonstrated that monocytes failed to completely replenish the LC network in most mice 10 weeks after transplant, suggesting a prolonged reduction in LC density. However, this decrease in LC numbers was also evident in mice that had received BMT alone, demonstrating that the slow-rate of division by mature LC ultimately dictated the speed at which the network was repaired, rather than LC origin. (Figure 7A). Confocal analysis of epidermal sheets revealed significant heterogeneity in the density of mLc in different fields of view (Figure 7B), but mLc tended to be smaller than eLC from BMT controls (Figures 7C). Notably, while mLc and dendritic epidermal T cells (DETC) were closely co-located within the epidermis, we found no difference in the frequency of mLc 3 weeks after BMT into Tcrbd- recipients that lacked all endogenous T cells (Figure S7A).
Given the smaller volume of mLC, we considered whether they were less integrated within the epidermis than eLC, and therefore migrated more readily in draining LN. To test this we topically applied FITC to the ear of mice that had received BMT 10 weeks earlier, with or without T cells. LN cells were divided into migratory and resident populations based on expression of CD11c and MHC II as published by others (40, 52, 53) and we determined the number of EpCAM-Langerin+LC in the migratory gate (Figure S7B). At this time point all migrating LC were derived from donor BM (Figure S7B), and therefore we compared their frequency to that of host eLC in mice that had received BMT without T cells. The frequency of FITC+ cells within LN LC was equivalent irrespective of LC origin (Figure 7D), demonstrating that, while donor mLC acquired the capacity to migrate to LN, this was not to a greater extent than eLC. It was evident from flow cytometry plots that mLC picked up less FITC than eLC from un-transplanted controls. Comparison of the median intensity of FITC+ cells demonstrated that this was indeed the case, but a similar effect was also observed by eLC in BMT controls (Figure 7E and F). Thus, mLC migrate to LN, but irradiation and BMT may also impact on the acquisition of topical antigen by LC.

We next considered how changes to the density of the LC network would impact on the entry of cells into the epidermis in absence of T cell-mediated injury. Thus, transplanted mice received a second round of total body irradiation with BMT (without T cells) and we tracked the origin of epidermal LC 8 weeks later (Figure 7G). Host eLC were replaced by donor mLC after BMT with T cells (Tx1 only), but not BMT alone (Tx2 only), as expected. However, the epidermis of mice that had received both transplants (Tx1 and 2) contained 3 populations of co-existing LC. These were identified as radio-resistant host eLC and donor mLC (Figure 7H Langerin.GFP−CD45.1− and Langerin.GFP+CD45.1− respectively) and new BM-derived LC (Langerin.GFP−CD45.1+)
Discussion

We show that a single wave of monocytes are recruited into the epidermis to replenish the LC network after T cell-mediated killing of eLC. We have used a murine model of haematopoietic stem cell transplantation and graft-versus-host disease to determine the mechanisms by which the resident LC network is replenished after T cell-mediated pathology in the epidermis. In this setting, monocytes become quiescent, self-renewing cells, that we have called monocyte-derived LC (mLC), and which acquire the capacity to migrate to draining LN. mLC are transcriptionally homologous to the eLC they replace, despite on-going T cell-mediated immune pathology in the epidermis. This finding suggests that the skin environment is atypical since monocytes that differentiate within other inflamed tissues remain transcriptionally distinct from their resident macrophage counterparts (33, 34), and microglia, which closely resemble LC in terms of their capacity to self-renew without contribution from circulating monocytes, are replaced by cells that retain a persistent monocytic signature (32).

Tissue-resident macrophages, including LC, are seeded from embryonic precursors before birth (5). While there is a clear consensus on the role of adult monocytes in maintaining and replenishing tissue macrophages in other organs, the nature of the precursor that repopulates LC in the skin has remained elusive and controversial. Previously, studies that addressed the nature of LC replacement in the epidermis have depended on the destruction of resident eLC by acute (15-30 minutes) exposure of ear skin to UV irradiation. However, these studies have produced conflicting data on whether Gr1+ monocytes persist in the epidermis of UV-treated mice to become LC (17)(18). This work has led to the concept that an alternative ‘long-term’ LC precursor was required to replenish the LC network after UV-induced damage, and studies using human cells have since invoked a role for blood DC as LC precursors (24, 25). By contrast, in our stem cell transplant model allogeneic T cells are recruited to the epidermis over a period of weeks, resulting in prolonged immune pathology and inflammation (35). Under these conditions monocytes can become long-term LC, and DC precursors do not contribute to the emerging LC network. It is
conceivable that transient exposure to UV irradiation compared to the prolonged inflammation caused by allo-reactive T cells may trigger different mechanisms of LC repopulation in the skin. However, the adoptive transfer experiments previously used to define monocytes as LC precursors after UV irradiation are challenging, and require injection of large numbers of cells into Ccr2 Ccr6 mice to reduce competition from endogenous cells (17, 18). We suggest that the physiological recruitment of monocytes from the BM in our model has revealed their role in the repair of the damaged eLC network after immune pathology.

Statistical analysis of our mathematical models favors the linear differentiation of CD11b− monocytes into EpCAM− precursors of mLC. However, this conclusion is specific to the models that we considered; the transient nature of the EpCAM− population and similar kinetics of CD11b− and EpCAM− cells mean that it remains possible that the branched pathway may also occur with EpCAM− cells as a developmental endpoint. We think this is unlikely based on our evidence that EpCAM− cells express an intermediate phenotype and LDTF compared to CD11b− monocytes and mLC, and culture of EpCAM− cells induces up-regulation of Langerin. Despite the surge of monocytes entering the epidermis, and proliferation of CD11b− cells in situ, we have identified a bottleneck with only 4% of CD11b−/EpCAM− cells becoming mLC. The reasons for this inefficiency remain to be determined. One possibility is that monocyte differentiation is an intrinsically inefficient process and may also occur for the generation of tissue-resident macrophages at other sites. Alternatively, monocyte-derived EpCAM−/Langerin− cells can be identified within the oral mucosa, wherein inflammation blocks the transition to mucosal LC (13, 54). Therefore, it is possible that continued inflammation in our model blocks differentiation of EpCAM− cells.

