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MYB-activated models for testing therapeutic agents in adenoid cystic carcinoma

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Abstract

Objective: There are no effective systemic therapies for adenoid cystic cancer (ACC) and lack of tumor lines and mouse models have hindered drug development. We aim to develop MYB-activated models for testing new therapeutic agents.

Materials and Methods: We studied new ACC patient-derived xenograft (PDX) models and generated a matched cell line from one patient. In addition, we generated a genetically-engineered *MYB-NFIB* mouse model (GEMM) that was crossed with $Ink4a^{+/-}/Arf^{+/-}$ mice to study tumor spectrum and obtain tumor lines. Using human and murine ACC-like tumor lines, we analyzed *MYB* expression by RNA-Seq and immunoblot and tested efficacy of new MYB inhibitors.

Results: We detected *MYB-NFIB* transcripts in both UFH1 and UFH2 PDX and observed tumor inhibition by MYB depletion using shRNA *in vivo*. We observed rapid loss of MYB expression when we cultured UFH1 *in vitro*, but were able to generate a UFH2 tumor cell line that retained MYB expression for 6 months. RNA-Seq expression detected an ACC-like mRNA signature in PDX samples and we confirmed an identical KMT2A/MLL variant in UFH2 PDX, matched cell

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line, and primary biopsy. Although the predominant phenotype of the *MYB-NFIB* GEMM was B-cell leukemia, we also generated a MYB-activated ACC-like mammary tumor cell line. We observed tumor inhibition using a novel MYB peptidomimetic in both human and murine tumor models.

Conclusions: We generated and studied new murine and human MYB-activated tumor samples and detected growth inhibition with MYB peptidomimetics. These data provide tools to define treatment strategies for patients with advanced MYB-activated ACC.

Keywords

adenoid cystic cancer; MYB gene; peptidomimetic inhibitor; B-cell leukemia

Introduction

Management of malignant salivary gland tumors is a challenge for clinicians due to lack of effective therapy for patients with unresectable disease and also due to the large number of distinct clinico-histologic subtypes that must be studied separately as unique biologic entities. Adenoid cystic cancer (ACC) is the most common malignant tumor arising from major or minor salivary glands and can also present as primary tumors within breast, lung, skin, and other tissues (1–7). The identification of a recurrent *MYB-NFIB* fusion oncogene in most ACC cases provided a new tool for diagnosis and insight into a molecular pathogenesis for this malignancy. This chromosomal rearrangement typically truncates carboxy-terminal MYB domains while fusing the terminal NFIB exon encoding a long 3' untranslated region (8). The detection of MYBL1 fusions in cases of MYB fusion–negative ACC confirmed importance of MYB/MYBL1 activation as the signature oncogene driving pathogenesis of ACC (9–12). Further, there is evidence suggesting a role for the translocated NFIB locus to redirect a 3' super-enhancer to generate positive feedback activation sustaining deregulated MYB expression (13).

An obstacle to study the role of MYB activation in ACC and to develop effective MYB inhibitor compounds as therapeutic agents is the lack of validated human or mouse tumor cell lines. Although ACC PDX models have been studied, these are cumbersome for drug discovery projects (14–16). Until recently, there were no validated MYB-activated tumor cells adapted for *in vitro* growth to allow drug development screening and, currently there is only one report of a human cell line with MYB activation (17).

Methods and material:

Peptidomimetics.—The MYBMIM peptidomimetic and its TG3 inactive analogue were produced by solid phase synthesis and validated by mass spectrometry as previously published (24). Peptidomimetics were dissolved in PBS as previously established (24).

Cell viability analysis.— $5x10^3$ cells were seeded in 96-well plates overnight and treated with indicated compounds for 48 hours prior to addition of 40 µL 2 mg/mL MTT reagent (Promega) in PBS for 4 hours, and analyzed using SpectraMax M5 microplate reader (Molecular Devices, Sunnyvale, CA).

Cell line generation and culture conditions.—Patient tumor biopsies were minced and incubated with HBSS digestion solution (Thermo Fisher Scientific Inc), HEPES (Mediatech, Inc), CaCl₂, FBS, DNase I(Sigma Aldrich) and collagenase A(Sigma Aldrich), at 37°C for 60 min. Cell suspensions were obtained by passing mixture through 40-µm sieve (Fisher Scientific, Pittsburgh, PA). Washed cells were cultured in high-glucose Dulbecco's modification of Eagle medium (DMEM; Invitrogen), supplemented with Penicillin-Streptomycin (Sigma-Aldrich), 10% Fetal Bovine Serum (Invitrogen).

