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Research Article

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Cancer susceptibility and embryonic lethality in *Mob1a/1b* double-mutant mice

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Mps one binder 1a (MOB1A) and MOB1B are key components of the Hippo signaling pathway and are mutated or inactivated in many human cancers. Here we show that intact *Mob1a* or *Mob1b* is essential for murine embryogenesis and that loss of the remaining WT *Mob1* allele in *Mob1a^{Δ/Δ}1b^{tr/+}* or *Mob1a^{Δ/+}1b^{tr/tr}* mice results in tumor development. Because most of these cancers resembled trichilemmal carcinomas, we generated double-mutant mice bearing tamoxifen-inducible, keratinocyte-specific homozygous-null mutations of *Mob1a* and *Mob1b* (*kDKO* mice). *kDKO* mice showed hyperplastic keratinocyte progenitors and defective keratinocyte terminal differentiation and soon died of malnutrition. *kDKO* keratinocytes exhibited hyperproliferation, apoptotic resistance, impaired contact inhibition, enhanced progenitor self renewal, and increased centrosomes. Examination of Hippo pathway signaling in *kDKO* keratinocytes revealed that loss of *Mob1a/b* altered the activities of the downstream Hippo mediators LATS and YAP1. Similarly, YAP1 was activated in some human trichilemmal carcinomas, and some of these also exhibited MOB1A/1B inactivation. Our results clearly demonstrate that MOB1A and MOB1B have overlapping functions in skin homeostasis, and exert their roles as tumor suppressors by regulating downstream elements of the Hippo pathway.

Introduction

Tissue homeostasis requires a balance of cell proliferation, apoptosis, and differentiation. During tissue development, progenitor cells divide for a limited number of times before exiting the cell cycle and undergoing terminal differentiation (1). When progenitor cells proliferate inappropriately and/or undergo abnormal differentiation, cancer can result (2).

Skin is a tissue that begins as a single-layered epithelium made up of keratinocytes that later stratify and differentiate into hair follicles (HFs) or interfollicular epidermis (IFE). Disruption of signals balancing growth and differentiation in keratinocytes can initiate skin cancers. The most common skin cancer is basal cell carcinoma (BCC), the nodular subtype of which originates from HFs. Trichilemmal carcinomas are also HF-derived skin cancers, but are much more rare and show characteristic abnormalities of outer hair root sheath differentiation (3). Although impairment of the Sonic Hedgehog (SHH) intracellular signaling pathway is now well established as the cause of BCC (4, 5),

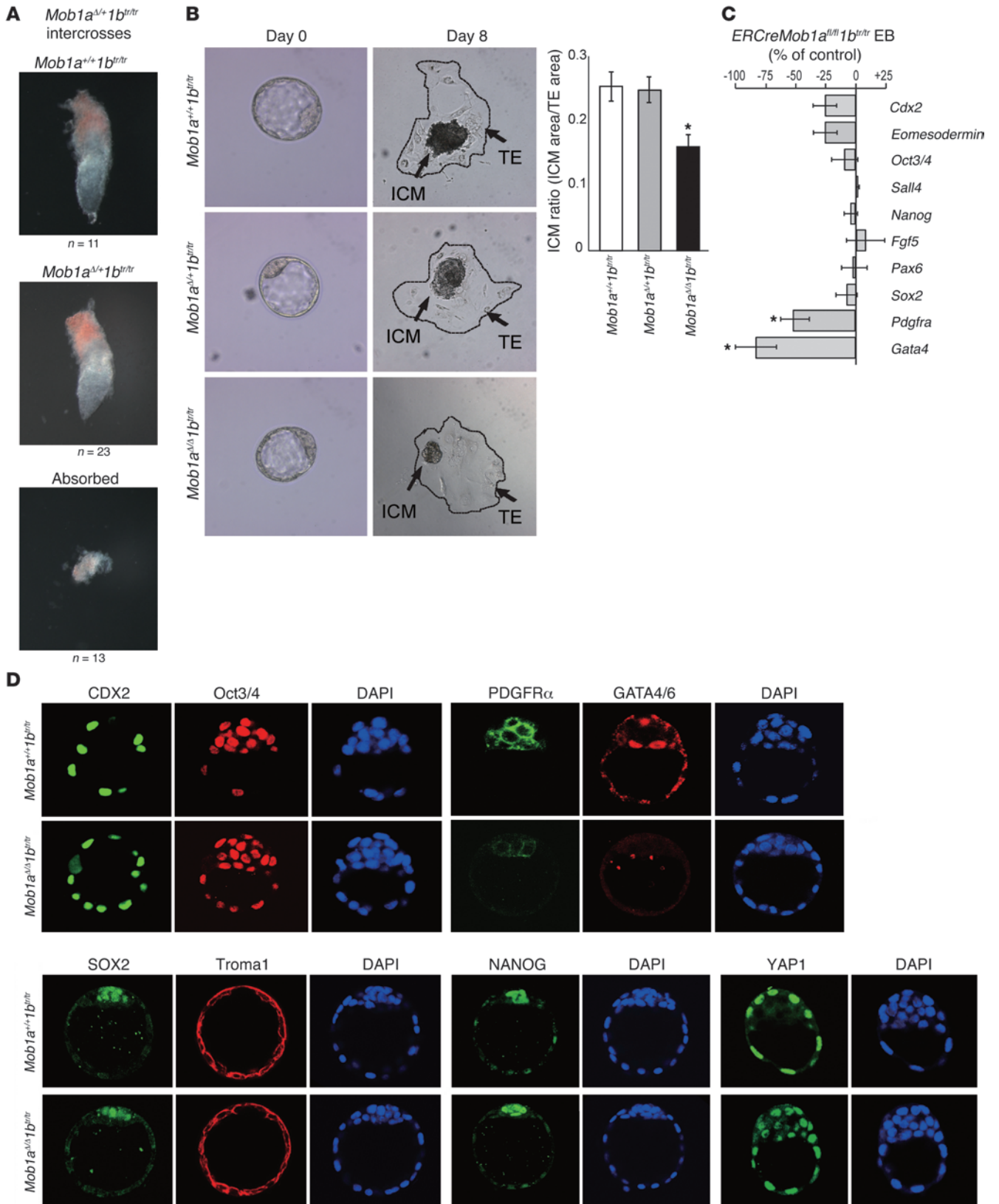
the molecular mechanism underlying the development of trichilemmal carcinomas is unknown.

The evolutionarily conserved Hippo signaling pathway was first identified as regulating the cell proliferation and apoptosis-controlling organ size in *Drosophila* (6, 7). In mammals, the canonical Hippo pathway includes the following: Neurofibromin 2 (NF2) (8), the Mammalian STE20-like protein (MST) kinases (9), large tumor suppressor homolog (LATS) and Nuclear Dbf2-related (NDR) kinases (10), the adaptor proteins SAV1 (WW45) (11) and MOB (12), and their downstream transcriptional coactivators YAP1 and its paralog transcriptional coactivator with PDZ-binding motif (TAZ, WWTR1) (13). All of these molecules (except YAP1 and TAZ) have been implicated as tumor suppressors. For example, mice deficient in *Lats1* or *Mst1/2* develop various cancers (14, 15), *SAV1* and *MOB* are mutated in human cancer cell lines (16, 17), and Hippo components are reduced in human cancer samples (18). Homozygous-null mutant mice lacking *Nf2*, *Mst1/2*, *Sav1*, *Lats2*, or *Yap1* are all embryonic lethal (15, 19–22), precluding study of these mediators in tumorigenesis. To complicate matters, multiple homologs of each mammalian Hippo element exist, and a noncanonical Hippo pathway influences the localization and activity of canonical Hippo components (15, 23). Thus, the precise physiological functions of Hippo elements in mammals remain obscure.

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**Figure 1**

Loss of *Mob1a/1b* impairs mouse embryo survival. (A) Morphologies of representative E6.5 embryos of the indicated genotypes from *Mob1a^{N/+}1b^{tr/tr}* intercrosses. *Mob1a^{N/Δ}1b^{tr/tr}* embryos were absorbed after implantation. (B) Left: photographs of E3.5 blastocysts of the indicated genotypes on day 0 and after culture for 8 days. The ICM is surrounded by TE giant cells. Right: mean ICM ratio (ICM area/TE area) ± SEM plotted for blastocysts of the indicated genotypes ($n = 8/\text{group}$); $*P < 0.01$. (C) Quantitative PCR determination of mRNAs for the indicated genes in EBs generated from *ERCreMob1a^{fl/fl}1b^{tr/tr}* ES cells. ES cells were cultured with/without tamoxifen for 2 days, and control and mutant EBs were generated using the hanging drop method after culture for 3 days (*Cdx2*, *Eomesodermin*, *Oct3/4*, *Sall4*) or 4 days (*Nanog*, *Fgf5*, *Pax6*, *Sox2*, *Pdgfra*, *Gata4*). mRNA levels in the mutant are expressed as the percentage increase over control values; $*P < 0.05$. (D) Immunostaining to detect the indicated proteins in blastocysts of the indicated genotypes. Nuclei were visualized with DAPI. Results shown are representative of at least 3 independent trials and at least 3 mice/group. Data are presented as the mean ± SEM, and *P* values were determined using the 2-tailed Student's *t* test.

