

Topical tissue nano-transfection mediates non-viral stroma reprogramming and rescue

Daniel Gallego-Perez^{1,2,3,4†}, Durba Pal^{1,4†}, Subhadip Ghatak^{1,4†}, Veysi Malkoc^{3,5}, Natalia Higuera-Castro^{1,4}, Surya Gnyawali^{1,4}, Lingqian Chang^{2,3}, Wei-Ching Liao³, Junfeng Shi^{3,6}, Mithun Sinha^{1,4}, Kanhaiya Singh^{1,4}, Erin Steen¹, Alec Sunyecz^{1,4,5}, Richard Stewart^{1,4}, Jordan Moore^{1,4}, Thomas Ziebro⁶, Robert G. Northcutt⁶, Michael Homsy⁵, Paul Bertani⁷, Wu Lu⁷, Sashwati Roy^{1,4}, Savita Khanna^{1,4}, Cameron Rink^{1,4}, Vishnu Baba Sundaresan⁶, Jose J. Otero^{4,8,9}, L. James Lee^{3,4,5*} and Chandan K. Sen^{1,4*}

Although cellular therapies represent a promising strategy for a number of conditions, current approaches face major translational hurdles, including limited cell sources and the need for cumbersome pre-processing steps (for example, isolation, induced pluripotency)^{1–6}. *In vivo* cell reprogramming has the potential to enable more-effective cell-based therapies by using readily available cell sources (for example, fibroblasts) and circumventing the need for *ex vivo* pre-processing^{7,8}. Existing reprogramming methodologies, however, are fraught with caveats, including a heavy reliance on viral transfection^{9,10}. Moreover, capsid size constraints and/or the stochastic nature of status quo approaches (viral and non-viral) pose additional limitations, thus highlighting the need for safer and more deterministic *in vivo* reprogramming methods^{11,12}. Here, we report a novel yet simple-to-implement non-viral approach to topically reprogram tissues through a nanochannelled device validated with well-established and newly developed reprogramming models of induced neurons and endothelium, respectively. We demonstrate the simplicity and utility of this approach by rescuing necrotizing tissues and whole limbs using two murine models of injury-induced ischaemia.

Recent advances in nuclear reprogramming *in vivo* have opened up the possibility for the development of ‘on-site’, patient-specific, cell-based therapies. We have developed a novel yet simple to implement non-viral approach to topically and controllably deliver reprogramming factors to tissues through a nanochannelled device (Fig. 1). This tissue nano-transfection (TNT) approach allows direct cytosolic delivery of reprogramming factors by applying a highly intense and focused electric field through arrayed nanochannels^{11,13}, which benignly nanoporates the juxtaposing tissue cell membranes and electrophoretically drives reprogramming factors into the cells (Fig. 1a–d). Detailed information regarding the TNT system fabrication process and simulation results is provided in Supplementary Figs 1 and 2. In contrast to current *in vivo* transfection technologies (for example, viruses and conventional tissue bulk electroporation (BEP)), in which gene delivery is highly stochastic in nature and could lead to adverse side effects (such as inflammatory response and cell death)¹⁴, nanochannel-based transfection enables more focused (Fig. 1b,c) and ample (Fig. 1d) reprogramming factor

delivery at the single-cell level, thus making this a powerful tool for deterministic *in vivo* gene transfection and reprogramming^{11,13}.

Experiments with fluorescein amidite (FAM)-labelled DNA on C57BL/6 mice established that TNT can deliver cargo into the skin in a rapid (<1 s) and non-invasive/topical manner (Fig. 1e). We next tested whether TNT-based topical delivery of reprogramming factors could lead to successful skin reprogramming using a robust model where overexpression of *Ascl1/Brn2/Myt1l* (*ABM*) is known to directly reprogram fibroblasts into induced neurons (iNs) *in vitro*^{11,15}. Our findings showed that TNT can not only be used for topical delivery of reprogramming factors (Fig. 1f), but it can also orchestrate a coordinated response that results in reprogramming stimuli propagation (that is, epidermis to dermis) beyond the initial transfection boundary (the epidermis) (Fig. 1g–i), possibly via dispatch of extracellular vesicles (EVs) rich in target gene cDNAs/mRNAs (Fig. 1h,i)¹⁶, among other plausible mechanisms¹⁷. Exposing naive cells to *ABM*-loaded EVs isolated from TNT-treated skin (Fig. 1j–l) established that these EVs can be spontaneously internalized by remote cells and trigger reprogramming (Fig. 1k,l and Supplementary Fig. 3). Moreover, gene expression analysis indicated that intradermal *ABM* EV injection triggered changes in the skin consistent with neuronal induction (Supplementary Fig. 4), as evidenced by increased *Tuj1* expression. The neurotrophic effect of skin-derived *ABM*-loaded EVs was further confirmed in a middle cerebral artery occlusion stroke mouse model (Supplementary Fig. 5)¹⁸.

