p53 deficiency triggers dysregulation of diverse cellular processes in physiological oxygen

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The mechanisms by which TP53, the most frequently mutated gene in human cancer, suppresses tumorigenesis remain unclear. p53 modulates various cellular processes, such as apoptosis and proliferation, which has led to distinct cellular mechanisms being proposed for p53-mediated tumor suppression in different contexts. Here, we asked whether during tumor suppression p53 might instead regulate a wide range of cellular processes. Analysis of mouse and human oncogene-expressing wild-type and p53-deficient cells in physiological oxygen conditions revealed that p53 loss concurrently impacts numerous distinct cellular processes, including apoptosis, genome stabilization, DNA repair, metabolism, migration, and invasion. Notably, some phenotypes were uncovered only in physiological oxygen. Transcriptomic analysis in this setting highlighted underappreciated functions modulated by p53, including actin dynamics. Collectively, these results suggest that p53 simultaneously governs diverse cellular processes during transformation suppression, an aspect of p53 function that would provide a clear rationale for its frequent inactivation in human cancer.

Introduction

The transcription factor p53 is a critical barrier to the development of cancer, as evidenced by three key observations. First, more than half of all human cancers are associated with direct inactivating mutations in TP53 (Hollstein et al., 1991; Levine, 2018). Second, Li-Fraumeni patients, who inherit inactivating mutations in TP53, are predisposed to early-onset cancers, including breast cancers and sarcomas (Hollstein et al., 1991). Finally, p53-null mice succumb to cancer with 100% penetrance (Donehower et al., 1992; Jackson et al., 1994; Kaiser and Attardi, 2018). Intriguingly, despite 40 yr of research on p53, the specific mechanisms by which p53 suppresses tumorigenesis remain incompletely understood.

The best-characterized function of p53 is as a transcription factor. In this role, p53 can elicit widespread transcriptional changes in response to various cellular stresses such as DNA damage, oncogene activation, and hypoxia, to promote specific cellular responses (Mello and Attardi, 2018; Vousden and Prives, 2009). The most studied p53 cellular responses are apoptosis and cell cycle arrest. Apoptosis is triggered by p53 in response to stressors such as acute DNA damage, via transcriptional up-regulation of proapoptotic genes such as the Bcl-2 family members Puma and Noxa. p53 can also limit cellular proliferation by inducing G1 or G2 arrest.

For example, at the G1/S boundary, p53 promotes transient cell cycle arrest in response to genotoxic damage by up-regulating the p21 CDK inhibitor, a measure that is thought to facilitate DNA repair (Karimian et al., 2016). It is through these canonical responses that p53 was proposed to act as a “guardian of the genome,” ensuring that upon acquisition of DNA damage, cells would transiently arrest to repair this damage or, alternatively, undergo apoptosis to remove damaged cells from the organism, thereby suppressing oncogenic transformation (Lane, 1992).

Several studies have suggested that p53 suppresses tumorigenesis through responses other than acute DNA damage-induced cell cycle arrest or apoptosis. First, a few reports suggested that the pathological response to acute DNA damage by p53, involving widespread induction of apoptosis, is dispensable for p53-mediated tumor suppression (Christophorou et al., 2006; Efeyan et al., 2007; Hinkel et al., 2009). Expanding on this idea were seminal studies using either knock-in mice bearing alterations to p53 that partially impaired its transcriptional capacity (p5325,26 [Brady et al., 2011; Jiang et al., 2011] and p533KR [Li et al., 2012b]) or gene-targeted mice lacking the canonical p53 target genes p21, Puma, and Noxa (Valente et al., 2013). Cells from these...
mice displayed defective p53-dependent induction of apoptosis and cell cycle arrest upon acute DNA damage, but the mice were nonetheless resistant to spontaneously arising tumors (Li et al., 2012b; Valente et al., 2013) and cancer in oncogene-driven mouse cancer models (Brady et al., 2011; Jiang et al., 2011). Together, these studies suggest that processes other than p53 responses to acute DNA damage are critical for tumor suppression or, alternatively, that in the absence of these responses, other processes can compensate to impede tumor development (Mello and Attardi, 2018).

A natural result of this revised view of p53-mediated tumor suppression has been increasing emphasis on understanding noncanonical p53 functions. Specifically, beyond regulating proliferation and apoptosis, p53 has been reported in specific settings to regulate additional cellular processes such as metabolism and stemness (Charni et al., 2017; Kaiser and Attardi, 2018). However, it is unclear whether the ability of p53 to regulate each cellular behavior is context dependent, with a specific p53-regulated process being fundamental for tumor suppression only in a particular tumor type. For example, p53-mediated tumor suppression in large T antigen-driven choroid plexus tumors is associated with oncogene-triggered apoptosis, whereas p53-induced apoptosis and DNA repair programs are critical for the suppression of Eµ-Myc-driven B cell lymphomas (Eischen et al., 2001; Garrison et al., 2008; Hemann et al., 2004; Janic et al., 2018; Michalak et al., 2009; Yin et al., 1997). Although tumor suppression has been ascribed to one or two specific p53 cell biological functions in such cancer models, a systematic evaluation of a wide range of p53 functions has not been performed within one specific cellular context. Determining how broadly p53 regulates diverse cellular processes in a particular context is a first step toward understanding the cellular basis of p53-mediated tumor suppression. Indeed, the complexity of the p53 transcriptional program, with alterations in the expression of hundreds of target genes, suggests that p53 might simultaneously regulate a variety of cellular functions during transformation suppression (Andrysik et al., 2017; Fischer, 2019).

Here, we examined to what extent the effects of p53 loss are pleiotropic, using an in vitro oncogene-expressing primary mouse embryonic fibroblast (MEF) model system in which p53 plays a critical role in suppressing transformation. Although transformation suppression by p53 in this setting has been associated predominately with the induction of apoptosis (Lowe et al., 1993; Soengas et al., 1999), we sought to interrogate the role of p53 in regulating a range of other cell biological functions in this context. Importantly, we performed these experiments in physiological (5%) oxygen tensions to more closely model in vivo conditions. By leveraging a spectrum of assays for different cellular processes, we revealed that p53 regulates an array of diverse cellular processes in this context, several of which were apparent only under physiological oxygen conditions. These findings support the intriguing notion that p53-mediated tumor suppression is a complex coordinated process reliant on modulation of numerous cellular programs.

**Results**

**Establishing a platform to interrogate global p53 functions**

To interrogate the capacity of p53 to globally regulate a variety of cellular processes during transformation suppression, we examined EIA:HrasG12V oncogene-expressing MEFs, as they provide a tractable model in which different cell biological assays can readily be performed and in which p53 plays a potent role as a transformation suppressor. Moreover, unlike many human cancer cell lines, these are primary, early-passage cells into which oncogenes are retrovirally transduced, and they therefore retain intact p53 signaling pathways. Of note, although previous studies have suggested that transformation suppression in these cells is due to apoptosis in response to cellular stress signals (Lowe et al., 1994; Soengas et al., 1999), we sought to determine whether other cellular processes are also regulated simultaneously by p53. To best model the oxygen tensions that most cells encounter in vivo, which range between 2 to 8% for most cells, we cultured cells in physiological oxygen (5% O2) rather than the standard atmospheric 21% O2 conditions.

To establish isogenic WT and p53-deficient cell lines, we generated early-passage MEFs from HrAg109 mice, in which Cas9 is constitutively expressed from the H11 promoter (Chiou et al., 2015). Three independent MEF lines were transduced with EIA- and HRasG12V-expressing retroviruses, and then infected with lentiviruses expressing one of two sgRNAs targeting exon 4 of Trp53 (sgp53) or a nontargeting control (sgNTC) sgRNA (Fig. 1, A and B; and Fig. S1 A). We thus established a panel of early-passage, isogenic, polyclonal p53 WT (sgNTC) and p53-deficient (sgp53) EIA- and HRasG12V-expressing cell lines (three sgNTC and six sgp53 cell lines). We confirmed p53 deficiency by DNA sequencing of the p53 locus (Fig. S1 B), immunoblotting and immunofluorescence analyses of p53 protein levels (Fig. 1, C and D; and Fig. S1 C), and Western blot or quantitative RT-PCR (qRT-PCR) analysis of p53 target gene expression (Fig. 1, C and E; and Fig. S1, C and D).

To confirm known p53-dependent phenotypes in this model system, we first assayed transformation suppression using soft agar assays, a robust in vitro surrogate of in vivo tumorigenicity (Lin et al., 1998). In 5% O2, EIA:HrasG12V MEFs targeted with sgp53 exhibited significantly greater cell colony formation than sgNTC controls, confirming that p53 inactivation in these cells enhanced their tumorigenic capacity (Fig. 1, F and G; and Fig. S1 E). We next examined apoptosis in response to two distinct stresses: acute DNA damage and serum starvation. Whereas sgNTC EIA:HrasG12V MEFs exhibited significant apoptosis in response to acute DNA damage or serum starvation, sgp53-targeted cells were protected from cell death induced by these stimuli, validating the p53-dependent cell death response to different stressors in EIA:HrasG12V MEFs (Fig. 1 H and Fig. S1 F). We next sought to understand which additional p53 downstream pathways might also be regulated during transformation suppression by performing a panel of phenotypic assays examining various cellular processes.

