Genomic Copy Number Dictates a Gene-Independent Cell Response to CRISPR/Cas9 Targeting

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ABSTRACT

The CRISPR/Cas9 system enables genome editing and somatic cell genetic screens in mammalian cells. We performed genome-scale loss-of-function screens in 33 cancer cell lines to identify genes essential for proliferation/survival and found a strong correlation between increased gene copy number and decreased cell viability after genome editing. Within regions of copy-number gain, CRISPR/Cas9 targeting of both expressed and unexpressed genes, as well as intergenic loci, led to significantly decreased cell proliferation through induction of a G2 cell-cycle arrest. By examining single-guide RNAs that map to multiple genomic sites, we found that this cell response to CRISPR/Cas9 editing correlated strongly with the number of target loci. These observations indicate that genome targeting by CRISPR/Cas9 elicits a gene-independent antiproliferative cell response. This effect has important practical implications for the interpretation of CRISPR/Cas9 screening data and confounds the use of this technology for the identification of essential genes in amplified regions.

SIGNIFICANCE: We found that the number of CRISPR/Cas9-induced DNA breaks dictates a gene-independent antiproliferative response in cells. These observations have practical implications for using CRISPR/Cas9 to interrogate cancer gene function and illustrate that cancer cells are highly sensitive to site-specific DNA damage, which may provide a path to novel therapeutic strategies. Cancer Discov, 6(8):914–29. © 2016 AACR.

See related commentary by Sheel and Xue, p. 824.
See related article by Munoz et al., p. 900.

INTRODUCTION

Genome engineering using site-specific DNA endonucleases has operationalized functional somatic cell genetics, enabling precise perturbation of both coding and noncoding regions of the genome in cells from a range of different organisms. Zinc-finger nucleases (ZFN) and transcription activator-like effector nucleases (TALEN) are custom-designed endonucleases that enable site-specific genome editing, but their widespread application has been limited by reagent complexity and cost (1, 2). The bacterial CRISPR/Cas9 (clustered regularly interspaced short palindromic repeats/CRISPR-associated) system, which serves as an adaptive immune mechanism, has been shown to serve as a versatile and highly effective technology for genome editing (3–8). CRISPR/Cas9 applications require introduction of two fundamental components into cells: (i) the RNA-guided CRISPR-associated Cas9 nuclease derived from Streptococcus pyogenes and (ii) a single-guide RNA (sgRNA) that directs the Cas9 nuclease through complementarity with specific regions of the genome (3, 7–11).

Genome editing occurs through induction of double-stranded breaks in DNA by the Cas9 endonuclease in an sgRNA-directed sequence-specific manner. These DNA breaks can be repaired by one of two mechanisms: nonhomologous end joining (NHEJ) or homology-directed repair (HDR; refs. 3, 12). CRISPR/Cas9-mediated gene knockout results from a DNA break being repaired in an error-prone manner through NHEJ and introduction of an insertion/deletion (indel) mutation with subsequent disruption of the translational reading frame (11). Alternatively, HDR-mediated repair in the presence of an exogenously supplied nucleotide template can be utilized to generate specific point mutations or other precise sequence alterations. Furthermore, nuclease-dead versions of Cas9 (dCas9) can also be fused to transcriptional activator or repressor domains to modulate gene expression at specific sites in the genome (13–17). CRISPR/Cas9 technology has been effectively utilized in cultured cells from a myriad of organisms (12) and has also been successfully used for in vivo modeling in the mouse germline (18, 19) as well as for somatic gene editing to generate novel mouse models of cancer (20–24).

Recent studies have shown that CRISPR/Cas9 can be effectively used for loss-of-function genome-scale screening in human and mouse cells (9–11, 25–28). These approaches rely upon lentiviral delivery of the gene encoding the Cas9 nuclease and sgRNAs targeting annotated human or mouse genes. Multiple different CRISPR/Cas9 knockout screening libraries have been developed, including both single-vector (Cas9 and the sgRNA on the same vector) and dual-vector systems (9, 25, 29). Pooled CRISPR/Cas9 screening is typically performed through massively parallel introduction of sgRNAs targeting

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all genes into Cas9-expressing cells, with a single sgRNA per cell. Positive- or negative-selection proliferation screens are performed, and sgRNA enrichment or depletion is measured by next-generation sequencing (9, 10).

To date, only a limited number of genome-scale CRISPR/Cas9 knockout screens have been reported, and these screens have demonstrated a high rate of target gene validation (9–11, 25–28). Wang and colleagues recently reported an analysis of cell-essential genes using CRISPR/Cas9-mediated loss-of-function screens in four leukemia and lymphoma cell lines (28). Hart and colleagues also reported identification of core and cell line–specific essential genes in five cancer cell lines of differing lineages (25). This approach has enabled the identification of known oncogene dependencies as well as many novel essential genes and pathways in individual cancer cell lines (25, 28). In addition to knockout screens, proof-of-concept CRISPR-activator or CRISPR-inhibitor screens using dCas9 and genome-scale sgRNA libraries have also been successfully conducted (30, 31). Moreover, in vivo genome-scale screens with CRISPR/Cas9 have also been performed for cancer-relevant phenotypes (32).

To identify cancer cell vulnerabilities in a genotype- and phenotype-specific manner, we performed genome-scale loss-of-function genetic screens in 33 cancer cell lines representing a diversity of cancer types and genetic contexts of both adult and pediatric lineages (Supplementary Table S1; ref. 29). When we analyzed essential genes across the entire dataset, we unexpectedly found a robust correlation between apparent gene essentiality and genomic copy number (CN), where the number of CRISPR/Cas9-induced DNA cuts predict the cellular response to genome editing.