Successful skin cell reprogramming was verified by immunofluorescence, which showed increased *Tuj1* and neurofilament expression over time (Fig. 1m,n). Further characterization was conducted via genome-wide transcriptome array analysis comparing *in vitro* and *in vivo* derived iNs (Supplementary Fig. 6). Electrophysiological activity, indicative of neuronal excitability, was successfully detected and monitored (*in situ*) in ~50% of the *ABM*-transfected mice (Fig. 1o) through a novel polypyrrole (PPy)-based biosensing platform (Supplementary Fig. 7)^{19,20}. No such activity was detected in any of the control mice. Lineage tracing experiments with a K14-Cre reporter mouse model established that the newly induced neurons partly originated from K14⁺ skin cells (Supplementary Fig. 8). Hair follicles also

¹Department of Surgery, The Ohio State University, Columbus, Ohio 43210, USA. ²Department of Biomedical Engineering, The Ohio State University, Columbus, Ohio 43210, USA. ³Center for Affordable Nanoengineering of Polymeric Biomedical Devices, The Ohio State University, Columbus, Ohio 43210, USA. ⁴Center for Regenerative Medicine and Cell-Based Therapies, The Ohio State University, Columbus, Ohio 43210, USA. ⁵Department of Chemical and Biomolecular Engineering, The Ohio State University, Columbus, Ohio 43210, USA. ⁶Department of Mechanical and Aerospace Engineering, The Ohio State University, Columbus, Ohio 43210, USA. ⁷Department of Electrical and Computer Engineering, The Ohio State University, Columbus, Ohio 43210, USA. ⁸Department of Pathology, The Ohio State University, Columbus, Ohio 43210, USA. ⁹Department of Neuroscience, The Ohio State University, Columbus, Ohio 43210, USA. [†]These authors contributed equally to this work. *e-mail: chandan.sen@osumc.edu; lee.31@osu.edu

