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Abstract Specific mutations in the RET proto-oncogene are associated with multiple endocrine neoplasia type 2A, a hereditary syndrome characterized by tumorigenesis in multiple glandular elements. In rare instances, MEN2A-associated germline RET mutations have also occurred with non-MEN2A associated cancers. One such germline mutant RET mutation occurred concomitantly in a young adult diagnosed with alveolar rhabdomyosarcoma, a pediatric and young adult soft-tissue cancer with a generally poor prognosis. Although tumor tissue samples were initially unable to provide a viable cell culture for study, tumor tissues were sequenced for molecular characteristics. Through a hierarchical clustering approach, the index case sample was matched to several genetically similar cell models, which were transformed to express the same mutant RET as the index case and used to explore potential therapeutic options for mutant RET-bearing alveolar rhabdomyosarcoma. We also determined whether the RET mutation associated with the index case caused synthetic lethality to select clinical agents. From our investigation, we did not identify synthetic lethality associated with the expression of that patient’s RET variant, and overall we did not find experimental evidence for the role of RET in rhabdomyosarcoma progression.

INTRODUCTION

Rhabdomyosarcoma (RMS) is the most prevalent pediatric soft tissue sarcoma, with frequent incidence in adolescent and young adults (Rudzinski et al. 2017). The alveolar subtype (aRMS) accounts for 20%–30% of all RMS tumors and thus represents ~1% of all pediatric cancers. aRMS has an annual incidence within the United States of ~350 cases/yr (Amer et al. 2019). aRMS is commonly initiated by a gene fusion event, most prevalently the PAX3:FOXO1-positive (P3F) variant, less commonly the PAX7:FOXO1-positive (P7F) variant, and rarely the atypical fusions (PAX3:NCOA1, PAX3:INO80) or even fusion-negative (Parham and Barr 2013; Shern et al. 2014). A substantial fraction of patients who develop metastatic aRMS (25%–30%) have metastatic disease at the time of diagnosis, with aRMS most frequently spreading to bone marrow, distal nodes, and bone (Ognjanovic et al. 2009; Amer et al. 2019).

Standard clinical care for aRMS is a combination of surgery, radiation, and intensive chemotherapy (Hawkins et al. 2013). Therapeutic schedules have been developed for RMS patients based on their Oberlin risk group, with the goal of reducing therapy-related toxicity in patients in low risk groups. The Oberlin risk score is determined by several factors but is
strongly influenced by the number of metastatic lesions and the location of the metastatic disease. RMS patients with fusion-positive tumors are more likely to have a higher risk score than fusion-negative patients; however, fusion status has little additional prognostic value when patients are stratified by Oberlin score. The 5-yr event-free survival rates based on Oberlin score are Oberlin 0 or 1, 52.4%; Oberlin 2, 22.8%; Oberlin 3, 6.3%; and Oberlin 4, 0% (Rudzinski et al. 2017).

The research presented here complements our case report of a teenage male patient (denoted CF-00034) with a germline \textit{RET C634F} mutation diagnosed with aRMS bearing the canonical PAX3:FOXO1 genomic rearrangement (Crawford et al. 2020). \textit{RET} mutations occur in 1.8% of tumors across disparate disease types (Kato et al. 2017), and the PAX3:FOXO1 fusion occurs in 55% of ARMS cases (Marshall and Grosveld 2012). The \textit{RET C634F} mutation is associated with Multiple Endocrine (MEN) Type 2 and has well established oncogenic potential (Schuffenecker et al. 1998; Zhou et al. 2007; Masbi et al. 2014). Prior reports demonstrate metastatic PAX3:FOXO1 aRMS in a 13-yr-old male (Agarwal et al. 1997; Jones et al. 2010) with familial MEN Type 2A induced by \textit{RET V804M} mutation, a patient genotypically and phenotypically reminiscent of the index patient. MEN syndromes can also be caused by loss-of-function mutations in MEN1 or CDKN1B, such as a case of rhabdomyosarcoma with a MEN1 1280delG truncation mutation (Alpers and Jones 2010). Nonetheless, rhabdomyosarcoma bearing MEN syndromes is a rare occurrence with few published clinical cases.