Entry of CD11b− cells into the epidermis triggers a burst of proliferation that has also been reported when phagocytic monocytes enter the skin after UV-irradiation (17). Subsequent to this, mLC divide at a rate that matches that of steady state eLC (12, 48). This concordance both validates our modeling approach, and reveals a developmental convergence of quiescent mLC with
their embryonic counterparts. It has been suggested that the density of tissue macrophage populations is controlled by mechanisms of quorum sensing in response to CSF-1 (55), but this has not been directly demonstrated experimentally. We found that density-dependent proliferation was required to fit mathematical models to our data, supporting the concept of quorum sensing within the epidermal niche. CSF1 and IL-34 compete for the CSF1 receptor (56). Given the dominance of IL-34, rather than CSF1 in the epidermal environment (57), and our *in vitro* data showing that IL-34 increases expression of *Id2* and promotes BM-LC survival, it is possible that IL-34 fulfills this function in the skin.

We have shown that the epidermal compartment is not resealed after immune-mediated destruction of eLC, and BM-derived cells continue to be recruited into the epidermis. It is notable that small numbers of donor CD11b+ cells constitutively enter the epidermis of transplanted mice in the steady state, as observed 10 weeks post-transplant (see Figure 3D). One possibility is that the inefficient repair of the mLC network reduces competition between established and incoming cells (55). In this sense, the LC network in the epidermis more closely resembles that of the oral mucosa (1, 13) after exposure to immune injury.

The activation of auto- or allo-reactive T cells, and destruction of tissue-resident cells can have profound impacts on the balance of immune cells within tissue compartments with long-term consequences for the control of infection and cancer at these sites. The skin is highly sensitive to such changes and is a major target organ for T cells in patients suffering from graft-versus-host disease following HSCT, and those receiving immune checkpoint blockade (58). However, we know little about the impact of immune injury in these patients on the regulation of immunity in the skin. Here we provide insights into the cellular mechanisms by which the LC compartment is replenished and maintained after damage, and demonstrate that immune injury triggers an adaptive process that converges closely with mechanisms to regenerate other tissue-resident myeloid cells.

**Materials and Methods**
**Study design.** The aim of this study was to define the nature of long-term repopulating LC, and to identify the cellular processes by which the network was repaired after immune injury. We used an *in vivo* model of LC replacement and measured changes to cell populations using flow cytometry and confocal microscopy. This was combined with mathematical modeling of the flow between epidermal populations, and RNA sequencing to determine changes in gene expression. Sample sizes were based on previous experiments, and the availability of genetically engineered donors. No outliers were excluded and the number of replicates and independent experiments is given in each figure. There was no randomization or blinding. Recipients were co-housed were possible.

**Mice.** C57BL/6 (B6) mice were purchased from Charles River and bred in house by the UCL Comparative Biology Unit. Langerin.DTR.EGFP mice (59) were originally provided by Adrian Kissenpfennig and Bernard Malissen. C57BL/6 TCR-transgenic anti-HY MataHari mice (60) were provided by Jian Chai (Imperial College London, London, UK). Ccr2− mice were a gift from Frederic Geissmann (61). All strains were bred in house at UCL. TCRbd− mice were generated by crossing Tcrd− (62) and Tcrb− (63) lines, and bred in house at Imperial College London Hammersmith campus. CD45.1− OT-I TCR transgenic mice were bred in house. All procedures were conducted in accordance with the UK Home Office Animals (Scientific Procedure) Act of 1986, and were approved by the Ethics and Welfare Committee of the Comparative Biology Unit, Hampstead Campus, UCL, London, UK.

**Bone marrow transplants.** Recipient male CD45.2 B6 mice were lethally irradiated (11 Gy total body irradiation, split into 2 fractions over a period of 48 hours) and reconstituted 4 hours after the second dose with 5x10^6 female CD45.1 C57BL/6 T cell-depleted BM cells and 2x10^6 CD4 T cells, with 1x10^7 CD8 Thy1.1· Matahari (Mh) T cells, administered by intravenous injection through the tail vein. CD4 and CD8 T cells were isolated by magnetic selection of CD4 or CD8 BM cells or splenocytes using the Miltenyi MACS system (QuadroMACS Separator, LS columns, CD4 [L3T4] MicroBeads, CD8a [Ly-2] MicroBeads; Miltenyi Biotec) according to the manufacturer’s
instructions, either to be discarded from T cell-depleted BM, or selected for injection of splenic T cells. In some experiments Langerin.DTR.EGFP recipients or BM donors were used to track Langerin+ host or donor LC respectively.

For secondary irradiation experiments, BMT recipients initially received BM from Langerin.DTR.GFP donors with CD4 and CD8 Mh T cells. 8 weeks later, mice received 11 Gy split dose total body irradiation and CD45.1+ C57BL/6 female T cell-depleted BM alone.

**Mixed chimera experiments.** BM from Vav-Cre.Rosa26LSLCre and Clec9a-Cre.Rosa26LSLCre donors was a gift from Caetano Reis e Sousa (Francis Crick Institute). Irradiated B6 male mice received a 50:50 mix of BM from the reporter mice with T cells. For CCR2 competitive chimeras, irradiated CD45.2+ Langerin-DTR.EGFP recipients received a 50:50 mix of CD45.2+ Ccr2+/+ and CD45.1+ Ccr2−/− BM with T cells.

**FITC painting.** The dorsal side of ear pinna were coated with 25µl of a 1:1 mixture of 0.5% FITC (Fluorescein isothiocyanate, Sigma-Aldrich, UK) in acetone and dibutyl phthalate (Sigma-Aldrich, UK). 72 hours later the draining auricular and cervical LN were harvested and analyzed.

**Generation of tissue single cell suspensions.** Epidermal single cell suspensions were generated as described (35). Split dorsal and ventral sides of the ear pinna were floated on 2.5mg/ml Dispase II (Roche) in HBSS 2% FBS for 15 hours at 4°C, followed by mechanical dissociation of the epidermal layer in PBS/1mM EDTA/1% FBS by mincing with scissors or using the GentleMACS tissue dissociator (Miltneyi Biotech). Epidermal cells were passed sequentially through 70µM and 40µM cell strainers to remove clumps of cells. Numbers of cells were calculated by addition of counting beads (Invitrogen) before staining of samples for flow cytometry, and normalized to 0.1g weight for both complete ears before processing. After separation from the epidermis, the dermis was minced into small pieces and digested with 250 U/ml collagenase IV (Worthington, USA) and 800U/ml DNasel (AppliChem, USA) at 37°C for 1 hour. Dermal single cell suspensions were then generated using the GentleMACS tissue dissociator (Miltneyi Biotech).
LN were teased apart with needles and digested in HBSS/4000U/ml Collagenase IV (Worthington) for 40 min at 37°C. Digestion was quenched with 10mM EDTA, and the cells were passed through a 40 μM cell strainer, washed, and resuspended in PBS/1mM EDTA/1% FBS.

