Genetic modifiers of the BRD4-NUT dependency of NUT midline carcinoma uncovers a synergism between BETis and CDK4/6is

Sida Liao,1,2 Ophélia Maertens,1,3,4 Karen Cichowski,1,3,4 and Stephen J. Elledge1,2,4

1Division of Genetics, Department of Medicine, Brigham and Women’s Hospital, Boston, Massachusetts 02115, USA; 2Department of Genetics, Program in Virology, Howard Hughes Medical Institute, Harvard Medical School, Boston, Massachusetts 02115, USA; 3Harvard Medical School, Boston, Massachusetts 02115, USA; 4Ludwig Center at Harvard, Harvard Medical School, Boston, Massachusetts 02215, USA

Bromodomain and extraterminal (BET) domain inhibitors (BETis) show efficacy on NUT midline carcinoma (NMC). However, not all NMC patients respond, and responders eventually develop resistance and relapse. Using CRISPR and ORF expression screens, we systematically examined the ability of cancer drivers to mediate resistance of NMC to BETis and uncovered six general classes/pathways mediating resistance. Among these, we showed that RRAS2 attenuated the effect of JQ1 in part by sustaining ERK pathway function during BRD4 inhibition. Furthermore, overexpression of Kruppel-like factor 4 (KLF4), mediated BETi resistance in NMC cells through restoration of the E2F and MYC gene expression program. Finally, we found that expression of cyclin D1 or an oncogenic cyclin D3 mutant or RB1 loss protected NMC cells from BETi-induced cell cycle arrest. Consistent with these findings, cyclin-dependent kinase 4/6 (CDK4/6) inhibitors showed synergistic effects with BETis on NMC in vitro as well as in vivo, thereby establishing a potential two-drug therapy for NMC.

Keywords: BETi; BRD4; CDK4; genetic screens; JQ1; oncogenes; tumor suppressor

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NUT midline carcinoma (NMC) is an aggressive subtype of squamous cell cancer. Genetically, it is defined by the fusion of most of the coding sequence of the testis-specific gene NUT on chromosome 15 to several other genes—most frequently the BRD4 gene on chromosome 19 (BRD4-NUT) [French et al. 2003, 2008]. BRD4 is an epigenetic reader that binds to acetyl-histone. Fusion to the NUT protein results in recruitment of p300/CBP, a histone acetyltransferase, leading to regional histone hyperacetylation. Such acetylation further recruits BRD4-NUT in a feed-forward manner, eventually creating massive regions of acetylated chromatin that covers individual topologically associated domains across the genome [Alekseyenko et al. 2015]. The transcription of oncogenic proteins such as c-Myc [encoded by MYC] and its regulators within these regions is thus stimulated, blocking cellular differentiation and promoting uncontrolled growth of carcinoma cells [Grayson et al. 2014; Alekseyenko et al. 2015].

Small molecule bromodomain and extraterminal (BET) domain inhibitors (BETis) that target and inhibit BRD4’s association with chromatin have shown anti-cancer efficacy on a variety of cancers, including NMC, in preclinical models and are currently being evaluated in clinical trials [Stathis and Bertoni 2018]. A clinical proof of concept for BETis in NMC was demonstrated recently [Stathis et al. 2016]. Two of four NMC patients receiving BRD4 inhibition therapy showed increased overall survival [19 and 18 mo, respectively], which was notably longer than the median survival of 6.7 mo reported in the largest retrospective series of patients with NMC [Bauer et al. 2012]. However, as is often the case with other single-agent therapies, cancer cells can develop resistance to BETis through a variety of mechanisms [De Raedt et al. 2014; Fong et al. 2015; Kumar et al. 2015; Hogg et al. 2016; Kurimchak et al. 2016; Marcotte et al. 2016; Shi et al. 2016; Shu et al. 2016; Jang et al. 2017; Janouskova et al. 2017]. Despite the initial response, all of the NMC patients developed resistance to the BETis and eventually
relapsed. Therefore, there is an urgent need to identify the underlying mechanisms of resistance to BRD4 inhibition in NMC in order to develop combination therapies or alternative regimens after BRD4 inhibitor progression.

Cancer is driven by a number of distinct genetic alterations, including gain or loss of chromosomes and chromosomal segments, translocations, frameshifts, and point mutations that result in inactivation of tumor suppressor genes (TSGs) or activation of oncogenes. We previously developed an algorithm called TUSON Explorer to identify known and putative cancer drivers using human tumor sequencing data (Davoli et al. 2013). This method quantifies the likelihood that a gene is a cancer driver based on the distortion of its mutational signature from the pattern expected for a “neutral” gene. For example, TSGs will have a higher ratio of loss-of-function to benign mutations than neutral genes, while oncogenes will have a higher ratio of missense to benign mutations than neutral genes, and these missense mutations tend to cluster together (Fig. 1A). We and others have shown that cancer drivers can genetically interact and substitute for one another to drive proliferation and survival in EGFR mutant non-small cell lung cancers and other cancer models (Harbinski et al. 2012, Wilson et al. 2012, Sharifnia et al. 2014, Liao et al. 2017), suggesting that it might be a general notion that cancer drivers can form a function network and genetically modify the dependency of each other. Thus, the TSGs and oncogenes identified by TUSON Explorer could serve as an ideal list to survey for the genetic modifiers that can partially replace the cancer driver of interest—BRD4-NUT, in the case of NMC.

