Intermittent treatment of BRAFV600E melanoma cells delays resistance by adaptive resensitization to drug rechallenge

Andrew J. Kavrana,1,2, Scott A. Stuarta,1,3, Kristyn R. Hayashia, Joel M. Baskena,4, Barbara J. Brandhuberc,5, and Natalie G. Ahna,b,6

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Patients with melanoma receiving drugs targeting BRAFV600E and mitogen-activated protein (MAP) kinase kinases 1 and 2 (MEK1/2), or downstream targets, mitogen-activated protein (MAP) kinase kinases 1 and 2 (MEK1/2), show clinical benefit in about 65% of patients with BRAFV600E-positive melanoma when administered continuously (4–6). However, resistance invariably develops, limiting median overall survival to ~2 y (7, 8). Many resistance mechanisms reactivate the MAPK pathway providing a growth advantage in the presence of an inhibitor. Known mechanisms include BRAFV600E amplification, alternative splicing of BRAFV600E, and other oncogenic mutations in the RAF and MEK pathways (9–14). Adaptive resistance also occurs in the absence of genomic alterations and can involve transcriptional changes through epigenetic mechanisms that promote the epithelial-to-mesenchymal transition (EMT), melanocyte dedifferentiation, and neural crest stem cell–like reemergence (15–26).

An emerging body of evidence has suggested that intermittent dosing schedules, in which periods of treatment with targeted therapeutics are interrupted by periods of drug removal, might have advantages over continuous treatment (2, 27, 28). Preclinical studies with patient-derived xenograft (PDX) melanomas or xenografts from established human melanoma cell lines showed that intermittent dosing can delay drug resistance and tumor growth compared with continuous dosing (29–32). Clinical reports and a phase 2 clinical trial have shown dozens of cases where patients with melanoma develop resistance and progress when treated with BRAF or MEK1/2 inhibitors continuously, but then show further response when retreated after a drug holiday period (32–37). By contrast, phase 2 trials of intermittent dosing with BRAF and MEK inhibitor combinations showed worse progression-free survival and no difference in overall survival compared with continuous treatment (38, 39). The reasons for variability in patient responses and trial outcomes are unknown and may reflect an incomplete understanding of mechanisms underlying the response to intermittent treatment.

Significance

Preclinical studies of metastatic melanoma treated with targeted therapeutics have suggested that alternating periods of treatment and withdrawal might delay the onset of resistance. This has been attributed to drug addiction, where cells lose fitness upon drug removal due to the resulting hyperactivation of mitogen-activated protein (MAP) kinase signaling. This study presents evidence that the intermittent treatment response can also be explained by the resensitization of cells following drug removal and enhanced cell loss upon drug rechallenge. Resensitization is accompanied by adaptive transcriptomic switching and occurs despite the sustained expression of resistance genes throughout the intermittent treatment.

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1A.J.K. and S.A.S. contributed equally to this work.

2Present address: Oncology Discovery, Bristol Myers Squibb, Cambridge, MA 02142.

3Present address: Department of Medicine, University of Colorado Anschutz Medical Campus, Aurora, CO 80045.

4Present address: Platform Operations, Enveda Biosciences, Boulder, CO 80301.

5To whom correspondence may be addressed. Email: natalie.ahn@colorado.edu.

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The current model explaining the beneficial response to intermittent treatment postulates the importance of drug addiction. Here, drug removal allows for the hyperactivation of MAPK signaling, which in turn leads to cell death or cell cycle arrest (2, 29–31, 40–42). Intermittent scheduling is thought to alternate between selection pressure against drug-sensitive cells during periods of drug treatment and selection against drug-resistant cells during periods of drug withdrawal. However, there is limited evidence that patient or xenograft tumors significantly regress when the drug is withdrawn, as predicted by the drug addiction model. Instead, xenograft tumors usually increase in volume with drug withdrawal and decrease volume after drug rechallenge (29, 43, 44). This raises the possibility that other mechanisms besides drug addiction may contribute to improved outcomes seen with intermittent treatment.

