Cancer Cell

Treatment-Induced Tumor Dormancy through YAP-Mediated Transcriptional Reprogramming of the Apoptotic Pathway

Graphical Abstract



Authors

Kari J. Kurppa, Yao Liu, Ciric To, ..., Henry W. Long, Nathanael S. Gray, Pasi A. Jänne

Correspondence

pasi_janne@dfci.harvard.edu

In Brief

Kurppa et al. show that YAP activation mediates resistance to combined EGFR/ MEK inhibition by inducing dormancy in non-small-cell lung cancer cells. Targeting the YAP pathway, in part by using a newly developed covalent TEAD inhibitor, promotes apoptosis of the dormant therapy-resistant cancer cells.

Highlights

- Loss of EGFR signaling leads to senescence-like dormancy in *EGFR*-mutant lung cancer
- YAP promotes survival and dormancy in the absence of EGFR downstream signaling
- YAP/TEAD/SLUG suppress apoptosis through transcriptional repression of BMF
- A TEAD inhibitor enhances EGFR inhibitor-mediated apoptosis and prevents dormancy

Kurppa et al., 2020, Cancer Cell *37*, 104–122 January 13, 2020 © 2019 Elsevier Inc. https://doi.org/10.1016/j.ccell.2019.12.006





Treatment-Induced Tumor Dormancy through YAP-Mediated Transcriptional Reprogramming of the Apoptotic Pathway

Kari J. Kurppa,^{1,23} Yao Liu,^{2,3,23} Ciric To,¹ Tinghu Zhang,^{2,3} Mengyang Fan,^{2,3} Amir Vajdi,⁴ Erik H. Knelson,¹ Yingtian Xie,^{1,5} Klothilda Lim,^{1,5} Paloma Cejas,^{1,5} Andrew Portell,^{1,6} Patrick H. Lizotte,^{1,6} Scott B. Ficarro,^{2,7,8,21} Shuai Li,⁹ Ting Chen,⁹ Heidi M. Haikala,¹ Haiyun Wang,¹⁰ Magda Bahcall,¹ Yang Gao,^{11,12} Sophia Shalhout,^{13,22} Steffen Boettcher,^{1,14} Bo Hee Shin,¹ Tran Thai,¹ Margaret K. Wilkens,¹⁵ Michelle L. Tillgren,¹⁵ Mierzhati Mushajiang,¹ Man Xu,^{1,6} Jihyun Choi,¹

(Author list continued on next page)

¹Department of Medical Oncology, Dana-Farber Cancer Institute and Harvard Medical School, Boston, MA 02215, USA

²Department of Cancer Biology, Dana-Farber Cancer Institute, Harvard Medical School, Boston, MA 02215, USA

³Department of Biological Chemistry and Molecular Pharmacology, Harvard Medical School, Boston, MA 02215, USA

⁴Department of Informatics and Analytics, Dana-Farber Cancer Institute, Boston, MA 02215, USA

⁵Center for Functional Cancer Epigenetics, Dana-Farber Cancer Institute, Boston, MA 02215, USA

⁶Belfer Center for Applied Cancer Science, Dana-Farber Cancer Institute, Boston, MA 02215, USA

⁷Blais Proteomics Center, Dana-Farber Cancer Institute, Harvard Medical School, Boston, MA 02215, USA

⁸Department of Oncologic Pathology, Dana-Farber Cancer Institute, Boston, MA 02215, USA

⁹Division of Hematology & Medical Oncology, Laura and Isaac Perlmutter Cancer Center, New York University Langone Medical Center, New York, NY, USA

¹⁰School of Life Science and Technology, Tongji University, 200092 Shanghai, China

¹¹Department of Pediatric Oncology, Dana-Farber Cancer Institute, Boston, MA 02215, USA

(Affiliations continued on next page)

SUMMARY

Eradicating tumor dormancy that develops following epidermal growth factor receptor (EGFR) tyrosine kinase inhibitor (TKI) treatment of *EGFR*-mutant non-small cell lung cancer, is an attractive therapeutic strategy but the mechanisms governing this process are poorly understood. Blockade of ERK1/2 reactivation following EGFR TKI treatment by combined EGFR/MEK inhibition uncovers cells that survive by entering a senescence-like dormant state characterized by high YAP/TEAD activity. YAP/TEAD engage the epithelial-to-mesenchymal transition transcription factor SLUG to directly repress pro-apoptotic *BMF*, limiting drug-induced apoptosis. Pharmacological co-inhibition of YAP and TEAD, or genetic deletion of *YAP1*, all deplete dormant cells by enhancing EGFR/MEK inhibition-induced apoptosis. Enhancing the initial efficacy of targeted therapies could ultimately lead to prolonged treatment responses in cancer patients.

INTRODUCTION

Epidermal growth factor receptor (EGFR) tyrosine kinase inhibitors (TKIs) are the standard of care for patients with advanced *EGFR*-mutant non-small cell lung cancer (NSCLC) (Mok et al., 2009; Rosell et al., 2012; Soria et al., 2018). However, acquired resistance inevitably develops, limiting clinical efficacy (Cortot and Jänne, 2014). In most cases, resistance arises after a dramatic initial response followed by a stable minimal residual disease (MRD), or dormant, state, with subsequent gradual growth

Significance

Genotype-directed therapy rarely, if ever, leads to complete tumor eradication. The residual tumors, following drug treatment, can serve as reservoirs for the development of acquired drug resistance. Using *EGFR*-mutant lung cancer as a model system, we identify YAP activation as a major counter regulatory mechanism promoting cell survival and dormancy following sustained inhibition of EGFR and its downstream signaling, through transcriptional repression of the proapoptotic protein BMF. Targeting YAP, indirectly using a tankyrase inhibitor or directly using a TEAD inhibitor, enhances the apoptotic effects of EGFR/MEK inhibition. Our findings suggest that targeting YAP/TEAD following genotype-directed therapy may be one strategy to enhance initial drug-induced apoptosis, reduce residual disease, and as such lead to improved treatment outcomes. Arrien A. Bertram,¹ Benjamin L. Ebert,^{1,14,16} Rameen Beroukhim,^{1,14,17} Pratiti Bandopadhayay,^{11,14,18} Mark M. Awad,¹ Prafulla C. Gokhale,^{6,15} Paul T. Kirschmeier,^{1,6} Jarrod A. Marto,^{2,7,8,21} Fernando D. Camargo,^{13,22} Rizwan Haq,^{17,19} Cloud P. Paweletz,^{1,6} Kwok-Kin Wong,⁹ David A. Barbie,¹ Henry W. Long,^{1,5} Nathanael S. Gray,^{2,3} and Pasi A. Jänne^{1,6,20,24,*}

¹²Department of Pediatrics, Harvard Medical School, Boston, MA 02215, USA

¹³Stem Cell Program, Boston Children's Hospital, Boston, MA 02215, USA

¹⁴Broad Institute of MIT and Harvard, Cambridge, MA 02142, USA

¹⁵Experimental Therapeutics Core, Dana-Farber Cancer Institute, Boston, MA 02210, USA

¹⁶Howard Hughes Medical Institute, Dana-Farber Cancer Institute, Boston, MA 02215, USA

¹⁷Department of Medicine, Harvard Medical School, Boston, MA 02115, USA

¹⁸Dana-Farber/Boston Children's Cancer and Blood Disorders Center, Boston, MA 02115, USA

¹⁹Department of Molecular and Cellular Oncology, Dana-Farber Cancer Institute, Boston, MA 02215, USA

²⁰Lowe Center for Thoracic Oncology, Dana-Farber Cancer Institute, 450 Brookline Avenue, LC4114, Boston, MA 02215, USA

²¹Department of Pathology, Brigham and Women's Hospital and Harvard Medical School, Boston, MA 02215, USA

²²Department of Stem Cell and Regenerative Biology, Harvard University, Cambridge, MA 02138, USA

²³These authors contributed equally

24Lead Contact

*Correspondence: pasi_janne@dfci.harvard.edu https://doi.org/10.1016/j.ccell.2019.12.006

of a drug-resistant tumor. Previous preclinical studies suggest that, following EGFR TKI treatment, *EGFR*-mutant tumor cells can enter a drug-tolerant state, reminiscent of dormancy in patients, allowing cells to evade apoptosis and survive (Hata et al., 2016; Sharma et al., 2010). Over time, the drug-tolerant cells can acquire resistance through mutational or non-mutational mechanisms (Hata et al., 2016). While the establishment of this state seems to be largely stochastic and dictated mostly by epigenetic mechanisms (Guler et al., 2017; Sharma et al., 2010), the mechanistic bases of how cancer cells evade the initial apoptosis in response to drug treatment—the absolute requirement to enter the drug-tolerant state—or maintain tolerance in the presence of drug, are poorly understood.

Our previous work demonstrated that despite sustained EGFR inhibition following EGFR TKI treatment, reactivation of ERK1/2 occurs within just a few days (Ercan et al., 2012; Tricker et al., 2015). Concomitant inhibition of MEK effectively prevents reactivation of ERK1/2, results in a greater initial apoptotic response, and leads to a more durable tumor control in vitro and in vivo than single-agent EGFR inhibition (Ercan et al., 2012; Tricker et al., 2015). The EGFR (osimertinib) and MEK (selumetinib) inhibitor combination has been studied in patients resistant to previous EGFR TKIs and is also under evaluation as initial therapy for advanced EGFR-mutant NSCLC in a phase II clinical trial (NCT03392246; Ramalingam et al., 2019). However, even with this combination, acquired resistance can still develop (Tricker et al., 2015). In this study we set out to elucidate the mechanisms that allow cancer cells to evade apoptosis and survive despite combined EGFR/MEK inhibition.

RESULTS

Combined EGFR and MEK Inhibition Results in a Stable but Reversible Dormant State

Combined EGFR/MEK inhibition prevents the reactivation of ERK1/2 following EGFR inhibition and delays the onset of drug resistance *in vitro* and *in vivo* (Tricker et al., 2015) (Figure S1A). In PC-9 cells, treatment with single-agent osimertinib (O) leads

to re-colonization of wells within 8 weeks (Figure 1A). The combination of O and the MEK inhibitor trametinib (T) prevents any measurable regrowth (Figure 1A). However, few viable cells can still be detected after 15 weeks of treatment (Figure 1B). We used live-cell imaging and observed that the OT-treated cells surviving the initial apoptosis remained in a largely non-proliferative, or dormant, state throughout the treatment period. However, within days following drug withdrawal the cells began to proliferate and re-colonize the wells (Figure 1C, Video S1). This phenomenon was consistent across EGFR-mutant NSCLC cell lines (Figures 1C and S1B). These observations suggest that while combined EGFR/MEK inhibition eliminates cells in which reactivation of ERK signaling occurs following singleagent EGFR inhibition, a separate population enters a dormant state, surviving combined EGFR/MEK inhibition. There was no evidence of reactivation of EGFR and/or ERK signaling in the dormant cells during treatment (Figure 1D) and EGFR signaling was restored in cells that grew following drug washout (Figure 1D). These cells were still sensitive to OT, and morphologically indistinguishable from the untreated cells (Figure S1C), suggesting that we did not select out a subclone with a pre-existing resistance mutation (Hata et al., 2016). To formally address whether the establishment of dormancy following OT treatment is pre-determined or a stochastic process, we barcoded PC-9 cells using the EvoSeq library (Feldman et al., 2019), treated the cells with DMSO, gefitinib (G), O, or OT for 3 weeks, sequenced DNA from the remaining cells and analyzed the findings as described (Bhang et al., 2015) with some modifications. We observed a large fraction of shared barcodes within the G-treated (data not shown) and O-treated (Figures 1E, 1F, and S2A) cells, strongly suggesting selection of pre-existing clones, consistent (for G) with previous studies (Hata et al., 2016). In contrast, the vast majority of barcodes in the OTtreated cells were unique (Figures 1E, 1F, and S2A). Comparison of the shared barcodes between O and OT cells demonstrated that 89% of the barcodes identified in the O group are not present in the OT group (Figure S2B). These findings suggest that, while resistance to O likely occurs through a selection process of a pre-existing clone, the ability of cells to enter



Figure 1. Combined EGFR/MEK Inhibition Promotes a Senescence-like Dormant State

(A) Proliferation of PC-9 cells treated with DMSO, 100 nM osimertinib (O) alone, or in combination with 30 nM trametinib (T).

(B) Images of control cells (at 1 week) or dormant PC-9 cells (at 15 weeks). Scale bars, 200 $\mu m.$

(C) Cells were treated as in (A) for 6 weeks followed by drug washout.

(D) Western blot analysis of EGFR downstream signaling following treatment with OT for the indicated times or 21 days followed by drug washout (rebound).

(E) Fraction of barcodes shared among replicates following indicated treatments in barcoded PC-9 cells.

(F) Relative abundance of individual barcodes. Shared and unique indicate barcodes shared by > 2 or ≤2 replicates, respectively.

dormancy following OT is predominately driven by a stochastic process.

Dormant State following EGFR/MEK Inhibition Shares Characteristics with Cellular Senescence

To characterize the dormant state, we performed RNA sequencing (RNA-seq) in PC-9, HCC827, and HCC4006 cells following treatment with either DMSO or with OT for 2 weeks. Gene set enrichment analysis (GSEA) revealed an upregulation of gene expression signatures involved in inflammatory response, epithelial-to-mesenchymal transition (EMT), and protein secretion, while cell-cycle-associated gene expression signatures were robustly downregulated (Figure 1G). These findings along with the spread-out, flattened morphology of the dormant cells (Figure 1B) prompted us to guery similarities between the dormant state and cellular senescence. We stained DMSO-. O-. or OTtreated PC-9, HCC827, and HCC4006 cells for senescence-associated β-galactosidase activity (SA-β-Gal) (Debacq-Chainiaux et al., 2009), and noted that for all three cell lines the majority of cells surviving the combination treatment stained positive for SA-B-Gal (Figures 1H and 1I). Further GSEA revealed a significant enrichment of a senescence-associated gene expression signature (Fridman and Tainsky, 2008) in the dormant cells (Figures 1J and S3A). A significantly smaller proportion of O-treated cells demonstrated SA-B-Gal activity compared with those treated with the OT combination (Figures 1H and 1I). Senescent cells characteristically exhibit increased secretion of pro-inflammatory factors (senescence-associated secretory phenotype [SASP]) (Coppé et al., 2010). By analyzing conditioned medium from dormant cells, we observed an increased secretion of several cytokines and chemokines, including the prominent SASP factor interleukin-6, compared with untreated cells (Figure S3B). The expression of several classical SASP factors (Coppé et al., 2008) in the cytokine, chemokine, IGFBP, and MMP families was also upregulated in the dormant cells (Figure S3C), analogous to SASP.