MOB1 was originally shown in yeast to regulate mitotic exit and modulate cytokinesis (24–29). In *Drosophila*, *dMOB1* (*MATS*) overexpression is tumor suppressive, while loss of *dMOB1* triggers tumor development (17). In humans, 7 MOB homologs (hMOB1A-B, 2A-C, 3, 4, where MOB indicates Mps one binder) have been identified (30), and hMOB1A (hMATS1, MOBKL1B) and hMOB1B (hMATS2, MOBKL1A) share 95% amino acid identity. Although no apparent functional domain has been found in hMOB1A/1B, only these 2 MOB proteins can bind to and activate LATS1/2 (31). In vitro, hMOB1A overexpression inhibits cell proliferation, while suppression of hMOB1B or hMOB1A mediated by shRNA or siRNA increases proliferation (30) or impairs mitotic exit (32), respectively. In vivo, the *hMOB1A* gene is mutated in melanoma and breast cancer cell lines, and its expression is downregulated in human colorectal and non-small-cell lung cancers (17, 33, 34). However, the normal function of hMOB1 proteins in vivo is unknown. To assess whether MOB1 is an important functional component of mammalian Hippo signaling and to determine whether MOB1 is truly a tumor suppressor in vivo, we generated and characterized double-mutant mice lacking *Mob1a* and *Mob1b*.

Results

Embryonic lethality of *Mob1a/1b*-null mutant mice. We first generated single-mutant mice bearing a null mutation of *Mob1a* (*Mob1a^{N/Δ}*) or a trapped mutation of *Mob1b* (*Mob1b^{tr/tr}*) (Supplemental Figure 1, A–D; supplemental material available online with this article; doi:10.1172/JCI63735DS1) but no abnormalities were observed in morphology, body weight, histology, or life span (Supplemental Figure 1E and data not shown). We then attempted to generate double-homozygous-null mutant (*Mob1a^{N/Δ}1b^{tr/tr}*) mice, but no viable pups were obtained (Supplemental Table 1), indicating that complete loss of *Mob1* is embryonic lethal. We next analyzed embryos from *Mob1a^{N/+}1b^{tr/tr}* intercrosses at various time points during gestation. By E6.5, although decidua were formed, 28% of embryos were absorbed too severely to genotype (Figure 1A and Table 1). Thus, MOB1A and MOB1B have overlapping functions, and MOB1 is essential for postimplantation embryogenesis.

To pinpoint the embryonic defect, we obtained individual blastocysts (E3.5) from *Mob1a^{N/+}1b^{tr/tr}* intercrosses and cultured them

for 8 days. *Mob1a^{N/Δ}1b^{tr/tr}* blastocysts appeared normal at E3.5, but showed growth failure of the inner cell mass (ICM) by day 8 (Figure 1B). In contrast, the trophoctoderm (TE) of *Mob1a^{N/Δ}1b^{tr/tr}* blastocysts developed normally.

To create conditional tamoxifen-inducible *Mob1* mutants, we generated *ERCreMob1a^{fl/fl}1b^{tr/tr}* ES cells and established embryoid bodies (EBs) using the hanging drop method. Control and mutant EBs were cultured for 2 days with (or without) tamoxifen to delete *Mob1* and then grown without leukemia inhibitory factor (LIF) for 3–4 days. Quantitative RT-PCR examination showed that *Gata4* and *Pdgfra* mRNAs (essential for primitive endoderm formation) were markedly suppressed in *Mob1a^{N/Δ}1b^{tr/tr}* EBs, whereas *Cdx2* and *Eomesodermin* mRNAs (essential for TE) were only slightly inhibited (Figure 1C). Levels of *Oct3/4* (*Pou5f1*) and *Sall4* mRNAs (essential for ICM), as well as *Nanog*, *Fgf5*, *Pax6*, and *Sox2* mRNAs (essential for primitive ectoderm), were normal in the double-mutant EBs. Immunostaining of double-mutant blastocysts with antibodies recognizing CDX2 or Troma-1 (TE markers), NANOG or SOX2 (primitive ectoderm markers), or Oct3/4 confirmed that the levels of these proteins were also normal in the absence of *Mob1*. However, levels of GATA4 and PDGFRA proteins (primitive endoderm markers) were greatly reduced in the mutants (Figure 1D). Moreover, YAP1, which is usually expressed only in the nuclei of TE cells, was abnormally activated and expressed in the nuclei of ICM cells (Figure 1D). Thus, *Mob1a^{N/Δ}1b^{tr/tr}* embryos have a defect in primitive endoderm formation.

***Mob1a^{N/+}1b^{tr/tr}* and *Mob1a^{N/Δ}1b^{tr/+}* mice show increased susceptibility to tumorigenesis.** *Mob1a^{N/+}1b^{tr/tr}* and *Mob1a^{N/Δ}1b^{tr/+}* mice were born alive and were healthy and fertile, but 52% of them had dental malocclusion (Supplemental Figure 2A). In addition, 25% of these heterozygotes exhibited disequilibrium due to disorganized inner ear hair bundles in the cochlear organ of Corti (Supplemental Figure 2B and Supplemental Video 1), and 31% showed histological evidence of increased trabeculae in femurs (Supplemental Figure 2C).

Because mutation or reduced expression of *MOB1A* occurs in cancer patients (33), we monitored tumorigenesis in *Mob1a^{N/+}1b^{tr/tr}*, *Mob1a^{N/Δ}1b^{tr/+}*, and *Mob1a^{N/+}1b^{tr/+}* (control) mice over 70 weeks. Various tumor types arose spontaneously in 100% of *Mob1a^{N/+}1b^{tr/tr}* and *Mob1a^{N/Δ}1b^{tr/+}* mice, but in only 4% of controls (Figure 2, A and B, and Table 2). Southern blotting confirmed the loss of the WT *Mob1* allele in all tumors (Figure 2C). In most mutants, skin tumors in the form of facial swellings (Figure 2A) were observed. Benign exos-

Table 1

Viability of embryos of the indicated *Mob1a* genotypes from *Mob1a^{N/+}1b^{tr/tr}* intercrosses at the indicated stages of embryogenesis

	Total	Genotype of <i>Mob1a</i>			Absorbed
		+/+	Δ/+	Δ/Δ	
Viable	108	39	69	0	
E16.5	3	1	2	0	
E14.5	15	4	11	0	
E10.5	19	8	9	0	2
E9.5	27	11	13	0	3
E7.5	19	3	12	0	4
E6.5	47	11	23	0	13
E3.5	33	9	16	0	0