**p53 does not dampen cellular proliferation in oncogene-expressing MEFs**

The canonical responses of p53 to stress stimuli include not only induction of apoptosis, but also inhibition of cellular proliferation. Although the best-characterized p53 function in this regard is in driving cell cycle arrest or senescence in response to a specific stress signal, p53 can also simply dampen proliferation rates (Tyner et al., 1999). We thus sought to determine whether p53 can regulate oncogene-driven cell proliferation in this
cellular model. We found no significant difference in the proliferative rate between sgNTC- and sp53-targeted E1A:HrasG12V-expressing MEFs (Fig. 1, J and K). Although p53 is classically considered a regulator of the G1-S transition, expression of E1A in this cellular model inactivates retinoblastoma (Rb) family member-mediated inhibition of cell cycle progression, the primary mechanism by which p53 is thought to inhibit cell proliferation (Deng et al., 2005; Narita et al., 2003). Thus, while p53 is capable of triggering apoptosis in E1A:HrasG12V-expressing MEFs, it does not clearly inhibit cell proliferation in this model.

p53 regulates ploidy in E1A;HrasG12V MEFs

A hallmark of dysregulated G1-S progression triggered by p53 deficiency is genomic instability, an observation that earned p53 the title of “guardian of the genome” (Lane, 1992). p53 is thought to preserve genomic integrity by inducing G1 arrest when cells have sustained DNA damage or have undergone aberrant mitosis with consequent ploidy anomalies, known as the tetraploidy checkpoint (Fujiiwara et al., 2005; Ganem et al., 2014; Lanni and Jacks, 1998; Liu et al., 2004). Given the lack of cell cycle phenotype with p53 inactivation in E1A;HrasG12V MEFs, we sought to determine whether p53 deficiency still promotes genome destabilization. To this end, we first performed metaphase spreads on cells of each genotype grown in 5% O2 and quantitated chromosome number (Fig. 2, A–C). Cells targeted with sgp53 exhibited a significantly higher proportion of cells with >40 chromosomes than sgNTC controls, including both tetraploid and polyploid cells (Fig. 2, A–C). To confirm this finding in cells undergoing cell division (not metaphase arrested), we used the FUCCI (fluorescent ubiquitination-based cell cycle indicator)
polyploid cells in G1 (Fig. 2 D). We observed a significantly higher percentage of both G1 tetraploid and polyploid cells in G2/M phase (Sakaue-Sawano et al., 2008). Together with cell cycle marker system, which differentially marks cells in G1 and G2/M phase (Sakaue-Sawano et al., 2008), DNA content analysis, the proportion of normal diploid cells in G1 tetraploid and polyploid populations in E1A;HrasG12V MEFS can be distinguished from abnormal tetraploid cells and DNA content analysis, the proportion of normal diploid cells in G1 tetraploid and polyploid populations in E1A;HrasG12V MEFS. Scale bar, 25 μm.

Figure 2. p53 promotes genomic stability in E1A;HrasG12V;H11Cas9 MEFS. (A) Representative metaphase spreads from sgNTC and sgp53(2) E1A;HrasG12V MEFS. White arrows indicate BrdU-positive multinucleated cells. Scale bar, 25 μm. (B) Quantitation of metaphase spreads from sgNTC and sgp53(2) E1A;HrasG12V MEFS. (C) Quantitation of chromosomes per cell from metaphase spreads. n = 3 cell lines/sgRNA. >30 cells analyzed per cell line. Data are presented as mean ± SD; *, P < 0.05, nested t test. (D) Generation of Fucci cell cycle marker–expressing variants of E1A;HrasG12V MEFS. (E) Quantitation of G1 tetraploid and polyploid populations in sgRNA-targeted E1A;HrasG12V MEFS. Data represent the mean ± SD of n = 5 cell lines/sgRNA. *, P < 0.05, two-way ANOVA, Dunnett’s multiple comparison test. (F) Generation of Fucci cell cycle marker–expressing variants of E1A;HrasG12V MEFS. (G) Quantitation of normal and abnormal mitotic events in E1A;HrasG12V MEFS. Data are mean ± SD of n = 3 cell lines/sgRNA. >50 mitotic events analyzed per cell line. Data are presented as mean ± SD; *, P < 0.05, Mann–Whitney unpaired t test. (H) Quantitation of abnormal mitotic events in E1A;HrasG12V MEFS. Data are mean ± SD of n = 3 cell lines/sgRNA. >50 mitotic events analyzed per cell line. Data are presented as mean ± SD; *, P < 0.05, one-way ANOVA, Bonferroni’s multiple comparison posttest. (I) Representative image of a multinucleated sgp53(2)-targeted E1A;HrasG12V MEF. WGA marks cell membrane, and DAPI marks nuclei. Scale bar, 25 μm. (J) Representative BrdU proliferation analysis of multinucleated sgNTC and sgp53(2) E1A;HrasG12V MEFS. White lines indicate outlines of multinucleated cells. White arrows indicate BrdU+ multinucleated cells. Scale bar, 25 μm. (K) Quantitation of BrdU-positive multinucleated sgRNA-targeted E1A;HrasG12V MEFS. Data are mean ± SD of n = 3 cell lines/sgRNA. *, P < 0.05, two-way ANOVA, Dunnett’s multiple comparison test. For A–K, all experiments were performed in physiological (5%) oxygen.

Next, to investigate the types of abnormal mitotic events in sgp53-targeted cells that might drive altered ploidy, we performed time-lapse, live-cell imaging analysis of mitosis. E1A;HrasG12V sgp53-targeted MEFS exhibited an increased proportion of abnormal mitotic events (defined here as multipolar spindle formation or failed cytokinesis) relative to sgNTC cells (Video 1 and Fig. 2, F and G). One outcome arising from these mitotic defects is the generation of cells with more than one nucleus (Fig. 2 H). Specifically, while any cell with more than one nucleus in sgNTC MEFS tended to be only binucleate, p53-deficient cells exhibited both binucleate and multinucleated cells, with a small proportion of these multinucleated cells displaying >10 apparent nuclei (Fig. 2 I). Interestingly, after pulsing with BrdU, ~70% of these p53-deficient bi- and multinucleate cells displayed BrdU positivity, compared with only ~40% of the binucleate cells in sgNTC cells (Fig. 2 J and K). This finding suggests that in the absence of p53, these highly abnormal cells are more likely to attempt to continually divide, supporting the idea of a failed tetraploidy checkpoint in these cells. Together, these results demonstrate that, despite being unable to dampen proliferation, p53 remains capable of regulating mitotic fidelity in E1A;HrasG12V MEFS, perhaps by direct regulation of mitotic events and/or clearance of cells that have undergone abnormal mitosis. These studies thus underscore the importance of p53 in faithful genome propagation.
p53 promotes DNA repair in response to acute DNA damage in E1A;Hras<sup>GL2V</sup> MEFs

Beyond maintaining genomic integrity by regulating ploidy, p53 has also been shown to promote different types of DNA repair (Williams and Schumacher, 2016). Interestingly, the capacity of p53 to regulate DNA repair has recently been suggested to be critical for p53-mediated tumor suppression in lymphoid cells (Janic et al., 2018; Valente et al., 2013). To determine whether p53 function in E1A;Hras<sup>GL2V</sup> MEFs is also linked to the repair of damaged DNA, we harvested WT and p53-deficient E1A;Hras<sup>GL2V</sup> MEFs at various time points after treatment with 2 Gy γ-irradiation (IR) and immunostained cells with γH2AX antibodies to detect foci that form at sites of double-strand breaks (Fig. 3 A). We found no significant difference in the percentages of cells exhibiting high nuclear γH2AX fluorescence in either the basal state or the early response to acute DNA damage (1 h after IR) in cells with different p53 statuses. In contrast, sgp53-targeted E1A;Hras<sup>GL2V</sup> MEFs exhibited significantly higher γH2AX staining 6 h after IR than sgNTC cells, suggesting delayed induction of DNA repair (Fig. 3, B and C). Both sgNTC- and sgp53-targeted cells showed little evidence of cell death 6 h after IR (not depicted), suggesting that differences in γH2AX levels did not arise from differences in apoptosis after irradiation. By 24 h after IR, γH2AX levels in both sgNTC- and sgp53-targeted MEFs returned to baseline levels (Fig. 3, B and C), indicating that factors other than p53 may contribute to late DNA repair responses. These results indicate that p53 plays a critical role in enhancing early DNA repair responses to acute DNA damage in E1A;Hras<sup>GL2V</sup> MEFs.

p53 loss does not significantly impact induction of ferroptosis in E1A;Hras<sup>GL2V</sup> MEFs

Recently, the ability of p53 to promote ferroptosis, an iron-dependent form of cell death that culminates in toxic lipid peroxidation, has been suggested to be critical for p53-mediated tumor suppression (Jiang et al., 2015b). Ferroptosis can be induced by inhibiting cystine import via the cystine/glutamate antiporter system x<sup>−</sup> using the small molecule erastin2 and can be blocked by small-molecule inhibitors of lipid peroxidation, such as ferrostatin-1 (Dixon et al., 2012). We examined whether p53 deficiency in E1A;Hras<sup>GL2V</sup> MEFs altered sensitivity to erastin2-induced ferroptosis. Initially, we performed FACS-based cell viability assays in 5% O<sub>2</sub>, to determine the sensitivity of these cells to erastin2-induced ferroptosis. After 16 h of treatment, sgNTC E1A;Hras<sup>GL2V</sup> MEFs exhibited decreased viability relative to sgp53-targeted cells (Fig. 4 A). However, while cotreatment of cells with ferrostatin-1 reduced erastin2-induced lipid peroxidation, it failed to rescue erastin2-induced cell death in sgNTC cells (Fig. 4 A). These findings suggest that erastin2 induces nonferroptotic cell death in E1A;Hras<sup>GL2V</sup> MEFs in 5% O<sub>2</sub>.

To further probe the role of p53 in ferroptosis, we shifted our studies to atmospheric (21%) O<sub>2</sub>, under which erastin2-induced ferroptosis has been well characterized. We monitored ferroptosis in erastin2-treated E1A;Hras<sup>GL2V</sup> MEFs expressing the nuclear-localized mKate2 live-cell marker and incubated with the dead cell marker SYTOX Green (Fig. 4 B). Erastin2 induced robust cell death in a dose-dependent manner in both sgNTC- and sgp53-targeted E1A;Hras<sup>GL2V</sup> MEFs (Fig. 4, C and D), and cell death was rescued by cotreatment with ferrostatin-1, consistent with the induction of ferroptosis (Fig. 4 E). Upon comparing the dose–response curves over time (Fig. 4 F), and EC<sub>50</sub> values (Fig. 4 G), no significant difference was observed between sgp53 and sgNTC cells, suggesting that p53 does not regulate sensitivity to erastin2-induced ferroptosis in E1A;Hras<sup>GL2V</sup> MEFs.