Using immunofluorescence following a 10-day treatment with OT, we detected a robust increase in punctate, H3K9Me³-positive nuclear foci, another hallmark of senescence (Narita et al., 2003) (Figure 1K). Senescence is also invariably associated with the induction of p16^{INK4a}, p21^{Cip1}, and/or p27^{Kip} (Campisi and D'Adda Di Fagagna, 2007). Although p16^{INK4a} or p21^{Cip1} were not consistently induced, we observed a robust induction of p27^{Kip}, which was subsequently downregulated in growing cells following drug withdrawal (Figure S3D).

The Establishment of Dormancy following EGFR/MEK Inhibition Is Critically Dependent on Activation of YAP/TEAD

To explore potential epigenetic changes in the dormant cells, we employed an assay for transposase-accessible chromatin combined with next-generation sequencing (ATAC-seq). We observed a significant difference in the global epigenetic states between the OT-induced dormant versus DMSO-treated cells (Figure 2A), which reverted upon drug washout, demonstrating that the changes acquired at dormancy are reversible (Figure 2A). There was also a significant difference in epigenetic states between single-agent O- and OT-induced dormant cells (Figure 2A). We performed a motif analysis to interrogate transcription factor binding sites associated with the ATAC-seq peaks with higher signal (more accessible chromatin) in OT-induced dormant cells versus control cells (Figure 2B). The three most significantly enriched motifs were the consensus sites for TEAD family transcription factors (Figure 2C), suggesting that the OT-induced epigenetic state is associated with increased TEAD transcription factor binding. The TEAD transcription factors serve as canonical partners for the Hippo pathway effector YAP, which has been associated with resistance to targeted therapy in several contexts, including in resistance to EGFR TKIs in EGFR-mutant NSCLC (Chaib et al., 2017; Hsu et al., 2016). Indeed, we observed a significant enrichment of a YAP/ TEAD gene expression signature (Zhang et al., 2009) in dormant EGFR-mutant cells (Figure 2D). TEAD-binding motifs also scored as the most significant top hits separating the OT- and O-treatment-induced states (Figure 2E). In accordance with these findings, we observed significantly higher YAP/TEAD activity, as measured by CTGF and ANKRD1 expression, in OT-induced dormant PC-9 cells compared with O-treated cells (Figure 2F). Consistently, we also detected increased chromatin accessibility at putative distal enhancer sites upstream of CTGF TSS in OT-induced dormant cells compared with cells treated with O alone (Figure S4A). Taken together, these results demonstrate that dormant cells induced by combined EGFR/MEK inhibition adopt a distinct, reversible epigenetic state distinguished from the untreated or the O-treated state by increased YAP/TEAD activity.

To assess the role of YAP activity in the establishment of OT-induced dormant state, we treated *EGFR*-mutant NSCLC cell lines for 3 weeks with OT with or without the tankyrase inhibitor XAV939, an indirect inhibitor of YAP activity (Wang et al., 2015), and assessed the regrowth of cells after drug washout. Remarkably, the OT/XAV939 combination dramatically reduced the number of dormant cells (Figure S4B), diminishing regrowth (Figure 2G). Similar findings with three additional structurally divergent tankyrase inhibitors were observed (Figure S4B). In addition, we tested the effect of several inhibitors targeting putative resistance pathways to EGFR TKIs, or the effect of chemotherapy in PC-9 cells, but observed little to no effect on the establishment of the dormant cell population (Figure S4C). Notably, the combination of single-agent O and XAV939 was significantly inferior to the OT/XAV939 combination in all

⁽G) GSEA of Hallmark gene sets comparing dormant cells versus DMSO-treated control cells. Normalized enrichment scores (NES) for gene sets with false discovery rate (FDR) < 0.1 in at least two cell lines are shown.

⁽H) Senescence-associated β-galactosidase (SA-β-Gal) staining of cells treated as indicated for 10 days. Scale bars, 100 μm.

⁽I) Quantification of (H).

⁽J) GSEA of senescence signature comparing dormant, OT-treated PC-9 cells versus control cells.

⁽K) Immunofluorescence (IF) staining for H3K9Me³ in control cells or dormant cells treated with OT for 10 days. Scale bars, 20 µm.

Mean ± SEM are shown in all plots except (I) where mean ± SD are shown. ANOVA (I) or t test (K) were used for statistical analyses. ***p < 0.001, **p < 0.01. See also Figures S1–S3.





(B) ATAC-seq signal intensities centered on upregulated (UP) or downregulated (DOWN) peaks in dormant, OT-treated cells versus control cells. (C and D) Analysis for enriched transcription factor motifs (C) and GSEA of YAP/TEAD signature (D) (Zhang et al., 2009).

(E) Left: ATAC-seq signal intensities centered on UP or DOWN peaks in OT-treated versus O-treated cells. Right: analysis for transcription factor motifs enriched in upregulated peaks.

(F) qPCR analysis of YAP target gene expression.

EGFR-mutant NCSLC cell lines tested (Figure 2G), suggesting that the differences seen in YAP/TEAD activity between OT-induced dormant cells and single-agent O-treated cells may reflect different degrees of dependency on YAP.

To further validate the role of YAP in the establishment of the dormant state, we knocked out *YAP1* in three different *EGFR*mutant NSCLC cell lines, including a patient-derived cell line, DFCI243, using the CRISPR/Cas9 system (Figure 2H). Strikingly, *YAP1* knockout (KO) completely abolished the establishment of dormant cells in all cases, measured by lack of regrowth following drug withdrawal after a 3-week treatment with OT (Figure 2I). In contrast, two of three *YAP1* KO cell lines treated with O alone regrew following washout (Figure S4D).

In vivo, OT treatment of mice bearing xenograft tumors from *YAP1* KO PC-9, HCC4006, and DFCI243 cells, or from the corresponding control cells, led to a durable response for the entire 4-week treatment period (Figure 2J). However, the control tumors started to regrow soon after treatment cessation, consistent with the presence of a dormant cell population *in vivo*. In comparison, *YAP1* KO tumors had an increased tumor regrowth latency and significantly smaller tumors at the time of regrowth in all models (Figure 2J), consistent with a reduction in the dormant cell population. Collectively, these results demonstrate that the establishment of a drug-tolerant state following EGFR/MEK inhibition, but not single-agent EGFR inhibition, is critically dependent on YAP/TEAD activity.

YAP Activation Is Necessary for Cancer Cell Viability upon Combined EGFR/MEK Inhibition

To monitor YAP/TEAD activity following drug treatment, we introduced a fluorescent YAP/Hippo pathway reporter (Mohseni et al., 2014) into PC-9 cells (PC-9 YAP reporter cells) and used live-cell imaging to track YAP activity over time. The OT treatment robustly induced YAP activity (Figure 3A), which was completely blocked by the addition of XAV939 (Figure 3A). Consistently, we noted decreased phosphorylation of the main YAP upstream negative regulator, LATS1, and a decrease in YAP S127 phosphorylation, which regulates YAP cytosolic retention (Figure S5). Consequently, YAP nuclear localization was significantly increased in *EGFR*-mutant NCSLC cells following a 10-day OT treatment, but not following single-agent O treatment (Figure 3B), in agreement with the more prominent increase in YAP activity observed in the combination-treated cells (Figures 2F and 3A).

Treatment-induced activation of YAP suggested that active YAP protects cells from the initial apoptosis. To evaluate this possibility, we used non-invasive monitoring of caspase-3/7 activity over time in the PC-9 YAP reporter cells. An increase in YAP activation occurred proportionally to apoptosis (Figures 3C and 3D), and cells with high YAP activity (YAP^{high} cells) were significantly less likely to undergo apoptosis (Figure 3D; odds ratio 0.26 at 80 h, p < 0.0001 by Fisher's exact test). Consistently, XAV939/OT treatment increased apoptosis in *EGFR*-mutant NSCLC cells compared with OT or to O/XAV939 alone (Figures 3E and 3F). Also, the *YAP1* KO cells underwent heightened and accelerated apoptosis in response to OT (Figure 3G). Importantly, this hypersensitive phenotype was rescued by the re-expression of wild-type *YAP1* in the *YAP1* KO cells (Figure 3H).

The OT combination triggered significantly higher YAP reporter activity than single-agent O (Figure 3A), suggesting that the differences in YAP/TEAD activity seen in the drug-tolerant state (Figures 2E and 2F) reflect the cells' immediate responses to the different drug treatments. As the main consequence of concomitant MEK inhibition is the suppression of ERK1/2 reactivation following EGFR inhibition, our results suggest that ERK1/2 reactivation and YAP activation are two separate means by which EGFR-mutant NSCLC cells evade apoptosis following single-agent O treatment. We quantified the proportion of YAPhigh cells in single-agent O- and OT-treated populations using the PC-9 YAP reporter cells. Following a 10-day treatment, the O-treated cell population contained both YAP^{high} (40%) and YAP-negative (60%) cells, whereas the OT treatment largely depleted YAP-negative cells, leaving behind mostly YAPhigh cells (>80%) (Figure 3I). Taken together, these observations demonstrate that chronic downregulation of EGFR and its downstream signaling by concomitant EGFR and MEK inhibition selects for cells that induce high YAP activity upon treatment, creating a vulnerability that can be exploited to selectively promote apoptosis in these cells through simultaneous inhibition of YAP (Figure 3J).

YAP-High, Senescence-like Dormant State Also Occurs In Vivo

To study the dormant state in vivo, we performed single-cell RNA-seq (scRNA-seq) in cells from PC-9 xenograft tumors treated with OT until MRD (3 weeks) (Figures 4A, 4B, and S6A). We also performed scRNA-seq in dormant PC-9 cells following 3 weeks of OT treatment in vitro. We detected a significant increase in cells enriched for YAP, EMT, or senescence gene expression signatures in the OT-treated PC-9 cells in vitro and in vivo (Figure 4C). We also analyzed YAP expression and subcellular localization from the same PC-9 MRD in vivo samples using immunohistochemistry. Consistent with the scRNA-seq studies, we detected an increase in active, nuclear YAP in the MRD tumors, with more intense nuclear staining in the combination-treated tumors (Figure 4D and 4F). We further evaluated MRD tumors from genetically engineered EGFR^{L858R/T790M} mice (Zhou et al., 2009) following 2 weeks of O treatment and similarly noted an increase in YAP nuclear localization (Figures 4E and 4F). As these mice have an intact immune system, we evaluated T cell infiltration. We observed an increase in

⁽G) Regrowth of EGFR-mutant NSCLC cells after washout following a 3-week treatment with the indicated drug combinations.

⁽H) Western blot analysis of YAP protein levels in YAP1 knockout (KO) and control (CTRL) cells.

⁽I) Proliferation of cells in (H) treated as indicated for 21 days, followed by drug washout.

⁽J) Mice bearing CTRL or YAP1 KO cell xenograft tumors were treated with vehicle or OT followed by treatment cessation and follow-up. Data with at least 6/8 live mice per group are plotted. Right: tumor volumes at time of regrowth, indicated by an arrow.

Mean \pm SEM are shown in all plots except (F), where mean \pm SD are shown. ANOVA was used for statistical analyses in all but (J), where t test was used. ***p < 0.001, *p < 0.05. See also Figure S4.



Figure 3. YAP Activation Is Necessary for Cancer Cell Viability upon Combined EGFR/MEK Inhibition

(A) YAP activity following indicated treatments in PC-9 cells transduced with a fluorescent YAP/Hippo pathway reporter (PC-9 YAP reporter cells).
 (B) IF staining for YAP nuclear localization following the indicated treatments. Scale bars, 500 μm.

(C) YAP activity and apoptosis in PC-9 YAP reporter cells treated with OT.

(D) Analysis of overlap between YAP^{high} cells (red) and apoptotic cells (green) after 80 h of treatment in PC-9 YAP reporter cells. Scale bars, 150 µm.

(E) Apoptosis in PC-9 cells treated with the indicated drugs or drug combinations.

(F) Apoptosis in EGFR-mutant NSCLC cells treated as indicated. Peak apoptosis values over 72 h are shown.

(G) Apoptosis in YAP1 KO or CTRL cells treated as indicated.

infiltration of CD4+ and CD8+ T cells (Figures 4G and S6B), suggesting that the MRD tumors elicit an immune response, consistent with our findings of an increase in secreted inflammatory factors (Figures S2B and S2C) and with previous studies in lung cancer patients (Thress et al., 2017). However, despite the T cell response, neither O- nor O/selumetinib-treated EGFR^{L858R/T790M} mice could be cured by the treatment (Figure S6C) suggesting that the immune response is insufficient to eradicate the YAP^{high} residual cells.

We further studied tumors from EGFRL858R/T790M mice that had developed acquired resistance to WZ4002 (preclinical third-generation EGFR inhibitor)/T combination from our previous study (Tricker et al., 2015) and detected robust YAP nuclear staining and a lack of pERK1/2 expression in the resistant tumor nodules (Figure 4H). A significantly higher proportion of nuclear YAP-positive cells in WZ4002/T-resistant nodules was detected compared with single-agent WZ4002-resistant nodules (Figure 4H), consistent with our in vitro observations (Figures 2 and 3). Finally, we analyzed YAP nuclear staining and pERK1/2 expression in a tumor from a patient with advanced EGFR-mutant lung cancer treated with O/selumetinib (NCT03392246) who had a sustained partial response. The patient underwent surgery following 11 months of treatment while in a clinical MRD state. The residual tumor demonstrated intense YAP staining and an absence of pERK1/2 staining (Figure 4I).

YAP Mediates the Evasion of Apoptosis by Repressing the Induction of the Pro-apoptotic Protein BMF

We next sought to elucidate the mechanism by which YAP protects EGFR-mutant NSCLC cells from apoptosis. YAP1 KO had no effect on canonical EGFR signaling or on the induction of BIM, a known mediator of apoptosis following EGFR inhibition (Costa et al., 2007; Cragg et al., 2007; Gong et al., 2007) (Figure 5A). This suggests that YAP affects the apoptotic process independently of EGFR signaling and downstream of BIM. Unlike previous reports (Lin et al., 2015; Rosenbluh et al., 2012), we did not observe any significant changes in the levels of anti-apoptotic proteins BCL-XL, BCL2, BCL-w, or MCL-1 in YAP1 KO cells compared with control cells at baseline or following OT (Figure S7A). In contrast, we observed a substantial increase in BAX activity (Figure S7B), cytochrome c release (Figure S7C), and caspase activation (Figure 3G) in YAP1 KO cells in response to OT, suggesting that the increased apoptosis seen in YAP1 KO cells is mediated by the intrinsic apoptotic pathway.

