Nociceptive sensory neurons drive interleukin-23-mediated psoriasiform skin inflammation

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The skin has a dual function as a barrier and a sensory interface between the body and the environment. To protect against invading pathogens, the skin harbours specialized immune cells, including dermal dendritic cells (DDCs) and interleukin (IL)-17-producing γδ T cells, the aberrant activation of which by IL-23 can provoke psoriasis-like inflammation1–4. The skin is also innervated by a meshwork of peripheral nerves consisting of relatively sparse autonomic and abundant sensory fibres. Interactions between the autonomic nervous system and immune cells in lymphoid organs are known to contribute to systemic immunity, but how peripheral nerves regulate cutaneous immune responses remains unclear3,4. We exposed the skin of mice to imiquimod, which induces IL-23-dependant psoriasiform inflammation12. Here we show that a subset of sensory neurons expressing the ion channels TRPV1 and Nav1.8 is essential to drive this inflammatory response. Imaging of intact skin revealed that a large fraction of DDCs, the principal source of IL-23, is in close contact with these nociceptors. Upon selective pharmacological or genetic ablation of nociceptors9–11, DDCs failed to produce IL-23 in imiquimod-exposed skin. Consequently, the local production of IL-23-dependent inflammatory cytokines by dermal γδT17 cells and the subsequent recruitment of inflammatory cells to the skin were markedly reduced. Intradermal injection of IL-23 bypassed the requirement for nociceptor communication with DDCs and restored the inflammatory response2. These findings indicate that TRPV1+Nav1.8+ nociceptors, by interacting with DDCs, regulate the IL-23/IL-17 pathway and control cutaneous immune responses.

Repeated topical application of imiquimod (IMQ) to murine skin provokes inflammatory lesions that resemble human psoriasis7,9. This response is mediated by IL-23, which stimulates skin-resident γδ T cells to secrete IL-17 and IL-22, cytokines that induce inflammatory leukocyte recruitment and acanthosis10. Indeed, antibodies targeting the shared p40 subunit of IL-12 and IL-23 inhibit both IMQ-induced murine dermatitis and human psoriasis10,11. Frequent symptoms in human psoriasis, aside from the prominent skin lesions, include the sensations of itch, pain and discomfort in affected areas11. Clinical reports suggest that intraleosionally administered anaesthetics or surgical denervation of psoriatic lesions not only abrogate local sensation, but also ameliorate local inflammation12. Similarly, in mutant mice with disseminated psoriasisform dermatitis, peripheral nerve dissection attenuated skin inflammation13; however, cutaneous nerves are composed of sympathetic and several types of sensory fibres, and the role of individual types of nerve fibres remains unclear14. Using the IMQ model, we investigated whether and how specific subsets of peripheral nerves contribute to the formation of psoriasiform skin lesions.

Skin sensations perceived as inflammatory pain, noxious heat and some forms of itch are transmitted by sensory fibres that express the cation channel TRPV1. Most TRPV1+ fibres co-express the sodium channel Na+,1.8 (ref.9–11). Na+,1.8+ nociceptors can be identified in the dermis of Na+,1.8-TdTomato (TdT) mice by their red fluorescence15. Confocal microscopy of skin samples from Na+,1.8-TdT mice co-stained for tyrosine hydroxylase (TH), which identifies sympathetic fibres, and for β3-tubulin, a pan-neuronal marker, revealed that Na+,1.8+TH– nociceptors represent the vast majority of cutaneous nerve fibres, whereas Na+,1.8+TH+ sympathetic fibres are rare. To investigate the roles of sympathetic fibres and nociceptors in the IMQ model, mice were treated systemically with either 6-hydroxydopamine (6OHDA) or resiniferatoxin (RTX) to ablate TH+ sympathetic neurons or TRPV1+ nociceptors, respectively16,17 (Extended Data Figs 1, 2). Subsequently, IMQ was applied topically to one ear and the ensuing inflammatory response was assessed based on the change in ear thickness, size of the myeloid infiltrate (Extended Data Fig. 3a) and tissue contents of inflammatory cytokines.

After sympathetic denervation, IMQ-induced ear swelling was reduced compared to controls (Extended Data Fig. 1c); however, the inflammatory infiltrate was increased, whereas IL-17A, IL-17F, IL-22 and IL-12/23p40 production remained unchanged (Extended Data Fig. 1d–i). Thus, sympathetic innervation exerts little or no direct local control over the inflammatory skin response. The observed changes were probably due to cardiovascular effects and/or global immune dysregulation after systemic sympathectomy18 (Supplementary Information). By contrast, in RTX-treated mice both ear swelling and inflammatory infiltrates in IMQ-exposed ears were profoundly reduced (Fig. 1b–e and Extended Data Fig. 4a–d). RTX treatment did not alter the systemic supply of inflammatory cells19 (Extended Data Fig. 4e, f). Moreover, intravital microscopy of ear skin revealed similar leukocyte rolling in RTX-treated and control mice (Extended Data Fig. 5), indicating that the absence of nociceptors did not affect the baseline adhesiveness of dermal microvessels19. It is more likely that ablation of TRPV1+ sensory nerves reduced IMQ-induced inflammation through local, extravascular mechanisms. However, the attenuated inflammatory response was not limited to the skin, as the IMQ-induced enhancement in cellularity of the draining auricular lymph node was also blunted by RTX (Fig. 1f).