To systematically identify the TSGs and oncogenes that can partially replace BRD4-NUT in BRD4-NUT-dependent NMC tumor cells, we performed CRISPR and onco-gene/proto-oncogene expression screens in parallel using an in vitro NMC model. Here, we show that this genetic approach successfully recovered previously validated cancer drivers that mediate BETi resistance, such as MYC, in addition to novel cancer drivers not previously linked to BETi resistance in NMC. We further characterized the mechanisms underlying BETi resistance mediated by several novel oncogenes. Among these, we showed that mutation of RRAS2, a member of the R-Ras subfamily of Ras-like small GTPases, attenuated the effect of JQ1 in part by sustaining ERK pathway function during BRD4 inhibition. We also showed that overexpression of Kruppel-like factor 4 (KLF4), a transcriptional factor, mediated JQ1 resistance in NMC cells through restoring the E2F and MYC gene expression program upon JQ1 treatment. Finally, we showed that expression of cyclin D1 and mutant cyclin D3 or loss of RB1 protected the NMC cells from JQ1-induced cell cycle arrest. In accordance with this observation, cyclin-dependent kinase 4/6 (CDK4/6) inhibitors showed synergistic effects with JQ1 on NMC in vitro as well as in vivo, revealing the central role of cell cycle regulation in mediating JQ1 response. These

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**Figure 1.** Genetic screens identify modifiers of the cellular response to reduced BRD4-NUT signaling. (A) Schematic of the pipeline used by TUSON Explorer to predict TSGs and oncogenes (OGs). (B) Outline of the genetic screening strategy. (C,D) Log2 fold change comparing end time point (day 17) with start time point (day 0) of JQ1 treatment plotted against vehicle (DMSO) treatment for genes in the CRISPR and ORF screens, respectively. The top 10 genes of the CRISPR screen and the top 20 genes of the ORF screen [false discovery rate (FDR) <0.05] are highlighted. The asterisk indicates a mutant form of the gene.
findings provide new biochemical insight into the resistance mechanisms to BETis in NMC as well as a rationale for combination therapy of BETis and CDK4/6 inhibition on NMC.

Results

To investigate the question of which drivers could substitute for BRD4-NUT in NMC, we used a CRISPR library containing 10 guide RNAs (gRNAs) per gene to a list of ~500 putative proto-oncogenes [Liao et al. 2017] to a total of ~400 constructs that contained ~150 both wild-type proto-oncogenes and their recurring mutant alleles identified by TUSON. We also included genes that are frequently amplified in cancers [Santarius et al. 2010], identified in the Cancer Gene Census [Futreal et al. 2004], or implicated in cancer hallmarks such as cell proliferation [Sack et al. 2018], anchorage-independent growth [Pavlova et al. 2013], epithelial-to-mesenchymal transition [Taube et al. 2010], etc. as well as ~40 “neutral genes” that behaved in a neutral fashion in a previous genetic screen that looked for cell proliferation regulators [Sack et al. 2018]. We used these libraries to determine which alterations could substitute for BRD4-NUT signaling using a chemical inhibitor of BRD4: JQ1. We performed screens using a NMC cell line for BRD4-NUT signaling using a chemical inhibitor of proliferation regulators (Sack et al. 2018). We used these libraries to determine which alterations could substitute for BRD4-NUT signaling using a chemical inhibitor of BRD4: JQ1. We performed screens using a NMC cell line (NMC1015) that harbors a BRD4-NUT fusion and is sensitive to JQ1 [Grayson et al. 2014].

The schematic of the CRISPR and ORF screens is outlined in Figure 1B and described in detail in the Supplemental Material. In each screen, cells were treated with either DMSO or 200 nM JQ1 for ~17 d. We used the MAGeCK (model-based analysis of genome-wide CRISPR/Cas9 knockout) scoring algorithm (Li et al. 2014) and edgeR analysis (Robinson et al. 2010) to rank the performance of individual genes in the CRISPR and ORF screen, respectively, based on enrichment, comparing the JQ1 treatment group with the DMSO treatment group. The rank and false discovery rate (FDR) of each gene in the two screens are summarized in Supplemental Table S1. The top 10 hits (FDR < 0.05) from the CRISPR screen and top 20 hits (FDR < 0.05) from the ORF screen are shown in Figure 1, C and D. An immediate validation of our screen approach is that MYC, a major downstream target of BRD4-NUT that can replace its function to prevent differentiation/arrested proliferation induced by the knockdown of BRD4-NUT (Grayson et al. 2014), scored as the number one hit in the oncogene screen.

Notably, the novel cancer drivers that we identified largely fall into the following five categories: (1) MYC-related genes: MYCN (wild type and c.131C > T; p.P44L), MAX (c.179G > A; p.R60Q), and FBXW7; (2) the receptor tyrosine kinase (RTK) signaling pathway: ERBB2, SRC, RRAS2 (wild type and c.216A > C; p.Q72H), H-RAS, Neuronalubristatosis type 1 (NURST1), MAP2K1 (c.371C > A; p.P124Q), and PIK3CA (c.3140A > G; p.H1047R); (3) cell cycle regulation: CCND1, CCND3 (c.869T > G; p.I290R), and RB1; (4) the G-protein-coupled receptor (GPCR)/cyclic AMP (cAMP)/cAMP-dependent protein kinase A (PKA) signaling pathway: GNAS (c.2530C > T; p.R844C) and PRKAR1A; and (5) the transforming growth factor β (TGF-β) pathway: SMAD4 and TGFB2.

