

# RNAi screens in mice identify physiological regulators of oncogenic growth

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**Tissue growth is the multifaceted outcome of a cell's intrinsic capabilities and its interactions with the surrounding environment. Decoding these complexities is essential for understanding human development and tumorigenesis. Here we tackle this problem by carrying out the first genome-wide RNA-interference-mediated screens in mice. Focusing on skin development and oncogenic (Hras<sup>G12V</sup>-induced) hyperplasia, our screens uncover previously unknown as well as anticipated regulators of embryonic epidermal growth. Among the top oncogenic screen hits are *Mllt6* and the Wnt effector  $\beta$ -catenin, which maintain Hras<sup>G12V</sup>-dependent hyperproliferation. We also expose  $\beta$ -catenin as an unanticipated antagonist of normal epidermal growth, functioning through Wnt-independent intercellular adhesion. Finally, we validate functional significance in mouse and human cancers, thereby establishing the feasibility of *in vivo* mammalian genome-wide investigations to dissect tissue development and tumorigenesis. By documenting some oncogenic growth regulators, we pave the way for future investigations of other hits and raise promise for unearthing new targets for cancer therapies.**

Genome-wide cellular RNA interference (RNAi) screening has advanced the identification of genes involved in oncogenic growth control. To date, however, high-throughput screens in mammalian cells have been limited to cultures, in which even the best systems incompletely model physiological environments. We have overcome this impediment by devising methods to efficiently and selectively transduce murine epidermis through *in utero* lentiviral targeting of progenitors in embryonic day (E)9.5 embryos<sup>1</sup>. When coupled with short hairpin RNA (shRNA) expression, lentiviral transduction is stably propagated throughout skin epithelium, resulting in RNAi-mediated reductions in target transcript and protein levels. This enables rapid analysis of complex genetic pathways in mammals, something previously only possible in lower eukaryotes<sup>2–4</sup>.

The correlation between a tissue's growth and turnover rates and its susceptibility to cancer makes embryonic epidermis an attractive model for exploring how rapidly growing tissues balance proliferation and differentiation, and what prevents them from doing so in tumour progression. Given the efficacy of our system in single-gene studies, we have now expanded this scale by more than four orders of magnitude to conduct genome-wide RNAi screens. Our objectives were to first, demonstrate the feasibility of such screens in mammals; second, identify epidermal growth regulators in their native, physiological environment; third, uncover how epidermal growth control changes when it is propelled by a well-known oncogene; and fourth, demonstrate the implications of our findings for tumour progression in mice and humans.

## Epidermal growth is rapid and uniform

Following completion of gastrulation and continuing to birth, mouse surface ectoderm commences rapid growth to match embryo expansion (Fig. 1a). Beginning as a monolayer, E9.5 ectoderm differentiates into a stratified, multi-layered epidermis that by birth constitutes a barrier that retains fluids and excludes microbes. Mature epidermis maintains an inner progenitor layer, which fuels tissue homeostasis and wound repair.

To quantify epidermal growth, we randomly marked single cells at clonal density by infecting E9.5 *Rosa26*<sup>lox-stop-lox-(LSL)-yfp</sup> Cre-reporter embryos (*R26*<sup>yfp/+</sup>)<sup>5</sup> with an LV-Cre lentivirus<sup>1</sup>, and then monitored their expansion during development (Fig. 1b). By E18.5, single yellow

fluorescent protein (YFP)<sup>+</sup> cells at E10.5 had grown to clones constituting ~40 cells (Fig. 1b, c; ~5–6 divisions per cell). Variability in clone size ranged within 1–2 cell divisions, indicating strikingly uniform growth throughout the epidermis.

We next examined how growth is affected by oncogenic *Hras1*, which is found mutated in many cancers and is the primary target in skin carcinogenesis models<sup>6,7</sup>. *K14*-Cre<sup>8</sup>-driven, epidermal-specific expression of Hras<sup>G12V</sup> from its endogenous locus (*Hras*<sup>lox-wild-type-stop-lox-G12V</sup>)<sup>9</sup> resulted in mice whose skin displayed epidermal overgrowth as well as oncogenic Hras dose-dependency for one (*K14-cre; Hras*<sup>LSL-G12V/+</sup>; Hras<sup>oncoX1</sup>) or two (*K14-cre; Hras*<sup>LSL-G12V/LSL-G12V</sup>; Hras<sup>oncoX2</sup>) copies (Supplementary Fig. 1). Additional distinctions included the expansion of both progenitors (keratin 5<sup>+</sup>) and differentiating layers (spinous keratin 10<sup>+</sup>; granular filaggrin<sup>+</sup>) (Supplementary Figs 1–3).

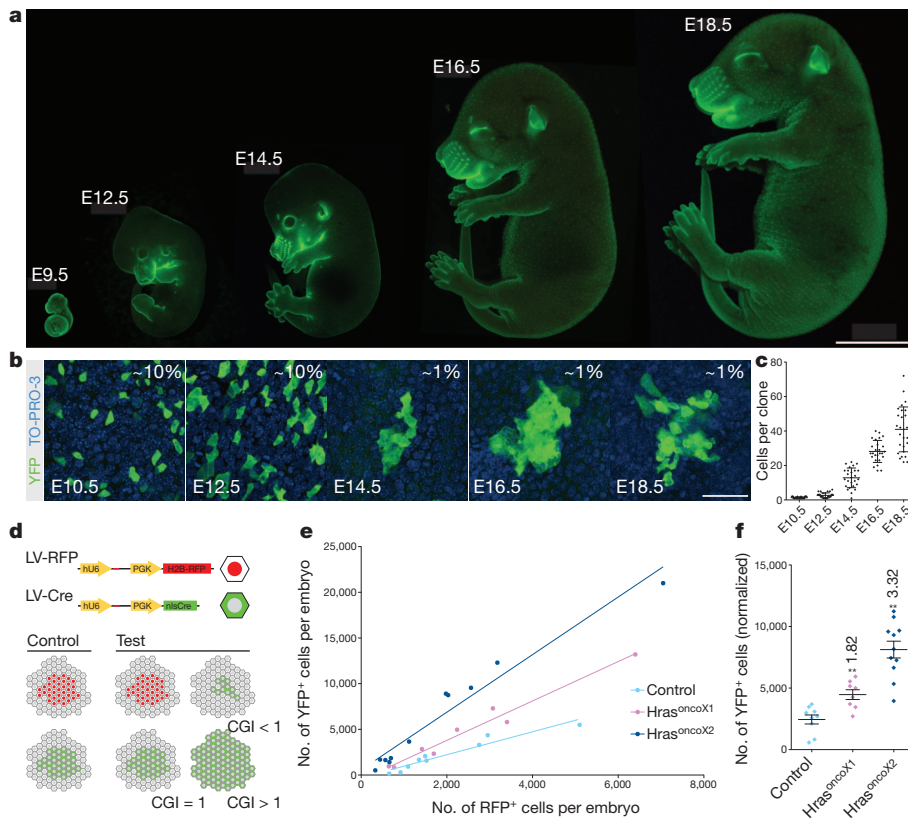
To quantify the impact of oncogenic Hras on epidermal growth, we used a cellular growth index (CGI) assay<sup>1</sup> (Fig. 1d). Cre-reporter (*R26*<sup>yfp/+</sup>) embryos, transduced with a LV-Cre and red fluorescent protein-expressing lentivirus (LV-RFP) mix showed similar relative numbers of YFP<sup>+</sup> to RFP<sup>+</sup> keratinocytes across several embryos, indicating that control YFP<sup>+</sup> and RFP<sup>+</sup> populations grew at comparable rates (Fig. 1e, f). Transduction of the same lentiviral mixture into test animals (where Cre-transduction induces Hras<sup>G12V</sup>) revealed consistently more YFP<sup>+</sup> (Hras<sup>G12V</sup>) than RFP<sup>+</sup> (control) cells (CGI = 1.82, Hras<sup>oncoX1</sup>; 3.32, Hras<sup>oncoX2</sup>; Fig. 1e, f). These findings demonstrate that Hras<sup>G12V</sup> confers a dose-dependent growth advantage to skin epidermis and that growth rates can be documented and quantified.