Here we use an in vitro strategy to examine cell autonomous responses of metastatic melanoma cells to intermittent treatment with the BRAF inhibitor, LGX818/encorafenib. Like vemurafenib and dabrafenib, LGX818 acts as a type I1/2 BRAF inhibitor (3), but its intermittent scheduling response has not been extensively examined. We report that an intermittent schedule with LGX818 substantially lowers cell viability compared with continuous treatment, in a manner that correlates with the degree of MAPK pathway activation. Both drug addiction following prolonged LGX818 treatment and drug resensitization following withdrawal can be observed over the multicyle time course. However, cell loss is greatest during periods of drug rechallenge, indicating that resensitization is the dominant mechanism underlying the efficacy of the intermittent schedule in this model. Transcriptome profiling through cycles of drug treatment and withdrawal reveals that resensitization is a reversible process that involves adaptive switching between states of drug resistance and drug sensitivity. Importantly, the transcriptome of the resensitized state can be distinguished from that of the initial, drug-sensitive state of naïve cells and occurs against a background of sustained elevation of MAPK signaling and known resistance mechanisms. Genes controlling adaptive switching between cell states may be useful targets to delay the onset of resistance in melanoma.

**Results**

**BRAF**<sup>V600E</sup> Amplification Confers Resistance to BRAF<sup>V600E</sup> and MEK1/2 Inhibitors. In order to generate cells with amplified BRAF/MAPK signaling, BRAF<sup>V600E</sup> was overexpressed in a human metastatic melanoma cell line (WM239A) under the control of a cumate-inducible promoter (SI Appendix, Fig. S1 A). The oncopogene was engineered as a fusion with green fluorescent protein (GFP) and used to confirm expression across the stable cell population after optimizing induction time and cumate concentration (SI Appendix, Figs. S1 B and D). A self-claving T2A sequence ensured complete separation of GFP from BRAF<sup>V600E</sup> following expression (SI Appendix, Fig. S1 E).

Induction of BRAF<sup>V600E</sup> increased the levels of active, phosphorylated ERK1/2 (ppERK), as measured by anti-ppERK immunoreactivity (Fig. 1A). Elevated MAPK signaling was confirmed by phosphorylation of the ERK substrate, RPS6KA (ppRSK, Fig. 1A). Both ERK and RSK phosphorylation were strongly blocked by 500 nM LGX818 (Fig. 1B). Partial BRAF<sup>V600E</sup> expression and pathway activation was apparent even in the absence of cumate, which reflected leakiness of the expression vector following selection (Fig. 1A and SI Appendix, Figs. S1 B and S2 A and B).

Dose–response experiments were used to measure the effect of BRAF<sup>V600E</sup> overexpression on drug resistance. Cells with amplified BRAF<sup>V600E</sup> increased the half-maximal inhibitory concentration (IC<sub>50</sub>) by 50-fold over control cells harboring empty vector (Fig. 1C). Similar increases in IC<sub>50</sub> were seen in dose–response measurements with MEK162/binimetinib (SI Appendix, Fig. S2 D). Thus, BRAF<sup>V600E</sup> overexpression strongly increased chemoresistance toward BRAF<sup>V600E</sup> and MEK inhibitors.

**Comparison of Intermittent and Continuous Treatment**

In order to compare intermittent and continuous treatment schedules, cells were seeded in 96-well dishes and cultured for 4 wk, monitoring cell viability at the end of each week (Fig. 2 A). Intermittent time courses followed a schedule of 7 d on the drug followed by 7 d off the drug, where cells were treated with 500 nM LGX818 during weeks 1 and 3 and the drug was removed in weeks 2 and 4. In parallel, cells were treated continuously with 500 nM LGX818 over the entire 4-wk period. Cells treated continuously grew slowly for the first 2 wk after which a drug-resistant population emerged (Fig. 2B). By contrast, cells treated intermittently yielded cell numbers that were initially similar to the continuous experiment for the first 2 wk, but then declined after drug rechallenge in week 3 (Fig. 2B). Thus, intermittent treatment inhibited the cell expansion seen with continuous treatment. Control cells expressing empty vector were strongly suppressed with either treatment schedule (Fig. 2C).

In order to compare responses to BRAF<sup>V600E</sup> against other oncogenes associated with resistance in melanoma, cells were engineered to individually express MEK2<sup>C125S</sup>, EGFR<sup>L858R</sup>, NRAS<sup>Q61K</sup>, or the p61-BRAF<sup>V600E</sup> splice variant (8–11). Like
assays and normalized to the initial number of cells seeded. Cell numbers (mean ± SEM) measured at each time point for (B) WM239A-BRAFV600E cells (n = 5 or 6) and (C) WM239A empty vector cells (n = 6) are shown for continuous and intermittent treatments. (D) Western blots of lysates separated by low-bis sodium dodecyl-sulfate polyacrylamide gel electrophoresis show ERK phosphorylation in cells expressing empty vector (lane 1), BRAFV600E (lane 2), and BRAFV600E cells treated continuously “C” or intermittently “I” across the 4 wk with 500 nM LGX818.