For blood cells, erythrocytes were removed by hypotonic lysis with distilled water, and cells then resuspended in PBS/1mM EDTA/1% FBS.

**Bone marrow cultures.** BM was flushed from femurs and tibias of donors, and red blood cells lysed in 1ml ammonium chloride (ACK buffer, Lonza UK) for 1 minute at room temperature. Cells were washed and resuspended at 2.5x10^6/ml in R5 medium (RPMI 1640, Lonza, Switzerland), 5% heat-inactivated FBS (Life Technologies, USA), 1% L-glutamine (2 mM; Life Technologies, USA), 1% Pen-strep (100 U/ml Life Technologies, USA) and 50μM β-ME (Sigma-Aldrich, UK). 1ml of cell suspension was plated per well in tissue culture-treated 24 plates and supplemented with 20ng/ml recombinant GM-CSF and 5ng/ml TGFβ (Peprotech, USA). Cells were cultured at 37°C. Media was partially replaced on day 2 of culture, completely replaced on day 3, and the cells harvested on day 6. Some cultures were supplemented with combinations of 8μg/ml IL-34 (Generon, UK), 100 μg/ml BMP7 (R&D systems, USA and 10μg/ml CSF-1 (Biolegend, USA) for the duration of the culture.

**Flow cytometry.** Cells were distributed in 96 well conical bottom plates and incubated in 2.4G2 hybridoma supernatant (containing αCD16/32) for at least 10 min at 4°C to block Fc receptors. For cell surface labeling, cells were incubated with fluorochrome-conjugated antibodies diluted in 100μl FACS buffer (PBS/1mM EDTA/1% FBS) at 4°C for at least 20 min in the dark: EpCAM (G8.8, eBioscience, USA), CD11b (M1/70, eBioscience, USA), CD45.1 (A20, BD Biosciences, Germany), CD45.2 (104, eBioscience, USA), MHC II I-A/I-E (M5/114.15.2, eBioscience, USA), CD11c (HL3, BD Pharningen, USA), CD24 (M1/69 BD Biosciences or Biolegend), CD205 (205yekta, eBioscience), B220 (RA3-6B2, BD Biosciences), Vβ8.3 TCR (1B3.3, BD Biosciences, Germany). To exclude lineage+ cells, we used a cocktail of CD3 (145-2C11, BD Biosciences,
Intracellular staining with αLangerin (CD207) antibodies (eBioL31, eBioscience, USA) was performed after cell surface immunolabeling. Samples were washed with FACS buffer, fixed in 100μl fixation solution (BD Cytofix/Cytoperm solution, BD Biosciences, UK) for 15 min at 4°C, washed twice with permeabilization buffer (BD Perm/Wash, BD Biosciences, UK) and incubated with 100μl αLangerin diluted in permeabilization solution at 4°C for 30 min in the dark.

Live cells were identified by exclusion of propidium iodide (unfixed cells) (Life Technologies, USA), or a fixable viability dye (eBioscience, USA or Life Technologies, USA). Multicolor flow cytometry data were acquired with BD LSRFortessa and BD LSR II cell analyzers equipped with BD FACSDiva v6.2 software (BDBiosciences, Germany). Fluorescence activated cell sorting was performed on a BD FACSARia equipped with BD FACSDiva v5.0.3 software (BD Biosciences, Germany). All samples were maintained at 4°C for the duration of the sort. Cells were sorted into PBS/2% FBS before resuspension in Buffer RLT (QIAGEN, USA) or directly into Buffer RLT with 1% 2-β-mercaptoethanol (Sigma, UK), disrupted through vortexing at 3200 rpm for 1 min, and immediately stored at -80°C until further processing. 2-3 biological replicates were obtained for each sample from at least 2 independent experiments, each containing a minimum of 4000 cells (pooling where necessary from multiple mice from individual experiments). Flow cytometry data were analyzed with FlowJo X v9 and 10 (LLC, USA), and cells were pre-gated on singlets (FSC-A versus FSC-H), and a morphological FSC/SSC gate.

**Measurement of cell proliferation.** In vivo. Epidermal single cell suspensions were immuno-labeled with surface antibodies, then fixed and permeabilized using the eBioscience intra-nuclear staining kit, before incubation with αKi67-v450 antibodies (SolA15, eBioscience, USA). Gates were set on non-proliferating cells and unstained cells. Alternatively, mice were injected with 100μg 5-Ethynyl-2’-deoxyuridine (EdU) i.p (Invitrogen, USA) and euthanized 4 hours later.

In vitro. Cells were pulsed with 10μM EdU on day 2 or day 5 of culture and the medium replaced...
For both *in vivo* and *in vitro* studies, cells were labeled for flow cytometry using the Click-iT Plus EdU Flow Cytometry Assay Kit (Invitrogen, USA), according to the manufacturer’s instructions.

**Immunohistochemistry.** 4mm biopsy punches were excised from the dorsal and ventral sides of split ears and incubated in 0.5M ammonium thiocyanate for 30 min at 37°C to remove the epidermis. Epidermal sheets were collected in eppendorf tubes, washed twice with PBS, and fixed with cold (stored at -20°C) acetone for 10 min. Sheets were washed twice with PBS, and blocked using 0.25% fish gelatin, 10% normal goat serum in PBS for 1 hour at room temperature. Sheets were then incubated with primary rat αLangerin (eBioscience), and αCD45.2-biotin (eBioscience) antibodies (both diluted 1:100 in blocking buffer), and incubated for 1 hour at room temperature. Sheets were washed 4 times in PBS, and incubated with goat αRat-Alexa 647 (Jackson ImmunoResearch Laboratories, 1:1000) and Streptavidin-e570 (eBioscience, 1:200) secondary antibodies diluted in blocking buffer for 1 hour at room temperature. Stained sheets were washed 4 times with PBS, and mounted on slides with ProLong™ Diamond anti-fade mountant (Invitrogen). Samples were imaged on a Nikon Ti inverted microscope, through a 20X objective for epidermal sheets (Plan Apochromat N.A. 0.75 W.D. 1mm) or 40X for sorted EpCAM+ cells (Plan Apochromat N.A. 0.95 W.D. 0.21mm), using a C2 confocal scan head with 488nm and 561/568nm optimized fluorescence filter cubes (Nikon Instruments, Tokyo, Japan). Multiple Z-stacks were acquired for each sample. Data was saved as nd2 files using FIJI/ImageJ for quantification.

