# **Cell Reports**

# Human Pluripotent Stem Cell-Derived Tumor Model Uncovers the Embryonic Stem Cell Signature as a Key Driver in Atypical Teratoid/Rhabdoid Tumor

### **Graphical Abstract**



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### In Brief

Terada et al. present *SMARCB1*-deficient human pluripotent stem cell-derived atypical teratoid/rhabdoid tumor (AT/RT) models and show that ESC-like signature is a critical driver of malignant phenotypes of AT/RT. Genetic ablation targeting the maintenance of pluripotency inhibits AT/RT cell growth, suggesting that the ESC-like signature could be a promising therapeutic target for AT/RT.

## **Highlights**

- SMARCB1-deficient human iPSCs give rise to AT/RT-like tumors
- ESC-like signature induces rhabdoid histology and causes a poor prognosis
- AT/RT exhibits ESC-like transcriptional signature and DNA methylation landscape
- ESC-like signature could be a therapeutic target for AT/RT





# Human Pluripotent Stem Cell-Derived Tumor Model Uncovers the Embryonic Stem Cell Signature as a Key Driver in Atypical Teratoid/Rhabdoid Tumor

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#### SUMMARY

Atypical teratoid/rhabdoid tumor (AT/RT), which harbors SMARCB1 mutation and exhibits a characteristic histology of rhabdoid cells, has a poor prognosis because of the lack of effective treatments. Here, we establish human SMARCB1-deficient pluripotent stem cells (hPSCs). SMARCB1-deficient hPSCderived neural progenitor-like cells (NPLCs) efficiently give rise to brain tumors when transplanted into the mouse brain. Notably, activation of an embryonic stem cell (ESC)-like signature confers a rhabdoid histology in SMARCB1-deficient NPLC-derived tumors and causes a poor prognosis. Consistently, we find the activation of the ESC-like gene expression signature and an ESC-like DNA methylation landscape in clinical specimens of AT/RT. Finally, we identify candidate genes that maintain the activation of the ESC-like signature and the growth of AT/RT cells. Collectively, SMARCB1-deficient hPSCs offer the human models for AT/RT, which uncover the role of the activated ESC-like signature in the poor prognosis and unique histology of AT/RT.

#### INTRODUCTION

Atypical teratoid/rhabdoid tumor (AT/RT) is an extraordinarily lethal malignant CNS tumor that occurs mainly in early childhood. The representative morphological feature of AT/RT is a population of cells with classic rhabdoid features: eccentrically located nuclei containing vesicular chromatin, prominent eosinophilic nucleoli, abundant cytoplasm with an obvious eosinophilic globular cytoplasmic inclusion, and well-defined cell borders (Louis et al., 2016). Loss of SMARCB1 (also known as INI1, SNF5, or BAF47) expression at the protein level is observed in almost all AT/RTs, and current consensus holds that immunohistochemical staining for SMARCB1 is a sensitive and specific test for the diagnosis of AT/RT (Louis et al., 2016). Several mouse models of Smarcb1 ablation have been developed to model AT/RT and rhabdoid tumor (Ng et al., 2015; Han et al., 2016). Genetic ablation of Smarcb1 at different developmental stages revealed that intracranial tumors with the rhabdoid histology develop only when Smarcb1 is deleted at embryonic day 6-10 (E6-E10), whereas ablation after birth results in lymphoma development, indicating that a restricted early developmental window is required for the initiation of intracranial rhabdoid tumors (Han et al., 2016). Furthermore, compound deletion of Smarcb1 and p53 at GFAP-expressing neuronal progenitor cells resulted in the development of AT/RT with rhabdoid histological features. Moreover, the expression profile of mouse and human intracranial rhabdoid tumors revealed the highest correlation with neural progenitors and stem cells (Han et al., 2016). Taken together, these findings suggest that AT/RT may arise from neural stem or progenitor cells.

Patients with AT/RT show a rapid clinical deterioration and extremely worse outcome than those with other CNS tumors, despite aggressive surgical and adjuvant radiochemotherapy. Retrospective and epidemiological studies have demonstrated a mean overall survival ranging from 6 to 18 months (Hilden et al., 2004; von Hoff et al., 2011; Chen et al., 2006). Although intensive multimodality regimens have improved the survival rates for AT/RT patients, AT/RT is still refractory to most treatments (Frühwald et al., 2016). Several preclinical studies attempting to identify molecular targets of AT/RT have taken place in recent years. Particularly, the inhibition of enhancer of zeste homolog 2 (EZH2) has been reported to suppress rhabdoid tumor cell growth, and a clinical trial with a specific inhibitor for EZH2 has been in progress against SMARCB1-defective tumors (Knutson et al., 2013; Alimova et al., 2013; Wilson et al., 2010). Besides expectation of these preclinical studies, the development of effective therapeutic approaches has been desired for this deadly cancer.





# Figure 1. Generation of an Atypical Teratoid/Rhabdoid Tumor Model using hPSCs Lacking SMARCB1 and TP53 (A) A schematic illustration of the establishment of hiPSCs <sup>SMARCB1-/-; TP53-/-</sup> and NPLCs <sup>SMARCB1-/-; TP53-/-</sup>. (B) Representative morphology of hiPSCs, hiPSCs <sup>TP53-/-</sup> and hiPSCs <sup>SMARCB1-/-; TP53-/-</sup>. Scale bars, 200 μm.

 (C) The lack of TP53 and SMARCB1 proteins in hiPSCs <sup>SMARCB1-/-; TP53-/-</sup> was confirmed by western blot analysis.
 (D) Immunofluorescent staining for NESTIN in hiPSCs <sup>SMARCB1-/-; TP53-/-</sup> and NPLCs <sup>SMARCB1-/-; TP53-/-</sup>. NESTIN-expressing cells emerged after neural induction.

(E) A schematic illustration of the xenograft transplantation of hiPSCs or NPLCs into the brain of immunocompromised mice. (F) MRI and representative histological images of a NPLC <sup>SMARCB1-/-; TP53-/-</sup>-derived tumor (NPLC-tumor). NPLCs <sup>SMARCB1-/-; TP53-/-</sup> give rise to brain tumors, which consist of densely packed undifferentiated small blue round cells with rosette formation. Scale bars, 500 µm (left) and 20 µm (right).

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Previous studies have shown that poorly differentiated tumors in humans often exhibit activation of an embryonic stem cell (ESC)-like gene expression signature, which is correlated with a worse prognosis in adults (Ben-Porath et al., 2008; Wong et al., 2008). Consistently, SALL4, one of the key factors in the maintenance of pluripotency, is re-expressed in a subset of hepatocellular carcinoma cells, especially in patients who have an unfavorable prognosis, suggesting that the acquisition of ESC-like features plays a role in cancer progression (Yong et al., 2013). Takahashi and Yamanaka (2006) succeeded to confer ESC properties to somatic cells upon the transient expression of four reprogramming factors. Recently, an in vivo reprogramming strategy, which enables the fate conversion of differentiated somatic cells to pluripotent stem cells (PSCs) in vivo, has been developed (Taguchi and Yamada, 2017). Notably, a premature termination of the in vivo reprogramming in mice causes the development of pediatric cancer-like tumors with activation of the ESC-like signature (Ohnishi et al., 2014). In addition, the transient expression of reprogramming factors in Kras mutant mice causes the development of alfa-fetoprotein (AFP)-producing cancers, which simultaneously express pluripotency-associated genes and exhibit activation of the ESC-like signature (Shibata et al., 2018). Collectively, activation of the ESC-like signature is involved in the development and progression of particular types of cancer.

Given that PSCs can give rise to various cell types while preserving genetic information, human PSCs (hPSCs) have provided an attractive platform for disease modeling in a genetically defined background in human cells. Indeed, previous studies tried to model human cancers by using hPSCs (Kim et al., 2013; Stricker et al., 2013; Sancho-Martinez et al., 2016), and hESCs with genetic mutation at histone H3.3 succeeded to model diffuse intrinsic pontine glioma in the proper cellular context after neural differentiation (Funato et al., 2014). Here, we established an AT/RT model using human induced pluripotent stem cells (iPSCs) lacking *SMARCB1*. The human iPSC-derived AT/RT model uncovered an unappreciated feature of AT/RT, which could be a therapeutic target.

#### RESULTS

#### Generation of an Atypical Teratoid/Rhabdoid Tumor Model using Human iPSCs Lacking SMARCB1 and TP53

A previous mouse study demonstrated that AT/RT-like tumors develop when mutations for both *Smarcb1* and *p53* are intro-

duced into neural progenitor cells. In the present study, to establish a human AT/RT model, we introduced genetic mutations at SMARCB1 and TP53 into 201B7 human iPSCs (hiPSCs) (hiPSCs SMARCB1-/-; TP53-/-) using the CRISPR/Cas9 system. We first established hiPSCs deficient of TP53 alone (hiPSCs TP53-/-), and then introduced an additional mutation at SMARCB1 to generate hiPSCs SMARCB1-/-; TP53-/- (Figures 1A-1C). Subsequently, we induced neural differentiation in hiPSCs SMARCB1-/-; TP53-/-, hiPSCs TP53-/- and control 201B7 hiPSCs (Yan et al., 2013). iPSCs with all genotypes efficiently changed their morphology into neural progenitor-like cells (NPLCs) (Figure 1D). Consistently, NANOG and OCT4 expression were reduced in NPLCs at day 14 of the neural induction (Figure S1A). In contrast, an increased expression of neural progenitor cell (NPC)-related genes, such as NESTIN, NCAM, and PAX6, was observed in these NPLCs (Figures S1A-S1C). However, the expression levels of NPC-related genes were lower in NPLCs with SMARCB1 deficiency when compared with wild-type control NPLCs (Figure S1A). Further induction of neuronal differentiation in NPLCs confirmed that neuronal differentiation is impaired in SMARCB1-deficient cells in vitro (Figure S1B). We also examined the effect of SMARCB1 deficiency on the cell growth of hPSCs and NPLCs in vitro. The TP53 ablation promoted cell proliferation in both hPSCs and NPLCs, whereas the SMARCB1 ablation inhibited the cell growth of hiPSCs (Figure S1D). Although SMARCB1 deficiency inhibited neuronal differentiation, it did not increase cell proliferation in NPLCs in vitro (Figure S1D).

To test the effect of *SMARCB1* deficiency on the tumorforming ability of NPLCs *in vivo*, we performed xenograft transplantation of NPLCs *SMARCB1-/-*; *TP53-/-* and control NPLCs into mouse brain (Figure 1E). The orthotopic transplantation of NPLCs *SMARCB1-/-*; *TP53-/-* caused the development of aggressive tumors (n = 26/26) (Figures 1F and S1E). In contrast, control 201B7 NPLCs and NPLCs *TP53-/-* gave rise to only microscopic tumors (n = 2/2, 4/5, respectively) (Figures S1F and S1G), and no macroscopic tumor was developed after the transplantation (n = 0/4, 0/6, respectively) (Figure S1E). Together, *SMARCB1* deficiency markedly promoted tumor formation from NPLCs after orthotopic transplantation *in vivo*.

The histological analysis demonstrated that NPLC <sup>SMARCB1-/-; TP53-/-</sup>-derived tumors (hereafter NPLC-derived tumors) were medulloblastoma-like or embryonal tumor with multilayered rosettes (ETMR)-like tumors, which mainly consisted of densely packed undifferentiated small, blue, round

<sup>(</sup>G) Percentage of tumors with rhabdoid cells. hiPSC-tumors frequently contained rhabdoid cells, whereas a majority of NPLC-tumors did not. \*\*p < 0.01 (Fisher's exact test).

<sup>(</sup>H) Quantification of the rhabdoid area within a tumor. Note that the rhabdoid area is significantly larger in hiPSC-tumors than NPLC-tumors. Data are represented as the median with interquartile range. \*\*p < 0.01 (Mann-Whitney U test).

<sup>(</sup>I) MRI and representative histological images of a hiPSC <sup>SMARCB1-/-; TP53-/-</sup>-derived tumor (hiPSC-tumor). Note that the hiPSC-tumor contains a number of rhabdoid cells. Scale bars, 500 µm (left) and 20 µm (right).

<sup>(</sup>J) Immunostaining for Ki67, SMARCB1, GFAP, vimentin, synaptophysin, S100, SMA, CD99, and EMA in a hiPSC-tumor. Tumor cells lack staining for SMARCB1. The majority of hiPSC-tumor cells express Ki67, vimentin, and CD99, while a subset of tumor cells are positive for GFAP, synaptophysin, S100, SMA, and EMA. Scale bars, 500 µm (top) and 50 µm (bottom). Areas in the high-magnification images are shown in the low-magnification images.

<sup>(</sup>K) Clustering analysis using all probes except for those of lowly expressed genes in a microarray analysis. Global expression profiles in tumors from *SMARCB1*defcient hiPSCs and NPLCs exhibited similar expression patterns with an AT/RT cell line. Note that medulloblastoma cell lines and a glioblastoma cell line are independently clustered. Microarray data were obtained from GSE45265, GSE36947 (medulloblastomas), and GSE26313 (glioblastoma). See also Figure S1.



Figure 2. hiPSC-Derived Tumors Exhibit Activation of the ESC-like Gene Expression Signature, which Drives Rhabdoid Tumors *In Vivo* (A) Clustering analysis using microarray data revealed that both a hESC-like module genes (left) (Wong et al., 2008) and ESC Core module genes (right) (Kim et al., 2010) are similarly expressed in an hiPSC-tumor, a NPLC-tumor, an AT/RT cell line, and hESC lines, whereas medulloblastoma lines and a glioblastoma cell line are clustered with NSCs. The microarray data in Figure 1K were used. Data for NSCs were obtained from GSE18296 and GSE27667.

(B) Immunohistochemical analysis of SALL4, LIN28A, and LIN28B in hiPSC-tumors and NPLC-tumors. Scale bars, 50 μm.

(C) Quantification of the SALL4-positive cells in hiPSC-tumors and NPLC-tumors. Note that hiPSC-tumors contain SALL4-positive cells more frequently than NPLC-tumors. Data are represented as the mean with 95% confidence interval. \*\*p < 0.01 (unpaired t test with Welch's correction).

(D) A qRT-PCR analysis for SALL4 and LIN28 expression in hiPSC- and NPLC-tumors. Data are presented as the mean of technical triplicates. The mean expression level of hiPSC-tumors was set to 1. \*p < 0.05 (unpaired t test with Welch's correction).

(E) GSEA showing that the ESC-like module is enriched in hiPSC-tumors compared to NPLC-tumors.