To identify potential YAP target gene(s), we performed RNAseq on PC-9 and HCC4006 YAP1 KO or control cells with and without OT treatment (Figure 5B). Focusing on genes that mediate apoptosis through the activation of caspases (Hallmark Apoptosis gene set, 161 genes) (Liberzon et al., 2015), we identified *BMF* as one of the top upregulated genes in drug-treated YAP1 KO cells compared with drug-treated control cells in both cell lines (Figure 5C). The BMF gene encodes a proapoptotic BH3-only protein that can sequester anti-apoptotic proteins, but, unlike BIM, cannot directly activate BAX or BAK (Bhola and Letai, 2016; Kuwana et al., 2005). Together with BIM induction upon EGFR inhibition (Figure 5A), an increase in BMF in YAP1 KO cells would be expected to lead to increased activation of BAX and thus to enhanced apoptosis, consistent with our observations (Figures 3G, S7B, and S7C). We confirmed the RNA-seq results using qPCR; YAP inhibition by XAV939 or YAP1 KO significantly increased BMF expression in response to OT in EGFR-mutant NSCLC cell lines in vitro and in vivo (Figure 5D). Due to lack of high-affinity antibodies for BMF, we used CRISPR/Cas9 technology to produce N-terminally HAtagged BMF under the endogenous promoter in PC9 (Figure 5E and S7D). In these cells, YAP inhibition or YAP knockdown led to a robust increase in BMF protein levels in response to OT, while BIM levels remained unchanged (Figure 5F). Moreover, re-introduction of wild-type YAP, but not a TEAD-binding-deficient S94A mutant, to the YAP1 KO background completely abolished the increase in BMF expression (Figure 5G). Ectopic overexpression of BMF using a doxycycline-inducible vector in EGFR-mutant NSCLC cells (Figure S7E) induced rapid apoptosis, which was further increased by co-treatment with OT (Figure S7F), demonstrating that induction of BMF alone, without YAP activation, is sufficient to sensitize EGFR-mutant NSCLC cells to apoptosis. Downregulation of BMF expression using small interfering RNA (siRNA) significantly decreased apoptosis in YAP1 KO cells in response to OT (Figures 5H and S7G), demonstrating that the induction of BMF expression is necessary for the increased apoptosis in YAP1 KO cells. Hence, YAP facilitates evasion of apoptosis in EGFR-mutant NSCLC cells by repressing the expression of BMF upon combined EGFR/MEK inhibition, leading to the establishment of the dormant cell population (Figure 5I).

YAP Represses *BMF* Induction by Engaging EMT Transcription Factor SLUG

Next, we investigated the molecular mechanisms by which YAP represses the expression of *BMF*. Although YAP is mostly linked to transcriptional activation, it has also been shown to complex with transcription factors and modulators to drive transcriptional repression, often in association with TEAD (Beyer et al., 2013; Kim et al., 2015; Zaidi et al., 2004). Thus, we hypothesized that the YAP/TEAD complex is directly repressing *BMF* by forming a tertiary complex with a transcriptional repressor.

In search for such transcriptional repressors, we noted that an EMT gene expression signature was highly enriched in the dormant cells induced by OT treatment (Figure 1G and 4C). In addition to EMT being a known mechanism of drug resistance in *EGFR*-mutant lung cancer (Byers et al., 2013; Sequist et al., 2011; Shibue and Weinberg, 2017; Zhang et al., 2012), YAP has been reported to mediate EMT and to directly bind to

⁽H) Left: western blot analysis of YAP protein levels in YAP1 KO cells transduced with wild-type YAP1. Right: cells were treated with OT and analyzed as in (G). Only data from drug-treated cells is shown.

⁽I) Proportions of YAP^{high} cells in PC-9 YAP reporter cell populations treated as indicated. Scale bars, 150 µm.

⁽J) Different means for EGFR-mutant NSCLC cells to avoid apoptosis following EGFR inhibition.

Mean ± SEM are shown in all plots except (I), where SD is shown. ANOVA was used for statistical analyses in all but (D), where Fisher's exact test was used. ***p < 0.001. See also Figure S5.



Figure 4. YAP-High, Senescence-like Dormant State Also Occurs In Vivo

(A) Growth curves of PC-9 xenograft tumors harvested for single-cell RNA-seq (scRNA-seq) and immunohistochemistry (IHC).

(B) Fluorescence-activated cell sorting scheme used to obtain scRNA-seq samples from the dissociated xenograft tumors.

(C) YAP, EMT, and senescence signature enrichments in single cells from the xenograft tumors.

(D and E) IHC staining for YAP in the xenograft tumors (D) or in residual tumors from EGFR^{L858/T790M} mice (E) following 2-week treatment with vehicle or O. Scale bars, 100 μm.

(F) Quantification of (D) and (F).

(G) Quantification of infiltrating T cells in the same tumors as in (E) based in CD4/CD8 IHC.

(H and I) IHC staining for YAP and pERK in WZ4002- or WZ4002/T-resistant tumors from EGFR^{L856/T790M} mice (H) or in a residual tumor of an *EGFR*-mutant NSCLC patient following treatment with O/selumetinib for 11 months (I). Scale bars, 800 μm (H), 100 μm (I).

Kolmogorov-Smirnov test (C), ANOVA (F) when more than two groups, (H) or t test (F) when two groups, (G) and (I) were used for statistical analyses. ***p < 0.001, **p < 0.01; n.s., not significant. See also Figure S6.



Figure 5. YAP Mediates the Evasion of Apoptosis by Repressing the Induction of Pro-apoptotic BMF

(A) Western blot analysis of EGFR downstream signaling following 24 h treatment as indicated.

(B) RNA-seq samples used in (C).

- (C) Expression of genes regulating apoptosis in OT-treated YAP1 KO cells versus OT-treated CTRL cells. Colors indicate log2 fold change values with p < 0.001. (D) qPCR analysis of *BMF* expression in CTRL or YAP1 KO cells treated as indicated for 24 h *in vitro* or 3 days *in vivo*.
- (b) deck analysis of Birr expression in CTRE of TAPT KO certs treated as indicated for 24 min vitro of

(E) Schematic representation of the endogenous *BMF* locus in PC-9 HA-BMF cells.

(legend continued on next page)

canonical EMT transcription factors, including SNAIL, SLUG, and ZEB1 (Lehmann et al., 2016; Tang et al., 2016). We therefore explored the possibility that EMT, the development of a dormant state, and the requirement for YAP in mediating evasion of apoptosis through repression of BMF following drug treatment, were all linked. In PC-9 and HCC4006 YAP1 KO cells following 24 h OT treatment, compared with control cells, the EMT signature was negatively enriched, suggesting that YAP is triggering the EMT program in EGFR-mutant NSCLC cells (Figure 6A). gPCR analysis revealed that SNAI2, encoding SLUG, was the dominantly expressed EMT transcription factor in EGFR-mutant NCSLC cell lines (Figure 6B). We further observed that endogenous YAP, TEAD, and SLUG proteins co-immunoprecipitate in both PC-9 and HCC4006 cells upon 48 h OT treatment (Figure 6C). Knockdown of SLUG by siRNA in PC-9 and HCC4006 cells resulted in a significant increase in BMF expression following treatment with OT, similar to that observed following YAP knockdown (Figure 6D and 6E), and the increase in BMF expression translated into a robust increase in apoptosis upon treatment (Figure 6F). These results suggest that members of the YAP/TEAD/SLUG complex co-operate to repress BMF expression and thus suppress apoptosis in response to OT treatment. To confirm that the YAP/TEAD/SLUG complex directly binds the BMF locus to mediate repression, we performed chromatin immunoprecipitation followed by next-generation sequencing (ChIP-seq) using antibodies against endogenous YAP, TEAD4, and SLUG in PC-9 cells treated either with DMSO or OT. We detected a robust increase in YAP and SLUG binding to chromatin after 48 h of OT treatment, while TEAD4 chromatin binding was less affected (Figure 6G). Specifically, we observed overlapping YAP, TEAD, and SLUG peaks at the BMF promoter region as well as at nearby H3K27Ac-positive enhancer regions upon treatment (Figure 6H), demonstrating that the YAP/TEAD/SLUG repressor complex directly binds to the BMF locus. Taken together, these results provide a mechanistic explanation for YAP-mediated suppression of proapoptotic signaling through the engagement of TEAD and the EMT program to directly repress the induction of BMF expression upon combined EGFR/MEK inhibition (Figure 6I).

Development of a Novel Covalent TEAD Inhibitor to Target YAP Dependency upon Combined EGFR/MEK Inhibition

The strict dependency of OT-treated cells on YAP presents an attractive target for drug development. Although our results point toward TEAD as the main mediator of YAP effects in this context (Figures 2C, 2D, and 6H), we wanted to further confirm whether other effector pathways downstream of YAP might also play a role. The YAP protein has several functional domains, many of which mediate protein-protein interactions (Piccolo et al., 2014). We systematically mutated the key functional domains

in YAP (Figure 7A), and determined which of the mutants could rescue the apoptotic phenotype imparted by *YAP1* deficiency following OT treatment in PC-9 cells. We observed that introduction of *YAP1* with a mutation in the TEAD-binding domain (S94A) (Zhao et al., 2008), or with a deleted transactivation domain (TAdel; Figure 7A), had the least ability to rescue *YAP1* deficiency (Figure 7B), confirming that YAP-mediated evasion of apoptosis is absolutely dependent on TEAD, as well as on intact transactivation domain.

TEAD, as a transcription factor, is presumed undruggable. However, recent studies revealed a hydrophobic pocket for the post-translational palmitoylation of TEAD (Chan et al., 2016; Noland et al., 2016), and flufenamic acid as a molecule binding to this pocket (Pobbati et al., 2015). Flufenamic acid/ TEAD2 co-crystal revealed extensive hydrophobic interactions as its main binding mode (Pobbati et al., 2015), providing a structural basis for the rational design of a covalent TEAD inhibitor through an acrylamide as a covalent warhead to react with the conserved Cys380 on TEAD2. We reasoned that the trifluoromethylated phenyl ring forms extensive hydrophobic interactions, whereas the carboxylic acid of flufenamic acid, proximal to Cys380, might be replaced by an acrylamide warhead to react with the cysteine. Hence, MYF-01-37 (Figure 7C) was developed as a covalent binder to TEAD, targeting Cys380 when incubated with the TEAD2 protein (C359 in TEAD1) (Figures S8A-S8C). Pretreating cells with MYF-01-37 led to loss of direct TEAD pull-down by biotin-MYF-01-037 (Figure S8D) from whole-cell lysate (Figure S8E), confirming MYF-01-037 binding to TEAD in cells. This target engagement of TEAD resulted in inhibition of direct YAP/TEAD interaction (Figure 7D) in HEK293T cells and in the reduction in canonical YAP target gene CTGF expression in PC-9 cells (Figure 7E). This reduction was abrogated by the overexpression of a TEAD1 C359S mutant that disrupts the covalent binding of the drug to TEAD, but not by overexpression of wild-type TEAD1 (Figure 7E), demonstrating that the observed inhibition of YAP activity is due to on-target covalent binding of the compound to TEAD. Importantly, XAV939, which inhibits YAP activity via a TEAD-independent mechanism (Wang et al., 2015), was still able to inhibit CTGF expression also in cells expressing the TEAD1 C359S mutant (Figure 7E). As a single agent, MYF-01-37 had minimal impact on cell viability of several EGFR-mutant NCSLC cell lines (Figure S8F), which is consistent with the apparent dispensability of YAP activity in EGFRmutant NSCLC cells at steady state (Figure 2I). When combined with OT, MYF-01-37 completely suppressed the increased YAP activity induced by OT treatment in PC-9 YAP reporter cells (Figure 7F), led to a robust increase in BMF expression (Figure 7G), and to subsequent increase in apoptosis in PC-9 and HCC4006 cells compared with OT alone (Figure 7H), thus phenocopying the effects of tankyrase inhibition or YAP1 KO

⁽F) Western blot analysis of BMF, BIM, and YAP expression in PC-9 HA-BMF cells transfected with non-targeting (NT) or YAP siRNA and treated as indicated for 24 h.

⁽G) qPCR analysis of BMF expression in CTRL or YAP1 KO cells transduced as indicated, and following treatment with either DMSO or OT for 24 h.

⁽H) Peak apoptosis over 72-h treatment in PC-9 and HCC4006 cells transfected with NT or BMF siRNA.

⁽I) The mechanism of YAP/TEAD-mediated suppression of apoptosis in EGFR-mutant NSCLC cells following EGFR/MEK inhibition.

Mean \pm SD are shown in all plots except (H), where mean \pm SEM is shown. ANOVA was used for statistical analyses. ***p < 0.001, **p < 0.01; n.s., not significant (p > 0.05). See also Figure S7.



Figure 6. YAP Represses BMF Induction by Engaging EMT Transcription Factor SLUG

(A) GSEA of EMT signature in YAP1 KO versus control cells treated with OT for 24 h.

(B) qPCR analysis of EMT transcription factor expression in untreated EGFR-mutant NSCLC cells.

(C) Co-immunoprecipitation analysis of the interaction between YAP, TEAD, and SLUG in PC-9 cells following treatment with DMSO or OT for 48 h. (D) Western blot analysis of YAP and SLUG protein levels in PC-9 or HCC4006 cells transfected with non-targeting (NT), YAP, or SLUG siRNA.

(E) qPCR analysis of BMF expression in cells in (D) following 24 h treatment with DMSO or OT.

(Figures 3A, 3E, 3G, and 5D). Importantly, a 10-day treatment with MYF-01-37 in combination with OT led to a dramatic decrease in dormant cells compared with OT alone (Figure 7I).

DISCUSSION

Genotype-directed therapy is the standard of care for many cancers that harbor an activated oncogene (Blanke et al., 2008; Drilon et al., 2018; Peters et al., 2017). While this treatment approach has transformed cancer care for many genomic subtypes of cancer, these therapies are rarely, if ever, curative. One explanation for such observations is the inability of genotype-directed therapies to eradicate all tumors cells. In *EGFR*-mutant NSCLC, a complete response is observed only in a small minority (<5%) of patients following treatment with EGFR TKI (Mok et al., 2017; Soria et al., 2018). As *EGFR* mutations are truncal mutations (i.e., in every cell of a tumor) (Jamal-Hanjani et al., 2017), it is not clear why a proportion of tumor cells can survive initial EGFR inhibitor-induced apoptosis and subsequently persist in the presence of drug treatment.

The development of EMT, as a drug-resistant state following EGFR inhibitor treatment, has been observed both in model systems and in lung cancer patients (Byers et al., 2013; Sequist et al., 2011; Zhang et al., 2012). In addition, hyperactivation of the Hippo pathway effector YAP has been shown to dampen the efficacy of targeted treatment in several contexts (Zanconato et al., 2016), including the efficacy of EGFR TKIs in *EGFR*-mutant NCSLC (Hsu et al., 2016; Chaib et al., 2017). However, the mechanistic basis for these observations remains largely unexplored. Here we provide a mechanistic link for these two different observations and demonstrate that a critical transcription factor mediating EMT, SLUG, and YAP together leads to transcriptional repression of *BMF* following EGFR/MEK treatment and as such limits the initial drug-induced apoptotic effect allowing the formation of a dormant state.