Lymph nodes have a critical function in dermal antigen presentation to naive T cells and in the generation of migratory effector cells. They also possess peripheral innervation20, which could have been altered by RTX; however, the role of skin-draining lymph nodes during psoriatic inflammation is unclear. To address this issue, we tested the effect of IMQ in lymphotxin-α-deficient (LTα−/−) mice, which are devoid of lymph nodes. Compared with wild-type mice, there was no statistical difference in ear thickness, frequency or composition of the inflammatory infiltrate in ears of LTα−/− mice (Fig. 1g, h and data not shown). Additionally, we treated wild-type mice with FTY720, which blocks T-cell egress from lymph nodes, preventing trafficking of effector cells to peripheral tissues21. Again, IMQ elicited full-fledged inflammation in exposed ears (Fig. 1i, j and data not shown), indicating that T-cell priming in skin draining lymph nodes is dispensable for the acute induction of psoriasisform inflammation. Together, these findings imply that TRPV1+ nociceptors promote local immune responses directly in the skin.

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Neutrophil recruitment to the skin and keratinocyte hyperproliferation, hallmarks of IMQ-induced inflammation, are driven by IL-17 and IL-22, respectively\(^2,3\). We asked whether nociceptors regulate the production of these cytokines. Indeed, after IMQ treatment, protein levels of IL-17A, IL-17F and IL-22 had markedly increased in the ears of control mice, whereas in RTX-treated mice IL-17A was very low and IL-17F and IL-22 remained below the detection limit (Fig. 2a–c).

Thus, TRPV1\(^+\) nociceptors control the generation of several key effector cytokines in psoriasiform dermatitis.

As our findings in FTY720-treated mice suggested that recruitment of lymph-node-derived effector T cells is not needed for IMQ-induced skin inflammation, we asked whether nociceptors regulate cytokine production by skin-resident lymphocytes, particularly γδ T17 cells, which produce IL-17F and IL-22 (refs 3, 22, 23). In the ears of IMQ-treated control mice, most IL17F\(^+\) lymphocytes were dermal γδ T cells, whereas few conventional γβ T cells expressed IL-17F (Fig. 2d and Extended Data Figs 3b, 6a). In RTX-treated mice, both IL-17F\(^+\) and IL-22\(^+\) dermal γδ T cells were significantly reduced, indicating that TRPV1\(^+\) neurons drive the local production of IL-17F and IL-22 primarily by γδ T17 cells (Fig. 2e–g and Extended Data Fig. 6b, c). These T cells seed the skin in early life and are poised for rapid IL-17 production upon stimulation by IL-23 (ref. 24). Indeed, in the ears of IL-23R\(^-/-\) mice\(^5\), ∼60% of dermal γδT17 and ∼20% of γβ T cells expressed IL-23R (Fig. 2h, i), whereas T cells in auricular lymph nodes and epidermal γδ T cells expressed little or no IL-23R (Extended Data Fig. 6d, e).

In light of the preferential expression of IL-23R on dermal γδ T17 cells, the known role of IL-23 as a driver of IL-17 and IL-22 generation...
in psoriasiform inflammation\(^{12}\) and our finding that RTX pre-treatment profoundly diminished IMQ-induced cytokine production by γδT17 cells, we proposed that TRPV1\(^{+}\) nociceptors might control dermal IL-23 production. Topical IMQ treatment of control mice markedly increased p40 protein levels, as well as messenger RNA levels of Il12b (IL-12/23p40) and Il23a (IL-23p19), but not Il22a (IL-12p35). These effects were nearly abolished after RTX treatment (Fig. 3a and Extended Data Fig. 7a–c), suggesting that TRPV1\(^{+}\) nociceptors are essential for cutaneous IL-23 production. It seems unlikely that IL-12 had a major impact in the IMQ model, because IMQ-induced inflammatory parameters were considerably reduced in IL-23RGFP/GFP mice, which respond to IL-12 but not IL-23 (Fig. 3b–f). However, IL-12-dependent skin inflammation induced using a chemical irritant, 2,4-dinitrofluorobenzene (DNFB)\(^{26}\), was profoundly reduced in RTX-treated mice, suggesting that RTX-sensitive fibres also have a role in IL-12-driven dermatitis (Extended Data Fig. 7d, e).

Of note, although IL-17F and IL-22 production as well as myeloid infiltrates were virtually abolished in IL-23RGFP/GFP mice, ear swelling was only partially reduced (Fig. 3b–f), suggesting that topical IMQ promoted modest tissue oedema through an IL-23-independent pathway. This activity was independent of nociceptors, as RTX treatment of IL-23RGFP/GFP mice had no effect on the IMQ-induced swelling (Extended Data Fig. 7f).

Although the above findings clearly demonstrate that nociceptors are indispensable for IMQ-induced dermal IL-23 production, it remained possible that nociceptive fibres exert additional pro-inflammatory functions, for example, by directly regulating γδT17 cells. To address this, we performed intradermal IL-23 injections, which trigger psoriasiform skin inflammation in murine skin\(^{12}\). Regardless of whether nociceptors were ablated or left intact, IL-23 administration resulted in profound ear swelling and fully rescued IL-17A and IL-17F production by Thy1\(^{+}\) cells (Fig. 3h, i) and γδT17 cells (Extended Data Fig. 7g, h). Together, these results suggest that the pro-inflammatory function of TRPV1\(^{+}\) neurons is rooted exclusively in the promotion of IL-23 production, at least in this experimental setting.