\textit{RET} is a transmembrane coreceptor tyrosine kinase involved in neurite guidance and outgrowth (Hezam et al. 2018). In neurons, \textit{RET} stimulates a multitude of biological functions including cell survival, differentiation, proliferation, and migration/chemotaxis. In cancer cells \textit{RET} is associated cell migration and proliferation, autophagy, and chemoresistance (Mulligan 2019). \textit{RET} mutations occur regularly in other cancer types such as medullary thyroid cancer (43%–71% of patients) and lung carcinomas (16.7%) (Kato et al. 2017), which has led to clinical trials of highly selective \textit{RET} inhibitors with promising results (Ackermann et al. 2019). \textit{RET} signaling requires the stepwise assembly of a signaling complex that first requires binding of a dimerized ligand of the glial cell line–derived neurotrophic factor (GDNF) family (GDNF, neurturin [NTRN], artemin [ARTN], persephin [PSPN]) by two GDNF family receptors (GFRα1–4) (Takahashi 2001), which forms a tetrameric complex that binds and induces dimerization of \textit{RET} (Supplemental Fig. 1A). The active signaling complex in a 2:2:2 ratio results in trans-autophosphorylation of \textit{RET} that can lead to activation of several signaling pathways including MAPK, PI3K/AKT, and Src. However, mutant \textit{RET} does not have the same signaling complex requirements. Mutations in the kinase domain or \textit{RET} rearrangements render the kinase constitutively active (Cranston et al. 2006), whereas mutations in the \textit{RET} extracellular domain, as in the case of the index patient (C634F), render the receptor ligand-independent (Kjær et al. 2006).

Given the association of mutant \textit{RET} to cancer oncogenesis and metastasis, and previous reports of mutant \textit{RET} in aRMS patients (Agarwal et al. 1997; Jones et al. 2010; Crawford et al. 2020), we investigated if the germline \textit{RET} mutation coincident in a subset of aRMS patients may have contributed to the clinical course of CF-00034, and if the \textit{RET C634F} mutation subsequently induces atypical susceptibilities in RMS tumor cells for patients with elevated \textit{RET} expression or germline \textit{RET} mutations.

\section*{RESULTS}

\subsection*{RET Signaling Components in aRMS}

To determine the importance of \textit{RET} activity in aRMS, we evaluated the survival data and paired gene expression data from sporadic cases of RMS (\textit{n} = 23 cases) archived by the
Children’s Oncology Group (Fig. 1). RET expression demonstrated a trend associated with decreased long term survival over 10 years in an expression-dependent manner when adjusted for known clinical covariates (age, stage, sex, fusion status), although cohort sizes were small (four to 14 patients/group) and results did not reach statistical significance ($P = 0.069$). We next evaluated whole-transcriptome sequencing data from human aRMS biopsy samples to determine the expression of the RET signaling complex members (Supplemental Fig. 1). Although RET and GFRα were expressed in several samples, no single sample expressed both coreceptors RET and GFRα simultaneously. The evaluation of this diverse collection of RMS tumors indicates that RET is rarely expressed (∼5%), whereas coreceptors GFRA1–4 are more commonly expressed (∼34%), and all samples transcribe at least one RET ligand, with ATRN and NRTN being near-universally expressed.

Endotype Analysis and Compound Screen of Endotypically Similar Cell Models

The lack of available RET-mutant aRMS cell models coupled with lack of patient-derived primary cultures at the time of our studies necessitated identifying cellular models genetically and transcriptionally similar to the index case for in vitro experiments. Similarity of cell models was quantified using an aRMS-focused dendrogram endotype computational model (Ricker et al. 2020). Multiple aRMS exome and transcriptomes sequencing data sets were obtained from internal cc-TDI CuRe-FAST samples, the Database of Genotypes and Phenotypes (dbGaP), the Gene Expression Omnibus (GEO) database, Foundation Medicine sequencing panels, the Jackson Laboratory PDX sequencing data, and Champions Oncology PDX sequencing data as well as statistical comparison of RET and RET signaling partner expression between ARMS and ERMS cohorts (Supplemental Tables 1, 2). Using DNA mutation data and RNA expression data from the accumulated aRMS sample cohort, endotype clustering analysis was performed on the aRMS sample sequencing data cohort. Hierarchical clustering identified multiple out-groups (self-clustering consistent with expectations) such as Foundation Medicine panel data (which identifies mutations in a subset of tumor-associated genes), non-PAX3 fusion aRMS samples, and fusion-negative samples.

