The circadian molecular clock creates epidermal stem cell heterogeneity

Peggy Janich1, Gloria Pascual1, Anna Merlos-Suárez2, Eduard Batlle1,2, Jürgen Ripperger4, Urs Albrecht4, Karl Obrietan5, Luciano Di Croce1,3 & Salvador Aznar Benitah1,3

M murine epidermal stem cells undergo alternate cycles of dormancy and activation, fuelling tissue renewal. However, only a subset of stem cells becomes active during each round of morphogenesis, indicating that stem cells coexist in heterogeneous responsive states. Using a circadian–clock reporter–mouse model, here we show that the dormant hair–follicle stem cell niche contains coexisting populations of cells at opposite phases of the clock, which are differentially predisposed to respond to homeostatic cues. The core clock protein Bmal modulates the expression of stem cell regulatory genes in an oscillatory manner, to create populations that are either predisposed, or less prone, to activation. Disrupting this clock equilibrium, through deletion of Bmal (also known as Arntl) or Per1/2, resulted in a progressive accumulation or depletion of dormant stem cells, respectively. Stem cell arrhythmia also led to premature epidermal ageing, and a reduction in the development of squamous tumours. Our results indicate that the circadian clock fine-tunes the temporal behaviour of epidermal stem cells, and that its perturbation affects homeostasis and the predisposition to tumorigenesis.

Epidermal stem cells ensure that skin homeostasis is maintained. Murine epidermal stem cells are located either at the permanent portion of the hair follicle—termed the bulge—and are exclusively responsible for hair cycling1–3, or at the junction between the epidermis and the hair follicle (isthmus), and feed into the epidermis and sebaceous glands4–7. In addition, a continuous proliferation of basal interfollicular epidermal cells ensures daily epidermal maintenance8. The circadian molecular clock creates an oscillatory pattern of clock activity with a ratio of 95:5 venusbright:venusdim cells, which could be subdivided into cells with highest mean fluorescence intensity of venus (venusbright), and those with no venus fluorescence (venusdim), as determined by immunohistochemistry (Supplementary Fig. 1), and fluorescence-activated cell sorting (FACS) (Fig. 1a, b). At this stage, the bulge contained approximately equal numbers of venusbright and venusdim stem cells (Fig. 1a, b). As hair follicles synchronously progressed into anagen (from P20 to P31), the proportion of venusbright bulge cells steadily increased, reaching a proportion of 90% venusbright to 10% venusdim at the peak of follicle growth (Fig. 1a, b). Conversely, the basal layer of the interfollicular epidermis (6 integrinbright/CD34−) showed a homogeneous pattern of clock activity with a ratio of 95:5 venusbright:venusdim cells, irrespective of the day analysed (Supplementary Fig. 2). Heterogeneity of the clock phase in bulge cells during the telogen to anagen transition was also observed with another independently generated reporter line, Per1–GFP, as shown by three-dimensional whole-mount GFP fluorescence imaging of tail epidermis (Supplementary Fig. 4a, b)21. This circadian variation was further confirmed.

Circadian heterogeneity in hair–follicle stem cells

We monitored the activity of the clock in epidermal stem cells by means of a reporter bacterial artificial chromosome (BAC) transgenic mouse, in which the expression of fluorescent venus is under the regulation of the full-length promoter of the Per1 gene22. Venus expression mirrors the endogenous oscillation of the clock in the suprachiasmatic nuclei, where the central pacemaker resides, thereby establishing its bona fide circadian reporter activity22. We first studied the behaviour of the clock in the dorsal skin of Per1–venus mice collected between postnatal days (P)19 and P31, when hair follicles synchronously transit from the dormant to the growth phase.