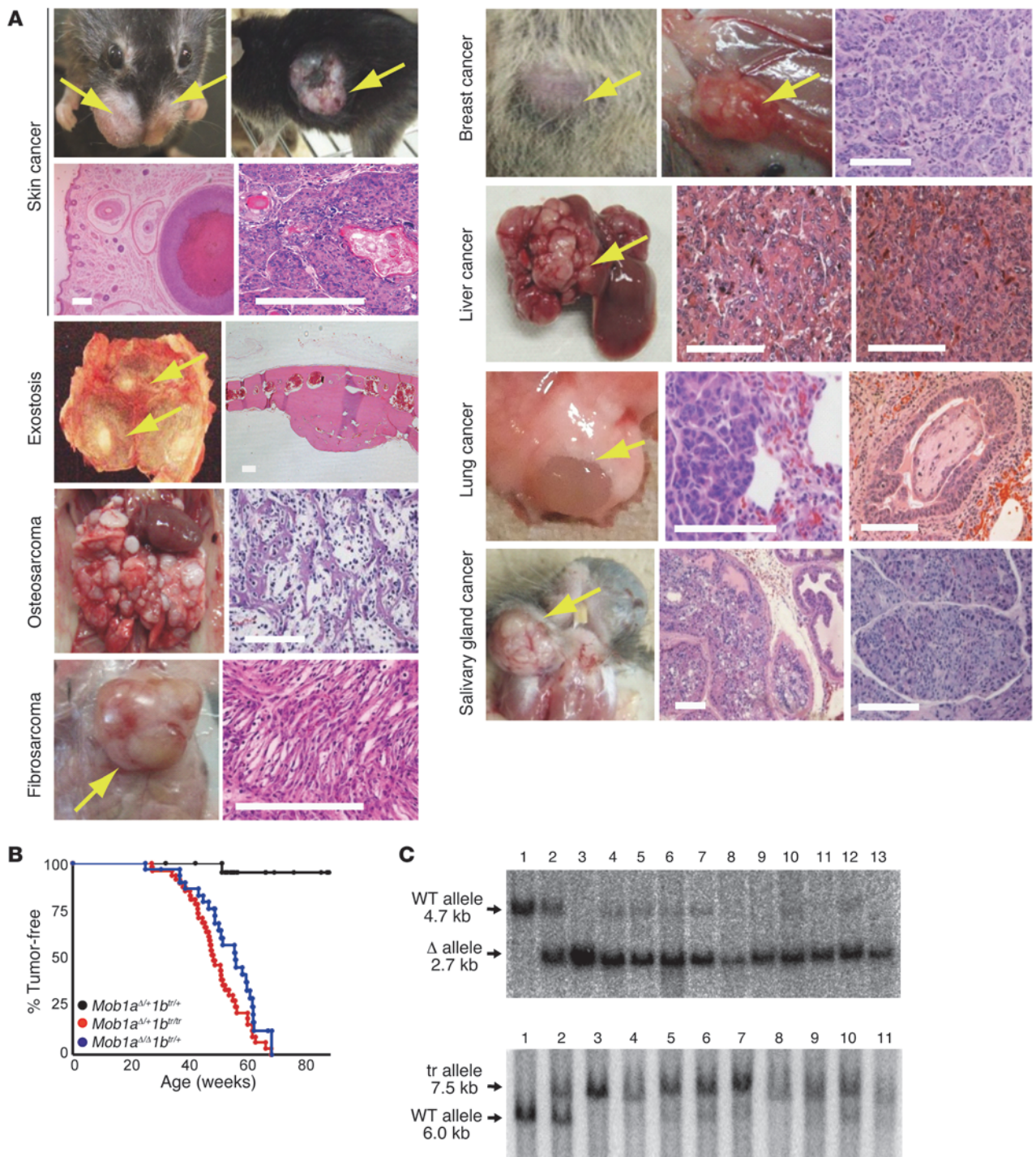


Figure 2

Loss of *Mob1a/1b* promotes tumorigenesis. **(A and B)** Tumor types arising in *Mob1a*^{Δ/Δ}*1b*^{tr/+} or *Mob1a*^{Δ/+}*1b*^{tr/tr} mice. **(A)** Macroscopic or microscopic photographs of the indicated gross or H&E-stained tumors in mutant mice. Yellow arrows, tumor masses. Scale bars: 100 μm. **(B)** Kaplan-Meier analysis of tumor onset and survival for the *Mob1a*^{Δ/Δ}*1b*^{tr/+} (*n* = 31), *Mob1a*^{Δ/+}*1b*^{tr/tr} (*n* = 48), and control *Mob1a*^{+/+}*1b*^{tr/+} (*n* = 22) mice in **A**. **(C)** Southern blots of tumor DNA showing loss of the WT *Mob1* allele. Top (hybridized to probe A of Supplemental Figure 1A): lanes 1–3, control thymic DNA from *Mob1a*^{+/+}, *Mob1a*^{Δ/+}, and *Mob1a*^{Δ/Δ} mice, respectively; lanes 4–5, osteosarcomas; 6–7, fibrosarcomas; 8–9, skin cancers; 10–11, breast cancers; 12–13, salivary gland cancers of *Mob1a*^{Δ/+}*1b*^{tr/tr} mice. Bottom (probe C): 1–3, control tail DNA of *Mob1b*^{+/+}, *Mob1b*^{tr/+}, and *Mob1b*^{tr/tr} mice, respectively; 4–5, osteosarcomas; 6–7, fibromyosarcomas; 8–9, skin cancers; 10–11, liver cancers of *Mob1a*^{Δ/Δ}*1b*^{tr/+} mice. Results shown are representative of at least 3 independent trials and at least 3 mice/group. Data are presented as the mean ± SEM, and *P* values were determined using the 2-tailed Student's *t* test.



Table 2
Total tumor numbers (incidence) for the mice in Figure 2A

	<i>Mob1a^{Δ/+}1b^{tr/tr}</i>	<i>Mob1a^{Δ/Δ}1b^{tr/+}</i>	Overall
Skin cancer	23 (100%)	14 (100%)	37 (100%)
Exostosis	20 (87%)	14 (100%)	34 (92%)
Osteosarcoma	7 (30%)	2 (14%)	9 (24%)
Fibrosarcoma	6 (26%)	2 (14%)	8 (22%)
Liver cancer	0 (0%)	7 (50%)	7 (19%)
Breast cancer	5 (22%)	1 (7%)	6 (16%)
Lung cancer	0 (0%)	2 (14%)	2 (5%)
Salivary gland cancer	1 (4%)	1 (7%)	2 (5%)

tosis in calvaria occurred in 92% of animals with just 1 WT *Mob1* allele as well as extraskeletal osteosarcomas (24%), subcutaneous fibrosarcomas (or myofibrosarcomas) (22%), breast cancers (16%), lung cancers (5%), and salivary gland cancers (5%). Liver cancers were found only in *Mob1a^{Δ/Δ}1b^{tr/+}* mutants (50%). Interestingly, none of our *Mob1*-deficient mice developed colon tumors or melanomas, unlike MOB1A-deficient humans (17, 33). Nevertheless, MOB1 is a powerful tumor suppressor in multiple tissues in vivo.

Keratinocyte-specific Mob1a/1b-deficient mice show gross skin abnormalities and die during lactation. Because the dominant tumor in *Mob1a^{Δ/+}1b^{tr/tr}* and *Mob1a^{Δ/Δ}1b^{tr/+}* mice arose in the skin, we generated tamoxifen-inducible, keratinocyte-specific *Mob1a/1b* double-homozygous mutant mice (*Krt14CreERMob1a^{fl/fl}1b^{tr/tr}*). The *Keratin-14* (*Krt14*) promoter directs gene expression in the basal layer of epidermal and follicular keratinocytes, including stem cells in the HF bulge. Thus, tamoxifen-induced, *Krt14*-controlled Cre-mediated deletion of a floxed gene disrupts expression of that gene throughout the IFE and HFs (35). When we administered tamoxifen to *Krt14CreERMob1a^{fl/fl}1b^{tr/tr}* mice at P1, we created double-homozygous conditional mutant mice [*kDKO*(P1)] that initially appeared healthy. Single-mutant mice (*Krt14CreERMob1a^{fl/fl}1b^{tr/tr}*) without tamoxifen or *Krt14CreERMob1a^{fl/fl}1b^{+/+}* mice with tamoxifen were indistinguishable from the WT in gross appearance, histology, and survival (data not shown), allowing us to choose *Krt14CreERMob1a^{fl/fl}1b^{tr/tr}* mice without tamoxifen as controls. Southern blotting confirmed that tamoxifen disrupted the *Mob1a^{fllox}* allele in almost all keratinocytes of *kDKO*(P1) mice, leaving the *Mob1a^Δ* allele (Supplemental Figure 3A). In addition, immunostaining using an Ab recognizing both MOB1A and MOB1B (Supplemental Figure 3B) confirmed that the MOB1A and MOB1B proteins are expressed both in IFE cells (especially in the granular layer) and in HF cells (especially in matrix cells and outer root sheath cells) in normal mice and humans (Supplemental Figure 3, C and D). In contrast, MOB1A and MOB1B proteins were not detected in the epidermis of *kDKO*(P1) mice (Supplemental Figure 3C).

By 13 days of age, *kDKO*(P1) mice were significantly smaller than control littermates and readily identified by their “wrinkled-bear” facial skin (Figure 3A) and ruffled hair (Figure 3A). These mutants’ front paws (Figure 3A), ears, and lips were hyperplastic and enlarged. All *kDKO*(P1) mice died of malnutrition within 10–30 days of birth (Figure 3), likely due to dysphagia caused by hyperplasia of the oral cavity epithelium.

We next analyzed the architecture of *kDKO*(P1) skin at various time points after birth. In WT mice, dividing keratinocytes are restricted to the basal epidermal layer. As these cells exit the cell

cycle, they differentiate to form the spinous and granular skin layers as well as the dead, enucleated stratum corneum layers. The anagen phase of hair development thickens the skin until day 17, when the first catagen phase is triggered and skin thickness regresses until day 19. Hair remodeling then begins its lifelong cycle of spontaneous regrowth and regression (36). We saw this pattern clearly in our biopsy series from control mice (Supplemental Figure 4). However, the IFE and HFs of *kDKO*(P1) mice showed hyperplastic multilayered epithelium at day 16 (Figure 3B), and impaired epidermal regression during catagen (Supplemental Figure 4; day 19). In addition, parakeratosis (reduced enucleation) was evident in the stratum corneum layer of *kDKO*(P1) mice (Figure 3C).

To investigate keratinocyte differentiation, we used immunostaining to detect keratin markers in the fat pad epidermis, which allows easy analysis of epidermal layers. Keratin-15⁺ (KRT15⁺) cells, the most primitive in IFE, normally form a single basal layer attached to the IFE basement membrane (BM). This configuration was clearly visible in our controls, whereas the KRT15⁺ cells of *kDKO*(P1) mice, although nearly normal in number, were scattered inside the IFE without attachment to the BM (Figure 3D). In control mice, KRT14⁺ and KRT5⁺ cells were present in 1 or 2 IFE layers (as expected), but appeared in multiple layers in *kDKO*(P1) mice (Figure 3D). Moreover, there were increased numbers of irregularly organized KRT10⁺ cells in the suprabasal layers of *kDKO*(P1) epidermis (Figure 3D). Filaggrin⁺ cells, the most highly differentiated keratinocytes, were present in *kDKO*(P1) epidermis but at a reduced percentage (Figure 3D). Similar IFE alterations were observed in the back skin of *kDKO*(P1) mice (Figure 3E and Supplemental Figure 5). KRT17⁺ outer root sheath cells were significantly increased in back skin HFs of *kDKO*(P1) mice, and Trichohyalin⁺ inner root sheath cells were elevated slightly (Figure 3E). Thus, the phenotype of *kDKO*(P1) mice may be largely due to abnormal expansion of immature stem/progenitor skin cells.

To try to overcome the early lethality of *kDKO*(P1) mice, we waited until *kDKO* mice reached 28 days of age before treating them with tamoxifen for 7 days. However, like *kDKO*(P1) mice, all of these *kDKO*(P28) mice showed keratinocyte hyperplasia (Figure 3F) and died within 15–55 days after treatment (Supplemental Figure 6). Thus, MOB1 deficiency in epidermal cells severely disrupts normal IFE and HF development and homeostasis.