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cells with rosettes (Figure 1F). However, the majority of NPLC-derived tumors did not contain rhabdoid cells (Figures 1G and 1H), a characteristic histology of AT/RT. Although NPCs have been suggested to be a cell-of-origin for AT/RT, other studies proposed that pluripotent fetal cells, which are more immature than NPCs, could be an origin based on the morphology in an electron microscope analysis and the unique gene expression in AT/RTs (Bouffard et al., 2004; Deisch et al., 2011). Therefore, we next transplanted hiPSCs SMARCB1-/-; TP53-/- without neural induction (Figure 1E). The transplantation of hiPSCs SMARCB1-/-; TP53-/- resulted in aggressive tumor formation (n = 18/18) (Figures 11 and S1E), which was similarly observed after the transplantation of NPLCs SMARCB1-/-; TP53-/-. In contrast, the transplantation of control 201B7 hiPSCs and hiPSCs TP53-/caused only microscopic tumor formation (n = 2/5 and 1/8, respectively), and no macroscopic tumor formation was observed after the transplantation (Figures S1E and S1F). Notably, the histological analysis demonstrated that hiPSC SMARCB1-/-; TP53-/--derived tumors (hereafter hiPSC-derived tumors) frequently contained a large number of rhabdoid cells with vesicular chromatin, prominent nucleoli, and eosinophilic globular cytoplasmic inclusions, which are representative histological features of AT/RT (Figures 1G and 1I). The tumor area with rhabdoid histology was significantly larger in hiPSC-derived tumors than NPLC-derived tumors (Figure 1H). Immunohistochemical analyses revealed that hiPSC-derived tumors exhibit shared features with AT/RT, including high proliferative activity, lack of SMARCB1 expression, and positive staining for vimentin, glial fibrillary acidic protein (GFAP), synaptophysin, CD99, S-100, EMA, and smooth muscle actin (SMA) (Figure 1J). Consistent with the histological observations, clustering analysis of global gene expression revealed that a hiPSC-derived tumor exhibited a similar expression pattern with an AT/RT cell line, whereas medulloblastoma and glioblastoma cell lines fell into a different cluster (Figure 1K). To exclude the possibility of a PSC clone-specific phenotype, we also established an SMARCB1-deficient PSC line by using a different iPSC line, 1383D6 (Figure S1H). The orthotopic transplantation of both 1383D6 iPSCs SMARCB1-/- and 1383D6 NPLCs SMARCB1-/caused aggressive tumors (5/7 and 5/5, respectively) (Figure S1I), which phenocopied the tumors from 201B7 iPSCs and NPLCs lacking SMARCB1 and TP53. Notably, four out of five 1383D6 iPSC SMARCB1-/--derived tumors contained rhabdoid cells, although the area of the rhabdoid cells was smaller than in 201B7 hiPSC SMARCB1-/-; TP53-/--derived tumors (Figures S1I and S1J). In sharp contrast, no rhabdoid

cells were detected in 1383D6 NPLC <sup>SMARCB1-/-</sup>-derived tumors (Figures S1I and S1J). Collectively, we succeeded to model human AT/RT by inoculation of hiPSCs lacking *SMARCB1* into mouse brain.

# hiPSC-Derived Tumors Exhibit an ESC-like Gene Expression Signature

Our results demonstrating that iPSC-derived tumors exhibited a robust rhabdoid phenotype raised the possibility that a pluripotency-related program is associated with the characteristic rhabdoid histology. Therefore, we next investigated the expression of ESC-related modules in hiPSC- and NPLC-derived tumors together with various brain tumor cell lines. Notably, both hiPSCand NPLC-derived tumors together with an AT/RT cell line exhibited the ESC-like activation patterns of ESC-related modules, including the ESC Core module, which is composed of genes co-occupied by multiple factors in the core pluripotency network (Figure 2A) (Kim et al., 2010; Wong et al., 2008). We next analyzed the expression of SALL4, LIN28A, and LIN28B, which are associated with the maintenance and acquisition of pluripotency, in hiPSC- and NPLC-derived tumors. Immunohistochemical analvsis revealed that both hiPSC- and NPLC-derived tumors expressed SALL4, LIN28A, and LIN28B, but the expression was particularly pronounced in hiPSC-derived tumors (Figures 2B and 2C). We also confirmed the elevated expression of SALL4 and LIN28B in hiPSC-derived tumors by qRT-PCR (Figure 2D). Consistently, gene set enrichment analysis (GSEA) showed an enrichment of the ESC-like module (Wong et al., 2008) in hiPSC-derived tumors compared to NPLC-derived tumors (Figure 2E) (Subramanian et al., 2005). Together, we confirmed the ESC-like signature is activated in hiPSC-derived tumors.

# Induction of the ESC-like Signature Leads to Rhabdoid Phenotypes in NPLC-Derived Tumors

To further investigate the role of the pluripotency-related signature in the histogenesis of rhabdoid cells, we next tried to induce the ESC-like signature during tumor development from NPLCs <sup>SMARCB1-/-; TP53-/-</sup> by the forced expression of four reprogramming factors, namely, *OCT4*, *SOX2*, *KLF4*, and *c-MYC* (*OSKM*-NPLCs) (Figure S2A). It was reported that the removal of polysialylated-neural cell adhesion molecule (PSA-NCAM)-negative cells could prevent mesodermal tumor formation in hiPSC-derived NPC transplantation, indicating that PSA-NCAM antibody is useful for excluding undifferentiated, tumor-forming PSCs after neural induction (Lee et al., 2015). To eliminate the tumorigenic undifferentiated PSCs after neural induction into NPLCs <sup>SMARCB1-/-; TP53-/-</sup>, we sorted neural-differentiated cells with the PSA-NCAM antibody using

(I) Quantification of the rhabdoid area in OSKM-NPLC-tumors. Note that the rhabdoid areas in OSKM-NPLC-tumors are significantly larger than those in the control NPLC-tumors. Data are represented as the mean with 95% confidence interval. \*\*\*\*p < 0.0001 (unpaired t test with Welch's correction).

(J) Survival curve of mice transplanted with hiPSCs <sup>SMARCB1-/-; TP53-/-</sup> or NPLCs <sup>SMARCB1-/-; TP53-/-</sup>. The overall survival of hiPSC <sup>SMARCB1-/-; TP53-/-</sup>-transplanted mice is shorter than NPLC <sup>SMARCB1-/-; TP53-/-</sup>-transplanted mice. Kaplan-Meier analysis was performed. \*p < 0.05 (log-rank test).

See also Figure S2.

<sup>(</sup>F) A schematic illustration of the xenograft transplantation study. PSA-NCAM-positive NPLCs <sup>SMARCB1-/-; TP53-/-</sup> were sorted by MACS, followed by the retroviral transduction of *GFP* (*GFP*-NPLCs) or *OSKM* (*OCT4*, *SOX2*, *KLF4*, and c-*MYC*: *OSKM*-NPLCs) and orthotopic transplantation into the mouse brain. (G) GSEA showing that the ESC-like module is activated in *OSKM*-NPLC-tumors compared to control NPLC-tumors.

<sup>(</sup>H) Representative histological images of an OSKM-NPLC-tumor. A number of rhabdoid cells are observed in the tumor. Scale bars, 500 µm (top) and 20 µm (bottom).

magnetic-activated cell sorting (MACS). We confirmed the high efficiency of MACS of PSA-NCAM-positive cells by a flow cytometric analysis (Figure S2B). OCT4, SOX2, KLF4, and c-MYC were retrovirally transduced into NPLCs or PSA-NCAM-positive NPLCs (Figure S2C), and the OSKM-NPLCs were inoculated into the mouse brain to obtain NPLC-derived tumors (Figures 2F and S2A). All mice transplanted with OSKM-NPLCs developed aggressive brain tumors (n = 9). Although OSKM-NPLCs did not exhibit a prominent upregulation of pluripotency-related genes in vitro (Figure S2D), an enrichment of the ESC-like module was observed in OSKM-NPLC-derived tumors when compared to control NPLC-derived tumors (Figure 2G), suggesting that OSKM induction conferred the ESC-like gene expression signature in NPLC-derived tumors (Wong et al., 2008). Of particular note, OSKM-NPLC-derived tumors exhibited an apparent rhabdoid histology in most areas (Figure 2H). Consistently, the rhabdoid area in OSKM-NPLC-derived tumors was significantly larger than that in control NPLC-derived tumors (Figure 2I). Collectively, we concluded that activation of the ESC-like signature is responsible for the characteristic rhabdoid histology.

### Orthotopic Transplantation of hiPSCs <sup>SMARCB1-/-; TP53-/-</sup> Results in Poor Survival Compared to

### NPLC SMARCB1-/-; TP53-/- Transplantation

Previous studies demonstrated that activation of the ESC-like signature in tumors is associated with a worse prognosis of adult cancer patients (Ben-Porath et al., 2008). Therefore, we next investigated the survival period of mice after the orthotopic transplantation of hiPSCs SMARCB1-/-; TP53-/- and NPLCs SMARCB1-/-; TP53-/-. Consistent with a positive correlation between activation of the ESC-like signature and the poor prognosis, the overall survival of hiPSC SMARCB1-/-; TP53-/--transplanted mice that developed tumors with rhabdoid histology was substantially shorter than NPLC SMARCB1-/-: TP53-/--transplanted mice (Figure 2J). Similarly, mice inoculated with OSKM-NPLCs, which also developed tumors with rhabdoid histology, exhibited poor survival when compared to mice inoculated with control NPLCs (Figures S2E and S2F). Together, these results indicate that activation of the ESC-like signature is associated with a worse prognosis of SMARCB1-defecient tumors.

#### *c-MYC* Induces Activation of the ESC-like Signature in NPLC-Derived Tumors and Drives Rhabdoid Tumor Development *In Vivo*

Our results indicate that activation of the ESC-like signature is associated with the unique rhabdoid histology and poor prognosis of AT/RT. However, how AT/RT cells acquire the ESC-like signature remains unclear. A previous study demonstrated that *c-Myc* activates the ESC-like signature in adult epithelial cells and cancer cells and increases the fraction of tumorinitiating cells (Wong et al., 2008). Considering that *c-MYC* is overexpressed in a subset of AT/RTs (Johann et al., 2016), we next investigated the effect of *c-MYC* induction on the activation of the ESC-like signature and the rhabdoid phenotype in NPLC-derived tumors. *c-MYC* was retrovirally transduced in NPLCs <sup>SMARCB1-/-; TP53-/-</sup> or PSA-NCAM-positive NPLCs <sup>SMARCB1-/-; TP53-/-</sup> (MYC-NPLCs), and then MYC-NPLCs were inoculated into mouse brain (Figures 3A, S3A, and S3B). Notably, *MYC*-NPLCs gave rise to aggressive tumors containing a large number of rhabdoid cells (Figures 3B, 3C, S3C and S3D). Moreover, these tumors exhibited an activation of ESC-like module genes (Figure 3D) and increased expression of *SALL4* and *LIN28* (Figure 3E) compared to control tumors.

To investigate the effect of *c-MYC* overexpression on the disease outcome, we next investigated the survival period of mice transplanted with *MYC*-NPLCs. The overall survival of these mice was significantly shorter than that of controls (Figures 3F and S3E). Collectively, these results demonstrate that *c-MYC* overexpression alone is sufficient for activation of the ESC-like signature and rhabdoid histology in NPLC-derived tumors, which was linked to a worse prognosis.

# Activation of the ESC-like Gene Expression Signature in Human AT/RT Specimens

Above, we showed that activation of the ESC-like signature is related to the emergence of rhabdoid cells, a unique histological feature of AT/RT, and the worse prognosis of NPLC SMARCB1-/-; TP53-/--transplanted mice. We next examined the gene expression profile of clinical samples of human AT/RTs, medulloblastomas, and glioblastomas by using previously published datasets. Clustering analysis revealed that ESC-like module genes (Wong et al., 2008) are similarly activated in AT/RT samples and ESCs and iPSCs, whereas medulloblastomas and glioblastomas fall into a different cluster (Figure 4A). Similar results were obtained in the clustering analysis using the ESC Core module (Figure 4A) (Kim et al., 2010). We also examined activation of the ESC-like signature in ETMRs, which often show LIN28A immunoreactivity. Notably, some ETMRs also exhibited ESC-like activation of the ESC Core module genes (Figure S4A). Consistent with the ESC-like gene expression signature, the expression of SALL4, LIN28A, and LIN28B were elevated in AT/RTs compared to glioblastomas (Figure 4B). SALL4 was highly expressed in AT/RTs compared with medulloblastomas, but the expression level of LIN28B was higher in medulloblastomas (Figure 4B). The increased expression of SALL4, LIN28A, and LIN28B were also confirmed in AT/RT clinical specimens by immunohistochemistry (Figure S4B) (Deisch et al., 2011) (Weingart et al., 2015). Taken together, we confirmed that human AT/RTs harbor the ESC-like signature.

## ESC-like DNA Methylation Landscape in Human AT/RT Specimens

Given that human AT/RTs exhibit the ESC-like gene expression signature, we next examined DNA methylation patterns between AT/RTs and PSCs. We performed a comprehensive genomewide methylation analysis of CpG islands (CGIs) in AT/RTs together with other brain tumors by using previously published datasets. In this analysis, we first extracted CpG sites within CGIs or around transcription start sites (TSSs;  $\pm$  1,500 bp), which are differentially methylated between PSCs and adult brain tissues (DNAme difference [Dif], >0.6), and then examined DNA methylation levels in AT/RTs, medulloblastomas, ETMRs, glioblastomas, fetal brain tissues, and neural stem cells (NSCs). We found that CGI methylation levels at brain-unmethylated sites (PSC-methylated sites) are often increased in AT/RTs but



#### Figure 3. c-MYC Induces Activation of the ESC-like Signature and Drives Rhabdoid Tumors In Vivo

(A) A schematic illustration of the xenograft transplantation study. PSA-NCAM-positive NPLCs <sup>SMARCB1-/-; TP53-/-</sup> were sorted by MACS and then retrovirally transduced with *GFP* (*GFP*-NPLC) or c-*MYC* (*MYC*-NPLC), which were transplanted into the mouse brain. The control NPLCs are the same as Figure 2F.
 (B) Representative histological images of *GFP*-NPLC-tumors and *MYC*-NPLC-tumors. *MYC*-NPLC-tumors contain rhabdoid cells. Scale bars, 500 μm (top) and 20 μm (bottom).

(C) Quantification of the rhabdoid area in NPLC-tumors. Note that the rhabdoid area in MYC-NPLC-tumors is significantly larger than in control NPLC-tumors. Data are represented as the mean with 95% confidence interval. \*\*\*\*p < 0.0001 (unpaired t test with Welch's correction).

(D) GSEA showing that the ESC-like module is enriched in MYC-NPLC-tumors compared to control NPLC-tumors.

(E) Expression levels of *LIN28B* and *SALL4* in *MYC*-NPLC-tumors, control NPLC-tumors, iPSCs, and human dermal fibroblasts (HDF) by RNA sequencing (RNA-seq) analysis. Lines indicate mean expression values. RPKM, reads per kilobase of exon per million mapped sequence reads.

(F) Survival curves of mice transplanted with MYC-NPLCs or control NPLCs. The overall survival of mice inoculated with MYC-NPLCs is significantly shorter than of controls. Kaplan-Meier analysis was performed. \*\*\*\*p < 0.0001 (log-rank test).

See also Figure S3.

not so much in other tumors or NSCs (Figures 4C and S5). Although ETMRs exhibited the ESC-like gene expression signature, PSC-methylated sites remain unmethylated in ETMRs (Figure S5). In contrast, CGI methylation levels at adult brainspecifically methylated sites tended to be decreased in AT/RTs but not in other tumors, including ETMRs compared to adult



Figure 4. Human AT/RT Specimens Exhibit Activation of the ESC-like Gene Expression Signature and ESC-like DNA Methylation Landscape (A) Clustering analysis using microarray data revealed that both hESC-like module genes (left) and ESC Core module genes (right) are similarly expressed in AT/RT samples and ESCs and iPSCs. Note that medulloblastoma (MB) and glioblastoma (GBM) samples are clustered separately from ESCs and iPSCs. Microarray data of hPSCs, AT/RTs, medulloblastomas, and glioblastomas were obtained from GSE22392 (hESC/hiPSC), GSE70678 (AT/RT), GSE37418 (MB), and GSE53733 (GBM).