Apoptosis in response to EGFR TKIs in EGFR-mutant NCSLC is executed by the intrinsic apoptotic pathway and invariably associated with the upregulation of BIM (Costa et al., 2007; Cragg et al., 2007; Gong et al., 2007). BIM levels are suppressed by the MAPK pathway, both transcriptionally and post-transcriptionally (Ley et al., 2005), and thus mechanisms which uncouple EGFR inhibition from ERK1/2 inhibition would be expected to block EGFR inhibitor-mediated upregulation of BIM, promoting cell survival (Ercan et al., 2012; Tricker et al., 2015). O can also activate YAP, allowing drug-induced cell survival through a completely different mechanism (Figures 3A, 3I, 3J, and 4F). Thus, single-agent EGFR TKI treatment can lead to both ERK1/ 2 reactivation and YAP activation, whereas, upon combined EGFR/MEK inhibition, activation of YAP becomes the dominant survival mechanism in EGFR-mutant NSCLC cells (Figures 3A, 3I, 3J, 4F, 4H, and 4I). These results suggest that YAP can maintain the viability of EGFR-mutant NSCLC cells in the chronic absence of EGFR and its downstream signaling. Intriguingly,

the ability of YAP to compensate for the loss of dominant oncogene in MAPK-dependent cancers has been previously shown in the context of mutant *KRAS*-driven NSCLC and pancreatic ductal adenocarcinoma (Kapoor et al., 2014; Shao et al., 2014). In these studies, *YAP1* overexpression (Shao et al., 2014) or *YAP1* amplification (Kapoor et al., 2014) rescued the loss of *KRAS* in a MAPK pathway-independent mechanism.

Overexpression of YAP and its paralog TAZ has been shown to induce EMT in a TEAD-dependent manner (Lei et al., 2008; Zhang et al., 2009; Zhao et al., 2008). Considering that YAP activation seems to be a specific adaptation mechanism in cells that cannot re-activate EGFR downstream signaling, the YAP/ TEAD/SLUG interplay repressing BMF may be the immediate response protecting cells undergoing a YAP-dictated global change in cellular state. Analogously, Shao et al. (2014) also found that the YAP-mediated bypass of KRAS loss was associated with the acquisition of a mesenchymal state, suggesting that YAP may drive the EMT program as a mechanism to adapt to loss of oncogene signaling in other cancer contexts as well. Whether our observations on YAP-mediated suppression of apoptosis through transcriptional repression of BMF also extend to other genotype-directed cancer therapies remains unknown and will need to be evaluated in future studies. Also, we cannot rule out a possibility that, in addition to YAP/TEAD/ SLUG, other factors are involved in the long-term survival of EGFR-mutant cancer cells treated with OT.

Interestingly, the dormant state resulting from YAP/TEAD activation shared several characteristics with cellular senescence. Unlike treatment-induced senescence in response to DNA-damaging chemotherapeutic agents (Ewald et al., 2010), *EGFR*-mutant NSCLC cells seem to only reversibly adopt the senescence program upon EGFR/MEK inhibition to tolerate the lethal drug exposure, and revert back to the normal steady state upon drug withdrawal. Consequently, the senescent-like population, at least in this context, can serve as a reservoir of dormant cells that are later, upon acquisition of additional resistance mechanisms, capable of re-establishing the tumor and drive clinically observed drug resistance.

The therapeutic vulnerability of *EGFR*-mutant NSCLC to YAP/ TEAD antagonism identified in this study led us to develop a novel covalent TEAD inhibitor, MYF-01-37. As YAP is widely associated with resistance to cancer therapies, we also tested the effect of TEAD inhibition and/or YAP1 KO in other genotype or TKI combination contexts within the NSCLC space, including in *ALK*-rearranged, *MET*-amplified, and *EGFR*-mutant *MET*-amplified models, and observed increased apoptosis following YAP/TEAD co-targeting in most models (Figures 8A–8C). These data suggest wide potential for co-targeting YAP/TEAD with genotype-directed therapy. In accordance with our observations that *YAP1* loss has negligible consequences in *EGFR*-mutant NSCLC cells at steady state, MYF-01-37 did not demonstrate single-agent toxicity (Figure S8F). This is in stark contrast to a recently published, structurally similar covalent TEAD inhibitor,

(I) The mechanism by which YAP/TEAD/SLUG complex represses *BMF* expression upon combined EGFR/MEK inhibition.

⁽F) Apoptosis in cells in (D) following treatment with DMSO or OT.

⁽G) Number of peaks called by MACS2 (FDR < 0.01).

⁽H) ChIP-seq signal traces in BMF locus. H3K27Ac was used to identify enhancer regions.

 $Mean \pm SD (E) \text{ or mean} \pm SEM (F) \text{ are shown. ANOVA was used for statistical analyses. } ***p < 0.001, **p < 0.01.$



Figure 7. Development of Novel Covalent TEAD Inhibitor to Target YAP Dependency upon Combined EGFR/MEK Inhibition (A) YAP1 mutants used in the rescue experiment in (B). BD, binding domain.

(B) Viability (CellTiter-Glo) of CTRL cells or PC-9 YAP1 KO cells transduced with YAP1 mutants (A) following 72 h treatment with OT.

(C) Top: the structure of MYF-01-37. Bottom: MYF-01-37 binding to the palmitoylation pocket in TEAD1 based on molecular docking. The cysteine 359 targeted by MYF-01-37 is indicated.

(D) Effect of MYF-01-37 or the corresponding reversible control on YAP/TEAD interaction measured using Split Gaussia Luciferase Assay.

(E) Left: western bot analysis of the expression of myc-tagged TEAD1 in PC-9 cells transduced as indicated. Right: qPCR analysis of CTGF expression after 24 h treatment with XAV939 or MYF-01-37 in the transduced PC-9 cells.

(F) YAP activity in PC-9 YAP reporter cells after 72 h treatment with OT or OT in combination with XAV939 (XAV) or MYF-01-37 (MYF).

(legend continued on next page) Cancer Cell 37, 104–122, January 13, 2020 117 TED-347 (Bum-Erdene et al., 2019), (Figure S8F), which is toxic most likely due to its covalent warhead. Unlike the highly reactive α -chloroketone covalent warhead in TED-347, the acrylamide warhead in MYF-01-37 is more suitable for covalent targeting of proteins in living cells, and thus most likely contributes to the low non-specific toxicity of MYF-01-37 as a single agent (De Cesco et al., 2017; Liu et al., 2013). Further development is needed to optimize the pharmacological properties of MYF-01-37 to enable preclinical testing of the compound using *in vivo* models of *EGFR*-mutant NSCLC.

Ultimately, the strategy of co-targeting EGFR, MEK, and YAP/ TEAD to enhance the initial treatment efficacy in EGFR-mutant NSCLC and limit the establishment of the dormant state, will need to be tested in a clinical trial. Although EGFR and MEK inhibitors can be administered together (NCT03392246; Ramalingam et al., 2019), three-drug combinations raise the concern for toxicity. Auspiciously, YAP appears dispensable for normal homeostasis in many adult organs, suggesting that targeting YAP might be well tolerated (Zanconato et al., 2016). Also, as our findings reveal, the main role of YAP1 loss is in enhancing the initial apoptotic effect of EGFR/MEK inhibition. Thus, it is plausible that a three-drug combination would be necessary only transiently, followed by a two-drug treatment, thus reducing potential toxicity. In support of this approach, we observed identical potency when we treated PC-9 cells for 1 week with OT/ XAV939 or MYF-01-37 followed by 2 weeks of OT compared with a 3-week continuous OT/XAV939 or MYF-01-37 treatment (Figure 8D). The potential different treatment approaches will need to undergo clinical evaluation to determine both their efficacy and toxicity.

STAR***METHODS**

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

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j. ccell.2019.12.006.

ACKNOWLEDGMENTS

The authors would like to thank the following for valuable help with this study: David Feldman and Paul Blainey from the Massachusetts Institute of Technology in Cambridge, MA, for providing research materials and expertise for the barcoding experiment; Zachary Herbert and Maura Berkeley from the Molecular Biology Core Facility at the Dana-Farber Cancer Institute for the sequencing services; Dana-Farber/Harvard Cancer Center in Boston, MA, for the use of the Specialized Histopathology Core, which provided histology and immunohistochemistry service. Dana-Farber/Harvard Cancer Center is supported in part by an NCI Cancer Center Support Grant no. NIH 5 P30 CA06516; the members of the NYU Experimental Pathology Research Laboratory, which is partially supported by the Cancer Center Support Grant P30CA016087 at NYU Langone's Laura and Isaac Perlmutter Cancer Center, for their expertise and immunohistochemistry support.

This work was supported by the National Cancer Institute grants R01CA135257 (to P.A.J.) and R35CA220497 (to P.A.J.), The American Cancer Society (CRP-17-111-01-CDD) (to P.A.J.), the Claudia Adams Barr Program for Innovative Cancer Research (to P.A.J.), the Ildiko Medve and Adria Sai-Halasz EGFR Lung Cancer Research Fund (to P.A.J.), Balassiano Family Fund for Lung Cancer Research (to P.A.J.), the Jane and Aatos Erkko Foundation (to K.J.K.), the Instrumentarium Science Foundation (to K.J.K.), and the Orion Research Foundation (to K.J.K.), S.B. was supported by fellowships from the Swiss Cancer League (KLS-3625-02 2015) and Swiss National Science Foundation (P300PB_161026/1 and P400PM_183862), Swizerland. J.A.M. was supported by NIH R01 CA222218 and CA233800.

AUTHOR CONTRIBUTIONS

Conceptualization, K.J.K., P.A.J., T.Z., N.S.G., and R.H.; Investigation, K.J.K., Y.L., C.T., M.F., E.H.K., K.L., A.P., P.H.L., S.B.F., S.L., T.C., H.M.H., M.B., Y.G., S.S., B.H.S., T.T., M.K.W., M.L.T., and M.M.; Resources, M.X., J.C.,

(H) Apoptosis in PC-9 and HCC4006 cells treated as indicated.

Mean ± SEM are shown in all plots except (E), where mean ± SD is shown. ANOVA was used for statistical analyses. ***p < 0.001, **p < 0.01. See also Figure S8.

⁽G) qPCR analysis of BMF expression in cells in (E) following 24 h treatment as indicated.

⁽I) Regrowth of PC-9 and HCC4006 cells after drug washout following a 2-week treatment as indicated.



Figure 8. Co-targeting YAP/TEAD with Genotype-Directed Therapy

(A and B) Apoptosis in NSCLC cell lines treated as indicated.

(C) Left: western blot analysis of YAP expression in control (CTRL) and YAP1 KO H3122 and EBC-1 cells. Right: apoptosis in CTRL and YAP1 KO H3122 and EBC-1 cells treated as indicated.

(D) PC-9 cells were treated as indicated in the scheme on the left, followed by drug washout. Regrowth of cells was monitored and quantified as in Figure 2G. Mean \pm SEM are shown. ANOVA was used for statistical analyses. *** p < 0.001.

P.T.K., D.A.B., F.D.C., P.B., R.B., A.A.B., K.-K.W., and M.M.A.; Methodology, S.B., B.L.E., P.C., H.W.L., and C.P.P.; Formal Analyses, Y.X., H.W., and A.V.; Writing, K.J.K. and P.A.J.; Funding Acquisition, P.A.J. and N.S.G.; Supervision, P.A.J., N.S.G., T.Z., P.C.G., and J.A.M.

DECLARATION OF INTERESTS

P.A.J. has received consulting fees from AstraZeneca, Boehringer Ingelheim, Pfizer, Roche/Genentech, Merrimack Pharmaceuticals, Chugai Pharmaceuticals, Ariad Pharmaceuticals, Eli Lilly and Company, Araxes Pharama, Ignyta, Novartis, Mirati Therapeutics, Takeda Oncology, Daiichi Sankyo, Biocartis, Voronoi, SFJ Pharmaceuticals, and LOXO Oncology; receives post-marketing royalties from DFCI-owned intellectual property on EGFR mutations licensed to Lab Corp; has sponsored research agreements with AstraZeneca, Daichi Sankyo, Boehringer Ingelheim, PUMA, Eli Lilly, Astellas Pharmaceuticals, and Takeda Oncology: and has stock ownership in Gatekeeper Pharmaceuticals and LOXO Oncology. N.S.G. is a founder, science advisory board member (SAB), and equity holder in Gatekeeper, Syros, Petra, C4, B2S, and Soltego. The Gray lab receives or has received research funding from Novartis, Takeda, Astellas, Taiho, Janssen, Kinogen, Voronoi, Her2llc, Deerfield, and Sanofi. R.H. has received research grants from Bristol-Myers Squibb and Novartis. K.K.W. is a founder and equity holder of G1 Therapeutics and has Consulting/Sponsored Research Agreements with AstraZeneca, Janssen, Pfizer, Array, Novartis, Merck, Takeda, Ono, Targimmune, and BMS, C.P.P. has received honoraria from Bio-Rad and AstraZeneca, is a co-founder of Xsphera Biosciences, and is on the scientific advisory board of DropWorks and XSphera Biosciences. P.B. receives grant funding from Novartis Institute of Biomedical Research for an unrelated project. J.A.M. serves on the SAB of 908 Devices.