Hras<sup>G12V</sup> not only conferred a dose-dependent increase in proliferation, but also in suppression of apoptosis (Supplementary Fig. 4). In addition, consistent with established pro-inflammatory effects of Hras<sup>G12V</sup>, innate and adaptive immune cells infiltrated underlying dermis. Last, real-time PCR revealed no evidence for oncogenic induction of cellular senescence-associated cyclin-dependent kinase (CDK) inhibitors<sup>10</sup>.

## Establishing pooled screen parameters

On the basis of our CGI assay principle, we expected that, after epidermal transduction with a pool of shRNA-expressing lentivirus, any

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**Figure 1 | Embryonic epidermal tissue growth is rapid and responsive to oncogenic Hras.**

**a**, Mouse embryogenesis, highlighted by propidium iodide (E9.5) or K14-actin-GFP (E12.5–18.5). **b**,  $R26^{yfp/+}$  Cre-reporter embryos infected at E9.5 with LV-Cre and analysed at days shown. Transduced cells are YFP<sup>+</sup>. Transduction levels (% YFP<sup>+</sup> cells) depend upon viral titre. **c**, Cell numbers in transduced YFP<sup>+</sup> clones at ages shown. **d**, Schematic of CGI assay. E9.5  $R26^{yfp/+}$  Cre-reporter (control) or  $gene^{lox/lox} R26^{yfp/+}$  (test) embryos are infected with a LV-Cre and LV-RFP mix. At E18.5, numbers of RFP<sup>+</sup>:YFP<sup>+</sup> cells in control and test animals are compared, and phenotypes scored as neutral (CGI = 1), growth advantaged (CGI > 1) or disadvantaged (CGI < 1). **e**, Numbers of RFP<sup>+</sup> and YFP<sup>+</sup> cells at E18.5 in control,  $Hras^{oncoX1}$  and  $Hras^{oncoX2}$  embryos. Upper shift is consistent with growth advantage. **f**, RFP<sup>+</sup> cell numbers normalized to YFP<sup>+</sup> cells in control,  $Hras^{oncoX1}$  and  $Hras^{oncoX2}$  animals. CGI assay suggests a 1.8-fold overgrowth ( $P = 0.002$ ) in  $Hras^{oncoX1}$  and 3.3-fold overgrowth ( $P < 0.0001$ ) in  $Hras^{oncoX2}$  epidermis. Error bars indicate  $\pm$  s.d. (**c**) and  $\pm$  s.e.m. (**f**). \*\* ( $P \leq 0.01$ ) in **f** indicates statistical significance of comparison to control. For CGI assay (**e**, **f**), data points are individual embryos: control ( $n = 9$ ),  $Hras^{oncoX1}$  ( $n = 8$ ) and  $Hras^{oncoX2}$  ( $n = 11$ ). Scale bars, 5 mm (**a**) and 50  $\mu$ m (**b**).

shRNA that targets an essential mediator of growth will be reduced or lost during development, whereas shRNAs targeting negative growth regulators will become overrepresented. By comparing relative shRNA abundance in the initial pool and at E18.5, we expected to identify shRNAs and their targets that confer either growth advantage or disadvantage.

The success of the approach depended upon our ability to: (1) modify growth at a low multiplicity of infection ( $MOI \leq 1$ ); (2) measure individual shRNA abundance in the pool; (3) transduce embryonic epidermis at a  $MOI \leq 1$ ; and (4) achieve complete screen coverage, in which every shRNA in the pool is tested. We set up a series of controls to ensure that these parameters were met. Underscoring the feasibility of pooled-formats for *in vivo* RNAi screens, we demonstrated that targeting of (1) anaphase promoting complex component *Anapc5* during normal growth, and (2) *Hras1* during oncogenic hyperplasia, reduced average clone sizes, even with transductions where most cells harboured only a single shRNA (Supplementary Fig. 5).

To quantify individual shRNA representation in a complex pool, we used the Illumina-based count-by-sequencing principle (Supplementary Fig. 6). We designed oligonucleotides to amplify the target sequence of each shRNA, and optimized pre-amplification and clean-up pipelines to yield a product to apply directly to the sequencing cell. We tested our protocol against a defined template generated by combining genomic DNAs from independently transduced cell lines, so that individual genome-integrated shRNAs were present in amounts corresponding from a single cell (6 pg) up to 2,048 cells (12.3 ng).

We amplified and sequenced this standard set, and showed that reactions were: (1) quantitative, with increased sequencing reads corresponding to shRNA abundance in the pool; (2) sensitive, detecting all three single-copy shRNAs; and (3) highly reproducible (Supplementary Fig. 6). Independent counts of the standard set showed identical sequencing bias for a given shRNA, and thus became neutralized in relative comparisons of absolute counts, especially with  $\geq 32$  copies of the shRNA. Indeed, a >30-fold screen coverage proved sufficient to sample all shRNAs in our pool (see below). At this level, growth-neutral

shRNAs were >1,000-fold represented in the E18.5 sequencing quantification reaction, because each E9.5 epidermal cell generates  $\sim 40$  cells by E18.5.

We next determined that at an infection level of 13–27%, most transduced epidermal keratinocytes carried a single lentivirus ( $MOI \leq 1$ ) (Supplementary Fig. 7). To ensure that at least 30 individual cells were infected with each shRNA at E9.5, a pool of  $\sim 78,000$  shRNAs required  $\sim 10^6$  cells to be targeted. We used high-resolution imaging of TO-PRO3-labelled embryos and established that at E9.5, surface ectoderm contained  $\geq 120,000$  cells per embryo. Together, this suggested that transducing  $\geq 90$  embryos would achieve the requisite coverage (Supplementary Fig. 8).

### Screens identify known growth regulators

We pooled the genome-wide collection of murine shRNA lentiviruses in roughly equal concentrations<sup>11</sup>, and profiled the starting composition of the pool ( $t = 0$ ) in transduced primary mouse keratinocytes (Fig. 2a). Physiological screens were performed in control and  $Hras^{oncoX2}$  embryos transduced at E9.5 *in utero* (Fig. 2b). Epidermal cells were collected after 24 hours (initial pool) and 9 days of development, and integrated lentiviral hairpins from genomic DNAs were sequenced and quantified (Fig. 2c).