(B) Expressions of SALL4 and LIN28 in AT/RTs, medulloblastomas, and glioblastomas compared to ESCs and iPSCs. Data are represented as the median with interquartile range. p < 0.05, p < 0.05, p < 0.01, p < 0.001, p < 0.001 (Kruskal-Wallis test and Dunn's multiple comparisons test). The same microarray data as Figure 4A were used.

(C) The DNA methylation landscape in AT/RTs analyzed using Infinium450K data. In this analysis, we first extracted differentially methylated CpG sites within CpG islands (CGIs) between hPSCs and adult brains. hPSC-specific methylated CpG sites and adult brain-specific methylated CpG sites within CGIs were analyzed (legend continued on next page)

brain tissues or NSCs (Figures 4C and S5). Similar patterns of DNA methylation alterations were observed at CpG sites around TSSs (Figure S5). Taken together, we concluded that AT/RTs harbor an ESC-like DNA methylation landscape.

#### Pediatric Cancers Exhibit Activation of the ESC-like Gene Expression Signature

Previous studies demonstrated that pluripotency-related genes, such as LIN28B, are frequently upregulated in other pediatric cancers that occur in early childhood, which raised the possibility that activation of the ESC-like signature is a shared feature in pediatric cancers. Therefore, we next examined the expression of the ESC-like module genes in three other pediatric cancers, namely, neuroblastoma (NB), Wilms' tumor (WT), and hepatoblastoma (HB), by comparing them with their corresponding adult cancer, namely, adrenocortical carcinoma (ACC), renal cell carcinoma (RCC), and hepatocellular carcinoma (HCC), respectively. Notably, a principal-component analysis (PCA) revealed that all three pediatric cancer types exhibited similar expression profiles with PSCs and activation of the ESC-like module genes compared to the corresponding adult cancers (Figures 5A and S6). Consistently, the pediatric cancers showed increased expression of SALL4 and LIN28B compared to their counterparts (Figure 5B). Together, our data indicate that activation of the ESC-like signature is a common characteristic of pediatric cancers.

#### Genetic Screening with CRISPR/Cas9 to Develop a Strategy for AT/RT Therapy

Our findings about activation of the ESC-like signature suggest the signature could make a promising therapeutic target for AT/RT. Therefore, we performed genetic screening to identify genes that play a role in the malignant features of AT/RT (Figure 6A). Accordingly, we picked out 110 target genes that are associated with the maintenance of ESC identity based on previous reports (Tables S1 and S2). A lentiviral CRISPR/Cas9 system was employed to disrupt the candidate genes in a cancer cell line, which was established from hiPSC-derived tumors (Figure 6A; Table S1). The high efficiency of non-homologous end joining by this lentiviral CRISPR/Cas9 system was confirmed 7 days after lentiviral transduction in the pediatric cancer cell line SK-N-BE(2) (Figure S7A). The genetic screening revealed decreased cancer cell growth after the transduction of CRISPR/Cas9 together with single guide RNA (sgRNA) for dozens of genes compared to non-targeting control (NTC) sgRNA (Figure 6B; Table S3). Notably, among the candidate genes was Enhancer of zeste homolog 2 (EZH2), which has been reported as a target of AT/RT treatment (Knutson et al., 2013; Alimova et al., 2013; Wilson et al., 2010), indicating that our screening successfully identified potential therapeutic targets. Indeed, we confirmed that GSK126, a specific inhibitor for EZH2, efficiently suppressed growth of the hiPSC-derived

cancer cell line. Moreover, GSK126 efficiently inhibited the ESC-like signature in the hiPSC-derived cancer cell line (Figure S7B).

Given that other pediatric cancers similarly exhibit activation of the ESC-like signature, we also performed the same genetic screening in two neuroblastoma cell lines, namely, SK-N-AS and SK-N-BE(2). We found that *RAD21* knock out efficiently reduced cell proliferation in the hiPSC-derived cancer cell line and neuroblastoma cell lines (Figures 6B and 6C). The inhibitory effect of cell growth was confirmed in another guide RNA targeting RAD21 (Figure S7C). Notably, *RAD21* as well as *EZH2* was highly expressed in AT/RTs compared to glioblastomas (Figure 6D). Moreover, the increased expression of *RAD21* and *EZH2* was similarly observed in other pediatric cancers, including neuroblastomas, Wilms' tumors, and hepatoblastomas, when compared to the corresponding adult cancers (Figure 6E). Therefore, we focused on *RAD21* and *EZH2* and conducted further experiments.

The inhibitory effect on tumor cell growth in the hiPSC-derived cancer cell line by the knockout of *RAD21* and *EZH2* was validated in a large-scale culture (Figure 6F). Xenograft experiments revealed that the lentiviral CRISPR/Cas9-mediated knock out of *RAD21* or *EZH2* extended the overall survival of mice after the inoculation of hiPSC-derived cancer cells (Figures 6G and 6H). Therefore, we concluded that *RAD21* and *EZH2* are potential therapeutic targets for AT/RTs.

RAD21 is a key central component within the multi-protein cohesin complex. A previous study demonstrated that histone deacetylase 8 (HDAC8) functions as a deacetylase of SMC3, another component of the cohesin complex, and plays a role in recycling cohesin during cell division (Deardorff et al., 2012). Moreover, it was shown that PCI34051, a HDAC8-specific inhibitor, reduces the localized cohesin, indicating that the inhibitor impairs cohesin function. Considering that RAD21 knock out inhibited cancer cell growth in the AT/RT model, we next investigated the effect of PCI34051 on cancer cell growth in this model. PCI34051 treatment resulted in a modest reduction in the cell growth of hiPSC-derived cancer cells (Figures 6I and S7D). The growth inhibitory effect was not obvious in the presence of siRAD21, which supports the notion that PCI34051 inhibits cell growth by impairing cohesin function (Figure S7D). Most notably, simultaneous treatment with PCI34051 and GSK126 markedly inhibited the cell proliferation (Figure 6I). Mechanistically, suppression of the ESC-like gene expression signature by GSK126 was more pronounced by the combination treatment of PCI34051 and GSK126 (Figures 6J and S7E), although PCI34051 alone did not significantly suppress the ESC-like signature (Figure S7F). Consistently, a gene ontology enrichment analysis revealed that the combination treatment induced genes associated with nervous system development (Figure 6K), suggesting that the combined inhibition of HDAC8 and EZH2 induced neuronal differentiation. We also confirmed that the

for AT/RTs and medulloblastomas as well as fetal brains and NSCs. AT/RTs harbor increased methylation at the PSC-specific methylated CpG sites, whereas adult brain-specific methylated CpG sites are less methylated in AT/RTs. Each dot indicates the median of DNA methylation at the analyzed CpG sites. Infinium450K data of hPSCs, normal brains, fetal brains, NSCs, AT/RTs, and medulloblastomas were obtained from GSE60821 (hPSC), GSE92462 (hPSC, normal brain, fetal brain, and NSC), GSE36278 (normal brain and fetal brain), GSE70460 (AT/RT), and GSE75153 (MB). See also Figures S4 and S5.



#### Figure 5. Pediatric Cancers Exhibit Activation of the ESC-like Gene Expression Signature

(A) Principal-component analysis of gene expressions in pediatric cancers and adult cancers using hESC-like module genes (top) and ESC Core module genes (bottom). Note that all pediatric cancers exhibit activation of the ESC-like module and the ESC Core module when compared to the corresponding adult cancers. AT/RT, atypical teratoid/rhabdoid tumor; NB, neuroblastoma; WT, Wilms' tumor; HB, hepatoblastoma; GBM, glioblastoma: ACC. adrenocortical carcinoma: RCC. renal cell carcinoma: HCC, hepatocellular carcinoma. Microarray data of hPSCs and pediatric and adult cancers were obtained from GEO: GSE22392 (hESC/hiPSC), GSE70678 (AT/RT), GSE16476 (neuroblastoma), GSE11151 (Wilms' tumor/renal cell carcinoma), GSE53224 (Wilms' tumor), GSE75271 (hepatoblastoma), GSE53733 (GBM), GSE10927 (adrenocortical carcinoma), GSE66272 (renal cell carcinoma), and GSE62232 (hepatocellular carcinoma).

(B) Expressions of SALL4 and LIN28B in pediatric cancers and adult cancers. Note that pediatric cancers show increased expression of SALL4 and LIN28B compared to the corresponding adult cancers. Data are represented as the median with interquartile range. \*\*\*\*p < 0.0001 (Mann-Whitney U test). The same microarray data as Figure 5A were used.

See also Figure S6.

mens of AT/RTs but not in medulloblastomas or glioblastomas. Of particular note, *SMARCB1*-deficient hPSC-transplanted mice showed poor survival compared to *SMARCB1*-deficient NPLC-

combination treatment remarkably suppressed proliferation in other rhabdoid and AT/RT cell lines (Figure 6L), as well as neuroblastoma cell lines (Figure 6M). In sharp contrast, the suppressive effect was not prominent in two of three glioblastoma cell lines (Figure S7G), which is consistent with the fact that most glioblastomas do not exhibit activation of the ESC-like signature.

#### DISCUSSION

In the present study, we established hPSCs deficient for *SMARCB1* to model AT/RT. Although previous studies suggested that NPCs are a cell-of-origin for AT/RT, the majority of tumors from *SMARCB1*-deficient NPLCs lacked typical rhabdoid cells. On the other hand, despite having the same genetic abnormality, the transplantation of *SMARCB1*-deficient hPSCs caused tumors containing a large number of rhabdoid cells, indicating that PSC-related embryonic cell properties are associated with the histogenesis of rhabdoid cells, which was further supported by the fact that forced activation of the ESC-like signature confers the rhabdoid histology in *SMARCB1*-deficient NPLC-derived tumors. Consistent with that notion, we found activation of the ESC-like signature in clinical specitransplanted mice, indicating a positive correlation between activation of the ESC-like signature and poor prognosis. Collectively, we developed a human AT/RT model using hPSCs and identified activation of the ESC-like signature as an important determinant of the unique histology and poor prognosis of AT/RT.

The fact that rhabdoid histology is prominent in PSC-derived tumors compared to NPLC-derived tumors suggests that an earlier embryonic program than the NPC program is involved in the unique histology. However, it remains unclear how AT/RT cells acquire the ESC-like signature in the postnatal brain. Notably, we found that ESC-methylated CpG sites are unmethylated in NSCs and fetal brains, suggesting that the ESC-like methylation patterns in AT/RTs do not simply reflect the DNA methylation patterns in NSCs or fetal cells. These findings may support the assumption that the ESC-like signature is acquired during AT/RT development. It is also interesting to note that genetic ablation of TP53 seems to enhance the emergence of the rhabdoid histology, although clinical AT/RTs hardly harbor the TP53 mutation. Given that the loss of TP53 greatly promotes somatic cell reprogramming into iPSCs (Hong et al., 2009), it is possible that TP53 deficiency contributed to the acquisition of the ESC-like signature by accelerating the reprogramming



(legend on next page)

process in this particular model. In contrast, we found that *TP53* deficiency alone did not promote the tumor formation *in vivo*, suggesting that the *TP53* deficiency enhances tumor formation in conjunction with *SMARCB1* deficiency.

Notably, we showed that *c-MYC* overexpression induces activation of the ESC-like signature in NPLC-derived tumors and drives tumor development with the rhabdoid phenotype. A previous study demonstrated that *c-MYC* activates the embryonic transcriptional program and causes stem cell-like phenotypes (Wong et al., 2008). Moreover, a recent study demonstrated that *c-MYC*-driven dedifferentiation supports the onset of a stem cell-like state and tumorigenesis in mammary epithelial cells (Poli et al., 2018). Together, we propose that *c-MYC* induces dedifferentiation and activates the ESC-like signature during tumor development from *SMARCB1*-deficient NPLCs. The fact that *c-MYC* is frequently amplified in AT/RT in patients at higher age (Johann et al., 2016) may further support the notion that *c-MYC* induces dedifferentiation during AT/RT development.

Taking advantage of our human AT/RT model, we showed that activation of the ESC-like signature is correlated with the poor prognosis, which raised the possibility that the ESC-like signature is a promising therapeutic target for AT/RT. Accordingly, we performed a CRISPR/Cas9 knock out screening targeting the maintenance of ESC identity and identified genes that potentially maintain the growth of tumor cells. Notably, the identified genes included *EZH2*, which was previously reported as a potential therapeutic target in AT/RT (Choi et al., 2016; Weingart et al., 2015). Importantly, we found that an EZH2 inhibitor efficiently inhibited the ESC-like signature in AT/RT cells, which is consistent with a previous study that demonstrated *Ezh2* maintains the stem cell-associated signature in *Smarcb1*-deficient mouse embryonic fibroblasts (Wilson et al., 2010).

We also found that knock out of *RAD21*, which encodes a component within the cohesin complex, significantly suppresses the growth of AT/RT cells, suggesting that the function of cohesin too could be a target for AT/RT treatment. Consistently, an HDAC8-specific inhibitor, which indirectly reduces localized cohesin, together with the EZH2 inhibitor synergistically inhibited activation of the ESC-like signature and markedly suppressed the proliferation of AT/RT cells but had minimal effect on glioblastoma cells. Although a mechanistic basis for the potent inhibition of the ESC-like signature by the combined inhibition of EZH2 and HDAC8 remains to be solved, we propose that inhibition of the ESC-like signature is an effective strategy for AT/RT treatment.

It is interesting that activation of the ESC-like signature was similarly detectable in other pediatric cancers, such as neuroblastomas, Wilms' tumors, and hepatoblastomas. Notably, the same knock out screening in neuroblastoma cell lines revealed that knock out of *RAD21* often suppressed the growth of neuroblastoma cells. Furthermore, the combined treatment with EZH2 and HDAC8 inhibitors synergistically reduced the proliferation of neuroblastoma cells. Together with previous findings that partial reprogramming *in vivo* induces activation of the ESC-like signature and causes the development of cancers that resemble pediatric cancers (Ohnishi et al., 2014), it is possible that activation of the ESC-like signature may be a general driver of pediatric

#### Figure 6. CRISPR/Cas9-Mediated Genetic Screening to Develop a Therapeutic Strategy for AT/RT

(A) A schematic illustration of the genetic screening with the lentiviral CRISPR/Cas9 system. An alamarBlue assay was performed to evaluate cell viability. The LentiCRISPR-transduced cell population was examined at days 3–5 and then used for the adjustment to determine the cell growth ratio at day 9. The average of the duplicate values was used for each assay.

(B) The results of a genetic screening of the hiPSC SMARCB1-/-: TP53-/--derived cancer cell line. The transduction of CRISPR/Cas9 with sgRNA targeting EZH2 and RAD21 inhibited cell growth.

(C) The results of a genetic screening of the neuroblastoma cell lines SK-N-AS and SK-N-BE(2). The LentiCRISPR-transduced cell population was examined at days 3 and 4, then used for the adjustment to determine the cell growth ratio at days 7–12. The transduction of CRISPR/Cas9 with sgRNA for *RAD21* inhibited cell proliferation in both cell lines.

(D) The expression of *EZH2* and *RAD21* in AT/RTs and glioblastomas. The increased expressions of both *EZH2* and *RAD21* were observed in AT/RTs compared to glioblastomas. The same microarray data as Figure 4A were used. \*\*\*\*p < 0.0001 (Kruskal-Wallis test and Dunn's multiple comparisons test).

(E) The expression of *EZH2* and *RAD21* in pediatric cancers and the corresponding adult cancers. Note that the increased expressions of *RAD21* and *EZH2* were similarly observed in other pediatric cancers. Data are represented as the median with interquartile range. \*\*\*p < 0.001, \*\*\*\*p < 0.0001 (Mann-Whitney U test). The same microarray data as Figure 5A were used.