To probe why E1A;Hras<sup>GL2V</sup> MEFs grown in 5% O<sub>2</sub> displayed different sensitivity to erastin2-induced ferroptosis from that of cells cultured in 21% O<sub>2</sub>, we analyzed basal lipid peroxidation. Interestingly, both p53-expressing and p53-deficient cells in 5% O<sub>2</sub> exhibited less basal lipid peroxidation than in 21% O<sub>2</sub> (Fig. 4 H). Thus, lower basal peroxidation in 5% O<sub>2</sub> may present a barrier for the induction of ferroptosis, leading cells to engage another cell death pathway upon cystine deprivation, a notion consistent with erastin2 triggering apoptosis in some conditions where ferroptosis is blocked (Huang et al., 2018a; Huo et al., 2016). These findings also underscore the importance of examining cellular phenotypes in physiological oxygen conditions.

p53 restrains glycolysis and regulates nucleotide metabolism in E1A;Hras<sup>GL2V</sup> MEFs

The capacity of p53 to inhibit the Warburg effect, by dampening glycolysis and promoting oxidative phosphorylation, has been proposed as a mechanism by which p53 suppresses cellular transformation (Vousden and Ryan, 2009). Most studies investigating the role of p53 in metabolism have been performed in atmospheric 21% O<sub>2</sub>, whereas oxygen availability in vivo ranges between 2% and 8% O<sub>2</sub> for most cells. Given our findings highlighting the importance of oxygen tension for the induction of ferroptosis, we sought to determine the role of p53 in metabolic regulation in both atmospheric and physiological oxygen tensions.

To examine the role of p53 in glucose metabolism in E1A;Hras<sup>GL2V</sup> MEFs, we performed [U-13C]<sub>6</sub>-glucose tracing experiments in either 5% or 21% O<sub>2</sub> followed by liquid chromatography (LC)/mass spectrometry (MS) to determine labeled metabolite levels (Fig. 5 A). In 5% O<sub>2</sub>, sgNTC E1A;Hras<sup>GL2V</sup> MEFs displayed enhanced glycolysis and decreased TCA cycle activity relative to sgNTC cells grown in 21% O<sub>2</sub>, with significantly higher abundance of total and 13<sup>C</sup>-labeled lactate (Fig. 5 B), decreased percentage of (m+2) ion abundance for several TCA intermediates like fumarate and malate (Fig. 5 C), and lower (m+2) citrate: sgNTC [5%] versus sgNTC [21%]; P < 0.05, indicating a relative suppression of pyruvate oxidation at 5% O<sub>2</sub>. Notably, p53 was able to enhance pyruvate oxidation to produce citrate in 5% O<sub>2</sub>, evidenced by a significantly higher (m+2) citrate:(m+3) lactate ratio in sgNTC cells than in sgp53(2) cells, whereas p53-dependent differences in 21% O<sub>2</sub> were not significant (Fig. 5, D and E). Together, these findings suggest that p53 demonstrates a more pronounced regulatory role in pyruvate oxidation and tumor cell metabolism in physiological oxygen and highlights the importance of using appropriate oxygen tensions during cell culture in vitro to accurately model in vivo cellular function.
To gain a more comprehensive view of how p53 modulates metabolism in 5% O₂, we performed targeted LC-MS/MS metabolomics analysis on sgNTC- and sgp53-targeted E1A; HrasG12V MEFs (Fig. 5 F) and identified 15 metabolites with significantly different abundances between genotypes (Fig. 5 Gand Table S1). Metabolite set enrichment analysis of metabolites with abundance changes of at least twofold upon p53 loss indicated a strong nucleotide signature, involving both purine- and pyrimidine-related metabolites (Fig. 5 G and H). Notably, in addition to decreased nucleoside/tide levels, sgp53-targeted cells exhibited significant increases in levels of glutamine and glycine, both of which are precursors for nucleotide metabolism but are also involved in other metabolic pathways such as serine and one-carbon metabolism (Fig. 5 G). Collectively, these results confirm that p53 restrains alterations in glucose metabolism typical of the Warburg effect, primarily through a rebalancing of glycolytic and oxidative metabolism. Moreover, p53 influences other aspects of metabolism, such as nucleotide availability, in E1A; HrasG12V MEFs.

### p53 restricts migration and invasion in E1A; HrasG12V MEFs

Although roles for p53 in inducing apoptosis, maintaining genomic stability, and regulating metabolism may be central for restraining initial tumor growth, p53 may also modulate tumor progression. The ability of WT p53 to modulate behaviors paramount for metastasis, including migration and invasion, has been shown only in limited contexts (Gadea et al., 2007; Muller et al., 2011). Interestingly, in live-cell imaging experiments of E1A; HrasG12V MEFs grown in 5% O₂ (Fig. 2 F), we observed that sgp53-targeted cells exhibit more pronounced lamellipodia (membrane protrusions at the leading edges of motile cells thought to be critical for driving cellular migration) than sgNTC cells (Video 2; Krause and Gautreau, 2014). Hence, we analyzed cellular migration by performing Boyden chamber Transwell...
assays on sgNTC- and sgp53-targeted cells grown in 5% O2. Although both sgNTC- and sgp53-targeted E1A;HRasG12V MEFs migrated through the Transwell filter, sgp53-targeted cells exhibited approximately threefold greater migration than sgNTC cells over 24 h (Fig. 6, A and B). Notably, these experiments were performed without any growth factor or nutrient gradient, suggesting that even without a specific cue, WT p53 restrained cellular migration in this context. In contrast, while a similar trend was observed in cells cultured in 21% O2, differences between sgNTC- and sgp53-targeted cell migration were not significant (see Fig. S2 A, no nutrient gradient), suggesting that the capacity of p53 to regulate migration is dampened under atmospheric oxygen conditions.

Next, we sought to determine whether WT p53 also modulates invasiveness in this model system. We performed 3D-collagen matrix invasion assays on sgNTC- and sgp53-targeted cells grown in 5% and 21% O2. In 5% O2, both sgNTC- and sgp53-targeted cells formed small colonies within the collagen matrix. However, while sgNTC cells remained as small spheroids, sgp53-targeted cells penetrated into the surrounding collagen matrix, forming star-like, invasive structures (Fig. 6 C). Quantitation of invading colonies indicated there was a dramatic increase in the invasive capacity of sgp53-targeted cells (∼70% invasive colonies) relative to sgNTC cells (<5% invasive colonies; Fig. 6 D). Together, these results demonstrate that WT p53 plays a critical role in dampening both the migratory and invasive potential of
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E1A;Hras<sup>G12V</sup> MEFs in physiological oxygen conditions. In striking contrast, p53-deficient cells grown in 21% O<sub>2</sub> failed to display any invasive behavior (Fig. 6, E and F), again reinforcing the need for appropriate oxygen tensions during cell culture to accurately model p53 function in vitro. Collectively, our results demonstrate that p53 deficiency in neoplastic cells can cause pleiotropic dysregulation of a variety of cellular processes.

The effect of oxygen tension on classic p53-dependent phenotypes

Although the ability of p53 activity to regulate metabolism and invasion was markedly different in 5% and 21% O<sub>2</sub>, it remained unclear whether classic p53 pathways were similarly impacted by altering oxygen tension. We therefore performed apoptosis and proliferation assays on E1A;Hras<sup>G12V</sup> MEFs cultured in 5% or 21% O<sub>2</sub>. We found that apoptosis induced by acute DNA damage and serum deprivation was similar in magnitude and p53 dependence in both 5% and 21% O<sub>2</sub> (Fig. S2 B). In contrast, whereas proliferation was accelerated in physiological conditions, p53 loss had no significant impact in either oxygen tension (Fig. S2 C).

Thus, the consequences of altering oxygen tension are phenotype specific, with p53-dependent noncanonical responses appearing more affected. It also remained unclear how oxygen tension might affect tumorigenicity in vitro. To address this question, we assessed the impact of altering environmental oxygen on tumorigenic potential using soft agar assays. Although sgp53(2) cells formed significantly more colonies than sgNTC cells in both 5% and 21% O<sub>2</sub>, sgp53(2) cells grown in 21% O<sub>2</sub> formed fewer colonies than cells in 5% O<sub>2</sub>. Thus, p53-deficient E1A;Hras<sup>G12V</sup> MEFs grown in 5% O<sub>2</sub> are significantly more tumorigenic than the same cells grown in 21% O<sub>2</sub>, suggesting that studies performed under atmospheric oxygen conditions may underestimate the contribution of p53 action in transformation suppression. Together, these findings further highlight the need for use of appropriate oxygen tension during cell culture in vitro to accurately model in vivo cellular function.

p53 deficiency induces pleiotropic alterations in HCT116 human colon carcinoma cells

As our studies clearly suggested that p53 deficiency drives pleiotropic changes in cell behavior in E1A;Hras<sup>G12V</sup> MEFs, we next sought to determine if such pleiotropy is observed in other cellular contexts. We thus examined a set of canonical and
p53 induces diverse transcriptional programs under physiological oxygen conditions

Collectively, our findings demonstrate an expansive role for p53 in regulating varied cellular processes in oncogene-expressing cells under physiological oxygen conditions (Fig. 7 A). To develop a broader understanding of the cellular pathways transcriptionally regulated by p53 in this setting, we performed gene expression analysis on early-passage sgNTC- and sgp53-targeted E1A;HrasG12V MEFs grown in 5% O2 (Fig. 7 B). RNA sequencing (RNA-seq) analysis revealed that p53 elicits widespread transcriptional changes in E1A;HrasG12V MEFs, with 2,520 genes differentially expressed between sgNTC- and sgp53-targeted cells (Fig. 7, B and C; and Table S2). Given that our RNA-seq dataset was generated in 5% O2, while most studies on p53 have been performed at 21% O2, we first sought to determine how well our dataset aligned with the classic p53 transcriptional program (Fig. 7 D). We indeed observed a strong p53 dependence for classic p53 target genes, such as Cdkn1a, Bax, and Mdm2.