Our pre-amplification and sequencing reactions did not bias shRNA quantification, and Illumina sequencing reads were of high quality. They mapped to the shRNA library with predictable efficiency and indicated complete coverage of the pool (Supplementary Figs 9, 10). Significantly altered shRNAs were identified and ranked on the basis of two independent methods. To ensure reproducibility, we used DESeq statistical package<sup>12</sup>, which accounted for biological variability among our replicates (sets of 30 embryos per condition) and is the best method to identify candidates per se (Fig. 2d–k). However, informative yet variable shRNAs can be excluded by the high stringency of DESeq, which reduces the ability to control for off-target effects by requiring that multiple shRNAs show consistent behaviour. We therefore also analysed pooled data sets (90 embryos per condition;

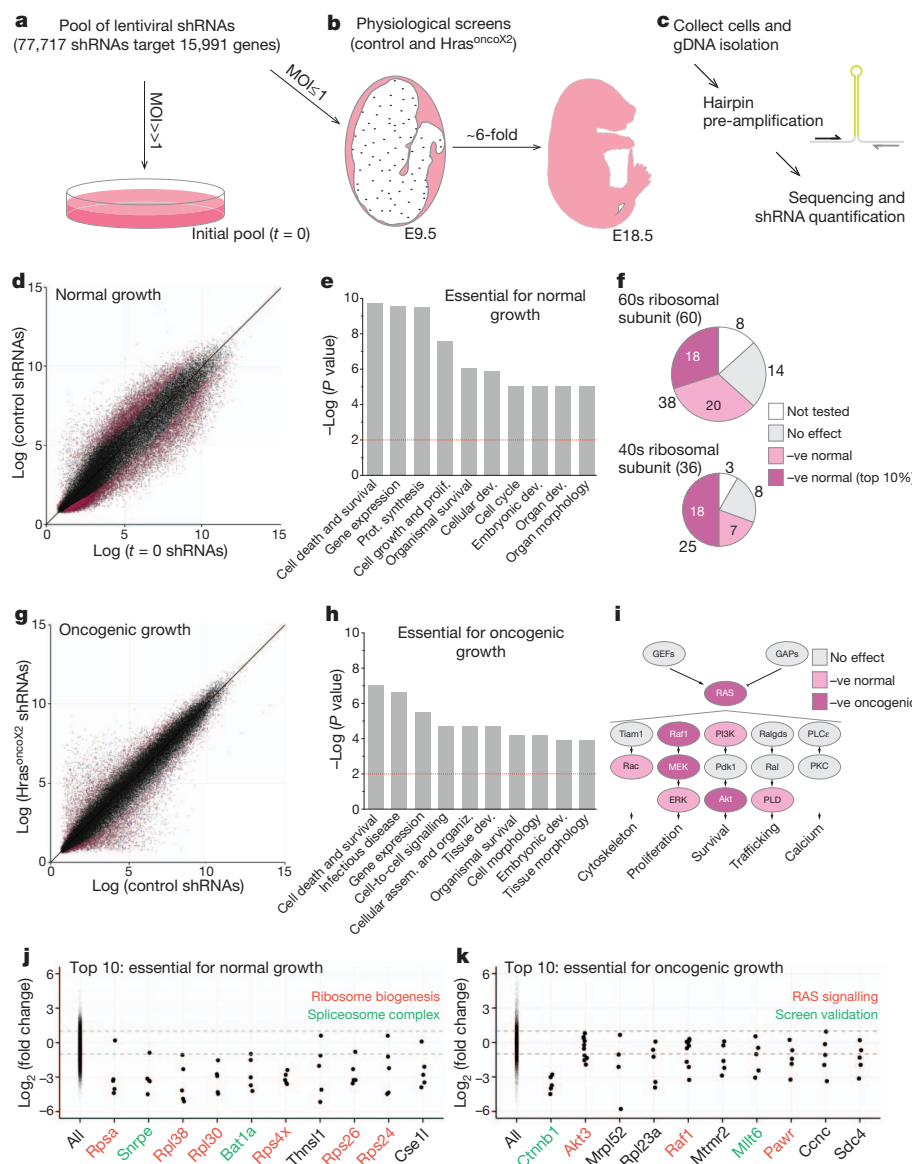
Supplementary Fig. 11) using Fisher's exact test, which reduces variability by averaging individual shRNA abundance. By maximizing screen coverage, this method produces a more inclusive list of significantly altered shRNAs, and hence was preferred for ranking candidates identified by DESeq (Fig. 2f, j, k). Importantly, the overlap between these approaches was extensive (Supplementary Fig. 11), underscoring the robustness of our data.

We identified ~1,800 genes as essential for normal growth (Fig. 2d, e and Supplementary Tables 1, 2) and significantly enriched for function in protein synthesis ( $P = 3.1 \times 10^{-10}$ ) and gene expression ( $P = 2.6 \times 10^{-10}$ ). Genes encoding 60S and 40S ribosomal proteins were also highly represented and among the top 10% of all hits for normal growth, underscoring our screen's power to identify regulators of normal growth/viability (Fig. 2f and Supplementary Fig. 12). Indeed, our top 10 candidates for regulators of normal growth featured six ribosomal genes and two genes essential for messenger RNA splicing (Fig. 2j).

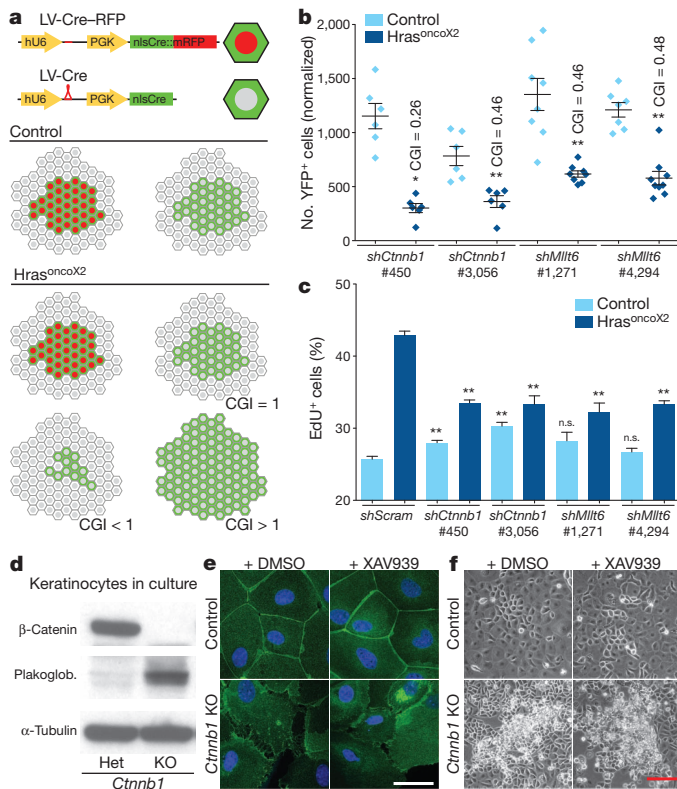
In our screen for specific regulators of Hras<sup>G12V</sup>-dependent oncogenic growth, ~160 genes surfaced as candidates (Fig. 2g and Supplementary Tables 3, 4). They diverged in identity and function from those implicated in normal growth regulation, as most housekeeping growth regulators were eliminated by pair-wise comparison of control and Hras<sup>oncoX2</sup> shRNA abundance (Fig. 2h). The oncogene-specific

regulators included Hras1, and downstream Ras pathway members Raf1, and Mek and Akt proteins (Fig. 2i). Equally notable was the absence of upstream oncogenic Ras regulators, for example, guanine nucleotide exchange factors (GEFs) and GTPase-activating proteins (GAPs), which are not expected to arise in a screen for hairpins suppressing Hras<sup>G12V</sup>-induced growth. Our top 10 hits for regulators of oncogenic growth included well-established Ras pathway members *Akt3* and *Raf1*, as well as the Ras-regulated *Pawr*<sup>13</sup> (Fig. 2k). Cyclin C (*Ccnc*), a putative regulator of haematopoietic stem cell quiescence<sup>14</sup>, was also on this list, as was *Mllt6*, encoding a myeloid/lymphoid, or mixed-lineage, leukaemia (MLL) translocation partner and a component of an epigenetic modifier complex<sup>15</sup>.