At P19, a stage in which the bulge is predominantly dormant, the bulge cells expressing CD34 and the highest levels of 6 integrin (6 integrinbright/CD34−) contained a continuum of venus-expressing populations, which could be subdivided into cells with highest mean fluorescence intensity of venus (venusbright), and those with no venus fluorescence (venusdim), as determined by immunohistochemistry (Supplementary Fig. 1), and fluorescence-activated cell sorting (FACS) (Fig. 1a, b). At this stage, the bulge contained approximately equal numbers of venusbright and venusdim stem cells (Fig. 1a, b). As hair follicles synchronously progressed into anagen (from P20 to P31), the proportion of venusbright bulge cells steadily increased, reaching a proportion of 90% venusbright to 10% venusdim at the peak of follicle growth (Fig. 1a, b). Conversely, the basal layer of the interfollicular epidermis (6 integrinbright/CD34−) showed a homogeneous pattern of clock activity with a ratio of 95:5 venusbright:venusdim cells, irrespective of the day analysed (Supplementary Fig. 2). Heterogeneity of the clock phase in bulge cells during the telogen to anagen transition was also observed with another independently generated reporter line, Per1–GFP, as shown by three-dimensional whole-mount GFP fluorescence imaging of tail epidermis (Supplementary Fig. 4a, b)21. We next verified whether these clock stem cells states showed circadian rhythmicity. FACS analysis indicated that the expression of venus in bulge stem cells in anagen (P27), or interfollicular epidermis basal cells, followed a circadian pattern, irrespective of whether the mice were maintained in 12 h light/dark cycles, or in constant darkness under free running conditions (Fig. 1c and Supplementary Fig. 4a, b)21. This circadian variation was further confirmed.
The clock regulates stem cell genes

We then compared the global transcriptomes of purified venusbright and venusdim bulge cells from the dorsal skin of P18–19 mice (6bright/CD34+/venusbright and 6bright/CD34+/venusdim). As expected, both populations showed differential expression of core circadian transcripts, such as CRY2, PER1, NR1D1, RORβ, DECO2 and E4BP4 (also known as Nfl3) (Fig. 1d and Supplementary Table 1). Intriguingly, although the bulge is inactive at P18–19, both populations differed in the expression of a significant number of genes previously shown to constitute the bulge signature, including CD34, Sox9, Dkk3, Fzd1, Fzd2, Fzd3, Fzd7, Chx11, Lgr5, Lgr6, Smad9, Smad7, Smad5, Ltbp2, Ltbp3, Ltbp4, Lrh1, Gfl2, Rgs16, Pde4a, and Pde4g5 (Fig. 1d, e, and Supplementary Table 1). Other pathways differentially expressed relevant for bulge behaviour related to integrins, Notch, TGF-β and Shh, among others (Fig. 1d, e and Supplementary Table 1).

Thus, the coexisting clock states of dormant bulge stem cells correlated with differential expression of key epidermal homeostasis genes. Promoter analysis revealed that several of these genes, encoding for proteins known to control bulge dormancy, activation, differentiation and niche interactions, contained several putative Bmal1/Clock-binding sites within their proximal and distal promoter regions (Supplementary Table 2). These included the Wnt signalling factors Dab2, Left1, Dkk3, Fzd2, Sox9, Lhx2 and Tcf4; TGF-β regulators such as Smad7, Lefty, Smurfl and Smad9; and Itgα6 as well as modulators of Bmp and Notch signalling (Supplementary Table 2). We confirmed by chromatin immunoprecipitation (ChIP) that Bmal1/Clock bound to these promoters in intact adult tail epidermis (Supplementary Fig. 5a), and that the binding of Bmal1 was circadian (Fig. 2). Chromatin occupancy of Bmal1 to these genes was also confirmed by time-lapse in vivo confocal microscopy of venus fluorescence in dorsal skin explants biopsied from adult Per1−/venus mice, and GFP fluorescence in whole mounts of tail epidermis of Per1−/GFP mice (Supplementary Fig. 4c, d).

Clock arrhythmia affects homeostasis

We next sought to study the biological significance of this clock mechanism in epidermal stem cells in vivo. To this end, we generated mice with a conditional deletion of Bmal1 in the keratin-14+ basal keratinocyte compartment (K14Cre/Bmal1LoxP/LoxP), hereafter referred to as Bmal1KO (Supplementary Fig. 6a, b). Deletion of Bmal1 causes circadian arrhythmicity without the need to perturb any other core circadian member, and its ubiquitous deletion causes premature ageing, including defects in adult skin morphogenesis. We crossed Bmal1KO and Per1−/venus mice, and verified that the circadian clock of bulge stem cells and basal interfollicular epidermal cells was arrhythmic, and permanently skewed towards a clocklow (venusdim) state (Supplementary Fig. 6c).