H&E and Immunohistochemistry.—The tissue specimens were fixed with 10% formalin, dehydrated using ethanol, and paraffin-embedded sections obtained. Images were taken by Leica DM6000B fluorescence microscope.

Generation of MYB-NFIB transgenic mice.—*MYB* exons 1-14 was fused with NFIB exon 12 and subcloned into Lox-stop-Lox conditional pCBR-based plasmid (Promega) and confirmed by nucleotide sequencing. Transgenic *MYB-NFIB* injection was performed using FVB mice. MMTV-Cre FVB mice and INK4/ARF knockout B6 mice were crossed with *MYB-NFIB* fusion mice to generate *MYB-NFIB/MMTV-Cre* and *MYB-NFIB/MMTV-CRE/Ink4a^{+/-}Arf^{+/-}* mice. Primer sequences used to identify the MYB-NFIB fusion; forward primer pCBR-BN-376F: 5'-CAGATGTGCAGTGCCAGCACCGATG-3'; reverse primer pCBR-BN-1268R: 5'-TGGCGAGGCGCTTTCTTCAGGTAGG-3'.

Flow cytometric analysis.—Peripheral blood (PB) was collected from indicated mice genotypes and subjected to an automated blood count (Element HT5® Hematology Analyzer, HESK, Colorado, USA). After lysing RBCs, total mononuclear cells were stained with fluorochrome-conjugated antibodies including B220 (PE-Cy7), IgM (APC-Cy7), CD3 (FITC), CD4 (PE), CD8 (Pacific Blue), CD11b (PE-Cy7) and Gr.1 (APC) for 45 minutes at 4°C. Cells were washed with PBS and acquired on a LSR Fortessa flow cytometer (BD Biosciences). All data were analyzed by FlowJo-V10 software (BD Biosciences).

Lentivirus production and cell transduction.—3 x 10^{6} 293T cells were seeded in 10cm dishes overnight and transfected with pLKO.1 vector as recommended (Qiagen, Germantown, MD). X-tremeGENETM HP DNA Transfection Reagent (Sigma Aldrich, St. Louis, MO) was used for transfection and viral supernatants were used to infect target cells in presence of 8 µg/mL polybrene (Sigma Aldrich, St. Louis, MO). shRNA clones TRCN0000040058, TRCN0000040059, TRCN0000040060 from TRC-Hs1.0 (Human) (Horizon Discovery Group Co) were used as indicated.

Xenograft Tumor Studies.— $2x10^{6}$ PDX tumor single cells transduced with shRNA vector control or MYB-shRNA lentivirus were mixed with 200 µl Matrigel (BD Biosciences, Franklin Lakes, NJ) and subcutaneously injected in 6-week-old NOD.CB17-Prkdcscid/ NcrCrl (NOD/SCID) (Jackson Laboratory, Bar Harbor, ME). Control and treatment groups were matched by sex and age. Tumors were measured every other week with a dial caliper, and tumor volumes were calculated using the formula: tumor volume= (width² × length)/2.

MYB dual color breakapart fluorescence in situ hybridization (FISH)

Four-micron paraffin-embedded sections were analyzed with MYB breakapart probe (ZytoVision, Bremerhaven, Germany). Using a Vysis VP2000TM slide pretreatment instrument and reagents, slides were mounted with probe solution, denatured at 80°C and hybridized for 24 hours at 37°C. (Abbott, Abbott Park, Illinois). Slides were washed, counterstained with DAPI and analyzed with a Leica DM6000B microscope and Ariol Software v4.0 (LeicaBiosystems, Wetzlar, Germany).

RNAseq Data Processing and Analysis.—Sequencing reads were mapped to the human transcriptome (hg 19) using bowtie2 (38), and gene expression levels were summarized into TPM values from uniquely mapped reads using RSEM (39). Clustered heatmaps of gene expression were generated with R package 'heatmap3' (40) based on a set of previously defined gene signature that was highly expressed in ACC tumors (21). We then applied single-sample gene get variation analysis (ssGSVA) (41) to determine the molecular phenotypes of each sample using log2(TPM+1) data. We obtained GSVA scores for hallmark gene sets or canonic pathways (MSiDB hallmark H set and canonic pathway C2 set) (42) for each sample. Comparison of GSVA scores was conducted using R package 'limma' (43). *MYB-NFIB* gene fusion detection was performed using Star-Fusion (44).

Statistical Methods

Statistical analyses of experimental data were carried out using Prism Graphpad 6.0. The two-sided t test was used to analyze data from cell viability and xenograft assays. Data were presented as mean and standard deviation (\pm SD). The log-rank test was used to determine the P value in the survival difference between *MMTV*-Cre/*MYB-NFIB* and *MMTV*-Cre/*MYB-NFIB*/Ink4a^{+/-}/Arf^{+/-} groups in Kaplan-Meier survival analysis. All statistical tests were two-sided and P values were considered statistically significant when not greater than .05.