Given the stronger phenotype (larger fold change) induced by the oncogenes compared with TSGs, we primarily focused our efforts on characterization of the novel oncogene hits.

Mutant RRAS2 attenuates the effect of JQ1 by sustaining ERK pathway activation during BRD4-NUT inhibition

One of the top hits identified in our oncogene screen is RRAS2, a Ras-related GTPase with transforming potential [Chan et al. 1994; Graham et al. 1994]. Downstream effectors of RRAS2 include three members of the mitogen-activated protein kinases (MAPKs); namely, ERK1/2, c-Jun N-terminal kinase, and p38 MAPK as well as phosphoinositide 3-kinase (PI3K) [Graham et al. 1994; Movilla et al. 1999; Rosario et al. 2001; Murphy et al. 2002]. The amino acid Q72 of RRAS2 is conserved in K-Ras, H-Ras, and N-Ras (Q61) in humans [Fig. 2A]. Q61 mutations in KRAS are commonly found in human cancers, and mutations such as Q61L have been shown to increase KRAS activity through higher affinity to RAF kinase compared with wild type [Hunter et al. 2015]. Consistent with this, both wild-type and Q72H mutant RRAS2 scored in our screen, and the mutant form showed a stronger phenotype (larger fold change) in the screen [Fig. 1D].

To explore how RRAS2 impacts JQ1 resistance in NMC cells, we first validated the effect of expression of Q72H mutant RRAS2 on JQ1 resistance using an independent NMC cell line, NMC797 cells [Toretsky et al. 2003]. As seen with NMC1015 cells, expression of mutant RRAS2 significantly increased the survival of NMC797 cells in response to JQ1 treatment, as measured by sulforhodamine B (SRB) assay [Fig. 2B]. To identify the downstream effectors of mutant RRAS2, we examined the two signaling kinases that have been reported previously to be responsible for RRAS2-induced cell transformation: ERK and PI3K.

Surprisingly, JQ1 treatment inhibited ERK signaling measured by ERK1/2 phosphorylation (p-ERK1/2) and phosphorylation of its downstream effector, P90RSK (p-P90RSK), starting as early as 6 h and reduced the phosphorylation of those two proteins to nearly undetectable levels by 24 h in NMC1015 cells [Fig. 2C]. However, expression of RRAS2Q72H abolished the effect of JQ1 on ERK signaling [Fig. 2C]. JQ1 also decreased the phosphorylation of AKT (p-AKT) and phosphorylation of its downstream effector, PRAS40 (p-PRAS40), at 6 h, and phosphorylation was barely detectable at 24 h [Fig. 2C]. Expression of RRAS2Q72H activated AKT and restored PRAS40 phosphorylation in the presence of JQ1, demonstrating that RRAS2Q72H can rescue both of these signaling arms.

We note that JQ1 reduced the protein level of c-Myc [a known JQ1 target] and cyclin D1 [another gene identified in our screen in NMC1015 cells] with kinetics similar to its effect on ERK and AKT signaling. Cells expressing mutant RRAS2 also showed higher levels of c-Myc and...
cyclin D1, providing an attractive mechanism of action (Fig. 2C). Because ERK and AKT can regulate both of these pathways, we investigated their contributions to c-Myc and cyclin D1 expression downstream from RRAS2. Cells were cotreated with JQ1 and an ERK inhibitor (SCH772984) or an AKT inhibitor (GDC-0068). As noted above, JQ1 suppressed p-ERK and p-P90RSK and dramatically reduced the expression of c-Myc and cyclin D1, whereas RRAS2 largely restored ERK and P90RSK phosphorylation and increased c-Myc and cyclin D1 expression levels. Notably, SCH772984 reversed the effects of RRAS2 on p-ERK and p-P90RSK (Fig. 2D). Interestingly, the higher levels of c-Myc and cyclin D1 in mutant RRAS2-expressing NMC1015 cells during JQ1 treatment were also reduced to that of control cells by SCH772984 treatment. SCH772984 did not further reduce the p-P90RSK level of control NMC1015 cells, suggesting that JQ1 alone potently inhibited ERK signaling in these cells.

As expected, GDC-0068 reduced p-PRA540 levels of mutant RRAS2-expressing cells to that of control cells treated with JQ1 alone. However, it had less of an effect on c-Myc and cyclin D1 in mutant RRAS2-expressing cells than the ERK inhibitor [Fig. 2E]. Together, these results suggest that the RRAS2 mutant attenuates the suppressive effect of JQ1 on c-Myc and cyclin D1 levels, primarily through sustained ERK signaling, although sustained AKT may also contribute to a lesser extent.

The observation that ERK inhibitors could effectively reverse the effects of RRAS2 on these important downstream signals in JQ1-treated cells suggested that ERK inhibitors might also negate the resistance conferred by RRAS2 in JQ1-treated cells. Consistent with signaling changes, while SCH772984 did not further reduce the viability of JQ1-treated control cells, SCH772984 resensitized mutant RRAS2-expressing cells to JQ1 [Fig. 2F]. The contribution of ERK signaling to JQ1 resistance is further supported by the fact that the upstream kinase of ERK1/2, MEK1 (encoded by MAP2K1), also scored highly in the oncogene screen [Fig. 1D]. Thus, sustained ERK signaling significantly contributes to the ability of RRAS2 to overcome JQ1’s effect on survival.