**RESULTS**

**High-Resolution CRISPR/Cas9 Screening in Cancer Cell Lines for Gene Dependencies**

Using the dual-vector GeCKOv2 CRISPR/Cas9 system, we performed genome-scale pooled screening in 33 cancer cell lines representing a wide diversity of adult and pediatric cancer types (Supplementary Table S1; Fig. 1A). Cancer cell lines were transduced with a lentiviral vector expressing the Cas9 nuclease under blastcidin selection. These stable cell lines were then infected in replicate (n = 3 or 4) at low multiplicity of infection (MOI < 1) with a library of 123,411 unique sgRNAs targeting 19,050 genes (6 sgRNAs per gene), 1,864 miRNAs and 1,000 nontargeting negative control sgRNAs (29). Infected cells were purified by selection with puromycin and then passaged with an average representation of 500 cells per sgRNA until an endpoint of 21 or 28 days. At the endpoint, the abundance of sgRNAs in these cells was quantitated from genomic DNA by massively parallel sequencing and compared with the abundance in the plasmid pool used for virus production to define the relative dropout or enrichment in the screen (Fig. 1A).

The log₂-normalized read counts of the 1,000 nontargeting sgRNAs show a slight enrichment in representation from the original plasmid DNA pool, indicating that on average nontargeting guides have no substantial detrimental effect on viability (Fig. 1B). As positive controls, we also compiled a list of 213 putative cell-essential genes that are part of the ribosome, proteasome or spliceosome complexes (Supplementary Table S2). In contrast to the nontargeting negative control guides, the read counts of these positive controls in late time point samples were substantially depleted compared with the initial reference pool (Fig. 1B). Replicate reproducibility after quality control for each cell line was consistently high (Fig. 1C).

We defined a CRISPR/Cas9 guide score for each sgRNA in the screen by first calculating the log₂ fold change in abundance from the screen endpoint compared with the pool of plasmid DNA, followed by subtraction of the median scores of the negative control sgRNAs (see Methods). Hence, in our dataset a guide score of zero equates to the median effect of negative control sgRNAs. Similarly, the second most depleted sgRNA for each gene was used to call a single “second-best” CRISPR/Cas9 guide score and therefore allow the representation of gene-level dependencies (33). Significant depletions of sgRNAs are denoted by negative CRISPR/Cas9 guide scores and correspond to decreased proliferation/survival after CRISPR/Cas9-mediated gene editing.

To identify genes essential for viability in each cell line across a variety of cancer contexts, we rank-ordered genes by second-best CRISPR/Cas9 guide score from most negative (most depleted) to positive (not depleted or enriched). For each cell line, we identified key vulnerabilities corresponding to both oncogenic driver lesions as well as nononcogene dependencies (Fig. 1D–F). For instance, we observed that KRAS, ESR1, and EGFR were essential genes in KRAS-mutant (Fig. 1D), estrogen receptor–positive (Fig. 1E), and EGFR-mutant cell lines (Fig. 1F), respectively. Moreover, we observed strong dependency on a number of other cancer-relevant genes and therapeutic targets in each cell line, including BRD4, MTOR, IGF1R, CCND1, and MYC (Fig. 1D–F). Thus, our approach to CRISPR/Cas9 screening yields high-quality reproducible data that enable the identification of cancer gene dependencies across many different cellular contexts.

**Genomic Copy-Number Variation Predicts the Response to CRISPR/Cas9 Genome Targeting Independent of Target Gene Expression**

Copy-number alterations (CNA) are the most common genetic alterations in human epithelial cancers (34) and lead to overexpression of driver oncogenes in cancer. To identify such driver oncogenes responsible for cancer cell proliferation and survival within regions of copy-number amplification, we mapped sgRNAs in CRISPR/Cas9 screens of each cell line to genomic coordinates and investigated the relationship of apparent gene essentiality with ABSOLUTE DNA CN data available from the Broad Institute–Novartis Cancer Cell Line Encyclopedia (CCLE; Methods; Supplementary Table S1; refs. 35, 36). We observed a striking enrichment of negative CRISPR/Cas9 guide scores for genes that reside in CN amplifications in several cancer cell lines (Fig. 2A and B; Supplementary Fig. S1A–S1C). Specifically, CRISPR/Cas9 targeting of genes that reside in amplifications conferred decreased proliferation/survival as compared with targeting genes that mapped outside of these amplifications. As expected, we found that known oncogenes, such as AKT2, MYC, or CDK4, scored as essential in cell lines that harbored...
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Figure 1. Genome-scale loss-of-function CRISPR/Cas9 screening in cancer cell lines. A, schematic of the pooled screening process. B, cumulative frequency of log$_2$-normalized read counts per million to 1,000 nontargeting sgRNA controls (red) and sgRNAs targeting 213 positive control genes (blue; Kyoto Encyclopedia of Genes and Genomes ribosome, proteasome, and spliceosome subsets; Supplementary Table S2) in both the initial DNA reference pool (dotted) and 28 days after transduction in the PANC-1 cell line (solid). C, a box plot of the Pearson correlation between replicates (y-axis) plotted for each cell line (x-axis) shows the range of replicate–replicate correlations after quality control (Methods). D–F, rank-ordered depiction of second-best CRISPR guide scores for each gene in the Panc 08.13 (D), T47D (E), and CORL105 (F) cell lines. Hallmark cancer-relevant oncogene and nononcogene dependencies are depicted in red for each cell line.
Figure 2. Genom-scale CRISPR/Cas9 screening identifies a strong correlation between CN and sensitivity to CRISPR/Cas9 genome editing. Two cell lines are shown: SU86.86 (A, C, E) and HT29 (B, D, F). A, chromosome 19q amplicon in SU86.86 and (B) chromosome 8q amplicon in HT29. Three tracks are plotted along genomic coordinates within the region defined by the red box on the chromosome schematic. Top, ABSOLUTE genomic CN from CCLE single-nucleotide polymorphism arrays, with red indicating CN gain above average ploidy and blue indicating CN loss below average ploidy; middle, CRISPR/Cas9 guide scores plotted according to the second most dependent sgRNA for each gene, with purple trend line indicating the mean CRISPR guide score for each CN segment defined from the above track; bottom, RNAi gene-dependency scores. AKT2 and MYC, known driver oncogenes at these loci, respectively, are highlighted in orange. For RNAi data, shRNAs targeting AKT2 used in Project Achilles were not effective in suppressing AKT2.