Tumorigenic anomalies in Mob1a/1b double-homozygous mutant keratinocytes. To determine whether loss of MOB1 conferred tumorigenic properties on keratinocytes, we first immunostained *kDKO*(P1) skin with anti-Ki67 Ab and found increased numbers of Ki67⁺ proliferating epithelial cells (Figure 4A). In control mice at P13, a few Ki67⁺ cells were located among basal IFE cells and outer root sheath cells, with more scattered among HF matrix cells. However, the incidence of Ki67⁺ cells in *kDKO*(P1) skin at P13 was 1.5 times higher than in controls, and Ki67⁺ cells were also found in the suprabasal layers normally quiescent in control epidermis. In addition, TUNEL-positive apoptotic cells were reduced in *kDKO*(P1) epidermis (Figure 4B). Thus, increased proliferation and repressed apoptosis may contribute to *kDKO*(P1) epidermal hyperplasia. Cell-plating studies revealed that the saturation density of *kDKO*(P1) keratinocytes was also increased (Figure 4C). Because the cell size (forward scatter) of control and mutant keratinocytes showed no difference by FACS analysis (data not shown), the increased saturation density of the mutant suggests impaired contact inhibition. Histological analysis confirmed that the mutant basal epidermal layer showed significantly increased cell density

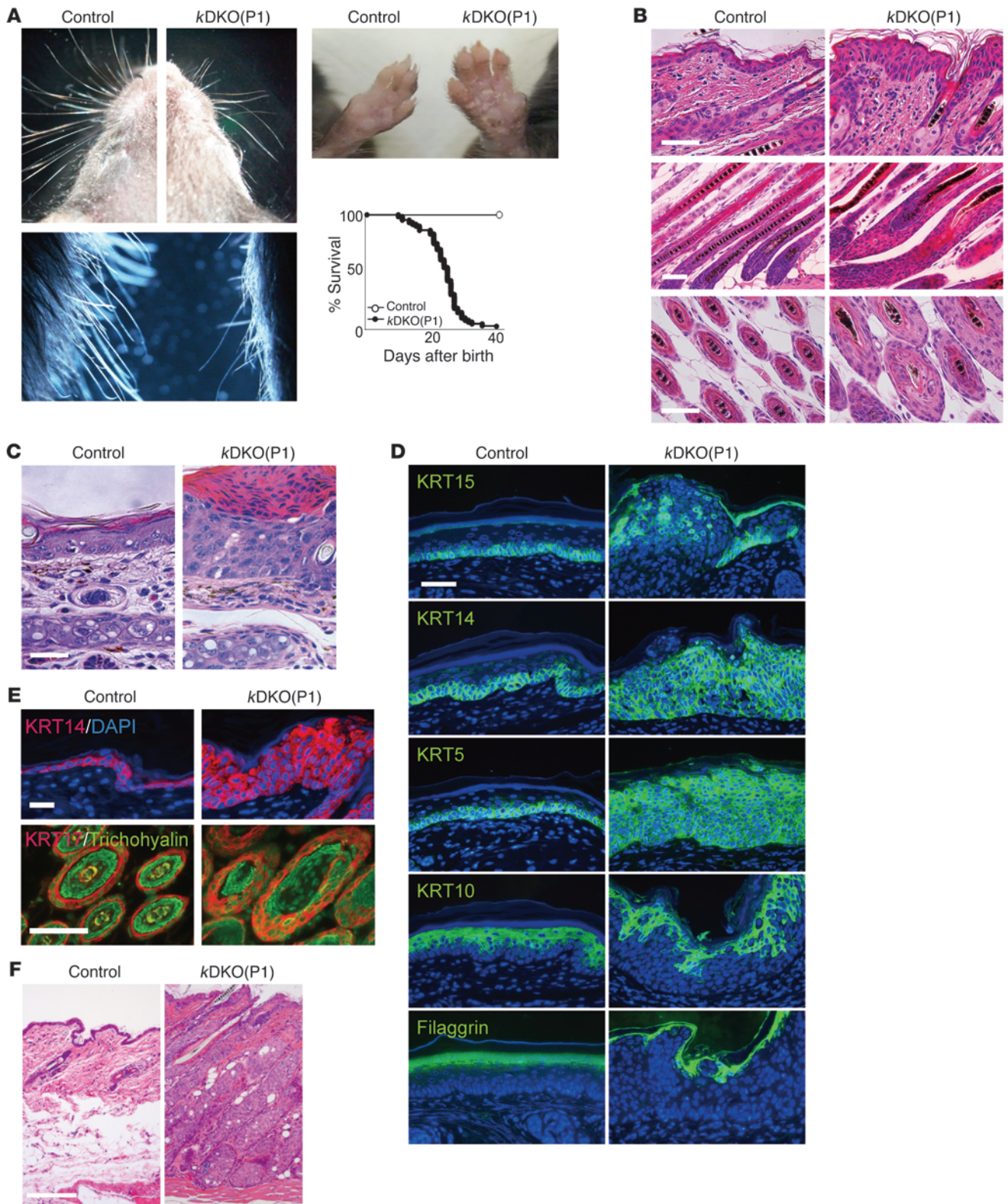


Figure 3

Characterization of keratinocyte-specific *Mob1a/1b* double-homozygous mutant mice. (A–C) Features of 16-day-old *Krt14CreERMob1a^{fl/fl}1b^{tr/tr}* mice that were originally left untreated (control) or treated with tamoxifen (*kDKO*) at P1. (A) *kDKO*(P1) mice have a wrinkled-bear facial appearance (top left), ruffled, shaggy body hair (bottom left), abnormally large front paws (top right), and decreased survival (bottom right). (B) H&E-stained longitudinal and transverse sections of the back skin of the mice in A. *kDKO*(P1) mice show hyperplasia of IFE and HF. Scale bars: 50 μ m. (C) *kDKO*(P1) mice show parakeratosis (reduced enucleation) in stratum corneum layers. Scale bar: 20 μ m. (D) Immunohistochemical analyses of fat pad epidermis of the mice in A using Alexa Fluor 488–tagged Abs recognizing the indicated differentiation markers corresponding to a specific skin layer. DAPI, nuclei. Scale bar: 50 μ m. (E) Immunohistochemical analyses of back skin of the mice in A using Abs recognizing the indicated skin layer markers. For top panels, cells were stained with Alexa Fluor 568–tagged anti-KRT14 Ab. DAPI, nuclei. Scale bars: 20 μ m. For bottom panels, cells were stained with Alexa Fluor 488–tagged anti-Trichohyalin Ab and Fluor 568–tagged anti-KRT17 Ab. Scale bar: 50 μ m. (F) H&E-stained longitudinal sections of skin from 42-day-old *Krt14CreERMob1a^{fl/fl}1b^{tr/tr}* mice that were left untreated (control) or treated with tamoxifen (*kDKO*) at P28. The *kDKO*(P28) mutant shows hyperplasia of IFE and HFs. Scale bar: 200 μ m. Results shown are representative of at least 3 independent trials and at least 3 mice/group. Data are presented as the mean \pm SEM.

(control: 7.6 ± 0.36 nuclei per 50 μ m BM; mutant: 8.9 ± 0.38 nuclei per 50 μ m BM; $n = 5$; $P < 0.05$). Morphologically, mutant basal cells were elongated and columnar rather than round like control basal cells (Figure 4D), also indicating defective contact inhibition.

Because yeast *MOB1* regulates mitotic exit and cytokinesis (24–29), we speculated that mammalian *MOB1* might affect processes that support cell division. Indirect immunofluorescence analysis of control and *kDKO*(P1) keratinocytes using anti- γ -tubulin and anti- α -tubulin Abs revealed an increase in centrosome number (>2 centrosomes/cell) (Figure 4E). Aberrant centrosome numbers are associated with multipolar spindle formation that impairs chromosome segregation. Indeed, a greater percentage of *kDKO*(P1) keratinocytes exhibited multi-polar spindles (Figure 4F). Micronuclei, which are cytoplasmic chromosomal fragments generated by aberrant mitosis, were also increased in *kDKO*(P1) keratinocytes (Figure 4G). Similar defects were observed in *Mob1a* and *Mob1b* double-homozygous mutant embryonic fibroblasts (MEFs) (Supplemental Figure 7, A–C).

To determine which mitotic stage was altered in *kDKO*(P1) keratinocytes, we divided the mitotic process into 4 parts based on cell morphology: part I, cells progressing from the round morphology of a resting cell to chromosome alignment (corresponding to prometaphase); part II, from chromosome alignment to ingression of the cleavage furrow (metaphase/late anaphase); part III, from ingression to furrow completion (late anaphase/telophase); and part IV, from furrow completion to cell division (cytokinesis). We used time-lapse video microscopy to measure the duration of each of parts I–IV in control and *kDKO*(P1) keratinocytes. Part IV was significantly accelerated in *MOB1*-deficient cells compared with controls, while part I was slightly faster in the mutant (Supplemental Figure 8, A and B). No significant differences were detected for parts II and III. Thus, loss of *MOB1* accelerates a cell's exit from mitosis.

We noted that *kDKO*(P1) mice had an apparent increase in the number of skin progenitors in the IFE (Figure 3D). We therefore performed RT-PCR to evaluate stem/progenitor cells in the HFs.