(F) Growth ratios (day 11/day 3) of hiPSC-derived cancer cells transduced with CRISPR/Cas9 with sgRNA for non-targeting control (NTC), *EZH2*, or *RAD21*. The transduction of CRISPR/Cas9 with sgRNA for *EZH2* and *RAD21* reduced the cell growth ratio compared to *NTC*. Data are presented as the mean of three biological replicates with standard deviation. The growth ratio of *NTC*-transduced cells was set to 1. \*p < 0.05, \*\*\*\*p < 0.0001 (Kruskal-Wallis test and Dunn's multiple comparisons test).

(G) A schematic illustration of the xenograft transplantation study of hiPSC-derived cancer cells transduced with lentiCRISPR/Cas9 together with sgRNA for NTC, EZH2, or RAD21.

(H) Survival curves of mice transplanted with hiPSC-derived cancer cells transduced with CRISPR/Cas9 and sgRNA for *NTC*, *EZH2*, or *RAD21*. Kaplan-Meier analysis was performed. \*p < 0.05, \*\*p < 0.01 (log-rank test).

(I) Cell viability of hiPSC-derived cancer cells treated with DMSO, PCI34051 (10  $\mu$ M), GSK126 (10  $\mu$ M), or a combination of PCI34051 and GSK126 (10  $\mu$ M each). Note that the inhibitory effect is markedly enhanced by the combination treatment. \*\*p < 0.01, \*\*\*p < 0.01 (one-way ANOVA and Dunnett's multiple comparisons test). (J) GSEA showing the suppression of the ESC-like module by the combined treatment of PCI34051 and GSK126 compared to DMSO treatment in hiPSC-derived cancer cells.

(K) Gene ontology enrichment analysis showing that the combined treatment of PCI34051 and GSK126 resulted in the induction of genes associated with neuronal development in hiPSC-derived cancer cells. Genes showing  $\geq$  1.5-fold changes compared to DMSO-treated cells were used in the analysis.

(L) Cell viability of rhabdoid cell lines treated with DMSO, PCl34051 (10  $\mu$ M), GSK126 (10  $\mu$ M), or the combination of PCl34051 and GSK126 (10  $\mu$ M each). \*p < 0.05, \*\*\*p < 0.001, \*\*\*\*p < 0.0001 (one-way ANOVA and Dunnett's multiple comparisons test).

(M) Cell viability of neuroblastoma cell lines treated with DMSO, PCI34051 (10  $\mu$ M), GSK126 (10  $\mu$ M), or the combination of PCI34051 and GSK126 (10  $\mu$ M each). \*\*\*p < 0.001, \*\*\*\*p < 0.0001 (one-way ANOVA and Dunnett's multiple comparisons test).

See also Figure S7 and Table S1.

cancer development and, therefore, could be a therapeutic target for pediatric cancers. Indeed, previous studies showed that EZH2 inhibitors are often effective for pediatric cancers (Chen et al., 2018) and that an HDAC8 inhibitor suppresses cell proliferation and induces differentiation in neuroblastoma cells (Oehme et al., 2009). Collectively, we propose that the combined inhibition of EZH2 and HDAC8 could be a promising strategy to treat pediatric cancers by targeting the ESC-like signature.

In summary, we established a human AT/RT model using *SMARCB1*-deficient hPSCs. Taking advantage of the AT/RT model, we unveiled that activation of the ESC-like signature plays a central role in the unique rhabdoid histology and poor prognosis of AT/RT. Finally, we showed that this signature could be a promising therapeutic target for AT/RT as well as other pediatric cancers.

#### **STAR**\***METHODS**

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- CONTACT FOR REAGENT AND RESOURCE SHARING
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
  - Mice
  - Cell culture and neural induction
  - Establishment of hiPSC <sup>SMARCB1-/-; TP53-/-</sup> -derived cancer cell line
- METHOD DETAILS
  - TP53 and SMARCB1 knockout using CRISPR/Cas9 system
  - Retroviral transduction
  - Xenograft tumor model
  - O Histological analysis and immunostaining
  - Immunofluorescent staining
  - Western blot analysis
  - Magnetic-based cell sorting (MACS) and flow cytometric analysis
  - $\odot~$  Genetic screening with lentiviral CRISPR/Cas9 system
  - Cell growth inhibition assays and cell proliferation assays
  - siRNA transfection
  - RNA Preparation, gRT-PCR and microarray analysis
  - Library preparation for RNA sequencing
  - RNA-seq data analyses
  - DNA methylation analyses for infinium methylation array data
  - Quantification and Statistical Analysis
- DATA AND SOFTWARE AVAILABILITY

#### SUPPLEMENTAL INFORMATION

Supplemental Information can be found with this article online at https://doi. org/10.1016/j.celrep.2019.02.009.

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#### **AUTHOR CONTRIBUTIONS**

Y.T., N.J., and Yasuhiro Yamada proposed the research project, designed the experiments, performed the experiments, and wrote the manuscript. M.S., Yosuke Yamada, and T.U. performed the experiments. T.Y., M.K., and S.O. analyzed the microarray and RNA-sequencing data. S.M., Y.A., M.N., Y.M., K.M., and Y.M. provided materials and technical instructions.

#### **DECLARATION OF INTERESTS**

The authors declare no competing interests.

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#### **STAR\*METHODS**

#### **KEY RESOURCES TABLE**

REAGENT or RESOURCE	SOURCE	IDENTIFIER		
Antibodies	_			
Anti-SALL4	Abnova	Cat# clone 6E3; RRID: AB_566160		
Anti-LIN28A	Cell Signaling Technology	Cat# 8706; RRID: AB_10896850		
Anti-LIN28B	Cell Signaling Technology	Cat# 5422; RRID: AB_10697489		
Anti-Ki67 (SP6)	Abcam	Cat# ab16667; RRID: AB_302459		
Anti-INI1/SNF5	Sigma-Aldrich	Cat# SAB4200202; RRID: AB_10697389		
Anti-Vimentin (V9)	SantaCruz	Cat# sc-6260; RRID: AB_628437		
Anti-GFAP	DAKO	Cat# IR524		
Anti-CD99	DAKO	Cat# clone 12E7; RRID: AB_2076419		
Anti-S100	DAKO	Cat# GA504		
Anti-EMA	Novocastra	Cat# NCL-L-EMA; RRID: AB_563531		
Anti-SMA	Nichirei	Cat# clone1A4		
Anti-Synaptophysin	Nichirei	Cat# clone27G12		
Anti-NESTIN	Millopore	Cat# MAB5326; RRID: AB_2251134		
Anti-TP53	Santa Cruz	Cat# sc-126; RRID: AB_628082		
Anti-β-actin	Santa Cruz	Cat# sc-47778; RRID: AB_626632		
ECL anti-mouse IgG, HRP-linked whole antibody from sheep	GE Healthcare	Cat# NA931; RRID: AB_772210		
ESC anti-rabbit IgG and HRP-linked whole antibody from donkey	GE Healthcare	Cat# NA934; RRID: AB_772206		
Anti-PSA-NCAM antibody conjugated with microbeads	Miltenyi Biotec	Cat# 130-92-981		
Anti-PSA-NCAM	Millipore	Cat# MAB5324; RRID: AB_95211		
StemFit AK03N	Ajinomoto	Cat# AK03N		
StemFit AK02N	Ajinomoto	Cat# AK02N		
Laminin-511	Wako	Cat# 892012		
PSC Neural Induction Medium	Life Technologies	Cat# A1647801		
Y27632	Wako	Cat# 253-00513		
Chemicals, Peptides, and Recombinant Proteins	1			
GSK126	Funakoshi	Cat# A-1275		
PCI34051	Selleck	Cat# S2012		
siGENOME SMARTpool siRNA, RAD21	Dharmacon	Cat# M-006832-01-0005		
siGENOME SMARTpool siRNA, Non-Targeting	Dharmacon	Cat# D-001206-13-05		
Critical Commercial Assays				
AlamarBlue cell viability reagent	Bio-Rad	Cat# BOF012B		
Cell Counting Kit-8	Dojindo	Cat# 341-07761		
Human Gene 1.0 ST Array	Affymetrix	Cat# 901086		
Truseq Stranded mRNA LT sample prep kit	Illumina	Cat# RS-122-2101, RS-122-2102		
Deposited Data				
Microarray data	This paper	GSE118653		
RNA-seq data	This paper	GSE118654		
Affymetrix Human Gene 1.0ST Array datasets	Gene Expression Omnibus	GEO: GSE26313, GSE45265, GSE36947, GSE18296 and GSE27667		

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Affymetrix Human Genome U133 Plus 2.0 Array datasets	Gene Expression Omnibus	GEO: GSE70678, GSE37418, GSE53733, GSE16476, GSE11151, GSE53224, GSE75271, GSE10927, GSE66272, GSE62232, GSE73038 and GSE22392
Illumina HumanMethylation450 BeadChip datasets	Gene Expression Omnibus	GEO: GSE60821, GSE92462, GSE70460, GSE75153, GSE36278 and GSE73801
Experimental Models: Cell Lines		
Human: 201B7 hiPSC	Laboratory of Masato Nakagawa	hiPS Cell Line: 201B7
Human: 1383D6 hiPSC	Laboratory of Masato Nakagawa	hiPS Cell Line: 1383D6
Human: KUP-ATRT-1	Kyoto University	N/A
Human: A204	ATCC	HTB-82
Human: G401	ATCC	CRL-1441
Human: G402	ATCC	CRL-1440
Human: SK-N-AS	ATCC	CRL-2137
Human: SK-N-BE(2)	ATCC	CRL-2271
Human:T98G	Cell Resource Center for Biomedical Research, Institute of Development Aging and Cancer, Tohoku University	TKG0471
Human:A172	Cell Resource Center for Biomedical Research, Institute of Development, Aging and Cancer, Tohoku University	TKG0183
Human:U87-MG	ATCC	HTB-14
Experimental Models: Organisms/Strains		
Mouse: NOD/ShiJic-scidJcl	CLEA Japan	N/A
Oligonucleotides		
Oligos for sgRNA, see Table S1	This study and Brunello Library	Human CRISPR Knockout Pooled Library (Brunello)
Primers for quantitative PCR, see Table S4	This study	N/A
Oligos for TP53 sgRNA: CGCTATCTGAGCAGCGCTCA	This study	N/A
Oligos for SMARCB1 sgRNA: TGAGAACGCATCTCAGCCCG	This study	N/A
Recombinant DNA		
lentiCRISPR v2	Sanjana et al., 2014	Addgene: #52961
pX330-U6-Chimeric BB-CBh-hSpCas9	Cong et al., 2013	Addgene: #42230
pMYs-IRES-GFP	Cell Biolads	Cat# RTV-021
pMYs-c-MYC-IRES-GFP	This study	N/A
pMx-GFP	Cell Biolads	Cat# VPK-302
pMxs-hOCT4	Takahashi et al., 2007	Addgene: #17217
pMxs-hSOX2	Takahashi et al., 2007	Addgene: #17218
pMxs-hKLF4	Takahashi et al., 2007	Addgene: #17219
pMXs-hc-MYC	Takahashi et al., 2007	Addgene: #17220
pCMV-VSV-G	Cell Biolads	Cat# VPK-302
pMD2.G	Addgene	Cat# 12259
psPAX2	Addgene	Cat# 12260
Software and Algorithms		
GSEA software (version 3.0)	Subramanian et al., 2005	http://software.broadinstitute.org/gsea/ index.jsp
DAVID bioinformatics database website	Huang et al., 2009	https://david.ncifcrf.gov/home.jsp

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Graphpad Prism 6	MDF	http://www2.usaco.co.jp/shop/c/cGPPrism/
GeneSpring GX software (version 12)	Agilent Technology	https://www.chem-agilent.com/contents. php?id=27881
Other		
Gene set: WONG_EMBRYONIC_ STEM_CELL_CORE	Wong et al., 2008	http://software.broadinstitute.org/gsea/ msigdb/cards/WONG_EMBRYONIC_ STEM_CELL_CORE
Gene set: hESC-like module	Wong et al., 2008	Table S5
Gene set: Core Human module	Kim et al., 2010	Table S3

#### **CONTACT FOR REAGENT AND RESOURCE SHARING**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Yasuhiro Yamada (yasu@ims.u-tokyo.ac.jp).

#### **EXPERIMENTAL MODEL AND SUBJECT DETAILS**

#### Mice

All animal experiments were approved by the Animal Experiment Committee at CiRA and IMSUT, and the care of the animals was in accordance with institutional guidelines. All mice used for the present study were on NOD/ShiJic-*scid*Jcl (NOD SCID) mice of both sexes (8-10 weeks). NOD SCID mice were purchased from CLEA Japan, Inc. Mice were kept in the animal facility with 12 hours of light and dark cycle with food and water *ad libitum*.

#### **Cell culture and neural induction**

201B7 and 1383D6 hiPSCs (provided by Masato Nakagawa, CiRA) were cultured in StemFit AK03N or AK02N (Ajinomoto) on cell culture plates coated with laminin-511 (Wako) at 37°C with 5%  $CO_2$ . For neural induction, culture medium was switched to GIBCO PSC Neural Induction Medium (Life Technologies) containing Neurobasal medium and GIBCO PSC neural induction supplement. At day 7 of the neural induction, these cells were dissociated and plated in neural expansion medium containing 50% Neurobasal medium, 50% Advanced DMEM/F12 and neural induction supplement. Cells were treated with 10  $\mu$ M ROCK inhibitor Y27632 (Wako) at the time of plating overnight to prevent cell death. To further induce differentiation, culture medium was switched to Neurobasal medium containing 2% B-27 Supplement (GIBCO) and GlutaMAX Supplement (GIBCO).

KUP-ATRT-1, an AT/RT cell line, was established at Kyoto University. Other tumor cell lines were purchased from ATCC or Cell Resource Center for Biomedical Research, Institute of Development, Aging and Cancer, Tohoku University. Tumor cells were cultured with RPMI 1640 (GIBCO) or MEM, GlutaMAX<sup>™</sup> supplement (GIBCO) containing 10% fetal bovine serum (FBS) at 37°C with 5% CO<sub>2</sub>.

#### Establishment of hiPSC SMARCB1-/-; TP53-/- -derived cancer cell line

A block of hiPSC <sup>SMARCB1-/-; TP53-/-</sup>-derived tumor was mechanically minced. The minced tissue was digested with StemPro Accutase Cell Dissociation Reagent (GIBCO) at 37°C for 15 minutes. After washing, the tissue was triturated and passed through a 100  $\mu$ m cell strainer. Cells were plated onto cell culture plates coated with laminin-511 in AK03N media. Passaging of the cultures was performed approximately once a week.

#### **METHOD DETAILS**

#### TP53 and SMARCB1 knockout using CRISPR/Cas9 system

The *TP53* and *SMARCB1* genes were knocked-out with P3 primary cell 4D-Nucleofector X Kit (Lonza). 201B7 cells were transfected with modified pX330-U6-Chimeric\_BB-CBh-hSpCas9 plasmid (Addgene: Plasmid #42230) (Cong et al., 2013) expressing a sgRNA against *TP53* (CGCTATCTGAGCAGCGCTCA) or *SMARCB1* (TGAGAACGCATCTCAGCCCG) with puromycin-resistance gene. To avoid genomic integration, puromycin selection (1  $\mu$ l/ml) was performed only for two days. Each single colony was picked-up and expanded. The knockout for TP53 and SMARCB1 was confirmed by sequencing and western blot.