The very top candidate for preferential regulation of oncogenic growth was *Cttnb1*, encoding the Wnt signalling and intercellular adhesion effector  $\beta$ -catenin. At first glance,  $\beta$ -catenin seemed obvious, as its over-activation has been implicated in a variety of cancers, including those of skin<sup>16-18</sup>. However,  $\beta$ -catenin is also thought to be essential for stem cells<sup>19</sup>. Hence, it was surprising that its hairpins surfaced in a screen for selective inhibitors of oncogenic and not normal growth. Even more paradoxical was that in our parallel genome-wide screen, *Cttnb1* was the top candidate for negative regulation of normal epidermal growth (Supplementary Tables 1, 2).



**Figure 2 | Genome-wide RNAi screens for physiological regulators of normal and oncogenic growth identify expected and surprising regulators.** **a–c**, Schematic of the RNAi screens based on relative enrichment/depletion of individual shRNAs over time. **a**, shRNAs against 15,991 mouse genes are combined into a lentiviral pool whose composition is determined from the 'initial pool' ( $t = 0$ ) experiment, in which transduced cells are analysed 24 h after infection. **b**, Genes that regulate normal and oncogenic growth are identified in two screens, in which E9.5 control or K14-Cre<sup>+</sup>; Hras<sup>oncoX2</sup> embryos are infected *in utero*, allowed to develop for 9 days, and processed. **c**, After epidermal tissues are collected and used in genomic DNA (gDNA) isolation, individual shRNAs are pre-amplified and quantified by sequencing unique hairpin regions. **d–k**, Significantly enriched/depleted shRNAs identified using DESeq analysis. **d**, Dot plot of relative abundance of 77,717 shRNAs at  $t = 0$  and in E18.5 epidermis. **e**, Putative normal growth regulators are significantly enriched ( $P \geq 0.01$ ) for gene function categories promoting cell viability and development. **f**, Normal growth regulators encode for ribosomal 60S and 40S subunit components ( $P = 1.12 \times 10^{-9}$  and  $P = 3.26 \times 10^{-7}$ , respectively), with many in the top 10% of all hits (maroon). **g**, Relative shRNA abundance in control and Hras<sup>G12V</sup> animals reveal oncogenic growth-altering shRNAs. **h**, Putative oncogenic growth regulators are enriched ( $P \geq 0.01$ ) in gene categories that support cell growth. **i**, Downstream effectors of Ras signalling score as essential for growth (pink), with many exhibiting an oncogene-specific requirement (maroon). **j**, **k**, shRNAs for the top ten essential regulators of normal (**d**) and oncogenic (**g**) growth are markedly depleted relative to the genome-wide pool (all).



**Figure 3 | Suppressing  $\beta$ -catenin and *Mllt6* selectively affects *Hras*<sup>G12V</sup>-dependent epidermal hyperplasia.** **a**, Modified CGI assay measures effect of shRNA-mediated gene knockdown in animals with Cre-activated transgene expression. Transduction with LV-Cre-RFP co-expressing scrambled (scram) shRNA, and LV-Cre co-expressing candidate-targeting shRNA, leads to generation of YFP<sup>+</sup> RFP<sup>+</sup> scrambled and YFP<sup>+</sup> knockdown clones in control or *Hras*<sup>G12V</sup> animals. Numbers of YFP<sup>+</sup> cells (normalized to YFP<sup>+</sup> RFP<sup>+</sup>) in control and *Hras*<sup>oncoX2</sup> animals reflect lentiviral mix composition after normal and oncogenic growth, respectively. **b**, Fewer YFP<sup>+</sup> cells are found in oncogenic animals upon knockdown of *Ctnnb1* and *Mllt6* with independent shRNAs. **c**, Reduced EdU incorporation following *Ctnnb1* knockdown in *Hras*<sup>G12V</sup> animals contrasts with increased proliferation in control epidermis. *Mllt6* depletion also reduces EdU labelling in oncogenic growth. **d**, Immunoblot of control (*Ctnnb1* heterozygous (Het)) and *Ctnnb1* knockout (KO) keratinocyte lysates shows upregulation of plakoglobin. **e**, Establishment of cell adhesion 48 h after Ca<sup>2+</sup> shift is unaffected in keratinocytes treated with a Wnt-signalling inhibitor (XAV939) but impaired in *Ctnnb1* knockout cells. E-cadherin (green) marks adherens junctions and DAPI (4',6-diamidino-2-phenylindole; blue) labels the nuclei. **f**, Unlike control cells or cells treated with a Wnt-inhibitor, *Ctnnb1* knockout keratinocytes form overgrown foci upon reaching confluence. Error bars (**b**, **c**) indicate  $\pm$  s.e.m. Data points (**b**, **c**) represent individual embryos with  $n = 6$  (*shCtnnb1* and *shScram* in control),  $n = 7$  (*shMllt6* no. 4,294 in control),  $n = 8$  (*shMllt6* no. 1,271),  $n = 9$  (*shMllt6* no. 4,294 in *Hras*<sup>G12V</sup>), or  $n = 10$  (*shScram* in *Hras*<sup>G12V</sup>), each scored through immunofluorescence analysis of ten 425.1  $\mu\text{m}^2$  images. NS, not significant ( $P > 0.05$ ); \* $P \leq 0.05$  and \*\* $P \leq 0.01$  indicate statistical significance. Scale bars, 50  $\mu\text{m}$  (**e**), 10  $\mu\text{m}$  (**f**).

### Validating oncogene-specific regulators

We chose *Ctnnb1* and *Mllt6* for further study. For both *Ctnnb1* and *Mllt6*, a direct correlation was observed between transcript knockdown *in vitro* and severity of growth defects *in vivo* (Supplementary Fig. 13), strongly arguing against off-target effects. We validated our candidates as oncogenic growth regulators with a modified *in vivo* CGI assay involving two lentiviral vectors (Fig. 3a). In one, Cre-recombinase fused to monomeric RFP (LV-Cre-mRFP) contained a scrambled shRNA control. In the other vector, untagged Cre was used; this vector encoded the test shRNA against the candidate. Transduction of E9.5 control or *Hras*<sup>oncoX2</sup> Cre-reporter embryos marked two separate populations: RFP<sup>+</sup> YFP<sup>+</sup> cells represented the

baseline rate of normal (in control) or oncogenic growth (in *Hras*<sup>oncoX2</sup>); YFP<sup>+</sup> cells represented the rate of growth that occurs when the target transcript is depleted.

The ratios of YFP<sup>+</sup> cells normalized to YFP<sup>+</sup> RFP<sup>+</sup> cells in *Hras*<sup>oncoX2</sup> and control animals revealed that two independent *Ctnnb1* shRNAs displayed reductions of  $\sim 2$ – $4$ -fold in oncogenic relative to normal growth. Similar  $\sim 2$ -fold reductions in YFP<sup>+</sup> cells were observed in the *Hras*<sup>oncoX2</sup> background when *Mllt6* transcripts were diminished (Fig. 3b). The physiological effects of *Ctnnb1* or *Mllt6* knockdown were profound: the neonatal oncogenic phenotype was significantly ameliorated, and epidermal proliferation in *Hras*<sup>oncoX2</sup> embryos was markedly and reproducibly suppressed. By contrast, no significant effects were seen on apoptosis (Fig. 3c and Supplementary Fig. 13).