Next, we sought to identify the IL-23-producing cell type(s) regulated by TRPV1\(^{+}\) nociceptors. In intestinal barrier tissues, Ly-6C\(^{hi}\) inflammatory monocytes were identified as a major source of IL-23 in a colitis model\(^{27}\). However, depletion of neutrophils and inflammatory monocytes with anti-Gr-1 did not affect IMQ-induced dermal IL-23p40 or IL-17F levels (Fig. 3j, k and Extended Data Fig. 8a), suggesting that skin-resident dendritic cells and/or macrophages rather than migratory myeloid cells were the critical source of IL-23. Thus, we injected CD11c-diphtheria toxin receptor (DTR) mice with diphtheria toxin (DTX), which depleted DDCs and Langerhans cells, but not macrophages (Extended Data Fig. 8b, c). After DTX treatment, IMQ-induced expression of Il23a mRNA was markedly reduced in treated ears (Fig. 3l). Although these results do not discriminate between the relative contributions of Langerhans cells and DDCs to IL-23 production, IL-34-deficient mice—which

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**Figure 3 | DDC-derived IL-23 is critical to drive psoriasiform skin inflammation and acts downstream of RTX-sensitive nociceptors.** a. After 3 days of IMQ challenge of vehicle (DMSO)- or RTX-treated mice, ears were harvested and total protein was prepared to quantify IL-23p40 by ELISA (n = 5 per experiment; ***P < 0.001). b. Ears of wild-type (WT) or IL-23RGFP/GFP mice (n = 5 per group) were treated daily with IMQ and ear thickness was measured relative to the contralateral ear at the indicated time points (**P < 0.001). c–f. After 3 days of IMQ challenge in wild-type (n = 5) or IL-23RGFP/GFP mice (n = 4) total protein was prepared from ear skin and IL-17F (c) and IL-22 (d) were quantified by ELISA (\*P < 0.05; **P < 0.001). Cell suspensions from exposed ears stained for inflammatory monocytes (e) and neutrophils (f) to assess total numbers (n = 5 per experiment; \*P < 0.05; **P < 0.01). g–i. IL-23 was injection intradermally every other day into the ears of DMSO- or RTX-treated mice (n = 8 per group). g. Ear thickness was measured as indicated. h. After 3 days, the number of IL-17A-producing (h) and IL-17F-producing (i) Thy1\(^{+}\) cells per ear was quantified by FACS (n = 5). j, k. Mice were treated with anti-Gr-1 to deplete neutrophils and inflammatory monocytes or isotype-matched control monoclonal antibody and challenged with IMQ for 3 days. Ear skin protein lysates were analysed for IL-23p40 (j) and IL-17F (k) by ELISA (n = 3). l. CD11c-DTR mice were treated with DTX or PBS 12 h before IMQ challenge. After 6 h, ears were harvested and processed for total RNA isolation and Il23a mRNA levels were analysed by quantitative polymerase chain reaction (qPCR) (n = 4; ***P = 0.001). AU, arbitrary units. ND, not detected; NS, not significant. Error bars show mean ± s.e.m.
mounts revealed that at steady state IL-23, we sought to characterize the spatial relationship between DDCs subsets of DDCs were FACS sorted (Extended Data Fig. 8c) from cell selectively lack Langerhans cells—are not impaired in IMQ-induced skin inflammation. We focused our analysis on DDCs, which can be further subdivided into CD11c+ and CD11b+ subsets. Both subsets, as well as macrophages, were sorted by fluorescence-activated cell sorting (FACS) from IMQ-treated and control ears to measure mRNA for Il23a and Il12a. Although CD103+ DDCs showed the most notable upregulation of Il23a mRNA, taking into consideration that CD11b+ DDCs are more abundant, we estimate that the latter subset produced ~75% of the Il23a mRNA, consistent with a recent report (Fig. 4a and Extended Data Fig. 8c, d).

Having identified DDCs as the principal source of IMQ-induced IL-23, we sought to characterize the spatial relationship between DDCs and cutaneous nerves. Remarkably, confocal microscopy of skin whole mounts revealed that at steady state ~75% of DDCs were either in direct contact or in close proximity to sensory nerves (Fig. 4b–d and Extended Data Fig. 9a). Interactions were apparent along the entire length of nerves, suggesting that DDCs may receive signals from unmyelinated nociceptor axons and not just from nerve terminals. However, given the high density of peripheral nerves in the skin, it was difficult to judge whether the association with DDCs occurred merely by chance or reflected a biased distribution. To address this possibility, we compared DDC localization relative to two other dense anatomical structures: blood and lymph vessels. In resting tissues, contacts of DDCs with these microvascular networks were only about half as frequent as with peripheral nerves (Fig. 4d).

Of note, even though inflammation enhances DDC motility and egress into draining lymphatics, IMQ challenge did not alter the frequency of DDCs contacting nerves (Extended Data Fig. 9b), suggesting that DDCs engaged in dynamic interactions with nociceptors. We reconstituted Na1.8-Tdt mice with yellow fluorescent protein (YFP)-tagged CD11c bone marrow and performed time-lapse multiphoton–intravital microscopy (MP–IVM) in ear skin to generate three-dimensional time-lapse videos of interactions between YFP+ DDCs and Tdt+ nociceptors. These interactions were highly diverse (Supplementary Video 1); some DDCs seemed to be anchored on nerves and sometimes extended protrusions to probe the surrounding tissue, whereas others appeared to use nerve fibres as a scaffold for directional migration (Supplementary Video 2).

Together, our results support the idea that DDCs can physically interact with a subset of nociceptors that regulate the production of IL-23. However, it should be cautioned that the pharmacological target of RTX, TRPV1, is also expressed on some non-neuronal cells. Consequently, we sought an alternative to confirm the role of nociceptors in an experimental system that does not rely on pharmacologically targeting TRPV1. To this end, we used Na1.8-diphertheria toxin (DTA) mice in which Na1.8+ fibres, which respond to mechanical pressure, inflammatory pain and noxious cold, are selectively deleted. Although nociceptor-associated transcripts in dorsal root ganglia were more profoundly reduced in this genetic model than after RTX treatment, Trpv1 mRNA levels were only reduced by ~80%, consistent with the fact that noxious heat sensing fibres express TRPV1, but not Na1.8 (Extended Data Figs 2c, 9c).