#### **Retroviral transduction**

To generate retroviral vectors, Plat-GP cells on 150 mm dishes were cultured with DMEM containing 100 U/ml penicillin, 100  $\mu$  g/ml streptomycin (P/S) (Nacalai tesque) and 10% FBS (GIBCO) until 70%–80% confluency and were transfected with 12.375  $\mu$ g retroviral

vector plasmid (pMx-GFP, pMxs-hOCT4, pMxs-hSOX2, pMxs-hKLF4, pMxs-hc-MYC and pMYs-IRES-GFP or pMYs-hc-MYC-IRES-GFP) (Addgene, Cell Biolabs, Inc.) (Takahashi et al., 2007) independently in combination with 5.625  $\mu$ g pCMV-VSV-G (Cell Biolabs, Inc.) using Lipofectamin 2000 (Life technologies). 24 hours after the transfection, the culture media was refreshed, and the supernatant was collected over 3 consecutive days. The filtered supernatant was concentrated by PEG-*it* (System Biosciences), re-suspended, aliquoted and stored at  $-80^{\circ}$ C. 1 ×  $10^{6}$  of NPLCs<sup>SMARCB1-/-; TP53-/-</sup> or PSA-NCAM-positive NPLCs<sup>TP53-/-</sup>; SMARCB1-/-; SMARCB1-/-</sup> were incubated with the concentrated virus-containing supernatant overnight. Five days after the infection, cells were harvested and transplanted into the mouse brain. At the same time, RNA was extracted from the infected cells.

#### Xenograft tumor model

1  $\mu$ l of 4 × 10<sup>5</sup> cells were injected into the left striatum of NOD SCID mice at the coordinates of 1.5 mm lateral from the bregma and 2 mm deep from the dura using a 10  $\mu$ l Hamilton syringe with a flow rate of 1  $\mu$ l per minute. MRI studies were performed on a 1.5-Tesla MRI scanner (MRmini SA1508; DS Pharma Biomedical). Transplanted mice were traced until they died or showed some neurological signs (observation period: 4-31weeks).

#### Histological analysis and immunostaining

Mice were transcardially perfused with 1 × PBS followed by 4% paraformaldehyde. Following dissection, brain tissues were transferred to PBS and subsequently embedded in paraffin and sectioned. Sections were stained with hematoxylin and eosin (H&E), and serial sections were used for the immunohistochemical analysis. The primary antibodies, which were incubated at 4°C overnight in blocking buffer, were as follows: anti-SALL4 (Abnova: #clone 6E3), anti-LIN28A (Cell Signaling Technology: #8706), anti-LIN28B (Cell Signaling Technology: #5422), anti-Ki67 (SP6) (Abcam: #ab16667), anti-IN1/SNF5 (Sigma-Aldrich: #91735), anti-Vimentin (V9) (SantaCruz: #sc-6260), anti-GFAP (DAKO: #IR524), anti-CD99 (DAKO: #clone 12E7), anti-S100 (DAKO: #GA504), anti-EMA (Novocastra: #NCL-L-EMA), anti-SMA (Nichirei: #clone1A4) and anti-Synaptophysin (Nichirei: #clone27G12). The sections were incubated with the appropriate species of HRP-conjugated secondary antibodies (Nichirei, Histofine) at room temperature for 30 minutes, and chromogen development was performed using DAB (Nichirei). The stained slides were counterstained with Meyer hematoxylin. The histopathological sections were reviewed by two different pathologists (Yo Y and Ya Y) who have been certified by the Japanese Society of Pathology.

#### Immunofluorescent staining

The samples were fixed with 4% paraformaldehyde for five minutes and soaked in PBS. They were then incubated with anti-NESTIN (Millipore: #MAB5326) at 4°C overnight in blocking buffer and were processed with × 500 DAPI (Invitrogen) and × 150 fluorescence-labeled secondary antibodies diluted with 0.5% BSA in PBS for 90 minutes at room temperature. After washing in PBS for 5 minutes twice, the samples were mounted and evaluated with a confocal laser-scanning microscope (Zeiss LSM700 or 710).

#### Western blot analysis

Cell lysates were prepared in RIPA buffer with protease inhibitors. Protein samples were carried out according to standard methods. The primary antibodies used were anti-INI1/SNF5 (Sigma-Aldrich: #91735), anti-TP53 (Santa Cruz Biotechnology: #sc-126) and anti-β-actin (Santa Cruz, sc-47778). The secondary antibodies used were ECL anti-mouse IgG, HRP-linked whole antibody from sheep (NA931, GE Healthcare), ECL anti-rabbit IgG and HRP-linked whole antibody from donkey (NA934, GE Healthcare). ImageQuant LAS4000 (GE Healthcare) was used for band detection.

#### Magnetic-based cell sorting (MACS) and flow cytometric analysis

Expanded NPLCs<sup>SMARCB1-/-; TP53-/-</sup> were exposed to 10 µM of Y27632 (Wako) for more than one hour to prevent cell death prior to the MACS procedure. After dissociation, the cells were briefly blocked in 0.5% BSA-PBS solution and then incubated with anti-PSA-NCAM antibody conjugated with microbeads (Miltenyi Biotec: #130-92-981) for 15 minutes at 4°C. After extensive washing, the cell suspension was loaded on a separation column (LS column) that was attached to a magnetic stand. Positively-labeled cells that remained in the column were eluted to a tube with culture medium for further manipulation and analysis. After magnetic cell separation, we further used flow cytometry to examine the purity of PSA-NCAM-positive cells. MACS-sorted cells were cultured for 24 hours and stained with anti-PSA-NCAM antibody (Millipore: #MAB5324) for 15 minutes at 4°C followed by CF488-conjugated secondary antibody for 10 minutes at 4°C. The positive fraction was evaluated by flow cytometry (Aria II, BD) using non-stained cells as a control.

#### Genetic screening with lentiviral CRISPR/Cas9 system

We utilized the lentiCRISPR v2 (Addgene #52961) (Sanjana et al., 2014) for one-by-one gene disruption. The candidate genes were selected based on the previous literatures and databases (Table S1). The lentiviral vector has both a sgRNA scaffold and a Cas9 to induce insertions and deletions (indels) at the genomic locus of the cells into which they are transduced.

The lentiviral cloning and production were performed as described previously (Sanjana et al., 2014). Briefly, the complementary oligonucleotides for sgRNAs (Table S1) were annealed by heating to 95°C for 3 minutes and subsequent cooling to 60°C for 3 minutes on ice. The lentiCRISPR v2 plasmid was digested with BsmBI (New England Biolabs) at 55°C overnight and purified by FastGene

Gel/PCR Extraction Kit (Genetics). The annealed oligonucleotides were ligated into the digested lentiCRISPR v2 plasmid by Ligation high Ver.2 (TOYOBO) at 16°C for 30 minutes. The reactants were transformed into Stbl3 chemically competent *E. coli* (Thermo Fisher Scientific). The grown colonies were analyzed by Sanger sequencing with ABI 3500xL (Applied Biosystems) to confirm correctly recombined clones. The lentiviral plasmids were extracted by GenElute Plasmid Miniprep Kit (Sigma).

To generate lentivirual vectors, HEK293T cells on 60 mm dishes were cultured with DMEM containing 100 U/ml penicillin, 100 µg/ml streptomycin (P/S) (Nacalai tesque), 10% FBS (GIBCO) until 70%-80% confluency and transfected with 2.4 µg lenti-CRISPR v2 plasmid, 0.8 µg pMD2.G (Addgene #12259) and 1.6 µg psPAX2 (Addgene #12260) using PEI max (Polysciences). 24 hours after transfection, culture media were refreshed, and the supernatant was collected over 3 consecutive days. The filtered supernatant was concentrated by PEG-it (System Biosciences), re-suspended, aliquoted and stored at -80°C. No lentiviral functional titration was performed. Instead, we measured base-line cell viability at days 3-5 during the screening process to adjust the transduction efficiency bias as explained below. The screening was performed with 96-well plate format in duplicate. Each cell line was plated to be 70%-80% confluent at the day of lentiviral transduction (day 0). We avoided using outer lanes, where the value could be variable because of the media evaporation. At day 0, virus-containing supernatant was added to each well, and 24 hours later, puromycin was added to eliminate non-transfected cells. At days 3-5, the baseline cell viability was assessed using alamarBlue (Bio-Rad). Fluorescence was detected with 2104 EnVision Multi Detection Microplate Reader (Perkin Elmer). The average fluorescence intensity value in blank wells was subtracted to determine the fluorescence intensity value of each well. Media were replaced every 2-3 days. When wells transduced with non-targeting control (NTC) sgRNA reached 70%-80% confluency, cell viability was measured again by alamarBlue to assess the effect of each knockout on cell growth. Cell growth rates were calculated as follows: the average of duplicate fluorescence intensity values was divided by the average at baseline. Cell growth rates were compared with NTC sgRNA-transduced cells.

To confirm the CRISPR/Cas9-mediated cleavage efficiency in this screening, we randomly selected 20 genes among the candidate genes and assessed the indel frequency at days 3 and 7 in the SK-N-BE(2) cell line. The indel frequency was calculated by TIDE software (https://tide.nki.nl) (Brinkman et al., 2014). The TIDE software parameters used in this study were as follows: left boundary, 100 bp; right boundary, -10 bp; decomposition window, 115-685 bp; indel size range, 20 bp.

The titration of puromycin was pre-determined for each cell line to efficiently eliminate non-transduced cells at days 3-5. The concentrations of puromycin used in this study were 1.0, 1.0 and 1.5  $\mu$ g/ml for hiPSC <sup>SMARCB1-/-; TP53-/-</sup>-derived cancer cell line, SK-N-BE(2) and SK-N-AS, respectively.

#### Cell growth inhibition assays and cell proliferation assays

In vitro drug sensitivity was determined using alamarBlue cell viability reagent (Bio-Rad) according to the manufacturer's protocol. Cells were plated into 96-well culture plates at a density of  $2 \times 10^3$  cells/well. Cells were treated with 10  $\mu$ M of DMSO, GSK126, PCI34051 or a combination of GSK126 and PCI34051 at day 0 and incubated for 7 days. The experiment was performed in triplicate, and each sample was measured three times. Fluorescence intensity was detected with 2104 EnVision Multi Detection Microplate Reader. The average fluorescence intensity value of blank wells was subtracted to determine the fluorescence intensity of each well at each time point. The average fluorescence intensity of DMSO-treated cells at day 7 was set to 1. Cell Counting Kit-8 (Dojindo) was also used to examine *in vitro* cell proliferation. Abosrbance was measured using iMark Microplate Absorbance Reader (Bio-Rad).

#### siRNA transfection

siRNA transfection was performed using Lipofectamine RNAi Max (Invitrogen). We performed knockdown assays with a siRNA targeting *RAD21* (Dharmacon). Nontargeting siRNA (Dharmacon) was used as a control. Culture medium was exchanged every 2 days, and cell proliferation was determined using Cell Counting Kit-8 (Dojindo).

#### **RNA Preparation, qRT-PCR and microarray analysis**

Total RNA was isolated using the RNeasy Plus Mini kit (QIAGEN). The qRT-PCR analysis was performed using GoTaq qPCR Master Mix (Promege). The specific primer pairs used for amplification are shown in Table S4. The transcript levels were normalized to the *GAPDH* level. The microarray analysis was performed using Human Gene 1.0 ST Array (Affymetrix) in accordance with the manufacturer's instructions, GeneSpring GX software program (version 12; Agilent Technology), GSEA software (version 3.0) and the DAVID bioinformatics database website (https://david.ncifcrf.gov/home.jsp). The gene sets of the hESClike module (Wong et al., 2008) and the Core Human module (Kim et al., 2010) were used for the clustering analysis. The gene set of "WONG\_EMBRYONIC\_STEM\_CELL\_CORE" in MSigDB (version 6.0) was used for the GSEA analysis.

#### Library preparation for RNA sequencing

200 ng of total RNA was prepared for the library construction. High-quality RNA (RNA Integrity Number value  $\geq$  7) assessed by Bioanalyzer was used for the library preparation. RNA-seq libraries were generated using the Truseq Stranded mRNA LT sample prep kit (Illumina). PolyA-containing mRNA was purified by poly-T oligo-attached magnetic beads, and the RNA was fragmented and primed for cDNA synthesis. Cleaved RNA fragments were reverse transcribed into first strand cDNA using transcriptase and random primers. Second strand cDNA was synthesized by the incorporation of dUTP, and ds cDNA was separated using AMPure XP beads (BECKMAN COULTER). A single 'A' nucleotide was added to the 3' ends of the blunt fragments, and then adapters with index were ligated to the ends of the ds cDNA. ds cDNA fragments were amplified by PCR with PCR primer Cocktail. The number of PCR cycles was minimized (15 cycles) to avoid skewing the representation of the libraries. RNA-seq libraries were sequenced on NextSeq 500 (75 bp or 86 bp single, Illumina).

#### **RNA-seq data analyses**

The sequenced reads were mapped to the human reference genome (hg38) using Tophat2 (version 2.1.1) with the GENCODE (version 27) annotation gtf file and the aligner Bowtie2-2.3.4 (Langmead and Salzberg, 2012) after trimming adaptor sequences and low-quality bases by cutadapt-1.16 (Martin, 2011). The uniquely mapped reads were used for further analyses. Using cufflinks-2.2.1 (Trapnell et al., 2010) with the human GENCODE (version 27, protein coding) annotation gtf file, reads per kilobase of exon per million mapped reads (RPKM) were calculated as the expression levels of each gene. RPKM values were used for the GSEA method.

#### DNA methylation analyses for infinium methylation array data

Infinium450K data were obtained from GSE60821 (hESC), GSE92462 (hESC, NSCs, normal brain and fetal brain), GSE36278 (normal brain, fetal brain and GBM), GSE75153 (MB), GSE73801 (ETMR) and GSE70460 (AT/RT). Previously described human CGIs (Illingworth et al., 2010) were used for the methylation analysis. The UCSC LiftOver tools (http://genome.ucsc.edu/) (Rosenbloom et al., 2015) were used to convert the coordinates of hg18 assembly into those of hg19 assembly. After the conversion of the CGI regions, overlap regions were merged into a single region. The UCSC refGene table was used to determine the TSS sites. The median signal value of the probes within each CGI and TSS  $\pm$  1,500 bp was calculated as the methylation signal of the region. Brain-methylated regions and brain-unmethylated regions were defined as the CGI and TSS  $\pm$  1,500 bp that represent higher (> 0.6) and lower (< 0.6) brain-methylation (median methylation signals in 7 brain samples), respectively, compared to PSC-methylation (median methylation signals in 6 PSC samples). The all probe methylation signals within the indicated regions were used in violin plots.

#### **Quantification and Statistical Analysis**

To quantify the dominance of rhabdoid cells in tumors, an H&E stained section was randomly photographed at  $10 \times$  magnification. Three or 4 pictures for each sample were processed with ImageJ software (NIH) to evaluate the positive area of rhabdoid cells. The positive area was determined by the area of rhabdoid cells divided by the area of the tumor in the histological image. To assess *SALL4*-positive cells, each section was randomly photographed at 200 × magnification. Positive nuclei in tumor cells were counted using five images. The number of positive nucleus was divided by the number of total nuclei of tumor cells in each image. These results were evaluated with Graphpad Prism 6 software.