Equally interesting to the selective effects of *Ctnnb1* knockdown on suppressing *Hras*<sup>G12V</sup>-dependent oncogenic growth were its positive effects on normal growth. These differences seemed to be physiologically relevant, as they were reflected at the level of EdU-incorporation and thickness of epidermal tissue (Fig. 3c and Supplementary Fig. 14). Although hitherto overlooked, the proliferative effects of *Ctnnb1* hairpins on normal epidermis are recapitulated upon conditional targeting of  $\beta$ -catenin<sup>20,21</sup>.

$\beta$ -Catenin is both an adherens junction component and a nuclear cofactor for Wnt regulators in the LEF/TCF family and other DNA-binding proteins<sup>22</sup>. However in contrast to its nuclear functions,  $\beta$ -catenin's role in adhesion has been assumed to be redundant with plakoglobin<sup>23</sup>. Given that intercellular defects can promote proliferation, we revisited this issue using a sensitive *in vitro* adhesion assay (Fig. 3d–f and Supplementary Fig. 15)<sup>24</sup>. Despite plakoglobin upregulation, *Ctnnb1*-null keratinocytes inefficiently formed cell–cell adhesions upon calcium induction. Moreover, the Wnt inhibitor XAV939 (ref. 25) failed to phenocopy these defects. Finally, consistent with the view that loss of  $\beta$ -catenin compromises contact inhibition and leads to cellular overgrowth, *Ctnnb1*-null cells were hyperproliferative and formed overgrown foci upon reaching confluence.

### Oncogenic growth and Wnt signalling

Although intercellular adhesion is often viewed as tumour suppressive, Wnt signalling is often associated with oncogenic growth. To test whether this might contribute to the negative effects of *Ctnnb1* knockdown on *Hras*<sup>G12V</sup> skins, we transduced embryos with both a Wnt-reporter and LV-Cre (Fig. 4a). In E18.5 control animals, reporter expression was predictably restricted to developing hair follicles<sup>26</sup> and largely abolished with concomitant *Ctnnb1* knockdown.

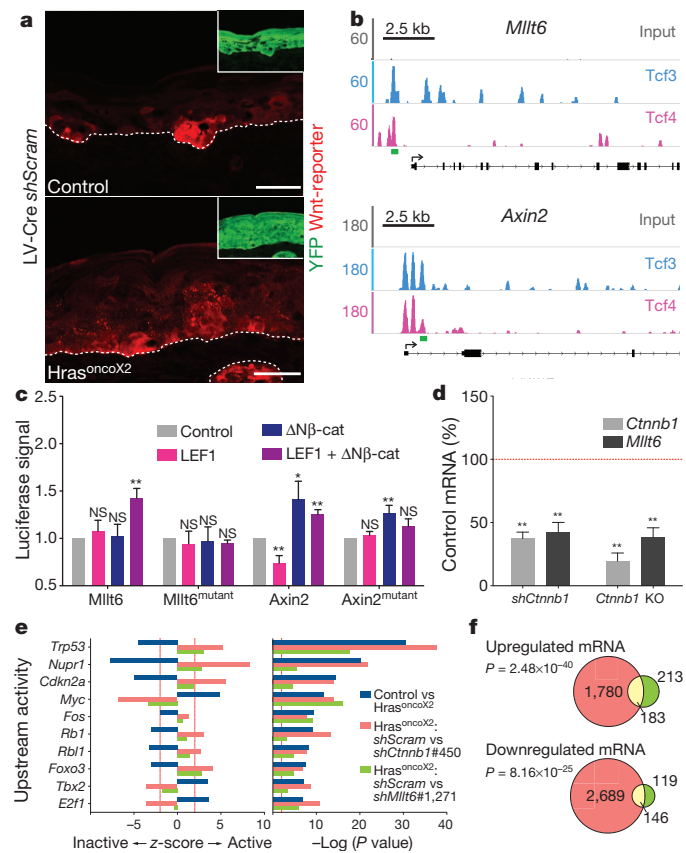
Intriguingly, Wnt-reporter expression was expanded throughout transduced *Hras*<sup>oncoX2</sup> interfollicular epidermis. In addition, *Hras*<sup>G12V</sup>-expressing epidermis displayed ectopic nuclear  $\beta$ -catenin and  $>6$ -fold upregulation of *Axin2* transcripts (Fig. 4a and Supplementary Fig. 16a–c). Conversely, bone morphogenetic protein signalling, which is antagonistic to Wnt signalling in skin<sup>27</sup>, was downregulated in *Hras*<sup>G12V</sup> epidermis, and was not rescued by  $\beta$ -catenin depletion, suggesting its independence of Wnt in this oncogenic context (Supplementary Fig. 18).

*Mllt6* expression paralleled  $\beta$ -catenin and Wnt-reporter activity, both in normal hair follicles and in evaginating structures of *Hras*<sup>oncoX2</sup> skin (Supplementary Fig. 16b). Chromatin immunoprecipitation followed by next-generation sequencing (ChIP-seq) analysis showed that Tcf3 and Tcf4 bound to a conserved *Leif/Tcf* motif upstream of *Mllt6* (Fig. 4b). A 299-base-pair (bp) segment (green line) encompassing this site drove LEF1/ $\beta$ -catenin-dependent luciferase reporter activity in a manner comparable to the 331-bp Tcf3/4 binding site of *Axin2*, an established Wnt-target gene (Fig. 4c and Supplementary Fig. 16d). In agreement, depletion or loss of  $\beta$ -catenin in embryonic epidermis *in vivo* reduced *Mllt6* transcript levels (Fig. 4d).

### $\beta$ -catenin and *Mllt6* in epidermal tumours

Although our screens were conducted on embryonic mouse skin, our findings showed relevance to cancer. RNA-seq analysis revealed that

*Cttnb1* or *Mllt6* depletion in oncogenic Hras epidermis affected a shared set of transcripts ( $P = 2.48 \times 10^{-40}$  and  $P = 8.16 \times 10^{-25}$ ) that globally suppressed pathways promoting tumorigenesis (for example, *Myc*, *E2f1*) and enhanced those restricting growth (for example, *Trp53*, *Cdkn2a*; Fig. 4e, f). Moreover, in human squamous cell carcinomas (SCC),  $\beta$ -catenin and MLLT6 were often upregulated and nuclear (notably in the basal layer, where cancer stem cells reside<sup>28</sup>). Our analysis of 75 different human skin SCCs showed that most tumours expressed both proteins, with significant correlation in their expression (Supplementary Fig. 18). It remains to be seen whether their co-expression in tumours and a shared effect on transcriptional profile during oncogenic growth reflects a functional interaction between our candidates, or is a result of their independent



effect/importance on the cellular machinery at the heart of oncogenic growth.

We next tested whether  $\beta$ -catenin and *Mllt6* are physiologically relevant to Hras<sup>G12V</sup>-dependent tumour initiation and maintenance. Whereas clonal LV-Cre-mediated activation of Hras<sup>G12V</sup> expression in mice resulted in squamous papilloma formation as early as 3 weeks of age, concomitant constitutive expression of *Cttnb1* or *Mllt6* shRNAs delayed tumour initiation (Fig. 5a). Moreover, growth of orthotopically transplanted SCC cells was significantly reduced following candidate depletion (Fig. 5b and Supplementary Fig. 19).