BRAFV600E, cells expressing p61-BRAFV600E or MEK2C125S substantially increased the phosphorylation of ERK and RSK as well as IC50 with LGX818 or MEK162 (SI Appendix, Fig. S2 A–D). Cells expressing p61-BRAFV600E displayed the greatest resistance to LGX818, with 10-fold higher IC50 than that of BRAFV600E or MEK2C125S. By contrast, EGFR1858RR or NRASQ61K only modestly increased ERK or RSK phosphorylation and IC50 (SI Appendix, Fig. S2 A–D). ERK and RSK phosphorylation were suppressed in all cells by 500 nM LGX818, MEK162, or the inhibitor combination (SI Appendix, Fig. S2 E–G).

During continuous treatment with LGX818, cells expressing MEK2C125S or p61-BRAFV600E remained static for the first week, after which a resistant population emerged (SI Appendix, Fig. S3 A and B). Like BRAFV600E, cells expressing MEK2C125S declined with intermittent treatment and remained inhibited for the duration of the time course (SI Appendix, Fig. S3 A). Intermittent treatment with either LGX818 or MEK162 only partially inhibited growth of cells expressing p61-BRAFV600E compared with continuous treatment (SI Appendix, Fig. S3 B and C). However, cells were substantially reduced by intermittent treatment with a combination of both LGX818 + MEK162 (SI Appendix, Fig. S3 D), indicating that strong inhibition of the MAPK pathway during periods of drug addition is important for maximal efficacy. Cells expressing EGFRL858R or NRASQ61K were strongly repressed by either intermittent or continuous treatment, both of which effectively inhibited expansion over the 4 wk time course (SI Appendix, Fig. S3 E and F).

Thus, intermittent treatment showed greater efficacy compared with continuous treatment, but only in cells with the highest levels of ERK activation and the strongest resistance to the inhibitor.

Studies invoking the drug addiction model for intermittent dosing have reported pronounced elevation of ppERK with BRAFV600E amplification (29). Therefore, we characterized the corresponding responses to continuous or intermittent treatment in our BRAFV600E cell system. Western blots showed that ppERK increased with BRAFV600E induction and decreased after the first week in the presence of LGX818, then increased with subsequent weeks of continuous treatment as resistant cell populations emerged (Fig. 2D, lanes 2–4, 6, and 8). By comparison, ppERK was elevated after weeks 2 and 4 of drug removal (lanes 5 and 9), decreasing when the drug was added back during week 3 (Fig. 2D, lane 7). The findings show that continuous treatment with LGX818 maintained levels of ERK activity that supported viability in the presence of the drug, while drug removal during intermittent treatment elevated ERK to levels equal to or greater than seen with the initial induction of BRAFV600E (Fig. 2D, lanes 2, 5, and 9).

**Intermittent Treatment Reverses Drug Addiction and Resensitizes Cells to LGX818.** To better understand the effect of the intermittent treatment schedule on drug sensitivity and drug addiction, we characterized the LGX818 dose–response of BRAFV600E cells collected at the end of each week of a continuous or intermittent time course. In this experiment, cells that were cumate induced to express BRAFV600E showed greater drug resistance (IC50 = 130 nM) than cells with empty vector (IC50 = 5 nM) (Fig. 3A). After continuous treatment with 500 nM LGX818 for 7 d, the IC50 increased to 860 nM and remained sustained at 1,000 to 1,200 nM over successive weeks (Fig. 3A). Continuous treatment also resulted in drug addiction, as evidenced by a 30 to 45% reduction in cell numbers at 0 nM LGX818 relative to their maximum levels at 100 nM (Fig. 3A). Thus, drug addiction accompanied resistance to LGX818 in our experimental system, consistent with previous models of resistance to MAPK pathway inhibitors (29, 30, 41–43).