All values and graphs are expressed as the mean with 95% confidence interval or the median with interquartile range, and statistical analyses were performed using unpaired t test with Welch's correction or Mann-Whitney U test for continuous variables and Fisher's exact test for categorical variables. One-way ANOVA or Kruskal-Wallis test was performed to compare multiple groups. Dunnett's or Dunn's multiple comparisons test was used for multiple comparisons. Kaplan-Meier survival curves were compared using the log-rank test. Statistical parameters including statistical significance and n values are described in the figures and figure legends. A value of p < 0.05 was considered significant. All analyses were conducted using Graphpad Prism 6 software.

#### DATA AND SOFTWARE AVAILABILITY

All data analyzed by microarray and RNA-seq have been deposited in the Gene Expression Omnibus (GEO) under accession numbers GSE118653 and GSE118654.

Cell Reports, Volume 26

## **Supplemental Information**

## Human Pluripotent Stem Cell-Derived Tumor Model

### Uncovers the Embryonic Stem Cell Signature as a

## Key Driver in Atypical Teratoid/Rhabdoid Tumor

Yukinori Terada, Norihide Jo, Yoshiki Arakawa, Megumi Sakakura, Yosuke Yamada, Tomoyo Ukai, Mio Kabata, Kanae Mitsunaga, Yohei Mineharu, Sho Ohta, Masato Nakagawa, Susumu Miyamoto, Takuya Yamamoto, and Yasuhiro Yamada

#### **Supplemental Information**

Human pluripotent stem cell-derived tumor model uncovers the embryonic stem cell signature as a key driver in atypical teratoid/rhabdoid tumor

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Figure S1: Generation of an atypical teratoid/rhabdoid tumor model using hiPSCs lacking *SMARCB1*, related to Figure 1

# Figure S1: Generation of an atypical teratoid/rhabdoid tumor model using hiPSCs lacking *SMARCB1*, related to Figure 1

(A) A qRT-PCR for pluripotency genes and neural differentiation genes. Data are presented as the mean  $\pm$  SD of biological triplicates. The mean expression level of hiPSCs <sup>201B7</sup> or NPLCs <sup>201B7</sup> was set to 1. (B) A qRT-PCR for neuronal differentiation genes after the neuronal differentiation of NPLCs for 7 days. Data are presented as the mean  $\pm$  SD of biological triplicates. The mean expression level of NPLCs (differentiation 7 days) was set to 1.

(C) Gene ontology enrichment analysis of NPLCs SMARCB1-/-; TP53-/- compared to undifferentiated hiPSCs SMARCB1-/-; TP53-/-. Analysis was performed using DAVID (Huang da et al., 2009). Genes showing  $\geq$  2-fold changes in NPLCs SMARCB1-/-; TP53-/- compared to hiPSCs TP53-/-; SMARCB1-/-; Were used in the analysis.

(D) Cell proliferation assays of hiPSCs and NPLCs. \*\*p<0.01, \*\*\*p<0.001 (One-way ANOVA and Dunnett's multiple comparisons test).

(E) Macroscopic tumor incidence after xenograft transplantation of hiPSCs and NPLCs into the brain of immunocompromised mice.

(F) Microscopic tumor incidence after xenograft transplantation of hiPSCs and NPLCs into the brain of immunocompromised mice.

(G) Representative histological images of microscopic tumors. The iPSC <sup>201B7</sup>-derived tumor exhibits a teratoma-like histology, which contains ciliated epithelial cells (left), while the NPLCs <sup>*TP53-/-*</sup>-tumor shows a scar-like histology (right). Scale bars, 500 μm (top) and 50 μm (bottom).

(H) Western blot analysis of 1383D6 iPSCs SMARCB1-/- for SMARCB1 and  $\beta$ -actin shows the lack of SMARCB1 in 1383D6 iPSCs SMARCB1-/-.

(I) Representative histological images of a 1383D6 iPSC *SMARCB1-/-*-derived tumor and 1383D6 NPLC *SMARCB1-/-*-derived tumor. Note that the 1383D6 iPSC *SMARCB1-/-*-derived tumor contains rhabdoid cells. Scale bars, 500 μm (top) and 20 μm (bottom).

(J) Quantification of the rhabdoid area within 1383D6 iPSC <sup>SMARCB1-/-</sup>-derived tumors and 1383D6 NPLC <sup>SMARCB1-/-</sup>-derived tumors. Data are represented as the median with interquartile range. \*p<0.05 (Mann-Whitney U test).



Figure S2: Induction of the ESC-like signature leads to rhabdoid phenotype in tumors, related to Figure 2

# Figure S2: Induction of the ESC-like signature leads to rhabdoid phenotype in tumors, related to Figure 2

(A) A schematic illustration of a xenograft transplantation study of NPLCs *SMARCB1-/-*; *TP53-/-* transduced with *GFP* (*GFP*-NPLCs) or four reprogramming factors (*OCT4, SOX2, KLF4* and *c-MYC*) (*OSKM*-NPLCs).
(B) A flow cytometric analysis of PSA-NCAM expression in NPLCs *SMARCB1-/-*; *TP53-/-* after sorting with

PSA-NCAM antibody by magnetic activated cell sorting (MACS).

(C) A qRT-PCR for *OCT4*, *SOX2*, *KLF4* and *c-MYC* at 5 days after the transduction of *GFP* or *OSKM* in NPLCs *SMARCB1-/-*; *TP53-/-*. Data are presented as the mean of technical triplicates. The mean expression level of *OSKM*-NPLCs was set to 1.

(D) A qRT-PCR for pluripotent genes and neural differentiation genes at 5 days after the transduction of *GFP, c-MYC* or *OSKM* in NPLCs <sup>*SMARCB1-/-*; *TP53-/- in vitro*. Data are presented as the mean of three biological replicates with standard deviation. The mean expression level of *GFP*-NPLCs was set to 1. \*p<0.05, \*\*p<0.01 (One-way ANOVA and Dunnett's multiple comparisons test). (E) (F) Survival curve of mice transplanted with *GFP*-NPLCs or *OSKM*-NPLCs. Note that mice inoculated with *OSKM*-NPLCs exhibit poor survival when compared to mice inoculated with control NPLCs. Kaplan-Meier analysis was performed. \*p<0.05, \*\*p<0.01 (Log-rank test).</sup>



Figure S3: *c-MYC* induces activation of the ESC-like signature and drives rhabdoid tumors *in vivo*, related to Figure 3

# Figure S3: *c-MYC* induces activation of the ESC-like signature and drives rhabdoid tumors *in vivo*, related to Figure 3

(A) A schematic illustration of a xenograft transplantation study of NPLCs <sup>SMARCB1-/-; TP53-/-</sup> transduced with *GFP* (*GFP*-NPLCs) or *c-MYC* (*MYC*-NPLCs). Control *GFP*-NPLCs are the same as Figure S2. (B) A qRT-PCR for *c-MYC* after the transduction of *GFP* or *c-MYC* in NPLCs <sup>SMARCB1-/-; TP53-/-</sup>. Data are presented as the mean of technical triplicates. The mean expression level of *MYC*-NPLCs was set to 1. (C) Representative histological images of a *MYC*-NPLC-derived tumor. Note that a number of rhabdoid cells are observed in the tumor. Scale bars, 500  $\mu$ m (upper) and 20  $\mu$ m (lower). (D) Quantification of the rhabdoid area in *GFP*-NPLC- and *MYC*-NPLC-derived tumors. Data are represented as the median with interquartile range. \*\*p<0.01 (Mann-Whitney U test). (E) Survival curves of mice transplanted with *GFP*-NPLCs or *MYC*-NPLCs. Note that the overall survival of mice inoculated with *MYC*-NPLCs is significantly shorter than of control. Kaplan-Meier analysis was performed. \*\*\*\*p<0.0001 (Log-rank test).



Figure S4: Human AT/RT specimens exhibit activation of the ESC-like gene expression signature, related to Figure 4

# Figure S4: Human AT/RT specimens exhibit activation of the ESC-like gene expression signature, related to Figure 4

(A) Clustering analysis using microarray data revealed that both hESC-like module genes (left) and ESC Core module genes (right) are similarly expressed in AT/RT samples and ESCs/iPSCs. Note that some ETMRs were clustered with AT/RT samples and ESCs/iPSCs. Medulloblastoma (MB) and glioblastoma (GBM) samples are basically clustered separately from ESCs/iPSCs. The microarray data of hPSCs, AT/RTs, ETMRs, medulloblastomas and glioblastomas were obtained from GSE22392 (hESC/hiPSC), GSE73038 (ETMR), GSE70678 (AT/RT), GSE37418 (MB) and GSE53733 (GBM).

(B) Immunohistochemical analysis of SALL4, LIN28A and LIN28B in clinical AT/RT specimens. Scale bars, 50  $\mu$ m.



Figure S5: DNA methylation landscape in human AT/RT specimens, related to Figure 4

#### Figure S5: DNA methylation landscape in human AT/RT specimens, related to Figure 4

The DNA methylation landscape in AT/RTs analyzed using Infinium450K data. In this analysis, we first extracted differentially methylated CpG sites within CGIs (left) or around TSSs (right) between hPSCs and adult brains. hPSC-specific methylated CpG sites and adult brain-specific methylated CpG sites were analyzed for AT/RTs, ETMRs, medulloblastomas and glioblastomas as well as fetal brains and NSCs. AT/RTs harbor increased methylation at the PSC-specific methylated CpG sites, while adult brain-specific methylated CpG sites are less methylated in AT/RTs. Each dot indicates the median of DNA methylation at the analyzed CpG sites. Infinium450K data of hPSCs, normal brains, fetal brains, NSCs, AT/RTs, ETMRs, medulloblastomas and glioblasomas were obtained from GSE60821 (hPSC), GSE92462 (hPSC/NSC/normal brain/fetal brain), GSE36278 (normal brain/fetal brain/GBM), GSE73801 (ETMR), GSE75153 (MB) and GSE70460 (AT/RT).



Figure S6: Pediatric cancers exhibit activation of the ESC-like gene expression signature, related to Figure 5

# Figure S6: Pediatric cancers exhibit activation of the ESC-like gene expression signature, related to Figure 5

Principal component analysis of the global gene expression profiles in pediatric cancers and adult cancers. Pediatric cancers exhibit ESC-like gene expression patterns when compared to the corresponding adult cancers. AT/RT: atypical teratoid/rhabdoid tumor, NB: neuroblastoma, WT: Wilms tumor, HB: hepatoblastoma, GBM: glioblastoma, ACC: adrenocortical carcinoma, RCC: renal cell carcinoma, HCC: hepatocellular carcinoma. The same microarray data as Figure 5A were used.



Figure S7: Genetic screening with CRISPR/Cas9 to develop a novel therapeutic strategy for AT/RT, related to Figure 6

# Figure S7: Genetic screening with CRISPR/Cas9 to develop a novel therapeutic strategy for AT/RT, related to Figure 6

(A) The cleavage efficiency at Day 3 and Day 7 after lentiviral-mediated transduction of CRISPR/Cas9 together with sgRNAs in SK-N-BE(2) *in vitro*. sgRNAs for 18 randomly selected genes were transduced with CRISPR/Cas9, and the efficiency of non-homologous end joining was measured by TIDE software 3 and 7 days after the lentiviral transduction. Data are represented as the median with interquartile range.
(B) GSEA showing that EZH2 inhibition by GSK126 suppresses the ESC-like module in hiPSC-derived cancer cells.

(C) Growth ratios (Day 14/Day 3) of hiPSC-derived cancer cells, SK-N-AS and SK-N-BE(2) transduced with CRISPR/Cas9 with sgRNA for non-targeting control (NTC) or *RAD21*. Two independent sgRNAs for *RAD21* reduced the cell growth ratio compared to *NTC*. Data are presented as the mean of three biological replicates with standard deviation. The growth ratio of *NTC*-transduced cells was set to 1. \*p<0.05, \*\*p<0.01 and \*\*\*\*p<0.0001 (One-way ANOVA and Dunnett's multiple comparisons test). (D) Cell viability of hiPSC-derived cancer cells treated with PCI34051 (10 µM) or the combination of PCI34051 (10 µM) and *siRAD21*. Note that the growth inhibitory effect of PCI34051 is not obvious in the presence of *siRAD21*. \*p<0.05, \*\*\*p<0.0001 (One-way ANOVA and Dunnett's multiple comparisons test).

(E) GSEA revealed that suppression of the ESC-like module by GSK126 is further pronounced by the combination treatment of GSK126 and PCI34051.

(F) Sole inhibition of HDAC8 by PCI34051 does not suppress the ESC-like module in hiPSC-derived cancer cells.

(G) Cell viability of glioblastoma cell lines treated with DMSO, PCI34051 (10  $\mu$ M), GSK126 (10  $\mu$ M) or the combination of PCI34051 and GSK126 (10  $\mu$ M each). \*p<0.05, \*\*\*p<0.001 (One-way ANOVA and Dunnett's multiple comparisons test).