Ethics statement.—IACUC guidelines and regulations followed U.S. National Institute of Health guidelines, U.S. Public Health Service policy, and U.S. Food and Drug Administration regulations to minimize stress and pain on research animals.

Results:

Characterization of human ACC tumor models

To develop new ACC tumor models, we obtained informed consent to collect excess ACC tumor from two patients undergoing medically indicated procedures using an IRB-approved protocol. Although we are able to maintain UFH1 ACC as a PDX model (UFH1PDX) by passaging through immunodeficient SCID mice, we were unable to adapt tumor cells to *in vitro* growth. This observation has been noted in previous reports where ACC tumor samples can only be studied *in vitro* as short-term cultures (8). Interestingly, while UFH1 tumor cells could be maintained by passaged through SCID mice, these tumor cells remained viable without proliferation in cell culture *in vitro* for > 6 months with no passaging and infrequent changing of media (data not shown). We tested *MYB* mRNA and protein expression from UFH1 cells immediately after excision from a PDX xenograft and plating onto plastic tissue

culture dishes (5-hour time period to allow for attachment) and compared this to MYB expression after 24 hours in cell culture. We observed loss of MYB mRNA by quantitative reverse transcription polymerase chain reaction (RT-PCR) and protein expression within 24 hours that correlated with loss of tumor cell growth in vitro (Supplemental Figure 1a). The mechanism for this MYB inactivation is unknown and we were unable to rescue this by supplementing with conditioned media, ROCK inhibitor, or murine feeder layers (36). We studied a second tumor sample from a palliative debulking procedure in a patient with locally recurrent and metastatic parotid gland ACC. The patient required multiple resections and post-operative photon radiation therapy and salvage palliative proton beam radiation therapy for unresectable recurrent tumor over several years. The patient presented with recurrent tumor (Fig. 1a) and lung metastases that progressed despite dual nivolumab/ ipilumumab therapy. A biopsy sample of the recurrent tumor showed poorly differentiated ACC with elevated nuclear MYB expression (Fig. 1b). A complex MYB-NFIB rearrangement with 3' MYB duplication was detected by hybrid-capture technology on an earlier tumor biopsy (Foundation Medicine, Cambridge, MA). Tumor cells from the biopsy were cultured *in vitro* and the UFH2 cell line has been maintained using high-glucose DMEM, supplemented with 1% Penicillin-Streptomycin antibiotic and 10% FBS with a doubling time of 36 hours (Fig. 1c). UFH2 identity was matched to the patient using CLIAcertified NGS assay sequencing performed on genomic DNA extracted from UFH2 cell line with detection of both ATRX E1433*frameshift and MLL/KTM2A S1325N variants detected in earlier tumor surgical sample (data not shown). To test for MYB expression, we performed protein immunoblot analysis using a control ACCX22 PDX (gift from Christopher Moskaluk, University of Virginia), the new UFH1 and UFH2 PDX models, the matched UFH2 cell line (in vitro passage 9 and late passage 34), and a control human mucoepidermoid tumor cell line (H3118). We observed elevated MYB expression in the PDX ACC samples and the matched UFH2 cell line (passage 9) but absence of MYB at passage 34 (Fig. 1d). Multiple MYB bands seen on protein immunoblot have been attributed to alternate splicing or protein processing (10) and the non-ACC salivary gland tumor (H3318) showed absent MYB expression. The early and late passage UFH2 tumor cell line were tumorigenic in SCID mice and the early passage excised xenograft showed poorly differentiated, solid histology on H&E staining similar to the primary tumor surgical biopsy (Fig. 1e). In summary, tumor cells obtained from UFH1 and UFH2 ACC patients exhibited distinct growth properties in vitro, perhaps suggesting that the ability to generate the UFH2 line was associated with the undifferentiated histology. Despite this difference in growth pattern, we confirmed MYB rearrangements by fluorescent in situ hybridization using breakapart probes in sections from both UFH1 and UFH2 PDX (data not shown).

RNA-Seq analysis of new ACC PDX models

We also extracted RNA from UFH1 PDX and UFH2 PDX and from the late passage 34 UFH2 cell line for RNA-Seq analysis. We confirmed detection of distinct *MYB-NFIB* fusion transcripts in both UFH1 PDX (*MYB* exon14 fused to either *NFIB* exon10 or exon11) and UFH2 PDX (*MYB* exon14 fused to either *NIFIB* exon11 or exon12) (Fig. 2a) but not in the late passage UFH2 tumor cell line. We compared RNA-Seq expression pattern of UFH1 and UFH2 with gene expression signature previously identified in a microarray study of 12 primary ACC tumors with matched adjacent normal tissues (21). While UFH1

and UFH2 were collected from recurrent metastatic disease compared to the microarray gene signature primarily collected from primary ACC cases, we noted clustering of UFH1, UFH2 PDX tumor samples, as well as the late passage UFH2 cell line, with similar expression patterns within the ACC signature (Fig 2b).