By contrast, cells treated intermittently varied in their dose–response to LGX818, depending on whether they were collected after periods of drug addition or drug removal. Removing LGX818 in week 2 reduced the IC50 to 300 nM, a threefold decrease from cells treated with the drug in week 1 (Fig. 3B). At the same time, removing LGX818 decreased the extent of drug addiction, as shown by the recovery of cell viability at 0 nM LGX818 (Fig. 3B). Rechallenge with LGX818 in week 3 reversed this behavior, increasing both IC50 and drug addiction back to the levels seen with continuous treatment. Removing LGX818 in week 4 decreased the IC50 and decreased drug addiction back to levels comparable to week 2. Therefore, each cycle of drug removal switched cells to a state with resepsitization to the BRAF inhibitor and each cycle of drug addition produced a state of drug addiction.

**Cell Loss during Intermittent Treatment Primarily Involves Resensitization after Drug Removal.** Conceivably, either drug addiction or drug resepsitization could account for the loss of cell viability observed with intermittent treatment. In order to
explore the contribution from each mechanism, we quantified cell death at different times during periods of drug addition or removal. On one hand, if drug addiction were the dominant mechanism, cell loss should be highest during drug-off weeks when ERK hyperactivation would be predicted to promote cell death. On the other hand, if drug resensitization were more important, cell loss should be highest during drug-on weeks when cells would be susceptible to rechallenge with LGX818.

To examine this, cells were seeded in 10-cm dishes and treated with continuous or intermittent schedules over 5 wk. Samples were taken on days 3 and 7 of each week, collecting all adherent and floating cells for analysis of cell viability by propidium iodide (PI) staining using flow cytometry (Fig. 3C). Propidium iodide was used as a marker of cell death, which is caused by parthanatos following drug withdrawal in drug-addicted melanoma cells (30). Continuous treatment led to substantial cell loss during week 1, which fell to lower levels over successive weeks (Fig. 3C). Parallel flow cytometry measurements showed BRAFV600E expression increased over the first 2 wk of continuous treatment, then maintained over successive weeks (SI Appendix, Fig. S4).

In cells treated intermittently, the percentages of PI-positive cells during drug-off periods (days 10, 14 and 24, 28) were comparable to those in cells treated continuously (days 10 through 28). However, cell death dramatically increased during periods of drug rechallenge (days 17, 21 and 31, 34) where the percentage of PI-positive cells reached levels as high as 60% (Fig. 3C). These results indicate that the reduction in cell fitness with intermittent treatment is best explained by the occurrence of resensitization during periods of drug withdrawal, followed by cell loss when cells are rechallenged with LGX818. This was consistent with the measurements of cell numbers during intermittent treatment, which decreased only when the drug was readded in week 3 (Fig. 2B).

**Transcriptomic Responses to Continuous and Intermittent Treatment.** In order to explore gene expression changes that accompany drug resensitization, RNA sequencing (RNA-seq) was used to examine cells expressing BRAFV600E after each week of intermittent or continuous treatment with LGX818 (Fig. 4A). Cell viability measurements conducted in parallel matched those observed previously, where resistant cells emerged after 2 wk of continuous treatment with LGX818 and
intermittent treatment delayed outgrowth (SI Appendix, Fig. S5). Datasets were also collected on cells with empty vector, BRAFV600E induced for 72 h, and BRAFV600E + 20 h LGX818 (Fig. 4B, group 1). A second group contained all cells that were continuously treated (weeks 1 through 4), as well as cells from intermittent week 3, when the drug was reintroduced (Fig. 4B, group 2). The third group included cells treated intermittently in weeks 2 and 4, when the drug was removed (Fig. 4B, group 3). The analysis revealed a striking effect of intermittent treatment, in which cells with the drug removed in weeks 2 and 4 were grouped together and well separated from cells with the drug readded in week 3.

Significantly, the separation between groups 1 and 2 mainly occurred along the PC1 axis, suggesting transcriptomic changes resulting from long-term treatment with LGX818. The separation between cells collected in successive weeks of intermittent treatment occurred primarily along the PC2 axis, corresponding to reversible movement between groups 2 and 3 (Fig. 4B). This suggested that PC2 largely recapitulates the transcriptomic state of continuously treated cells. Thus, reversible changes in the transcriptome accompany the ability of cells to switch between drug-resistant and -resensitized states during the intermittent time course.

**Transcriptome Changes Associated with Drug Resistance.** We asked whether the transcripts altered in response to continuous LGX818 treatment might reflect genes that function in cancer drug resistance, using gene set enrichment analysis (45, 46). The PC1 genes that increased expression with prolonged treatment were enriched in molecular signatures associated with resistance to BRAF and MEK inhibitors, including markers of the EMT, NF-κB signaling, inflammatory markers, and acute activation or inhibition of the MAPK pathway without treatment, in which cells with the drug removed in weeks 2 and 4 were grouped together and well separated from cells with the drug readded in week 3.