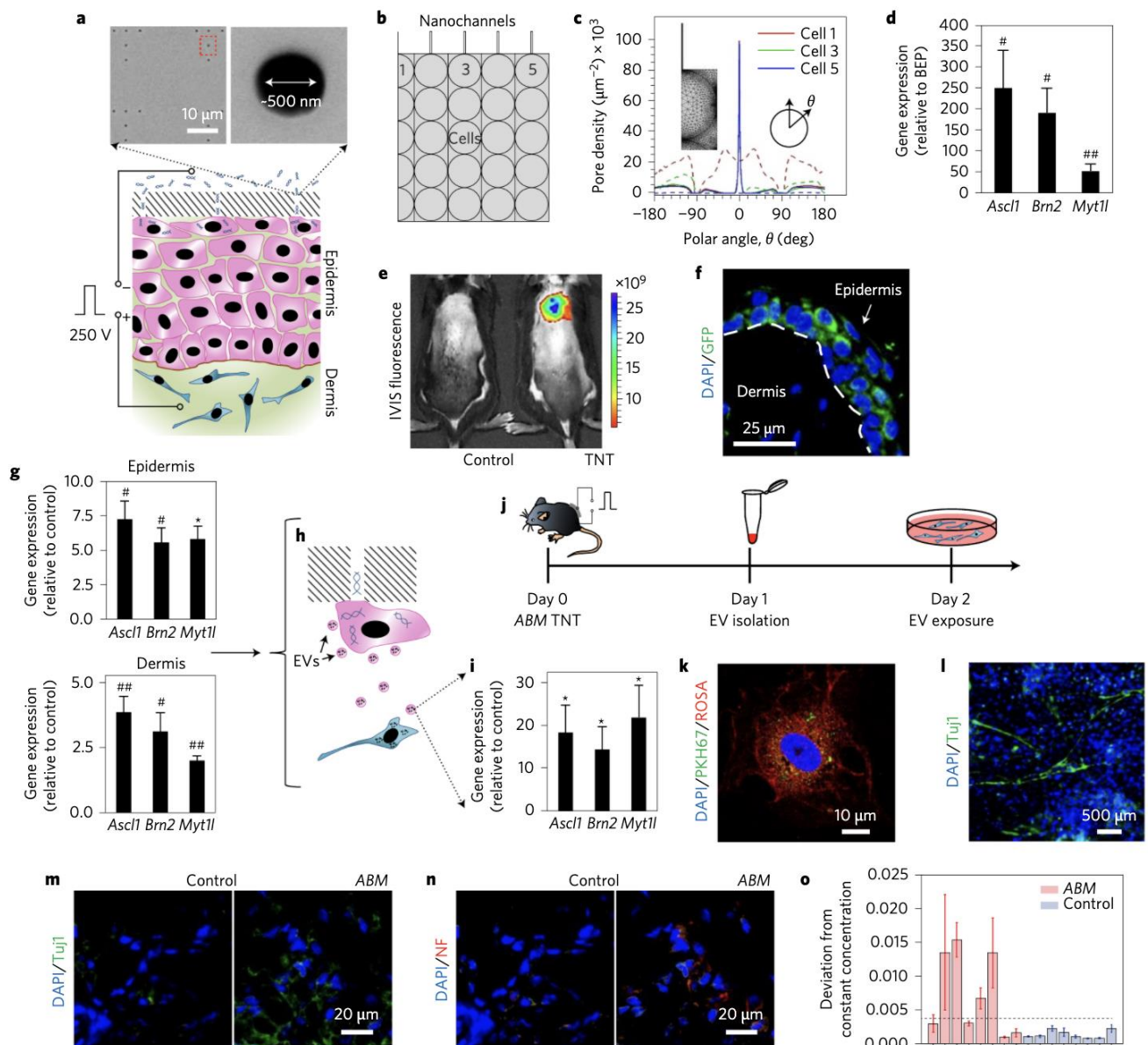


Figure 1 | TNT mediates enhanced reprogramming factor delivery and propagation beyond the transfection boundary. **a**, Schematic diagram of the TNT process on exfoliated skin tissue. Exfoliation is required to remove dead cells from the skin surface. The positive electrode is inserted intradermally, and the negative electrode is put in contact with the cargo solution. A pulsed electric field (250 V, 10 ms pulses, 10 pulses) is then applied across the electrodes to nanoporate exposed cell membranes and inject the cargo directly into the cytosol. Scanning electron micrographs (top) of the TNT platform surface show the nanopore array. **b**, Schematic diagram showing the boundary conditions for simulation purposes. Nanochannels are in direct contact with the outermost cell layer. **c**, Simulation of the poration profile for different cells (cells 1, 3 and 5 from **b**) undergoing TNT (solid lines) versus BEP (dashed lines). The plot shows that TNT leads to focused poration, and BEP results in widespread poration. **d**, ABM expression results for TNT versus BEP 24 h after transfection (n = 5). TNT resulted in superior ABM expression. BEP was conducted via intradermal injection of the ABM plasmids followed by a pulsed electric field. Controls for BEP experiments involved intradermal injections of ABM plasmids with no electric field implementation. **e, f**, Representative *in vivo* imaging system (IVIS) fluorescence (**e**) and confocal microscopy (**f**) images of mouse skin after TNT treatment with labelled DNA and the ABM factors, respectively. GFP is the reporter gene in the *Ascl1* plasmid. **g**, Laser capture microdissection (LCM) and qRT-PCR results of gene expression in the epidermis and dermis (t = 24 h) showing that gene expression propagated beyond the epidermal transfection boundary (n = 5–6). **h**, Schematic diagram illustrating the concept of EV-mediated transfection propagation from epidermis to dermis. **i**, qRT-PCR analysis of the EV cargo showing significant loading of ABM cDNAs/mRNAs (n = 6–8). **j**, Experimental design to confirm whether EVs are a viable vehicle for propagating transfection and reprogramming. **k**, Confocal micrograph showing a mouse embryonic fibroblast (red) that has spontaneously internalized the EVs (green) isolated from TNT-treated skin. **l**, Mouse embryonic fibroblast cultures showing iNs at day 7 after a 24 h exposure to ABM-laden EVs isolated from ABM-transfected skin. **m, n**, Immunostaining results (week 4) showing increased Tuj1 (**m**) and neurofilament (NF) (**n**) expression in the skin after ABM transfection. **o**, Electrophysiological activity shown as a statistically representative bar plot indicating changes in ionic concentration (quantified as average standard deviations from the norm per insertion site) of the extracellular niche as a result of neuronal cell cluster excitability (n = 8, P < 0.05, Fisher's exact test). This average was calculated for 5–10 trials (with 100 sequential discrete measurements per trial) for each ABM or control mouse. Activity was defined as changes in ionic concentration in excess of the baseline (dashed line: experimental noise measured in physiological saline solution). Each bar shows the results collected in individual mice. *P < 0.01 (Dunn's), #P < 0.05 (Tukey test), ##P < 0.05 (Holm-Sidak method).