ACC mouse model development

To further study ACC biology and investigate new drug development, we also generated a genetically-engineered mouse model (GEMM) with a conditional LoxP-Stop-LoxP MYB-NFIB transgene (Fig. 3a). MYB-NFIB mice were crossed with MMTV-Cre mice to obtain bi-allelic mice that express MYB-NFIB in secretory epithelium of virgin and lactating mammary gland, the salivary gland, seminal vesicle, skin, erythrocytes, B cells and T cells (Fig. 3b) (18). To test Cre activation of MYB-NFIB transgene, HEK293 cells were transfected with the MYB-NFIB pCBR-based transgene construct alone or co-transfected with Adeno-Cre. We confirmed by immunoblotting that the fusion transgene was activated only in presence of Adeno-Cre (Fig. 3c). To verify fusion transgene expression in mice crossed with MMTV-Cre, we performed RT-PCR and immunoblot on samples collected from murine tissues. Although, we detected expression of the transgene in whole salivary gland tissues by RT-PCR (Fig. 3d), we did not observe MYB expression by protein immunoblot in extracts from whole normal submandibular glands (n=9 mice) (data not shown). In addition, we did not detect epithelial salivary gland tumors in the GEMM (n=42 mice), however there was shorter overall survival compared to historical controls (37) due to development of a B-cell neoplasm in majority of mice (see below).

MYB-activated mammary breast tumor generated in transgenic mice

Since the p16 pathway is reported to undergo inactivation in a subset of salivary gland cancers including ACC (20, 21,22), we also crossed *MYB-NFIB/MMTV-CRE* transgenic mice with *Ink4a^{+/-}/Arf^{+/-}* mice and generated tri-allelic mice that expressed heterozygous Ink4a^{+/-}/Arf^{+/-} alleles. Within *MYB-NFIB/MMTV/Ink4a^{+/-}/Arf^{+/-}* cohort (n=20 mice) we detected a large mammary tumor with cribriform ACC-like adenocarcinoma histology at 6 months of age (Fig. 3e). We generated a murine tumor cell line from this breast tumor (UFBT) and observed elevated steady-state MYB protein expression (Fig 3f). RT-PCR and nucleotide sequencing confirmed expression of the human *MYB-NFIB* transgene (data not shown) suggesting this tumor arose as a MYB-dependent event. Although, we did not detect salivary gland tumors in the *MYB-NFIB/Ink4a/Arf^{+/-}* mice (Fig. 3g), we again observed predominantly B cell leukemia phenotype. While this work was in progress, we were aware of another report studying a similar MMTV-CRE driven *MYB-NFIB* GEMM crossed with mutant p53 that did not generate salivary gland tumors (32).

These observations may be explained by negative selection of MYB expression in murine salivary gland tissues via unknown mechanism or due to shortened survival for *MYB-NFIB* transgene mice that exhibited a predominant phenotype of leukocytosis, thrombocytosis, and splenomegaly (Supplemental Fig. 1b, 1c). To define the hematopoietic phenotype, we performed flow cytometry on peripheral blood (PB) cells of control *MMTV-Cre* mice and compared to both *MYB-NFIB* and *MYB-NFIB/Ink4a*^(+/-)/*Art*^(+/-) mice. We analyzed the B220⁺ cell population and identified a statistically significant expanded B-cell population

with an increase in immature B220⁺IgM⁺ B-cells in *MMTV-Cre/MYB-NFIB* transgene mice compared to *MMTV-Cre* controls. In contrast, we found a decreased proportion of myeloid CD11b⁺Gr.1⁺ and Gr.1⁻ cells and a decreased CD3⁺CD4⁺CD8⁻ and CD3⁺CD8⁺CD4⁻ T lymphocyte populations in PB of *MMTV-CRE/MYB-NFIB* transgene positive mice compared to *MMTV-Cre* control mice. In addition, we observed inhibition of other hematopoietic lineages in *MMTV-Cre/MYB-NFIB* mice (Supplemental Fig. 1d). Histologic analysis also showed disrupted splenic architecture and infiltration of clusters of clonal lymphocytes in spleen, liver, murine femur bone marrow sections, and also within salivary glands (Supplemental Fig. 2a). For all parameters, we observed a more aggressive B-cell leukemia phenotype in *MMTV-Cre/MYB-NFIB/Ink4a^(+/-)/Arf^(+/-)* mice compared with *MYB-NFIB/MMTV-Cre* or *MMTV-Cre* alone, including higher leukocyte and lower platelet counts (Supplemental Fig. 2b).