An endotype emerged which focused on RET expression as a key differentiator between hierarchical clusters (Fig. 2A). The index case was determined to be endotypically most similar to CF-00001X (a pediatric PAX3–FOXO1 aRMS with an established PDX model and PDX-derived cell line), U31431 (a PAX3–FOXO1 GEM model–derived cell culture), and four...
additional GEM cell models (U48484, U23674, U28285, U34279), all bearing \( \text{PAX3} - \text{FOXO1} \) fusions (Fig. 2B). Lacking a CF-00034-originating cell culture, cell models identified through endotype analysis were used as surrogates for the index case in subsequent experiments.

The human cell model (CF-00001X) and murine cell model (U31314) endotypically most similar to index patient CF-00034 were screened in a 60-compound drug screen (Supplemental Table 3) to identify therapeutic strategies potentially applicable to the index case (e.g., synthetic lethality). U31314 supplemented with RET ligand ARTN (100 nM) was assayed a second time using the 60-compound screen; no significant difference in drug sensitivity was observed.

Although CF-00001X was sensitive to a greater number of agents than U31314, few drugs showed growth inhibition activity in both models. Of note, we observe CF-00001X was sensitive to IGFR1 inhibitors (OSI-906 and BMS-754807), HDAC inhibitors (panobinostat, CUDC101, CUDC907 and entinostat), and inhibition of several targets in the PI3K pathway (AKT, PI3K, and mTOR).

Figure 2. Index case endotype and patient avatar cell models. (A) Endotype dendrogram of \( \text{PAX3} - \text{FOXO1} \) and \( \text{PAX7} - \text{FOXO1} \) alveolar rhabdomyosarcoma patient samples, cell lines, patient-derived xenograft (PDX) models, and genetically engineered mouse (GEM) models. Data sets were obtained from internal cc-TDI CureFAST samples, database of genotypes and phenotypes (dbGaP) and gene expression omnibus (GEO) sequencing data, Foundation Medicine sequencing panel data, Jackson Laboratory PDX sequencing data, and Champions Oncology PDX sequencing data. Based on DNA and RNA expression of genes in both aRMS variants, multiple aRMS endotypes exist. Validation outgroups include Foundation Medicine samples (out-grouped because of focused sequencing of a subset of target genes) and \( \text{PAX7} - \text{FOXO1} \) samples (out-grouped because of expected molecular differences from \( \text{PAX3} - \text{FOXO1} \) samples). (B) Cell models identified as endotypically most similar to the index patient CF-00034 through endotype analysis and hierarchical clustering of aRMS samples and cell models.
Phenotypic Analyses

The presence of a known, characterized oncogenic germline RET mutation (Schuffenecker et al. 1998; Masbi et al. 2014) led to the hypothesis that clinical RET inhibitors, as monotherapy or in combination, would be promising treatment options for RET-mutant aRMS. To test this hypothesis, we created synthetic mutant RET cell models mimicking the genotype of the index case patient. Human and murine aRMS cells selected as representative models from the dendrogram (Fig. 2B) were transiently transfected with commercially available RET C634F expression plasmid and assayed with a panel of clinically available agents relevant to aRMS biology and the germline RET mutation (Fig. 3). Transfection of cell models was confirmed by western blot analysis (Fig. 4A), confirming expression of RET C634F and expression of MYC–DDK (FLAG) tagged protein.