$CD34^+$ and $LGR6^+$ cells are the most immature HF stem cells located in the bulge and isthmus. We found that *Cd34* and *Lgr6* mRNAs were normal or reduced in *kDKO*(P1) keratinocytes, but that markers of activated stem/progenitor cells, such as *Sox9* and *Lgr5*, were elevated (Figure 4H). These findings were confirmed by FACS analysis and immunohistochemistry (Figure 4H). However, despite their normal numbers, $CD34^+$ cells in mutant HFs could not localize to the proper bulge area (Supplemental Figure 9). This result mirrors the failure of HF stem cells to localize to the basal cell layer in *MOB1*-deficient IFE and HFs (Figure 3D). In addition, $SOX9^+$ cells (which are normally located in the bulge at P21) were markedly increased among *kDKO*(P1) keratinocytes and scattered throughout the HFs (Figure 4H). To determine how *MOB1* inactivation affected HF stem cell self renewal, we quantified the ability of the total *kDKO*(P1) keratinocyte population to form colonies in culture. A lack of *MOB1* induced a 2.1-fold increase in colony-forming efficiency (Figure 4I). When these primary colonies were replated to test their ability to form secondary colonies, a 3.0-fold increase in secondary colony-forming efficiency was observed in the absence of *MOB1* (Figure 4I). Thus, *MOB1* inactivation enhances the self renewal of activated stem/progenitor keratinocytes. This property, along with hyperproliferation, apoptotic resistance, impaired contact inhibition, increased centrosome number, and accelerated mitotic exit, may account for the frequent cancers observed in *Mob1*-deficient mice.

MOB1-mediated regulation of the LATS1/2-YAP1 pathway controls skin homeostasis. To characterize the mechanisms driving the *kDKO*(P1) phenotype, we investigated the effects of *MOB1* loss on signaling mediated by YAP1 and its upstream kinases LATS1/2. In vitro, *MOB1* is important for the full activation of LATS kinases (37), and activated LATS1/2 phosphorylate YAP1 and inhibit its nuclear localization (38). Using immunohistochemistry, we found that, in control mice, YAP1 was localized in the nuclei of most outer root sheath cells, in some HF matrix cells, and in some IFE basal cells (Figure 5A). However, *kDKO*(P1) mice exhibited not only greater numbers of nuclear YAP1⁺ cells in these sites but also some nuclear YAP1⁺ cells in the upper IFE layer (Figure 5A). Immunoblotting of keratinocytes confirmed that the phosphorylation of YAP1 on Ser127 was reduced, but that total YAP1 was increased in the absence of *MOB1* (Figure 5B), especially in the nucleus (Figure 5C). When we cultured keratinocytes under high cell density conditions, YAP1 was inactivated and localized in the cytoplasm in control cells (Figure 5D). In contrast, in *kDKO*(P1) keratinocytes, YAP1 remained activated and positioned in the nucleus even at high-cell density (Figure 5D). Thus, *MOB1* is required for density-induced subcellular localization of YAP1.

We next investigated LATS1/2 phosphorylation in control and mutant keratinocytes. Immunoblotting demonstrated that, in the absence of any stimulation, levels of phospho-LATS1(T1079) and phospho-LATS2(T1041) were below the assay detection limit even in control keratinocytes (Figure 5E). We then stimulated control and *kDKO*(P1) keratinocytes with okadaic acid (OA), which activates the MST kinases acting upstream of LATS1/2 (39). While OA-stimulated control keratinocytes showed vigorous phosphorylation of LATS1/2, OA-stimulated *kDKO*(P1) keratinocytes did not. Strikingly, total LATS1/2 protein levels were also reduced in the mutant. This inhibition of LATS1/2 phosphorylation was not due solely to the reduction in total LATS1/2 proteins because levels of phospho-LATS1/2 were also markedly decreased when total LATS1/2 protein levels in each sample were adjusted to equality prior to electropho-

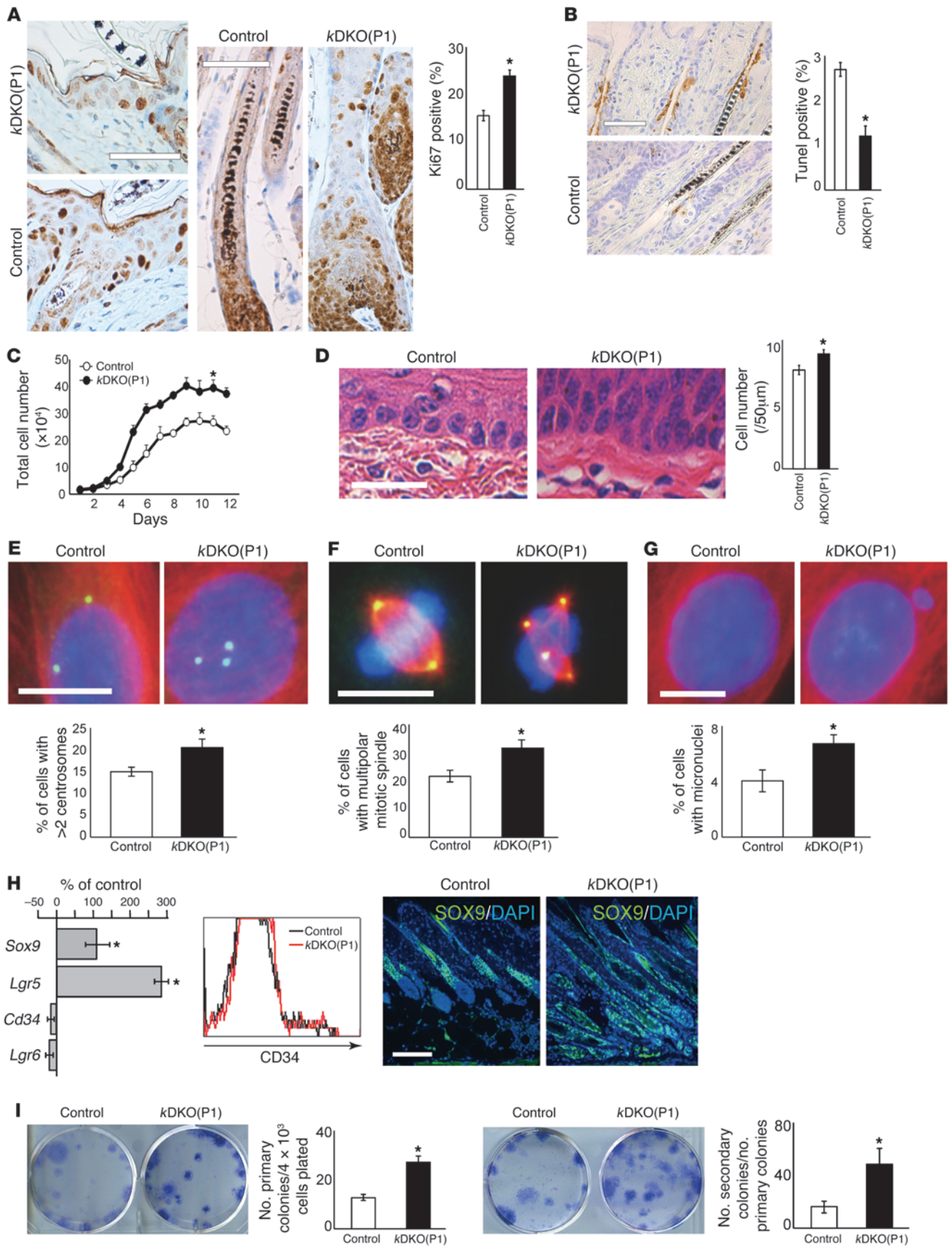




Figure 4

Tumorigenic anomalies in *Mob1a/1b* double-homozygous mutant keratinocytes. (A) Anti-Ki67 immunostaining of IFE (left) and HF (middle) of control and *kDKO*(P1) mice at P13. Scale bars: 50 μ m. Quantitation of Ki67⁺ cells (right); **P* < 0.01. (B) Histology (left) and quantitation (right) of TUNEL-stained cells in epidermis from control and *kDKO*(P1) mice at P16. Scale bar: 50 μ m; **P* < 0.01. (C) Keratinocytes from control and *kDKO*(P1) mice at P4 were cultured for the indicated number of days, and total cell numbers were counted. *kDKO*(P1) keratinocytes achieved higher saturation plating density; **P* < 0.01. (D) Left: H&E-stained epidermal basal layer of control and *kDKO*(P1) mice at P19. Scale bar: 20 μ m. Right: quantitation (cell number/50 μ m BM); *n* = 5/group; **P* < 0.05. (E–G) Immunostaining to detect γ -Tubulin (green) and α -Tubulin (red) in control and *kDKO*(P1) keratinocytes. DAPI, nuclei. Mutant keratinocytes showed excess centrosomes (E), multi-polar spindles (F), and micronuclei (G). Scale bars: 20 μ m; **P* < 0.05. (H) Identification of keratinocyte stem cells in HFs of control and *kDKO*(P1) mice at P19 (*n* = 4/group) using quantitative RT-PCR (left), **P* < 0.01; flow cytometry (middle) to detect CD34; immunostaining (right) to detect SOX9. Scale bar: 100 μ m. (I) Freshly isolated control and *kDKO*(P1) keratinocytes were plated to generate primary colonies (left) and secondary colonies (right). Giemsa staining (top) and colony counts (bottom) were performed on day 14 after plating; *n* = 4/group; **P* < 0.02. Results shown are representative of at least 3 independent trials and at least 3 mice/group. Data are presented as the mean \pm SEM, and *P* values were determined using the 2-tailed Student's *t* test.

resis (Figure 5E). These data show that MOB1 regulates LATS1/2-YAP1 signaling essential for mammalian skin homeostasis.