C and D, box plots of CRISPR guide scores for both expressed and not expressed genes located on (red) or off (black) the chromosome 19q amplicon in SU86.86 (C) and the chromosome 8q amplicon in HT29 (D). For the SU86.86, the amplicon represented in C red box plots ranges from 39.3–41.4 Mb on the corresponding plot in A. The number of represented genes is noted above each box plot. E and F, for each CN-defined genomic segment, median CRISPR/Cas9 guide score is plotted against CN. Each circle represents a single genomic segment of defined CN for the indicated cell line. The size of the circle corresponds to the number of sgRNAs targeting that segment. Nontargeting negative control sgRNAs are shown with a blue box plot, and known cell-essential genes (defined as positive controls) are shown as a red box plot embedded within the plot.
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We have demonstrated that there is a gene-independent antiproliferative effect of CRISPR/Cas9 targeting that occurs with even a single target locus, increases with increasing genomic CN, and is independent of the type of structural alteration that leads to increased CN. Thus, we hypothesized amplifications involving these genes. However, we also noted that sgRNAs targeting other genes in these same amplified regions appeared similarly detrimental to cell proliferation or survival (Fig. 2A and B; Supplementary Fig. S1A–S1C).

When we compared these observations with those derived from genome-scale RNA interference (RNAi) screens performed in the same cell lines (37), we failed to observe enrichment of apparently essential genes within amplifications and instead identified a small number of genes in each region of CN gain that scored as essential (Fig. 2A and B; Supplementary Fig. S1B and S1C). Moreover, we found that sensitivity to CRISPR/Cas9 targeting within amplified genomic regions was also observed for genes that failed to show significant mRNA expression (Fig. 2C and D; Supplementary Fig. S1D–S1F). These observations suggested that the observed dependency of cancer cells to CRISPR/Cas9 targeting of genes resident in amplifications was not the direct consequence of deleting the target gene.

We next sought to determine if this “CRISPR–CN relationship” also extends to lower levels of CNAs. For all 33 cancer cell lines screened, we defined genomic segments by their CN and labeled those segments with their median CRISPR/Cas9 guide score across all sgRNAs targeting within the segment (Fig. 2E and F; Supplementary Fig. S2). We found a striking correlation between CN and median CRISPR/Cas9 guide score across even low ranges of CNAs. The 1,000 “negative control” sgRNAs in the CRISPR/Cas9 library exhibited minimal effects on cell proliferation and viability, and the majority of other data points had lower CRISPR/Cas9 guide scores than the median of these negative controls (Fig. 2E and F; Supplementary Fig. S2). Strikingly, targeting a locus with an ABSOLUTE CN of 1, which corresponds to a single and F; Supplementary Fig. S2). Strikingly, targeting a locus with an ABSOLUTE CN of 1, which corresponds to a single CRISPR/Cas9-induced DNA cut, also resulted in reduced proliferation/viability in comparison with the negative controls (Fig. 2E–F; Supplementary Fig. S2), indicating that even a discrete instance of CRISPR/Cas9 genome modification significantly affects cell proliferation/viability. For each incremental increase in DNA CN, we observed a progressive decrease in CRISPR/Cas9 guide scores in nearly all of the cell lines that we screened (Fig. 2E–F; Supplementary Fig. S2). Moreover, we observed this CRISPR–CN correlation among both low-level CN gains (e.g., 1–2 extra copies) and high-level amplifications, and both focal and arm-level CNAs (Fig. 2; Supplementary Figs. S1 and S2).

Amplified Genes Rank among the Top Dependencies in Genome-Scale Negative-Selection CRISPR/Cas9 Screens

Given the profound impact of genomic CN on apparent essentiality in CRISPR/Cas9 screens as well as the high rate of CNAs in cancer cells, we reasoned that this effect could result in a high false-positive rate for the identification of essential genes. To characterize the impact of these false positives on CRISPR/Cas9 screening data, we compared the apparent essentiality of amplified genes with that of all other genes within each of the 33 cell lines. Specifically, we examined CRISPR guide scores for all genes and observed that genes residing in focal high-level amplifications consistently rank among the most highly essential genes identified for each cell line (Fig. 3A and B; Supplementary Fig. S3A–S3C).

We then performed an aggregate analysis of apparent essentiality due to amplified genes across the entire CRISPR/Cas9 dataset. For this analysis of all genes and all cell lines, we accounted for differences in Cas9 activity/efficacy across cell lines using cell line–specific Z-score normalization (see Methods). To investigate cnv gene dependencies within the dataset, we calculated composite CRISPR/Cas9 gene scores using the ATARiS algorithm, as previously described (Methods; ref. 38). We next calculated a global z-score for gene dependency values, representing the number of standard deviations from the mean of the distribution. In parallel, we performed a similar analysis of an available RNAi dataset (Fig. 3C). Thus, this analysis enables a global examination of apparent relative gene dependencies and their relationship to genomic CN amplification. Strikingly, we observed that increasingly essential genes (lower z-scores) were more likely to reside on CN amplifications in CRISPR/Cas9 data but not in RNAi data (Fig. 3C). For genes with a z-score of less than or equal to −5, 28.2% (87 of 308) of those genes reside within a CN amplification, defined as a CN ratio (ABSOLUTE/average sample ploidy) greater than two. Thus, CN amplification is a strong determinant of apparent essentiality in CRISPR/Cas9 screening data, and if not properly accounted for, this CRISPR–CN relationship will likely contribute to a higher false-positive rate for calling gene dependency. When we inspected results from another recently published study that screened five human cancer cell lines with a different CRISPR/Cas9 library (25), we found that gene CN also predicted essentiality (Supplementary Fig. S4A–S4E), thus indicating that the CRISPR–CN correlation occurs independently of the specific sgRNA library used.

The CRISPR–CN Relationship Is Observed across Multiple Different Chromosome Structural Alterations

To investigate the CRISPR–CN relationship across a spectrum of different chromosomal structural alterations, we performed whole-genome sequencing (WGS) on three cell lines harboring CN gains and amplifications and showed a strong correlation between CN and CRISPR/Cas9 guide scores (HT29, CAL120, and Panc-1). We observed the CRISPR–CN relationship in the context of several different structural amplification patterns, including near arm-level CN gain (Supplementary Fig. S5A), simple tandem duplication (Supplementary Fig. S5A), low-level copy gain from interchromosomal translocation (Supplementary Fig. S5B), and complex amplicon structure involving breakage–fusion–bridge cycles and chromothripsis (Supplementary Fig. S6). These observations suggest that the CRISPR–CN correlation occurs at both low- and high-amplitude CN changes and does not relate to specific types of chromosomal structural variation.