We extended this physiological relevance by performing xenografts of human SCC cells transduced with lentivirus harbouring scrambled or shRNAs targeting human *CTNNB1* or human *MLLT6*. Tumour initiation was significantly delayed, with *MLLT6* showing stronger effect than *CTNNB1* (Fig. 5c and Supplementary Fig. 19).

Finally, to assess whether *Cttnb1* and *Mllt6* are required for tumour maintenance, we engineered an LV-Cre vector that allows for doxycycline-regulated shRNA expression, thereby enabling induced depletion of *Mllt6* and *Cttnb1* following tumour formation in adult animals (Fig. 5d and Supplementary Fig. 20). Both had negative effects on tumour maintenance, with some tumours showing partial regression. Thus, the tumour-suppressive effects of *Cttnb1* and *Mllt6* shRNAs as first revealed in embryogenesis appeared to be functionally relevant to adult tumorigenesis.

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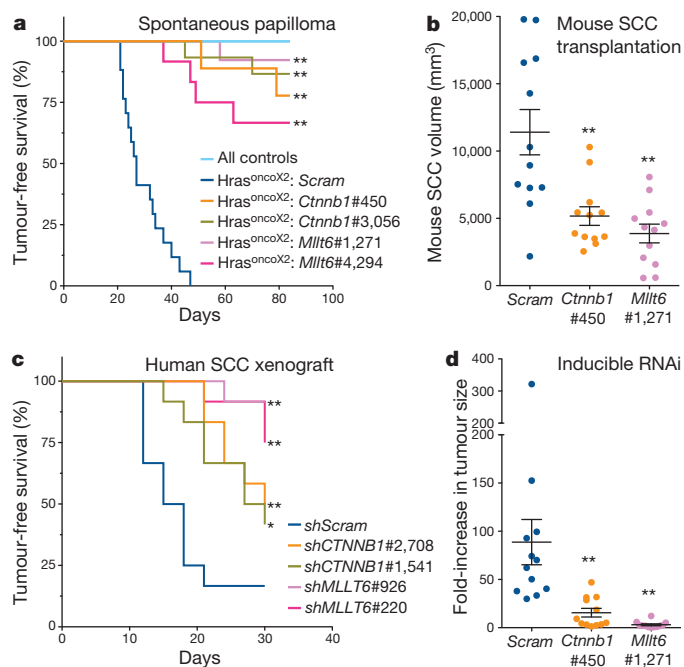
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## Discussion

The urgent need to understand cancer has fuelled human cancer genome sequencing and *in vitro* RNAi-based screening efforts to



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identify genes that preferentially affect cancer cells but not their normal counterparts. Although promising in concept, assay conditions and cell-line histories can profoundly affect genes identified in these screens<sup>29,30</sup>. Although xenogeneic transplantations of transduced human cells offer improvements<sup>31</sup>, they often incompletely simulate carcinoma ontogeny, which depends upon complex interactions with local and systemic environments. By targeting cells in their normal physiological context, we correct these deficits and abrogate many caveats, including epigenetic, genetic and stress-induced alterations in gene expression, all of which introduce heterogeneity and increase coverage requirements when cells are grown on plastic.

Our study accentuates a particular importance of  $\beta$ -catenin in promoting oncogenic effects, as it surfaced at the top of >15,000 genes in our screen. Moreover, increased Hras-mitogen-activated protein kinase signalling drove  $\beta$ -catenin's effects from negative to positive, as normal epidermal growth was actually impeded by  $\beta$ -catenin. In this regard, it is interesting that leukaemias also seem to be more sensitive to activated  $\beta$ -catenin than their normal counterparts<sup>32</sup>. Our findings further suggest that  $\beta$ -catenin's ability to balance tissue growth is exerted through its antagonistic functions in intercellular adhesion and transcriptional activation.

A myriad of new candidates from our screen await further investigation. Among them are chromatin modifiers, which have been increasingly implicated in human cancers<sup>33</sup>. In this regard, our validation of Mllt6 is intriguing, as MLL proteins are known to associate with DOT1L H3K79-methyltransferase complexes<sup>34</sup>. Given Mllt6's selective effects on oncogenic growth, it is tempting to speculate that this protein might function by guiding its histone modifier complex to a key cancer target gene(s). Although detailed understanding of this and other candidates awaits experimentation, our methodology paves the way for future studies aimed at uncovering mechanisms of SCC progression, with the hope of identifying targets that selectively compromise growth of one of the world's most prevalent and life-threatening cancers.

## METHODS SUMMARY

Animals were on a C57BL/6 background. Lentiviral production and ultrasound-guided injection into E9.5 amniotic space are as described<sup>1,35</sup>. Transduced embryos were developed to E18.5, after which epidermal suspensions were prepared for gDNA isolation. gDNAs from sets of 30 transduced embryos were combined and used as template for a 21-cycle pre-amplification PCR. For identification and quantification of shRNAs, clean pre-amplification product was sequenced using Illumina HiSeq2000, and the sequencing output was aligned to the TRC 2.x library with Burrows-Wheeler Aligner (BWA) with a maximum edit distance of three. Bioinformatics analyses of RNA-seq data and candidates identified by our screens were performed using IPA software (Ingenuity Systems). Figures were prepared using Adobe Photoshop and Illustrator CS5. Graphing and statistical analyses were performed using Prism 5 (GraphPad Software). Descriptions of antibodies and mouse strains are provided in Methods.

**Full Methods** and any associated references are available in the online version of the paper.

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**Author Contributions** S.B., P.J. and E.F. designed the experiments. S.B. and P.J. made shRNA pools and lentivirus, and performed the screens. Illumina sequence analysis was done by E.H. and S.B. RNAseq and IPA analyses were performed by S.B., B.E.K. and P.J. Imaging was done by S.B. and N.O., and image analysis by S.B. and E.H. CHIP-seq data was generated by W.-H.L., and P.J. performed luciferase assays. S.B. and E.F. wrote the paper. All authors provided intellectual input, vetted and approved the final manuscript.

**Author Information** Raw RNAseq data can be accessed at Gene Expression Omnibus under accession number GSE48480 and permissions information is available at [www.nature.com/reprints](http://www.nature.com/reprints). The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to E.F. ([fuchslb@rockefeller.edu](mailto:fuchslb@rockefeller.edu)).

## METHODS

**Lentivirus production and *in vivo* and *in vitro* transductions.** Large-scale production and concentration of lentivirus were performed as previously described<sup>1</sup>. Male and female animals was used in equal numbers, and all mice were on the C57BL/6 background, including *Gt(Rosa)26Sor<sup>tm1(eYFP)Cosl+</sup>* (Jackson Laboratories, donated by A. McMahon), *FR-Hras<sup>G12V</sup>* (9) and *Tg(K14-cre)1Efl<sup>u</sup>*. Mice were housed and cared for in an AAALAC-accredited facility, and all animal experiments were conducted in accordance with IACUC-approved protocols. Randomization and blinding were not used in this study. Detailed description of the *in vivo* lentiviral transductions can be found elsewhere<sup>1,35</sup>. For lentiviral infections in culture, cells were plated in 12-well dishes at 70,000 cells per well and incubated with lentivirus in the presence of polybrene (100 mg ml<sup>-1</sup>) overnight. After 2 days, infected cells were sorted on the basis of RFP expression (mouse and human SCC cells) or positively selected with puromycin (1 mg ml<sup>-1</sup>) for 4 days and processed for mRNA analysis. **mRNA quantifications.** Total RNAs were isolated from FACS-sorted cells from E18.5 epidermis or from flash-frozen, pulverized kidney, using the Absolutely RNA Microprep kit (Stratagene). Complementary DNAs were generated from 1 µg of total RNA using the SuperScript Vilo cDNA synthesis kit (Life Technologies). Real-time PCR was performed using the 7900HT Fast Real-Time PCR System (Applied Biosystems) and gene-specific and *Ppib* control primers. Real-time experiments were done on cells isolated from three transduced animals, or three independently transduced cell culture plates, and all reactions were performed in triplicate and in two separate runs. BRE-ZsGreen activity was measured using real-time PCR with ZsGreen-specific primers on cDNA from transduced epidermal cells as previously described<sup>4</sup>.