Received: May 17, 2019 Revised: October 11, 2019 Accepted: December 10, 2019 Published: January 13, 2020

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STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rabbit monoclonal (D7A5) anti-phosphorylated EGFR	Cell Signaling	Cat#3777; RRID: AB_2096270
Rabbit anti-EGFR	Cell Signaling	Cat#2232; RRID: AB_331707
Rabbit monoclonal (193H12) anti-phosphorylated AKT	Cell Signaling	Cat#4058; RRID: AB_331168
Rabbit anti-AKT	Cell Signaling	Cat#9272; RRID: AB_329827
Rabbit monoclonal (D13.14.4E) anti- phosphorylated ERK	Cell Signaling	Cat#4370; RRID: AB_2315112
Rabbit anti-ERK	Cell Signaling	Cat#9102; RRID: AB_330744
Rabbit anti-phosphorylated S6	Cell Signaling	Cat#2215; RRID: AB_331682
Rabbit monoclonal (5G10) anti-S6	Cell Signaling	Cat#2217; RRID: AB_331355
Mouse monoclonal (DM1A) anti-Tubulin	Sigma	Cat#T9026; RRID: AB_477593
Rabbit anti-H3K9Me3	Abcam	Cat#Ab8898; RRID: AB_306848
Rabbit monoclonal (D8H1X) anti-YAP	Cell Signaling	Cat#14074; RRID: AB_2650491
Rabbit monoclonal (D7D2Z) anti-CD4	Cell Signaling	Cat#25229; RRID: AB_2798898
Rabbit monoclonal (D4W2Z) anti-CD8α	Cell Signaling	Cat#98941; RRID: AB_2756376
Rabbit monoclonal (EPR8190-6) anti-TTF1	Abcam	Cat# ab133638; RRID: AB_2734144
Rabbit monoclonal (C34C5) anti-BIM	Cell Signaling	Cat#2933; RRID: AB_1030947
Mouse monoclonal (6E2) anti-HA	Cell Signaling	Cat#2367; RRID: AB_10691311
Rabbit monoclonal (C19G7) anti-SLUG	Cell Signaling	Cat#9585; RRID: AB_2239535
Rabbit monoclonal (D3F7L) anti-pan-TEAD	Cell Signaling	Cat#13295;' RRID: AB_2687902
Rabbit monoclonal anti-TEAD4	Abcam	Cat#ab58310; RRID: AB_945789
Rabbit anti-H3K27Ac	Diagenode	Cat#C15410196; RRID: AB_2637079
Rabbit monoclonal (9B11) anti-Myc-tag	Cell Signaling	Cat#2276; RRID: AB_331783
Rabbit anti-HSP90	Santa Cruz	Cat#sc-7947; RRID: AB_2121235
Rabbit monoclonal (D7C1M) anti-p16	Cell Signaling	Cat#80772; RRID: AB_2799960
Rabbit monoclonal (12D1) anti-p21	Cell Signaling	Cat#2947; RRID: AB_823586
Rabbit monoclonal (D69C12) anti-p27	Cell Signaling	Cat#3686; RRID: AB_2077850
Rabbit monoclonal (D9W2) anti-phosphorylated YAP	Cell Signaling	Cat#13008; RRID: AB_2650553
Rabbit anti-phosphorylated LATS1 (S909)	Cell Signaling	Cat#9157; RRID: AB_2133515
Rabbit monoclonal (D57D3) anti-phosphorylated LATS1 (T1079)	Cell Signaling	Cat#8654; RRID: AB_10971635
Rabbit monoclonal (C66B5) anti-LATS1	Cell Signaling	Cat#3477; RRID: AB_2133513
Rabbit monoclonal (54H6) anti-BCL-XL	Cell Signaling	Cat#2764; RRID: AB_2228008
Rabbit monoclonal (D55G8) anti-BCL-2	Cell Signaling	Cat#4223; RRID: AB_1903909
Rabbit monoclonal (D5V5L) anti-MCL1	Cell Signaling	Cat#39224; RRID: AB_2799149
Rabbit monocklonal (31H4) anti-BCL-w	Cell Signaling	Cat#2724; RRID: AB_10691557
Rabbit monoclonal (D2E11) anti-BAX	Cell Signaling	Cat#5023; RID: AB_10557411
ApoTrack™ Cytochrome c Apoptosis WB Antibody Cocktail	Abcam	Cat#ab110415
Mouse anti-activaed BAX	Incvitrogen	Cat#MA5-14003; RRID: AB_10979735
Rabbit anti-MEK		Cat#9122; RRID: AB_823567
Goat anti-rabbit HRP-linked antibody	Cell Signaling	Cat#7074; RRID: AB_2099233
Horse anti-mouse HRP-linked antibody	Cell Signaling	Cat#7076; RRID: AB_330924
Goat anti-Rabbit IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa Fluor 488	Life Technologies	Cat# A-11008; RRID: AB_143165

(Continued on next page)

Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Chemicals, Peptides, and Recombinant Proteins		
Osimertinib (<i>in vivo</i>)	MedChemExpress	Cat#HY-15772A
Osimertinib (<i>in vitro</i>)	Selleck Chem	Cat#S7297
Trametinib	Selleck Chem	Cat#S2673
XAV939	Selleck Chem	Cat#S1180
Selumetinib	Selleck Chem	Cat#1008
MYF-01-37	Synthesized at DFCI	N/A
Reversible control of MYF-01-37	Synthesized at DFCI	N/A
TED-347	Synthesized at DFCI	N/A
Reversible control of TED-347	Synthesized at DFCI	N/A
ZSTK474	Selleck Chem	Cat#S1072
NVP-BEZ235	Cayman Chemicals	Cat#10656
AZD2014	Selleck Chem	Cat#S2783
Ruxolitinib	Selleck Chem	Cat#S1378
Saracatinib	Cayman Chemicals	Cat#11497
BMS-345541	Cayman Chemicals	Cat#16667
Sotrastaurin	Cayman Chemicals	Cat#16726
Galunisertib	Cayman Chemicals	Cat#15312
Gefitinib	Selleck Chem	Cat#S1025
Crizotinib	Selleck Chem	Cat#S1068
Cisplatin	Selleck Chem	Cat#S1166
DAPT (GSI IX)	Selleck Chem	Cat#S2215
LY451039	Selleck Chem	Cat#S1594
Alt-B® S.n. Cas9 Nuclease V3	Integrated DNA Technologies	Cat#1081059
Critical Commercial Assays		
Critical Commercial Assays Quantitect Reverse Transcription Kit	Qiagen	Cat#205313
Critical Commercial Assays Quantitect Reverse Transcription Kit TaqMan Gene expression MasterMix	Qiagen Thermo Fisher Scientific	Cat#205313 Cat#4369016
Critical Commercial Assays Quantitect Reverse Transcription Kit TaqMan Gene expression MasterMix RNeasy Mini Kit	Qiagen Thermo Fisher Scientific Qiagen	Cat#205313 Cat#4369016 Cat#74104
Critical Commercial Assays Quantitect Reverse Transcription Kit TaqMan Gene expression MasterMix RNeasy Mini Kit Gateway™ LR Clonase™ II Enzyme mix	Qiagen Thermo Fisher Scientific Qiagen Invitrogen	Cat#205313 Cat#4369016 Cat#74104 Cat# 11789020
Critical Commercial Assays Quantitect Reverse Transcription Kit TaqMan Gene expression MasterMix RNeasy Mini Kit Gateway™ LR Clonase™ II Enzyme mix Gateway™ BP Clonase™ II Enzyme mix	Qiagen Thermo Fisher Scientific Qiagen Invitrogen Invitrogen	Cat#205313 Cat#4369016 Cat#74104 Cat# 11789020 Cat# 11791020
Critical Commercial Assays Quantitect Reverse Transcription Kit TaqMan Gene expression MasterMix RNeasy Mini Kit Gateway [™] LR Clonase [™] II Enzyme mix Gateway [™] BP Clonase [™] II Enzyme mix Phusion® High-Fidelity PCR Master Mix with HF Buffer	Qiagen Thermo Fisher Scientific Qiagen Invitrogen Invitrogen New England Biolabs	Cat#205313 Cat#4369016 Cat#74104 Cat# 11789020 Cat# 11791020 Cat# M0531
Critical Commercial Assays Quantitect Reverse Transcription Kit TaqMan Gene expression MasterMix RNeasy Mini Kit Gateway [™] LR Clonase [™] II Enzyme mix Gateway [™] BP Clonase [™] II Enzyme mix Phusion® High-Fidelity PCR Master Mix with HF Buffer Senescence β-Galactosidase Staining Kit	Qiagen Thermo Fisher Scientific Qiagen Invitrogen Invitrogen New England Biolabs Cell Signaling	Cat#205313 Cat#4369016 Cat#74104 Cat# 11789020 Cat# 11791020 Cat# M0531 Cat#9860
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REAGENT or RESOURCE	SOURCE	IDENTIFIER
HCC2279; human EGFR-mutant NSCLC, female	Dr. Adi Gazdar (UT Southwestern,	Fingerprinted; RRID: CVCL_5131
	Dallas, TX)	
H1975; human EGFR-mutant NSCLC, female	ATCC	(CRL-5908); RRID: CVCL_1511
HCC827 GR6; human EGFR-mutant NSCLC, female	Established in the Jänne laboratory	N/A
DFCI243; human EGFR-mutant NSCLC, female	Established in the Jänne laboratory	N/A
H3122; human ALK-rearranged (EML4-ALK) NSCLC, sex unknown	Dr. Bruce Johnson (DFCI, Boston, MA)	Fingerprinted; RRID: CVCL_5160
EBC-1; human MET-amplified NSCLC, male	JCRB	(JCRB0820); RRID: CVCL_2891
293T/17	ATCC	(CRL-11268); RRID: CVCL_1926
Experimental Models: Organisms/Strains		
Mouse: NCr nude mice	Taconic Biosciences	Model# NCRNU-F
Mouse: NSG mice	The Jackson Laboratory	Cat# 005557
Mouse: EGFR ^{L858R/T790M} mice	Zhou et al., 2009	N/A
Oligonucleotides		
CTGF Taqman assay	Thermo Fisher Scientific	Cat# Hs01026927_m1
ANKRD1 Taqman assay	Thermo Fisher Scientific	Cat# Hs00173317_m1
BMF Taqman assay	Thermo Fisher Scientific	Cat# Hs00372937_m1
SNAI1 Taqman assay	Thermo Fisher Scientific	Cat# Hs00195591_m1
SNAI2 Taqman assay	Thermo Fisher Scientific	Cat# Hs00161904_m1
TWIST1 taqman assay	Thermo Fisher Scientific	Cat# Hs00361186_m1
TWIST2 taqman assay	Thermo Fisher Scientific	Cat# Hs02379973_s1
ZEB1 Taqman assay	Thermo Fisher Scientific	Cat# Hs00232783_m1
ZEB2 Taqman assay	Thermo Fisher Scientific	Cat# Hs00207691_m1
ACTB taqman assay	Thermo Fisher Scientific	Cat# Hs01060665_g1
ON-TARGETplus <i>BMF</i> siRNA	Dharmacon	L-004393-00-0005
ON-TARGETplus YAP1 siRNA	Dharmacon	L-012200-00-0005
ON-TARGETplus SNAI2 siRNA	Dharmacon	L-017386-00-0005
ON-TARGETplus Non-Targeting siRNA Pool	Dharmacon	D-001810-10-05
Recombinant DNA		
JP1722-YAP1	This paper	N/A
JP1722-YAP1-S94A	This paper	N/A
JP1722-YAP1-TAdel	This paper	N/A
JP1722-YAP1-PDZdel	This paper	N/A
JP1722-YAP1-WWmut	This paper	N/A
JP1722-YAP1-SH3bm	This paper	N/A
JP1722-YAP1-Y357F	This paper	N/A
TBS-mCherry (YAP/Hippo reporter plasmid)	Mohseni et al., 2014	N/A
pLEX307-TEAD1	This paper	N/A
pLEX307-TEAD1 C359S	This paper	N/A
Software and Algorithms		
GraphPad Prism	GraphPad	https://www.graphpad.com/scientific- software/prism/
ImageJ	ImageJ	https://imagej.net/Fiji
Starcode v1.3	Zorita et al., 2015	https://github.com/gui11aume/starcode/ releases
VIPER snakemake pipeline	Cornwell et al., 2018	https://bitbucket.org/cfce/viper/src/master/
Burrows-Wheeler Aligner	Li and Durbin, 2010	http://bio-bwa.sourceforge.net
MACS2	Zhang et al., 2008b	https://github.com/taoliu/MACS/
Deeptools	Ramírez et al., 2016	https://github.com/deeptools/deepTools

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
HOMER	Heinz et al., 2010	http://homer.ucsd.edu/homer/
CellRanger	10X Genomics	www.10xgenomics.com
R toolkit Seurat v.3.0	Butler et al., 2018	https://satijalab.org/seurat/
R package AUCell	Aibar et al., 2017	https://bioconductor.org/packages/ release/bioc/html/AUCell.html
Qupath	Bankhead et al., 2017	https://qupath.github.io

LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Pasi A. Jänne (Pasi_Janne@dfci.harvard.edu). All unique/stable reagents generated in this study are available from the Lead Contact with a completed Materials Transfer Agreement.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animal Models

Xenograft studies: Female NCr nude mice, 7-weeks old (for PC-9 studies) and female NSG mice, 6-weeks old (HCC4006 and DFCI243) were purchased from Taconic Biosciences, Inc. and The Jackson Laboratory, respectively. Animals were allowed to acclimate for at least 5 days before initiation of the study. All in vivo studies were conducted at Dana-Farber Cancer Institute with the approval of the Institutional Animal Care and Use Committee in an AAALAC accredited vivarium. The cells were harvested, and 5 x 10⁶ cells with 50% Matrigel (Fisher Scientific) were implanted subcutaneously in the right flank of the NCr nude or NSG mice. For efficacy studies, tumors were allowed to establish to 200 ± 50 mm³ in size before randomization into various treatment groups with 8 mice per group. Osimertinib (10 mg/kg once daily) and trametinib (1 mg/kg once daily) were administered orally as a suspension using 0.5% hydroxypropyl methylcellulose (HPMC) or 0.5% HPMC with 0.2% Tween 80 as vehicle, respectively. Control vehicle treated mice received 0.5% HPMC with 0.2% Tween 80 administered orally. Tumor volumes were determined from caliper measurements by using the formula V = (length x width²)/2. Tumor sizes and body weight were measured twice weekly. Mice were treated for 28 days, followed by measuring for re-growth of tumors. For the single-cell RNA-sequencing and immunohistochemical analysis of in vivo MRD tumors, PC-9 cells were implanted as above. When the tumors reached an average of $200 \pm 50 \text{ mm}^3$, the mice were randomly assigned to receive either vehicle, 10 mg/kg osimertinib, or 10 mg/kg osimertinib and 1 mg/kg trametinib (3 mice / group). The mice were treated orally once daily for 21 days. After treatment, the tumors were harvested and kept on ice in RPMI-1640 (Gibco), 10% FBS, and 1% penicillin/streptomycin (Gibco) until processing for single-cell RNA-sequencing, or formalin fixed for IHC. For the analysis of BMF expression in vivo, 6 mice/cell line were implanted as above. When the tumors reached an average of 350 ± 50 mm³, the mice were randomly assigned to receive either vehicle or 10 mg/kg osimertinib and 1 mg/kg trametinib (3 mice / group). The mice were treated orally once daily for 3 days, and tumors were harvested 3 hours after the final dose. Tumors were snap-frozen and kept in -80°C until analysis.