After IMQ challenge, the ears of Na1.8-DTA mice contained very low IL-17A, IL-17F, IL-22 and IL-23p40 as compared to littermate controls (Fig. 4e–h). We conclude that Na1.8+ TRPV1+ nociceptors are insufficient to induce IL-23 production; rather, NaV1.8+ TRPV1+ nociceptors are driving the response.

In light of these results, we propose a model of cutaneous neuroimmunomodulatory interactions (Extended Data Fig. 10) whereby dermal Na1.8+TRPV1+ nociceptors are essential to induce IL-23 production by nearby DDCs. IL-23 then acts on IL-23R and IL-22 secretion, which precipitates the recruitment of circulating neutrophils and monocytes driving psoriasiform skin inflammation. The fact that both RTX and Na1.8-DTA mice largely preserve a dense meshwork of dermal nerves (Extended Data Fig. 9d, e) implies that DDCs do not simply rely on nerves as a scaffold from which to produce IL-23. It is more likely that Na1.8+TRPV1+ nociceptors actively induce and control IL-23 production.

Although further studies will be needed to determine the precise molecular underpinnings of neuroimmunomodulation in the skin (Supplementary Information), the present findings indicating nociceptor-mediated control of DDCs and the IL-23/IL-17 axis open new avenues for the treatment of inflammatory diseases in the skin and perhaps elsewhere. Intriguingly, recent work has shown that Na1.8+ nerve fibres exert immunosuppressive activity during Staphylococcus aureus infection, raising the possibility that some pathogens have evolved mechanisms to subvert the pro-inflammatory function of nociceptors. Together, this recent work and the present study suggest an emerging new paradigm whereby TRPV1+Na1.8+ nociceptive fibres integrate environmental

**Figure 4** | DDCs are closely associated with cutaneous nerves and depend on Na1.8+ nociceptors for IMQ-induced IL-23 production. a, Mice were challenged with IMQ (n = 20 pooled mice per condition) and, after 6 h, myeloid cell populations comprising dermal macrophages (mac.) and two subsets of DDCs were FACS sorted (Extended Data Fig. 8c) from cell suspensions to measure Il23a mRNA by qPCR. b, Representative confocal micrographs of ear skin whole mounts from CD11c–YFP mice stained for β3-tubulin (peripheral nerves, red) and Lyve-1 (lymphatics, blue) or CD31 (blood and lymphoid endothelial cells, blue). Original magnification, ×200. c, Close-up confocal micrograph of a CD11c–YFP cell in contact with a nerve (see also Supplementary Videos 1, 2). d, Quantification of three-dimensional DDC proximity to peripheral nerves, lymphatics and blood vessels in normal ear skin. The frequency of DDCs (n = 330) in contact, proximal (0–7 μm) and distal (>7 μm) to nerve fibres was determined as described in Methods and a Chi-squared test showed bias of DDCs to nerves relative to lymphatics and blood vessels (***P < 0.0001). e–h, Ears of Na1.8-DTA or control littersmates were treated daily with IMQ. Total protein was prepared from ear skin after 3 days and IL-17A (e), IL-17F (f), IL-22 (g) and IL-23p40 (h) were quantified by ELISA (n = 4 ears per condition; *P < 0.05; **P < 0.01). ND, not detected. Error bars show mean ± s.e.m.
signals to modulate local immune responses to a variety of infectious and inflammatory stimuli.

METHODS SUMMARY

Published mouse strains used in this study are detailed in Methods. Na1.8-Cre mice were bred with Rosa26-DTA and Rosa26-TdT mice to generate Na1.8-DTA mice for functional studies and Na1.8-TdT mice for imaging, respectively. TRPV1+ nociceptors were deleted using three escalating doses (30 μg kg–1, 70 μg kg–1 and 100 μg kg–1) of RTX as described previously7. To induce psoriasis-like inflammation, 8–12-week-old mice were treated topically with 5% IMQ cream or injected with 500 ng per ear recombinant (r)IL-23. Ear thickness was measured using an engineer’s micrometer (Mitutoyo). Cytokines were quantified from skin protein extracts by ELISA (Biolegend, R&D). For flow cytometric analysis of tissue leukocyte markers and intracellular cytokines, single-cell suspensions from ear skin were prepared by enzymatic digestion4. Imaging of fixed skin tissue was performed using an Olympus Fluoview BX50WI inverted microscope, and MP–IVM in live anesthetized mice was performed using an upright microscope (Prairie Technologies) with a MaiTai Ti:sapphire laser (Spectra-Physics). All animal studies were approved by the Institutional Animal Care and Use Committee of Harvard Medical School and complied with National Institutes of Health guidelines.

Online Content

Any additional Methods, Extended Data display items and Source Data are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Supplementary Information

Available in the online version of the paper.

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Author Contributions

L.R.-B., J.O.-M., and U.H.v.A. designed the study. L.R.-B., J.O.-M., and U.H.v.A. wrote the manuscript. J.O.-M., L.R.-B., J.O.-M., and U.H.v.A. contributed to the experimental design and discussion and advice. This work was supported by National Institutes of Health (NIH) grants A090299, A078897, A095261 and A111595 (to U.H.v.A.), NIH 5F31AR063548-02 (to J.O.-M.) and the Human Frontiers Science Program, Charles A. King Trust and National Psoriasis Foundation (to L.R.-B.).