KLF4 mediated JQ1 resistance in NMC cells through restoring the E2F and MYC gene expression program upon JQ1 treatment

Another novel hit identified in our oncogene screen is the Yamanaka factor KLF4. KLF4 is a zinc finger-containing
transcription factor that plays a critical role in regulating a variety of cellular processes such as proliferation, differentiation, development, maintenance of normal tissue homeostasis, and apoptosis. It can function as a TSG or oncogene in different cancers, depending on the cellular context (Rowland and Peeper 2006). As an oncogene, KLF4 has been shown to mediate resistance to a variety of therapies such as chemotherapy and Her2 inhibition (Farrugia et al. 2015; Lund et al. 2017). However, how it might modulate BRD4-NUT dependency is unclear.

We first validated the effect of KLF4 expression on JQ1 resistance in NMC cells using a 3-d SRB assay. Surprisingly, expression of KLF4 did not provide NMC cells with resistance to JQ1 as measured by this short-term viability assay (Supplemental Fig. S1A,B). However, when we examined the effect of KLF4 expression on JQ1 resistance using a longer-term colony formation assay, which is more reflective of the initial screen, KLF4 expression significantly enhanced the colony formation of NMC1015 cells in the presence of JQ1 (Fig. 3A). One possible explanation is that KLF4 may take substantial time to remodel the cell state like Yamanaka factors have been suggested to work (Polo et al. 2012). Therefore, we pretreated KLF4-expressing NMC cells in the presence of JQ1 (Fig. 3B). The full analyses of differentially expressed genes (DEGs) across different conditions are shown in Supplemental Table S2. Among the top genes that are differentially expressed comparing KLF4-expressing and control cells under JQ1 treatment, MYC, its downstream target, TERT; and several long non-coding RNAs (lncRNAs) such as CCAT1 and CASC19 stood out (Fig. 3C). Notably, the transcriptional down-regulation of those genes by JQ1 is attenuated by KLF4 expression only at late [7 d] but not early [6 and 24 h] time points, which is consistent with the observation that KLF4-expressing cells manifest a resistant phenotype only in long-term, but not short-term, survival assays.

CCAT1 and CASC19 are near the MYC locus, and CCAT1 has been reported to play oncogenic roles through transcriptional up-regulation of MYC (Xiang et al. 2014; McCleland et al. 2016). Previously, Aleksyenko et al. (2015) had shown that CCAT1 and CASC19 are among the few genes that are within BRD4-NUT megadomains.
that overlap in multiple NMC cell lines, including NMC797 and NMC1015 cells, and whose expression is significantly reduced by JQ1 treatment. We validated the RNA-seq results using RT-qPCR (Fig. 3D). Our data indicate that expression of KLF4 prevents the down-regulation of CCAT1 and CASC19 by JQ1, which might result in higher MYC levels and thus resistance to JQ1. Indeed, gene set enrichment analysis (GSEA) revealed that E2F targets, MYC targets, and G2M checkpoint genes comprised the top-ranking gene sets when comparing the KLF4-expressing NMC1015 cells with control cells under JQ1 treatment at day 7 [Fig. 3E–G]. In agreement with this, KLF4-expressing cells exhibited higher levels of E2F1, c-Myc, and cyclin D1 during JQ1 treatment compared with control cells [Fig. 3H]. Taken together, these data suggest that KLF4-expressing cells are able to sustain the MYC and E2F gene expression programs under JQ1 treatment and bypass the cell cycle arrest induced by JQ1, providing a second distinct mechanism of resistance involving these same pathways.

**The central role of cell cycle regulators in mediating JQ1 resistance**

One recurring theme from our screens is cell cycle regulation. RB1 is a TSG that negatively regulates E2F-dependent transcription and cell cycle entry (Dyson 2016). During G1/S cell cycle progression, CDK4 and CDK6 form a complex with their regulatory subunit, D-type cyclins, which phosphorylates the Rb protein, resulting in its dissociation from E2F and activation of transcription of genes that are important for S phase, such as cyclin A (Schulze et al. 1995). Two central G1/S transition regulators—cyclin D1 and a mutant form of cyclin D3 (c.869T > G; p.I290R) that is highly stable (Schmitz et al. 2012)—scored as top 10 hits in the oncogene screen. In addition, the downstream inactivation target of the CDK4/6–Cyclin D complex, the RB1 gene, scored in the top 10 of TSG screens, and FRXW7, an F-box protein that targets cyclin E and c-Myc for degradation (Koep et al. 2001; Welcker et al. 2004; Yada et al. 2004), also scored highly in the TSG screen, further implicating the cell cycle.

We first validated the effect of expression of cyclin D1 and mutant cyclin D3 on JQ1 resistance using an independent NMC cell line, NMC797 cells. As seen with the NMC1015 cells in the screen, expression of cyclin D1 or mutant cyclin D3 significantly increased the survival of NMC797 cells under JQ1 treatment [Fig. 4A,B]. To validate that loss of RB1 mediates JQ1 resistance, we used CRISPR to knock out RB1 in NMC797 cells. Cells expressing Cas9 and two independent gRNAs targeting RB1 showed significantly increased survival in response to JQ1 compared with cells expressing Cas9 and control gRNAs targeting AAUS1 or GFP [Fig. 4C]. Thus, these cell cycle regulators mediate JQ1 resistance in multiple NMC cell lines.