#### Table S1: List of 110 target genes and their oligos for sgRNA, related to Figure 6

maintenance of ESC identity	Oligos for sgRNA	
hNTC	CACCCTGAAGGTGTCTGGCAGAGC	
mile	AAACGCTCTGCCAGACACCTTCAG	
LIN28B	CACCGCATCGACTGGAATATCCAAG	
EIIIZOB	AAACCTTGGATATTCCAGTCGATGC	
TCF3	CACCCACGCCTGCGTCTCTCCCGA	
1015	AAACTCGGGAGAGACGCAGGCGTG	
MADAKA	CACCTCAGGGTCGACATGTTAAAA	
MAI 4K4	AAACTTTTAACATGTCGACCCTGA	
	CACCACGGAAGGGGTCCCGAAACA	
GLUL	AAACTGTTTCGGGACCCCTTCCGT	
	CACCCTGATCATAAAACCCGTCCA	
ETV5	AAACTGGACGGGTTTTATGATCAG	
RND2		
MYCN		
	AAACAGGAGGACGCGTTCGGCCTG	
JARID2	CACCCTTCATGCCAGTCGACCCCC	
0111102	AAACGGGGGGTCGACTGGCATGAAG	
NIDBI	CACCGCTTGTCCATAGCCTCAACC	
NIPBL	AAACGGTTGAGGCTATGGACAAGC	
	CACCCACCGCAGCGCGGTTCGAAA	
KLF9	AAACTTTCGAACCGCGCTGCGGTG	
MYBL2		
	AACCIGUICGGCACAICIGAAI	
IFITMI		
	AAACGTTACIGGTATTCGGCTCTG	
APOC1	CACCTGGAGACGTCTGGGGTCCCC	
in oct	AAACGGGGACCCCAGACGTCTCCA	
UCK2	CACCGGACACGGTAGAAGCTATCC	
UCK2	AAACGGATAGCTTCTACCGTGTCC	
	CACCGGGCCTAGTCGGCATCATCG	
HLA-DPA1	AAACCGATGATGCCGACTAGGCCC	
EYAI		
	AAACACGGGICIIIAAACAAIIIC	
HTATIP2	CACCAGTCTTCCCGAAGCTTCGAC	
	AAACGTCGAAGCTTCGGGAAGACT	
PL 42G16	CACCATCATCGCTGCAAGCGTTGC	
1 LA2010	AAACGCAACGCTTGCAGCGATGAT	
	CACCCCGTGTATGCTTCGAGATGA	
HNRNPULI	AAACTCATCTCGAAGCATACACGG	
	CACCCGCCATCTACCATACGTTTG	
SLC15A1	AAACCAAACGTATGGTAGATGGCG	
ENAH		
MYBPH	CACCCCGTCAGACCTACATCCGCC	
	AAACGGCGGATGTAGGTCTGACGG	
IGFRP2	CACCGCTCCTTCATACCCGACTTG	
101/D1 2	AAACCAAGTCGGGTATGAAGGAGC	
6012	CACCATTATAAATACCGGCCCCGG	
SOX2	AAACCCGGGGGCCGGTATTTATAAT	
	CACCTTCTTACCAGTCTCCGTGTG	
NANOG	AAACCACACGGAGACTGGTAAGAA	
	CACCACTTGCAGGTGGTCCGAGTC	
POU5F1		
EYA2	CACCGCTIGTCGTACTCCACTTTG	
	AAACCAAAGTGGAGTACGACAAGC	
SETDS	CACCGACGGAGCGCCATGAAGTCCG	
SEIDO	AAACCGGACTTCATGGCGCTCCGTC	
	CACCGTGGCCGTCTGCGTGCGAGTG	
CENPE	AAACCACTCGCACGCAGACGGCCAC	
	CACCGCTCGGGCCACTCGTCACCTC	
CHAFIA		
CHAFIA	CACCGCTCGGGCCACTCGTCAGCTC AAACGAGCTGACGAGTGGCCCGAGC	
CHAF1A INCENP	CACCGCTCGGGCCACTCGTCAGCTC AAACGAGCTGACGAGTGGCCCGAGC CACCGAGGCCGAGCGCATGTTCACC	
CHAF1A INCENP	CACCGCTCGGGCCACTCGTCAGCTC AAACGAGCTGACGAGTGGCCCGAGC CACCGAGGCCGAGCGCATGTTCACC AAACGGTGAACATGCGCTCGGCCTC	
CHAF1A INCENP	CACCGCTCGGGCCACTCGTCAGCTC AAACGAGCTGACGAGTGGCCCGAGC CACCGAGGCCGAGCGCATGTTCACC AAACGGTGAACATGCGCTCGGCCTC CACCGAGGAAGCTCACCTTCGACG	
CHAF1A INCENP MSH6	CACCGCTCGGGCCACTCGTCAGCTC AAACGAGCTGACGAGTGGCCCGAGC CACCGAGGCCGAGCGCATGTTCACC AAACGGTGAACATGCGCTCGGCCTC CACCGAGGAAGCTCACCTTCGACG AAACCGTCGAAGGTGAGCTTCCTC	
CHAF1A INCENP MSH6	CACCGCTCGGGCCACTCGTCAGCTC AAACGAGCTGACGAGTGGCCCGAGC CACCGAGGCCGAGCGCATGTTCACC AAACGGTGAACATGCGCTCGGCCTC CACCGAGGAAGCTCACCTTCGACG AAACCGTCGAAGGTGAGCTTCCTC CACCGACCTACCTAGGCAATGCGT	
CHAF1A INCENP MSH6 FN1	CACCGCTCGGGCCACTCGTCAGCTC AAACGAGCTGACGAGTGGCCCGAGC CACCGAGGCCGAGCGCATGTTCACC AAACGGTGAACATGCGCTCGGCCTC CACCGAGGAAGCTCACCTTCGACG AAACCGTCGAAGGTGAGCTTCCTC CACCGACGTACCTAGGCAATGCGT AAACACGCCATGCCTAGGTAGGTC	
CHAF1A INCENP MSH6 FN1	CACCGCTCGGGCCACTCGTCAGCTC AAACGAGCTGACGAGTGGCCCGAGC CACCGAGGCCGAGCGCATGTTCACC AAACGGTGAACATGCGCTCGGCCTC CACCGAGGAAGCTCACCTTCGACG AAACCGTCGAAGGTGAGCTTCCTC CACCGACCTACCTAGGCAATGCGT AAACACGCATTGCCTAGGTAGGTC CACCGTAGCCCTCAGCATCGCACTC	
CHAF1A INCENP MSH6 FN1 ANP32A	CACCGCTCGGGCCACTCGTCAGCTC AAACGAGCTGACGAGTGGCCCGAGC CACCGAGGCCGAGCGCATGTTCACC AAACGGTGAACATGCGCTCGGCCTC CACCGAGGAAGCTCACCTTCGACG AAACCGTCGAAGGTGAGCTTCCTC CACCGACCTACCTAGGCAATGCGT AAACACGCATTGCCTAGGTAGGTC CACCGTAGCCCTCAGCATCCGAGTC	
CHAF1A INCENP MSH6 FN1 ANP32A	CACCGCTCGGGCCACTCGTCAGCTC AAACGAGCTGACGAGTGGCCCGAGC CACCGAGGCCGAGCGCATGTTCACC AAACGGTGAACATGCGCTCGGCCTC CACCGAGGAAGCTCACCTTCGACG AAACCGTCGAAGGTGAGCTTCCTC CACCGACCTACCTAGGCAATGCGT AAACACGCATTGCCTAGGTAGGTC CACCGTAGCCTCAGCATCCGAGTC AAACGACTCGGATGCTGAGGGCTAC	
CHAF1A INCENP MSH6 FN1 ANP32A NFKBIA	CACCGCTCGGGCCACTCGTCAGCTC AAACGAGCTGACGAGTGGCCCGAGC CACCGAGGCCGAGCGCATGTTCACC AAACGGTGAACATGCGCTCGGCCTC CACCGAGGAAGCTCACCTTCGACG AAACCGTCGAAGGTGAGCTTCCTC CACCGACCTACCTAGGCAATGCGT AAACACGCATTGCCTAGGTAGGTC CACCGTAGCCTCAGCATCCGAGTC AAACGACTCGGATGCTGAGGGCTAC CACCGCCTGAGCTCCGAGACTTTCG	
CHAF1A INCENP MSH6 FN1 ANP32A NFKBIA	CACCGCTCGGGCCACTCGTCAGCTC AAACGAGCTGACGAGTGGCCCGAGC CACCGAGGCCGAGCGCATGTTCACC AAACGGTGAACATGCGCTCGGCCTC CACCGAGGAAGCTCACCTTCGACG AAACCGTCGAAGGTGAGCTTCCTC CACCGACCTACCTAGGCAATGCGT AAACACGCATTGCCTAGGTAGGTC CACCGTAGCCCTCAGCATCCGAGGCTAC CACCGCCTGAGCTCCGAGACTTTCG AAACCGAAAGTCTCGGAGCTCAGGC	

UJA1	AAACTGCGACCAGTGGTGCGCTGAC
CRMP1	CACCGCTCCCGGAGAGCGCGACGAG
	CACCGTCACTCCAGCTTATACACC
DST	AAACGGTGTATAAGCTGGAGTGAC
CAMK2NI	CACCGACCAACAACTTCTTCGGCGC
CAIMIRZIVI	AAACGCGCCGAAGAAGTTGTTGGTC
IFITM2	CACCGCTCATCATCATCCAGTGT
SIX2	AAACCGTGCGCCGCAAATTCCCGC
D 4 V9	CACCGATGCGCTGGCGGACCACTTC
PAA8	AAACGAAGTGGTCCGCCAGCGCATC
PHOX2B	CACCGAGGTCCTTACCTGCGGCGTA
-	AAACTACGCCGCAGGTAAGGACCTC
IGF2	
	CACCGGCCCCTACCCGACGTGGTG
PAX2	AAACCACCACGTCGGGTAGGGGCC
LMOL	CACCGTGTGCCGACGCGACTACCTG
EMOI	AAACCAGGTAGTCGCGTCGGCACAC
CRABP2	CACCGCACAACGTCATCCGCCGTCA
	CACCGAACAATTATGAGCCCCGCAG
CIQB	AAACCTGCGGGGGCTCATAATTGTTC
	CACCGTCTACGCAGATTAATCATC
PRAME	AAACGATGATTAATCTGCGTAGAC
RGS4	CACCGACTTACATTCATGACTAATC
	AAACGATTAGTCATGAATGTAAGTC
GPR64	
RGS5	AAACCGTCCAGCGAGGTTCTACATC
	CACCGAACGTTGAGGGGCATCGTCG
MIC	AAACCGACGATGCCCCTCAACGTTC
HMGA1	CACCGGGCATCTCGCAGGAGTCCT
	AAACAGGACTCCTGCGAGATGCCC
HMGA2	
	CACCGGTCCCAATTAACCTAGCAA
EZH2_1	AAACTTGCTAGGTTAATTGGGACC
F7H2 2	CACCGTGGTGGATGCAACCCGCAA
	AAACTTGCGGGTTGCATCCACCAC
ARID1A	CACCGCAGCAGAACTCTCACGACCA
ARID4A	AAACACGGTATACCAACTAGCATC
CDV2	CACCGAGAGCCTGAAGAATTTGTCG
СВАЗ	AAACCGACAAATTCTTCAGGCTCTC
CDK8	CACCGAGGGCTGCAAAGTTGGCCG
	AAACCGGCCAACTTIGCAGCCCTC
CHD7	AAACACGTGCCATATAGCTGCCCAC
	CACCGACATTCTATTACCAAGAACA
EDFI	AAACTGTTCTTGGTAATAGAATGTC
FUBP1	CACCGTGATTGTAACAGGAACGGGC
	AAACGCCCGTTCCTGTTACAATCAC
H1FX	AAACCGGTGTAGATCTTGGCCAGC
	CACCGACAACTACTGACCTGCAAGC
H2AFZ	AAACGCTTGCAGGTCAGTAGTTGTC
MAX	CACCGTATTCCAGGAAGAGCAACCG
	AAACCGGTTGCTCTTCCTGGAATAC
MGA	LAUCGAUIGGAATCAACAACAATCG
	CACCGACATGACACGAATTGCCCTG
RIF1	AAACCAGGGCAATTCGTGTCATGTC
SETIDI	CACCGCCTTACCTGAATCAATACTG
SEIDBI	AAACCAGTATTGATTCAGGTAAGGC
SMAD2	CACCGCTAAATGTGTTACCATACCA
	AAACTGGTATGGTAACACATTTAGC
SMAD3	AACCTCCAATTCGGAGCGCTTCTC
	CACCGGGCGTACGAGTTTGACAAG
SMARCA4	AAACCTTGTCAAACTCGTACGCCC
SMARCCI	CACCGTGTTAAAGGAGGAACTGTAG
SMARCCI	AAACCTACAGTTCCTCCTTTAACAC

STAT2	CACCGACGCCGGTCTTGATGACGAG
STATS	AAACCTCGTCATCAAGACCGGCGTC
SUPTIAL	CACCGAATGTCAGTACCTTGATTG
50111011	AAACCAATCAAGGTACTGACATTC
TIMD	CACCGTTTATGGCAACCCTATCAAG
111/11/2	AAACCTTGATAGGGTTGCCATAAAC
ZNE136	CACCGAAAGATCACTACAAACACCG
2111130	AAACCGGTGTTTGTAGTGATCTTTC
711542	CACCGATGAGTGTAAGGTGCACAG
ZNF43	AAACCTGTGCACCTTACACTCATC
	CACCGCCAGTAATAGTACTCTCTCG
MEDI	AAACCGAGAGAGTACTATTACTGGC
	CACCGTCTGTTCAGACTCTAATAGG
RAD21_1	AAACCCTATTAGAGTCTGAACAGAC
	CACCGTGTAATTTAGAGAGCAGCG
RAD21_2	
CBXI	
CHD1	
	AAACCGTCLCTTAGGCGAATTAAC
YYI	
	AAACCGGTGACCAGCGTTTGTTCAC
CDC5L	
	AAACCGGCAGTTTGACGTGCAATTC
DDX47	CACCGICCICTATTGGAGTGCAGAG
	AAACCICIGCACTCCAATAGAGGAC
DHX15	CACCGTGGGTCTGGTAAAACAACAC
	AAACGTGTTGTTTTACCAGACCCAC
LUC7L3	CACCGTAGAACGTAGGATCAGACG
	AAACCGTCTGATCCTACGTTCTAC
MCM6	CACCGCTGCGGGGGGATACTCCCTCG
шешо	AAACCGAGGGAGTATCCCCCGCAGC
MCM7	CACCGAAGCCCAACTACTTACTGCG
	AAACCGCAGTAAGTAGTTGGGCTTC
RTFI	CACCGTGTCCGAGTCGATCACGACG
	AAACCGTCGTGATCGACTCGGACAC
SMC1A	CACCGATTGCCACAAGCATACTGCA
	AAACTGCAGTATGCTTGTGGCAATC
SMC4	CACCGCATTGATGAAAAATACGACG
	AAACCGTCGTATTTTTCATCAATGC
SNRPD2	CACCGACCCAGCACTCACCTATCGA
	AAACICGATAGGIGAGIGCIGGGIC
SSRP1	
	AAACIIGICAACIIIGCICGIGGIC
THAP11	
	AAACGCGTCAGTGGGTGCTTCTCC
TPR	CACCGACCIGGGAIGAAACIAACIG
	AAACCAGIIAGIIICAICCCAGGIC
WDR82	
	AAACICICGCCGTIGGGGCIGAAAC
SUV39H2	
TRIM28	
	AAACCAGIGGIACIGCCIGGGAAGC
EIF4A3	
EED	
	AAACIGIAIGAAIGICAIICACAAC
SOX11	
FOXM1	
E2F7	AAACCTAGTGTATGTCCCCTCTCC
ETV1	
TCF4	
BCL11A	A A A C C G C G G G T T G G T A T C C C T T C A C
WHSC1	
	CACCGACACCTTCGCCTAGTCACAC
KDM5B	AAACGTGTGACTAGGCGAAGGTGTC
	CACCGTGTGGGAGAGTATCCATGGCG
MTA1	
	In an electron in the reacted at ACAC

## Table S2: References for picking out 110 target genes, related to Figure 6

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#### Table S3: Results of genetic screening with lentiviral CRISPR/Cas9 system, related to Figure 6

hiPSO

Relative growth ratio compared to contorol 0.47371321 0.630024575884 0.630024575 0.689734173 0.721125506 0.737204194 0.743813879 0.74718664 0.764182888 -derived cancer cells 
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 Gene Name