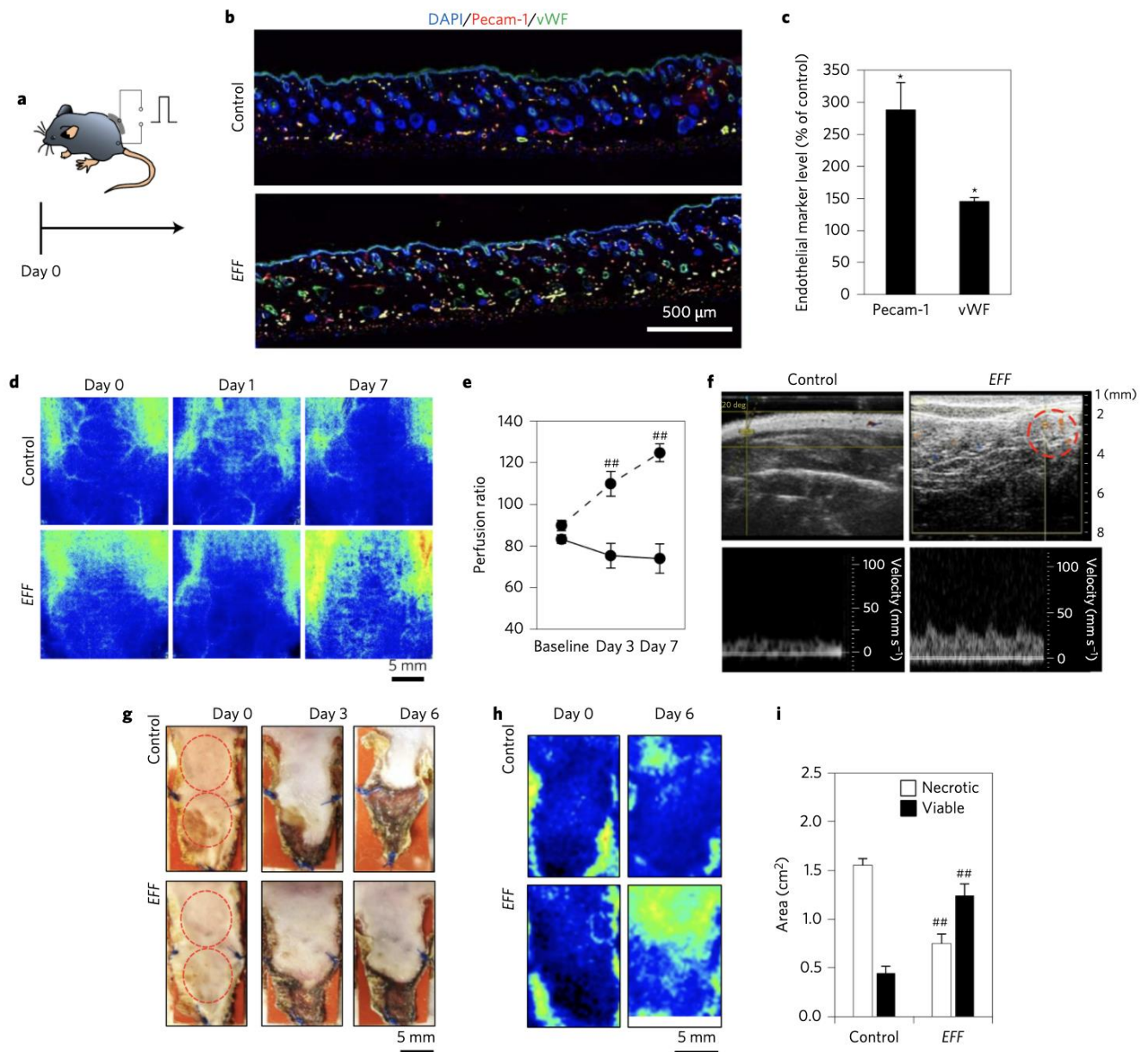


Figure 2 | EFF TNT leads to increased vascularization and rescue of skin tissue under ischaemic conditions. a–c, A one-time treatment of dorsal skin lasting only a few seconds led to increased angiogenesis (Pecam-1, vWF) of skin tissue (day 7) ($n = 3$). **d,e,** High-resolution laser speckle imaging shows enhanced perfusion to the EFF-treated area over time ($n = 5$). **f,** Ultrasound imaging of EFF-treated skin confirmed the presence of superficial blood vessels (dashed circle) with pulsatile behaviour, which suggests successful anastomosis with the parent circulatory system. **g,** Monopodicle flap experiment showing increased flap necrosis for controls compared to EFF-treated skin. **h,** Laser speckle imaging showing increased blood flow to the flapped tissue treated with EFF TNT. **i,** Quantification of flap necrosis ($n = 6$). * $P < 0.05$ (t-test), ** $P < 0.05$ (Holm-Sidak method).

consistently showed marked Tuj1 immunoreactivity, suggesting that follicular cells could participate in the reprogramming process^{21,22}. Additional experiments with a Col1A1-enhanced green fluorescent protein (eGFP) mouse model (Supplementary Fig. 8), where cells with an active Col1A1 promoter (for example, dermal fibroblasts) express eGFP, showed a number of collagen/eGFP⁺ cells in the dermis in a transition phase to Tuj1⁺, thus suggesting a fibroblastic origin for some of the reprogrammed cells in the skin.

Having validated the TNT platform for successful *in vivo* reprogramming using iNs as a case study, we then set out to develop a robust and simple non-viral methodology that would be capable of reprogramming skin cells into induced endothelial cells (iECs). To this end, we first identified and validated (*in vitro*) a set of reprogramming factors—*Etv2*, *Foxc2* and *Fli1* (EFF)—to promote

more rapid and effective reprogramming of somatic cells into iECs (Supplementary Figs 9 and 10) compared to previous reports¹⁰. *In vitro* non-viral transfection and reprogramming experiments¹¹ showed that EFF could reprogram human and mouse primary fibroblasts into iECs rapidly (<1 week) and efficiently (Supplementary Fig. 9).

Once the efficacy of EFF to induce direct endothelial cell reprogramming was established *in vitro*, we then proceeded to test this paradigm *in vivo*. Co-transfection of these three genes into the dorsal skin of C57BL/6 mice resulted in marked stroma reprogramming within a week, as evidenced by a significant increase in Pecam-1 and vWF expression compared to control skin (Fig. 2a–c), in addition to enhanced proliferative activity (Supplementary Fig. 11). Experiments with K14-Cre reporter and Col1A1-eGFP