**Confocal analysis.** Quantification of confocal records was performed using Definiens Developer software. Each channel in a record was processed with Gaussian filter followed by application of multi-resolution segmentation. Individual cells were detected based on their relative intensity in Langerin and CD45.2 channels. The cell volume (μm³ based on total number of voxels occupied by a cell) was measured for each cell type.

**Statistics.** All data, apart from RNAseq data, were analyzed using GraphPad Prism Version 6.00 for Mac OsX (GraphPad Software, USA). All line graphs and bar charts are expressed as means ±
SD. Significance was determined using a one-way ANOVA to measure a single variable in 3
groups, or two-way ANOVA for experiments with more than 1 variable, with post-tests as specified
in individual figures. A paired t-test was used in figure 6 to compare cells cultured under different
conditions. Significance was defined as *P < 0.05, **P < 0.01, ***P < 0.001. The false discovery
rate (fdr) was used to compare ontogeny pathways enriched in dermal or epidermal cells.

**Supplementary materials.**

Figure S1. Immune injury leads the gradual replenishment of the epidermis with LC-like cells.

Figure S2. Sorting strategies used to isolate cells for RNAseq.

Figure S3. EpCAM: monocyte-derived cells are distinct from donor LC.

Figure S4. Proliferation of monocytes and LC in situ combine to replenish the LC network.

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Table S1. Raw data file

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Supplemental Materials and Methods including mathematical modeling methods.
References:


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**Competing interests:** The authors declare that they have no competing interests.

**Data and materials availability:** The RNA sequencing data for this study have been deposited at NCBI Gene expression omnibus, GSE130257.

**Figure legends.**

**Figure 1. Immune injury leads the gradual replenishment of the epidermis with LC-like cells.**  
A. Male recipients received female BM alone (BMT) or with CD4 and CD8 (Matahari) T cells. Chimerism was measured within the mature CD11b-Langerin- LC population at different time points. Representative flow plots show the relative frequency of host (CD45.2) and donor (CD45.1)-derived cells.  
B. Graph showing the frequency ± SD of donor LC in mice receiving BMT with (circles) or without (triangles) T cells. Significance was determined with a 2-way ANOVA, ***P<0.001. Data are pooled from 2 independent experiments for each time point (n=5-10).  
C. Graph shows the number ± SD of Vβ8.3 Matahari (Mh) T cells in the epidermis over time, per 0.1g total ear tissue (n=7-8).  
D. Top - representative histogram overlays show the expression of LC-associated proteins on donor-derived LC (from mice that received BMT + T cells) or host eLC (BMT alone), 10 weeks post-transplant. Bottom - summary data showing the median fluorescent intensity (MFI) for each sample. H = host, D = donor, each symbol is one mouse. Data are pooled from 2 independent experiments (n=8), and representative of >3 different experiments.
**Figure 2.** DC lineage cells do not become long-term replacement LC.  
A. Schematic showing the experimental procedure. Male mice received female BMT with T cells. BM was composed of a 1:1 mixture of cells from syngeneic female Clec9a−/− and Vav−/− mice. 10 weeks later splenocytes and epidermal LC were assessed for the relative contribution of cells expressing Tomato (Tom) or YFP. B. Representative contour plots showing gated CD11c+MHCII+ cells in the spleen or CD11b+EpCAM−Langerin− LC in the epidermis of mice that received BMT with or without T cells. C. Summary bar graphs showing the frequency of red Tom- or yellow YFP- cells within splenic CD11c+MHCII+ (left) or epidermal EpCAM−Langerin− (right) cells in mice receiving BMT alone (open bars) or BMT with T cells (filled bars). Bars show the mean and range of data points. Data are pooled from 2 independent experiments, and analyzed using a 2-way ANOVA, ***P<0.001 (n = 5-6).

**Figure 3.** LC repopulation is preceded by a single wave influx of CD11b- cells. 
Mice received BMT with T cells, and the epidermis was analyzed at different time points. A. Left - dot plot shows the gating of single CD11b+−−−CD45.1+ donor myeloid cells. Right - summary graph showing the frequency ± SD donor CD11b- cells in mice receiving BMT alone (triangles) or BMT + T cells (circles) (n= 6-7). B. Graph shows the number ± SD per 0.1g total ear weight of donor CD11b- cells (n=5-10). C. Representative contour plots at 3 weeks showing 3 distinct subpopulations within single CD11b-−−−CD45.1+ cells. D. Summary graphs showing the frequency ± SD (left) and number ±SD (right) of cells within each of the gated populations shown in C: Circles CD11b−− (EpCAM−Langerin−); squares EpCAM; triangles donor LC (EpCAM−Langerin−) (n=7-8). Data are pooled from 2 independent experiments and analyzed by 2-way ANOVA for frequency; significance for numbers was calculated with a 2-way ANOVA with Tukey’s multiple comparisons test. E. Top - representative histogram overlays show surface expression levels of LC-defining proteins in the gated donor populations 3 weeks post-transplant. Bottom - graphs show summary data for the median fluorescent intensity (MFI). Symbols represent individual samples, analyzed
using a repeated measurements 1-way ANOVA. Data are pooled from 2 independent experiments per time point (n=6). *P<0.05, **P<0.01, ***P<0.001.