Bulge stem cells and primary keratinocytes isolated from the dorsal skin of Bmal1KO mice expressed lower transcript levels of Wnt-related

Figure 1 | The molecular clock regulates the expression of the bulge stem cell signature. a, b. Percentages of venusbright and venusdim cells in the bulge by FACS (n = 6). c. Venus mean fluorescence intensity of bulge cells in mice kept under 12 h light/12 h dark (LD) and 12 h dark/12 h dark (DD) conditions (n = 2). d. Heatmap of selected genes from arrays of venusbright and venusdim bulge cells of P19 mice (n = 3). e. Validation of microarray data by real-time PCR. Fold change is shown as relative to venusdim cells after normalization to pumilio 1 (Pum1) (n = 2; pool of 6 mice per replicate). Results in b, c and e are shown as mean ± s.e.m. *P < 0.05, **P < 0.01, ***P < 0.001 (two-tailed Student’s t-test). Clock-binding sites within their proximal and distal promoter regions (Supplementary Table 2). These included the Wnt signalling factors Dab2, Left1, Dkk3, Fzd2, Sox9, Lhx2 and Tcf4; TGF-β regulators such as Smad7, Lefty, Smurf2 and Smad9; and Itgα6 as well as modulators of Bmp and Notch signalling (Supplementary Table 2). We confirmed by chromatin immunoprecipitation (ChIP) that Bmal1/Clock bound to these promoters in intact adult tail epidermis (Supplementary Fig. 5a), and that the binding of Bmal1 was circadian (Fig. 2). Chromatin occupancy of Bmal1 to these genes was also confirmed in FACS-sorted bulge stem cells (Supplementary Fig. 5b).
genes including Dab2, Dkk3, Lef1 and Wnt10a, than control stem cells (Fig. 3a and Supplementary Fig. 5c). In addition, they contained lower messenger RNA levels of TGF-β inhibitors, and higher amounts of Tgbr2 and Smad3 (Fig. 3a and Supplementary Fig. 5c). The expression of Wnt and TGF-β-related factors in purified bulge stem cells varied within a 12 h period in wild-type mice, but not Bmal1KO mice (Supplementary Fig. 5d). Accordingly, the hair follicles of Bmal1KO mice showed the same differences at the protein level, as exemplified by immunohistochemical analysis of Sox9, Lef1, phospho-Smad2 and TGFβR2 in dorsal skin sections, and western blot from primary keratinocytes (Fig. 3b, d).

Altogether, these results indicated that the clock machinery might endow subpopulations of epidermal stem cells with different predispositions to respond to dormancy or activation stimuli, such as TGF-β and Wnt. In line with this, the hair follicle bulge and inter-follicular epidermis of Bmal1KO mice contained higher levels of active phospho-Smad2 (Fig. 3b, d). Moreover, Bmal1KO keratinocytes were more responsive to TGF-β treatment than control keratinocytes (Fig. 3c). We could not detect any Wnt activity using the Wnt-specific reporter containing binding sites for TCF/Lef proteins (TOP-Flash) in our primary cultures of mouse keratinocytes stimulated with the GSK3β inhibitor 6-bromoindirubin-3-oxime (BIO), or purified Wnt3a, in accordance with previous reports5; this prevented us from further studying the effect of Bmal1 deletion on Wnt responsiveness (Supplementary Fig. 7).

When plated at clonal density, Per1–venus<sup>bright</sup> bulge stem cells and basal epidermal cells showed a higher growth potential than the corresponding venus<sup>dim</sup> population, further suggesting that the clock<sup>bright</sup> state is more prone to become activated than the clock<sup>low</sup> counterpart (Fig. 3e). Additional data confirmed this hypothesis. First, the hair follicle bulges of Bmal1KO mice, which are permanently locked in the clock<sup>low</sup> state, contained fewer proliferative cells and a higher number of long-term label (BrdU)-retaining dormant stem cells (LRCs), from 10 months of age and onwards (Fig. 4a). Second, the epidermis of another model of circadian arrhythmia, Per1 and Per2 double-mutant mice (Per1/2dKO), which lacks the negative limb of the molecular clock and is therefore locked in the clock<sup>bright</sup> state, showed the opposite effects; that is, enhanced bulge proliferation, reduced numbers of bulge LRCs and sustained expression of epidermal clock target genes29 (Fig. 4b and Supplementary Fig. 8). The hair follicles of Bmal1KO were less efficient in becoming active upon depilation (Supplementary Fig. 9a). Bmal1KO bulge cells were also less hyperproliferative than wild-type bulge cells in response to treatment with the phorbol ester 12-O-tetradecanoylphorbol-13-acetate (TPA), thus delaying the entry of the hair follicles into anagen (Supplementary Fig. 9b). Lastly, epidermal stem cells purified from Bmal1KO and Per1/2dKO mice were less and more clonogenic, respectively, in vitro than wild-type cells (Supplementary Fig. 10a, b). Of note, deletion of Bmal1 or Per1/2 did not affect the proportion of bulge stem cells in adulthood compared to their controls (Supplementary Fig. 11a, b).