Figure 3. Monotherapy treatment of null plasmid and RET C634F plasmid transformed endotypically similar cell models. Four cell models identified as endotypically similar to the index patient CF-00034 (CF-00001X, RH30, U23674, U31431) were treated with six different monotherapies (cabozantinib, entinostat, vincristine, cyclophosphamide prodrug, mafosfamide, temsirolimus) to determine agent response and unique sensitives induced by RET C634F protein expression.
The clinical RET inhibitor cabozantinib was selected as the experimental RET inhibitor because of off-label clinical use in the previous case report (Crawford et al. 2020). Cabozantinib has been investigated clinically for RET-rearranged non-small-cell lung cancer (Drilon et al. 2016) and has shown significant clinical efficacy in RET M918T mutant medullary thyroid cancer (Krajewska et al. 2016). For combination therapies, we selected two clinical chemotherapy agents (vincristine and cyclophosphamide [tested as the prodrug mafosfamide]) not used to treat the index case patient described in the preceding manuscript (Crawford et al. 2020) and two clinical targeted inhibitor compounds (HDAC inhibitor entinostat [recently in a Phase I pediatric clinical trial, ADVL1513/NCT02780804] and mTOR inhibitor temsirolimus). Each agent was first tested as a monotherapy to determine overall single agent efficacy across RET C634F transfected and null plasmid cell models (Fig. 3). Drug response to monotherapy and cabozantinib-focused combination therapy of RET

![Figure 4. Cabozantinib-focused drug combinations in synthetic RET-mutant cell lines. (A) Western blots of null plasmid and RET C634F transformed endotypically similar cell models, confirming successful transformation and expression of RET C634F. The western blot image in this panel was generated from the grouping of two individual western blot images. No other modifications were made to the western blot image. (B) Representative 3D response manifold of RET C634F transformed RH30 cell model treated with combinations of cabozantinib and entinostat. (C) Representative 3D response manifold of null transformed RH30 cell model treated with combinations of cabozantinib and entinostat. (D) Representative Chou–Talalay Combination Index values from RET C634F cabozantinib plus entinostat combination. (E) Representative Chou–Talalay Combination Index values from null plasmid cabozantinib plus entinostat combination.](image-url)
C634F transfected versus null plasmid transfected cell models were tested for statistically significant differences. For monotherapy, absolute IC$_{50}$ values were used for statistical comparison when available or dose-level response vectors were used when single agents did not achieve 50% growth inhibition. In RET C634F transfected versus null cell models, no monotherapy treatment showed statistically significant change in sensitivity across all cell models (Fig. 3).

Combination therapies were tested in checkboard dosage design, with cabozantinib tested in combination with the remaining five monotherapy agents. Combination Index values and 3D dose response manifolds were generated for each cell model/drug combination (representative manifold presented in Fig. 4B,C). Several combinations exhibited synergy at specific combination doses and/or at dose regions in RET C634F synthetic cell models (representative combination index plots are presented in Fig. 4D,E). All manifold and combination index results are provided in Supplemental Table 4.

We also tested for statistically significant differences in single agent response based on IC$_{50}$ values (Table 1) and overall response curve (Table 2), as well as differences in drug combination response manifolds in RET C634F transfected versus null cell models (Table 3). Although no monotherapy or combination therapy showed statistically significant change across all cell models, monotherapy entinostat showed statistically significant difference in three of four models. However, although the statistically significant human model (RH30) showed increased sensitivity to entinostat, both mouse models showed statistically decreased entinostat sensitivity. Additionally, both RH30 and U31431 showed statistically

### Table 1. Statistical analysis of monotherapy and combination drug screens with and without synthetic RET mutation

|                | Human   | Mouse       |
|----------------|---------|-------------|
|                | RH30    | CF1X        |
| Cabozantinib   | -       | -           |
| Entinostat     | 0.7298  | 1 × 10^{-04}|
| Vincristine    | -       | -           |
| Mafosfamide    | 0.9604  | 0.7746      |
|                | U23674  | U31431      |
| Cabozantinib   | -       | 1 × 10^{-04}|
| Entinostat     | 1 × 10^{-04} | 1 × 10^{-04}|
| Vincristine    | -       | 0.0719      |
| Mafosfamide    | 1 × 10^{-04} | -          |

*Statistical comparison of RET WT and RET C634F monotherapy IC$_{50}$ values. Comparison is provided as P-values, with green text indicating statistically significant differences.