Studies using the EGFR^{L858/T790M} mouse model: All breeding, mouse husbandry, and *in vivo* experiments were performed with the approval of Dana-Farber Cancer Institute Animal Care and Use Committee. Tumors in the EGFR^{L858/T790M} mice (Zhou et al., 2009) were induced by 5 x 10⁷ pfu adenovirus expressing Cre Recombinase protein (Cat # VVC-U of Iowa-5, University of Iowa adenoviral core) at 6-8 weeks old and monitored by MRI to quantify lung tumor burden before being assigned to various treatment study cohorts. Mice were treated with either osimertinib or in combination with selumetinib and the lung tumors burden were quantified by MRI imaging before and after the drug treatment. Osimertinib was administered as 5 mg/kg once daily through oral gavage and selumetinib was administered twice daily at 50 mg/kg through oral gavage using 0.5% HPMC as vehicle. For efficacy study, the treatment was continued until 4 weeks then withdrawn. The mice were maintained and monitored by MRI for tumors relapse and humanely euthanized at endpoint. For short term study in order to acquire residual tumors samples, mice were euthanized and samples harvested after treating with osimertinib until MRI imaging showed no visible tumor (2 weeks).

Cell Line Authentication

293T cells and the NSCLC cell lines PC9, HCC827, HCC4006, HCC2279, H1975, H3122, EBC-1 and the patient-derived DFCl243 cell line were grown in RPMI-1640 (Gibco), 10% FBS, and 1% penicillin/streptomycin (Gibco). The HCC827 and HCC2279 cells were obtained from Dr. Adi Gazdar (UT Southwestern, Dallas, TX) in 2004. The PC9 cells were obtained from Dr. Kazuto Nishio (Kindai University, Osaka, Japan) in 2005. The H3122 cells were a kind gift from Dr. Bruce Johnson (Dana-Farber Cancer Institute, Boston, MA) in 2002. HCC4006 (CRL-2871), H1975 (CRL-5908), and 293T/17 (CRL-11268) cells were purchased from ATCC. EBC-1

cells were purchased from Japanese Collection of Research Bioresources (JCRB0820). DFCl243 and HCC827 GR6 (Engelman et al., 2007) cell lines were established in the Jänne laboratory. Cell line identity was confirmed by fingerprinting for the following cell lines: HCC4006, PC9, HCC827, HCC2279, and H3122. EBC-1, H1975 and 293T cells were purchased in 2015, 2016, 2017 respectively, and were not fingerprinted.

Patient Specimen

The patient included in the study provided written informed consent for the use of the specimen and the studies were performed in accordance with the Declaration of Helsinki, and approved by the Dana-Farber Cancer Institute Institutional Review Board. The patient, a 52-year old male, was diagnosed with stage IV EGFR exon 19 deletion NSCLC from a thoracentesis. He was enrolled in clinical trial (NCT03392246) combining osimertinib and selumetinib as his first systemic treatment for advanced *EGFR*-mutant NSCLC. Eleven months later, following a sustained partial response, the patient underwent surgical resection (left upper lobe wedge resection (the sample analyzed in this study)).

METHOD DETAILS

All *in vitro* experiments were performed in sub-confluent cell cultures to avoid artefactual observations due to YAP regulation by cell-cell contacts.

Expression Vectors

All YAP1 constructs used in this study harbor the cDNA encoding 488 amino acid YAP1 isoform (Sudol, 2012). Wild-type YAP1 and YAP1-WWmut cDNAs were amplified (Phusion® High-Fidelity PCR Master Mix with HF Buffer, New England Biolabs) from p2xFlagCMV2-YAP2 and p2xFlagCMV2-YAP2-1st&2nd WW mutant plasmids (gifts from Marius Sudol, Addgene plasmids #19045 and #19048, respectively) and subcloned into pDNR-dual (BD Biosciences) using Sall and Xbal restriction sites. pDNRdual-YAP1-S94A and pDNR-dual-YAP1SH3bm were created by amplifying the mutation sites from pLX304-YAP1(S94A) and pLX304-YAP1 SH3bm plasmids (gifts from William Hahn, Addgene plasmids #59145 and #59141, respectively) using primers 5'-AT-CAACGGGACTTTCCAAAATGTCG-3' and 5'-TTTTTTCTAGACTATAACCATGTAAGAAAGCTTTCTTTA-3' and subcloning the amplified regions into pDNR-Dual-YAP1 using BamHI and Xbal restriction sites. Because the pLX304-YAP1(S94A) and pLX304-YAP1_SH3bm contain the YAP1-504 isoform, a 48 base pair region from pDNR-Dual-YAP1S94A and pDNR-Dual-YAP1SH3bm was subsequently deleted to create the YAP1-488 isoforms. The deletions were done by PCR using primers 5'-GAGT TAGCCCTGCGTAGCCA-3' and 5'-CTGCCGAAGCAGTTCTTGCT-3' followed by re-ligation of the PCR product. The PDZdeletion mutant of YAP1 was created by PCR from p2xFlagCMV2-YAP2 using primers 5'- TTTTTGTCGACCAGAATTGATCTAC CATGGACT-3' and 5'-TTTTTTTCTAGACTAGCTTTCTTTATCTAGCTTGGTG-3' and subcloning the PCR product into pDNR-dual using Sall and Xbal restriction sites. The YAP1-TAdel cDNA was amplified from pLX304-YAP1_TA (gift from William Hahn, Addgene plasmid #59143) using primers 5'- ATCAACGGGACTTTCCAAAATGTCG-3' and 5'- TTTTTTCTAGACTATAACCATGTAAGAAA GCTTTCTGGGCT-3' and subcloned into pDNR-dual-YAP1 using BamHI and Xbal restriction sites. All YAP1 cDNAs were subsequently shuttled into JP1722 expression vector using the BD Creator System (BD Biosciences).

The TEAD1 C359C mutation was generated into pRK5-myc-*TEAD1* backbone (a gift from Kunliang Guan, Addgene plasmid #33109) by PCR using primers 5'-TCCCCAATGAGTGAATATATGATCAAC-3' and 5'-GCGGTTTATTCGGTATACAAATCG-3'. Both wild-type *myc-TEAD1* and the *myc-TEAD1* C359S mutant cDNAs were amplified from the pRK5-backbone using primers 5'-GGGGACAAGTTTGTACAAAAAGCAGGCTTCGCCACCATGGAGCAAAAGCTCATCTCAG-3' and 5'-GGGGACCACTTTGTACAAAAGCAAGCTCGCACCATGGAGCAAAAGCTCATCTCAG-3' and 5'-GGGGACCACTTTGTACAAAGCAAGCTCGTCAAATATG-3' and shuttled into the pLEX307 lentiviral vector (a gift from David Root, Addgene plasmid #41392) using the Gateway cloning technology (Invitrogen).

The TBS-mCherry vector has been described previously (Mohseni et al., 2014).

Cell Growth and Viability Assays

For Figures 1A and 1C, 350 cells/well were plated into 96-well plates and treated as indicated in the figures (n=60 wells / condition). Medium with fresh drugs was changed every 3-5 days. The confluency of the wells was determined weekly using the Incucyte FLR live cell analysis system (Essen Bioscience). For Figure S1B, cells were plated and treated as above, and the wells were manually scored as positive when the confluence was above 50 % and assessed weekly (Tricker et al., 2015). For Figure S1C, 78 000 PC-9 cells were plated into T25 flasks and treated the next day as indicated. Cell proliferation was monitored using Incucyte HD live cell analysis system (Essen Bioscience) by imaging 32 sectors in the T25 flask. For all other long-term growth assays (\geq 10 days), 1000 cells/well were plated into 96-well plates and treated as indicated in the figures (n=5-12 wells / condition). The confluency of the wells was determined daily using Incucyte HD. Endpoint cell viability assays were performed using Cell Titer Glo (Promega) according to manufacturer's instructions.

To determine number of dormant cells after treatment, viable cells were manually counted from the Incucyte images. A total of 10-12 wells with 3 images per well was analyzed for each condition.

Western Blotting and Antibodies

If not specified below, cells were plated at 15×10^4 cells / cm², treated the next day (if applicable) and lysed at specified timepoints in RIPA buffer (Boston Bioproducts) supplemented with cOmplete Mini EDTA-free Protease inhibitor cocktail (Roche) and PhoSTOP phosphatase inhibitor cocktail (Roche). Twenty micrograms of total protein was used for immunoblotting according to the antibody manufacturer's recommendations. For the assessment of protein levels in dormant cells, 13×10^4 cells / cm² were plated into 2×15 cm dishes. Cells were treated the next day and medium with fresh drugs was changed every 3-5 days. Cells were trypsinized at specified timepoints, washed with ice-cold PBS, and the cell pellets were lysed and immunoblotted as above.

Cellular Barcoding

PC-9 cells were transduced with the EvoSeq barcode library (Feldman et al., 2019) and bottlenecked to a complexity of approximately 500,000 barcodes. The barcoded cells were plated into five replicates per treatment, 5x10⁶ cells per replicate. The cells were then treated with 300 nM gefitinib, 100 nM osimertinib, or 100 nM osimertinib + 30nM trametinib for 3 weeks to establish the residual cell populations. After treatment, the cells were harvested and the genomic DNA was extracted, the barcode-containing sequences were amplified from the genomic DNA, and prepared for sequencing as described (Feldman et al., 2019). Each library was quantified by Qubit fluorometer, Agilent TapeStation 2200, and RT-qPCR using the Roche Kapa Biosystems library quantification kit according to manufacturer's protocols. Uniquely indexed libraries were multiplexed in equimolar ratios into two pools — one pool of twelve libraries and the other of thirteen libraries — and sequenced on two Illumina NextSeq500 runs with paired-end 75bp reads by the Dana-Farber Cancer Institute Molecular Biology Core Facilities. Quantification of the barcode abundance and barcode clustering were performed within each replicate using the software package Starcode (v1.3) with default parameters (Zorita et al., 2015). Only clusters with abundance 2 or larger were retained for further analysis. The 500 most abundant barcodes in each sample were used for the downstream analyses (Bhang et al., 2015).

RNA Extraction and Quantitative PCR (QPCR)

Cells were plated at 15×10^4 cells / cm², treated the next day, and RNA samples were extracted at specified timepoints using the RNeasy Mini kit (Qiagen). The RNA concentrations were measured with Nanodrop (Thermo Fisher Scientific) and 1 µg of total RNA was used for cDNA synthesis using the QuantiTect Reverse Transcription Kit (Qiagen). The QPCR reactions were set up in 20 µl using Taqman Gene Expression Master Mix (Thermo Fisher Scientific, cat. 4369016), Taqman Gene Expression Assays (Thermo Fisher Scientific) as per manufacturer's instructions, and 2 µl of 1:10 diluted cDNA. The following Taqman Gene Expression Assays were used in the study: *CTGF* (Hs01026927_m1), *ANKDR1* (Hs00173317_m1), *BMF* (Hs00372937_m1), *SNAI1* (Hs00195591_m1), *SNAI2* (Hs00161904_m1), *TWIST1* (Hs00361186_m1), *TWIST2* (Hs02379973_s1), *ZEB1* (Hs00232783_m1), *ZEB2* (Hs00207691_m1), and *ACTB* (Hs01060665_g1). The reactions were run in StepOne Plus Real-time PCR System (Applied Biosystems) using default reaction settings. The gene expression levels were normalized to *ACTB* housekeeping gene expression levels in each sample.

For the analysis of *BMF* expression *in vivo*, RNA was extracted from 25-30 mg of snap-frozen tumor samples using RNeasy Mini kit according to the kit protocol. Reverse transcription and gene expression analyses were performed as above.

RNA-sequencing

To analyze gene expression changes associated with dormancy, PC-9, HCC827 and HCC4006 cells were plated at 15×10^4 cells / cm² into 10 cm plates (DMSO treated control cells) or into 15 cm plates (dormant cells). The next day, cells were treated either with DMSO or with the combination of 100 nM osimertinib and 30 nM trametinib in duplicate. DMSO-treated control cells were harvested 24h later, and the dormant cells after 2 weeks of treatment. At these timepoints, cells were lysed into TRIzol and RNA extraction was performed according to the manufacturer's protocol.

In order to analyze YAP1 KO –associated gene expression changes, PC-9 and HCC4006 CTRL and YAP1 KO cells were plated into 10 cm dishes at 15 x 10⁴ cells / cm². The next day, the cells were treated with DMSO or with the combination of 100 nM osimertinib and 30 nM trametinib in triplicate. After 24 hours, the cells were lysed into TRIzol and RNA extraction was performed according to the manufacturer's protocol.

Libraries were prepared using Illumina TruSeq Stranded mRNA sample preparation kits from 500 ng of purified total RNA according to the manufacturer's protocol. The finished dsDNA libraries were quantified by Qubit fluorometer, Agilent TapeStation 2200, and RT-qPCR using the Kapa Biosystems library quantification kit according to manufacturer's protocols. Uniquely indexed libraries were pooled in equimolar ratios and sequenced on an Illumina NextSeq500 with single-end 75bp reads by the Dana-Farber Cancer Institute Molecular Biology Core Facilities. Sequenced reads were aligned to the UCSC hg19 reference genome assembly and gene counts were quantified using STAR (v2.5.1b). Differential gene expression testing was performed by DESeq2 (v1.10.1) and normalized read counts (FPKM) were calculated using cufflinks (v2.2.1). RNAseq analysis was performed using the VIPER snakemake pipe-line (Cornwell et al., 2018).

Gene set enrichment analyses from the RNA-seq data were performed according to the instructions (http://www.broadinstitute. org/gsea/index.jsp).

Senescence-Associated β-galactosidase Staining

PC-9, HCC827 and HCC4006 were plated into 6-well plates at 50 000 cells / well, and treated the next day with DMSO, 100 nM osimertinib or with the combination of 100 nM osimertinib and 30 nM trametinib in triplicate. DMSO-treated control cells were stained after 72h, and osimertinib and osimertinib/trametinib -treated cells were stained after 10-day treatment using Senescence β -Galactosidase Staining Kit (Cell signaling #9860) according to manufacturer's protocol. After staining, cells were imaged (5 images / well), and stained cells were manually counted from the images.

Cytokine Profiling

Multiplex assay was performed using the Human Cytokine/Chemokine Magnetic Bead Panel (Millipore cat# HCYTMAG-60K-PX30) on a Luminex MAGPIX system (Millipore). Conditioned media concentration levels of each protein were derived from 5-parameter curve fitting models. Protein levels were normalized to cell number in each well.