We next returned to the NMC1015 cells to explore how JQ1 affects the CDK4/6–cyclin D/Rb signaling axis by examining the levels of these proteins in NMC1015 cells. JQ1 treatment reduced cyclin D1 levels, Rb phosphorylation (p-Rb), and E2F1 levels in uninfected parental or control infected NMC1015 cells [Fig. 4D]. Like MYC, ectopic expression of D-type cyclins restored p-Rb and E2F1 levels in the presence of JQ1. Together, our data suggest that JQ1 down-regulates cyclin D1, whose reduction results in hypophosphorylation of Rb and cell cycle arrest. Blocking Rb activation by JQ1 through either RB1 loss or up-regulation of cyclin D1 or D3 is sufficient to induce JQ1 resistance.

**The CDK4/6 inhibitor synergizes with JQ1 on NMC in vitro**

Given that c-Myc can activate expression of D-type cyclins and the ability of ectopic expression of D-type cyclins to mediate JQ1 resistance, we hypothesize that the endogenous CDK4/6–cyclin D/Rb axis may play an important role in modulating JQ1 sensitivity and that CDK4/6 inhibition should synergize with JQ1. To test this hypothesis, we cotreated NMC1015 cells with JQ1 and the CDK4/6 inhibitor palbociclib and measured cell viability. We found that the combination of JQ1 and palbociclib achieved significantly higher cell proliferation inhibition than either drug alone across a range of doses [Fig. 5A]. We observed such synergism also on NMC797 cells [Fig. 5B]. To further quantify the synergistic effect between JQ1 and palbociclib and rule out the possibility of additive effect, we calculated the combination indices (CIs) of a range of different combinations of JQ1 and palbociclib on NMC1015 cells. A CI between 0 and 1 indicates a

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synergistic effect between two drugs, and the closer the index is to 0, the stronger the synergism [Chou and Talalay 1984]. We found that palbociclib showed strong synergism (CI < 0.3) with JQ1 across a range of doses [Fig. 5C]. To determine whether and how the combination of the two drugs may affect cell cycle, we examined cell cycle distribution under single or combinations of drug treatment using EdU labeling. While a small percentage of cells was still able to incorporate EdU and progress through the cell cycle with treatment by single agents, cotreatment of palbociclib and JQ1 induced complete cell cycle arrest and abolishment of EdU incorporation in both NMC797 and NMC1015 cells [Fig. 5D]. Consistent with this, cotreatment of palbociclib and JQ1 profoundly reduced the levels of E2F1 and its downstream target, cyclin A [Fig. 5E], which accumulates in S phase and is necessary for initiation and completion of DNA replication. Cotreatment of palbociclib and JQ1 did not induce cell death through mechanisms such as apoptosis, necrosis, or autophagic cell death, marked by cleaved-PARP (c-PARP) and Caspase-Glo 3/7 assay (data not shown), phosphorylation of RIP1 (p-RIP1), or LC3-II/LC3-I ratio, respectively [Fig. 5E], suggesting that the profound cell viability reduction induced by the combination treatment in vitro was primarily due to cell cycle arrest.

The CDK4/6 inhibitor synergizes with JQ1 on NMC in vivo

To validate the effectiveness of the combination therapy on NMC in vivo, we used a xenograft mouse model. Tumors generated from NMC1015 cells were established, and once they reached 200–300 mm³, mice were randomized into one of the four following treatment groups and treated for 21 d using well-established preclinical doses and regimens: (1) vehicle, (2) JQ1 alone, (3) palbociclib alone, and (4) JQ1 plus palbociclib. Despite a small initial weight loss seen in the palbociclib and combination treatment groups, the weight of mice of both groups recovered to the start point by the end of treatment, and no other signs of toxicity were observed [Supplemental Fig. S2A]. A spider plot of tumor growth is shown in Figure 6A. The tumors in the vehicle group grew rapidly to the maximum size threshold within 8 d, recapitulating the aggressive nature of NMC in human patients. While single treatment of JQ1 or palbociclib caused no tumor shrinkage and only slowed tumor growth, cotreatment of JQ1 and palbociclib caused tumor regression in the majority of tumors, and tumor size remained stable throughout the course of treatment. In only 8 d, when tumor size necessitated the sacrifice of vehicle-treated mice, the drug combination therapy exerted substantial effects on every tumor and induced frank regression in five out of seven tumors up to 45.1% [Fig. 6B,C]. During the course of treatment [21 d], all animals treated with vehicle or single agents had to be euthanized with the exception of one in the palbociclib-alone group; however, no animals died in the presence of the combination [Fig. 6D]. Even when drug treatment ceased, this drug combination significantly improved the median survival of the mice over each agent or vehicle alone [combination: 34 d; palbociclib: 16 d; JQ1: 12 d; vehicle: 8 d] [Fig. 6D]. Finally, as seen in vitro, palbociclib and JQ1 cotreatment profoundly reduced the expression of the cell growth marker Ki-67 and the levels of E2F1 and its downstream target, cyclin A, in vivo as measured by immunohistochemical staining and immunoblot analysis, respectively [Fig. 6E,F], suggesting that these agents induce a more complete cell cycle arrest compared with each agent alone. To investigate whether the tumor shrinkage induced by the combination treatment in vivo...
was due to apoptosis, we examined the c-PARP and cleaved caspase substrate levels in different treatment groups. Surprisingly, there was more apoptosis in the vehicle-treated tumors (Supplemental Fig. S2B). Such increased proliferation along with an increased apoptotic rate has been observed in other cancer types (Liu et al. 2001). However, compared with each agent alone, palbociclib and JQ1 cotreatment significantly increased the levels of p-RIP1 (Fig. 6G), suggesting the tumor shrinkage induced by the combination treatment in vivo was due to increased necroptosis.