### Loss of Bmal1 induces epidermal ageing

Reduced and enhanced proliferation, respectively, was also evident in the basal layer of the interfollicular epidermis of Bmal1KO and Per1/2dKO mice (Supplementary Fig. 12). Bmal1KO mice showed signs of inefficient epidermal self-renewal, with premature signs of ageing as early as 5 months of age, such as the accumulation of terminally differentiated cornified cells (Fig. 4c and Supplementary Fig. 9c). This was accompanied by increased expression of p16, which has been previously associated with increased epidermal ageing26, but not p19 or apoptosis (Supplementary Fig. 13). Because bulge stem cells do not contribute to epidermal maintenance in steady-state conditions29, we sought to understand the molecular mechanisms underlying the defects of the interfollicular epidermis of Bmal1KO mice. We performed

---

**Figure 2** | Circadian binding of Bmal to the promoters of genes involved in adhesion, cell cycle, TGF-β and Wnt pathways. ChIP from tail epidermis of wild-type mice collected every 3 h during 24 h (white and black bars represent day and night, respectively). Graph shows percentage of immunoprecipitated DNA over an input control from one representative experiment (n = 2).

---

**Figure 3** | Bmal1 modulates the response of bulge stem cells to activation and dormancy cues. a. Differential expression of genes in bulge cells of Bmal1WT and Bmal1KO mice. Fold-change values are shown as relative expression to Bmal1WT cells after normalization to Pum1 (n = 2). b. Western blot analysis for phospho-Smad2 (p-Smad2), Lef1 and Sox9 in tail epidermis of 9-month-old Bmal1WT and Bmal1KO mice. c. Primary mouse keratinocytes of Bmal1KO show enhanced responsiveness to TGF-β after 48 h of treatment (n = 3). d. Immunostaining for phospho-Smad2, TGFβR2, Lef1 and Sox9 in back skin of Bmal1WT and Bmal1KO mice (n ≥ 5). Scale bars 25 μm. e. Clonogenic assay of FACs-purified venus<sup>bright</sup> and venus<sup>dim</sup> bulge and interfollicular epidermis (IFE) keratinocytes from the back skin of P19 Per1–venus mice (7 × 10<sup>4</sup> bulge and 1 × 10<sup>5</sup> epidermal cells). Results in a and c are shown as mean ± s.e.m., *P < 0.05, **P < 0.01, ***P < 0.001 (two-tailed Student’s t-test).
microarray analysis of purified basal interfollicular epidermal progenitors (CD34<sup>6bright</sup>/CD45<sup>neg</sup>), from 10-month-old Bmal1KO mice and their control littermates (Supplementary Table 3). As expected, cells from Bmal1KO mice showed strong differential expression of most of the core circadian transcripts, including the microRNA miRNA-122 (ref. 32), some of which were validated independently by real-time quantitative PCR with reverse transcription (RT–qPCR) (Supplementary Fig. 14a and Supplementary Table 3). Interestingly, Gene Ontology (GO) analysis indicated that the cell cycle, energy and drug metabolism, calcium-sensing proteins, the epidermal barrier response and chromatin compaction were significantly affected upon deletion of Bmal1 (Supplementary Fig. 14b). Intriguingly, although Bmal1KO mice showed a hyperkeratotice phenotype, the viable epidermal layers expressed lower levels of terminal differentiation markers, including Flg, Lor, Sprr1, Lce genes and Tgf (Supplementary Fig. 14c and Supplementary Table 3). Interestingly, Bmal1KO cells also expressed lower amounts of the miRNA-23b/-27b/-24-1 cluster, which targets TGFβR2 and Smad proteins<sup>31</sup> (Supplementary Fig. 14d). The reduced levels of epidermal differentiation genes probably reflects the lower efficiency of activation of basal interfollicular epidermal cells in Bmal1KO mice, suggesting that the hyperkeratotic phenotype developed as a compensatory mechanism to ensure a certain degree of epidermal barrier protection.