### Table 2. Statistical analysis of monotherapy and combination drug screens with and without synthetic RET mutation

|                | Monotherapy response vector comparison (P-value) |
|----------------|-----------------------------------------------|
|                | Human       | Mouse       |
|                | RH30        | CF1X        |
| Cabozantinib   | 0.37        | 0.7706      |
| Entinostat     | 0.1363      | 0.0215      |
| Vincristine    | 0.0512      | 0.5773      |
| Mafosfamide    | 0.0604      | 0.3234      |
|                | U23674      | U31431      |
| Cabozantinib   | 0.5068      | 0.1772      |
| Entinostat     | 0.0618      | 0.1375      |
| Vincristine    | 0.0136      | 0.5747      |
| Mafosfamide    | 0.3652      | 0.1196      |

*Statistical comparison of RET WT and RET C634F monotherapy response vectors. Comparison is provided as P-values, with green text indicating statistically significant differences.*
significantly different response to the cabozantinib + mafosfamide combination, as well as increased overall synergy by combination index values (Supplemental Table 4). Nonetheless, the overall lack of statistically significant differences between RET\textsuperscript{C634F} trans-fected versus null plasmid cell models across all models suggests RET\textsuperscript{C634F} mutation in rhabdomyosarcoma may not induce sensitivity to RET inhibition, nor confer additional sensitivity to chemotherapy agents or targeted agents.

**DISCUSSION**

This research began with an unusual clinical observation, an aRMS patient with aggressive metastatic disease has an underlying germline RET\textsuperscript{C634F}-activating mutation. RET activation is known to be the ligand-independent, self-activated driver mutation of MEN2A syndrome cancers and thought to be involved in lung adenocarcinoma (Wang et al. 2019), breast cancer (Paratala et al. 2018), non-small-cell lung cancer, and colorectal cancer (Gozgit et al. 2018). Given the unique genotype of the patient, the poor prognosis associated with metastatic aRMS, and previous success of RET inhibitors in RET mutant tumors (Ackermann et al. 2019), we investigated the lethality of a clinical RET inhibitor as a monotherapy or in combination with other clinically available agents when treating RET-mutant expressing cell models. Sensitivity to monotherapies and combination therapies were consistent in both null plasmid transformed and RET\textsuperscript{C634F} transformed cell models, suggesting the RET mutation borne by the patient did not induce new susceptibility to RET inhibition. Although RET mutation does not necessarily guarantee complete response to a RET inhibitor (Valenciaga et al. 2018), lack of any significant susceptibility to RET inhibition is surprising and speaks to the clinical challenge aRMS presents even when a genomically relevant target is readily druggable.

Promisingly, the recently U.S. Federal Drug Administration (FDA)-approved RET inhibitor selpercatinib has been approved for the use in RET fusion-positive non-small-cell lung cancer (Drilon et al. 2020) and medullary thyroid cancer with RET mutations or RET fusions (Wirth et al. 2020), including both kinase-domain mutations and extracellular cysteine mutations. Notably, extracellular cysteine mutations show overall reduced response to selpercatinib compared to kinase domain mutations. Given the use of cabozantinib as a maintenance therapy in the index case patient, the potential impact of selpercatinib in the index case or similar patients is unclear.

We also tested a panel of multiple agents relevant to rhabdomyosarcoma biology against two of the endotype-based cell models, identifying susceptibility to IGFR1 and PI3K inhibitors in the endotypically closest human cell model, consistent with previous

### Table 3

| Combination therapy response vector comparison (P-value) | Human RH30 | CF1X | Mouse U23674 | U31431 |
|---------------------------------------------------------|------------|------|--------------|--------|
| cab-ent                                                 | 0.8663     | 0.529| 0.0252       | 0.7756 |
| cab-vnc                                                 | 0.1905     | 0.1199| 0.4231       | 0.1532 |
| cab-maf                                                 | 0.0062     | 0.1105| 0.3825       | 0.0319 |

Statistical comparison of RET WT and RET\textsuperscript{C634F} combination therapy response vectors as P-values. Comparison is provided as P-values, with green text indicating statistically significant differences.

Statistical analysis of monotherapy and combination drug screens with and without synthetic RET mutation

Statistical comparison of RET WT and RET\textsuperscript{C634F} combination therapy response vectors as P-values. Comparison is provided as P-values, with green text indicating statistically significant differences.
aRMS studies (Crosswell et al. 2006; Abraham et al. 2011; Berlow et al. 2019), again supporting RET status playing a minimal role in altering response to therapeutic agents. Although rarely occurring concomitantly, metastatic aRMS with RET C634F germline mutation presents a difficult phenotype and a dire clinical prognosis and will likely remain as intractable as classical aRMS cases.