Next, we sought to determine if our dataset could provide new insight into how p53 directly regulates the phenotypes observed in E1A;HrasG12V MEFs. By overlapping this RNA-seq dataset with a p53 chromatin immunoprecipitation sequencing (ChIP-seq) dataset we previously generated in MEFs (Kenzelmann Broz et al., 2013), we identified 569 p53-dependent genes that are also bound by p53. Of these likely direct targets, 226 are induced or repressed by p53 >1.5-fold (Fig. 7 E and Table S2). Literature analysis of these 226 genes uncovered strong p53-dependent transcriptional signatures relating to p53-dependent phenotypes in E1A;HrasG12V MEFs, including apoptosis, genomic fidelity, metabolism, and migration (Fig. 7 F). Some categories, such as apoptosis, largely comprised classic p53 target genes (Fig. 7 F), suggesting that these pathways are well characterized. For genome fidelity, we identified genes whose characterized functions relate to processes that p53 is thought only to impact indirectly, such as mitosis (e.g., Pocl, Mapre3, and Dyrk3; Fig. 7 G; Ban et al., 2009; Jang and Fang, 2011; Rai et al., 2018). For other processes, such as metabolism and migration/invasion, we identified various genes that might contribute to
The phenotypes we observed but not previously annotated as direct p53 targets (Fig. 7, H and I; Fischer, 2019). The EIA;HrasG12V MEF platform can therefore serve as a tool to identify novel p53-regulated genes in diverse p53-dependent processes.

p53 regulates actin dynamics through RhoD

We next sought to determine whether our platform could reveal underappreciated aspects of p53 biology using functional annotation of the 569 p53-bound and regulated genes. Using Enrichr, we identified canonical p53 signatures, as well as noncanonical signatures, such as chromatin/nucleosome remodeling and endosome formation (Fig. 8 A; Chen et al., 2013; Kuleshov et al., 2016). Of relevance to our findings demonstrating a role for p53 in inhibiting cell migration and invasion, we observed numerous signatures relating to actin dynamics, stress fibers, and the cytoskeleton, and identified a list of 57 p53-bound and regulated genes implicated in actin-related processes (Fig. 8, A and B). To determine whether these signatures reflected a clear functional outcome in these cells, we examined F-actin in sgNTC- and sgps3(1)-targeted EIA;HrasG12V MEFs. We observed gross alterations in F-actin structure, with a significantly increased percentage of cells with more than five stress fibers per cell in sgps3(1)-targeted cells (40–50%) than in sgNTC cells (<5%; Fig. 8 C and D). Notably, no significant difference was observed in the overall level of F-actin fluorescence between sgNTC- and sgps3(1)-targeted EIA;HrasG12V MEFs (Fig. 8 E). Instead, functional p53 is associated with reduced stress fibers.

To identify p53 target genes that might be involved in regulating this stress fiber phenotype, we inspected the list of actin-related genes (Fig. 8 B). On this list, we identified three members of the Rho GTPase family known to modulate stress fiber formation (Fig. 8 A; Chen et al., 2013; Kuleshov et al., 2016). Of relevance to our findings demonstrating a role for p53 in inhibiting cell migration and invasion, we observed numerous signatures relating to actin dynamics, stress fibers, and the cytoskeleton, and identified a list of 57 p53-bound and regulated genes implicated in actin-related processes (Fig. 8, A and B). To determine whether these signatures reflected a clear functional outcome in these cells, we examined F-actin in sgNTC- and sgps3(1)-targeted EIA;HrasG12V MEFs. We observed gross alterations in F-actin structure, with a significantly increased percentage of cells with more than five stress fibers per cell in sgps3(1)-targeted cells (40–50%) than in sgNTC cells (<5%; Fig. 8 C and D). Notably, no significant difference was observed in the overall level of F-actin fluorescence between sgNTC- and sgps3(1)-targeted EIA;HrasG12V MEFs (Fig. 8 E). Instead, functional p53 is associated with reduced stress fibers.

To identify p53 target genes that might be involved in regulating this stress fiber phenotype, we inspected the list of actin-related genes (Fig. 8 B). On this list, we identified three members of the Rho GTPase family known to modulate stress fiber formation: RhoD, RhoV (up-regulated), and RhoE (down-regulated). To hone in on the most relevant family member, we performed meta-analysis of five mouse and five human published RNA-seq and ChIP-seq datasets and observed that RhoD was the Rho GTPase most consistently bound and regulated by p53 across cell types and species (Fig. S4). Inspection of mouse and human p53 ChIP-seq datasets from doxorubicin-treated MEFs and human keratinocytes (Kenzelmann Broz et al., 2013; McDade et al., 2014) revealed p53-binding peaks containing p53
consensus sites within 10 kb of both the mouse and human RhoD loci (mouse, in the promoter; human, within intron 1; Fig. 9, A and B). Moreover, we validated p53-dependent expression of RhoD in E1A;HrasGLV MEFS by qRT-PCR analysis (Fig. 9 C). Together, these results suggest that RhoD is a bona fide p53 target gene in both human and mouse cells. To determine whether p53-mediated expression of RhoD in E1A;HrasGLV MEFS can indeed inhibit stress fiber formation, we overexpressed RhoD in p53-deficient E1A;HrasGLV MEFS. Overexpression of FLAG-hRHOD significantly decreased the number of stress fibers relative to cells overexpressing HA-GFP (Fig. 9, D and E). These studies thus highlight the regulation of actin dynamics, specifically inhibition of stress fiber formation mediated by RhoD, as an underappreciated aspect of p53 function in cells undergoing cellular transformation.

Collectively, our results suggest that p53 governs a complex network of cellular processes in E1A;HrasGLV MEFS, with p53 loss impacting apoptosis, DNA repair, genomic stability, multiple aspects of metabolism, migration, invasion, and actin dynamics (Fig. 9 F). The ability of p53 to coordinate these processes may therefore be integral to its capacity to suppress oncogenic transformation.

**Discussion**

Here, to gain insight into global p53 function, we examined the ability of p53 to govern a host of cellular processes during transformation suppression. Previous studies characterizing p53 cellular functions have typically described one particular cellular function for p53 in a given context, without a comprehensive analysis to determine whether p53 can act broadly to modulate numerous cellular behaviors in that particular setting. Thus, it remained unclear whether each cellular function that p53 regulates is relevant in a specific context or whether p53 might regulate a range of cellular processes in a particular tumor suppression setting. To test the latter model, we used oncogene-expressing MEFS, expressing or lacking p53, as an in vitro platform to perform a systematic analysis of the impact of p53 loss. Importantly, by using this primary cell-based system, we...
mitigated any effects of accrued mutations typical of cancer cell lines that might confound analyses, and we helped to ensure that p53 signaling pathways remain intact. Consistent with previous studies, we found that p53 loss enhanced anchorage-independent cell growth and protected cells from apoptosis induced by different stressors, validating this model as a tractable system to investigate p53 functions associated with transformation suppression. Beyond these established phenotypes, we now reveal dramatic pleiotropy with p53 deficiency, with widespread alterations in cellular behavior, supporting the idea that in tumor suppression, p53 acts through coordinate regulation of many processes.

An important facet of our study was the analysis of phenotypes at physiological oxygen. With the exception of p53-driven senescence, which is rescued in physiological oxygen (Parrinello et al., 2003), p53 cellular responses in vitro in physiological oxygen have not been well described. Remarkably, some dramatic phenotypes were revealed specifically in low oxygen conditions, including migration and invasion phenotypes observed with p53 deficiency. Thus, by cataloging phenotypes in 5% O2, which more closely models in vivo conditions, we observe that the contribution of p53 to tumor suppression may be more complex than previously thought.

Among the cellular phenotypes we examined were those described in recent studies implicating specific noncanonical p53 functions in tumor suppression. For example, ferroptosis was proposed as a central mechanism in p53-mediated tumor suppression (Jiang et al., 2015b; Li et al., 2012b). This notion is, however, controversial, with studies showing that p53 is capable of both potentiating and suppressing ferroptosis (Stockwell et al., 2017). Interestingly, in our model, p53 expression did not affect the induction of ferroptosis at 21% O2. Our observations therefore reinforce the idea that the role of p53 in ferroptosis is cell type specific and suggest further that ferroptosis does not universally explain p53 tumor suppression. Similarly, the capacity to promote DNA repair was proposed to be a critical mechanism by which p53 suppresses tumorigenesis (Janic et al., 2018; Valente et al., 2013). Supporting the idea that p53 promotes DNA repair, we observed delayed DNA double-strand break repair in response to γ-IR in p53-deficient cells. While not necessarily affecting cellular proliferation or survival per se, defective DNA repair upon p53 loss could fuel malignancy by facilitating the accrual of additional mutations during tumor progression.