The Response of Cells to CRISPR/Cas9 Genome Targeting Correlates with the Total Predicted Number of DNA Cuts at Target Loci

To investigate the CRISPR–CN relationship across a spectrum of different chromosomal structural alterations, we performed whole-genome sequencing (WGS) on three cell lines harboring CN gains and amplifications and showed a strong correlation between CN and CRISPR/Cas9 guide scores (HT29, CAL120, and Panc-1). We observed the CRISPR–CN relationship in the context of several different structural amplification patterns, including near arm-level CN gain (Supplementary Fig. S5A), simple tandem duplication (Supplementary Fig. S5A), low-level copy gain from interchromosomal translocation (Supplementary Fig. S5B), and complex amplicon structure involving breakage–fusion–bridge cycles and chromothripsis (Supplementary Fig. S6). These observations suggest that the CRISPR–CN correlation occurs at both low- and high-amplitude CN changes and does not relate to specific types of chromosomal structural variation.
Figure 3. Amplified genes represent the strongest perceived dependencies in pooled CRISPR/Cas9 screening data. A and B, rank-ordered plots showing the second-best CRISPR/Cas9 guide score for each gene in the indicated cell lines. sgRNAs targeting genes within the amplicons represented in Fig. 1 are highlighted in red for SU86.86 19q amplicon (A) and HT29 8q amplicon (B). These amplicon-targeting sgRNAs are significantly enriched as apparent dependencies relative to the other sgRNAs targeting genes outside these amplicons (one-sided Kolmogorov-Smirnov test: $P = 1.04E^{-41}$, A; $P = 5.57E^{-33}$, B). C, the cumulative fraction of amplified genes at or below a given dependency score is shown for both CRISPR/Cas9 and RNAi pooled screening datasets. Amplified genes are defined as those genes with a CN ratio $>2$. Gene dependency scores are shown as global z-scores for both CRISPR/Cas9 and RNAi screening datasets, with z-scores representing standard deviations from the mean of all genes evaluated in all cell lines screened (CRISPR/Cas9, $n = 33$ cell lines; RNAi, $n = 503$ cell lines).
that this gene-independent response reflects the total number of CRISPR/Cas9-induced DNA cuts at target loci. The GeCKOv2 library contains 3,593 sgRNAs that have multiple perfect match alignments along with a protospacer-adjacent motif (PAM) sequence within the hg19 reference genome. We typically remove these sgRNAs prior to analyzing cancer cell line dependencies. However, these promiscuous sgRNAs provided an opportunity to perform a comparative analysis of the response of cells to CRISPR/Cas9 editing and the relationship to the predicted number of CRISPR/Cas9-induced DNA cuts based on either CN or number of perfect-match on- and off-target alignments (“multiple alignment analysis”). For the CN analysis, we used only sgRNAs mapping to a single genomic locus. For the multiple alignment analysis, we reintroduced these multitargeting sgRNAs and used only sgRNAs targeting nonamplified regions, thus allowing segregation of the impact of CRISPR/Cas9-induced DNA cuts due to CN or promiscuous multiple genome alignments.

We observed that sgRNAs that target multiple sites in the unamplified genome yield a strong antiproliferative effect, similar to that observed for sgRNAs targeting genomic amplifications (Fig. 4A–D). We found that the number of predicted DNA cuts correlated strongly with the observed depletion of sgRNAs, whether mediated by CN (Fig. 4A and C) or multiple alignments (Fig. 4B and D). To quantify this effect, we calculated the slope coefficient for a linear regression of CRISPR guide scores versus predicted number of cuts for both singly and multiply targeted sets of sgRNAs within each cell line. We term these coefficients the CRISPR-Cut Index (CCI) for single-targeting sgRNAs where the amount of cutting depends on copy number (CCI-CN; Fig. 4A and C) and for multiple-targeting sgRNAs where the amount of cutting depends on the number of multiple alignments (CCI-MA; Fig. 4B and D). We observed that the CCI-CN and the CCI-MA for each individual cell line are comparable, suggesting that the decreased proliferation/survival response of cells increases in the number of loci targeted by CRISPR/Cas9. Whether this is similar, whether the number of target loci is driven by CNA of a single target locus or multiple different target loci within the genome (Fig. 4E).

We further investigated whether there was a difference in the cell response to CRISPR/Cas9-induced DNA cuts targeted to different chromosomes or multiple cuts within a single chromosome. Using the multiple alignment analysis described above, we further split multiple-targeting sgRNAs into sets that either targeted multiple chromosomes (interchromosomal) or targeted sites within only a single chromosome (intrachromosomal). We observed, on average, lower guide scores for sgRNAs targeting multiple interchromosomal loci as compared with sgRNAs targeting a comparable number of intrachromosomal loci (Fig. 5A and B; Supplementary Fig. S7). Moreover, the most promiscuous sgRNAs targeting more than 10 interchromosomal loci rank among most depleted sgRNAs in pooled screening data for each cell line (Fig. 5C and D). Thus, the response of cancer cells to multiple CRISPR/Cas9-induced DNA cuts is greater when multiple loci are targeted across several chromosomes. Beyond the effects of target gene disruption, these observations further suggest that CRISPR/Cas9 gene editing also yields an anti-proliferative response that is truly gene independent.

Variation in Cell Response to CRISPR/Cas9 Targeting

Because CCI-CN and CCI-MA are correlated across cell lines, we next calculated a net index for each cell line by integrating the number of targeted sites and genomic CN to predict the total number of cuts for all sgRNAs. We observed a plateau in CRISPR guide scores beyond a certain number of cuts for each cell line, typically ranging from 10 to 50 cuts, suggesting an important limitation in the resolution of sgRNA depletion for sgRNAs targeting many genomic sites (Supplementary Fig. S8A and S8B). Informed by this observation, we fit a segmented least-squares model composed of a general linear regression below a breakpoint (estimated by the model) and a flat segment above this breakpoint. The slope coefficient of the first segment of the model is used as the net index (CCI-Total), reflecting the magnitude of the effect of cutting on CRISPR guide scores.