 1 FNI

 2 [FITM2

 3 PP&ME

 4 SETD8

 5 SNRPD2

 6 SMC4

 7 CLQB

 8 TCF3

 9 EZH2

 10 MVCN

 11 EIF4A
 SD 0.064243312 0.079338947 0.104260527 0.032886971 0.067667121 0.130264191 0.027325322 0.017689598 0.031221687 0.049478903 ink 0.049226372 0.01162078 0.01262078 0.07100474 0.03822440 0.03822440 0.0382040 0.0382040 0.038205 0.05173804 0.035505960 0.05181753 0.06088300 0.051817538 0.051817538 0.0518533 0.0518533 0.0518533 0.0518533 0.0518533 0.0518533 0.0228240 0.03883153 0.0518533 0.0518533 0.0518533 0.0518533 0.0228240 0.03288353 0.047308820 0.014730842 0.014730842 0.014730842 0.014730842 0.014730842 0.014730842 0.014730842 0.014730842 0.014730842 0.014730842 0.014730842 0.014730842 0.014730842 0.014730842 0.014730842 0.01473084 0.014730842 0.014730842 0.014730842 0.014730842 0.014750 11 EFF-43 12 EFF-43 13 MAP4K4 14 GUU 15 JARD2 16 ETV3 16 ETV3 17 MYBL2 18 UCR2 19 APOCI 20 BCL11A 20 BCL11A 21 KLF9 22 RAD21 21 KLF9 22 RAD21 23 HNRNPULI 24 HLA-DPA1 25 CAMK2NI 26 EXA2 27 DHX15 28 MYBPH 26 GAKA2NI 26 EXA2 27 DHX15 28 MYBPH 27 DHX15 28 GAKA2NI 28 CAKA2NI 29 CAKA2NI 20 CAK 0.93434419 0.93449140 0.015416376 0.039180125 0.0282592 0.033700882 0.033700882 0.031203097 0.037066058 0.0314203078 0.031497718 0.039588074 0.008702249 0.018531997 0.002767853 0.030467343 0.93957937 0.94784047 0.956009234 0.958993037 0.96444836 0.964944735 49 SU/39/E 50 LUC7L3 51 EYA1 52 SSRP1 53 E2F7 54 RTF1 55 PAX2 56 ANP32A 57 LIN28B 58 hNTC 59 DST 60 GJA1 61 MCM7 62 PAX8 0.964944733 0.967701384 0.973212321 0.9843569955 0.992455009 0.99305148 0.9966289055 0.997706827 0.03228208 0.044369644 0.048744042 0.038391653 0.054704041 0.052812528 0.048452086 0.053726003 0.024087719 0.130331852 0.020890819 0.014895802 0.022904104 0.092297571 0.037264447 0.030482975 1.001685756 1.001685/56 1.002986873 1.009580956 1.011599431 1.015925545 1.018768979 1.019569065 62 PAX8 63 TRIM28 64 IGF2 65 CHAF1A 1.019569065 1.019614249 1.022850516 1.025198368 1.026718922 1.027048617 1.028057096 1.030018251 1.030509761 65 CHAFL 66 DDX47 67 IGFBP2 68 MCM6 69 ZNF43 70 LMO1 71 CBX1 72 MTA1 73 GPR64 74 CRMP1 75 CBX3 76 SMAD3 1.032984939 0.06951402 0.0245984 
 10
 SixtaD3

 77
 RCS4

 77
 RCS4

 78
 SUP16H

 79
 PHOX2B

 80
 MAX

 81
 CDK8

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 SMARC44

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 SMARC44

 84
 ARID4A

 85
 NFKBBA

 86
 H7LA78

 87
 MGA

 88
 SMARC41

 89
 EED

 90
 TIMP2

 91
 SMAD2

 92
 H1FX

 93
 CHD7

 94
 YV1

 95
 FCBP1

 96
 MSH6

 97
 MED1

 98
 EDF1

 100
 ZKF136

 94
 YV1

 95
 FCBP1

 100
 ZKF136

 101
 ARD1A

 102
 SETDB1

 103
 REF1

 104
 WD642

 105
 HIMG42
 0.027596984 0.064922005 0.011919653 0.014977001 0.01553768 0.034368434 0.007792492 0.040821165 0.011797434 0.040821165 0.011797434 0.031733782 0.033288636 0.018569008 0.018569008 0.018569008 0.018569008 0.018690098 0.02847620 0.037121991 0.039026623 0.022879782 0.022879782 0.02327629 0.022879782 0.022879782 0.022879782 0.022879782 0.022879782 0.022879782 0.022879782 0.022879782 0.022879782 0.022879782 0.022879782 0.022879782 0.022879782 0.022879782 0.022879782 0.022879782 0.02288743 0.02288743 0.02288743 0.02288743 0.0259886743 1.035063859 1.035576599 1.039592187 1.047577482 1.049063812 1.0503878 1.050994915 1.055854942 1.057425882 1.057425882 1.057981618 1.060835877 1.061182685 1.063659763 1.065043176 1.066063103 1.070264506 1.072841563 1.0706817451 1.079569319 1.080830931 1.081696089 1.080830931 1.081696085 1.097654637 1.109372871 1.110216144 1.117405554 1.120265621 1.126121047 1.127288957 1.141975854 1.148405364 1.150264716 0.051350721 0.028086986 0.031637066 0.02488206 0.010969745 0.046816292 0.046444506 0.020486602 0.035342853 0.008364431 1.162237302 1.219780016 0.00836443 0.12850196 109 ETV1 110 HTATI 111 KDM51 1.271766098 0.278748792

Rank	Gene Nama	Relative growth ratio	SD
Kank 1	EIF4A3	0.169565387	0.00980538
2	INCENP DDX47	0.527787752	0.01336574
4	MCM7	0.66923558	0.01685298
5	WDR82 LUC7L3	0.675463706	0.0371681
7	PHOX2B	0.68995379	0.00892269
8	CDC5L	0.693336429	0.00567363
10	CHAF1A	0.702607795	0.05587078
11	SSRP1	0.716823563	0.03671914
12	ETA2 TPR	0.737353215	0.0720527
14	RAD21	0.747631663	0.12126180
15	DHX15 CIOB	0.755507112 0.765545571	0.07577447 0.03731176
17	MCM6	0.770667755	0.03456806
18	DST NANOG	0.779968272	0.04649380
20	RTFI	0.788666151	0.01807272
21	SMC1A MYC	0.804984617	0.07338099
23	NFKBIA	0.814613849	0.07255601
24	IFITM2	0.814791199	0.02250499
25	MYBL2	0.82054/8/4	0.05093220
27	HMGAI	0.823551052	0.01930168
28	POU5F1 CRABP2	0.834678327	0.0793396
30	THAP11	0.840812994	0.02167771
31	TRIM28 SUPT16H	0.840892971	0.09766716
32	EED	0.845130685	0.04016357
34	LIN28B CAMK2N1	0.85108255	0.13512832
35	CENPE	0.875624545	0.05464093
37	SETD8	0.877680583	0.05741292
38	CHD1	0.8/932/529 0.89024884	0.05359496
40	TIMP2	0.910001826	0.01917063
41 42	SUV39H2 IGF2	0.911920696 0.912607631	0.06810177
43	IGFBP2	0.925028892	0.09329476
44	NIPBL EDF1	0.926839353	0.06938949
46	CHD7	0.958289333	0.03123068
47	PAX2 SMARCAA	0.961687727	0.02620402
49	FUBP1	0.964785045	0.02982196
50	SMC4	0.969079091	0.06977141
51	MAX	0.970574745	0.02272796
53	LMO1	0.973947123	0.03027747
54	RIFT SMAD3	0.978593715	0.07460903
56	PAX8	0.992282333	0.05922396
57	UCK2 SMARCC1	0.997291056	0.03744133
59	PRAME	0.999415342	0.04570719
60	hNTC SFTDR1	1 003438736	0.06977284
62	EZH2	1.004972622	0.03126583
63	CRMP1	1.008382645	0.02459358
65	ANP32A	1.010391678	0.03376189
66	SIX2	1.010971216	0.03382673
67	SMAD2	1.015686882	0.01210773
69	FNI	1.019482023	0.04311387
70	MGA CDK8	1.032093197 1.033383322	0.0534093
71	GJAI	1.035622325	0.06518786
73	MED I ZNF43	1.052961765	0.02386793
75	CBXI	1.060030611	0.08945915
76	PLA2G16 RGS4	1.060361987	0.01615454
78	MSH6_2	1.061137934	0.04083514
79	ARID4A	1.065230337	0.02824208
80	TCF3	1.0/4440/3/ 1.085152051	0.05554078
82	H2AFZ	1.088555744	0.02590183
83	CNF150 GPR64	1.098326367	0.03247323
85	HMGA2	1.10477116	0.04686357
86	STAT3 FOXMI	1.106066933	0.0397636
88	APOCI	1.201390586	0.06197617
89	HNRNPUL1 IFITM1	1.211376924	0.0313003
90 91	SOX11	1.265689384	0.04995011
92	BCL11A TCEA	1.269249166	0.07263045
93 94	SLC15A1	1.290443945 1.329457088	0.01384482
95	MAP4K4	1.345945572	0.12185391
96	WHSCI JARID2	1.358066892 1.392056741	0.02166840
97	MTAI	1.411135363	0.13029318
99	MYCN GLUI	1.441390666	0.04671890
100	E2F7	1.4420/40/1 1.45236403	0.05813790
102	ENAH	1.512990909	0.04171505
103	KDM5B MYBPH	1.518139597 1.579735763	0.05118251 0.05401290
104	ETV5	1.620459123	0.10836600
106	HLA-DPA1	1.744922707	0.15809322
107	RND2	1./5509556/ 1.80329171	0.16343047
109	HTATIP2	2.087640063	0.18659896
110	KLF9 EYAI	2.333761653 2.53060758	0.17499401 0.22459075

Rank		Gene Name	Relative growth ratio compared to contorol	SD
	1	MYCN	0.30760042	0.00668181
	2	SUPT16H EIF4A3	0.348697118 0.41083802	0.04708912
	4	SETDB1	0.525616952	0.05812066
	6	RAD21	0.673008303	0.004/4242
	7	TIMP2 CDC51	0.698883957	0.04764318
	9	LUC7L3	0.768067499	0.08131742
	10	EZH2 SMC14	0.797878906	0.091969870
	12	MCM6	0.807855081	0.04468380
	13	YYI SNPPD2	0.819126043	0.02252361
	14	RTF1	0.8249977130	0.08463342
	16	THAP11	0.839345055	0.04594584
	18	EED	0.850379696	0.06095225
	19	DHX15	0.859217002	0.05165476
	20	TPR	0.872510077	0.01491935
	22	MCM7	0.891909855	0.09569188
	23	WDR82 MYBL2	0.90256298	0.02017487
	25	SMC4	0.920549927	0.07157945
	26	EDF1	0.932464322	0.03225453
	28	LIN28B	0.934563892	0.02097685
	29	ANP32A MGA	0.940729803 0.94145195	0.03243555 0.12311429
	31	TRIM28	0.950994798	0.02313490
	32	HMGA1 SMAD3	0.966226807	0.10418342
	34	CHDI	0.975064385	0.1010516
	35	ZNF136 hNTC	0.998032135	0.34927862
	37	GJAI	1.000052694	0.02294124
-	38	CENPE PAY8	1.010773235	0.01790157
	39 40	SMAD2	1.012416037	0.04516802
	41	PHOX2B	1.015792563	0.02596021
	42	MAX	1.018297675	0.03383061
	44	IGF2	1.038329382	0.09435346
	45	SIX2 UCK2	1.043/9/056	0.029/1844
	47	CBX3	1.049024649	0.12517310
	48	SSRP1 CHD7	1.056972145	0.05792007
	50	SUV39H2	1.08837556	0.03453856
	51	PAX2 SOV2	1.090052358	0.01364757
	53	MSH6_2	1.098389799	0.04754616
	54	RIF1 CDK8	1.103591236	0.17066794
	56	HMGA2	1.121950403	0.13267212
	57	RGS5 CBV1	1.124248181	0.03860409
	59	FOXMI	1.128212499	0.02437302
	60	TCF4	1.147873706	0.04417714
	62	BCL11A	1.153907184	0.04543775
	63	LMOI	1.154653788	0.09179200
	64	MIAI EYA2	1.155682319	0.07154952
	66	CHAFIA	1.182820702	0.02351364
	67 68	GPR64 IGFBP2	1.189148551 1.200468476	0.0118783 0.06772624
	69	POU5F1	1.207771613	0.12241312
	70	ZNF43 MYC	1.208109021 1.211936704	0.38761077 0.06492653
	72	NIPBL	1.225542162	0.02407846
	73	DST H2AFZ	1.227785041	0.088312
	75	E2F7	1.240871988	0.08289953
_	76	CIQB NANOC	1.250473319	0.0924942
	78	NFKBIA	1.259577244	0.04585829
	79	CAMK2NI	1.268463505	0.02181439
	80	SLCIŠAI MEDI	1.278713428 1.303813441	0.06188193 0.02292317
	82	ARIDIA	1.305570971	0.22102712
	83 84	WHSCI KLF9	1.307680009 1.314040001	0.03663004
	85	HIFX	1.316887364	0.05993977
	86	SMARCA4 SOX11	1.329257585	0.55785237 0.10778225
	88	APOCI	1.335807366	0.06517905
-	89	GLUL SMARCC1	1.350770148	0.09038867
	90	IFITM2	1.361292041	0.02981560
	92	CRABP2	1.38470777	0.23103817
	93 94	MYBPH	1.400569057	0.14934736
	95	PLA2G16	1.412178723	0.06900855
	96 97	ARID4A RGS4	1.415399709 1.424994737	0.05916976
	98	HLA-DPA1	1.425461723	0.06816078
	99	KDM5B ENAH	1.428454545 1.49774191	0.05464664
	101	FNI	1.507854005	0.05536446
	102	IFITM1 IARID2	1.593387131	0.02117667
	103	RND2	1.623044227	0.09377848
	105	ETV5	1.652729228	0.41800659
	106	ETVI	1.792898283 1.823783525	0.20789986
	108	TCF3	1.850068304	0.05694204
	109	MAP4K4 PRAME	1.890700021 1.930211441	0.08746465
	111	HTATIP2	1.965732453	0.16430709

#### Table S4: Primer list, related to STAR Methods

Genes	Forward	Reverse	
Quantitative PCR			
GAPDH	ATGGGGAAGGTGAAGGTCG	GGGGTCATTGATGGCAACAATA	
NANOG	TGAGATGCCTCACACGGAGA	GTTTGCCTTTGGGACTGGTG	
OCT4	CGTGAAGCTGGAGAAGGAGAAGCTG	CAAGGGCCGCAGCTTACACATGTTC	
NESTIN	CAGCTGGCGCACCTCAAGATG	AGGGAAGTTGGGCTCAGGACTGG	
NCAMI	GAGGAGGAGAGGACCCCAAA	CTGGCACTCTGGCTTTGCTT	
PAX6	CAGCTCGGTGGTGTCTTTGTCA	CTGCGCCCATCTGTTGCTTTTC	
SALL4	CAACATTTGTGGGCGAGCTT	GCGGGCTGAGTTATTGTTCG	
LIN28B	GTGGGAGCCCCTGTTTAGGA	CATGATGATCAAGGCCACCA	
SOX2	CGGAAAACCAAGACGCTCA	GCCGTTCATGTAGGTCTGCG	
c-MYC	CCTGGTGCTCCATGAGGAGA	GCCTGCCTCTTTTCCACAGA	
MAP2	AGAGGGTGCCTTTGGAGAGC	TGCAGACACCTCCTCTGCTG	
TUJI	GGCCAAGGGTCACTACACG	GCAGTCGCAGTTTTCACACTC	
CDKN2A	GTGGACCTGGCTGAGGAG	CTTTCAATCGGGGATGTCTG	
E2F1	TCCAAGAACCACATCCAGTG	CTGGGTCAACCCCTCAAG	
CCND1	GAAGATCGTCGCCACCTG	GACCTCCTCCTCGCACTTCT	
GFAP	CGTGCAGACCTTCTCCAACC	GCATCTCCACGGTCTTCACC	
Quantitative PCR for transduction of			
pMxs-OCT4, SOX2, KLF4 and c-MYC			
c-MYC	ATACATCCTGTCCGTCCAAGCAGA	GACATGGCCTGCCCGGTTATTATT	
SOX2	TTCACATGTCCCAGCACTACCAGA	GACATGGCCTGCCCGGTTATTATT	
KLF4	CCACCTCGCCTTACACATGAAGA	GACATGGCCTGCCCGGTTATTATT	
OCT4	GCTCTCCCATGCATTCAAACTGA	CTTACGCGAAATACGGGCAGACA	
Direct sequencing			
SMARCB1	CTCGCTGACTGTTGCTTCCA	ACGGGACTGTTCCCACGTAA	
TP53	TGTGCAGCTGTGGGTTGATT	CGCAAATTTCCTTCCACTCG	