**Figure 4.** EpCAM+ monocyte-derived cells are distinct from donor LC. A. Schematic showing the populations of cells and phenotypic markers used to isolate cells for sequencing. B. Dendrogram showing clustering of samples. C. Schematic illustrating competitive chimera experiments to test the requirement for monocyte-derived cells. Male mice received female BMT with T cells. BM was composed of a 1:1 mixture of cells from congenic wild-type (CD45.1>Ccr2+) or CCR2-deficient (CD45.2>Ccr2-) mice. Epidermal cells were analyzed 3 weeks later. D. Representative contour plots showing the frequency of wild-type or knock-out cells within gated donor epidermal myeloid cells (host cells were excluded at this time point by the use of Langerin.EGFP recipients, and exclusion of GFP+ LC from our analyses). E. Summary data showing the frequency of Ccr2-, donor cells within each population. Bar graphs show the mean and range of data points, data are pooled from 2 independent experiments (n=6). Percent of Ccr2- cells versus Ccr2+ in each population ***P<0.001, 1-way ANOVA. F. Heat maps showing relative gene expression of defined genes grouped into panels according to distinct functional processes. Blood monocytes (grey) n = 2, EpCAM+ cells (cyan) n = 3, donor LC (magenta) n = 3, dermal monocytes (grey) n = 3.

**Figure 5. Proliferation of monocytes and LC in situ combine to replenish the LC network.** Mice received BMT with T cells. Total numbers and Ki67 expression of epidermal cells were analyzed at different time points and described with mathematical models. A. Representative histograms show gating of Ki67+ cells in the EpCAM+ and donor LC populations 3 weeks after BMT with T cells. B. Graphs show the frequency ± SD (left) and number ± SD per 0.1g total ear tissue (right) of Ki67+ cells within gated epidermal populations. Circles CD11b-; squares EpCAM+; triangles donor LC. Data are pooled from 2 independent experiments per time
point ($n = 7-8$) and significance calculated using a 2-way ANOVA, **P<0.001. C. Data from the experiments shown in B. were described with mathematical models. Upper panels - Fitted, empirical descriptions of the timecourses of Ki67 and Ki67 CD11b+ cells. Middle panels - fits to the total numbers and Ki67- fraction of EpCAM+ cells, using the empirical descriptions of the CD11b- cell kinetics as a source. Lower panels - fits to timecourses of mature LC numbers and the Ki67- fraction using either CD11b+ (P1) or EPCAM+ cells (P2) as a source. D. The model of a linear development pathway had the strongest statistical support. Numbers indicate parameter estimates from the model. E. Graph showing the relative contribution of proliferation in donor LC to influx (with 95 percent confidence interval) over time. F. Graph showing the estimated mean interdivision time (with 95 percent confidence interval) of donor LC at different times post-BMT with T cells. Parameter estimates are displayed in full in Table 1. G. Mice received EdU 3 weeks after BMT with T cells. 4 hours later, the skin and blood were harvested and cells analyzed for incorporation of EdU. Representative contour plots show overlaid gated CD11b+ Langerin- (yellow) or CD11b- Langerin+ (magenta) populations in the epidermis, or Ly6C-CD115- monocytes in the blood. FMO is the fluorescent minus one stain without the EdU detection reagent. H. Summary graph showing the mean ± SD frequency of EdU+ cells in the different groups. Circles are individual mice, n=6. Data are pooled from 2 independent experiments. Mo. = blood monocytes, dLC = donor LC.

Figure 6. Long-term LC are homologous to eLC and up-regulate Id2. A. Correlation matrix comparing differentially expressed genes between and blood monocytes, mLc 10 weeks post-BMT + T cells, and eLC from age-matched controls. B. Graphs show the relative FPKM count normalized to the maximum value for different transcription factors from the RNAseq data. Significance was calculated with a 1-way ANOVA, blood Ly6C- monocytes n = 2, epidermal EpCAM+ cells n = 3, donor LC n = 3, age-matched eLC n = 3. C. BM cells were cultured for 6 days with GM-CSF, TGFβ and different combinations of BMP7, CSF1 and IL-34. Box and
whiskers graph shows mean ± min. to max. numbers of DEC205·EpCAM· cells in the cultures. Significance was calculated with a 1-way ANOVA for non-parametric samples with Dunn’s multiple comparisons test. Each symbol is data from one culture, n = 5 independent BM donors, in 3 independent experiments. **D.** The bar graph shows the mean expression ± SD of Runx3 or Id2 relative to GAPDH in sorted DEC205·EpCAM· cells. Symbols are cells from 4 independent BM donors, in 3 independent experiments. Id2 expression in LC generated in the absence versus the presence of IL-34 was analyzed by paired t-test. **E.** Bar graph shows the mean frequency ± SD of EdU·cells on day 6 of culture after cells where pulsed with EdU for 24 hours on day 2 or day 5. Symbols are cells from independent cultures (n = 2-4). Data was analyzed using a 1-way ANOVA. **F.** Line graph shows the frequency of viable DEC205·EpCAM· LC in GM-CSF / TGFβ cultures with, or without, IL-34. Symbols represent paired individual BM cultures, and were analyzed using a paired t-test, n = 5. *P<0.05, **P<0.01, ***P<0.001.

**Figure 7. Immune damage and loss of eLC opens the epidermal compartment.** Male mice received BMT with or without T cells. **A.** Graph shows the number ± SD of total CD11b·Langerin· LC in mice receiving BMT alone (triangles) or BMT with T cells (circles). Data are pooled from 2 independent experiments (n=5-13). The white square and dotted line shows LC numbers ± SD in untreated controls (n=5). **B-C.** Epidermal sheets were stained with anti-Langerin and anti-CD45.2, and confocal images processed and quantified using the Definiens Developer software: eLC (cyan) are Langerin·CD45.2·; DETC (red) Langerin−CD45.2·; and mLC (yellow) are Langerin·CD45.2−. **B.** shows example images from mice receiving BMT + T cells, and the graph in **C.** is the volume of mLC compared to eLC from BMT controls. Data are from 1 transplant experiment with 3 BMT (20 fields of view analyzed) and 2 BMT+T cell recipients (14 fields of view analyzed) (n= 162 cells from BMT mice and 356 LC from BMT+ T cell recipients). **D-F.** Topical FITC was painted on the ear skin of control un-transplanted mice (No Tx), or BMT and BMT + T cell recipients 10 weeks post-transplant. 3 days later uptake of FITC was analyzed.
within MHCII-EpCAM-Langerin+ LC in draining LN. D. Bar graph showing the frequency ± SD of FITC+ cells within LC. E. Representative contour plots show FITC uptake in gated LC. F. Bar graph showing the FITC median fluorescent intensity ± SD within FITC+ LC. Data are pooled from 2 independent experiments (n=4-7), significance was analyzed using a 1-way ANOVA, ***P<0.001. G. BMT + T cell recipients received a second round of irradiation and BMT alone 8 weeks later. The schematic illustrate the experimental set-up. H. Flow plots show the outcome in the epidermis of independent mice, who have received the 1st transplant only (Tx 1), the 2nd transplant only (Tx 2) or both transplants (Tx 1 and 2). Contour plots are pre-gated on EpCAM-Langerin(PE-labelled)+ LC.
Table 1: Parameter estimates from the best-fitting model describing the linear flow from incoming monocytes to CD11^b^ cells, EpCAM^+^ cells and to mature donor LC. Data show the estimated value with 95% confidence interval (CI).