The CCI-Total showed considerable variability across cell lines. Although the limited sample size of this CRISPR/Cas9 screening dataset restricts the power for a full multivariate analysis of the genetic and biological influences on the CCI-Total, we found two variables that affect this index. Investigating the median CRISPR/Cas9 guide score for “positive control” cell-essential genes as a surrogate for CRISPR/Cas9 efficacy in the screens, we identified a strong correlation of this metric with the CCI-Total (Supplementary Fig. S8C), suggesting that Cas9 efficacy influences the strength of the CRISPR-CN relationship. We also identified that TP53 mutation status also correlates with the CCI-Total (Supplementary Fig. S8D). Although both TP53 mutant and wild-type cells clearly demonstrate the CRISPR-CN relationship, wild-type cells on average show a more pronounced effect, therefore suggesting that the p53 pathway may play a role in mediating the gene-independent response of cells to CRISPR/Cas9 targeting.

CRISPR/Cas9 Targeting of Amplified Regions Induces DNA Damage and a G2 Cell-Cycle Arrest

To interrogate a specific amplification example, we introduced sgRNAs targeting genes and intergenic regions inside and outside of the 19q13 amplicon in the PANC-1 pancreatic cancer cell line (Fig. 6A) and measured viable cell number in a short-term, arrayed format CellTiter-Glo luminescent assay (Fig. 6B; Supplementary Fig. S9A). We observed a significant reduction in cell proliferation for sgRNAs targeting loci inside the amplicon as compared with outside the amplicon at 6 days after expression of each sgRNA. We noted that the observed effect was equally strong for sgRNAs targeting both amplified genes and intergenic regions and was at least as potent as those sgRNAs targeting nonamplified known essential genes, such as RPL4, U2AF1, and MYC (Fig. 6B). Furthermore, we noted that CRISPR/Cas9 targeting of other loci that are not highly amplified resulted in decreased cell proliferation compared with LacZ and Luciferase targeting negative controls. In addition to interrogating sgRNAs targeting amplified regions, we also investigated the effect of two multitargeted sgRNAs on cell proliferation in this 6-day assay, including one sgRNA with multiple perfect match alignments as well as an sgRNA previously shown to target the genome at 151 different loci (Fig. 6B; ref. 39). Here, we
Figure 4. CRISPR/Cas9 sensitivity correlates with the number of predicted cuts for both guides targeting single loci and multiple loci. Data from two representative cell lines are shown (PA-TU-8902, A–B; Panc 08.13, C–D). A and C, CRISPR/Cas9 sensitivity for sgRNAs targeting only a single locus is plotted against CN of that locus. The black hash marks represent the median CRISPR guide score for all guides targeting a locus at that CN. The linear trendline is shown. B and D, CRISPR/Cas9 guide scores for sgRNAs targeting multiple loci are plotted against the predicted number of cuts for each sgRNA. Only sgRNAs targeting nonamplified regions are included, thus allowing segregation of the impact of multiple CRISPR/Cas9-induced DNA cuts due to either CN or number of target loci. The influence of the number of predicted DNA cuts on CRISPR/Cas9 guide scores was modeled for each cell line as the slope of the trend line in A–D and termed the CCI. The CCI was determined for both CN-driven (CCI-CN; A and C) and multiple alignment-driven effects (CCI-MA; B and D). E, scatterplot of CCI-MA versus CCI-CN showing strong correlation of the effect on CRISPR/Cas9 guide scores for either multiple alignment-driven or CN-driven DNA cuts across the cell lines.
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Figure 5. sgRNAs targeting multiple chromosomes show greater sensitivity to CRISPR/Cas9-induced cutting. Data from two representative cell lines are shown (PA-TU-8902, A, Panc 08.13, B). A and B, box plots of CRISPR/Cas9 sensitivity to the predicted number of CRISPR/Cas9-induced DNA cuts. CRISPR/Cas9 guide scores are shown on the y-axis and the predicted number of DNA cuts is shown on the x-axis. sgRNAs are divided into three groups. In red are sgRNAs that target a single locus, and therefore the total number of predicted cuts is based on CN. In yellow are sgRNAs that target multiple loci within a single chromosome (intrachromosomal). In blue are sgRNAs that target multiple loci across multiple chromosomes (interchromosomal). The analysis demonstrates a more potent detrimental influence on cell viability for multiple CRISPR/Cas9-induced DNA cuts across multiple chromosomes (interchromosomal) as compared with those restricted to a single chromosome (intrachromosomal). Multiple linear regression accounting for difference in the total number of cuts for interchromosomal versus intrachromosomal: A, \( a = -0.27, P = 2.64 \times 10^{-22} \); B, \( a = -0.16, P = 4.97 \times 10^{-22} \). C and D, waterfall plots showing CRISPR guide scores for all sgRNAs in the pooled screens performed on the indicated cell lines. sgRNAs from the multiple alignment analysis targeting multiple chromosomes with >10 predicted target sites are shown in red and are significantly enriched with negative CRISPR/Cas9 guide scores relative to all other sgRNAs in the library (one-sided Kolmogorov-Smirnov test: \( P = 2.13 \times 10^{-159} \); C, \( P = 9.17 \times 10^{-88} \); D). These data highlight the potent detrimental effect that these sgRNAs have on cell proliferation and viability within the screen.

also observed a potent reduction in cell proliferation with these multitargeted sgRNAs.