Author Information

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Mice. C57BL/6 mice, 4–8 weeks old, were purchased from Charles River or the Jackson Laboratory and female mice were used in experiments. IL-23 GFP/GFP mice, in which GFP is knocked into the cytoplasmic tail of IL-23R and the homozygous GFP mouse acts as a functional receptor knockout, were provided by M. Oukka and both male and female mice were used in experiments. CD11c–YFP mice were a gift from M. Nussenzweig and both male and female mice were used in experiments. LTα/ α mice were purchased from Jackson Laboratory and male mice were used in experiments. CD11c–DTRm mice were purchased from Jackson Laboratory and both male and female mice were used in experiments. Na.,1,8-Cre mice were previously described, Rosa26-DTA mice and Rosa26-TdT mice, which express a floxed-STOP cassette upstream of the ubiquitously expressed Rosa26-DTA or Rosa26-TdT construct, respectively, were purchased from Jackson Laboratory. Na.,1,8-Cre male mice were bred with Rosa26-DTA and Rosa26-TdT female mice to generate Na.,1,8-DTA for functional studies and Na.,1,8-TdT for imaging experiments, respectively, of which both matched male and female litters were used in experiments. Gliarial fibrillary acidic protein (GFAP)–TdT mice were generated by crossing heterozygous GFAP–Cre mice purchased from Jackson Laboratory with Rosa26-TdT mice. MHC-II–GFP mice were provided by M. Boes. Bone marrow chimaeras were generated by irradiating Na.,1,8-TdT mice or GFAP–TdT mice with two split doses totaling 1,300 rad and reconstituting with CD11c–YFP or MHC-II–GFP unfractionated and bone marrow injected intravenously, respectively. Bone marrow chimaeras were allowed to rest for 12 weeks before use. Mice were all housed in specific pathogen-free conditions in accordance with the National Institutes of Health and all experimental animal protocols were approved by the Institutional Animal Care and Use Committee at Harvard Medical School. For most animal experiments, because the contralateral ear served as a control for the ear in which an inflammatory stimulus was applied, no randomization was used. Investigators were blinded for the initial ear-swelling experiments and subsequently no blinding was used.

Depletion. RTX, a capsainica analogue, was injected subcutaneously into the flank of 4-week-old mice in three escalating (30 μg kg⁻¹, 70 μg kg⁻¹ and 100 μg kg⁻¹) doses on consecutive days. Control mice were treated with vehicle solution (DMSO in 0.3% Triton X-100, 10% goat serum and Fc Block and also stained with the following antibodies in the same buffer. Unconjugated antibodies used include: anti-neuronal class III β-tubulin (clone TU1; Covance), anti-TH (clone A285-105; Millipore), anti-Ly-1 (clone ALY7; Biosciences), anti-peripherin (Polyclonal; Abcam) and anti-NeuN (A60; Millipore). Alexa488-conjugated antibodies include: anti-neuronal class III β-tubulin (clone TU1); Covance, goat anti-mouse IgG (Invitrogen) and goat anti-rabbit IgG. Alexa647–conjugated antibodies include: goat anti-rat IgG and goat anti-rabbit IgG. Washing steps were performed in PBS with 0.2% BSA, and 0.1% Triton X-100. Ears were mounted with the dermis facing the imaging plane in FluorSave reagent (Calbiochem).

Confocal microscopy. Confocal images were acquired on an Olympus Fluoview BX50WI inverted microscope with ×10/0.4, ×20/0.5 and ×40/1.3 magnification/numerical aperture objectives. For images used in three-dimensional analysis, image planes were acquired at 0.5 μm intervals through the imaging volume. Intravitral two-photon microscopy. Anaesthetized mice were placed on a custom-built stage and the ear was fixed to a temperature-controlled metallic support to facilitate exposure of the dorsal aspect to the water-immersion ×20 objective (0.95 numerical aperture) of an upright microscope (Prairie Technologies). A MaiTai Ti:sapphire laser (Spectra-Physics) was tuned between 860 nm and 900 nm for multiphoton excitation and second-harmonic generation. For dynamic analysis of cell interaction in four dimensions, several x-y planes (512 × 512 with 2 spacing ranging from 2 μm to 4 μm were acquired every 15–20 s with an electronic zoom varying from 1× to 3×. Emitted light and second-harmonic signals were directed through 450/80 nm, 525/50 nm and 630/120 nm bandpass filters and detected with non-descanned detectors. Post-acquisition image analysis, volume rendering and four-dimensional time-lapse videos were performed using Imaris software (Bitplane scientific software).

Intravitral microscopy and image analysis. Intravitral microscopy of skin was performed as previously described. In brief, control or RTX-treated mice were anaesthetized and the left ear was exposed and positioned for epifluorescence intravitral microscopy. Preparations were transferred to an intravitral microscope (IV-500; Mikron Instruments), equipped with a Rapp OptoElectronic SP-20 xenon flash lamp system and QImaging Rolera-M GiEMCCD camera. The fluorescent dye rhodamine-6G (20 mg kg⁻¹ in PBS) was administrated through the catheterized right carotid artery to visualize circulating leukocytes. Cell behaviour in skin venules was recorded in 10 min recordings through >10 ×40 water-immersion objectives (Achromplan; Carl Zeiss). Rolling fractions in individual vessel segments were determined offline by playback of digital video files. The rolling fraction was determined as the percentage of cells interacting with skin venules in the total number of cells passing through a vessel during the observation period.

Cytokine quantification by ELISA. Skin biopsies from ears were obtained using 10 mm diameter skin biopsy punches (Acuderm). Samples were homogenized in Tissue Extraction Reagent I (Invitrogen) in the presence of protease inhibitor cocktail (Roche) using a gentleMACS dissociator (Miltenyi Biotech). Skin protein extracts were assayed for IL-17A, IL-22, IL-23, IL-23p (Bioleng) and IL-17F (R&D Systems) in accordance with the manufacturer’s instructions.