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<th>Parameter</th>
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<td><code> </code> week 10</td>
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Figure S1. Immune injury leads the gradual replenishment of the epidermis with LC-like cells.

Figure S2. Sorting strategies used to isolate cells for RNAseq.

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Table S1. Raw data file

Table S2. Differentially expressed genes between eLC and mLC

Supplemental materials and methods including mathematical modeling methods.
Figure S1. Immune injury leads the gradual replenishment of the epidermis with LC-like cells. Male CD45.2 C57BL/6 mice were irradiated and received congenic (CD45.1) BM and polyclonal CD4 T cells, with $10^6$ CD8 Matahari T cells. Recipients were analyzed at different time points for LC chimerism.
Figure S2. Sorting strategies used to isolate cells for RNAseq. A. Schematic showing the transplant strategy using Langerin.DTR.eGFP BM to identify and sort EpCAM⁺ cells and donor LC. Flow plots show pre-gated single epidermal cells. Top - donor EpCAM⁺ cells were identified by gating on CD11b⁺CD24⁻ cells within the CD11b⁺MHCII⁺EpCAM⁻Langerin.GFP⁻ gate. Alternatively, donor LC were identified either 3 weeks or 10 weeks post-transplant by expression of EpCAM and Langerin.GFP. Bottom - Male Langerin.DTR.eGFP mice received female congenic bone marrow. Host eLC were isolated 3 weeks later based on co-expression of EpCAM and Langerin.GFP. Un-transplanted, age-matched male Langerin.DTR.eGFP mice were used to sort control LC populations. B. Contour plots show the gating strategy used to sort activated monocytes from dermal single cell suspensions. Cells were identified as CD11b⁺CD24⁺CCR2⁺Ly6C⁻MHCII⁻, as described by (44). C. Contour plots showing gating of CD11b⁺Ly6C⁺CD115⁻ blood monocytes.
Figure S3. EpCAM\(^+\) monocyte-derived cells are distinct from donor LC.  
A. Principal components analysis of gene expression. 67% of the variance within the expression data is orthogonal to PC1.  
B. Heat map showing the relative expression of the top 50 genes that contributed to the differences between blood monocytes, EpCAM\(^+\) cells (3 weeks post-transplant) and donor LC (10 weeks post-transplant) along the PC1 axis.
Figure S4. Proliferation of monocytes and LC in situ combine to replenish the LC network.  
A. Schematic to illustrate possible pathways of LC development from CD11b^{high} cells within the epidermis.  
B. Plot from a gene set enrichment analysis comparing EpCAM^{+} cells to donor LC at week 10, showing enrichment of genes related to apoptotic pathways in EpCAM^{+} cells. Normalized enrichment score 1.457, FDR q-value 0.024.  
C and D. Graphs showing the ratio of enrichment (bars) and false discovery rate (FDR) q values (line) for pathways predicted by WebGestalt to be over-expressed by (C) EpCAM^{+} cells and (D) dermal monocytes compared to blood monocytes. Over-expressed genes had a fold change more than or equal to 2, q < 0.05.
Figure S5. Long-term mLC are homologous to eLC and up-regulate Id2.  
A. Heat map comparing gene expression between eLC, donor mLC and blood monocytes.  
B. Bar chart showing the number of differentially expressed genes (DEG) in each population compared to blood Ly6C⁺ monocytes.  
C. Plot from gene set enrichment analysis comparing host eLC to donor LC at week 3, showing enrichment of genes related to the inflammatory response in host eLC. Normalized enrichment score 1.438, FDR q-value 0.064.  
D. Host eLC or donor (d)LC were sorted from the epidermis 3 weeks post-BMT + T cells, or eLC isolated from untreated controls. Langerin.DTR.GFP mice were used to identify host or donor cells, as described in supplemental figure 3, and co-cultured with antigen-specific CD8 T cells. The graph compares proliferation ± SD of for cultures with 100 LC per well. Symbols are technical replicates, and data are representative of LC sorted in 2 independent experiments.  
E. Heat map showing the relative expression of genes associated with TGFβ signaling by different myeloid cell populations.  
F. Microscopy images of representative sorted EpCAM⁺ cells after culture with primary syngeneic keratinocytes and TGFβ for 36 hours. Cells were fixed and stained for CD45.2 and Langerin. Images are representative of cells from 2 independent experiments.  
G. Graphs show the relative gene expression (FKPM) of different receptors from the RNAseq data. Blood Ly6C⁺ monocytes n = 2, epidermal EpCAM⁺ cells n= 3, donor LC n = 3, age-age-matched host LC n = 3. There were no statistically significant differences between groups. Statistical analysis was performed with a 1-way ANOVA with Holm-Sidak’s test ***P<0.001.
Figure S6. Gating of BM-derived LC in vitro. Murine C57BL/6 BM was cultured for 6 days with GM-CSF and TGFβ. Representative contour plots show the phenotype of cells on day 6 of culture. EpCAM+DEC205+ cells were identified as BM-LC.
Figure S7 Immune damage and loss of eLC opens the epidermal compartment. A. Bars show the number (mean and range) of host eLC (purple) or donor LC (magenta) in the skin of WT or DETC-deficient recipients 3 weeks after BMT with T cells. Data are from 1 representative experiment of 2, n = 3-4. B. FITC was painted onto the ears of mice 10 weeks after BMT with or without T cells. Representative plots show the gating strategy. B220<sup>+</sup> cells were excluded from live, single cells. Of the B220<sup>neg</sup> cells, populations were identified as CD11c<sup>+</sup>MHCII<sup>high</sup> migrating cells and CD11c<sup>high</sup>MHCII<sup>+</sup> resident cells. LC were identified within the migrating cells by co-expression of Langerin and EpCAM, and these cells were confirmed to be derived from donor CD45.1<sup>+</sup> BM. The histogram overlay shows that up-take of FITC by EpCAM<sup>+</sup>Langerin<sup>+</sup> cells, compared to total migrating DC.
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<thead>
<tr>
<th>Gene name</th>
<th>Suggested function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rps4l</td>
<td>Ribosomal</td>
</tr>
<tr>
<td>Tmsb10</td>
<td>Cytoskeleton organisation</td>
</tr>
<tr>
<td>Acot7</td>
<td>Regulation of lipid metabolism</td>
</tr>
<tr>
<td>Rbfox1</td>
<td>RNA-binding protein</td>
</tr>
<tr>
<td>Polg</td>
<td>DNA polymerase subunit gamma</td>
</tr>
<tr>
<td>Stat1</td>
<td>Interferon signaling</td>
</tr>
<tr>
<td>Mfge8</td>
<td>Cell adhesion</td>
</tr>
<tr>
<td>Arhgap31</td>
<td>GTPase-activating protein</td>
</tr>
<tr>
<td>Map4k1</td>
<td>JNK activation</td>
</tr>
<tr>
<td>Itim2</td>
<td>Interferon-induced gene</td>
</tr>
<tr>
<td>Iigp1</td>
<td>Target for Stat1 signaling</td>
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<tr>
<td>Unc50</td>
<td>RNA-binding</td>
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<tr>
<td>Cyfip2</td>
<td>Cell adhesion</td>
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<tr>
<td>Clec4n</td>
<td>Dectin-2. Binds mannose-containing structures</td>
</tr>
<tr>
<td>Fgl2</td>
<td>Fibrinogen-like</td>
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<td>Whsc1f1</td>
<td>Chromatin organisation</td>
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<tr>
<td>Oasl2</td>
<td>Type I IFN signaling</td>
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<tr>
<td>Csf2rb</td>
<td>CSF-2 receptor β common sub-unit</td>
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<tr>
<td>Fmn1l</td>
<td>Cell motility and survival in macrophages</td>
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<tr>
<td>P2ry6</td>
<td>G-protein-coupled receptor</td>
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<tr>
<td>Cbfa2t3</td>
<td>Transcriptional co-repressor with CBFA2/Runx1</td>
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<tr>
<td>Elov5</td>
<td>Fatty acid synthesis</td>
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<tr>
<td>Ifit1</td>
<td>Interferon-induced gene</td>
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<td>Cep83</td>
<td>Organelle biogenesis</td>
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<td>Csf2rb2</td>
<td>Csf2rb pseudogene</td>
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<td>Gls</td>
<td>Glutaminase - metabolism</td>
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<td>Lgals3bp</td>
<td>Galectin-binding</td>
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<tr>
<td>Eif2s3y</td>
<td>Translation initiation - ribosome recruitment</td>
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</table>