**Loss of Bmal1 reduces skin tumorigenesis**

Because perturbation of the clock machinery affects the predisposition of certain tissues to carcinogenesis<sup>30</sup>, we next studied whether epidermal deletion of Bmal1 had any impact on the development of cutaneous squamous tumours. To this end, we crossed Bmal1KO mice with a transgenic line expressing oncogenic Sos, an activator of Ras, under the regulation of the Krt5 promoter (K5-SOS)<sup>35</sup>. In an EGFR mutant-heterozygous background, K5-SOS mice spontaneously developed squamous tumours, primarily in the tail, with 100% penetrance, as previously described<sup>35</sup>. Bmal1KO/K5-SOS mice developed significantly fewer neoplastic lesions at early-, mid- and late-stages of carcinoma development than control mice (Fig. 5a and Supplementary Fig. 15a). The skin lesions of Bmal1KO/K5-SOS mice were more differentiated—as determined by increased expression of involucrin and loricrin—contained large cornified islands and a higher number of apoptotic areas, as compared to control tumours (Supplementary Fig. 15b). Control mice had to be killed by two months of age, a time at which no Bmal1KO mice had developed the number, or size, of tumours. The growth of cutaneous squamous tumours has been shown to depend on Wnt activity in a population of CD34<sup>+</sup> tumour-initiating cells<sup>36</sup>. However, we could not detect any nuclear β-catenin, either in control or in Bmal1KO neoplastic lesions, suggesting that, in our model, tumour growth did not primarily depend on misregulated Wnt signalling (Supplementary Fig. 16a, b). We did observe a significant reduction in the percentage of CD34<sup>6bright</sup>/CD45<sup>neg</sup> tumour-initiating cells in Bmal1KO tumours with respect to wild-type tumours (Fig. 5b and Supplementary Fig. 17). Notably, whereas wild-type tumours expressed β6 integrin in basal and suprabasal cells, which has been previously associated with increased malignancy<sup>37</sup>, Bmal1KO tumours only expressed it in basal mouse were analysed). c, Histological analysis and immunostaining for filaggrin and loricrin (red) in 10–12-months-old Bmal1WT and Bmal1KO mice. Graph shows thickness quantification of the cornified layer (n = 13). Scale bars, 100 μm. Results are shown as mean ± s.e.m., *P < 0.05, **P < 0.01, ***P < 0.001 (two-tailed Student’s t-test). NS, not significant.

**Figure 4** | Clock perturbation in vivo results in changes in the number of dormant bulge stem cells, and premature epidermal ageing. a, b, Quantification of BrdU<sup>6</sup> LRCs and K67<sup>6</sup> cells in the bulge of 10-month-old Bmal1WT and Bmal1KO mice (a) and 10-week-old wild type (WT) and Per1/2dKO mice (b), showing opposite phenotypes (n = 4; 9 follicles per mouse were analysed).
immunostaining for this clock-controlled mechanism over stem cell regulation in humans stopped by adding 1.25 M glycine to a final concentration of 125 mM. Cells were in profound defects in tissue function39–41, and haematopoietic stem Conditional deletion of lishes a correct timing of stem cell activation and differentiation. Our results indicate that the molecular clock establishes stem cell renewal, shows a much more homogenous clock activity (albeit con- trary to the previous43. For the isolation of epidermal cells from back or tail skin, the skin was incubated in 1% formaldehyde. Cross-linking reactions were performed using the MAGnify Chromatin Immunoprecipitation System (Invitrogen).

Figure 5 | Loss of Bmal1 reduces the development of squamous tumours. a. Quantification of tumour weight of Bmal1WT/K5-SOS and Bmal1KO/K5-SOS littermates (n ≥ 5). b. FACS analysis and quantification of tε<sup>cre</sup>/CD4<sup>+</sup>/CD34<sup>+</sup> tumour-initiating cells (P36 mice). Numbers represent percentage of tε<sup>cre</sup>/CD34<sup>+</sup> cells from gated epithelial cells (n ≥ 3) c. Histological analysis and immunostaining for tε-intrins (arrows indicate suprabasal expression). Scale bars, 50 μm. Results in a and b are shown as mean ± s.e.m., *P < 0.05 (two-tailed Student’s t-test).