RNA isolation and qPCR. RNA from sorted and pelleted cells was isolated using RNAeasy Plus Minikit (Qiagen) including a gDNA elimination step. Ear skin and dorsal root ganglia (DRGs) were harvested and placed immediately in RNAlater (Ambion) before homogenization and RNA isolation using Qiagen RNAeasy Plus Minikit (Qiagen). DRGs for comparison of the efficacy of depletion were harvested from equivalent anatomical locations, typically consisting of the cervical and thoracic ganglia from C1–T2. cDNA synthesis was done using Superscript Vilo cDNA synthesis kit (Invitrogen) following the manufacturer’s instructions. Relative quantification of transcripts was done using validated Quantitect Primer Assays (Qiagen) combined with the QuantTect SYBR Green Detection Kit (Qiagen) on a LightCycler 480 (Roche). Expression of gene was calculated to Gapdh using the AC method.

Histology. Ears were embedded in paraffin and submitted for histological analysis by haematoxylin and eosin staining to the Harvard Rodent Histopathology Core. Whole-mount immunofluorescence of ear skin. Ears were harvested and split into dorsal and ventral halves. After fixation in 4% formalin A (B) and B2 collagen cartilage was removed under a dissecting microscope to expose the dermis easily for imaging. Tissue was blocked in blocking buffer containing PBS with 0.5% BSA, 0.3% Triton X-100, 10% goat serum and Fc Block and also stained with the following antibodies in the same buffer. Unconjugated antibodies used include: anti-neuronal β-tubulin, anti-TH (1:400 dilution), DAPI (1:1000 dilution) and Alexa Flour 488 (1:1000 dilution).
antibodies used include: anti-CD103 (clone 2E7; Biolegend). PE-conjugated antibodies used include: anti-δ-TCR (clone GL3; Biolegend), anti-Ly-6C (clone HK1.4; Biolegend). PerCPCy5.5-conjugated antibodies used include: anti-CD45.2 (clone 104; Biolegend). PE-Cy7-conjugated antibodies used include: anti-TCR-β (clone H57-597; Biolegend), anti-CD11c (clone HL3; BD Pharmingen). Alexa647-conjugated antibodies used include: anti-CCR6 (clone I4D706; BD Pharmingen), anti-CD11b (clone M1/70; Biolegend). APC-Cy7-conjugated antibodies used include: anti-Thy1.2/CD90.2 (clone 30-H12; Biolegend), anti-1A/1E Class-II (clone M5/114.15.2; Biolegend). Cells were then washed with PBS and resuspended in MACS buffer for immediate acquisition or fixed in Cytofix (BD Pharmingen) as per the manufacturer’s instructions for later acquisition. For analysis, cells were acquired on a BD FACS CANTO (BD Pharmingen) and analysed using FlowJo software (TreeStar). For intracellular cytokine staining, cells were not restimulated and rather harvested and digested in the presence of Brefeldin A (Biolegend). After staining for surface antigens, cells were then fixed and permeabilized using BD Cytofix/Cytoperm kit (BD Pharmingen) as per the manufacturer’s instructions. Cells were stained in Perm/Wash buffer. Alexa488-conjugated antibodies used include: anti-IL-17A (clone TC11-18H10.1; Biolegend), anti-IL-17F (clone 9D3.1C8; Biolegend). Alexa647-conjugated antibodies used include: anti-IL-17A (clone TC11-18H10.1; Biolegend), anti-IL-17F (clone 9D3.1C8; Biolegend) and anti-IL-22 (clone Poly5164; Biolegend). For determining total counts of cell subsets, an aliquot of the same cell suspension used for flow cytometry was stained for CD45.2 and acquired on an Accuri Cytometer (BD Biosciences) with a known amount of CountBright counting beads (Invitrogen). The total CD45^+ cell number was then determined in the original cell suspension and used for total quantification of the cell number of each subset of interest defined by multi-parameter flow cytometry based on that subset’s frequency relative to the CD45^+ population. For cell sorting, cells were stained for indicated surface markers and sorted using a BD FACS Aria (BD Biosciences) into complete DMEM media and used for total quantification of the cell number of each subset of interest defined by multi-parameter flow cytometry based on that subset’s frequency relative to the CD45^+ population. For cell sorting, cells were stained for indicated surface markers and sorted using a BD FACS Aria (BD Biosciences) into complete DMEM media before RNA extraction. Image analysis. Images from confocal microscopy and from two-photon microscopy were analysed on either Velocity software (Improvision) or Imaris (Bitplane). For determining the distance of CD11c-YFP cells from nerves, lymphatics or blood vessels, the centroid of dendritic cells was calculated by an unbiased approach, and the distance between a given dendritic cell centroid and the closest nerve, blood vessel or lymphatic vessel was calculated using Imaris software. To bin cells into contact (<0 μm), proximal (0–7 μm) and distal (>7 μm) fractions, the calculated average radius of a dendritic cell (7.04 μm) was subtracted from each measured distance. Statistical analyses. Precise experimental numbers of animals are reported in the figure legends. Experiments were repeated at least three times except in Figs 2e and 3e–g in which two replicates were done assessing ten mice total in each experimental group overall. Some data, such as ear-swelling curves, represent pooled averages of the sum total of animals used in experiments whereas other data consist of a representative experiment of the independent experiments. All statistical analyses were performed using Prism (GraphPad Software) and results are calculated as means with error bars representing the s.e.m. Means between two groups were compared by using a two-tailed t-test. Means between three or more groups were compared by using a one-way or two-way ANOVA. A Chi-squared statistical analysis was performed for Fig. 4d comparing the total number of dendritic cells in contact, proximal and distal bins relative to nerves versus lymphatic vessels or versus blood vessels.