**Immunostaining and histological analyses.** The following primary antibodies were used: chicken anti-GFP (1:2,000; Abcam); mouse anti-β-catenin (15B8, 1:1,000; Sigma); guinea pig anti-K5 (1:500; E. Fuchs); rat anti-CD34 (RAM34, 1:100; eBioscience), anti-Ecad (ECCD-1, 1:200; M. Takeichi) and anti-nidogen (ELM1, 1:2,000; Santa Cruz); rabbit anti-caspase 3 (AF835, 1:1,000; R&D), anti-RFP (PM005, 1:2,000; MBL), anti-K10 (PRB-159P, 1:1,000; Covance), anti-filaggrin (PRB-417P, 1:2,000; Covance), anti-pSmad1/5/8 (AB3848, 1:1,000; Millipore) and anti-Mllt6 (NBP1-89222, 1:100; Novus Biologicals). Secondary antibodies were conjugated to Alexa-488, 546 or 647 (1:1,000, Life Technologies). Detection of pSmad1/5/8 was enhanced using the Tyramide Signal Amplification (Perkin Elmer). Cells and tissues were processed as previously reported<sup>1</sup>, and mounted in ProLong Gold with DAPI (Life Technologies). Skin squamous cell cancer tissue array (SK802a) was obtained from US Biomax Inc. Immunohistochemistry preparations were developed using ImmPRESS Universal Antibody Polymer Detection method (Vector Laboratories). Confocal images were captured by a scanning laser confocal microscope (LSM510 and LSM780; Carl Zeiss) using Plan-Apochromat 20×/0.8 oil and C-Apochromat 40×/1.2 water lenses. Images were processed using ImageJ and Adobe Photoshop CS3. To quantify the number of ectodermal cells at E9.5, embryos were fixed in 4% paraformaldehyde, permeabilized in PBS + 0.1% Triton (Sigma), and the nuclei were labelled with TO-PRO-3 as recommended (Life Technologies). Tiled Z-stack images of were collected on a Zeiss LSM780 using a Plan-Apochromat 63×/1.4 oil lens. Stacks and metadata were imported into MATLAB (Mathworks) using the LOCI Bio-Formats Importer<sup>36</sup>. For each stack, the surface was located by finding the first Z position with an average intensity threefold above background, and the stack was cropped to 6 µm corresponding to the surface epithelium. The resulting images were segmented in three dimensions using Imaris (Bitplane AG) to obtain counts of nuclei. **Flow cytometry.** Primary epidermal keratinocytes were isolated<sup>37</sup> and then purified by fluorescence activated cell sorting (FACS) using BD FACSAria II (BD Biosciences). Nucleotide analogue EdU (50 mg per g body weight) was injected intraperitoneally 2 h before processing, and EdU (Life Technologies) incorporation and active caspase 3 (BD Pharmingen) assays were performed as recommended. Immune cell infiltration was analysed in whole skin dissociated with a sequential incubation in collagenase (Sigma; 0.25% in HBSS for 90 min) and trypsin (Gibco; 0.25% in PBS for 15 min) at 37 °C. The following biotin-conjugated rat antibodies (1:100; Pharmingen) were used: anti-CD11b (MI/70), anti-CD103 (M290), anti-Ly-6G/C (RB6-8C5), anti-CD3e (145-2C11), anti-CD45 (30-F11) and anti-CD45R (RA3-6B2). YFP/RFP quantification was based on detection of the native protein in unfixed cells. Flow cytometry analysis was performed on BD LSR II.

**Cell culture assays.** Cells were cultured in 0.05 mM Ca<sup>2+</sup> (E18.5 mouse epidermal keratinocytes and SCC cells) or 1.5 mM Ca<sup>2+</sup> (human SCC) E-media supplemented with 15% serum. Cell adhesion in primary epidermal keratinocytes seeded at low confluence was assayed by replacing their growth medium with a 1.5 mM Ca<sup>2+</sup> E-media, and fixing them at different times thereafter. Nucleotide analogue EdU (10 µM) was added to cell culture media 90 min before processing, and contact inhibition was analysed in cells 3 days after reaching confluence. Inhibition of Wnt signalling was achieved by addition of 5 µM tankyrase inhibitor

XAV939 (IC<sub>50</sub> values 11 nM (Tnks1) and 4 nM (Tnks2)) to the media 12 h before the start of the experiments.

**Lentiviral constructs.** Sequences of RNAi constructs are listed in Supplementary Table 5. Design of LV-RFP, LV-GFP and LV-Cre has been previously reported<sup>1</sup>. Lentiviral construct for inducible shRNA expression is a modification of tet-pLKO-puro<sup>38</sup> (Addgene plasmid 21915), where the IRES-Puro cassette was replaced between the XmaI and KpnI sites with the ligation of PCR-amplified XmaI/NheI-flanked *P2A* fragment and NheI/KpnI-flanked *nlsCre* cDNA. The lentiviral Wnt-reporter was fashioned after the lentiviral Beta-catenin Activated Reporter<sup>39</sup>. It includes 12 *Tcf/Lef* binding sites followed by a minimal TK promoter and an *mRFP1* transgene that were subcloned into a pLKO.1 backbone between KpnI and NheI sites. Lentiviral bone morphogenetic protein-reporter that contains a pair of bone morphogenetic protein response elements is a derivative of BRE-ZsGreen<sup>4</sup>, where the *reporter* cassette between XhoI and NheI sites has been placed between SalI and NheI sites of the pLKO-nlsCre-MCS vector.

**Tumour-free survival.** Control and *Hras<sup>oncX2</sup>* animals were transduced at E9.5 with low-titre LV-Cre containing constitutively expressing or inducible shRNA against scrambled control or test *Ctnnb1* and *Mllt6* shRNAs. Transductions were confirmed by real-time PCR of P7 (newborn) littermates, and the remaining animals were monitored for an additional 12 weeks. Animals were assessed every 2–3 days, and scored positive when tumours were larger than 2 mm in diameter. Animals transduced with an LV-Cre containing inducible shRNAs were allowed to form tumours for 60 days, at which point individual tumours were measured along their short and long axis using a digital caliper ( $t = 0$ ). Next, tumour-bearing animals were treated by a single intraperitoneal injection of doxycycline (100 µl of 50 mg ml<sup>-1</sup>) and maintained on doxycycline-containing chow for 8 weeks, and tumour size was assayed every 7 days. Because the tumour volumes at  $t = 0$  showed a range between 4–20 mm<sup>3</sup>, the assayed tumour size was normalized to the initial tumour volume, and expressed as fold-change over time. Transplantation of SCC cells transduced with control shRNA, or shRNAs targeting mouse and human *Ctnnb1* and *Mllt6*, into immunocompromised nude recipients were performed as previously described<sup>28</sup>, and animals were monitored every 3 days for a month. Tumour size was measured using a digital caliper, and tumour volume was calculated using the formula [(length × width)<sup>2</sup> × π]/6.