Discussion

Our results indicate that the molecular clock establishes stem cell states that are differentially predisposed to respond to activation and dormancy stimuli. The clock machinery controls the expression of essential epidermal stem cell regulators to anticipate the requirements of the tissue. In this sense, in the epidermal compartment where quiescent stem cells exist (that is, the bulge), the clock establishes a population of ‘ready-to-go’ cells that can rapidly and efficiently respond to activation stimuli, while simultaneously preventing all stem cells within the niche from becoming responsive. Future studies will be necessary to determine when, and how, this stem cell heterogeneity is established. On the other hand, the murine interfollicular epidermis, which primarily depends on continuously cycling basal progenitors rather than dormant stem cells<sup>60</sup> for its renewal, shows a much more homogenous clock activity (albeit containing approximately 5% of cells antiphasic with the majority). In this compartment the clock machinery might predominantly establish a correct timing of stem cell activation and differentiation. Conditional deletion of Bmal1 in liver, retina and pancreas results in profound defects in tissue function<sup>61</sup>, and haematopoietic stem cells show a Bmal1/Clock-dependent circadian release to the periphery<sup>2</sup>. Altogether, these findings indicate that the clock machinery may constitute a fine-tuning homeostatic mechanism in tissues in which dormant and active populations of stem cells coexist. Unbalancing the epidermal stem cell clock not only substantially affected long-term tissue homeostasis, but also the predisposition of the tissue to undergo neoplastic transformation. It is likely that perturbations of this clock-controlled mechanism over stem cell regulation in humans may have long-term consequences on tissue homeostasis, ageing and carcinogenesis.

METHODS SUMMARY

For the isolation of epidermal cells from back or tail skin, the skin was incubated in 0.25% trypsin for 2 h at 37°C, or overnight at 4°C, to separate the dermis from the epidermis. Back and tail keratinocytes were extracted as described previously<sup>46</sup>. For ChIP assays, cells in suspension were cross-linked for 10 min at room temperature (22–25°C) in 1% formaldehyde. Cross-linking reactions were stopped by adding 1.25 M glycine to a final concentration of 125 mM. Cells were centrifuged at 300g for 10 min at 4°C and washed in cold PBS. Cell lysis, sonification and ChIP assays were performed using the MAGnify Chromatin Immunoprecipitation System (Invitrogen).

For FACS analysis or sorting of bulge and epidermal stem cell populations, the skin was incubated for 30 min on ice with PE-conjugated anti-tε-integrin (CD49f clone NKi-GoH3, Serotec) and biotin-conjugated anti-CD34 (clone RAM34, BD Pharmingen), and/or followed by APC-conjugated streptavidin (BD Pharmingen) for 20 min. Dead cells were excluded by 7,6-diamidino-2-phenylindole (DAPI) incorporation. FACS analysis and sorting were performed using LSRII FACS Analyzers, FACSariaII, FACSdiva (BD Biosciences) and Flowjo software.

For microarray analysis, total RNA was isolated from FACS-sorted cells using Triozol extraction and RNeasy Micro Kit (Qiagen). Transcriptional profiling was performed using GeneChip Mouse Gene 1.0 ST Array (Affymetrix) and functional analysis was performed using DAVID Bioinformatics Resources 6.7.

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

Acknowledgements We thank the AICR (Association for International Cancer Research), the Spanish Ministry of Health (FIS) and AGAUR (Agència de Gestió d’Ajuts Universitaris i de Recerca; Government of Catalunya) for financial support. P.J. is the recipient of an AGAUR PhD Fellowship, and G.P. of a FIS fellowship. We thank D. McMahon (Vanderbilt University) for providing us with the Per1–GFP mice; E. Wagner (CNIO) for the K5–SOS mice; B. Kübler, the FACS and Genomics units of the IRB (Institut de Recerca Biomedica), the CRG (Center for Genomic Regulation) core facilities and the Animal Unit (Juan Martin Caballero) for technical support.

Author Contributions P.J. performed the experiments, and P.J. and S.A.B. analysed the results and wrote the manuscript. G.P. performed the analysis of K5–SOS mice, and A.M. and E.B. assisted P.J. in the initial FACS sorts. L.D.C. helped us with the initial ChIP experiments. K.O. provided the Per1–venus mice. J.R. and U.A. provided the Per1/Per2dKO mice.