MOB1 is involved in the onset of malignant outer root sheath tumors resembling trichilemmal carcinomas. Microscopic analysis of the skin tumors of *Mob1a^{Δ/Δ}1b^{+/+}* and *Mob1a^{Δ/+}1b^{tr/tr}* mice revealed lesions consisting of invasive epithelial lobules with atypical cytology, an increased number of mitotic cells, and horn cysts (Figure 6A). Trichilemmal keratinization and continuity with the follicular epithelium (Figure 6A) were focally evident. Because these features are characteristic of trichilemmal carcinomas of outer root sheath origin, we immunostained these tumors with Abs recognizing KRT17, KRT5, KRT15, KRT10, Trichohyalin, or AE13. All tumors were positive for KRT17 (outer root sheath marker), KRT5 (basal cells and outer root sheath), and KRT15 (basal cells, outer root sheath, and bulge), but negative for KRT10 (upper basal IFE) and Trichohyalin (inner root sheath) (Figure 6B). Some tumors were also positive for AE13 (hair shaft marker), confirming that these malignancies were derived from the HF outer root sheath. Thus, deficiency for the Hippo signaling element MOB1 in mice leads to tumors with the features of trichilemmal carcinomas.

In humans, the most common HF-derived carcinomas are nodular BCCs, which are driven by hyperactivated SHH signaling (5). However, immunoblotting of our MOB1-deficient keratinocytes showed that neither GLI2 (downstream effector in the SHH pathway) nor the LEF1 or HES1 proteins important for HF morphogenesis were elevated (Figure 6C). In addition, while AKT was not activated in the mutant keratinocytes, phospho-ERK was increased (Figure 6C). To rule out the involvement of the SHH pathway in tumorigenesis linked to MOB1 deficiency, we immunostained control and mutant keratinocytes with Abs validated to detect GLI2 or GLI1 (Supplemental Figure 10, A and B) in patterns consistent with a previous report (40). However, when we immunostained control and *kDKO*(P1) keratinocytes with these Abs, no differences were detected (Figure 6D and Supplemental Figure 10C). Furthermore, GLI2 levels in 4 trichilemmal carcino-

ma samples from *kDKO* (P1) mice were not obviously increased compared with GLI2 levels in cultured control keratinocytes (Figure 6E). Thus, the SHH pathway involved in nodular BCCs does not appear to be driving the development of the trichilemmal carcinomas arising in the absence of MOB1A/1B.

In humans, trichilemmal carcinomas are very rare and composed mainly of basaloid and squamous cells, with some clear cells (Figure 6F). We performed immunostaining to detect YAP1 in samples of 14 human trichilemmal carcinomas with accompanying noncancerous epidermis. We found that, in noncancerous skin, YAP1 was expressed in the nuclei of IFE basal cells and HF outer root sheath cells (Figure 6G). Strikingly, a significant proportion (11/14, 79%) of trichilemmal carcinomas showed an increase in total YAP1⁺ cells (Figure 6G and Table 3). Almost as many (10/14, 71%) trichilemmal carcinomas exhibited elevated nuclear YAP1 within the tumor mass (Figure 6G and Table 3). In addition, when immunostained to detect MOB1A/1B, half of trichilemmal carcinomas examined (5/10, 50%) showed decreased MOB1A/1B expression (Figure 6G and Table 3). Moreover, 10/10 (100%) of these cases in which MOB1 was decreased displayed elevated nuclear YAP1 within the tumor. These data support our hypothesis that it is the mammalian Hippo pathway, rather than the SHH pathway, that is involved in the onset of non-BCC follicular cancers such as trichilemmal carcinomas.

Discussion

We have demonstrated that mice completely deficient for *Mob1* have the most severe phenotype among strains lacking a Hippo signaling component. *Mob1a/1b* double-homozygous-null mutant mice die at gastrulation, much earlier than mice lacking *Mst1/2* (15), *Lats2* (21), *Lats1* (14), *Sav1* (20), or *Nf2* (19). Our data further indicate that MOB1 is essential for embryogenesis and that functions of MOB1A and MOB1B overlap. In addition, the tumor spectrum observed in heterozygous *Mob1*-deficient mice is the broadest among mutants lacking Hippo components. These findings suggest both that MOB1 is the key molecule in the Hippo signaling pathway and that MOB1 may have molecular target(s) other than the Hippo pathway. This notion is consistent with the reduced expression or mutation of MOB1 frequently observed in a variety of human cancers (17, 33, 34), and with MOB1's reported binding to a range of molecules, including TSSC1, NUP98, HDAC3, and DIPA (CCDD85B) (41, 42). Our results also imply that MOB1A may be more important than MOB1B, at least for embryogenesis and liver homeostasis, because *Mob1b* heterozygotes lacking *Mob1a* show partial embryonic lethality (Supplemental Table 1) and develop liver cancers (Table 2), whereas *Mob1a* heterozygotes lacking *Mob1b* are all viable and free of liver tumors.

The lethality of *Mob1*-deficient mice may stem from their failure to form primitive endoderm. The endodermal markers *Pdgfra* and *Gata4* were markedly reduced in *Mob1*-deficient cells, whereas the primitive ectoderm markers *Nanog*, *Sox2*, *Fgf5*, and *Pax6* were normal. Our mutant showed abnormal YAP1 activation, and activated YAP1 normally activates the transcription factor TEAD2. TEAD2 regulates primitive endoderm-specific genes such that sustained inhibition of TEAD2 enhances primitive endoderm-specific gene expression (43). YAP1 and TEADS reportedly increase the expression of pluripotency genes such as *Oct3/4* and *Sox2* (43, 44), as well as the trophoblast gene *Cdx2* (45, 46). However, levels of *Oct3/4*, *SOX2*, and *CDX2* proteins were normal in our *Mob1*-deficient embryos. Thus, although we found YAP1 to be activated in the

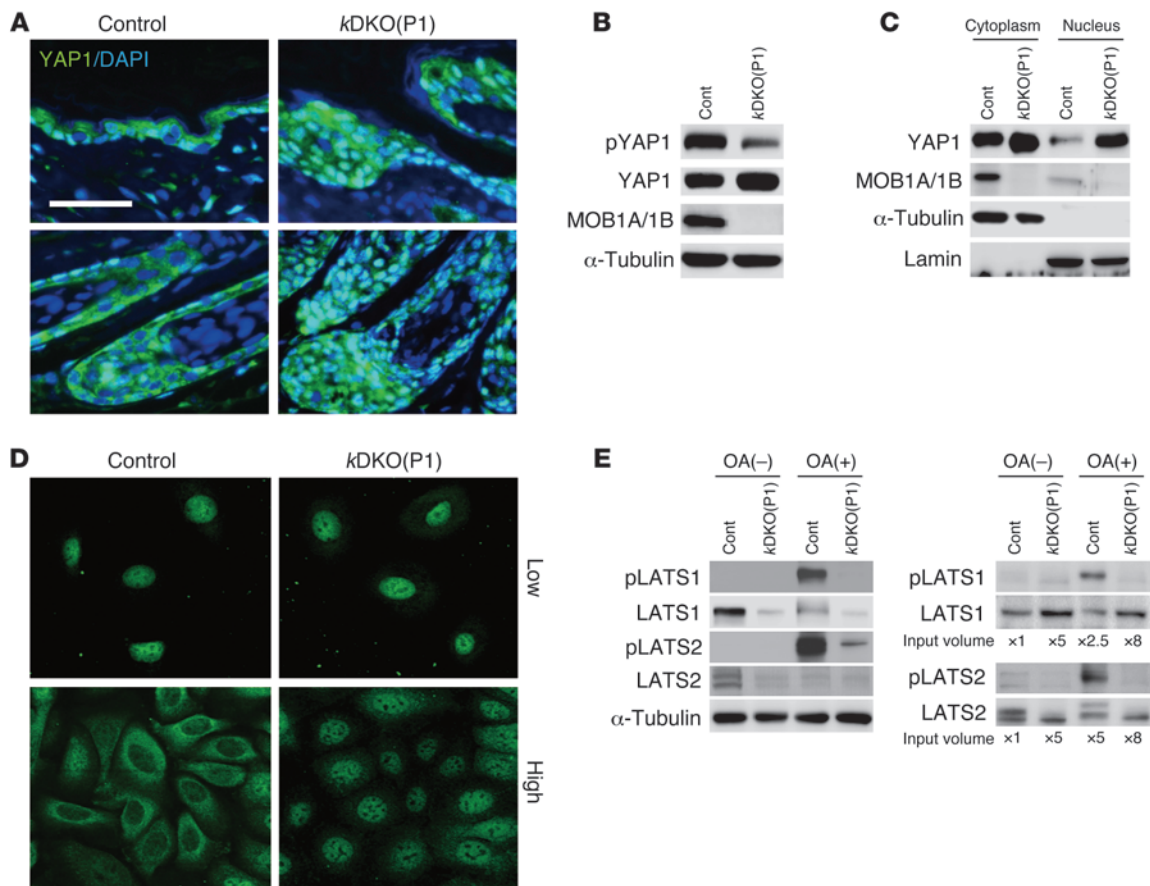


Figure 5 MOB1-mediated regulation of the LATS1/2-YAP1 pathway controls skin homeostasis. **(A)** Immunostaining of keratinocytes from control and *kDKO*(P1) mice at P19 to detect YAP1 in the IFE (top) and HF (bottom). Scale bar: 50 μ m. **(B)** Immunoblot of total extracts of control and *kDKO*(P1) keratinocytes to detect the indicated proteins. α -Tubulin, loading control. **(C)** Immunoblot of cytoplasmic and nuclear fractions of control and *kDKO*(P1) keratinocytes to detect the indicated proteins. α -Tubulin and Lamin, cytoplasmic and nuclear loading controls, respectively. **(D)** Immunostaining to detect YAP1 in keratinocytes from control and *kDKO*(P1) mice plated at low or high cell density. YAP1 is localized in the nucleus in mutant keratinocytes even at high cell density. **(E)** Immunoblot to detect the indicated proteins in total extracts of control (Cont) and *kDKO*(P1) keratinocytes that were left untreated (OA⁻) or treated with OA (OA⁺). Left: unadjusted lysates. Right: levels of LATS1 and LATS2 proteins in each sample were adjusted to equality before electrophoresis. Results shown are representative of at least 3 independent trials and at least 3 mice/group.