Immunofluorescence Staining and Imaging

Cells grown on coverslips were washed with PBS and fixed with 4 % PFA for 10 minutes. The cells were then permeabilized with 0.1 % Triton-X-PBS, followed by a blocking step in 1 % BSA-PBS. The cells were incubated for 60 minutes with Anti-Histone H3 (tri methyl K9) antibody (Abcam ab8898, 1:300) (Figure 1K) or with anti-YAP (Cell Signaling #14074, 1:200) (Figure 3B), washed 3 times with PBS, incubated with Alexa Fluor 488® -conjugated secondary antibody (A-11008, Life Technologies, 1:300) for 45 minutes, and washed 3 times with PBS. The nuclei were counterstained with 1 µg/ml DAPI (Cell Signaling #4083). The coverslips were mounted using Immu-Mount reagent (Thermo Scientific). Images from H3K9Me³-stainings were acquired using Nikon eclipse 80i microscope. Image analysis was performed using ImageJ software. For H3K9Me³, images were segmented using standard thresholding parameters and objects were automatically counted using ImageJ Analyze particles- plugin. For the analysis of YAP nuclear localization, the Intensity Ratio Nuclei Cytoplasm Tool –plugin for ImageJ was used.

ATAC-sequencing

PC-9 cells were plated at 15 x 10⁴ cells / cm² into 15 cm plates, and treated the next day with DMSO, 100 nM osimertinib or with the combination of 100 nM osimertinib and 30 nM trametinib in triplicate. DMSO-treated control cells were harvested 24h later. Osimertinib and osimertinib/trametinib -treated cells were harvested after 2 weeks of treatment. Rebound samples were obtained by withdrawing drugs from 3 additional osimertinib/trametinib -treated plates and harvesting the cells once the plates reached 60-70% confluence. Cells were trypsinized at timepoints, and cryopreserved in FBS + 8% DMSO in -80C until processing. After all samples were harvested, 50000 cells / sample were resuspended in 1 ml of cold ATAC-seq resuspension buffer (RSB; 10 mM Tris-HCl pH 7.4, 10 mM NaCl, and 3 mM MgCl2 in water). Cells were centrifuged at max speed for 5 min in a pre-chilled (4 C) fixed-angle centrifuge. After centrifugation supernatant was carefully aspirated. Cell pellets were then resuspended in 50 µl of ATAC-seg RSB containing 0.1% NP40, 0.1% Tween-20, and 0.01% digitonin by pipetting up and down 3 times. This cell lysis reaction was incubated on ice for 3 min. After lysis, 1 ml of ATAC-seq RSB containing 0.1% Tween-20 (without NP40 or digitonin) was added, and the tubes were inverted to mix. Nuclei were then centrifuged for 5 min at max speed in a pre-chilled (4 C) fixed-angle centrifuge. Supernatant was removed and nuclei were resuspended in 50 µl of transposition mix (Corces et al., 2017): 2.5 µl transposase (100 nM final), 16.5 µl PBS, 0.5 µl 1% digitonin, 0.5 µl 10% Tween-20, and 5 µl water) by pipetting up and down six times. Transposition reactions were incubated at 37 C for 30 min in a thermomixer with shaking at 1,000 r.p.m. Reactions were cleaned up with Qiagen columns. Libraries were amplified as described previously (Buenrostro et al., 2015). 36-bp paired-end reads were sequenced on a Nextseq instrument (Illumina).

ChIP-sequencing

PC-9 cells were plated at 15×10^4 cells / cm² into 15 cm plates, and treated the next day either with DMSO or with the combination of 100 nM osimertinib and 30nM trametinib in duplicate. Cells were trypsinized at after 48h of treamtent, and cryopreserved in FBS + 8% DMSO in -80C until processing. Cells were washed in phosphate-buffered saline (PBS) and crosslinked with 1% Formaldehyde for 10 minutes (H3K27Ac) or crosslinked with two agents starting with 2 mM DSG (Pierce) for 45 min at RT, followed by 1 ml 1% Formaldehyde for 10 min (YAP and TEAD4). Crosslinked Cell lines were quenched with 0.125 M glycine for 5 min at room temperature. After quenching, the material was resuspended in 1% SDS (50 mM Tris-HCl pH8, 10 mM EDTA) and sonicated for 5 minutes with a Covaris E220 instrument, 5% duty cycle, 140 Peak Incident Power, 200 Cycles per burst, in 1ml AFA Fiber milliTUBEs. Soluble chromatin was immunoprecipitated with 10 µg of H3K27ac antibody (Diagenode cat# C15410196), 7 µg of YAP antibody (Cell signaling #9585), or 1.5 µg of TEAD antibody (ab58310, Abcam). 5 µg of chromatin was used for H3K27Ac ChIP, and 40 µg for YAP, TEAD4 and SLUG ChIPs. ChIP-seq libraries were constructed using Accel-NGS 2S DNA library kit from Swift Biosciences. Fragments of the desired size were enriched using AMPure XP beads (Beckman Coulter). 36-bp paired-end reads were sequenced on a Nextseq instrument (Illumina).

ATAC-seq and ChIP-seq Analyses

The raw data from ATAC-seq and ChIP-seq was first ran through the ChiLin 2.0.0. pipeline (Qin et al., 2016) for initial quality control and preprocessing. Reads were mapped to human genome (hg19) using Burrows-Wheeler Aligner (Li and Durbin, 2010) and peak calling was performed using MACS2 (Zhang et al., 2008b). The output bedgraph files from MACS2 were converted to bigwig files

for visualization in the Integrative Genomics Viewer. Deeptools (Ramírez et al., 2016) was used to create heatmap plots. The PCA plot was generated by using the R package 'ggbiplot'. HOMER (Heinz et al., 2010) was used for the motif analysis.

CRISPR/CAS9 Gene Editing

YAP1 knock-outs were performed by CRISPR/CAS9 genome editing using the Alt-R CRISPR-CAS9 system (Integrated DNA Technologies, IDT) and Lonza 4D-Nucleofector (Lonza), following previously described protocol (Richardson et al., 2016). Guide sequences for YAP1 were designed using Deskgen (deskgen.com), and the corresponding Alt-R CRISPR-Cas9 crRNAs (crRNA) were ordered from IDT. The crRNA was hybridized with Alt-R CRISPR-Cas9 tracrRNA (IDT) by mixing 120 pmol of crRNA with 120 pmol of tracrRNA in 5 µl of CAS9 buffer (20 mM HEPES (pH 7.5), 150 mM KCl, 1 mM MgCl2, 10% glycerol and 1 mM TCEP), incubating the mixture at 95°C for 5 minutes and then letting the mixture cool to room temperature on benchtop (5-10 minutes). 100 pmol of Alt-R® S.p. Cas9 Nuclease V3 (IDT) in 5 µl of CAS9 buffer was slowly added to the crRNA:tracrRNA duplex and the subsequent solution was incubated for 20 minutes in room temperature to allow ribonucleoprotein complex (RNP) formation. The RNP complex was then added to 20 µl of cell suspension containing 300 000 cells suspended in Nucleofector SE Cell line solution (Lonza, cat. V4XC-1032), mixed and 20 μl of the cell/RNP mix was pipetted into one well of a Nucleocuvette Strip (Lonza, cat. V4XC-1032). The reaction mixtures were nucleofected using cell line -specific programs (see below) in the 4D-Nucleofector, and finally transferred to 6-well plates. After 72 hours, the nucleofected cells were single-cell cloned, and loss of YAP protein expression was analyzed from the single-cell clones by western blotting. The guide sequence 5'- TAATAGGCCAGTACTGATGC-3' was used to create PC-9, HCC4006, and DFCI243 YAP1 KOs. H3122 and EBC-1 YAP1 KOs were created using two guides with sequences 5'- TAATAGGC CAGTACTGATGC-3' and 5'-GAATGAGCTCGAACATGCTG-3' simultaneously to ensure high knock-out efficiency. The nucleofected H3122 and EBC-1 cells were not single-cell cloned, and bulk populations were used in the experiments. The nucleofection conditions were optimized using the Cell Line Optimization 4D-Nucleofector X Kit (Lonza, cat. V4XC-9064) following the kit protocol. The optimized programs used were: EN-138 for PC-9 and EBC-1 cells; CA-137 for H3122 cells; CM-137 for HCC4006 and DFCI243. All cell lines were nucleofected in SE Cell line solution.

Monitoring Caspase-3/7 Activity

Cells were plated into 96-well plates at 3000 cells/well in 100 µl of growth medium. The next day, drugs were added onto cells in 50 µl containing CellEvent[™] Caspase-3/7 Green ReadyProbes[™] Reagent (Molecular Probes) as per manufacturer's instructions (n=5-6 wells / condition). The wells were subsequently scanned every 2 hours using the Incucyte ZOOM live cell analysis system (Essen Bioscience) typically for a total of 72 hours. The acquired fluorescent signal for activated caspase-3/7 was normalized with well confluency at each timepoint (=normalized apoptosis). Peak apoptosis was determined as the highest normalized caspase-3/7 activity value during the assay.

Determining YAP Activity and Apoptosis in PC-9 YAP/Hippo Reporter Cells

3000 cells / well were plated into 96-well plates and treated the next day with the indicated drugs (n=5-6 wells / condition). YAP activity -induced mCherry expression was quantified using the Incucyte ZOOM live-cell analysis system. The mCherry signal was normalized to well confluency at each time point. For simultaneous detection of YAP activity and apoptosis, the cells were plated as above, and treated in the presence of CellEvent[™] Caspase-3/7 Green ReadyProbes[™] Reagent (Molecular Probes) as per manufacturer's instructions. The mCherry signal as well as the green fluorescence signal was quantified every 2 hours using Incucyte ZOOM.

To determine the odds ratio for YAP^{high} cells undergoing apoptosis in response to osimertinib/trametinib treatment, the number of YAP^{high} cells (cells with higher mCherry signal than untreated cells), apoptotic cells (positive for green fluorescence) and apoptotic YAP^{high} cells (YAP^{high} cells positive for green fluorescence) was determined using the Incucyte ZOOM software. The analyses were done at a single timepoint corresponding to the peak in apoptosis in response osimertinib/trametinib treatment (72-80 hours after the start of treatment, depending on the experiment), and 5-6 wells with 3 images / well were analyzed. From the same images, the total number of cells per image was manually determined. Using these metrics, a contingency table was built for the average number of YAP^{high} caspase-3/7 positive, YAP^{high} caspase-3/7 negative, YAP^{low}, caspase-3/7 positive, and YAP^{low}, caspase-3/7 negative cells. The odds ratio was computed in GraphPad Prism 7.04 software, and two-sided Fisher's exact test was used to analyze statistical significance.

The proportion of YAP^{high} dormant cells (Figure 3I) was determined manually from the Incucyte images after 10-day treatment. 5-6 wells with 3 images per well were analyzed.

Viral Transductions

For stable expression of YAP1 or YAP1 mutants, PC-9 YAP1 KO and HCC4006 YAP1 KO cells were transduced with lentivirus according to previously described standard protocol (Bahcall et al., 2016). Transduced cells were selected with 2 µg/ml puromycin. PC-9 YAP/Hippo reporter cells were created by lentiviral transduction of TBS-mCherry YAP/Hippo reporter construct (Mohseni et al., 2014). The subsequent cell pool was flow sorted for EGFP expression to select transduced cells.

Single-cell RNA Sequencing

For *in vitro* samples: 1.5 x 10⁶ PC-9 cells were plated onto T75 flasks. Cells were treated either with DMSO or with 100 nM osimertinib and 30 nM trametinib for 3 weeks. After treatment, the cells were washed with PBS, trypsinized, and loaded onto a 10X Chromium instrument (10X Genomics) per the manufacturer's instructions. For *in vivo* samples: Fresh tumor specimens were pooled and minced in a 15 ml conical tube with media (DMEM + 10% FBS), penicillin–streptomycin (Fisher Scientific), 100 U/mL collagenase type IV (Life Technologies) and 2.5 mg/mL DNAse I (Sigma Aldrich), then incubated for 45 min at 37°C. Single cell suspensions were isolated by straining through a 40 μm filters. Cells were incubated with Zombie Green[™] Fixable Viability Kit (BioLegend), blocked with Human TruStain FcX[™] (BioLegend), and stained with human anti-EpCAM (clone 9C4). Viable EpCAM+ tumor cells were isolated via FACS Melody instrument (BD Biosciences) according to gating schema (Figure S6A). Cells were loaded onto a 10X Chromium instrument (10X Genomics) per the manufacturer's instructions.

Single-cell RNA libraries were generated using the Single Cell 3' Reagent Kit (10X Genomics) per user guide. Quality control of the completed libraries was performed using Bioanalyzer High Sensitivity DNA Kit (Agilent) and then sequenced using the Illumina Next-Seq 500 platform by Novogene. The single-cell RNA-Seq data were processed with CellRanger software package (v.3.0.2). Briefly, the bcl files were converted to fastq files, which were aligned to human transcriptome (build GRCh38). After initial filtering with default parameters, the feature matrix generated by Cell Ranger was used to perform downstream analysis using R toolkit Seurat (v.3.0) (Butler et al., 2018). At this step, the cells with mitochondria percentages greater than 20 or expressing less than 200 genes were filtered out.

To characterize cell subpopulations in the samples, we performed the gene signature enrichment analysis for "YAP signature", "HALLMARK EMT signature" (HALLMARK_EPITHELIAL_MESENCHYMAL_TRANSITI

NSITION, MSigDB, http://software.broadinstitute.org/gsea/msigdb/index.jsp), and "FRIDMAN SENESCENCE UP" (MSigDB) signatures. The YAP signature was curated from gene sets obtained from multiple studies (Cordenonsi et al., 2011; Dupont et al., 2011; Wang et al., 2018; Zhang et al., 2009, 2008a). The YAP signature was filtered to include only those genes that were associated with strong YAP binding upon osimertinib/trametinib treatment in PC-9 cells based on the ChIP-Seq data (peaks with ChIP/input enrichment fold-change greater than 10) (Table S1). The enrichment scores for a given gene signature for each cell in a sample were calculated using R package AUCell (Aibar et al., 2017).

Immunohistochemistry

Staining of tumors from EGFR^{L858/T790M} mice (Figures 4E and S6B): Five-micron paraffin sections were stained on a Leica BondRX® autostainer, according to the manufacturers' instructions, with primary antibodies against F480 (Cell Signaling, cat # 70076S, 1:500), CD4 (Cell Signaling; cat # 25229; 1:100), CD8a (Cell Signaling; cat # 98941S; 1:400), YAP (Cell Signaling; Cat # 14074, 1:200) and TTF1 (Abcam, AB133638, 1:50). Prior to antibody incubation, the sections were heat-retrieved with ER1 buffer (pH 6; Leica AR9961) for 20 minutes (YAP), ER2 (pH 9; Leica AR9640) for 20 minutes (CD4, CD8a, F480) or ER2 for 60 mins (TTF1) (Leica, AR9640) at 100° and then treated for 5 minutes with hydrogen peroxide. Sections were incubated with primary antibodies for 30 minutes (CD4, CD8a, F480, YAP) or 60 minutes (TTF1), followed by Leica anti-rabbit HRP-conjugated polymer, and then developed with DAB, counterstained with hematoxylin (Leica DAB KIT, Cat # DS9800) and mounted with permount.