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**Fig. 4.** Transcriptomic profiling of continuously and intermittently treated cells. (A) Schematic of RNA samples collected from WM239A-BRAFV600E cells comparing continuous and intermittent drug treatment with 500 nM LGX818. Each condition was collected in biological triplicates, except for week 4 continuous (sample 7), which was collected in duplicates, for a total of 29 samples. (B) Principal component analysis (PCA) of the 6,000 highest variance genes. PC1 and PC2 account for almost 70% of the variance in these genes. The 10 different experimental conditions visibly cluster into three separate groups. (C) Heatmap of the 400 genes most positively or negatively associated with PC1 or PC2. Each row is mean centered and scaled to unit variance.
and hypoxia, while genes that decreased expression were enriched in gene sets associated with a proliferative, melanocytic phenotype (Fig. 5A). These signatures are characteristic of the invasive, dedifferentiated melanoma phenotype associated with melanoma malignancy (15–17, 23). The PC2 genes that reversibly decreased upon drug removal were negatively associated with gene signatures characteristic of the invasive phenotype, suggesting partial reversal of invasion/EMT-like processes.

We next assessed how the expression levels of genes responsive to continuous LGX818 treatment correlated with drug resistance across human melanoma cell lines. RNA-seq data

![Fig. 5. Overview of gene expression changes associated with EMT-like responses.](https://doi.org/10.1073/pnas.2113535119 pnas.org)
from 34 BRAFV600 melanoma cell lines available from the cancer cell line encyclopedia (CCLE) (47) were used to perform hierarchical clustering based on the expression levels of transcripts highly associated with PC1. These cell lines formed two main clusters, denoted cluster A and cluster B in Fig. 5B. The gene expression patterns of cell lines in cluster A resembled transcript levels in our drug-naive cells (group 1 in Fig. 4B), while cell lines in cluster B showed greater similarity to our cell system after long-term treatment with LGX818 (groups 2 and 3 in Fig. 4B). The degree of resistance among CCLE cell lines, measured by IC50 for the BRAFV600E inhibitor, PLX4720, or the MEK inhibitor, AZD6244, correlated well with cluster membership, largely separating drug-sensitive cells in cluster A from drug-resistant cells in cluster B (Fig. 5B). The analysis reveals that the transcript changes associated with LGX818 resistance in WM239A cells can explain the variance in baseline IC50 across many other BRAFV600E cell lines. Therefore, the transcriptome responses to continuous treatment in our experimental model are consistent with those associated with drug resistance across melanoma cell systems.

It was noteworthy that the transcriptomes of cells in group 3, characterized by drug removal, were distinct from those of cells with empty vector or induced BRAFV600E in group 1 (Fig. 4B and C). This implies that the resensitized cell state generated by LGX818 removal differs from the initial sensitive state of drug-naive cells. Prominent among the PC2 transcripts that switched reversibly following drug removal and subsequent rechallenge were those that were weakly responsive or in some cases nonresponsive to BRAFV600E induction in drug-naive cells (Fig. 4C). This implies that many PC2 genes normally respond weakly to MAPK signaling but acquire greater responsiveness to this pathway after a week of continuous exposure to the drug. Thus, the resensitized cell state includes new transcriptional responses that only occur after MAPK signaling is activated by drug removal.

We also compared transcript changes corresponding to known resistance genes in melanoma. To do this, we curated a list of 73 genes that were reported to control resistance toward BRAFV600E and/or MEK inhibitors in melanoma cell lines and tumors (genes and references in Dataset S4). These included signaling effectors, transcription factors, and mediators and markers of EMT-like responses, neural crest specification, and melanocyte differentiation. Forty-eight of the curated genes were significantly altered with continuous drug treatment, consistent with the emergence of resistant cell populations. For example, growth factor receptors and ligands, EMT-like and neural crest markers known to promote resistance (e.g., PDGFRB, EGFR, AXL, NGFR, and WNT5A) (12, 21, 42, 48, 49), increased with continuous drug treatment, while genes associated with differentiation to the melanocyte lineage (e.g., MITF, TYR, MLANA, and SOX10) (17, 21) decreased (Fig. 5C).