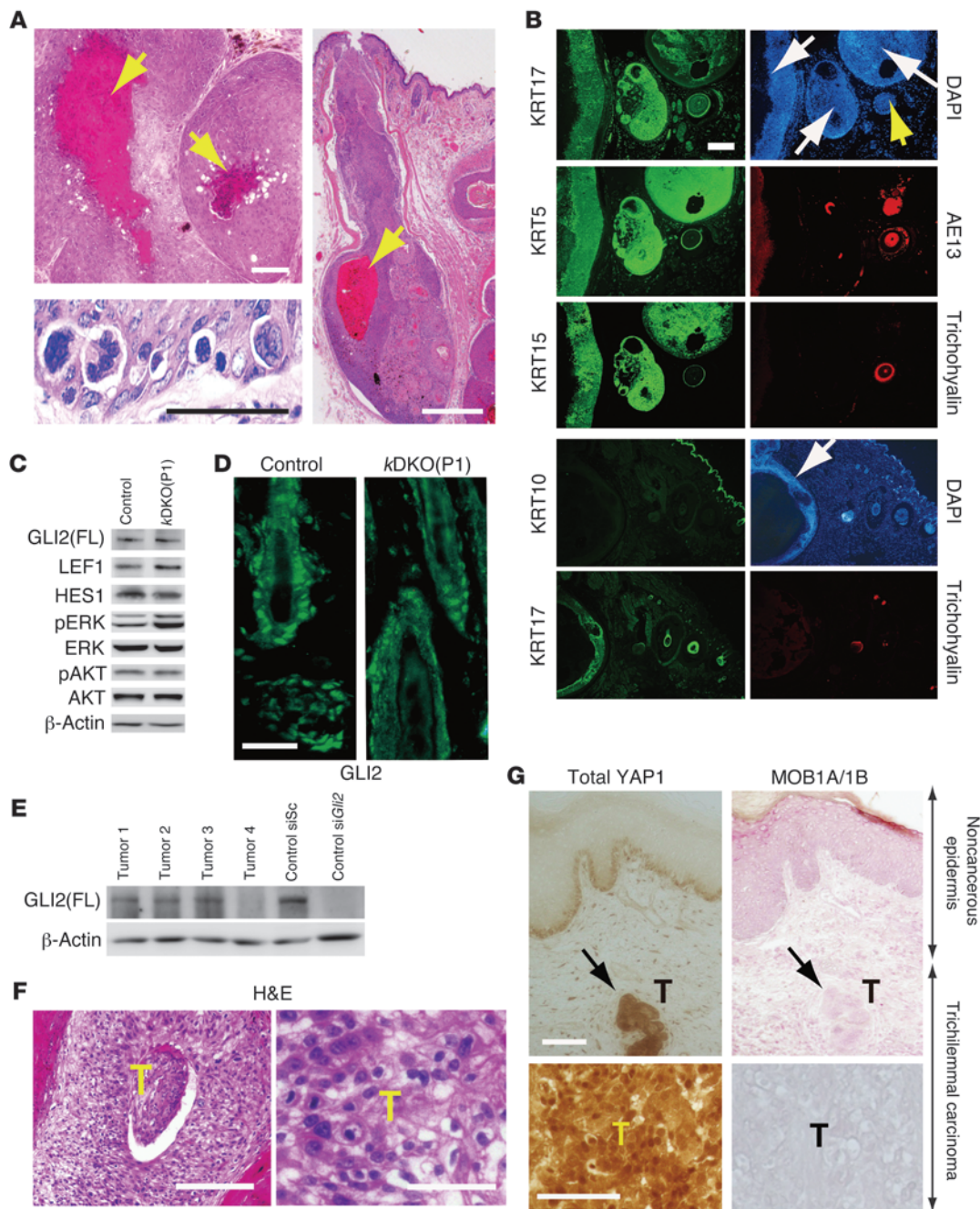
mutant ICM, a much stronger alteration of YAP1 activity may be necessary to induce a detectable effect on these genes.

We have shown that, in contrast to *MST1/2* and *LATS1/2*, *MOB1* is a potent tumor suppressor in a range of tissue types that includes the skin. All cancers examined in *Mob1a^{Δ/Δ}1b^{tr/tr}* and *Mob1a^{Δ/Δ}1b^{tr/+}* mice had lost the WT *Mob1* allele, suggesting increased genetic instability. *Mob1a/1b* mRNAs and proteins are frequently cosuppressed in tumor cell lines (data not shown), and the *MOB1A* and *MOB1B* amino acid sequences are 95% identical. We are currently clarifying whether a common mechanism regulates these 2 genes.

In our *Mob1a^{Δ/Δ}1b^{tr/tr}* and *Mob1a^{Δ/Δ}1b^{tr/+}* mice, the most frequent tumors were malignant outer root sheath tumors resembling trichilemmal carcinomas (Figure 6A). Histologically, these malignancies were not BCCs because they lacked the cellular palisading typical of BCCs (4). Moreover, cultured *Mob1a/1b* double-mutant keratinocytes did not show the SHH pathway activation important for BCC onset (Figure 6C). Benign trichilemmomas are frequently observed in Cowden disease patients with hereditary *PTEN* muta-

tions (47), but these tumors seldom become malignant. The trichilemmal growths in our *Mob1*-deficient mutants were clearly cancerous but showed no activation of the *PTEN* effector *AKT* (Figure 6C). Importantly, like our mutant mouse tissues, our human trichilemmal carcinoma samples exhibited frequent *MOB1A/1B* inactivation and YAP1 activation (Figure 6G and Table 3). These findings suggest that impaired Hippo signaling may drive trichilemmal carcinoma onset in humans.

Mob1-deficient keratinocytes exhibit enhanced proliferation, apoptotic resistance, impaired contact inhibition, increased centrosomes, accelerated mitotic exit, and enhanced progenitor self-renewal. In addition, polarity must be defective without *MOB1* because (a) *KRT15⁺* cells were scattered inside the IFE and not localized to the basal layer; (b) *CD34⁺* bulge stem cells were not localized in the bulge; and (c) hair bundles in the organ of Corti were disorganized (Supplemental Figure 2B). To date, 2 transgenic mouse strains overexpressing *Yap1* in the skin have been described (23, 48). Like our *Mob1* mutants, *Yap1* transgenic mice show hyperplastic IFE. However, these latter animals also have a severe defect in HF

**Figure 6**

Characterization of skin cancers of *Mob1*-deficient mice and human trichilemmal carcinomas. (A) H&E-stained sections of representative tumors from *Mob1a*^{NΔ1b}^{tr/+} or *Mob1a*^{N+1b}^{tr/tr} mice showing: (top left) characteristic trichilemmal keratinization (yellow arrows; also in right); (bottom left) atypical and highly mitotic cells; and (right) continuity with the epidermis. Scale bars: 50 μm (left); 500 μm (right). (B) Immunostaining of tumors from *Mob1a*^{NΔ1b}^{tr/+} or *Mob1a*^{N+1b}^{tr/tr} mice to detect the indicated skin markers. White arrows, cancerous lesions; yellow arrow, nontumorous HF. Scale bar: 200 μm. (C) Immunoblot of total extracts of control and kDKO(P1) keratinocytes to detect ERK and AKT activation as well as the HF morphogenesis proteins GLI2 (FL; full length), LEF1 and HES1. β-Actin, loading control. (D) Immunostaining to detect GLI2 in HFs from control or kDKO(P1) mice. Scale bar: 25 μm. (E) Immunoblot to detect GLI2 in total extracts of control keratinocytes transfected with scramble siRNA (Control siSc) or *Gli2* siRNA (Control si*Gli2*), as well as in samples of 4 trichilemmal carcinomas (Tumor 1, 2, 3, 4) from kDKO(P1) mice. (F) H&E staining of a human trichilemmal carcinoma viewed at low (scale bar: 200 μm) or high magnification (scale bar: 50 μm). (G) Immunostaining to detect YAP1 and MOB1A/1B in human trichilemmal carcinomas and nearby noncancerous tissues. Top: Both noncancerous and trichilemmal carcinoma (T) tissues can be seen in low-magnification images. Bottom: High-magnification images of human trichilemmal carcinomas. Scale bars: 100 μm. Results shown are representative of at least 3 independent trials and at least 3 mice/group.

**Table 3**

Quantitation of human trichilemmal carcinoma samples showing an increase in YAP1⁺ cells and/or elevation in nuclear YAP1, and a decrease in MOB1A/1B

	Increased YAP1	Nuclear accumulation of YAP1	Decreased MOB1A/1B
Trichilemmal carcinoma	11/14 (79%)	10/14 (71%)	5/10 (50%)

formation. This difference may account for the development of IFE-derived squamous cell carcinomas in *Yap1* transgenic mice, but HF-derived trichilemmal carcinomas in *Mob1*-deficient mutants.