A key challenge in the investigation of RET C634F mutant aRMS is the lack of readily available cell models and the initial biopsy material from the patient not generating a viable cell culture. Overcoming the lack of patient-derived cell model by using a cohort of sequenced aRMS samples to identify available cell models most molecularly similar to the index case is a strategy for research-level functional analysis, which may be applicable to other patients with atypical or challenging genotypes. Development of rare disease cohorts will beneficial both in understanding the disease as a whole but also in exploring the unique genotypes possessed by individual patients.

**METHODS**

**Cell Lines**

The RH30 cell line (Douglass et al. 1987) (RRID:CVCL_0041) was shared by the Children’s Oncology Group Repository (https://www.cccells.org/). RH30 was cultured in RPMI-1640 (11875-093; Thermo Fisher Scientific) supplemented with 10% fetal bovine serum (FBS) (26140079; Thermo Fisher Scientific) plus 1% penicillin-streptomycin (P/S) (15140-122; Thermo Fisher Scientific) maintained at 5% CO2 at 37°C.

**Murine Cell Model Establishment**

The mouse primary tumor cell cultures U31431 and U23674 were established from a site-of-origin conditionally induced aRMS tumor in a genetically engineered Myf6Cre,Pax3:Foxo1, p53 mouse as previously described (Abraham et al. 2014). In brief, extracted tumor was minced and digested with 10 mg/mL collagenase type IV (17104019; Thermo Fisher Scientific) overnight at 4°C. Dissociated cells were then incubated in Dulbecco’s modified Eagle’s medium (DMEM) (11995-073; Thermo Fisher Scientific) supplemented with 10% FBS and 1% P/S in 5% CO2 at 37°C. Cultures were maintained at low passage to minimize biological variation from the original tumor. All experiments were performed at or below passage 10.

**Patient-Derived Xenograft (PDX) Model Development**

All aspects of cancer tissue sharing for model development were reviewed and approved by the Children’s Cancer Therapy Development Institute Institutional Review Board. The CF-00001 PDX model and subsequent cell culture (denoted CF-00001X) were described previously (Bharathy et al. 2018). Fusion status of CF-00001 PDX-derived cells (denoted CF-00001X) was confirmed by RNA sequencing for PAX3–FOXO1 fusion reads and western blotting of CF-00001X culture lysate for PAX3–FOXO1 (Bharathy et al. 2018). Cultures were maintained at low passage to minimize biological variation from the original tumor. All experiments were performed at or below passage 10. CF-00001X was cultured in RPMI-1640 supplemented with 10% fetal bovine serum (FBS) plus 1% P/S and maintained at 5% CO2 at 37°C. The CF-0001 PDX model is maintained at the Jackson Laboratory (http://tumor.informatics.jax.org/mtbwi/pdxDetails.do?modelID=J000099761).
Data Integration and Dendrogram Generation

To define endotypes (disease subgroups defined by combined genomic, clinical, and pathobiological feature sets unique to each patient) within aRMS, we applied hierarchical agglomerative clustering analysis to a cohort of aRMS samples including human patient samples with PAX3–FOXO1 fusions (n = 23), patient samples with PAX7–FOXO1 fusions (n = 6 samples), patient samples with PAX3–NCOA1 fusions (n = 2 samples), patient samples with PAX3–INO8 fusions (n = 1 sample), fusion-negative patient samples (n = 8), patient-derived aRMS cell lines (n = 12), Foundation Medicine panel sequencing result from human aRMS (n = 15), genetically engineered mouse models (n = 17), and PDX models of human aRMS from Champions Oncology (n = 2: CTG-0329 and CTG-1419) and the Jackson Laboratory (n = 4: CF1_00327, CF2_06843, CF4_01466, PCB380_03416).