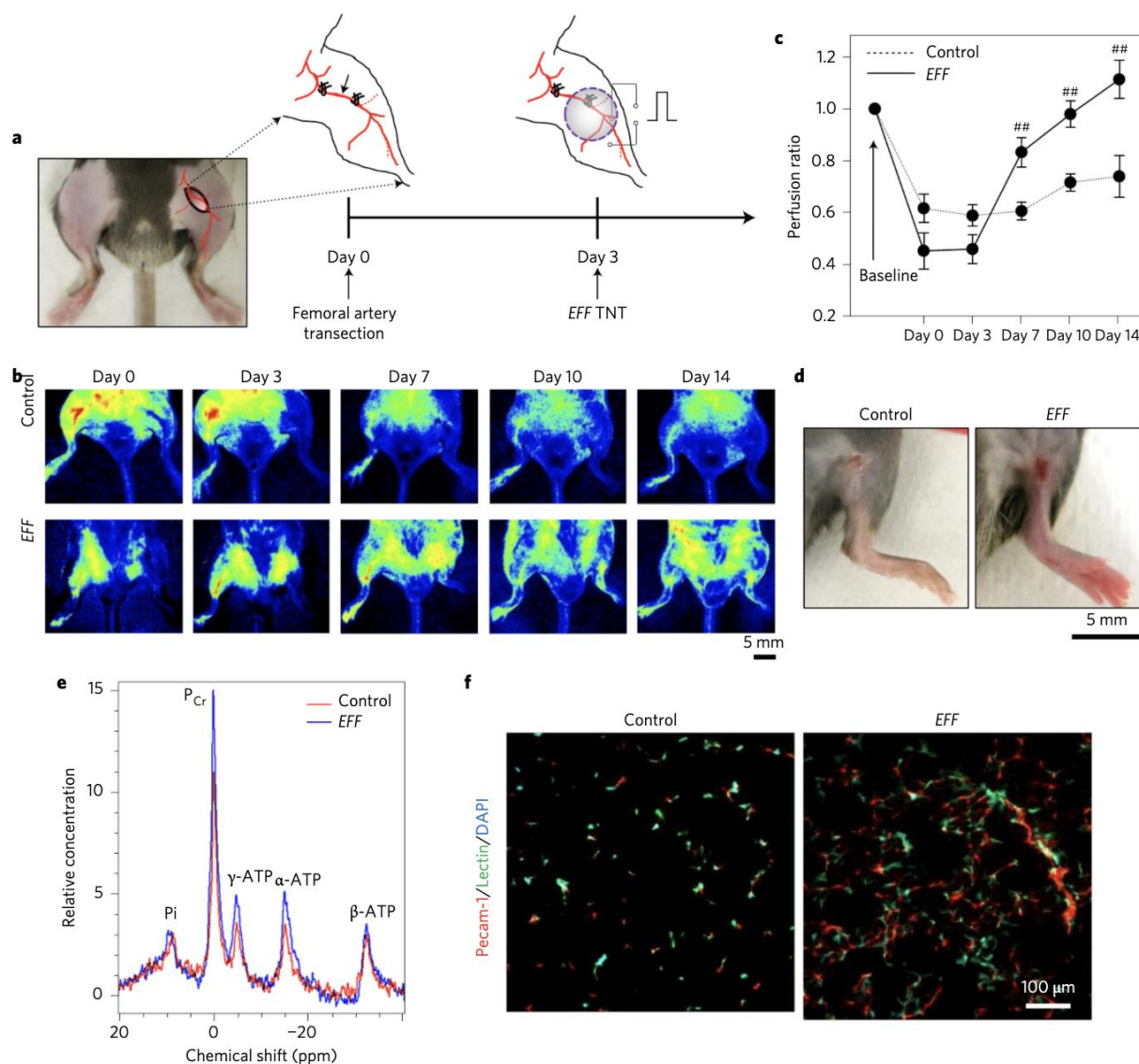


Figure 3 | EFF TNT rescues whole limbs from necrotizing ischaemia. a–c, A one-time treatment of thigh skin lasting only a few seconds led to increased limb reperfusion following transection of the femoral artery. Perfusion was calculated based on the ratio of the ischaemic versus normal/contralateral limb ($n = 5–7$). **d,** Control limbs showing more pronounced signs of tissue necrosis compared to EFF-treated limbs (day 14). **e,** NMR-based measurements of muscle energetics confirmed increased ATP and P_{Cr} levels for EFF-treated limbs compared to controls. **f,** Immunofluorescence analysis of the gastrocnemius muscle showing enhanced angiogenesis. ## $P < 0.05$ (Holm–Sidak method).

mouse models demonstrated that the reprogrammed cell population had, for the most part, a dermal origin (Supplementary Fig. 12). High-resolution laser speckle (HRLS) imaging of dorsal skin showed that TNT-based delivery of EFF enhanced blood flow to the treated area within 3 days (Fig. 2d,e). Ultrasound imaging detected unexpected pulsatile blood flow only 3 mm away from the surface of the skin (Fig. 2f, right), demonstrating successful anastomosis of the newly formed blood vessels with local functional cutaneous arteries. Note that, in control mice, blood vessels were not typically detected near the skin surface (Fig. 2f, left).

Once the robustness of the EFF cocktail to induce vascular endothelium was demonstrated both *in vitro* and *in vivo*, we studied whether EFF TNT-mediated topical skin reprogramming could lead to functional reperfusion of ischaemic tissues. We first tested this concept with a full-thickness $2 \times 1 \text{ cm}^2$ monopedicle dorsal skin flap in C57BL/6 mice, whereby blood supply to the flapped tissue only came from the cephalad attachment (Fig. 2g). Laser speckle monitoring after EFF treatment showed higher blood

perfusion compared to control flaps (Fig. 2h). As expected, control flaps showed significant signs of tissue necrosis (Fig. 2g,top,i). This tissue damage was significantly limited in response to EFF transection. Thus, TNT-mediated EFF delivery and subsequent stroma reprogramming effectively counteracted tissue necrosis under ischaemic conditions.

Finally, to verify whether TNT-based delivery of EFF could lead to whole limb rescue, we tested TNT in a hindlimb ischaemia C57BL/6 mouse model (Fig. 3a). EFF TNT was conducted on the inner thigh skin three days after transection of the femoral artery. Laser speckle monitoring recorded a significant reduction in blood flow to the limb immediately after surgery (Fig. 3b). Compared to control ischaemic limbs, EFF-treated limbs showed improved perfusion as early as day 7 post-TNT (Fig. 3b,c). HRLS imaging demonstrated an increased incidence of small collaterals in the EFF-treated limbs compared to controls (Supplementary Fig. 13). Macroscopic analysis showed more pronounced signs of tissue necrosis in the control limbs compared to the EFF-treated

ones (Fig. 3d). Additional experiments in BALB/c mice, which have a tendency to experience more deleterious side effects from injury-induced limb ischaemia^{23,24}, showed that *EFF* transfection also led to successful limb perfusion and minimized the incidence of necrosis and auto-amputation (Supplementary Fig. 14). Muscle energetics testing by NMR imaging showed increased levels of adenosine triphosphate (ATP) and phosphocreatine (P_{Cr}) in *EFF*-treated limbs compared to controls (Fig. 3e). Immunofluorescence analysis revealed marked revascularization far beyond the treatment area. Angiogenesis was also induced in more distal locations within the limb, such as the gastrocnemius muscle (Fig. 3f and Supplementary Fig. 15). Although the underlying mechanisms for this response need to be elucidated further, we found that autologous EVs, isolated from *EFF*-treated dorsal skin, have the potential to induce blood vessel formation when injected directly into the gastrocnemius muscle in a hindlimb ischaemia mouse model (Supplementary Fig. 16). Parallel *in vitro* experiments demonstrated that these EVs can induce reprogramming in naive cells (Supplementary Fig. 17). We thus propose that EVs dispatched from *EFF*-treated tissue serve as a mediator of the propagation of pro-iEC reprogramming signals. PCR analysis revealed that in addition to the transduced *EFF* cDNAs/mRNAs, the EVs also appeared to be preloaded with pro-angiogenic vascular endothelial growth factor and basic fibroblast growth factor mRNAs (Supplementary Fig. 16). This suggests that EVs derived from *EFF*-treated skin not only represent a viable mechanism for propagating *EFF* reprogramming signals throughout the target tissue, but may also play a role in niche preconditioning by spreading pro-angiogenic signals within the first hours after transfection.