To investigate the mechanism of decreased cell proliferation observed with sgRNAs targeting amplified regions or multiple genomic loci, we utilized a high-content imaging assay to interrogate cell-cycle kinetics in multiple sgRNAs in parallel (40). At 48 hours after expression of these sgRNAs, we observed decreased incorporation of the modified thymidine analogue 5-ethynyl-2′-deoxyuridine (EdU), with diminished S-phase suggestive of decreased DNA synthesis (Fig. 6C). We also observed an accumulation of cells in the G2 phase of the cell cycle with sgRNAs targeting amplified regions or multiple genomic loci (Fig. 6C). Moreover, we observed an increased number of \( \gamma \)-H2AX foci in cells infected with these amplicon-targeting or multitargeted sgRNAs as compared with control sgRNAs, suggesting that increased DNA damage leads to a G2
Figure 6. CRISPR/Cas9 targeting of amplified regions or multiple genomic loci induces DNA damage and a G₂ cell-cycle arrest. A, schematic of the PANC-1 19q13 amplicon (top) demonstrating ABSOLUTE DNA CN (middle) and CRISPR guide scores (bottom) mapped by genomic position. Schematic and color scheme are similar to that detailed in Fig. 2. B, in vitro validation experiment measuring arrayed proliferation and viability response of PANC-1 cells at 6 days after infection with sgRNAs targeting regions inside (red) and outside (blue) of the demonstrated amplicon. sgRNAs targeting intergenic regions are labeled by chromosomal locus and columns are given a checkered pattern. Multitargeted sgRNAs (MT-1 and MT-2) are indicated by black bars. sgRNAs targeting an alternative unamplified locus (12q, orange) and known essential genes (green) are also shown. Nontargeting negative control sgRNAs are shown in yellow. Dots placed below the CN plot (A) correspond to the validation sgRNAs targeting the indicated genes or intergenic regions on the locus, and are matched by color and left-to-right genomic position. CellTiter-Glo was performed at 6 days after infection. Error bars indicate SD of biological replicates (n = 3). P < 0.0001 for two-tailed t test comparing sgRNAs inside (red) versus outside (blue and orange) the amplicon. C, plot of the percentage of PANC-1 cells in each phase of the cell cycle at 48 hours after infection with the indicated sgRNAs targeting inside (red) or outside (blue) the amplicon. Data for a multitargeted sgRNA (MT-2) and a control sgRNA targeting an alternative locus (12q-5), as well as for control genes are also shown. Fraction of cells in each phase of the cell cycle is indicated by a unique pattern within the column corresponding to each cell-cycle phase. Color scheme is as indicated above, with coloration of the G₂ and S phases for emphasis. Error bars represent the standard deviation for the mean of three replicates. D, plot of the number of γH2AX foci present in PANC-1 cells at 48 hours after infection with the indicated sgRNAs. Color scheme is as indicated above, with checkered pattern corresponding to sgRNAs targeting intergenic regions.

cell-cycle arrest in these cells (Fig. 6C and D). Notably, we did not observe significant levels of apoptosis at this same time point by measuring cleaved PARP by immunoblotting (Supplementary Fig. S9B). We have performed similar experiments with the chromosome 12 amplicon in the CAL120 breast cancer cell line and confirmed that these observations are not restricted to the chromosome 19 amplicon in PANC-1 (Supplementary Fig. S10A–S1E).

Overall, these observations suggest that CRISPR/Cas9 genome targeting of amplified regions induces a potent early DNA-damage response and cell-cycle arrest that is proportional to the number of target loci. Notably, this antiproliferative effect is independent of targeting expressed protein coding genes and does not depend on target gene disruption and protein loss, which typically occurs on a longer time scale (10).
Increased Genomic CN of Cell-Essential Genes May Protect from Complete Gene Knockout

Although we found that an increased number of target loci for each sgRNA generally leads to increased gene-independent CRISPR/Cas9-mediated cytotoxicity, we reasoned that because CRISPR/Cas9 genome editing is often incomplete within a cell population, more copies of a target gene could also make a cell resistant to complete gene disruption and protein loss through CRISPR/Cas9 targeting of that locus. Therefore, we hypothesized that certain cell-essential gene sets may show the opposite correlation with DNA CN in pooled negative-selection screening. When we examined the CRISPR–CN correlation across all genes screened in all cell lines from the dataset, we first found an overall negative correlation, as expected. However, we also observed that cell-essential genes from the Kyoto Encyclopedia of Genes and Genomes (KEGG) gene sets for the proteasome, ribosome, and spliceosome exhibit a CRISPR–CN correlation significantly shifted in the positive direction relative to the rest of the genes in the genome, i.e., higher CN correlated with higher CRISPR gene scores and less observed gene essentiality (Fig. 7). These observations suggest that increased DNA CN for target genes with strong underlying essentiality may protect cells from complete CRISPR-induced knockout of these genes, and thus manifest as relatively less apparent essentiality compared with other essential genes in CN normal regions of the genome. Together, these data further highlight the importance of considering target gene CN and gene function in the interpretation of negative selection pooled screening data.

DISCUSSION

Using data from the genome-scale interrogation of essential genes in 33 cancer cell lines by CRISPR/Cas9, we found that targeting sequences within CN amplification regions of the genome induces increased cell proliferation/viability that is independent of target gene expression or the structure of the targeted amplicon. The magnitude of the effect increases with the amplitude of CN amplification, and CRISPR/Cas9 targeting within high-level amplifications shows some of the most profound antiproliferative effects observed in the screens. Moreover, analysis of sgRNAs targeting multiple genomic sites also revealed a strong correlation of cell proliferation/viability with the number of predicted CRISPR/Cas9 DNA cuts. Thus, we propose that there are two types of responses to CRISPR/Cas9 targeting in cancer cell lines: (i) an early antiproliferative effect of CRISPR/Cas9-induced DNA cuts that increases with the number of cuts conferred by each sgRNA and that is independent of the target gene, and (ii) the gene essentiality resulting from CRISPR/Cas9-induced knockout of the target gene and subsequent loss of normal protein expression.