Among the curated genes associated with drug resistance in melanoma, some transcripts changed reversibly with LGX818 removal in a manner consistent with resensitization (Fig. 5D). For example, growth factors NGF and GAS6 decreased during drug-off weeks, which might predict lower signaling through their respective receptors, NGFR and AXL. Likewise, the transcription factor ID3, which promotes resistance to BRAF inhibitor (50), decreased after drug removal, while LTF1, which associated with greater sensitivity (19, 26, 51), increased. Negative feedback regulators of growth factor signaling, DUSP4/6 and SPRY2/4, reversibly increased after drug removal, consistent with MAPK pathway reactivation (52, 53).

However, the majority of resistance genes from our curated list were inconsistent with resensitization and in fact predicted sustained resistance with intermittent scheduling. These included growth factor receptors/ligands (AXL, NGFR, EGFR, PDGFRB, PDGFB, and WNT5A) and transcription factors (JUN, FOSL1/FRA1, and ZEB1), which remained elevated following drug removal, and differentiation markers (MLANA/MART1, TYR, MITF, and SOX10), which remained repressed (Fig. 5C). Paradoxically, other responsive transcripts predicted increased resistance upon drug removal. For example, FGFR1, MET, AREG, TGFB1, IL1B, ITGB3, and CSPG4 were all up-regulated during drug-off weeks (Fig. 5D), which would be expected to increase receptor signaling and diminish sensitivity to BRAF inhibitors (42, 54–60). Also included in this group were the transcription factors TWIST1, SNAI1, and RELA, which promote EMT-like responses via epigenetic repression of histone marks (61, 62), but increased when the drug was removed (Fig. 5C and E) and SNAI2, which promotes the dedifferentiated, noninvasive phenotype (61), but increased with drug removal.

Taken together, the gene expression analyses revealed distinct molecular responses to the continuous and intermittent treatment regimes. Many transcripts altered by prolonged drug treatment corresponded to genes implicated in drug resistance in melanoma and were regulated in a manner consistent with the drug-resistant state observed with continuous treatment. Other transcripts switched expression between states of resistance and sensitivity, but only a subset of known resistance genes reversed in a manner consistent with resensitization upon drug removal.

Adaptive Responses to Intermittent Treatment. As noted above, many transcript changes that were significant after the first 7 d of LGX818 exposure were sustained when the drug was removed in weeks 2 and 4 (Figs. 4C and 5C). Their irreversibility may be due to slow rates of reversal or to an initial selection for drug-resistant cell subpopulations. On the other hand, nearly all genes that changed reversibly during the course of intermittent treatment were found to recover the transcriptome of continuously treated cells during week 3, when the drug was readded (Figs. 4C and 5D). This implies an adaptive mechanism, responsive to epigenetic regulation. However, bulk RNA-seq cannot distinguish between mechanisms involving adaptive transitions in cells switching from drug-resistant to drug-sensitive states, and mechanisms for selection pressure for sensitive cell subpopulations within a resistant majority.

To investigate these possibilities, we examined the dynamics of expression of the neural cell adhesion protein, L1CAM. Bulk transcript levels of L1CAM were elevated in cells treated for prolonged periods with LGX818, consistent with its role in promoting EMT-like responses and as a marker for melanoma drug resistance (18, 63, 64). Its expression decreased following drug removal in a manner that was reversible in subsequent weeks of drug rechallenge and withdrawal (Fig. 5D). The changes in L1CAM protein expression at the single-cell level were measured by flow cytometry at several time points during continuous and intermittent treatment using a fluorescently labeled primary antibody (Fig. 6A). We selected these time points in order to capture the cell population dynamics corresponding to reversal of expression during early stages of drug withdrawal. In drug naive cells (day 0), many cells expressed low levels of L1CAM, some overlapping with the nonspecific isotype control (Fig. 6B). After treating cells with LGX818 for 7 d, the median expression level increased by more than 10-fold over that of drug-naive cells and remained high for 14 d of continuous drug treatment. In the intermittent
schedule, L1CAM expression during the first 3 d of drug withdrawal gradually decreased in a unimodal fashion (Fig. 6B). The cells transiently expressed intermediate levels of L1CAM, which then decreased further to match the drug-naive population by the end of the week. The unimodal populations with intermediate L1CAM were inconsistent with selection pressure against a subpopulation of drug-addicted cells, which would have predicted bimodal cell populations shifting from high to low L1CAM. Therefore, the transcript responses following drug withdrawal largely follow an adaptive mechanism for transcription switching, instead of selective depletion of drug-addicted cells.