Beyond contributing to DNA double-strand break repair in our system, p53 exhibits an important role in maintaining chromosomal stability, as we observed significant increases in the numbers of tetraploid, polyploid, and aneuploid cells upon p53 inactivation. The capacity of p53 to regulate mitotic fidelity is thought to largely rely on p21-mediated cell cycle arrest (Kummer et al., 2013; Lanni and Jacks, 1998). However, we observed no clear effect of p53 loss on proliferation in our system.
presumably because expression of E1A blocks Rb-mediated arrest, the pathway through which p21 acts (Narita et al., 2003). Our findings therefore suggest that the regulation of polyplody in E1A;HrasG12V MEFs occurs through Rb- and cell cycle arrest-independent mechanisms. Intriguingly, our RNA-seq analysis identified several highly up-regulated genes whose encoded proteins have reported roles in mitosis, suggesting that p53 may influence mitotic events directly in these cells. Alternatively, p53 might regulate ploidy through elimination of polyploid cells by apoptosis. Indeed, the Hippo pathway component LATS2 promotes cell death in polyploid E1A- and HrasG12V-expressing human fibroblasts by enhancing p53 activity (Aylon et al., 2010; Ganem et al., 2014). Together, these results highlight an important cell cycle arrest-independent contribution of p53 to the maintenance of genomic fidelity during transformation suppression, another means of impeding malignant progression.

Our study has revealed p53-regulated metabolic processes that may contribute to limiting malignancy under physiological conditions. Previous reports in 21% O2 have shown that p53 regulates glucose metabolism primarily by inhibiting the Warburg effect, a reprogramming from oxidative respiration to glycolysis characteristic of cancer cells (Voussden and Ryan, 2009). Similarly, we observed that sgp53-targeted E1A;HrasG12V MEFs in 5% O2 exhibited enhanced lactate accumulation and a decreased propensity to convert pyruvate to citrate relative to sgNTC cells. Notably, we observed marked differences in glucose metabolism between 5% and 21% O2 tensions, with lactate production from glucose being highly favored under physiological oxygen conditions and citrate production favored in atmospheric oxygen. Moreover, the capacity of p53 to inhibit metabolic changes associated with the Warburg effect was greatly dampened in 21% O2. Thus, previous studies in 21% O2 may have underestimated p53’s contribution to inhibition of the Warburg effect, again highlighting the value of using physiological oxygen tensions during cell culture to accurately model in vivo cell behavior.

In addition to glucose metabolism, p53 regulates other metabolic pathways such as fatty acid oxidation and serine and one-carbon metabolism (Jiang et al., 2018a; Voussden and Ryan, 2009). Metabolomics analysis on E1A;HrasG12V MEFs in 5% O2 revealed that pyrimidine and purine metabolism were severely impacted by p53 loss, supporting several studies documenting altered nucleotide levels upon p53 loss (Huang et al., 2018b; Maddocks et al., 2013). The decreased pyrimidine and purine levels we observe with p53 loss may reflect impaired nucleotide synthesis, defective salvage and uptake pathways, or enhanced utilization of nucleotides—an observation consistent with the need to synthesize an entire genome’s worth of extra DNA in tetraploid cells. Together, our results underscore the complexity of p53’s regulation of metabolism, with p53 rewiring multiple aspects of glucose and nucleotide metabolism in E1A;HrasG12V MEFs.

Although dysregulation of the aforementioned cell processes might contribute to the growth of tumors, we also identified functions regulated by p53 that could be more relevant for metastatic spread. Mutation of TP53 in human cancers has been correlated with enhanced aggressiveness and metastatic capacity, but the role of p53 in modulating cell migration and invasion has more consistently been linked to p53 gain-of-function rather loss-of-function (Muller et al., 2011). However, some reports suggest a role for WT p53 in dampening both cell migration and invasion, through multiple mechanisms, including inhibition of podosome and filopodia formation and regulation of cell spreading (Kawauchi and Wolf, 2014; Muller et al., 2011). Of note, in our RNA-seq and ChIP-seq analyses, where actin-related functions were among the top molecular signatures, the previously described p53 target genes Rad and Rap2B (Di et al., 2015; Hsiao et al., 2011) encoding actin regulators were either absent or only mildly enriched. Instead, we identified the RhoGTPase, RhoD, as a key component of p53-dependent regulation of the actin cytoskeleton, with p53-dependent induction conserved between mouse and human cells and RhoD overexpression potently suppressing stress fiber formation in p53-deficient E1A;HrasG12V MEFs. Regulation of different types of actin filament assemblies is critical for multiple cell processes, such as mitosis and cytokinesis, establishing polarity, and intracellular trafficking (Pollard and Cooper, 2009; Tojkander et al., 2012). Altered expression of actin-binding proteins, including those regulating stress fiber formation, has been associated with enhanced proliferation, invasion, and metastasis (Liang et al., 2011; Stevenson et al., 2012; Tavares et al., 2017). Transcriptional regulation of RhoD and genes encoding other actin-binding proteins by p53, as identified in our RNA-seq analyses, could thus form a critical node in tumor suppression, with the complex interplay between these processes promoting an antitumor program.

It will be important in the future to determine how broadly p53 deficiency drives global phenotypic changes in different cellular contexts. As a proof-of-concept, we show that HCT116 human colon carcinoma cells exhibit dysregulation of multiple cellular processes upon p53 loss, including apoptosis, maintenance of ploidy and glucose metabolism. However, the spectrum of cell processes impacted by p53 deficiency was substantially dampened in HCT116 cells, suggesting selection against p53 signaling during long-term culture. These observations therefore reinforce the value of using a primary cell-based transformation model system, such as E1A;HrasG12V MEFs, where the immediate effects of p53 deficiency can be directly assessed. Our HCT116 studies nonetheless support the notion that pleiotropy in p53 function is conserved between species and differentiation states.

Collectively, our findings provide clear evidence that, beyond merely regulating cellular expansion, in this case through apoptosis, p53 can simultaneously regulate numerous other processes, including genome stabilization, DNA repair, metabolism, migration, invasion, and actin dynamics. These observations suggest that during tumor suppression, p53 may govern myriad processes to maintain tissue homeostasis. It will be important to determine whether cooperative regulation of all or only some of these cellular processes is essential for p53 to block tumorigenesis. Such concerted effects may explain why p53, rather than its target genes, is the most frequently mutated gene in human cancer: a single missense mutation in p53 can trigger an array of pro-tumorigenic changes in nascent cancer cells. It is
interesting to consider the interplay between defects in different processes that could amplify the negative outcomes of p53 loss. For example, loss of p53 increases ploidy and decreases nucleotide availability, which could affect DNA replication by inducing replication fork stalling, leading to increased DNA damage (Bester et al., 2011). These effects would then be further amplified due to impaired DNA repair in cells lacking p53, leading to enhanced mutational rates and promoting more aggressive disease. In addition, although our study uncovers the potential of p53 to regulate many diverse aspects of cellular behavior, the relevance of loss of these functions with p53 mutation in cancer may depend on tumor stage or microenvironmental cues. Understanding the complex spectrum of cellular activities regulated by p53 in each cell type during transformation suppression will provide insight into the cellular pathways that are best targeted in drug development for different cancer types, either alone or in a synthetic lethal approach.

Materials and methods

Mice

All animal experiments were performed in accordance with the Stanford University Administrative Panel on Laboratory Animal Care. H12Cas9 mice in which expression of Cas9 from the H11 promoter is under control of a Lox-Stop-Lox (LSL) cassette were obtained from Monte Winslow (Stanford University, Stanford, CA; Chiou et al., 2015) and crossed with CMV-Cre mice (in which expression of Cre-recombinase is under control of the ubiquitous CMV promoter; Schwenk et al., 1995) to recombine the LSL element and generate progeny constitutively expressing Cas9 under control of the H11 promoter (H12Cas9,CMV-Cre), H12Cas9/; CMV-Cre/+ mice were then crossed with WT females to generate H12Cas9 mice, and embryos were harvested at embryonic day 13.5 (E13.5) to prepare MEFs as described. Briefly, embryo body wall tissue was minced using a scalpel. Tissue was dissociated in trypsin for 30 min at 37°C, then dispersed by pipetting, before cells were plated into complete DMEM and allowed to grow in 37°C, 5% CO2, and 5% O2 (Johnson et al., 2005). Genotyping to confirm the presence of the H12Cas9 transgene and absence of the CMV-Cre transgene was performed using the following forward and reverse primers: Cas9-H11 forward, 5'-GGGCGCTCAAGTCTTGCAGAT-3'; H11 forward, 5'-TGAGCAGTTGAGCTGCTTCCGAGAT-3'; H11 reverse, 5'-CTGACGCTGGAGGCACTGTGTTCC-3'; Cas9 internal forward, 5'-CTGACGCTGCTTCC-3'; and Cre reverse, 5'-CGCACCAGTGGAGCTGCTTCC-3'.

Generation of isogenic, polyclonal p53 WT, and p53-null EIA; HrasG12V MEFs

Cells were cultured in high glucose (4.5 g/liter), sodium pyruvate (1 mM), DMEM (Gibco) containing 10% FBS (Sigma-Aldrich), and 100 µg/ml penicillin/streptomycin (Gibco), referred to herein as “complete medium” for all experiments unless otherwise stated. During cell line generation and subsequent experiments (except where noted) cells were maintained at 5% O2, 5% CO2, and 37°C. Primary MEFs that were generated from E13.5 H12Cas9 mice were transduced with EIA- and HrasG12V-expressing retroviruses (pWZL hygro-125-EIA and pBabe neo-HrasG12V, respectively). Briefly, DNA solutions containing 2 µg of VSV-Env and GAG-POL (retroviral packaging vectors) and 4 µg of plasmid vector in 500 µl of 250 mM CaCl2 were prepared. DNA was then precipitated via addition of 2× Heps-buffered saline (HBS; 274 mM NaCl, 10 mM KCl, 1.4 mM Na2HPO4, 15 mM β-glucose, and 42 mM Heps, pH 7.05) in a dropwise fashion while aerating the solution. Precipitated DNA was added dropwise to plates of 293AH cells in complete DMEM supplemented with 100 mM chloroquine. Cells were incubated at 37°C, 5% CO2, and 5% O2. After 48 h, viral supernatant from these cells was collected and filtered through a 0.4-µm filter. Viral supernatants were mixed at a 1:1 ratio and used to infect primary MEFs (three infections over 48 h). Hygromycin and neomycin selection (~72 h) was used to isolate positively transduced cells. To generate sgRNA-targeted EIA; HrasG12V cell lines, passage 2 EIA; HrasG12V cell lines were transfected with viral supernatants prepared from 293AH cells transduced with 2 µg VSV-G and delta8.2 (lentiviral packaging vectors) and 4 µg of the lentiviral sgRNA constructs (Transomics, pCLIP-sgRNA-EFS-Blast V74) expressing sgRNAs directed against p53 (sgp53(1), TEVM-11152982, and sgp53(2), TEVM-1220124; see Fig. S1 A for sgRNA sequences) or nontargeting control (sgNTC, pClip-All-EFS-Blast, CAMS1001-22059), as described above. Cells were grown for 7 d to allow for Cas9-mediated cutting of the p53 locus. Use of multiple guides directed toward p53 allows for identification and exclusion of phenotypes arising from off-target effects.