The mechanism of the early antiproliferative response to CRISPR/Cas9-mediated gene editing likely relates to induction of multiple double-strand DNA breaks and subsequent G$_2$ cell-cycle arrest. Wang and colleagues also recently reported an analysis of cell-essential genes using CRISPR/Cas9-mediated loss-of-function screens in four leukemia and lymphoma cell lines (28). They found that CRISPR/Cas9-mediated targeting of several genes within the BCR–ABL amplification in the K562 leukemia cell line and JAK2 amplification in the HEL erythroleukemia cell line induced decreased cell viability associated with increased levels of phosphorylated histone H2AX, a marker of DNA damage. Hart and colleagues also recently reported that guide RNAs targeting greater than 20 sites appear similar to known essential genes (25). Here, we present a comprehensive global analysis of this CRISPR–CN correlation in a large and diverse array of cancer cell lines and demonstrate that this phenomenon is pervasive across many different genetic and phenotypic contexts. Moreover, we provide the first evidence that this CRISPR–CN correlation occurs across a wide range of CNAs and chromosome structures, including those with low-level CN gain. Importantly, we demonstrate that targeting sequences within regions of high-level CN gain induces among the strongest observed viability phenotypes of all sgRNAs in the screen. Because this effect is not related to specific genes, these observations have important practical implications for utilizing CRISPR/Cas9 technology for cancer dependency profiling and for studying gene essentiality in general.

When we analyzed the effects on cell proliferation/viability induced by increased numbers of cuts, we noted that even a single CRISPR/Cas9-induced DNA cut resulted in decreased cell proliferation when compared with sgRNAs that do not target any human sequence. Thus, choice of negative controls for CRISPR/Cas9 experiments is critically important to interpret the consequences of CRISPR/Cas9-mediated genome
edit. Although nontargeting sgRNAs may best represent truly neutral negative controls, it may be more appropriate to use a targeting sgRNA directed at a non-genic and CN-normal region of the genome to better model the baseline impact of non-specific DNA targeting with CRISPR/Cas9. The observation that off-target CRISPR/Cas9 cuts likely also cooperate with on-target cuts to effect a cumulative toll on the cell highlights the paramount importance of optimal library design for better on-target and less off-target activity. Improved sgRNA libraries would thus allow better prediction of the total number of CRISPR/Cas9-induced DNA cuts according to baseline CN and therefore enable enhanced resolution of actual gene-based dependencies within the data.

Moreover, the observation that targeting the CRISPR/Cas9 endonuclease to even a single locus induced decreased proliferation/viability indicates that this approach to targeting genes induces a cellular response in the majority of cases. As such, the effects of this response should be considered in the interpretation of any phenotype observed after targeting a specific gene. Indeed, this observation may also affect efforts to use the CRISPR/Cas9 approach to perform genome editing for therapeutic purposes.

We also observed that for high-level genomic amplification, the cellular responses to CRISPR/Cas9 cutting toxicity overwhelm the signal from underlying gene essentiality, thus complicating efforts to use CRISPR/Cas9 for the identification of essential genes in amplified regions. Hence, it may be most prudent in individual cell line screening data to exclude certain reagents from consideration for the identification of essential genes, including sgRNAs targeting genomic amplifications as well as those predicted to confer multiple CRISPR/Cas9 DNA cuts. Failure to properly account for CNAs may lead to confounding effects and a higher rate of false-positive identification of cell-essential genes. Because CNAs are the most common genetic alteration found in human epithelial cancers, these observations have practical implications on both individual experiments as well as systematic efforts to interrogate the consequences of gene depletion. These observations also highlight the need to perform CRISPR/Cas9 screens across a large collection of diverse cancer cell lines to represent a variety of cancer gene dependencies while accounting for specific confounding genomic structural alterations within individual cell lines.

We propose that this observation extends beyond merely a confounding artifact of CRISPR/Cas9 technology and uncovers an important underlying biological concept that cancer cells are vulnerable to induction of site-specific double-stranded DNA breaks within regions of genomic amplification. Genome-scale CRISPR/Cas9 screening has provided an unprecedented resolution of the degree of DNA damage necessary to effect an antiproliferative or cytotoxic response in cancer cells, revealing an unappreciated susceptibility to even a small number of site-specific DNA breaks. Our observations support the notion that CRISPR/Cas9 targeting of amplified regions of the genome leads to increased DNA damage and a significant consequent antiproliferative response. Although these findings complicate the study of amplified regions with CRISPR/Cas9-based approaches, this early antiproliferative cell response may enable sequence-specific therapeutic approaches to target cancer. Many chemotherapy agents (e.g., cisplatin), as well as ionizing radiation, achieve their effects by inducing DNA damage that is not adequately repaired by cancer cells (41, 42). Although many cancer cells are more susceptible than normal cells to chemotherapy and radiation, a major limitation of these treatment approaches is the non-specific nature of these modalities and the narrow therapeutic window for preferential killing of cancer cells versus normal cells. Our observations suggest that targeting nonessential genes or even noncoding, intergenic regions of amplified DNA with CRISPR/Cas9 technology may unveil critical vulnerabilities in cancer cells that could be harnessed for cancer-specific therapy. A precision medicine approach using simultaneous combination of CRISPR/Cas9 reagents to target multiple amplified loci or tumor-specific mutated sequences within a cancer cell may enable development of cancer-specific treatments with an optimal therapeutic window.

**METHODS**

**CRISPR/Cas9 Screening**

Cancer cell lines were transduced with a lentiviral vector expressing the Cas9 nuclease under blasticidin selection (pXPR-311Cas9). Each Cas9-expressing cell line was subjected to a Cas9 activity assay (see below) to characterize the efficacy of CRISPR/Cas9 in these cell lines (Supplementary Table S1). Cell lines with less than 45% measured Cas9 activity were considered ineligible for screening. Stable polyclonal lines to represent a variety of cancer gene dependencies while accounting for specific confounding genomic structural alterations within individual cell lines.