Here, we show for the first time that TNT can be used to deliver reprogramming factors into the skin in a rapid, highly effective and non-invasive manner. Such TNT delivery leads to tailored skin tissue reprogramming, as demonstrated with well-established and newly developed reprogramming models of iNs and iECs, respectively. TNT-induced skin-derived iECs rapidly formed blood vessel networks that successfully anastomosed with the parent circulatory system and restored tissue and limb perfusion in two murine models of injury-induced ischaemia. TNT-based tissue reprogramming has the potential to ultimately enable the use of a patient's own tissue as a prolific immunosurveilled bioreactor to produce autologous cells that can resolve conditions locally/on site or distally upon harvesting. This simple to implement TNT approach, which elicits and propagates powerfully favourable biological responses through a topical one-time treatment that only lasts seconds, could also find applications beyond plasmid DNA-based reprogramming strategies, including oligo RNA (for example, miRs and siRNAs)-mediated reprogramming (Supplementary Fig. 18)²⁵, gene modulation, editing and so on.

Methods

Methods and any associated references are available in the [online version of the paper](#).

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Author contributions

TNT platform design, implementation and optimization (for different applications) was performed by D.G.-P., L.J.L., D.P., S.G. and C.K.S. TNT chip fabrication, transductions,

cell/tissue imaging, transgenic mouse experiments and histology were performed by D.G.-P., D.P., S.G., N.H.-C., V.M., S.G., L.C., M.S., E.S., A.S., J.M., P.B., W.L., J.J.O., L.J.L. and C.K.S. Electrophysiological activity measurements were conducted by T.Z., R.G.N., V.B.S. and J.J.O., with support from S.G. and N.H.-C. Global gene expression analyses were conducted by S.R., S.K., K.S. and C.K.S. TNT chip simulations were conducted by W.-C.L., J.S., L.C., D.G.-P. and L.J.L. S.R. oversaw and participated in the LCM work. Stroke recovery experiments were conducted by D.G.-P., V.M., A.S., R.S., M.H., S.K., C.R. and C.K.S. The manuscript was written by D.G.-P., L.J.L., D.P., S.G. and C.K.S. C.K.S. and L.J.L. jointly supervised this work.

Additional information

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Competing financial interests

The authors declare no competing financial interests.

Methods

TNT platform fabrication. TNT devices were fabricated from thinned (~200 µm) double-side-polished (100) silicon wafers (Supplementary Fig. 1). Briefly, ~1.5-µm-thick layers of AZ5214E photoresist were first spin-coated on the silicon wafers at ~3,000 r.p.m. Nanoscale openings were subsequently patterned on the photoresist using a GCA 6100C stepper. Up to 16 dies of nanoscale opening arrays were patterned on each 100 mm wafer. These openings were then used as etch masks to drill ~10-µm-deep nanochannels on the silicon surface using deep reactive ion etching (DRIE) (Oxford Plasma Lab 100 system). Optimized etching conditions were as follows: SF₆ gas, 13 s/100 s.c.c.m. gas flow/700 W inductance-coupled plasma (ICP) power/40 W RF power/30 mT automatic pressure controller (APC) pressure; C₄F₈ gas, 7 s/100 s.c.c.m. gas flow/700 W ICP power/10 W RF power/30 mT APC pressure. Microscale reservoirs were then patterned on the back side of the wafers by means of contact photolithography and DRIE. Finally, a ~50-nm-thick insulating/protective layer of silicon nitride was deposited on the TNT platform surface.

Animal husbandry. C57BL/6 mice were obtained from Harlan Laboratory. B6.129 (Cg)-Gt(ROSA)26Sortm4(ACTB-tdTomato,-EGFP)Luo/J mice obtained from Jackson Laboratories were bred with K14cre to produce K14cre/Gt(ROSA)26Sortm4(ACTB-tdTomato-EGFP)Luo/J mice. pOBCol3.6GFPTz mice were gifts from T. Wilgus (The Ohio State University). repTOP *mitoIRE* mice were obtained from Charles River Laboratories. Fsp1-Cre mice were donated by A. Deb (University of California, Los Angeles). Fsp1-Cre mice were crossed with the B6.Cg-Gt(ROSA)26Sor^{(tm9(CAG-tomato)Hze/j)} mice (Jackson Laboratories) to generate mice with tdTomato expression specific to fibroblasts. All mice were male and 8–12 weeks old at the time of the study. Genotyping PCR for ROSAmT/mG mice was conducted using primers oIMR7318-CTC TGC TGC CTC CTG GCT TCT, oIMR7319-CGA GGC GGA TCA CAA GCA ATA and oIMR7320-TCA ATG GGC GGC GGT CGT T, and the K-14 Cre transgene was confirmed using primers oIMR1084-GCG GTC TGG CAG TAA AAA CTA TC and oIMR1085-GTG AAA CAG CAT TGC TGT CAC TT. Genotyping PCR for Fsp1-Cre mice was conducted using the following primers: forward-CTAGGCCACAGAATTGAAAGATCT and reverse-GTAGGTGGAATTCTAGCATCATCC (for wild type, product length = 324 bp) and forward-GCGGCTGGCAGTAAAAACTATC and reverse-GTGAACAGCATTGCATTGCTGCTACTT (for Cre transgene, product length = 100 bp). td tomato was confirmed using the following primers: forward-AAGGGAGCTGCAGTGAGTA and reverse-CCGAAAATCTGTGGGAAATC (for wild type, product length = 196 bp) and forward-GGCATTAAGCAGCGTATCC and reverse-CTGTTCTGTACGGCATGG (mutant type, product length = 297 bp). All animal studies were performed in accordance with protocols approved by the Laboratory Animal Care and Use Committee of The Ohio State University. No statistical method was used to predetermine the sample size. Power analysis was not necessary for this study. The animals were tagged and grouped randomly using a computer-based algorithm (www.random.org).