To confirm CRISPR/Cas9 targeting of p53, forward and reverse primers were designed to amplify the region flanking both sgRNAs’ targeting sites in Exon 4 (p53 forward 5’-GGACTGCAG GTTCTCAGAAG-3’; and p53 reverse 5’-CTGAAGAGGACCCCAATT-3’), Sanger sequencing was performed. Sequencing reads were analyzed using interference of CRISPR edits (ICE) algorithm (Synthego) to determine the success of targeting. All sgp53-targeted cell lines exhibited knockout scores of >80 (Fig. S1 B). One sgp53(1) targeted cell line (line 2, embryo 1,588.5) exhibited expression of a protein migrating with p53 (Fig. S1 C) but was unable to induce expression of p21 and Mdm2 (Fig. S1 D), indicating it was functionally null for p53. Moreover, the cell line did not significantly differ from the two other completely null sgp53(1) cell lines in any assay performed (Fig. S1 A–F). To confirm on a broader level that this cell line was functionally p53 null, we chose this sgp53(1) guide for our RNA-seq analyses and observed no or minimal differences from the two completely null sgp53(1) cell lines when assessing p53-target gene expression (Fig. 7 D), again confirming the functionally null status of this line.

If prioritization of a single sgRNA was required, e.g., metabolomics, glucose-tracing, and actin and oxygen tension analyses, we chose to use the sgp53(2) guide given that three of three cell lines were completely p53 null, thus providing the cleanest system to assess the impacts of p53 loss (Fig. S1 C). Early passages of targeted cells (passages 2–3) were grown in bulk and frozen for future experiments. All phenotypic analyses were performed within 3 wk of thawing cells.
(<passage 15) to limit acquisition of phenotypes that were secondary to p53 deficiency.

Physiological versus atmospheric oxygen tension analyses
Frozen vials of sgNTC and sgp53(2) E1A:HrasG12V MEFs (generated under 5% O₂ conditions) were thawed and pelleted. Cells were resuspended in medium, and each vial was split in two separate plates. Cells were then incubated in 5% or 21% O₂ incubators (5% CO₂, 37°C) and allowed to equilibrate for a minimum of 72 h (at least one passage) before phenotypic analyses.

Immunoblotting
Protein was extracted from cells using NP-40 lysis buffer (50 mM Tris, pH 8.0, 150 mM NaCl, 1% NP-40, 0.5 mM EDTA, and 10% glycerol) containing Roche cComplete protease inhibitors. Protein was quantitated using the bicinchoninic acid assay kit (Pierce). 20 µg of protein was resolved on a 10% SDS-PAGE gel, blotted onto polyvinylidene difluoride membranes (Millipore), and blocked in 5% nonfat dry milk prepared in TBS with 0.1% Tween 20 (TBST). Washes were performed in TBST, and the following primary and secondary antibodies were used: rabbit anti-p53 (CM5; Novogene, i:2,000), rabbit anti-p21 (ab188224; Abcam, 1:1,000), mouse anti-Mdm2 (ab16895; Abcam, 1:500), mouse anti-Gapdh (10R-G109A; Fitzgerald, 1:10,000), peroxidase AffiniPure goat anti-rabbit (111-035-144; Jackson Immuno Research, 1:5,000), or goat anti-mouse IgG HRP-conjugated (115-035-003; Jackson Immuno Research, 1:5,000). Immunodetection was performed using ECL Western Blotting substrate (Pierce).

Immunofluorescence
Cells were grown on coverslips for 24 h before the beginning of the experiment. Coverslips were harvested and fixed with 4% PFA, permeabilized with 0.02% Triton X-100 in PBS, and blocked in 5% BSA with the primary and secondary antibodies listed in the relevant methods sections. Coverslips were then mounted using ProLongGold antifade reagent with DAPI (P36931; Thermo Fisher Scientific). qRT-PCR (in triplicate) was performed using a 7900HT Fast Real-Time PCR machine (Applied Biosystems). Power SYBR Green master mix (Thermo Fisher Scientific) using 1:10,000 dilutions of RNA was extracted from cells using Trizol reagent (Thermo Fisher Scientific). qRT-PCR and secondary Alexa Fluor 546 goat anti-rabbit IgG (A-11035; Invitrogen, 1:250) antibodies were used.

qRT-PCR
RNA was extracted from cells using Trizol reagent (Thermo Fisher Scientific), and cDNA synthesis was performed using Moloney murine leukemia virus reverse transcription (Thermo Fisher Scientific). qRT-PCR (in triplicate) was performed using Power SYBR Green master mix (Thermo Fisher Scientific) using a 7900HT Fast Real-Time PCR machine (Applied Biosystems). The following forward and reverse primers were used: Cdk4/2 (5'-CACAGCTCAGTGATGGAA-3', 5'-ACCATGACCACCAATCGAC-3'), Bbc3/Puma (5'-GGGGGAGAAGAAGAAGGA-3', 5'-AGTCCCTAGAAGATTTAGTACATG-3'), RhoD (5'-ATTGTGTGTGGCTTGCAAGATA-3', 5'-CGACGCTGAACACTCAAGATAGG-3') and Actb (5'-TCTTACCGACATGAAATGACGAT-3', 5'-CTGCTTGCATGCCGACATCG-3'). Transcript abundance was calculated using a standard curve and normalized to Actb.

Soft agar assay
Anchorage-independent growth was assessed using soft agar assays. Briefly, 1.5 ml of phenol-free complete DMEM supplemented with 0.5% low melt agarose (BP160; Fisher) and 10 µg/ml gentamicin was aliquoted into six-well plates (in triplicate) and allowed to set at RT. Next, trypsinized, washed, and pelleted cells were resuspended in phenol-free complete DMEM supplemented with 0.3% noble agar and 10 µg/ml gentamicin at a final density of ~5,000 cells/1.5 ml. 1.5 ml of cells were then overlaid, in triplicate, onto prepared wells and allowed to set at RT before being overlaid with 1 ml complete DMEM. Cells were incubated at 37°C and 5% CO₂ and either 5% or 21% O₂ for 2 wk, and medium was refreshed once a week. Colonies were visualized by incubation with Giemsa stain (48900; Sigma-Aldrich) for 15 min at RT, then incubation overnight at 4°C. Plates were scanned using an Epson Perfection 3490/3590 scanner, and images were captured at 600-dpi resolution using EpsonScan software. To quantitate colonies, a 400 x 400-pixel square was isolated from the center of each well, and the number of colonies counted using ImageJ (National Institutes of Health).

Cell viability assays
Cells were treated with doxorubicin (0.2 µg/ml) or low-FBS DMEM (0.1% FBS) for 24 h. For 0.1% FBS experiments, cells were washed 3× with warm PBS to remove any residual FCS before addition of 0.1% FBS medium. Cells were harvested by trypsinization and stained with Annexin-V-FITC (BioLegend, 1:40) and propidium iodide (PI; Promocell, 1 µg/ml). Flow cytometric analysis on a BD LSR Fortessa flow cytometer was used to assess cell viability, with events captured using BD FACSDiva software and sample data analyzed using FlowJo 10.5.3 analysis software (FlowJo). Cell viability is expressed relative to DMSO or 10% FBS medium-treated controls.

Proliferation analysis
50,000 cells were plated in six-well plates in duplicate. At 24, 48, 72, and 96 h, duplicate wells were trypsinized, pelleted, re-suspended in 1 ml of PBS, and counted using a LUNA II Automated cell counter. The t = 24 h time point served as a starting cell count to control for any differences in seeding potential between cell lines. Fold change in cell number was calculated by dividing the average cell count at each time point (t = 48, 72, and 96 h) by the t = 24 h cell count.