Staining of the PC-9 xenograft tumors, WZ4002 and WZ4002/trametinib –resistant EGFR^{L858/T790M} mice tumors, and the patient sample (Figures 4D, 4H, and 4I): All IHC was performed on the Leica Bond III and RX automated staining platforms. Phospho-ERK (Cell Signaling # 4370) and YAP (Cell Signaling #14074) antibodies were run at 1:150 and 1:200 dilutions, respectively using the Leica Biosystems Refine Detection Kit with citrate antigen retrieval, and chromogenically stained with DAB (Leica Biosystems Refine Detection Kit).

All IHC stainings were quantified using the QuPath software (0.2.0-m4) (Bankhead et al., 2017). The Positive Cell Detection –analysis was used with default settings to detect and quantify cells staining positive for pERK, CD4, CD8, TTF1, or the nuclear staining of YAP. Five individual, randomly selected fields per tumor were quantified. Quantified values from each individual field from all tumors are shown in the graphs to represent the heterogeneity in the tumor samples.

Detection of Activated BAX

For the analysis of BAX activation, cells $(15 \times 10^4 \text{ cells / cm}^2)$ growing on 10 cm plates were treated for 24 hours with DMSO or with the combination of 100 nM osimertinib and 30 nM trametinib, and lysed in CHAPS lysis buffer (50 mM Tris-HCl, 1 % CHAPS, 150 mM NaCl, 5 mM EDTA) supplemented with protease and phosphatase inhibitors. 1000 µg of total protein was used for immunoprecipitation with 1 µg of conformation-specific BAX antibody (clone 6A7; MA5-14003, Invitrogen) and 20 µl of Protein A/G PLUS-Agarose beads (Santa Cruz Biotechnology). The lysates were incubated with the antibody and beads overnight at +4°C, after which the beads were washed four times with 1000 µl of CHAPS buffer, resuspended to 50 µl SDS Sample Buffer (Boston Bioproducts), and incubated

at +95°C for 5 minutes. Activated BAX was detected by immunoblotting using an antibody detecting total BAX (Cell Signaling #5023). For control, total BAX levels were also determined from total cell lysates.

Detection of Mitochondrial Cytochrome c Release

Cells (15 x 10⁴ cells / cm²) growing on 15 cm plates were treated with the combination of 100 nM osimertinib and 30 nM trametinib for 24 hours, and fractionated into mitochondrial and cytosolic fractions using Cell Fractionation Kit – Standard (ab109719, Abcam) according to the manufacturer's instructions. Cytochrome c and ATP synthase subunit alpha (mitochondrial marker) was detected from the fractions by immunoblotting using ApoTrack[™] Cytochrome c Apoptosis WB Antibody Cocktail (ab110415, Abcam). MEK 1/2 was used as the cytosolic marker (Cell Signaling #9122).

Gene Knock-down by siRNA

Cells were plated on 6-well plates at 15×10^4 cells / cm². The next day, the cells were transfected with 10 nM siRNAs using DharmaFECT 1 (Dharmacon) according to manufacturer's protocol. Forty-eight hours later, the cells were trypsinized and plated into experiments. The following Dharmacon SMARTpool ON-TARGETplus siRNA pools were used in the analyses: *BMF* (L-004393-00-0005), *SNAI2* (L-017386-00-0005), *YAP1* (L-012200-00-0005), and ON-TARGETplus Non-Targeting Pool (D-001810-10-05). Gene knock-down was controlled by western blotting or QPCR 72h after transfection.

Co-immunoprecipitation

PC-9 and HCC4006 cells treated for 48h with either DMSO or with the combination of 100 nM osimertinib and 30 nM trametinib were lysed in IP buffer (1% Triton-X100, 50 mM Tris, 150 mM NaCl, pH 7.4) supplemented with cOmplete Mini, EDTA-free Protease inhibitor cocktail (Roche) and PhoSTOP phosphatase inhibitor cocktail (Roche). 1500 µg of total protein was used for immunoprecipitation with Cell Signaling antibodies recognizing endogenous YAP (#14074), TEAD (pan-TEAD) (#13295), or SLUG (#9585) and 20 µl of Protein A/G PLUS-Agarose beads (Santa Cruz Biotechnology; sc-2003). Immunoprecipitations were carried out overnight in +4C, followed by four washes with 1 ml IP buffer. After washes, the beads were re-suspended in SDS Sample Buffer (Boston Bioproducts), and boiled for 5 min. Co-immunoprecipitated proteins were analyzed by western blotting.

Chemistry

Unless otherwise noted, reagents and solvents were obtained from commercial suppliers and were used without further purification. 1H NMR spectra were recorded on 500 MHz (Varian AS600), and chemical shifts are reported in parts per million (ppm, δ) downfield from tetramethylsilane (TMS). Coupling constants (J) are reported in Hz. Spin multiplicities are described as s (singlet), br (broad singlet), d (doublet), t (triplet), q (quartet), and m (multiplet). Mass spectra were obtained on a Waters Micromass ZQ instrument. Preparative HPLC was performed on a Waters Sunfire C18 column (19 mm × 50 mm, 5 μ M) using a gradient of 15–95% methanol in water containing 0.05% trifluoroacetic acid (TFA) over 22 min (28 min run time) at a flow rate of 20 mL/min. Purities of assayed compounds were in all cases greater than 95%, as determined by reverse-phase HPLC analysis (Scheme 1).



Scheme 1. Synthesis of MYF-1-37

Reagents and conditions: (a) Pd(OAc)2, XPhos, NaOtBu, toluene, 100°C; (b) HCl/dioxane, MeOH; (c) DIEA, MeCN, 0°C.

Tert-butyl 3-methyl-3-((3-(trifluoromethyl)phenyl)amino)pyrrolidine-1-carboxylate (2)

To a solution of 1-bromo-3-(trifluoromethyl)benzene (223 mg, 1.0 mmol) and tert-butyl 3-amino-3-methylpyrrolidine-1-carboxylate (200 mg, 1.0 mmol) in 5 mL of toluene was added Pd(OAc)₂ (22 mg, 0.1 mmol), XPhos (58 mg, 0.1 mmol) and NaOtBu (192 mg, 2 mmol) under N₂. The mixture was stirred at 100°C overnight. The mixture was filtered. The filtrate was concentrated in vacuo, then purified by flash chromatography on silica gel (hexane: ethyl acetate = 4:1) to provide compound **2** (240 mg, 70%). LC/MS (ESI) m/z = 345 (M + H)+.

3-methyl-N-(3-(trifluoromethyl)phenyl)pyrrolidin-3-amine (3)

To a solution of *t*ert-butyl 3-methyl-3-((3-(trifluoromethyl)phenyl)amino)pyrrolidine-1-carboxylate (240 mg, 0.7 mmol) in 3 mL of methanol was added 4N HCl/dioxane (1 mL) solution. The result solution was stirred at room temperature for 1 h, and then concentrated in vacuo to obtain the product as HCl salt, which was used into next step without any purification. LC/MS (ESI) m/z = 245 (M + H)+. **1-(3-methyl-3-((3-(trifluoromethyl)phenyl)amino)pyrrolidin-1-yl)prop-2-en-1-one (MFY-1-37)**

To a solution of 3-methyl-N-(3-(trifluoromethyl)phenyl)pyrrolidin-3-amine (28 mg, 0.1 mmol) and DIEA (33 uL, 0.2 mmol) in 1 mL of acetonitrile was added acryloyl chloride dropwise at 0°C until the reaction completed. The mixture was diluted with dichloromethane, washed with 1 N NaHCO3 solution and brine. The organic layer was dried over sodium sulfate, concentrated in vacuo and then

purified by prep-HPLC (MeOH/H2O, 0-100%) to provide the title compound (23.4 mg, 79%). LC/MS (ESI) m/z = 299 (M + H)+. 1H NMR (500 MHz, DMSO-d6) δ 7.28 (t, J = 8.0 Hz, 1H), 6.98 – 6.87 (m, 2H), 6.83 (d, J = 7.4 Hz, 1H), 6.56 (ddd, J = 18.1, 16.8, 10.3 Hz, 1H), 6.21 (d, J = 4.3 Hz, 1H), 6.12 (ddd, J = 16.8, 6.6, 2.4 Hz, 1H), 5.66 (ddd, J = 10.3, 8.8, 2.4 Hz, 1H), 3.87 – 3.73 (m, 1H), 3.71 – 3.59 (m, 1.5H), 3.53 – 3.43 (m, 1H), 3.39 (d, J = 12.3 Hz, 0.5H), 2.36 – 2.16 (m, 1H), 2.08 – 1.85 (m, 1H), 1.42 (s, 3H).

MYF-01-37 Docking to TEAD2

MYF-01-037-02 was docked into TEAD2 crystal structure (pdbcode: 5HGU) using Glide covalent docking program (version 2019 release 1). The Cys380 was defined as the reactive residue for Michael addition reaction. Default parameter values were used for docking calculation. Prior to docking, the protein structure was processed and energy optimized using protein preparation protocol in Schrodinger suite software.

MYF-01-37 Competition Pulldown

Cells were treated with test compounds for 6 h at the indicated doses. Total cell lysates were prepared using pulldown buffer (50 mM Tris, pH 7.4, 250 mM NaCl, 5 mM EDTA, 50 mM NaF, 1 mM Na3VO4, 1% NP40, 0.02% NaN2, 1 mM PMSF and protease inhibitor cocktail). To pull down TEADs, 1 mg of total protein was combined with biotinylated MYF-01-37 at 1 µM and rotated at 4°C for 6 hrs followed by RT for 1 h. Subsequently, streptavidin agarose resin (30 µl 50% slurry) was added and samples rotated for another 2 h at 4°C. The resin was subsequently washed 3x with pulldown buffer, and TEADs released from the resin by boiling for 10 min in 2x gel loading buffer and resolved by western blotting. As a loading control, 25 µg of total protein was used.

Mass Spectrometry Analysis

TEAD2 protein was incubated with DMSO or a 20-fold molar excess of MYF-01-37 for 6 hours at 37°C. Reactions were then analyzed by LC-MS using a Shimadzu autosampler and LC (Marlborough, MA) coupled to an LTQ ion trap mass spectrometer (ThermoFisher Scientific, San Jose, CA). Protein was injected onto a self packed column (0.5 mm I.D., packed 5 cm POROS 50R2 from Applied Biosystems, Framingham, MA), desalted for 4 minutes with 100% A (A=0.2 M acetic acid in water), eluted with a gradient (0-100% B in 1 minute; A=0.2 M acetic acid in water, B=0.2 M acetic acid in acetonitrile), and introduced to the mass spectrometer by electrospray ionization (spray voltage=4.5 kV). The mass spectrometer acquired full scan MS data (m/z 300-2000). Mass spectra were deconvoluted using MagTran version 1.03b2 (Zhang and Marshall, 1998).

To identify the site of modification, labeled protein was diluted 1:1 with 100 mM ammonium bicarbonate, reduced with 10 mM DTT at 56°C for 30 minutes, alkylated with 22.5 mM IAA for 30 minutes at room temperature, and then digested with trypsin overnight at 37°C. Tryptic peptides were desalted by C18 (SOLA, ThermoFisher Scientific), dried by vacuum centrifugation, reconstituted in 5% MeCN, 0.1% trifluoroacetic acid, and analyzed by nanoLC-ion mobility MS/MS using a NanoAcquity UPLC system (Waters Corp., Milford, MA) interfaced to a timsTOF Pro mass spectrometer (Bruker, Billerica, MA). Peptides were injected onto a self-packed pre-column (4 cm POROS10R2, Applied Biosystems), resolved on an analytical column (30 μ m I.D. x 50 cm Monitor C18, Orochem, Naperville, IL; 10-60% B in 40 minutes; A = 0.2 M acetic acid in water, B = 0.2 M acetic acid in acetonitrile) and introduced to the mass spectrometer by electrospray ionization using a captive spray ion source (spray voltage = 2 kV). The mass spectrometer collected ion mobility MS spectra over a mass range of m/z 100-1700 and 1/k0 of 0.6 to 1.6, and then performed 10 cycles of PASEF MS/MS with a target intensity of 20k and a threshold of 250. Active exclusion was enabled with a release time of 0.4 minutes. Raw data was converted to .mgf using the tdf to mgf converter (Bruker), and searched using Mascot 2.6.1 against a forward reversed human refseq database (NCBI). Search parameters specified a precursor mass tolerance of 20 ppm, a product ion tolerance of 50 mmu, fixed carbamidomethylation of cysteine, and variable oxidation of methionine as well as variable MYF-1-37 modification of cysteine. Search results were downloaded and converted to xls using multiplierz software (Alexander et al., 2017) , and peptide fragment ions were assigned using mzStudio (Ficarro et al., 2017). Inhibitor related fragment ions were assigned as described (Ficarro et al., 2016).

YAP-TEAD Split Gaussia Luciferase (SGL) Assay

The SGL assay monitors real-time cell-based protein-protein interaction between YAP and TEAD, as physical proximity between both proteins would result in luciferase signal reconstitution. 293T cells were plated in 24-well plates and transfected with N-GLuc-YAP and C-GLuc-TEAD using TransIT-293 (Mirus Bio). The pCMV-Red Firefly Luc Vector (Life Technologies) was used as an internal control. Cells were treated with indicated concentrations of compounds or vehicle control (DMSO) in duplicates. Luciferase activity was measured by Dual-Luciferase Assay (Promega, Madison, WI) according to the manufacturer's manual.

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical Analyses

When comparing two groups, statistical significance was calculated by two-tailed unpaired *t*-test with Welch's correction. One-way ANOVA with Dunnett's multiple comparisons test was used when comparing 3 or more groups. Fisher's exact test (two-sided) was used to analyze statistical significance for the odds ratio for YAP^{high} cells undergoing apoptosis in response to osimertinib/trametinib treatment (Figure 3D). Kolmogorov-Smirnov Test was used to assess significance in single-cell RNA-seq analyses (Figure 4C). The statistical tests used are indicated in the figure legends. All statistical analyses were generated using GraphPad Prism 7.04 software, except for Figure 4C where statistical analyses were run in R. Data are presented as mean ± standard deviation or standard error, as indicated in the figure legends. A p-value that was less than 0.05 was considered significant in all analyses. Any cutoffs used for the p-value or FDR are indicated in the figure legends. All analyses were performed with 3-12 replicates as indicated in the Method Details. In animal experiments n represents the number of animals in treatment groups.

DATA AND CODE AVAILABILITY

The datasets generated during this study are available at GEO under accession number GEO: GSE131604.