MYB depletion inhibits ACC cell viability in vivo and in vitro

To study if MYB depletion affects growth proliferation of the new ACC UFH1 and UFH2 samples, we obtained three different LKO.1 lentiviral *MYB* shRNA clones and tested the efficiency of each constructs using a MYB-activated hematopoietic leukemic cell line, K562. Using immunoblot analysis we validated that transient expression of shRNA clones 59 and 60 depletes MYB expression (Fig. 4a). Therefore, we used lentiviral clones 59 and 60 for transient transfection into the UFH2 and UFBG cell lines. We observed a statistically significant decrease in cell viability/metabolism by MTT assay (Fig. 4b). We also tested the effect of MYB inhibition *in vivo* using MYB shRNA or control vectors on PDX xenograft growth from tumor cells obtained from UFH1PDX or ACCX22. MYB depletion significantly inhibited the tumor volume of xenograft tumors (Fig. 4c) suggesting that sustained MYB expression is required for efficient tumor growth of these cell lines.

Novel MYB inhibitors suppress MYB-activated ACC cell lines viability

Recent studies have proposed that Celastrol, a natural low-molecular-weight compound, can inhibit MYB function through disruption of its interaction with the KIX domain of p300/ CBP, a transcription co-activator of MYB protein (22). To test the efficacy of Celastrol on suppressing tumor growth/metabolism of the human UFH2 early passage cell line and the murine ACC-like mammary tumor cell line (UFBT), we performed MTT assay after incubation with increasing concentrations of either the control DMSO solvent or Celastrol. We observed that Celastrol inhibits viability at 1.25 μ M (Fig. 5a) that was similar to the effective Celastrol dose in MYB-activated leukemia cells (23). However, we detected a similar reduction in cell viability following exposure of Celastrol to tumor lines that were not associated with MYB activation (3t3 and AsPC1) suggesting a more general cytotoxicity of Celastrol (Fig. 5b). As an alternate strategy to test for MYB-specific cytotoxicity, we tested the effect of a peptidomimetic inhibitor (MYBMIM) that was recently developed using a novel structure-guided molecular design to target and interfere with the assembly of the MYB:CBP/P300 co-transcriptional protein complex (24). To test the effect of MYBMIM on our new human and murine MYB-activated tumor cell lines (UFH2 early passage tumor cell line and murine mammary UFBT tumor cell line), we incubated tumor cells with varying concentrations of either MYBMIM and performed MTT assay for growth viability/ metabolism. To control for non-specific peptidomimetic effects, we also treated cells with

the inactive MYBMIM analogue termed TG3, which lacks 3 key amino acid sidechains that are responsible for specific binding to CBP/P300 (24). We observed a dose-dependent tumor cell viability inhibition in both MYB-activated tumor cells by MYBMIM but not TG3 (Fig. 5c). To test for MYB-specific toxicity or growth inhibition of the MYB peptides, we incubated MYBMIM or TG3 with 3T3, Panc-1 and AsPC1 tumor cell lines that are not associated with MYB activation. We did not detect cell viability inhibition on these MYBnegative tumor cell lines using MYBMIM (Fig. 5d). This suggests that MYBMIM may have limited toxicity on normal tissues and therefore represents a promising probe for the development of selective MYB inhibitors for ACC patients and tumors with MYB activation.

Discussion:

It has been 40 years since the first report describing isolation and characterization of a defective avian tumor virus encoding the MYB oncogene (25). Over the following decades, the MYB gene has been identified as a key transcriptional regulator essential for the normal maturation for all hematopoietic elements (26-29). However, defining the role of MYB in tumorigenesis has been challenging due to the inability to develop functional oncogene assays using non-hematopoietic cells or tissues and, until recently, the lack of recurrent structural somatic mutations that provide further support for a 'driver oncogene' role in human cancer. The identification of recurrent chromosomal translocations resulting in either MYB or MYBL1 fusion transcripts in the majority of human adenoid cystic cancer (30) provides the strongest support for MYB activation as an etiologic event in carcinogenesis that also including the important subset of breast, colon, and hematopoietic tumors associated with MYB overexpression. Despite the direct evidence for structural MYB rearrangements, it has been difficult to generate validated human ACC tumor cell lines (31) and there are no activated MYB mouse models that recapitulate ACC salivary gland tumors. These obstacles have limited the ability to study MYB biology and develop new therapeutic agents. For example, we observed elevated steady-state MYB protein expression in extracts from excised human ACC xenografts but rapid loss of MYB protein expression within 5 hours of plating cell culture *in vitro* limits the ability to study MYB and to develop therapeutic strategies (Supplemental Fig. 2c). The mechanism for MYB down-regulation following plating tumor cells on plastic dishes *in vitro* is unexplained but may be linked with the difficulty to adapt xenograft ACC tumors to cultured tumor cell lines in vitro. We also generated and studied a new MYB-NFIB GEMM under MMTV-Cre regulation. As recently reported using a similar conditional MYB-fusion transgene strategy (32) we did not observe epithelial salivary gland tumors in either MYB-NFIB alone or in mice crossed with $Ink4^{+/-}/Arf^{+/-}$ null allele gene. While this might be partly explained by shortened survival due to the predominant phenotype of lethal B-cell leukemia in majority of mice, MMTV may be a suboptimal CRE promoter as we were unable to detect steady-state MYB-NFIB expression in murine salivary gland tissues while the human transgene was expressed in murine leukemia tumor cells. Similar to the prior report (32), we observed mammary tumors and were able to successfully generate a murine breast cell line associated with constitutive MYB transgene activation that allowed us to pursue drug testing for growth inhibition.

There are 7 candidate MYB inhibitors that have been recently proposed: celastrol and naphthoquinone (23,33), crizotinib (MET/ALK inhibitor) and linstinib (dual IGF1/insulin receptor inhibitor (34), MYBMIM, a novel MYB peptidomimetic (24), and the related semisynthetic epipodophyllotoxins chemotherapy agents, teniposide and etoposide (35). We selected for testing Celastrol, a natural product from *Tripterygium wilfordii*, and MYBMIM, a novel MYB peptidomimetic. In contrast to several of the other proposed pharmacological agents which had different known specificities and appeared to targeted MYB indirectly, Celastrol and MYBMIM offered two different pharmacologic strategies to interfere with the ability of MYB to achieve a functional protein complex in cancer cells (22, 23, 24). We have now observed specific effects of MYBMIM to preferentially suppress the growth and survival MYB-activated tumor cell lines validating this strategy for further pharmacologic development. Recently, there is a report of another human ACC tumor cell line (17) and the activated tumor cell lines reported here will offer new opportunities for drug candidate screening and development.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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We have isolated and studied two new ACC patient-derived xenograft (PDX) models and developed one matched ACC tumor cell line. We also generated and analyzed the tumor phenotype of a conditional *MYB-NFIB* transgene genetically engineered mouse model (GEMM). We did not detect salivary gland tumors with this GEMM or with *MYB-NFIB* mice crossed to Ink4a^{+/-}/Arf^{+/-} mouse model. However, we were able to isolate a murine mammary carcinoma associated with elevated *MYB-NFIB* transgene expression and ACC-like histology. We observed reduced survival of mice due to immature B-cell leukemia consistent with a role for MYB deregulation in blocking hematopoietic cell lineage differentiation. We have focused on characterizing these new MYB-activated human and murine cell lines to test candidate MYB inhibitors. These data will provide tools to study the biology of ACC and help develop new therapeutic strategies for both ACC and other common adult malignancies associated with MYB activation.

Highlights

We studied a new ACC PDX sample that express MYB-NFIB with a matched cell line

We studied a MYB-NFIB mouse model and generated an ACC-like mammary tumor cell line

We show MYB-specific inhibition in ACC tumor lines using a peptidomimic inhibitor

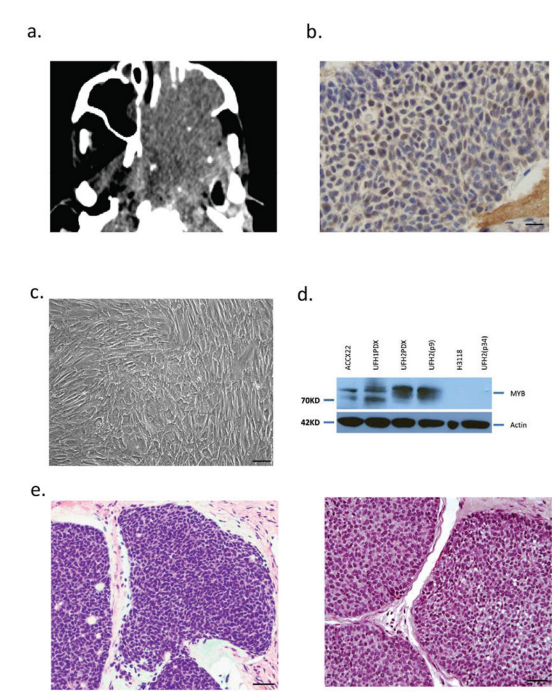


Figure 1.