Metaphase spreads
Cells were treated with 10 µg/ml KaryoMAX Colcemid Solution ( Gibco) for 2 h. Trypsinized cells were then incubated in 0.56% KCl at RT for 20 min. Cells were fixed in ice-cold 3:1 methanol:glacial acetic acid solution (5 min) and then pelleted on low speed (500 rpm) three times. Cells were dropped on glass slides from ~1-m height, dried and stained with 3% Giemsa stain (Promega), and coverslips were mounted. Metaphase spreads were captured using a Leica DM6000B bright-field microscope using a Leica 100×/1.4 oil-immersion objective. Images were captured with a Hamamatsu C11440-42U digital camera and Leica Application Suite X (LASX) software. Chromosome number was quantitated in >20 cells per sgRNA per cell line.
Analysis of ploidy using the FUCCI cell cycle marker system

To generate sgNTC- and sgps3-targeted Eia;HrasG12V MEF lines expressing the FUCCI system, 293AH cells were transfected with 2 µg retroviral packaging vectors and 4 µg hCDT1-mKO2 or hGeminin-mAG retroviral constructs (Sakauke-Sawano et al., 2008) in 500 µl Opti-MEM (Gibco) mixed with 20 µl Lipofectamine 2000 reagent (Invitrogen) in 480 µl Opti-MEM. A 1:1 hCDT1/hGEM ratio of viral supernatant was used to transduce sgNTC- and sgps3-targeted Eia;HrasG12V cells. Transduced cells were cultured for 1 wk before analysis. 100,000 cells were plated into six-well plates, and after 24 h cells were trypsinized, pelleted, resuspended in 1 ml complete DMEM containing DyeCycle Violet reagent (Genesee, 1:1,000), and incubated at 37°C for 30 min before flow cytometric analysis on a BD LSRFortessa X-20 flow cytometer using BD FACSDiva software to capture events. To quantify G1, tetraploid and polyploid, samples were analyzed using FlowJo 10.5.3 software. Live cell populations were gated on forward/side scatter scatterplots; live cells were then plotted on a mKO2/mAG scatterplot, and cells positive for either marker were gated (FUCCI+ cells). DNA content of FUCCI+ cells was then plotted on a histogram using DyeCycle Violet fluorescence, and gates were drawn around cells exhibiting 2N DNA content (>2N cells). Finally, >2N cells were plotted on an mKO2/mAG scatterplot. G1 tetraploid cells were defined as cells having >2N DNA expressing hCDT1-mKO2, and polyploid cells were defined as having >4N DNA content.

Live-cell imaging of mitosis and migration

Cells in complete DMEM minus phenol red were plated in Ibidi µ-slide eight-chamber glass-bottom culture slides precoated with poly-D-lysine (5 µg/cm2). Cells were imaged by phase µ-slide eight-chamber glass-bottom culture slides precoated with poly-D-lysine (5 µg/cm2). Cells were imaged by phase

Quantitation of DNA repair by γH2AX foci resolution assay

Cells were irradiated with 2 Gy IR using a 137Cs source. Unirradiated cells served as controls. At 1, 6, and 24 h after IR, cells were subjected to immunofluorescence analysis (see above) using antibodies against phosphorylated histone H2AX-Ser139 (JBW-301; Millipore, 1:1,000) and fluorescein goat anti-mouse IgG (F1-2000; Invitrogen, 1:250). Cells were visualized using a Leica DM6000B fluorescent microscope with a 40×/0.85 dry objective. Images were captured using a Hamamatsu C1440-42U digital camera and LASX acquisition software. BrdU+ cells were analyzed using FlowJo 10.5.3 analysis software (FlowJo). Ferrostatin-1 was obtained from Cayman Chemicals. Both drugs was synthesized by Acme Bioscience (Palo Alto, CA), and Ferroptosis analysis

Erastin2 (compound 35MEW28, reported in Dixon et al., 2014) was synthesized by Acme Bioscience (Palo Alto, CA), and Ferrostatin-1 was obtained from Cayman Chemicals. Both drugs were resuspended in DMSO and stored at −20°C before use. For ferroptosis analysis using PI at 5% O2 tension, cells were treated with 10 or 100 nM erastin2 ± 1 µM Ferrostatin-1 for 16 h. Cells were harvested by trypsinization and then stained with PI (Promocell, 1 µg/ml). Flow cytometric analysis of cell viability was performed using a BD LSR Fortessa flow cytometer with events captured using BD FACSDiva software. Sample data were analyzed using FlowJo 10.5.3 analysis software (FlowJo). Ferroptosis susceptibility at 21% O2 tension was assayed using STACK (scalable time-lapse analysis of cell death kinetics;
Lipid peroxidation flow cytometric analysis

*EIA;Hras*G12V* sgNTC- or sgp53-targeted MEFs were cultured in 5% or 21% O₂ for 72 h before harvesting by trypsinization. Cells were then stained with the BODIPY 581/591 C11 lipid peroxidation sensor (D3861 Thermo Fisher Scientific), according to manufacturer’s instructions. Metabolomics analysis

3 × 10^6 sgNTC-targeted and 2 × 10^6 sgp53(2)-targeted *EIA; Hras*G12V,HiFα/α MEFs were plated in 10-cm plates in complete DMEM. Different seeding densities were used to account for differences in seeding efficiency and cell size between sgNTC and sgp53(2) cells, ensuring that cells were analyzed when growing exponentially. Cells were analyzed when 80% confluent. At 24 h, cells were fixed and lysed by incubation in 80% methanol on dry ice. Methanol-extracted samples were then processed by the Children’s Medical Center Research Institute Metabolomics Core Facility at UT Southwestern. Targeted LC/MS/MS analysis were performed as previously described (Kim et al., 2017; Mullen et al., 2014). Relative metabolite abundances were determined by normalizing peak areas to total ion current. Unpaired Student’s t tests were used to determine any statistically significant difference in metabolite concentrations. Pathway enrichment analysis on significant hits was performed using Metaboanalyst 3.0 (Chong et al., 2018).

Transwell Boyden chamber migration assay

Boyden chamber assays were performed with complete DMEM (10% FBS) in the lower compartment of the Transwell plate and 25,000 cells in 500 µl of complete DMEM (10% FBS) in the top compartment (Corning 12-well control inserts, 8-µM pores). Cells were incubated for 24 h before nonmigrating cells were removed, and inserts were fixed in 4% PFA (15 min), washed three times with PBS, and stained with 0.1% crystal violet (Sigma-Aldrich) for 30 min. Washed (H₂O) and dried membranes were then covered with coverslips, and cells were counted using a LEITZ inverted microscope. Tumor cell migration was scored as nonmigrating (cells in the upper compartment), 1 ≤ 10 cells/membrane, 3 ≤ 100 cells/membrane, and 5 ≥ 100 cells/membrane. Mobility of MEFs with sgp53(2) mutations was compared with sgNTC MEFs, and each condition was performed in triplicate or quadruplicate. The data were plotted on PE-Texas Red versus FITC dot plots, with PE Texas Red+ as the gating control.
visualized by bright-field microscopy on a Leica DM6000B microscope using a 20×/0.8 dry objective. Images were captured using a Hamamatsu C11440-42U digital camera and LASX acquisition software. Five images in random positions were captured per insert (excluding the regions closest to the insert edge). Cells per 20× objective field were then quantified.

Collagen 3D matrix invasion assay
A single-cell suspension of 5,000 cells in 100 µl DMEM with 20% FBS was mixed with 100 µl rat tail collagen I (Corning) on ice. 50 µl cell/collagen suspension was then plated into the wells of a 96-well plate in triplicate. Collagen was allowed to polymerize for 30 min at RT before plates were incubated at 37°C, 5% CO₂, and 5% O₂ for 1 h. Collagen plugs were overlaid with 100 µl complete DMEM and incubated for 5 d. Excess media was removed, and collagen plugs were fixed in 4% PFA for 1 h before plugs were permeabilized in 0.02% Triton X-100 in PBS for 30 min. Plugs were washed with PBS three times before being stained with WGA and DAPI (BioLegend, 1:1,000) in PBS overnight at 4°C. Collagen plugs were washed three times in PBS before being plated onto glass slides on which paraffin wax was used to create a hydration barrier, and excess PBS was then added to maintain hydration of the collagen plug during imaging. Collagen plugs were overlaid with a coverslip and sealed with nail polish. Quantitation of invading/non-invading colonies was performed by manually scanning the entire collagen plug by eye using a Leica DMi8 inverted fluorescence microscope, set on a 40×/0.85 dry objective. To create representative images, collagen plugs were imaged on a DM6000B microscope using a 40×/0.85 dry objective, with a Z-stack of the entire colony captured. Images were captured using a Leica DFC9000 GT digital camera and processed using LASX software and the 3D-deconvolution and maximum-projection algorithms.

RNA-seq expression analysis
For RNA-seq, 10⁶ cells (three sgNTC and three sgp53(1) cell lines) were plated in complete DMEM and incubated for 24 h at 37°C, 5% CO₂, and 5% O₂. The sgp53(1) sgRNA was used for this RNA-seq analysis to determine the functionality of the protein expressed in embryonic line 2, which migrated at the same size as p53 on SDS-PAGE. Analysis focusing on canonical p53 target genes indicated that this line did not significantly differ from the other two completely null sgp53(1) lines, indicating it was functionally null. RNA-seq libraries were prepared using the Illumina TruSeq Kit (v.2), using 1 µg total RNA for each library. RNA-seq libraries were submitted for sequencing on an Illumina HiSeq 4000 at the Stanford Functional Genomics Facility. The sequence data were filtered for sequence and alignment quality according to the Stanford Functional Genomics Facility’s pipeline. Filtered FASTQ files were then submitted to Basespace’s RNA Expression analysis pipeline (v1.0.0). Briefly, the STAR aligner (Dobin et al., 2013) was used to align the reads to the mouse genome (mm10), and DESEQ2 (Love et al., 2014) was used for differential expression analysis. A volcano plot showing highly significant genes (q value ≤0.05, fold-change >1.5) was generated. The q value refers to the P value adjusted for false discovery rate. Genes with a q value of ≤0.05 were used for all subsequent analysis. All RNA-seq data are available in the GEO database, accession no. GSE136355. Previously published p53 ChIP-seq results from primary MEFs treated with 0.2 µg/ml doxorubicin were used to define p53-bound genes and were defined as those that display p53 binding within 10 kb of the gene. (Kenzelmann Broz et al., 2013), available on the GEO database, accession no. GSE46240. To identify novel p53-direct target genes, we cross-referenced our RNA-seq dataset with the TargetGeneReg database, derived from a meta-analysis of human and mouse p53 expression profiling datasets (Fischer, 2019). Heatmaps were generated using Heatmapper (Babicki et al., 2016). To determine the cell processes in which the 226 genes up- or down-regulated by p53 >1.5-fold had previously been implicated, annotations on GeneCards, the human gene database (Stelzer et al., 2016), were used to bin genes into functional categories. Categorization was then refined by literature analysis on PubMed. Pathway enrichment analysis was performed using Enrichr (Chen et al., 2013; Kuleshov et al., 2016) using the KEGG 2019 Human, KEGG 2019 Mouse, and GO Cell Component databases. Meta-analysis of RhoGTPase expression in human and mouse RNA-seq and ChIP-seq datasets was derived from the following studies: Allen et al., 2014; Kenzelmann Broz et al., 2013; Lee et al., 2010; Li et al., 2012a; McDade et al., 2014; Menendez et al., 2013; Nikulsen et al., 2012; Tanikawa et al., 2017; Tonelli et al., 2015; Wang et al., 2014.