Data from disparate human data sets were aligned to a common set of genes across DNA sequencing data and RNA sequencing data. Mouse-origin samples were converted from mouse gene identifiers to homologous human gene identifiers via BioMart (Smedley et al. 2009) and merged with the human DNA or RNA sequencing data sets, respectively. Gene expression values were normalized to read depth across samples using edgeR (Robinson et al. 2009) and then across individual genes by assigning gene-level z-scores. Zero expression values were assigned to all DNA sequencing data sets lacking RNA sequencing data (i.e., Foundation panel sequencing data sets) to allow for clustering analysis. For mutation data, samples were assigned the value 1 for each gene bearing a mutation, and 0 otherwise. The resulting quantified mutation and expression data sets were concatenated as feature vectors for downstream clustering analysis. The Euclidean distance metric was used to define distance between feature vectors, with average-linkage clustering used to merge nonsingleton feature vectors into groups. Clustering analysis was performed using RStudio Version 3.6.1 using the hdist package.

Sequencing Data Analysis

All sequencing data from patient samples, patient-derived cell lines, genetically engineered mouse models, and the Jackson Laboratory PDX models were analyzed by the previously published analysis pipeline (Kats et al. 2019). In brief, gene mutations from whole-exome sequencing experiments were identified by aligning fastq files to the GRCh38 human reference genome or GRCm38 mouse reference genome via BWA aligner, processing with GATK best practices (DePristo et al. 2011), and called with MuTect2 (Benjamin et al. 2019). Gene expression from whole transcriptome sequencing experiments was analyzed by aligning fastq files to the GRCh38 human reference genome or GRCm38 mouse reference genome using STAR aligner (Dobin et al. 2013) and quantified by RSEM (Li and Dewey 2011).

Drug Screen of Endotypically Similar Cell Models

CF-00001X and U31431, were evaluated in a proliferation assay against a 60-compound library which was previously described and reported (Kats et al. 2019). In brief, cells were cultured and harvested, then plated via BioTek MultiFlo liquid dispenser into thawed preprinted compound library 384-well opaque white wall plates (164610; Thermo Fisher Scientific). Cells in plates were incubated for 72 h then assayed for cell viability via CellTiter Glo 2.0 (G9241; Promega) luminescent assay. Drug response and IC50 values were determined using Microsoft Excel.

Transfection of PAX3–FOXO1 Alveolar Rhabdomyosarcoma Cell Models

The RET mutant plasmid RET C634F built into pCMV6 Entry. carboxy-terminal Myc/DDK FLAG Tagged cloning vector was purchased from OriGene (RC403308; OriGene) and
expanded by Aldevron (http://www.aldevron.com/). The plasmid was transfected into four cell models: RH30, CF-00001X, U23674, and U31431.

Cells were plated in tissue-culture treated six well plates (140675; Thermo Fisher Scientific) at initial cell populations at 150,000 cells/well. After a 24-h acclimation period, the plated cells were transfected with DNA plasmid (lipofectamine 6 µL + 2 µg DNA) for 2 h in serum-free media (RPMI-1640 for human cells, DMEM for murine cells), and then washed and replaced with growth media. Transient transfection of RET C634F plasmid via the above methodology was confirmed for all cell models via western blot for Myc/DDK FLAG antibody (F3165; Millipore Sigma).

Immunoblotting
Following transfection, all four cell models were cultured for 24 h then harvested for protein via RIPA Lysis and Extraction Buffer (Thermo Fisher Scientific 89900) + Halt Protease and Phosphatase Inhibitor Cocktail (Thermo Fisher Scientific 78441). Null plasmid cells were cultured for 48 h, and protein was isolated in the same manner. RET expression in transfected and null plasmid cell models was analyzed using anti-RET antibody (1:1000 dilution; Cell Signaling Technologies 3220). Presence of RET C634F protein was confirmed using anti-FLAG M2 antibody (1:5000 dilution; Sigma-Aldrich F1804).