Table S2. List of all differentially expressed genes between eLC from age-matched controls and mLC 10 weeks post-BMT with T cells. Genes up-regulated in mLC are highlighted in bold. Suggested functions are inferred from gene ontology terms on the GeneCard® database (https://www.genecards.org).
Supplemental Materials and Methods.

Sample preparation for gene expression analysis. RNA was extracted using the RNeasy Micro Kit (QIAGEN, USA) following the manufacturer’s protocol.

RT-PCR. cDNA was synthesized using a QuantiTect Reverse Transcription Kit (Qiagen, USA). The quantitative reverse transcription PCR (qRT-PCR) was run on a CFX96 Touch Real Time PCR detection system (Bio-Rad, USA), using a Quantifast SYBR Green PCR kit (Qiagen, USA). Primer sequences for Id2 (F AAAACAGCCTGTGGACCAC and R CTGGGCACCAGTTCCTTGAG) and Runx3 (F CAGGTTCAACGACCTTCGATT and R GTGGTAGGTAGCCACTTGGG) were obtained from published sources (18, 20). mRNA expression was normalized to Gapdh (F ACCTGCCAAGTATGATGACATCA and R GGTCCTCAGTGTAGCCCAA m) by calculating 2^{-ΔCt}.

RNA sequencing. RNA yield, quality and integrity were evaluated using the RNA 6000 Pico kit on an Agilent 2100 Bioanlyser (Agilent Technologies, USA). Only samples with a RNA Integrity Number (RIN) above 8.0 were included in the study. cDNA amplification from total RNA was performed using the SMART-seq® v4 Ultra® Low Input RNA Kit (Clontech, USA). Paired-end sequencing libraries were prepared from the amplified cDNA according to the Nextera® XT DNA library prep protocol (43 bp reads, approx. 23 million reads/sample), and sequenced using an Illumina NextSeq 500 (Illumina, USA).

RNA sequencing analysis. Expression of transcript counts was quantified using kallisto (64) software with a GRCh38 transcript model. The data were imported to the R statistical environment and summarized at the gene level (that is transcript counts summed per gene) using tximport (65). Statistical transformations for visualization (for example, the rlog) and preprocessing and then analyzes of differential expression were performed using DESeq2 (66). Multiple testing adjustments of differential expression utilized the Benjamini–Hochberg (False Discovery Rate) and independent hypothesis weighting (IHW) adjustments (67).

GeneSet Enrichment Analysis. We used GeneSet Enrichment Analysis (GSEA) (68) together with the MSigDB pathways database (69). Briefly the statistical output from DESeq2 analysis was used to rank the differential expression of all genes from most upregulated to most
downregulated. Mouse genes were then converted to the nearest human homologue and then the ranking used as input for the GSEA tool and the MSigDB "Hallmark" GeneSets - a curated set of well defined biological processes.

**Gene expression analysis for heat maps.** FASTQ Toolkit, version 1.0.0 (BaseSpace, Illumina), was used for adapter trimming of the reads. Alignment and mapping of all libraries were performed using TopHat Alignment, version 2.0.0, and Cufflinks Assembly & DE, version 2.0.0 (BaseSpace, Illumina, USA). The dataset was filtered to remove genes where two or more samples in each group had an FPKM count of <1. Gene expression levels were calculated using the Cufflinks Assembly & DE, version 2.0.0 (BaseSpace, Illumina, USA), employing a geometric library normalization method and a fragment bias correction algorithm. Differential gene expression was calculated using a threshold of fold change > 2 and false discovery rate (FDR) < 0.05. Gene expression graphs are shown as relative FPKM counts were normalized to the maximum.