Discussion

NMC is a particularly lethal cancer that does not benefit from conventional chemotherapy and has an overall survival of only 6.9 mo (French et al. 2008; Bauer et al. 2012). Thus, it represents a model disease for the clinical testing of BETis given the direct oncogenic driver role that BRD4-NUT plays in NMC and the urgent need for novel therapeutic approaches. Despite the promising responses seen in some patients treated with BETis, why most patients fail to respond or why even the responders eventually relapse is unknown. Understanding which signaling pathways can modulate the sensitivity of NMC tumors to BETis could inform new therapeutic approaches to treat this deadly cancer. Here, using high-throughput loss-of-function and gain-of-function screening technologies, we systematically explored ~900 known drivers of tumorigenesis for their ability to impact NMC’s response to BETis with the hope that some of these pathways might contribute to NMC tumorigenesis in a basal state and that inhibiting these pathways might improve the efficacy of BETi therapy. We identified multiple cancer driver networks that can genetically modify BRD4-NUT dependency to partially compensate for it and drive proliferation and survival of NMC cancer cells during BETi treatment. Unlike the MYC locus, the regulatory regions and gene bodies of those cancer drivers do not fall in BRD4-NUT megadomains (Supplemental Fig. S3A), suggesting that those cancer drivers are not the direct targets of BRD4-NUT in this disease and compensate for BRD4-NUT inhibition in an indirect manner. Overall, those
cancer drivers that we identified largely fall into six categories [Fig. 7].

The first category encompasses MYC and MYC-related genes such as MYCN, encoding N-Myc. Structurally, the coding regions of both MYC and MYCN are highly homologous. Both c-Myc and N-Myc form heterodimers with Max at consensus E-box sequences (CANNTG) and regulate transcription. Functionally, c-Myc and N-Myc are partially redundant but have several distinct spatiotemporal expression patterns and functions [Rickman et al. 2018]. In addition to the previously validated expression patterns and functions (Rickman et al. 2018), we recovered wild-type MYCN and a mutant form: MYCN [c.131C > T; p.P44L]. The N-MycP44L mutation is recurrently identified in different cancers but remains to be characterized biochemically [Rickman et al. 2018]. Our results suggested that N-MycP44L is indeed an activating mutation given its stronger phenotype (i.e., greater fold change) than wild-type N-Myc in our screen. The fact that many of the mutant alleles picked up by TUSON Explorer achieved stronger phenotypes than their corresponding wild-type alleles in our oncogene screen [Supplemental Table S1] confirmed the robustness of the TUSON Explorer algorithm to identify bona fide activating oncogenic mutations from passenger mutations. We also recovered a mutant form of MAX [c.179G > A; p.R60Q] in our oncogene screen. The MaxR60Q mutation is recurrently identified in many cancers and is suggested to be an activating mutation through inhibition of Max homodimerization but not disruption of c-Myc/Max heterodimerization, thus shifting the balance toward c-Myc activation in proliferating cells [Dela Cruz et al. 2016]. We also recovered the tumor suppressor FBXW7—the F-box substrate adaptor component of the SCFFbx7 E3 ubiquitin ligase that targets c-Myc for degradation [Welcker et al. 2004; Yada et al. 2004]—in our loss-of-function screen. Together, our data confirmed the central role of c-Myc signaling in mediating the oncogenic function of BRD4-NUT and revealed that there are many potential ways for NMC cancer cells to sustain MYC function, thus bypassing the need for BRD4-NUT during BRD4 inhibition.

The second class of genes encompasses the RTK signaling pathway. We recovered ERBB2, SRC, PIK3CA (c.3140A > G; p.H1047R), and several RAS-related genes, including wild-type RRAS2, mutant RRAS2 [c.216A > C; p.Q72H], and wild-type H-RAS, in our oncogene screen as well as their negative regulator, NF1 [Ohba et al. 2000], in the TSG screen. We further showed that the RRAS2Q72H mutant mediates JQ1 resistance predominately through ERK signaling. The importance of ERK signaling was further supported by the fact that the upstream kinase of ERK1/2, MAP2K1, with an activating mutation [c.371C > A; p.P124Q] that harbors RAF-independent kinase activity [Carlino et al. 2015] scored highly in our screen. Previous work has demonstrated that a mechanism of resistance to BETis in ovarian cancers involves the activation of RTKs and their downstream signaling pathways [Kurimchak et al. 2016]. Our results suggest that activation of RTK signaling can mediate BETi resistance in NMC cancer cells as well.

The third general class of suppressors involves the GPCR/cAMP/PKA signaling pathway. We identified GNAS [c.2530C > T; p.R844C] in our oncogene screen. GNAS encodes the α subunit of the stimulatory G protein that transduces signals from GPCR signaling to the cAMP/PKA pathway. Activating mutations in GNAS promote tumorigenesis through activation of PKA, resulting in up-regulation of Wnt and MAPK/ERK signaling [Wilson et al. 2010; Regard et al. 2011; Nomura et al. 2014]. In agreement with the gain-of-function screen, a negative regulator of PKA, PRKARIA, was one of the top-scoring TSGs recovered in our loss-of-function screen. PRKARIA encodes the type 1α regulatory subunit [R1α] of PKA. Loss-of-function mutations of PRKARIA are responsible for the Carney complex [CNC] [Bertherat et al. 2009; Horvath et al. 2010], a multiple neoplasia syndrome. Its inactivation acts as a tumorigenic signal through increasing PKA activity, resulting in ERK signaling activation, Wnt signaling activation, c-Myc activation, up-regulation of D-type cyclins and E2F1, increased cell cycle rates, and decreased apoptosis [Robinson-White et al. 2003, 2006a,b; Nadella and Kirschner 2005; Nesterova et al. 2008; Almeida et al. 2010; Basso et al. 2014]. Given the important role that ERK signaling and cell cycle entry play in BETi resistance in our screens, their up-regulation through increased PKA activity could be a plausible mechanism of GNAS and PRKARIA loss-mediated BETi resistance in NMC and is worthy of further investigation.