Author Information Microarray data can be retrieved from the Gene Expression Omnibus under accession number GSE27079. Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of this article at www.nature.com/nature. Correspondence and requests for materials should be addressed to S.A.B. (salvador.aznar-benitah@crg.es).
METHODS

Animals. Bmal1WT and K14Cre mice purchased from The Jackson Laboratory were crossed each with either to obtain K14Cre/Bmal1WT and K14Cre/Bmal1KO (Bmal1KO), and with K5-SOS mice to generate K14Cre/Bmal1WT/K5-SOS (Bmal1WT/K5SOS) and K14Cre/Bmal1KO/K5-SOS (Bmal1KO/K5-SOS) littermate controls.25,26 Per1–GFP and Per1–venus mice have been described previously22,23. K14Cre/Bmal1wt/wt, Per1–GFP and Per1–venus mice were generated by crossing K14Cre/Bmal1WT and K14Cre/Bmal1KO mice, respectively. Per1 and Per2 double-mutant mice (Per1/2DKO) have been described previously, and were compared to age and sex-matched controls to avoid differences in hair cycle behaviour between males and females.27 Mice were housed in a AAALAC-I approved animal unit under 12 h light/12 h dark or 12 h dark/12 h dark cycles, and SPF conditions, and all procedures were approved by the CEEX (Ethical Committee for Animal Experimentation) of the Government of Catalonia. For experiments in constant darkness, light was turned off at Zeitgeber time (ZT)12 and animals were housed in 12 h dark/12 h dark conditions for 5 days. For 5-bromo-2-deoxyuridine (BrdU)-labelling experiments, 100 μg/kg BrdU ( Invitrogen) was injected intraperitoneally into the mice and chased for 10 weeks or up to 10 months, as indicated. To activate epidermal proliferation, back skin or tail skin of 10-week-old Bmal1WT and Bmal1KO mice was treated with three times with 20 nm TPA (Siga-Aldrich) during one week. For depilation experiments, dorsal skin of 15-month-old Bmal1WT and Bmal1KO mice in telogen phase was depilated with cold wax strips (Taky).

Primary mouse keratinocyte cultures. Primary keratinocytes from newborn mice or tail skin of adult mice were isolated as described previously28. Cells were plated in EMEM (Lonza) containing 4% chelated FBS, 1% penicillin/streptomycin, and 20 nM calcium for 24 h; medium was then changed to growth medium in 0.25% gelatin/PBS. Nuclei were stained with DAPI (1:5,000, Roche), and epidermal sheets were formed using the MAGnify Chromatin Immunoprecipitation System (Promega). Staining with haematocyanin or eosin was done according to a standard protocol. Samples were acquired with a Leica DMi 6000B or a Leica TCS SP5 confocal microscope.

Time-lapse microscopy and quantification. For time-lapse imaging, back skin of Per1–venus mice was fixed with 0.5% agarose in an imaging dish (ibidi) and over-layered with E-medium. Images were taken every 15 min for a period of 48 h using a Leica TCS SP5 confocal microscope equipped with a tempered chamber of 37 °C and 5% CO2. Mean fluorescence intensity of individual GFP-positive nuclei was quantified using ImageJ software.

FACS. Epidermal cells from back skin of Per1–venus, Bmal1WT and Bmal1KO mice, or tail skin of Bmal1WT/K5-SOS and Bmal1KO/K5-SOS mice, were isolated as described previously.25 Per1–venus mice were killed between ZT10–ZT12 or as specified in the figure legends. Cell suspensions were incubated for 30 min on ice with the following antibodies at the given dilutions: 1:100 for biotin- or APC-conjugated anti-CD34 (clone RAM34, BD Pharmingen), PE-conjugated anti-CD31 (clone MEC13.3, eBioScience), PE-conjugated anti-CD45 (clone 30F11, eBioScience), PE-conjugated anti-CD140a (clone APAS, BD Pharmingen) and FITC-conjugated anti-epcam (clone G8.8, Biologend); 1:200 for PE- or FITC-conjugated anti-β6-integrin (CD49f clone NKI-GoH3, Serotec); and 1:500 for APC-conjugated streptavidin (BD Pharmingen). DAPI staining was used to rule out that dead cells were present. FACS was performed using FACS AriaII and FACS Diva software (BD Bioscience). Sorted cells were collected in E-medium, either plated in culture or re-suspended in Trizol (Invitrogen), and subjected further to RNA isolation. FACS analysis was performed using LSRII FACS Analysers (BD Biosciences) and Flowjo software.