Discussion

Our study demonstrates that adaptive mechanisms for cell switching from drug-resistant to drug-resensitized states can explain the growth suppressive effects of treating melanoma cells intermittently with LGX818. The BRAF\textsuperscript{V600E}\textsuperscript{-}amplified cell model used in our study showed transcripts responsive to the drug that corresponded well with resistance genes known to function across melanoma cell lines and tumors. Importantly, when drug resistance was induced by prolonged treatment, the cells displayed both a drug addiction response after treatment and a drug resensitization response after drug withdrawal. Cell death was highest during periods of rechallenge, showing that cell viability was affected most by the resensitization to kinase inhibitor. Transcriptome analyses showed many gene expression changes that were readily reversible over the intermittent time course, nearly all which returned to levels seen in continuously treated cells when the drug was reintroduced. This indicated that the responses to intermittent treatment were not due to selection pressure, as supported by the expression dynamics of L1CAM in single-cell populations.

Previous studies have shown that drug addiction is caused by resistance mechanisms that maintain MAPK pathway signaling at submaximal levels in the presence of BRAF inhibitor, and hyperactivate MAPK signaling when the drug is removed, thereby triggering cell death or cell cycle arrest (29, 30, 40–42). Thus, it has been theorized that the intermittent treatment effect arises by alternating selection against drug-sensitive cells in the presence of BRAF inhibitor and counterselection against drug-addicted cells in the absence of the inhibitor (2). In our system, MAPK signaling was maximally activated following LGX818 withdrawal, based on increased phospho-ERK1/2 and the expression of downstream pathway targets such as DUSP and SPRY. But while drug addiction was evident, it played a secondary role in the response to intermittent treatment. In cells treated continuously, cell viability was highest at 100 nM LGX818 and at 500 nM fell to levels less than or equal to that in the absence of the drug (Fig. 3A). This may explain why drug addiction did not contribute significantly to growth suppression with intermittent treatment in our experiments, because it would have had its largest effect at drug concentrations lower than we used. While it is hard to extrapulate in vitro cell behavior to clinical outcomes, plasma concentrations of BRAF inhibitors reached in patients (65, 66) may affect the extent to which drug addiction occurs.

Our findings are consistent with a role for phenotypic plasticity in drug resistance, which has been well documented in BRAF\textsuperscript{V600E}\textsuperscript{-}mutated melanomas as well as other cancers (15–17). Current models postulate elevated expression of genes in rare cell populations, which enhance resistance in response to BRAF inhibitor by promoting epigenetic pathways for dedifferentiation and transcriptional reprogramming (18–26). Classes of transcripts that promote resistance include drivers of EMT and invasion (AXL\textsuperscript{high}, WNT5A\textsuperscript{high}, TGF\textsuperscript{high}, TWIST\textsuperscript{high}, and SNAI1\textsuperscript{high}) and markers of lineage development from differentiated melanocytes to neural crest stem cells (NGFR\textsuperscript{high}, MITF\textsuperscript{low}, and SOX10\textsuperscript{low}) (20–26). Exploiting the reversibility of these adaptive regulatory events during early, nonmutational phases of drug resistance has been proposed as a treatment strategy for cancer. Our findings concur and further suggest that adaptive mechanisms may underlie beneficial responses to intermittent scheduling.

Significantly, transcriptome profiling of our system revealed many resistance genes that changed with continuous drug treatment but were irreversible over the intermittent time course. These reflect adaptive responses to drug and/or the selection of cell populations able to persist and survive during the first week of LGX818 treatment. This set included most genes characteristic of the invasive, neural crest phenotype and EMT-like responses. This means that resensitization after drug removal occurred despite a large number of resistance genes that did not reverse. Altogether, only a few genes implicated in resistance both displayed reversibility and changed in a direction consistent with...
resensitization. Leading candidates included LEF1, which is repressed in drug resistance melanoma and strongly associated with phenotype switching to the proliferative, noninvasive state (19, 26, 51), and ID3, which is elevated in resistant cells and whose depletion in resistant cells confers resensitization (50). They did not include genes reported to regulate drug addiction in melanomas, such as JUNB, FOSL1, or CDKN1A (40, 41), consistent with a minimal influence of drug addiction on the intermittent treatment effect.