The fourth class involves the TGF-β pathway. The TGF-β pathway is known to play both anti-tumorigenic and protumorigenic roles in cancer biology, depending on the tumor type, stage, genetic background, and microenvironment as well as the status of Smad and other pathways [Inman 2011]. Up-regulation of TGF-β signaling has been proposed as a mechanism of BETi resistance in colon cancer cells [Shi et al. 2016]. However, we found that loss of SMAD4 or, with slightly less potency, TGFBR2, two major components of the canonical TGF-β signaling pathway, mediates BETi resistance in NMC in our screen. We also found that the TGF-β signaling is one of the top signaling pathways transcriptionally induced by JQ1 treatment in NMC1015 cells [Supplemental Fig. S4A]. Thus, it is likely that activation of TGF-β signaling could mediate part of the anti-cancer effect of BETis on NMC.

A fifth class is a catchall category of several unrelated drivers such as the Yamanaka factor KLF4, MDM4, and PCBP1 [c.299T > C; p.L100P] in the oncogene screen and
TRIP12, ARID2, and KEAP1 in the TSG screen. Of these, we characterized the mechanism of KLF4-mediated JQ1 resistance. We found that KLF4-expressing cells are able to sustain the E2F and c-Myc transcription program during BETi treatment. Given the central role that MYC plays in mediating the oncogenic function of the BRD4-NUT fusion protein, it is not surprising that KLF4 and other modifiers of BETi treatment might regulate MYC function. Whether and how the other novel hits recovered in our screens impact MYC function are worthy of future investigation.

The final class of suppressors involves cell cycle regulation. Our data revealed the central role of cell cycle regulators in mediating the oncogenic function of BRD4-NUT and demonstrated that there are multiple ways for NMC cancer cells to bypass the cell cycle arrest induced by BETIs—through RB1 loss or up-regulation of D-type cyclins. This suggested that reinforcing cell cycle arrest might improve the efficacy of BETIs. Critically, we showed that CDK4/6 inhibition synergizes with BETIs against NMC tumors in vitro and in vivo. Importantly, CDK4/6 inhibitors such as palbociclib, ribociclib, and abemaciclib have been approved in clinics for the treatment of certain breast cancers. Therefore, the pharmacokinetic and pharmacodynamic profiles of the relevant drugs have been established. Our results on synergy and the favorable clinical properties of the relevant inhibitors of these pathways suggest that this combination forms the basis of a clinical trial for combination therapy of BETIs and CDK4/6 inhibitors on NMC.

In summary, analysis of cancer drivers’ effects on BRD4-NUT dependency in NMC has provided a broad picture of the genetic landscape of resistance to BETIs in this cancer. These pathways now become candidates to explore in NMC tumors that evolve BETi resistance during therapy. In addition, beyond the synergism that we identified with CDK4/6 inhibitors, future therapies that reinforce other aspects of the mechanism of BETi inhibition of tumorigenesis might also show activity in combination therapy to further improve the efficacy of BETIs.

Materials and methods

Cell culture and reagents

NMC1015 and NMC797 cells were kindly provided by Dr. Christopher A. French [Brigham and Women’s Hospital and Harvard Medical School]. 293T cells were purchased from American Type Culture Collection. NMC1015, NMC797, and 293T cells were maintained in DMEM-Glutamax (Thermo Fisher Scientific) supplemented with 10% heat-inactivated fetal bovine serum (GE Healthcare HyClone), 100 IU/mL penicillin, and 100 µg/mL streptomycin in a humidified atmosphere of 95% air and 5% CO2 at 37°C. SCH772984 and GDC-0068 were purchased from Selleck Chemicals. JQ1 used in vitro was purchased from Sigma-Aldrich as described previously (Sanjana et al. 2014).

293T cells were transduced with vector DNA, pRev, pTag, pHV Gag/pol, and pVSVG. Viruses were harvested 48 h after transfection and filtered (45-µm pore size). Transit-293 (Mirus) was used to transfect 293T cells.

Cells were transduced with 4 µg/mL polybrene (Sigma), and infected cells were selected using 1 µg/mL puromycin (Clontech) for 3 d. pLentiCRISPR-V2-infected cells were subjected to cell viability assay or immunoblot analysis 1 wk after infection to allow for genome modification by Cas9. Cells expressing ORFs were treated with 1 µg/mL Dox for 3 d prior to subsequent experiments for ORF expression and maintained in medium containing 1 µg/mL Dox during subsequent experiments.