Establishment of human ACC PDX and cell line (UFH2). (a) CT image of unresectable ACC requiring palliative debulking surgery. (b) Nuclear anti-MYB immunohistochemical staining in tumor biopsy. (c) UFH2 tumor cells *in vitro* at confluence. (d) MYB protein immunoblot (Anti-Myb, abcam, #ab45150) using ACCX22 (positive control), H3118 mucoepidermoid (negative control) and UFH1 PDX and UFH2 PDX and cell line at indicated passage number. (e) H&E staining primary ACC tumor biopsy section (left) and excised matched UFH2 PDX xenograft section (right).

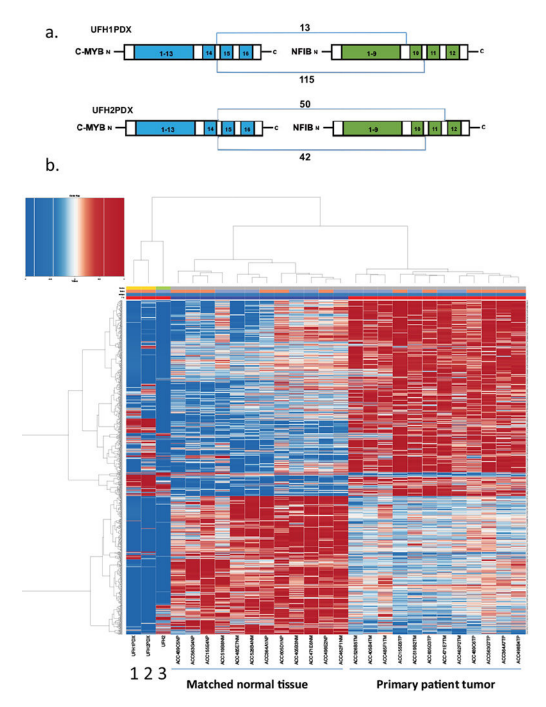
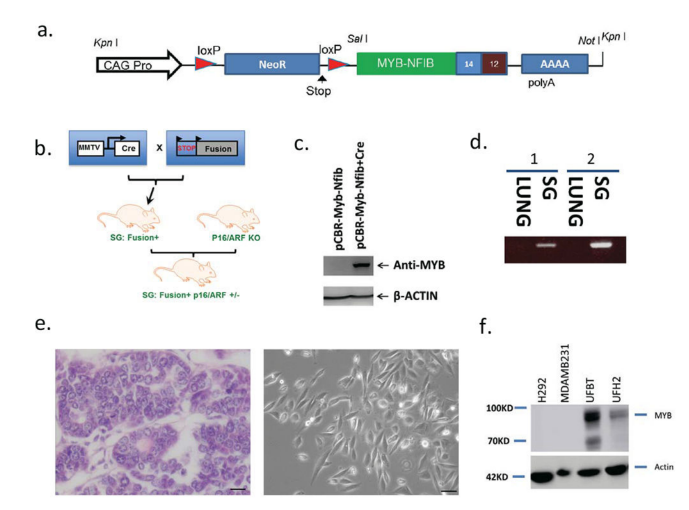


Figure 2.

MYB-NFIB fusion transcripts and gene expression pattern in ACC PDXs and ACC cell line. (a) Schematic of distinct *MYB* fusion transcripts detected in either UFH1PDX or UFH2PDX. (b) hierarchical clustering of mRNA expression by RNA-Seq using ACC UF PDX tumors and ACC cell lines compared with previously defined mRNA signature (21) from 12 ACC patient tumors and matched normal samples as indicated. Lane 1 (UFH1 PDX), Lane 2 (UFH2 PDX), Lane 3 (UFH2 cell line passage 34).

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g.

Modeling MYB-NFIB and Ink4a-Arf induced tumor model				
	lymphocytic leukemia/Total mice	%	ACC-like/Total mice	%
MYB-NFIB/MMTV-CRE	16/42	38	0/42	0
MYB-NFIB/MMTV-CRE/Ink4a-Arf+/	10/20	50	1/20	5

Figure 3.

Conditional *MYB-NFIB* transgene model. (a) Schematic lox-stop-lox design pCBR-based transgene driven by ubiquitously expressed CAG promoter. (b) Breeding strategy for *MYB-NFIB*/MMTV-Cre mice and mice crossed to Ink4a/ARF^{+/-}. (c) CRE-dependent expression of *MYB-NFIB* transgene verified by immunoblot analysis using 293 cells transfected construct alone or co transfected with adeno-Cre. (d) RT-PCR analysis of RNA extracted from *MYB-NFIB* transgene-positive normal salivary gland (SG) or lung. (e) H&E staining of resected mammary tumor from *MYB-NFIB/p16/ARF^{+/-}* animal (left) and photomicrograph image of subconfluent mammary tumor line (UFBT) (right). (f) MYB immunoblot of H292 mucoepidermoid tumor cell line and MDAMB231 triple-negative breast tumor line (negative controls) and mammary UFBT and UFH2 tumor cell lines. (g) Summary of predominant tumor phenotypes.