Interestingly, many transcripts that responded reversibly with each cycle of drug removal were elevated to levels well above those seen in drug-naive cells. In fact, many reversible transcripts showed little or no response to short-term induction of BRAFV600E. This suggests that genes that were not normally downstream of MAPK signaling became pathway targets after 7 d of drug treatment. Conceivably, drug treatment may have enriched a subpopulation of cells with an expanded range of transcriptional responses. Alternatively, new targets may have been triggered by the hyperactivated MAPK signaling that followed drug withdrawal. Further definition of the cellular mechanisms involved in reversible resensitization may lead to novel targets and the potential contribution of MAPK signaling thresholds to the resensitized cell state.

The partial reversal of resistance gene expression upon drug removal helped explain the partial resensitization response of only a threefold decrease in IC50. Therefore, the drug resistant state in our hands appears to represent an intermediate-to-late stage in transcriptome reprogramming, with a few rarely reversible and many slowly reversible resistance genes (21). Conceivably, a longer period of drug withdrawal might have eventually returned cells to the initial transcriptome and sensitivity level of drug-naive cells, as observed in other studies of transcriptome dynamics (25). However, in order for intermittent treatment to block cell expansion, the amount of time needed for resensitization must be balanced against the faster cell doubling time during the period of drug removal. Therefore, partial resensitization may be a practical condition needed for a successful intermittent treatment schedule.

So far, intermittent treatment of melanoma has been unproven, with clinical outcomes arguing for and against its potential effectiveness. On one hand, two recent phase II trials revealed worse progression-free survival in patients with melanoma treated intermittently with dabrafenib + trametinib or vemurafenib + cobimetinib compared with those treated with continuous therapy, with no difference in overall survival (38, 39). On the other hand, retrospective and prospective studies of dozens of patients who progress on BRAF or MEK inhibitor have reported that more than one-third show a second clinical response after a drug holiday period (27, 33–37). Preclinical studies using xenografts have been mixed as well. While some have reported delayed emergence or complete suppression of resistant tumors using intermittent scheduling (29–32), others have observed more rapid outgrowth of tumors (43, 44). Factors that could affect whether beneficial responses are seen with intermittent treatment, in vitro or in animals, may include drug concentration relative to IC50, doubling times for resistant vs. sensitive cells, and cell specificity in signaling pathway activation. A notable report showed a substantial benefit of intermittent treatment in xenografts from a drug-resistant melanoma cell line, but only when the dosing schedule was tailored individually for each mouse (32). Optimal scheduling was established using a predator–prey model for adaptive drug therapy, which postulates expansion and loss of sensitive cells during drug-off and drug-on periods, respectively (67, 68). Early stage clinical trials are ongoing to test these intriguing concepts for melanoma, breast, and prostate cancer, and the possibility that personalized scheduling may be optimized by dynamic measurements of cancer blood markers (68–70).

In summary, our findings show that intermittent treatment can suppress cell growth and delay the emergence of drug resistance by transitioning from a drug-resistant state to a more sensitive state through adaptive transcriptional mechanisms. Significantly, this can occur against a background of many resistance genes that either fail to reverse when the drug is removed or change in a manner that would paradoxically predict increased resistance. Thus, we propose that intermittent treatment generates a distinct cellular state that accompanies resensitization, which may include transcripts not normally regulated by MAPK signaling and are triggered by higher signaling thresholds. Genes controlling resensitization may be useful targets to augment or improve the response duration of current treatment strategies for melanoma.

Materials and Methods

The human metastatic melanoma cell line, WM239A, was a kind gift from Meenhard Herlyn, Wistar Institute, Philadelphia, PA. LGX818/encorafenib and MEK162/cobimetinib were obtained from Selleck Chemicals. The CellTiter-Glo 2.0 assay (Promega) was used to determine cell numbers. Detailed methods for construction of cell lines, biochemical- and cell-based assays, and RNA-seq measurements are described in detail in SI Appendix, which includes SI Appendix, SI Materials and Methods, Figs. S1–S6, and Datasets S1–S4.

Data Availability. RNA-seq data have been deposited in NCBI Gene Expression Omnibus (GSE117123). A runnable and editable version of the code used in this study can be found on code ocean (https://codeocean.com/capsule/9070543/tree/v1), and a code repository is available on GitHub at https://github.com/andykavan/Intermittent_Drug_Treatment. All other study data are included in the article and/or supporting information.

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Author affiliations: 4Department of Biochemistry, University of Colorado, Boulder, CO 80309; 5BioFrontiers Institute, University of Colorado, Boulder, CO 80309; and 6Structural Biology, Array BioPharma, Boulder, CO 80301

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