At the biochemical level, *Mob1*-deficient keratinocytes exhibited reductions in not only phospho-LATS1/2, but also total LATS1/2 proteins. *Lats1/2* mRNAs were comparable in control and mutant keratinocytes (data not shown), indicating that this surprising decrease in LATS1/2 proteins occurs posttranscriptionally. Like MOB1, SAV1 is a scaffolding protein in the Hippo pathway, but unlike *Mob1* deficiency, *Sav1* deficiency decreases phospho-MST, but does not affect total MST or LATS1/2 proteins (20). Another interesting biochemical observation was the activation of ERK in our *Mob1*-deficient keratinocytes: no connection between MOB1 and ERK has been reported to date. With respect to Hippo signaling, ERK can be activated by YAP1 (49) or suppressed by NF2 (50) or MST2 (51). Since NF2 and phospho-MST1/2 were not decreased in our *Mob1*-deficient keratinocytes (data not shown), we speculate that the increased YAP1 in these cells may have triggered their abnormal ERK activation.

Because all of our *Mob1a/1b*-deficient mice spontaneously developed tumors (and especially trichilemmal carcinomas), we believe that the loss of MOB1A/1B helps to both initiate and promote carcinoma onset. With respect to tumor initiation, the observed increase in the number of centrosomes and/or micronuclei in our mutant cells may have introduced detrimental alterations into their DNA. With respect to cancer promotion, the enhanced proliferation, apoptotic resistance, impaired contact inhibition, and increased progenitor self renewal associated with loss of MOB1A/1B may support the growth and progression of cells that have undergone tumor initiation events.

The very rare occurrence of trichilemmal carcinomas in humans has slowed the identification of genes involved in their development. Expression levels of MOB1A, LATS1/2, and SAV1 are more frequently reduced in the advanced stages of colon (33), breast (52), and renal cancers (53) than in the early stages of these malignancies. In lung cancers, the reverse is true, since MOB1A levels are frequently lower in the early pT1 stage of non-small-cell lung cancer as compared with later stages (34). These observations suggest that the loss of Hippo signaling molecules can be an important driver of cancer progression in humans. Further study of alterations to gene or protein expression or functions of Hippo signaling components in a broad range of human malignancies may increase our understanding of their involvement in tumorigenesis.

In conclusion, our results demonstrate that (a) MOB1 is a broadly-acting tumor suppressor in mice and (b) Hippo signaling drives trichilemmal carcinoma onset in the skin. Therapeutic strategies to control Hippo signaling or MOB1 expression might therefore benefit many cancer patients, particularly those with HF-derived cancers.

Methods

Generation of *Mob1a^{lox}* and *Mob1b^{trap}* mice. A conditional targeting vector based on the Cre-loxP system was constructed to delete a genomic fragment containing exon 2 of the murine *Mob1a* gene (Supplemental Figure 1). This deletion generates a frame shift and early stop codon in exon 3. We introduced 1 loxP site into *Mob1a* intron 1 and 2 loxP sites into intron 2 to flank *Mob1a* exon 2. The PGK-Hyg resistance cassette was inserted in antisense orientation between the 2 loxP sites in intron 2. The resulting

targeting vector was electroporated into E14K ES cells, and homologous recombinants were confirmed by Southern blotting using 5' flanking probe A and a Hyg-specific probe. Correctly targeted *Mob1a* mutant clones were transiently transfected with pMCI-Cre (54) to delete the loxP-flanked Hyg^r gene. Progeny clones sensitive to hygromycin were subjected to Southern blotting to identify those retaining exon 2 flanked by 2 loxP sites (the *Mob1a^{lox}* allele) and those lacking exon 2 (*Mob1a^Δ* allele; equivalent to a knockout mutation). For *Mob1b*-deficient mice, an ES cell line bearing a gene trap mutation of intron 2 of the *Mob1b* gene was obtained from the Sanger Institute (Hinxton, United Kingdom). The single integration of the trapped site was confirmed by PCR and genomic Southern blotting using 5' flanking probe C and a neospecific probe. *Mob1a^{lox}*, *Mob1a^Δ*, and *Mob1b^{trap}* mice were generated from ES cell clones using standard procedures and backcrossed to C57BL/6 mice 3 times before intercrossing to generate *Mob1a^{β/β}*, *Mob1a^{Δ/Δ}*, and *Mob1b^{tr/tr}* progeny. Primers used for genotyping are listed in Supplemental Table 2.

In vitro culture of preimplantation embryos. E3.5 embryos derived from the intercrossing of *Mob1a^{Δ/Δ}1b^{tr/tr}* males and females were analyzed as described (55). Briefly, blastocysts were individually cultured in 24-well plates in ES cell medium without LIF and photographed every 24 hours. On day 8, the morphology of each embryo was recorded and its genotype determined by PCR.

Embryo immunostaining. Immunofluorescent staining of embryos was performed as described (45). Briefly, embryos (E3.5) were cultured overnight prior to fixation in 4% paraformaldehyde. Fixed tissues were permeabilized with 0.2% Triton X-100, blocked with 2% goat serum, and incubated overnight at 4°C with Abs recognizing CDX2 (Bioscience Medical), Oct3/4 (gift of H. Niwa, Riken), PDGFRα (eBiosciences), GATA4/6 (Santa Cruz Biotechnology Inc.), NANOG (ReproCELL), Troma-1 (DSHB), SOX2 (Abcam), or YAP1 (Sigma-Aldrich). Secondary Abs were Alexa Fluor 488 goat anti-rabbit or Alexa Fluor 568 goat anti-mouse (Molecular Probes). Nuclei were visualized using DAPI (Dojindo). Stained embryos were examined by confocal microscopy.

Generation and differentiation of *Mob1a/1b* double-mutant ES cells. ES cells established from embryos derived from *Mob1a^{β/β}1b^{tr/tr}* intercrosses were transfected with linearized CMV-Mer-Cre-Mer plasmid DNA (56) plus pApuro2 (57). To generate ERCCre*Mob1a^{β/β}1b^{tr/tr}* ES clones, ES cells were selected by culture for 12 days in complete ES cell medium containing 1 μg/ml puromycin (Invivogen) on an irradiation-inactivated Puro MEF feeder layer. Tamoxifen (3.3 μM; Toronto Research Chemicals) was added (or not) for 2 days to completely delete the *Mob1a* gene (data not shown). Control and mutant EBs were generated from hanging drops of approximately 1,000 ES cells/20 μl medium as described (58).

Generation of keratinocyte-specific *Mob1a/Mob1b* double-mutant mice. *Mob1a^{β/β}1b^{tr/tr}* mice were mated to *Krt14CreER* transgenic mice (35) in which Cre can be activated by tamoxifen under the control of the *Krt14* promoter. Offspring carrying *Krt14CreER* plus 2 copies of the floxed *Mob1a* allele (*Mob1a^{β/β}*) and 2 copies of the trapped *Mob1b* allele (*Mob1b^{tr/tr}*) were administered tamoxifen and used in the analysis as homozygous double mutant (*Krt14CreERMob1a^{β/β}1b^{tr/tr}* +tamoxifen; kDKO) mice.



Krt14CreERMob1a^{β/β1b^{tr/tr}} mice without tamoxifen and *Mob1a^{+/-1b^{tr/tr}}* or *Mob1a^{+/-Mob1b^{tr/tr}}* mice with tamoxifen but without *Krt14CreER* were indistinguishable in pilot experiments examining histology and MOB1 protein levels. *Krt14CreERMob1a^{β/β1b^{tr/tr}}* mice without tamoxifen were arbitrarily chosen to serve as controls. To induce Cre in keratinocyte stem/progenitor cells, P1 mice were intraperitoneally administered 0.2 mg tamoxifen on that one day, or P28 mice received 0.5 mg tamoxifen each day for 7 days starting on P28.

Characterization of primary keratinocytes. Primary keratinocytes isolated from control and *kDKO* mice were cultured as described (59). For cell saturation density assays, keratinocytes at passage 2 were seeded in type I collagen-coated 24-well plates (0.3×10^5 /well) and cultured for up to 12 days before counting cell numbers. For colony-forming assays, keratinocytes at passage 1 were seeded in 6-well plates (0.4×10^4 /well). After 2 weeks, half the wells were fixed and stained with Giemsa (Muto Pure Chemicals), and the number of primary colonies was counted. Keratinocytes of an unfixed well were then trypsinized and reseeded so that each well of a 6-well plate received cells from 1 primary colony. Reseeded cells were cultured for 2 weeks to generate secondary colonies.

Clinical samples. Surgically obtained trichilemmal carcinoma samples containing noncancerous tissues were acquired from the Department of Regenerative Dermatology (Osaka University). Resected cancer tissues were fixed in formalin and stained with anti-YAP1 Ab (Cell Signaling) and anti-MOB1A/1B Ab (AP7031b; Abgent) using a standard protocol.

Statistics. KaleidaGraph software was used for statistical analyses. Data are shown as the mean \pm SEM, and *P* values were determined using the 2-tailed Student's *t* test unless otherwise stated. *P* < 0.05 was considered statistically significant.

Study approval. The clinical sample study design was approved by the Institutional Review Board of Osaka University, and written informed consent was

obtained from all patients. All animal experiments were approved by the Animal Experiment Review Boards of Kyushu University and Akita University.

See Supplemental Methods for additional details.

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