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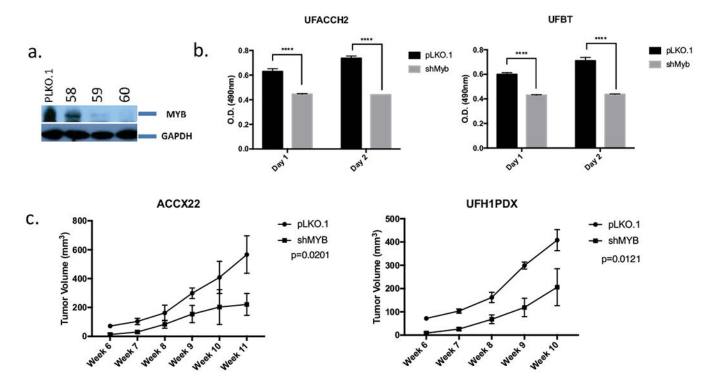


Figure 4.

MYB inhibitors suppress ACC viability *in vitro* and *in vivo*. (a) Immunoblot analysis testing efficiency of MYB depletion with PLKO.1 control and three different lentiviral MYB shRNAi clones. (b) Viability of UFBT and UFH2 cell lines after transduced with control or MYB shRNA lentiviruses (error bars represent standard deviation of UFBT (n=3 experiments) or UFH2 (n=5 experiments). (c) 2x10^6 ACCX22 (left) or UFH1PDX (right) cells were treated with shMYB or control and injected subcutaneously to dorsal flanks of NOD/SCID mice. ACCX22 PDX (left; n=3 mice control and n=3 mice shMYB), UFH1PDX (right; n=3 mice control and n=6 mice shMYB). Error bars represent median and standard deviations.

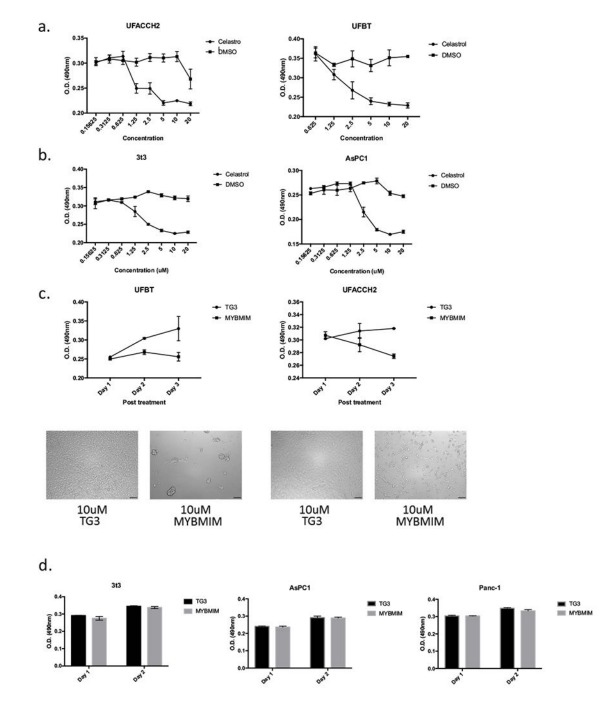


Figure 5.

MYB inhibitors suppress ACC cell lines viability *in vivo*. (a) Viability of early passage human UFACCH2 or murine ACC-like breast tumor UFBT cell lines after exposure to Celastrol at indicated concentrations or DMSO control for 48 hrs. Error bars represent standard deviations of four biological replicates. (b) Viability of control tumor cell lines that are not associated with MYB activation (3T3 and AsPC1) after 48 hrs exposure to Celastrol. (c) Viability (top panel) of UFBT (left) or UFH2 (right) cells, treated with 10 μ M of inactivated TG3 analogue peptide or 10 μ M MYBMIM. Error bars represent standard

deviations of three biological replicates. Photomicrographs (bottom panel) after incubation with indicated peptide mimic inhibitors (UFBT, left and UFH2, right). (d) Viability of non-MYB-activated control tumor cells (3T3, AsPC1 and Panc-1) after exposure to 10 μ M control TG3 or MYBMIM for 48 hrs. Error bars represent standard deviations of four biological replicates.