**Tcf3/4 ChIP-seq and luciferase assay.** Details of the Tcf3/4 ChIP-seq will be reported elsewhere. For luciferase assays, passage 9–14 293FT cells were seeded in 96-well culture plates and transfected at 60–70% confluence using standard calcium phosphate procedures. Cells were co-transfected with control Renilla pRL-TK, and combinations of 50 ng pGL3-Mllt6, Mllt6<sup>mutant</sup>, Axin2 or Axin2<sup>mutant</sup> and 200 ng of K14-expression vectors encoding Lef1, ΔNβ-cat or control (empty vector). After 44 h, cells were collected and luciferase activity was measured using the TD-20/20 luminometer (Turner Biosystems) and the Dual-Glo Luciferase Assay System (Promega). Each transfection was performed in duplicate and repeated seven times.

**Sample preparation and pre-amplification.** Epidermal cells were isolated from E18.5 mouse skin using previously established procedures<sup>37</sup>. Cells from individual embryos were used for genomic DNA isolation with the DNeasy Blood & Tissue Kit (Qiagen), and each sample was analysed for target transduction using real-time PCR. gDNAs from 30 transduced embryos were pooled, and 200 µg of the total was used as template in a 10-ml pre-amplification reaction with 21 cycles and Phusion High-Fidelity DNA Polymerase (NEB). PCR products were run on a 2% agarose gel, and a clean ~200-bp band was isolated using QIAquick Gel Extraction Kit as recommended by the manufacturer (Qiagen). Final samples were then sent for Illumina HiSeq 2000 sequencing.

**Sequence processing and relative shRNA quantification.** For each genotype, DNA from 30 embryos was pooled and independently sequenced using custom forward (5′-AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTCCGATCTATATCTTGTGGAAAGGACGAAACACC-3′) and reverse (5′-CAAGCAGAAGACGGCATACGAGCTCTCCGATCTAATTGTGGATGAATACTGCCATTGTGTC-3′) oligonucleotides. Illumina reads were trimmed to the 21-nucleotide hairpin sequence using the FASTX-Toolkit and aligned to the TRC 2.x library with BWA (v0.6.2)<sup>40</sup> using a maximum edit distance of 3. Putative growth regulators were identified by combining two methodologies. First, Illumina reads from 3 sets of 30 embryos were treated as independent biological replicates in the DESeq R package<sup>42</sup>. Dispersions (variability) for each hairpin were estimated using a local fit to the data for each genotype, and hairpins with a  $P$  value < 0.05 by the negative binomial test were considered for downstream analysis. Second, an analysis was carried out on a pooled data set in which the reads from three sets of embryos were combined to maximize screen coverage and average biological variability. Although this precludes estimation of within-group variability, it has the effect of reducing noise for poorly counted hairpins when operating close to the minimum required screen coverage. Fisher's exact test was applied on a per-hairpin basis using combined reads by assembling a 2 × 2 contingency table<sup>41</sup>. The columns of the table

are the treatment conditions (for example, control and Hras<sup>oncoX2</sup>), and the rows correspond to the sequencing counts for a given hairpin in the first, and the counts for all other hairpins in the pool in the second. The test thus calculates the probability of observing a difference in hairpin representation relative to the expected representation in the pool. We further adjust the *P* value for multiple testing using the Benjamini–Hochberg correction. Hairpins with a *P* value < 0.05 were considered for further analysis. A gene was considered significantly enriched or depleted if at least two hairpins exhibited a twofold or greater change in normalized reads with a significant *P* value, and no hairpins in the set exhibited a change of equal magnitude in the opposite direction. Hits common to both analyses were ranked by number of significant hairpins and the magnitude of their effect. All analyses were carried out in the R statistical environment<sup>42</sup>, with some plots produced using the ggplot2 package<sup>43</sup>. Gene lists were imported into the Ingenuity Pathway Analysis software (Ingenuity Systems), and analyses and graphic outputs of relative enrichment in functional gene categories were performed as recommended.

**Choice of statistical analysis of relative shRNA abundance.** The strength of the Fisher's exact test is that it can calculate a probability of observing a difference in shRNA representation in a comparison of pooled data sets. When operating close to the minimum required screen coverage or when it is not feasible to perform many independent replicates, this strategy can be advantageous to increase coverage and reduce noise if combined with additional stringent criteria (that is, requiring a gene to be targeted by multiple hairpins) and validation.

Because this methodology does not explicitly account for sample-to-sample variability (instead maximizing coverage and averaging out variability), we independently analysed our data using two additional statistical methodologies that directly address variability within biological replicates. Importantly, both of these methods used the same stringent set of thresholds (twofold change in hairpin count, and a requirement for least two hairpins to show a significant effect in same direction, and none in the opposite). First, we used DESeq<sup>12</sup>, an R package designed for the analysis of Illumina sequencing-based assays, which estimates and accounts for biological variability in a statistical test based on the negative binomial distribution. Second, we treated independently sequenced sets of 30 embryos as biological replicates, and generated replicate-specific lists of candidate genes. Comparison of hits shared between these replicates to the hits identified in our analysis of pooled samples yielded a highly conserved set of candidate genes consistent with strong reproducibility of our data. Both analyses identify a list of candidates that substantially overlaps with those identified by our pooling and ranking scheme, with nearly all of our top hits identified regardless of the methodology.

When conducting shRNA drop-out screens, perhaps the most important criteria in identifying potential candidates is that a gene be targeted by multiple, independent hairpins to avoid off-target effects. We thus felt our data would be best-served by combining an analysis of pooled data, which tends to be more

inclusive at the level of hairpins and enables ranking by number of independent hairpins, and the results of DESeq, which ensures reproducibility of hits.

**RNA-seq and IPA network analyses.** Epidermal progenitors were FACS sorted into TrizolLS (Invitrogen) and RNA was purified using Direct-zol RNA MiniPrep kit (Zymo Research) per manufacturer's instructions. Quality of the RNA was determined using Agilent 2100 Bioanalyzer, with all samples passing the quality threshold of RNA integrity numbers (RIN) > 8. Library preparation using Illumina TrueSeq mRNA sample preparation kit was performed at the Weill Cornell Medical College Genomic Core facility, and cDNA was sequenced on Illumina HiSeq 2000. Reads were mapped to mm9 build of the mouse genome using TopHat, and transcript assembly and differential expression were determined using Cufflinks<sup>44</sup>. Differentially regulated transcripts were analysed in IPA (Ingenuity Systems), and the upstream transcriptional regulators were predicted using the Upstream Regulator Analysis package, with a significant overlap between the data set genes and transcription factor targets set at *P* < 0.01, and the regulation direction (activated or inhibited) at *z*-score = 2.

**Statistics.** All quantitative data were collected from experiments performed in at least triplicate, and expressed as mean ± s.d. or s.e.m. The fits of cellular and tumour growth were compared using the extra sum-of-squares *F*-test, and expression of CTNBN1 and MLLT6 in human SCC tissue was analysed using a non-parametric (Spearman) correlation. Differences between groups were assayed using two-tailed student *t*-test using Prism 5 (GraphPad Software). Significant differences were considered when *P* < 0.05.

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