**Overrepresentation analysis.** The Web-based Gene Set Analysis Toolkit (WebGestalt), a suite of tools for functional enrichment analysis, was used to identify overrepresented gene ontology (GO) annotation categories and translate gene lists into functional profiles. Enrichment of GO terms and associated p-values were calculated based on hypergeometric distribution statistics, adjusting the false discovery rate using the Benjamini-Hochberg procedure.

**In vitro T cell assays.** LC were sorted from epidermal single cell suspensions and pulsed with 0.5µM SIINFEKL peptide in T cell medium (RPMI (RPMI 1640, Lonza, Switzerland), 5% heat-inactivated FBS (Life Technologies, USA), 1% L-glutamine (2 mM; Life Technologies, USA), 1% Pen-strep (100 U/ml Life Technologies, USA) and 50µM β-ME (Sigma-Aldrich, UK) for 30 min at 37°C. Splenic OT-I cells, were directly isolated using the Miltenyi MACS system (OctoMACS Separator, MS columns, CD8a [Ly-2] MicroBeads; Miltenyi Biotec), and labeled with 5µM cell trace violet (Molecular Probes) according to the manufacturer’s instructions. 5x10^4 T cells were co-cultured with 100 LC for 65 hours before analysis by flow cytometry. Cells were gated on live, single CD45.1^+CD8^+ T cells.
**Culture of EpCAM⁺ cells.** Coverslips were pre-coated with 2% (3-Aminopropyl)triethoxysilane (Sigma-Aldrich, UK) in acetone and placed in the wells of 6-well tissue culture plates. These were seeded with 5x10⁴ primary keratinocytes (live, CD45neg) sorted from the epidermis of untreated female C57BL/6 mice. Approximately 4 hours later, purified EpCAM⁺ (Langerin.GFP⁺EpCAM⁺ CD24low) cells were sorted from BMT+ T cell recipients and were added to the wells, with 5ng/ml TGFβ. Cells stained and imaged by confocal microscopy 36 hours later.
Mathematical modelling. As described in the text we modelled the flows, and dynamics within, three populations – CD11b$^{\text{high}}$ cells, EpCAM$^+$ intermediates, and mature LC. We refer to these populations as $P_1$, $P_2$ and $P_3$ respectively. We considered two developmental pathways – linear ($P_1 \rightarrow P_2 \rightarrow P_3$) and branched ($P_1 \rightarrow P_2$, $P_1 \rightarrow P_3$). To describe these, we fitted empirical descriptor functions to the timecourses of numbers of Ki67$^+$ and Ki67$^-$ CD11b$^{\text{high}}$ cells, which we refer to $P_1^+(t)$ and $P_1^-(t)$ respectively. These were both of the form

$$\log_{10}(f(t)) = c + \frac{at}{(b + t)^2}$$

Then, for the Ki67$^+$ and Ki67$^-$ EpCAM$^+$ cells (the $P_2$ population), the simplest, general model was

$$\frac{dP_2^+}{dt} = \mu_1 P_1^+ + \rho_2 (2P_2^- + P_2^+) - (\beta + \delta_2)P_2^-$$

$$\frac{dP_2^-}{dt} = \mu_1 P_1^- - \rho_2 P_2^- + \beta P_2^+ - \delta_2 P_2^-$$

where we use the superscripts $+/-$ to denote Ki67$^+$ and Ki67$^-$ cells respectively. Here $\rho_2$ is the rate of cell division; $\beta$ is the mean lifetime of Ki67 expression post-division, which we set at 3.5d (Gossel et al., eLife (2017)); $\mu_1$ is the rate of differentiation from $P_1$ to $P_2$; and $\delta_2$ is the rate at which $P_2$ cells are lost through death or onward differentiation, combined.

We fitted the above equations simultaneously to the total numbers of cells in $P_2 = P_2^+ + P_2^-$, and the Ki67 proportion $P_2 = P_2^+/(P_2^- + P_2^+)$, log- and logit-transformed respectively to normalise residuals. This fitting strategy, detailed in Hogan, Gossel et al., PNAS (2015), maximises the joint log likelihood of the datasets. Fitting on counts and Ki67 fractions rather than the timecourses of Ki67$^+$ and Ki67$^-$ cells assures independence of errors. We found no evidence for cell division within $P_2$ ($\rho_2 \simeq 0$), giving the simpler model

$$\frac{dP_2^+}{dt} = \mu_1 P_1^+ - (\beta + \delta_2)P_2^+$$

$$\frac{dP_2^-}{dt} = \mu_1 P_1^- + \beta P_2^+ - \delta_2 P_2^-.$$  

We then modelled $P_3$ as follows, assuming it is fed from a precursor population $X(t)$, which could be $P_1$ (the branched pathway) or $P_2$ (the linear pathway):

$$\frac{dP_3^+}{dt} = \mu X^+(t) + \frac{\rho_3}{1 + P_3(t)/\lambda} (2P_3^-(t) + P_3^+(t)) - (\beta + \delta_3)P_3^-(t)$$

$$\frac{dP_3^-}{dt} = \mu X^-(t) - \frac{\rho_3}{1 + P_3(t)/\lambda} P_3^-(t) + \beta P_3^+(t) - \delta_3 P_3^-(t)$$

where $P_3 = P_3^- + P_3^+$. The term proportional to $\rho_3$ is a density-dependent division rate that decreases with the size of the mature LC pool $P_3$ and is half-maximum when $P_3 = \lambda$. Mature
LC are generated from their precursor population at per capita rate $\mu$ and die at rate $\delta_3$. We estimated the parameters $\mu$, $\rho_3$, $\lambda$ and $\delta_3$ by fitting the model above simultaneously to the timecourses of total numbers of mature LC and the proportion expressing Ki67.

We found that $P_2$ as a source provided a superior description of the $P_3$ data than a $P_1$ source, with $\Delta AIC = 5$, equivalent to a degree of relative support of $1 - e^{-\Delta AIC/2} \approx 90\%$ for the linear over the branched pathway. The efficiency of differentiation from $P_2$ into mature LC is the proportion of $P_2$ cells that survive to become mature LC, $\mu/(\mu + \delta_2)$.

(70, 71).