Actin structure analysis
For actin structure analysis, cells were grown on coverslips for 24 h, fixed, and permeabilized as described above. Cells were stained with Alexa Fluor 488 Phalloidin (a kind gift from Matt Footer, Stanford University, Stanford, CA; 1:1,000) for 1 h at 37°C in a humidified chamber. Cells were visualized using a Leica DMi8 inverted microscope with a 63×/1.4 oil-immersion objective. Z-stack images were captured and processed using a Leica DFC9000GT digital camera, LASX acquisition software, and 3D-deconvolution and maximum-projection algorithms. Total cell Phalloidin fluorescence was quantified using ImageJ. Briefly, using the Alexa Fluor 488 channel, region of interest gates were drawn around individual cells, and total cell fluorescence and cell area were measured. To account for differences in cell size between sgNTC- and sgp53-targeted cells, phalloidin fluorescence values were normalized to total cell area. Stress fiber number was analyzed manually. Total cell phalloidin fluorescence and stress fiber analysis represents the combined analysis of >30 cells per cell line.

hRhoD overexpression
HA-tagged GFP (pcDNA3.1-3XHA-GFP plasmid; Brady et al., 2011) and FLAG (DYKDDDDK)-tagged hRhoD ORF clone (pcDNA3.1-C-(K)-DYK, ABIN4924381, Genomics-Online.com) overexpression constructs were transfected into sgp53(2) E1A;Hrdsqg¹HIf¹Cas9 MEFs using Lipofectamine 2000 (Invitrogen), as described above. After 24 h, cells were harvested, fixed, permeabilized, and stained with Alexa Fluor 647 Phalloidin (A22287; Thermo Fisher Scientific, 1:1,000) to mark F-actin, mouse anti-FLAG (clone M2 F1804; Sigma-Aldrich, 1:50), mouse anti-HA (12CA5;
Statistical analysis

Unless otherwise stated, all statistical analyses were performed using GraphPad Prism 8.

Online supplemental material

Fig. S1 shows sgp53 CRISPR targeting sites and validation studies on the nine cell lines used in this study. Fig. S2 shows comparative analysis of cell behavior under physiological (5%) or atmospheric (21%) oxygen conditions. Fig. S3 shows phenotypic analysis of the consequences of p53 loss in human HCT116 colon carcinoma cell lines. Fig. S4 shows metanalysis of RhoGTPase expression in mouse and human RNA-seq and ChIP-seq datasets. Video 1 shows normal and abnormal mitotic events in sgNTC- and sgp53-targeted E1A;HrasG12V MEFs grown in 5% O2. Video 2 shows 2D migration of sgNTC- and sgp53-targeted E1A;HrasG12V MEFs in 5% O2. Table S1 shows metabolomics analysis of sgNTC- and sgp53(2)-targeted E1A;HrasG12V MEFs generated in 5% O2. Table S2 shows the differentially expressed genes between sgNTC- and sgp53(i)-targeted E1A;HrasG12V MEFs grown in 5% O2.

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Figure S1. Generation and validation of isogenic, WT, and p53-null E1A;HrasG12V;H11Cas9 MEFs. (A) Trp53 exon structure and the binding sequences of the two sgRNAs used to target p53 in this study. PAM motifs are in red. Exons encoding different p53 domains are also indicated: transactivation domain (TAD), proline rich domain (PR), DNA binding domain (DBD), nuclear localization signal (NLS), and oligomerization domain (OD). (B) Summary of ICE analysis on E1A;HrasG12V;H11Cas9 MEFs used in this study. ICE score represents the percentage of sequences within the pool with non-WT sequences, whereas the KO score represents the percentage of sequences in the pool with either a frameshift or 21+-bp indel. R squared values refer to the Pearson correlation coefficient for the ICE score. (C) Western blot analysis of p53, p21, and Mdm2 after doxorubicin (dox) treatment of all nine E1A;HrasG12V MEF lines used in this study. Gapdh serves as a loading control. (D) qRT-PCR analysis of p53 target gene expression in all nine E1A;HrasG12V MEF lines used in this study. n = 3 cell lines/sgRNA (in triplicate). Data are presented as mean ± SD, after normalization to β-actin. (E) Representative images of soft agar assays of all nine sgRNA-targeted E1A;HrasG12V MEFs 24 h after DNA damage (doxorubicin treatment). n = 3 cell lines/sgRNA, three to five independent experiments. Data are presented as mean ± SD. For A–F, all experiments were performed in 5% O2.
Figure S2. The impact of physiological (5%) and atmospheric (21%) oxygen tension on cell behavior in vitro. (A) Analysis of migration in \textit{E1A;HrasG12V} MEFs in 21\% O$_2$. Data are mean fold change in migrating cells ± SD, expressed relative to counterpart sgNTC control cell line. \( n = 3 \) cell lines/sgRNA; ns, not significant (P > 0.05), Mann–Whitney unpaired t test. (B) Cell viability analysis by AnnexinV/PI staining of sgNTC and sgp53(2) \textit{E1A;HrasG12V} MEFs in 5\% or 21\% O$_2$ cotreated with 0.2 µg/ml doxorubicin or 0.1% FBS for 24 h. \( n = 3 \) independent experiments for each sgRNA. Data are presented as mean ± SD; ****, P < 0.0001; ns, not significant (P > 0.05), two-way ANOVA, Sidak’s multiple comparison posttest. (C) Proliferation of \textit{E1A;HrasG12V} MEFs in 5\% or 21\% O$_2$ measured by cell counting starting 24 h after plating. \( n = 3 \) cell lines/sgRNA. Data are mean fold change in cell number ± SD; **, P < 0.005; not significant (P > 0.05), linear regression analysis. (D) Representative soft agar assays measuring anchorage-independent growth of sgNTC and sgp53(2)-targeted \textit{E1A;HrasG12V} MEFs in 5\% or 21\% O$_2$. Scale bar, 3.5 mm. (E) Average colony number ± SD of sgNTC and sgp53(2) \textit{E1A;HrasG12V} MEFs in 5\% and 21\% O$_2$. \( n = 3 \) cell lines/sgRNA, in triplicate; *, P < 0.05; ****, P < 0.0001; ns, not significant (P > 0.05), one-way ANOVA, Tukey’s multiple comparison posttest.
Figure S3. Loss of p53 in human HCT116 colon carcinoma cells alters the behavior of multiple cellular pathways. (A) Cell viability analysis by AnnexinV/PI staining on TP53+/+ and TP53−/− HCT116 cells in 5% O2 after 0.2 μg/ml doxorubicin or 0.1% FBS treatment for 48 h. n = 3 independent experiments for each genotype. Data are presented as mean ± SD; *, P < 0.05, two-way ANOVA, Sidak’s multiple comparison posttest. 

(B) Proliferation analysis of TP53+/+ and TP53−/− HCT116 cells measured by cell counting starting 24 h after plating. n = 3 independent experiments. Data are presented as mean fold change in cell number ± SD.

(C) Ploidy analysis in TP53+/+ and TP53−/− HCT116 cells showing percentage of cells with >4N DNA content by PI staining. n = 3 independent experiments per genotype; data represent mean ± SD; *, P < 0.05, unpaired t test.

(D and E) [U13C]glucose tracing on TP53+/+ and TP53−/− HCT116 cells showing glycolytic (D) and TCA cycle (E) intermediates. Data represent mean ± SD of n = 3 cell lines per sgRNA; *, P < 0.05; **, P > 0.005; ***, P < 0.001; ****, P < 0.0001, one-way ANOVA, Tukey’s multiple comparison test.

(F) Citrate:lactate mass heavy ion ratio of TP53+/+ and TP53−/− HCT116 cells. Data represent mean ± SD of n = 2 samples per cell line; Student’s t test.

(G) Representative images of TP53+/+ and TP53−/− HCT116 cells grown in a 3D collagen matrix. WGA stains cell membranes, and DAPI marks nuclei. Scale bar, 0.2 mm.

(H) Percentage of invading colonies in 3D collagen assay. Data are mean ± SD, n = 2 independent experiments per genotype, P = 0.073, unpaired t test. All HCT116 experiments were performed in 5% O2.
Video 1. **Time-lapse live imaging of sgNTC- and sgp53-targeted E1A;HrasG12V;H11Cas9 MEFs undergoing normal and abnormal mitosis.** Normal and abnormal mitoses in sgNTC and sgp53(2)-targeted E1A;HrasG12V MEFs. 16-h phase-contrast live imaging, image capture every 5 min, frame rate = 5 frames/s.

Video 2. **Time-lapse live imaging of sgNTC- and sgp53-targeted E1A;HrasG12V;H11Cas9 MEFs migrating in 2D.** Cell migration in 2D of sgNTC and sgp53(2)-targeted E1A;HrasG12V MEFs. 16 h frames/s.

Two tables are provided online in Excel files. Table S1 shows the metabolomics dataset: sgNTC- and sgp53(2)-targeted E1A;HrasG12V MEFs grown in 5% O₂ for 24 h. Table S2 shows the RNA-seq dataset from E1A;HrasG12V MEFs in 5% O₂, annotated to show genes that had p53 ChIP peaks in the Kenzelmann Broz et al. (2013) ChIP seq dataset.