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Extended Data Figure 1 | 6OHDA treatment ablates sympathetic nerve function and reduces ear swelling, but does not ameliorate the inflammatory response to IMQ treatment. a, The experimental protocol was as follows. Mice were injected intraperitoneally with 6OHDA, resulting in a reversible chemical sympathectomy lasting for approximately 2 weeks. After a rest period of 3 days animals were challenged topically on the ear with IMQ. b, Representative section of splenic white pulp showing B cells (B220, white), T cells (CD3, red), and TH\(^+\) (green) nerve fibres in vehicle (ascorbic acid)-treated and sympathectomized (6OHDA) mice. c–i, Analysis of the inflammatory response in ears of vehicle (ascorbic acid)-treated and sympathectomized (6OHDA) mice after daily topical IMQ challenge: timecourse of change in ear thickness of IMQ-treated ear relative to the contralateral ear (n = 10; *\(P < 0.05\); **\(P < 0.01\)) (c) and total number of infiltrating monocytes (d) and neutrophils (e), and the amount of IL-17A (f), IL-17F (g), IL-22 (h) and IL-23p40 (i) in protein extracts of IMQ exposed ears at day 3 (*\(P < 0.05\); n = 5). NS, not significant.
Extended Data Figure 2 | RTX treatment diminishes noxious heat sensation and decreases the expression of nociceptor markers on dorsal root ganglia. a, Schematic protocol of nociceptor ablation and induction of psoriasiform skin inflammation. RTX was injected subcutaneously into the back in three escalating doses (30 μg·kg⁻¹, 70 μg·kg⁻¹ and 100 μg·kg⁻¹) on consecutive days and mice were allowed to rest for at least 4 weeks before IMQ treatment. b, Denervation was confirmed by immersing the tail of mice into a temperature-controlled water bath maintained at 52 °C and the latency to the first tail movement to avoid water was measured (n = 6). c, Total RNA was isolated from dorsal root ganglia (level C1–C7) of vehicle (DMSO)- and RTX-treated mice and the levels of *Trpv1*, *Scn10a* (Nav1.8), *Tac1* (substance P), *Mrgprd*, *Trpm8* and *Trpa1* mRNA relative to *Gapdh* were determined (n = 3).
Extended Data Figure 3 | Gating strategy for myeloid cells and T-cell subsets from digested ear skin. a, The ear skin of mice challenged for 3 days with IMQ was digested as described in Methods and, after doublet exclusion and gating on defined FSC-A, SSC-A parameters, infiltrating myeloid cells were gated as CD45$^+$ I-Ab$^+$ (Class-II)$^+$, CD11b$^+$ CD11c$^-$, and then subdivided into inflammatory monocytes and neutrophils based on Ly-6C and Ly-6G staining. b, The ear skin of naive mice was digested as described in Methods and, after doublet exclusion and gating on defined FSC-A, SSC-A parameters, cutaneous T cells were gated on CD45$^+$, Thy1$^+$, and then divided into subsets based on staining for $\delta$-TCR and $\beta$-TCR.
Extended Data Figure 4 | RTX treatment reduces the immune cell infiltrate upon IMQ treatment in the skin but does not affect reservoirs of inflammatory monocytes and neutrophils at steady state. **a**, The ear skin of vehicle (DMSO) or sensory denervated (RTX) mice was treated with topical IMQ cream daily and the total numbers of CD45⁺ cells were determined on day 3 as explained in Methods (n = 10). **b**, Representative histological sections of untreated and IMQ-treated ears at day 3 stained by haematoxylin and eosin (×20) (n = 5 per condition). **c, d**, Total inflammatory monocytes (CD45⁺, CD11b⁺, Ly-6Cʰigh) and neutrophils (CD45⁺, CD11b⁺, Ly-6Gʰigh) were determined by flow cytometry (n = 5–10 mice per time point). Two-way ANOVA was run to compare total numbers of inflammatory monocytes and neutrophils between DMSO and RTX conditions over days 3–6 (****P < 0.0001). One-way ANOVA was run to compare total inflammatory monocytes and neutrophil numbers over days 3–6 within DMSO or RTX conditions (**P < 0.003). **e**, Bone marrow was isolated from wild-type and RTX mice from one femur and the frequency of inflammatory monocytes (CD45⁺, CD11b⁺, Ly-6Cʰigh, Ly-6Gʰigh) and neutrophils (CD45⁺, CD11b⁺, Ly-6Cʰmid, Ly-6Gʰigh) relative to CD45⁺ cells was determined by flow cytometry (n = 5). **f**, Spleens from wild-type and RTX mice were processed for flow cytometry and the frequency of inflammatory monocytes (CD45⁺, CD11b⁺, Ly-6Cʰhigh, Ly-6G⁻) and neutrophils (CD45⁺, CD11b⁻, Ly-6Cʰmid, Ly-6G⁻) relative to CD45⁺ cells was determined (n = 5). NS, not significant.
Extended Data Figure 5 | Leukocyte rolling fractions in skin venules of control and RTX-treated mice analysed by intravital microscopy. Combined results are shown for 26 venules from 5 control mice and for 20 venules from 4 RTX-treated mice. Data are expressed as mean ± s.e.m. of four experiments. NS, not significant.