We propose that this observation extends beyond merely a confounding artifact of CRISPR/Cas9 technology and uncovers an important underlying biological concept that cancer cells are vulnerable to induction of site-specific double-stranded DNA breaks within regions of genomic amplification. Genome-scale CRISPR/Cas9 screening has provided an unprecedented resolution of the degree of DNA damage necessary to effect an antiproliferative or cytotoxic response in cancer cells, revealing an unappreciated susceptibility to even a small number of site-specific DNA breaks. Our observations support the notion that CRISPR/Cas9 targeting of amplified regions of the genome leads to increased DNA damage and a significant consequent antiproliferative response. Although these findings complicate the study of amplified regions with CRISPR/Cas9-based approaches, this early antiproliferative cell response may enable sequence-specific therapeutic approaches to target cancer. Many chemotherapy agents (e.g., cisplatin), as well as ionizing radiation, achieve their effects by inducing DNA damage that is not adequately repaired by cancer cells (41, 42). Although many cancer cells are more susceptible than normal cells to chemotherapy and radiation, a major limitation of these treatment approaches is the non-specific nature of these modalities and the narrow therapeutic window for preferential killing of cancer cells versus normal cells. Our observations suggest that targeting nonessential genes or even noncoding, intergenic regions of amplified DNA with CRISPR/Cas9 technology may unveil critical vulnerabilities in cancer cells that could be harnessed for cancer-specific therapy. A precision medicine approach using simultaneous combination of CRISPR/Cas9 reagents to target multiple amplified loci or tumor-specific mutated sequences within a cancer cell may enable development of cancer-specific treatments with an optimal therapeutic window.

**Data Quality Control**

Quality control measures were used to remove cell line replicate samples where (i) the single-nucleotide polymorphism (SNP) genotype fingerprint failed to match the reference cell line as previously described (37), (ii) the reproducibility between replicates was less than 80%, and (iii) principal component analysis showed a replicate or cell line to be an outlier.

**Data Processing**

Data were processed in a reproducible GenePattern pipeline and are provided on the Project Achilles portal (43). A fold change was calculated per sgRNA, and the median of nontargeting controls (n = 1,000) in the GeCKOv2 library was subtracted from each sgRNA to generate a CRISPR guide score. Given the gene-independent effect of CRISPR/Cas9 described in this article, we chose to use the second-best CRISPR/Cas9 guide score for the purpose of ranking gene-level dependencies in individual cell lines. See Supplementary Methods for further details.

**Cancer Cell Lines and Cas9 Activity Assay**

Cancer cell lines were obtained primarily from the CCLE, which obtained each line from the original source (Supplementary Table S1; ref. 35). All cell lines were Mycoplasma negative, and identity was confirmed through fingerprinting prior to screening using an Affymetrix SNP array as previously described (37). Prior to screening, cell lines were engineered to stably express Cas9 under Blasticidin selection, and Cas9 activity was assayed using a lentivirus with an EF1a-driven
puromycin-2A-GFP cassette, and a U6-driven sgRNA targeting GFP (pXPR_011; ref. 44). The initial level of GFP is measured with FACS and monitored over time as a measure of cells harboring modified alleles. Cells with GFP remaining are due to either modifications that do not inactivate GFP florescence or inactive Cas9.

**Essential Gene Controls**

Genes from the KEGG gene sets for ribosome, proteasome, and spliceosome subunits (Supplementary Table S2) were used as cell-essential (positive) controls in the analysis of negative selection CRISPR/Cas9 screening data. Guide sequences that were a perfect match to sgRNAs targeting any other gene or noncoding sequence were removed, except when specifically utilized in described analyses.

**CN Analysis**

DNA CN data were derived from SNP microarrays, and ABSOLUTE CN calls were made as previously described (35, 36). CRISPR/Cas9 screening data were mapped according to the genomic position of sgRNA sequence (guide-level data) or target gene (by ATARiS algorithm) to the human genome version 19 (hg19). CRISPR/Cas9 screening data were plotted in parallel to Project Achilles shRNA dependency data (43) or CCLE CN or gene expression data (35, 37).

**WGS and Analysis**

WGS was performed through the Broad Institute-Novartis CCLE, as previously described (45). Whole-genome DNA sequencing data of the cancer cell lines are aligned by the Burrows-Wheeler aligner (46) to the human genome reference 19. The aligned reads are filtered for PCR duplicates by MarkDuplicates from Picard. Read-depth coverage was computed and normalized using the previously described approach (47). Briefly, the number of aligned reads was counted for nonoverlapping 1-kb bins and then normalized for GC-content and mappability biases using the HMMeCopy R/Biocondutor package. The normalization was applied to both the cancer cell line and nontargeting negative control sgRNA. Error bars are the result of three biological replicates.

**Immunoblots**

Cells were infected at high MOI in 6-well plates, and protein was extracted at 48 hours after infection. Immunoblotting was performed using antibodies for PARP (Cell Signaling Technology, 46D11, #9552, 1:1,000) and β-actin (Sigma Aldrich, A5316, 1:5,000).

**High-Content Imaging Assay and Analysis**

PANC-1 and CAL120 cells constitutively expressing Cas9 protein were plated at a density of 4,000 cells per well, infected in replicates in 96-well plates at high MOI and analyzed at an endpoint of 48 hours after infection. Cells were labeled with EdU and fixed with paraformaldehyde and then labeled with anti-phH3 (S10) primary antibody (Rabbit: #9701, Cell Signaling Technology, 1:800), anti-phospho-histone H2A.X (Ser139, Mouse: 05-636, END Millipore, 1:1,250) and Hoechst 33342 (H3570, Thermo Fisher Scientific, 1 μg/mL). Imaging was performed with the OperaPhenix imaging system on 20x magnification, and data were analyzed using the PerkinElmer Harmony software (40). See Supplementary Methods for additional details.

**Disclosure of Potential Conflicts of Interest**

C.-Z. Zhang is a consultant/advisory board member for Pillar Biosciences and Ori Capital. A.D. Cherniack reports receiving a commercial research grant from Bayer AG. G. Kryukov is Senior Director, Computational Biology, at KSQ Therapeutics. L.A. Garraway reports receiving a commercial research grant from Novartis, has ownership interest (including patents) in Foundation Medicine, and is a consultant/advisory board member for Warp Drive, Novartis, Foundation Medicine, and Boehrirger Ingelheim. M. Meyerson reports receiving a commercial research grant from Bayer and is a consultant/advisory board member for the same. W.C. Hahn reports receiving a commercial research grant from Warp Drive, Novartis and is a consultant/advisory board member for the same. No potential conflicts of interest were disclosed by the other authors.

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Genomic Copy Number Affects CRISPR/Cas9 Screens


Genomic Copy Number Dictates a Gene-Independent Cell Response to CRISPR/Cas9 Targeting

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