Mammalian cell culture and *in vitro* reprogramming. Primary human adult dermal fibroblasts (ATCC PCS-201-012) were purchased, mycoplasma-free and certified, directly from ATCC. No further cell line authentication/testing was conducted. The cells were expanded in fibroblast basal medium supplemented with fibroblast growth kit–serum-free (ATCC PCS 201–040) and penicillin/streptomycin. E12.5–E14 mouse embryonic fibroblasts (MEFs) were cultured in DMEM/F12 supplemented with 10% fetal bovine serum. Non-viral cell transfection and reprogramming experiments were conducted via three-dimensional nanochannel electroporation (NEP), as described previously¹¹. Briefly, the cells were first grown to full confluency overnight on the 3D NEP device. Subsequently, a pulsed electric field was used to deliver a cocktail of plasmids (0.05 µg µl⁻¹) into the cells consisting of a 1:1:1 mixture of *Fli1:Etv2:Foxc2*. The cells were then collected 24 h after plasmid delivery, placed in EBM-2 basal medium (CC-3156, Lonza) supplemented with EGM-2 MV SingleQuot kit (CC-4147, Lonza) and further processed for additional experiments/measurements. *Etv2* and *Fli1* plasmids were donated by A. Ferdous (Department of Internal Medicine, UT Southwestern Medical Center, Texas). *Foxc2* plasmids were donated by T. Kume (Department of Medicine–Cardiology and Pharmacology, Northwestern University–FCVRI, Chicago).

***In vivo* reprogramming.** The areas to be treated were first depilated 24–48 h before TNT. The skin was then exfoliated to eliminate the dead/keratin cell layer and expose nucleated cells in the epidermis. The TNT devices were placed directly over the exfoliated skin surface. *ABM* or *EFF* plasmid cocktails were loaded in the reservoir at a concentration of 0.05–0.1 µg µl⁻¹. A gold-coated electrode (the cathode) was immersed in the plasmid solution, and a 24G needle counter-electrode (the anode) was inserted intradermally, juxtaposed to the TNT platform surface. Pulsed electrical stimulation (10 pulses, 250 V in amplitude, duration of 10 ms per pulse) was then applied across the electrodes to nanoporate the exposed cell membranes and drive the plasmid cargo into the cells through the nanochannels. *ABM* plasmids were mixed at a 2:1:1 molar ratio as described previously¹¹. Unless otherwise specified, control specimens involved TNT treatments with a blank, phosphate buffer saline (PBS)/mock plasmid solution (Supplementary Fig. 19).

Electrophysiological activity measurements. The general principle of extracellular recordings was used to detect electrophysiological activity in the skin. Chronoamperometric measurements were conducted using PPy-based probes to detect neuronal excitability through two small incisions on the skin of sedated mice. For details see Supplementary Fig. 7.

Middle cerebral artery occlusion stroke surgery and analysis. Transient focal cerebral ischaemia was induced in mice by middle cerebral artery occlusion (MCAO), achieved using the intraluminal filament insertion technique as previously described¹⁸. Magnetic resonance imaging (MRI) images were used to determine infarct size as a percentage of the contralateral hemisphere after correcting for oedema.

Ischaemic skin flaps. Monopodicle (that is, random-pattern) ischaemic flaps measuring 20 mm × 10 mm were created on the dorsal skin of C57BL/6 mice. Briefly, eight- to ten-week-old mice were anaesthetized with 1–3% isoflurane. The dorsa were depilated, cleaned and sterilized with betadine. A monopodicle flap was created on the dorsal skin of the mice by making 20-mm-long full-thickness parallel incisions 10 mm apart. The bottom part of the skin was cut to create a free hanging flap. Flap edges were cauterized. A 0.5 mm silicon sheet was placed under the flap and then sutured to the adjacent skin with 5–0 ethicon silk suture. Finally, a single dose of buprenorphine was administered subcutaneously to control pain. Laser speckle imaging (Perimed) was conducted 2 h post-surgery to confirm successful blood flow occlusion. TNT-based transfections were conducted 24 h before skin flapping.

Hindlimb ischaemia surgery. Unilateral hind-limb ischaemia was induced via occlusion and subsequent transection of the femoral artery²⁶. Briefly, eight- to ten-week-old mice were anaesthetized with 1–3% isoflurane, and placed supine under a stereomicroscope (Zeiss OPMI) on a heated pad. The femoral artery was exposed and separated from the femoral vein via a ~1 cm incision. Proximal and distal end occlusions were induced with 7–0 silk suture, which was then followed by complete transection of the artery. Finally, a single dose of buprenorphine was administered subcutaneously to control pain. Laser speckle imaging (PeriCam PSI High Resolution, PeriMed) was conducted 2 h post-surgery to confirm successful blood flow occlusion.

Isolation of EVs. EVs were isolated from 12-mm-diameter skin biopsies, which were collected in optimal cutting temperature compound (OCT) blocks and stored frozen for later use. Briefly, the blocks were thawed and washed with PBS to eliminate the OCT. Following removal of the fat tissue with a scalpel, the skin tissue was minced into ~1 mm pieces and homogenized with a microgrinder in PBS. After centrifugation at 3,000g, an Exoquick kit was used at a 1:5 ratio (Exoquick: supernatant) to isolate EVs from the supernatant for 12 h at 4 °C. EVs were precipitated via centrifugation at 1,500g for 30 min. Total RNA was then extracted from the pellets using a Mirvana kit (Life Technologies) following the recommendations of the manufacturer.