Extended Data Figure 6 | Dermal γδ T cells represent a major source of IL-17F and IL-22 in skin during IMQ challenge and already express IL-23R at steady state. **a**, Wild-type mice were challenged with IMQ and the total numbers of IL17Fγδ dermal T cells and αβ T cells at 3 days (n = 15) or 6 days (n = 10) were quantified. **b**, Representative flow plots (related to those depicted in Fig. 2g) of gating for IL-22γδ cells within dermal γδ T cells after 6 days of IMQ treatment. **c**, Ears of DMSO- or RTX-treated mice were exposed for 6 days to IMQ and the frequency of IL-17Fαβ and IL-22αβ cells within αβ T cells was determined (n = 5). **d**, Auricular lymph node (aLN) cells from IL-23R<sup>GFP</sup> mice were analysed by flow cytometry for expression of IL-23R<sup>GFP</sup> cells within the γδ T cells and αβ T cell compartment at steady state (representative FACS plot from eight mice analysed). **e**, The ear skin from IL-23R<sup>GFP</sup> mice was digested and analysed by flow cytometry and the distribution of T-cells subsets within IL-23R<sup>GFP</sup> and IL-23R<sup>GFP</sup> fractions of Thy1<sup>+</sup> cells determined (representative FACS plot from eight mice analysed). NS, not significant.
Extended Data Figure 7 | TRPV1\(^+\) nociceptors regulate IMQ- and DNFB-induced dermatitis and are upstream of IL-23.  a–c, After 3 days of IMQ challenge, ears were harvested and processed for total RNA isolation and \(\text{Il}12b\) (a), \(\text{Il}23a\) (b) and \(\text{Il}12a\) (c) mRNA levels were analysed by qPCR (\(n = 5\)).  d, DNFB (0.5% in acetone) was applied topically to DMSO and RTX mice. Time course of change in ear thickness of IMQ-treated ear relative to the contralateral ear is represented (\(n = 5\)). Two-way ANOVA was run to compare ear swelling under DMSO and RTX conditions over time (****\(P < 0.0001\)).  e, Representative FACS plots from ears harvested after 24 h of DNFB application.  f, IL-23\(^{\text{GFP/GFP}}\) mice were treated with RTX and then compared to wild-type and IL-23\(^{\text{GFP/GFP}}\) littermate controls during IMQ treatment. Ear thickness was calculated relative to the contralateral ear (\(n = 5\)).  g, After two IL-23 injections into the ear skin of wild-type and IL-23\(^{\text{GFP/GFP}}\) mice, the frequency of IL-17F\(^+\) cells within dermal \(\gamma \delta\) T cells was determined by flow cytometry (\(n = 5\)).  h, IL-23 was injected twice into the ear skin of vehicle- and RTX-treated mice and the total numbers of IL17A\(^+\) or IL-17F\(^+\) dermal \(\gamma \delta\) T cells per ear were determined by flow cytometry (\(n = 5\)).  NS, not significant.
Extended Data Figure 8 | Selective depletion of migratory and skin-resident myeloid cell subsets in ear skin and gating strategy used for sorting to isolate RNA from MHC-II⁺ cells in skin. **a**, Wild-type mice were treated with anti-Gr-1 (clone RB6-8C5 to deplete neutrophils and inflammatory monocytes) or matched isotype control, challenged with IMQ for 3 days and skin was digested to quantify the total numbers of inflammatory monocytes and neutrophils per ear. Shown are representative plots pre-gated on CD45⁺ cells and quantification of cell numbers from n = 3 mice. **b**, DTX treatment resulted in depletion of both subsets of DDCs as well as Langerhans cells (LCs) but not macrophages. Cells were gated as shown in Extended Data Fig. 8c and normalized to levels in wild-type mice based on the frequency within the CD45⁺ population from n = 4 mice. **c**, Ear skin from naive mice was digested and analysed by flow cytometry for the indicated subsets. Shown is a representative plot pre-gated on CD45⁺ Class II⁺ cells from which further subsets were divided based on CD11b and CD11c expression and then F4/80 and CD103 as indicated. **d**, Total RNA from sorted cells was isolated and qPCR for Il12a relative to Gapdh was performed from naive and IMQ-treated ears after 6 h from n = 20 pooled mice.
Extended Data Figure 9 | DDCs are found in close apposition to Na,1.8⁺ nociceptors and characterization of RTX-treated and Na,1.8-DTA mice.
a, Representative confocal micrographs of CD11c–YFP mice stained for β3-tubulin, Lyve-1 (lymphatics) and CD31 (blood and lymphatic endothelial cells). b, Three-dimensional quantification of DDC proximity to peripheral nerves in naive and 6 h post-IMQ treatment ears binned into contact (<0 µm), proximal (0–7 µm) and distal (>7 µm) fractions as explained in Methods (n of dendritic cells = 200). c, Total RNA from dorsal root ganglia (C1–C4) of littermate control and Na,1.8-DTA mice was isolated and levels of mRNA for Trpv1 (TRPV1), Scn10a (Na,1.8), Tac1 (substance P) and Trpa1 (TRPA1) were determined relative to Gapdh. This demonstrates the efficacy of the Na,1.8-DTA system and combined with the original reference characterizing the pain phenotype of these mice illustrates that a subset of peptidergic TRPV1⁺ nerve fibres is spared. d, Representative confocal micrograph of whole-mount ear skin of vehicle- and RTX-treated mice showing preserved nerve scaffold. e, Representative confocal micrographs of whole-mount ear skin of control and Na,1.8-DTA mice showing preserved nerve scaffold. Although dorsal root ganglia showed a loss of the hallmark ion channels of these nerve subsets (Extended Data Figs 1c, 9c), surprisingly we still observed that RTX mice and Na,1.8-DTA mice maintain a meshwork of nerves in the skin.
Extended Data Figure 10 | Summary diagram.