Arrays. Total RNA was isolated from FACS-sorted cells in Trizol by chloroform extraction, followed by the RNA clean-up protocol of RNeasy Micro or Mini Kit (Qiagen). Transcriptional profiling was performed using GeneChip Mouse Gene 1.0 ST Array (Affymetrix). Arrays of venusGFP and venusLum brite cells of P19 Per1–venus mice were performed as triplicates from a pool of n = 64 mice. Arrays of epidermal cells from 10-month-old Bmal1WT and Bmal1KO mice were performed as triplicates using three independent mice in each group. Functional analysis of microarray data was performed using DAVID Bioinformatics Resources 6.7.

RT–qPCR. Total RNA from cultured or FACS-sorted cells was purified as described above, or using either the RNeasy Micro and Mini Kit (Qiagen) or miRvana miRNA isolation Kit (Ambion). Equal amounts of RNA were reverse-transcribed using Superscript III (Invitrogen). RT–qPCR was performed with SYBR Green Master Mix (Roche) and gene-specific primers (as given in Supplementary Table 4)48 using a Light Cycler 480 Instrument (Roche). Relative levels of expression were determined by normalization to pumilio 1 (Pum1), using the ΔΔCt method.

CHIP. For ChIP assays from intact epidermis, mice were killed at ZT2, or for time-course analysis every 3 h during a period of 24 h, and tails were incubated in 0.25% trypsin for 2 h at 37 °C to separate the dermis from the epidermis. Tail keratinocytes were extracted as described.27 Cells in suspension or FACS-purified cells (2 × 10^6) were cross-linked for 10 min at room temperature in 1% formaldehyde. Cross-linking reactions were stopped by adding 1.25 M glycine to a final concentration of 125 mM. Cells were centrifuged at 300g for 10 min at 4 °C and washed in cold PBS. Cells lysis, sonification and ChIP assays were performed using the MAgnify Chromatin Immunoprecipitation System (Invitrogen). For each immunoprecipitation, 2 × 10^5–10^6 cells were incubated with 2 μl of anti-Bmal1 or anti-Clock antibody (provided by J. Ripperger, University of Fribourg), or 2 μl of rabbit IgG control antibody (Invitrogen). Luciferase activity was measured in a Centro LB 960 luminometer (Berthold) and expressed as above described using gene-specific primers (Supplementary Table 5).

Genotyping and western blots. Genomic DNA of primary mouse keratinocytes from Bmal1WT and Bmal1KO mice was isolated using a standard protocol. Multiplex PCR was performed as previously described to determine recombination efficiency.27 Protein extracts from newborn keratinocytes or from tail epidermis were analysed by SDS-PAGE and western blotting for Bmal1 (ref. 49), anti-phospho-Smad2 (1:1,000, #3801, Cell Signaling), anti-Lef1 (1:1,000, clone C12A5, Cell Signaling), anti-sox5 (1:400, H-90, Santa Cruz) and anti-tubulin (1:5,000, Sigma Aldrich).

Luciferase assay. Primary mouse keratinocytes were transiently transfected with either FOP-Flash or TOP-Flash and pCMV-Renilla plasmids using FuGene6 (Roche) according to the manufacturer’s instructions. The TOP-Flash construct reports Wnt activity by driving the expression of Luciferase under the regulation of TCF-Lef binding sites. FOP-Flash contains the same binding sites mutated so that they are no longer responsive to TCF-Lef. Cells were treated in the absence or presence of 1 μM BIO (Calbiochem) at 48 h post-transfection for a period of 24 h. Luciferase activity was measured in a Centro LB 960 luminometer (Berthold Technologies) using Dual-Luciferase Reporter Assay System (Promega).

Promoter analysis. Gene promoter analysis for potential Bmal1/Clock-binding sites (in general, from −5,000 to +1,000 bases from the transcriptional start site) were analysed using Genomatix Software.

Statistics. Results are presented as mean ± s.e.m. Statistical significance was determined by two-tailed Student’s t-test, one-way ANOVA, two-way ANOVA
with Bonferroni post-test, or Cosinor analysis. A $P$ value of $P \leq 0.05$ was considered to be statistically significant.