Western blot analysis

The Western blot method is described in detail in the Supplemental Material. The following antibodies were from Cell Signaling: CA4PDH [8884], phospho-ERK1/2 (T202/Y204; 4370P), total ERK1/2 [4695P], phospho-AKT [S473, 4060P], total AKT [2920S], phospho-p90RSK [Ser380, 9341S], c-Myc [5605S], cyclin D1 [2978S], phospho-PRAS40 [Thr246; 2997S], KLF4 [4038S], phospho-Rb [Ser780, 3590S], total Rb [9309P], E2F1 [3742S], c-PARP [9541S], cleaved caspase substrate motif [8698S], phospho-RIP1 [Ser166, 65746S], total RIP1 [3493T], and LC3B [3868T]. Other antibodies used were cyclin A (Santa Cruz Biotechnology, sc-751), vinculin (Sigma, V9131), and RRAS2 (Abcam, ab182264).

 Colony formation assay

Single-cell suspensions were seeded into six-well plates [200 cells per well] and incubated overnight before continuous treatment of DMSO or 200 nM JQ1 for the indicated times, with drug and medium replaced every 3 d. At the end of treatment, cells were fixed with 10% trichloroacetic acid, washed, and stained with methyl blue (Sigma-Aldrich). Colony numbers of each condition were quantified using ImageJ and normalized to control cells under DMSO treatment.

Cell viability assay

The effects of drugs on cell viability were determined by SRB (Sigma-Aldrich) as described previously [Vichai and Kittikara 2006].

Cell cycle analysis

Cells seeded in six-well plates [5 × 104 cells per well] were treated with DMSO or 200 nM JQ1 for 48 h. Cells were pulsed with 10 µM EdU final concentration for 1 h using the Click-IT Plus Edu Alexa fluor 647 flow cytometry assay kit (Thermo Fisher Scientific) and then processed according to the manufacturer’s instructions. Samples were analyzed by flow cytometry (BD LSR II).

RT-qPCR

Total RNA was isolated using the RNAsesy minikit (Qiagen), and cDNA was synthesized using SuperScript IV (Invitrogen).

The sequences targeted by the gRNAs used were GFP (GGGCG AGGACGGTGTACCCG), AAV5 (GGGGCCACTAGGGACA GGAT), RB1 gRNA1 (GGTTCTTTGACCAATGGG), and RB1 gRNA2 (GCAGTGTGATTATTCTGGAG).

cDNAs were picked from the ORFeome version 8.1 or Ultimate ORF libraries. Stop codons and mutations were introduced using the QuikChange II XL site-directed mutagenesis kit (Agilent Technologies). cDNAs were subcloned into the pHAGE-TREx-BC-DEST vector via LR recombinase reaction (Invitrogen).

To produce lentiviruses, 293T cells were transfected with vector DNA, pRev, pTag, pHV Gag/pol, and pVSVG. Viruses were harvested 48 h after transfection and filtered (45-µm pore size). Transit-293 (Mirus) was used to transfect 293T cells.

Cells were transduced with 4 µg/mL polybrene (Sigma), and infected cells were selected using 1 µg/mL puromycin (Clontech) for 3 d. pLentiCRISPR-V2-infected cells were subjected to cell viability assay or immunoblot analysis 1 wk after infection to allow for genome modification by Cas9. Cells expressing ORFs were treated with 1 µg/mL Dox for 3 d prior to subsequent experiments for ORF expression and maintained in medium containing 1 µg/mL Dox during subsequent experiments.

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RT-qPCR

Total RNA was isolated using the RNAsesy minikit (Qiagen), and cDNA was synthesized using SuperScript IV (Invitrogen).
according to manufacturer’s instructions. RT-qPCR was performed in triplicate using the TaqMan gene expression master mix (Invitrogen) with TaqMan gene expression assay (Life Technologies) on an Applied Biosystems Fast 7500 machine using GAPDH as the endogenous normalization control. The IDs for the TaqMan assays used were as follows: GAPDH (Hs09999905_m1), MYC (Hs00153408_m1), CCAT1 (Hs04402620_m1), and CASC19 (Hs04405851_g1).

**Gene expression profiling**

NMC1015 cells were transduced with pHAGE-TREx-KLF4 or empty vector. Infected cells were treated with 1 µg/mL puromycin [Clontech] for 3 d. Cells were then passaged for 1 wk and treated with 1 µg/mL Dox [Clontech] for 3 d to allow for gene expression. Cells were then treated with 200 nM JQ1 in medium containing 1 µg/mL Dox for 0 h, 6 h, 24 h or 7 d (two replicates for each condition). Total RNA was isolated using the RNeasy minikit [Qiagen]. RNA-seq libraries were generated using NEBNext Ultra RNA library preparation kit (New England Biolabs). Fifty-base-pair single-end sequencing was performed using an Illumina NextSeq. Reads were aligned to the hg19 genome using Hisat2 (Pertea et al. 2016), transcripts and frequencies were assessed from the aligned data by subread (Liao et al. 2013), and edgeR (Robinson et al. 2010) was used to identify DEGs and generate the signed LR scores used for GSEA (http://www.broadinstitute.org/gsea; Subramanian et al. 2005).

**Xenograft mouse model**

The xenograft mouse model is described in the Supplemental Material.

**Statistical analysis**

Data are presented as the mean ± SD, and significance was analyzed using the two-sided Student’s t-test unless specified otherwise. Differences were considered significant when $P < 0.05$.

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Author contributions: S.L. and S.J.E. developed the concept and wrote the manuscript. S.L. designed and performed experiments and analyzed the data, and S.J.E. supervised the study. O.M. and K.C. provided expertise in xenograft mouse models.

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