DNA plasmid preparation. Plasmids were prepared using a plasmid DNA purification kit (Qiagen Maxi-prep, cat. no. 12161 and Clontech Nucleobond cat. no. 740410). DNA concentrations were obtained from a Nanodrop 2000c Spectrophotometer (Thermoscientific). For a list of plasmid DNA constructs and their original sources see Supplementary Table 1.

LCM and quantitative real-time PCR. LCM was performed using a laser microdissection system from PALM Technologies (Zeiss). Specific regions of tissue sections, identified based on morphology and/or immunostaining, were cut and captured under a ×20 ocular lens. The samples were catapulted into 25 µl of cell direct lysis extraction buffer (Invitrogen). Approximately 1,000,000 µm² of tissue area was captured into each cap, and the lysate was then stored at –80 °C for further processing. Quantitative real-time (qRT)-PCR of the LCM samples was performed from cell direct lysis buffer following the manufacturer's instructions. For a list of primers see Supplementary Table 2.

Immunohistochemistry and confocal microscopy. Tissue immunostaining was carried out using specific antibodies and standard procedures. Briefly, OCT-embedded tissue was cryosectioned at 10 µm thick, fixed with cold acetone, blocked with 10% normal goat serum and incubated with specific antibodies (Supplementary Table 3). The signal was visualized by subsequent incubation with appropriate fluorescence-tagged secondary antibodies (Alexa 488-tagged α-guinea pig, 1:200; Alexa 488-tagged α-rabbit, 1:200; Alexa 568-tagged α-rabbit, 1:200) and counter-stained with DAPI. Lectin-based visualization of blood vessels was conducted via tail vein injection of FITC-labelled lectin 30 min before tissue collection. Images were captured by a laser scanning confocal microscope (Olympus FV 1000 filter/spectral).

IVIS imaging. The animals were imaged under anaesthesia using an IVIS Lumina II optical imaging system. repTOP *mitoIRE* mice were pre-injected with substrate luciferin (potassium salt of beetle luciferin, Promega) at a dose of 100 mg kg⁻¹,

5–10 min before imaging. Overlay images with luminescence images were made using Living Image software.

MRI of stroke brains. Magnetic resonance angiography was used to validate our MCAO model in mice and to optimize the occluder size and the internal carotid artery insertion distance for effective MCAO. T2-weighted MRI was performed on anaesthetized mice 48 h after MCA–reperfusion using a 9.4 T MRI (Bruker Corporation, Bruker BioSpin). Images were acquired using a Rapid Acquisition with Relaxation Enhancement (RARE) sequence using the following parameters: field of view (FOV) 30 × 30 mm², acquisition matrix 256 × 256, TR 3,500 ms, TE 46.92 ms, slice gap 1.0 mm, rare factor 8, number of averages 3. The resolution was 8.5 pixels per mm. Raw MR images were converted to the standard DICOM format and processed. After appropriate software contrast enhancement of images using Osirix v.3.4, digital planimetry was performed by a masked observer to delineate the infarct area in each coronal brain slice. Infarct areas from brain slices were summed, multiplied by slice thickness, and corrected for oedema-induced swelling as previously described to determine the infarct volume¹⁸.

Analysis of muscle energetics. Muscle energetics was evaluated by NMR spectroscopy measurements on a 9.4 T scanner (Bruker BioSpec) using a volume coil for RF transmission and a 31P coil for reception²⁷. *In vivo* imaging was conducted in a custom-made 1H/31P transceiver coil array. Data were acquired using a single pulse sequence. The raw data were windowed for noise reduction and Fourier-transformed to the spectral domain.

Ultrasound-based imaging and characterization of blood vessels. Blood vessel formation was parallel monitored via ultrasound imaging. Briefly, a Vevo 2100 system (Visual Sonics) was used to obtain ultrasound images on the B-mode with an MS 250 linear array probe²⁸. Doppler colour flow imaging was implemented to monitor and quantify the blood flow characteristics under systole and diastole.

GeneChip probe array and ingenuity pathway (IPA) analyses. LCM was used to prepare tissue isolates enriched for *in vivo*-derived iNs from ABM-transfected mouse skin^{29,30}. Tissue isolates were processed in lysis buffer using a PicoPure RNA Isolation Kit (ThermoFisher). RNA extraction, target labelling, GeneChip and data analysis were performed as described previously^{29–31}. The samples were hybridized to Affymetrix Mouse transcriptome Array 1.0 (MTA1.0). The arrays were washed and scanned with the GeneArray scanner (Affymetrix) at The Ohio State University facilities as described already^{29,31}. Expression data have been submitted to the Gene

Expression Omnibus (GEO, <http://www.ncbi.nlm.nih.gov/geo>) under series accession no. GSE92413. Raw data were normalized using RMA16 and analysed using Genespring GX (Agilent). Additional data processing was performed using dChip software (Harvard University)^{29,31}. Functional annotation of the similar genes across groups was performed using IPA analysis.

Statistical analysis. Samples were coded, and data collection was performed in a blinded fashion. Data are reported as mean ± standard error of 3–8 biological replicates. Unsuccessful transfections (for example, due to poor contact between the skin and the nanochannels, or nanochannel clogging) were excluded from the analysis. Experiments were replicated at least twice to confirm reproducibility. Comparisons between groups were made by analysis of variance (ANOVA). Statistical differences were determined using parametric/non-parametric tests, as appropriate, with SigmaPlot version 13.0.

Data availability. GeneChip expression data can be accessed through the Gene Expression Omnibus. Additional data are available from the corresponding authors upon reasonable request.

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