Monotherapy and Combination Therapy Drug Studies of Transformed Cell Models
Cell lines transfected with either null pCMV6 Entry plasmid or RET C634F pCMV6 Entry plasmid were collected and plated in growth media at 1500 cells/well into 384-well drug plates preprinted with six agents validated as a monotherapy and five cabozantinib-focused drug combinations based on the six monotherapy agents. The following drugs were tested as monotherapy: cabozantinib (Selleck Chemicals S4001), entinostat (Selleck Chemicals S1053), vincristine (Selleck Chemicals S1241), mafosfamide (Santa Cruz Biotechnology sc-211761), and temsirolimus (Selleck Chemicals S1044). Individual agents were tested in triplicate at the following doses: [10 nM, 100 nM, 1 µM, 10 µM]. The following two-drug combinations were selected for validation: cabozantinib + entinostat, cabozantinib + vincristine, cabozantinib + mafosfamide, and cabozantinib + temsirolimus. Each combination of agents was tested at the Cartesian product of the following dosages: cabozantinib [10 nM, 100 nM, 1 µM, 10 µM] × combination agent [10 nM, 100 nM, 1 µM, 10 µM] for a total of 16 data points per combination. Three technical replicates of each monotherapy/combination screen were performed for each null plasmid– and RET C634F plasmid–transfected cell model. Cell models in growth media were plated in the preprinted plates and incubated for 72 h in 5% CO2 at 37°C. At the end of the 72-h period, cell viability was assessed by CellTiter Glo 2.0 luminescent assay per manufacturer’s protocol and quantified via BioTek Synergy HT plate reader.

Drug sensitivity as IC50 value of each monotherapy was determined by Hill curve fitting using GraphPad Prism. Chou–Talalay Combination Index values (Chou 2010) for drug combinations were generated using the CompuSyn software tool. Effect values for CompuSyn monotherapy and combination therapy were determined by mean cell viability based on n = 3 independent technical replicates for each transfected cell model. Each null plasmid– and RET C634F plasmid–transfected cell model was compared for statistically significant differences of drug response to determine if RET C634F expression resulted in changes in drug sensitivity.

Sequencing Data
Sequencing data for endotype analysis and dendrogram generation were gathered from previously published data sets with the following accession IDs: GSE138269, GSM984615,
GSM758578, GSE142775 (Gene Expression Omnibus database), EGAS00001004359, EGAS00001003981 (European Genome-phenome Archive), PRJNA613152 (Short Read Archive), Champions TumorGraft database, Mouse Tumor Biology Database (http://tumor.informatics.jax.org/mtbwi/pdxSearch.do), the OncoGenomics Database (https://pob.abcc.ncifcrf.gov/cgi-bin/JK), and phs001121.v1.p1 (Database of Phenotypes and Genotypes).

**Statistics**

Student’s t-tests were used to compare IC₅₀ values from null and RET C634F transfected aRMS cell lines treated with different monotherapy agents. Hotelling’s T² test was used to compare one-dimensional drug response vectors from null and RET C634F transfected aRMS cell lines treated with different monotherapy agents, when 50% inhibitory concentrations were not reached. Hotelling’s T² test was also used to compare two-dimensional drug response manifolds from null and RET C634F transfected aRMS cell lines treated with different combination therapy agents.

**ADDITIONAL INFORMATION**

**Data Deposition and Access**

All sequencing data used to generate results in this manuscript have been previously deposited to online sequencing databases. The following data sets under the given accession IDs were used: GSE138269, GSM984615, GSM758578, GSE142775 (Gene Expression Omnibus database), EGAS00001004359, EGAS00001003981 (European Genome-phenome Archive), PRJNA613152 (Short Read Archive), Champions TumorGraft database, Mouse Tumor Biology Database (http://tumor.informatics.jax.org/mtbwi/pdxSearch.do), the OncoGenomics Database (https://pob.abcc.ncifcrf.gov/cgi-bin/JK), and phs001121.v1.p1 (Database of Phenotypes and Genotypes).

**Ethics Statement**

All human tissue samples were acquired through the Cancer Registry for Familial and Sporadic Tumors (CuRe-FAST) tumor banking program. All patients enrolled in CuRe-FAST provided informed consent. All aspects of the study were reviewed and approved by the Children’s Cancer Therapy Development Institute (cc-TDI) Institutional Review Board (IRB). Patient data and clinical and pathologic information are maintained in a de-identified database.

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**Author Contributions**

N.E.B. and K.A.C. designed the experiments, interpreted results, and wrote the manuscript; C.J.B., C.N., and I.S. provided materials and edited the manuscript; E.R.R. edited the manuscript; and C.K. wrote the manuscript.

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