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15 Abstract

Multiple high-throughput functional screens in cancer cell lines have generated large 16 amounts of information on drug efficacy in a variety of genomic contexts and cancer 17 lineages. Inconsistencies between datasets have led us to investigate whether these 18 large-scale screens reproduce established clinical drug-gene associations and if genomic 19 features particular to specific genes improve said reproducibility. We evaluated three 20 large-scale, small-molecule drug screens and one CRISPR/Cas9 gene essentiality screen, all 21 within the context of clinical interpretations derived from a new cancer variant annotation 22 resource published by the Variant Interpretation for Cancer Consortium (VICC). We identified low levels of concordance between the three drug screen datasets and clinical drug-gene associations using mutation status, gene expression, and copy number as genomic indicators. Less than half of the clinical drug-gene cancer associations from the

VICC resource were identified in these three drug screen datasets, suggesting a barrier to

translating findings from these large-scale screens into the clinic.

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32 Introduction

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Cancer cell lines are a long-standing model for systematic testing of candidate 33 therapeutics, beginning with the National Cancer Institute 60 (NCI60) assay from the late 34 1980s (M. C. Alley et al., 1988; R. H. Shoemaker, 2006; Stinson et al., 1992), which has 35 been used to screen over 100,000 compounds as of 2010 (Holbeck, Collins, & Doroshow, 2010). Since then, numerous small-molecule and gene essentiality screens of various scales 37 and study aims have been conducted in cell lines, from grouping drugs by the apeutic target similarity (Greshock et al., 2010) to screening only near-haploid cell lines to generate genome-level insights into gene essentiality (T. Wang et al., 2015) to broadly identifying cancer dependencies with large-scale screens in multiple cancer types (McDonald et al., 2017; J. M. McFarland et al., 2018; Meyers et al., 2017; Patel et al., 2017) or select lineages (Heiser et al., 2009; Marcotte et al., 2012; Patel et al., 2017). Despite their widespread use, cancer cell lines are known to have issues with inconsistent naming conventions and contamination (M. Yu et al., 2015), and some cell lines have been shown to vary widely at the genetic level and in response to drug treatment across strains (Ben-David et al., 2018). Additionally, comparisons between cancer cell lines and tumors indicate that cell lines have higher numbers of genomic aberrations (Domcke, Sinha, Levine, Sander, & Schultz, 2013; Mouradov et al., 2014; Neve et al., 2006) and tend to be hypermethylated (Paz et al., 2003; Varley et al., 2013), which could prove an impediment to translating cell line discoveries into the clinic. 51 There have also been debates over consistency between drug screen datasets, namely 52

There have also been debates over consistency between drug screen datasets, namely
the Broad Institute's and Novartis Institutes for Biomedical Research's Cancer Cell Line
Encyclopedia (CCLE) (J. Barretina et al., 2012; T. C. C. L. E. Consortium & Consortium,
2015) and the Genomics of Drug Sensitivity in Cancer (GDSC) from the Cancer Genome
Project at the Wellcome Sanger Institute and the Center for Molecular Therapeutics at

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Massachusetts General Hospital Cancer Center (Garnett et al., 2012; Yang et al., 2012).
   The GDSC has also been referred to as the Cancer Genome Project (CGP) and the Sanger
   dataset. Studies have shown that drug-gene interactions matched between CCLE and
   GDSC exhibited poor correlation and inconsistencies (Haibe-Kains et al., 2013; Jang, Neto,
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   Guinney, Friend, & Margolin, 2014), prompting other groups to join the debate on how
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   best correct for experimental and methodological variation between the original drug
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   screens and subsequent computational analysis (T. C. C. L. E. Consortium & Consortium,
   2015; Geeleher, Cox, & Huang, 2016; Geeleher, Gamazon, Seoighe, Cox, & Huang, 2016;
   Hatzis et al., 2014; Haverty et al., 2016; Safikhani et al., 2016, 2017).
        The prevalence and importance of cancer cell lines in large-scale therapeutic research
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   and the apparent inconsistencies between the GDSC and CCLE encouraged us to compare
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   the results from these functional screens to clinical drug-gene associations. An outstanding
   question concerning studies based on cancer cell lines is whether these the cell line systems
   can accurately model tumor dynamics or recapitulate clinical cancer vulnerabilities, and
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   many large-scale grants and clinical trials are fundamentally anchored by results from
71
   screens conducted in cancer cell lines. Thus, our goal was to evaluate how well these
   functional screens recapitulate known drug, gene, and tumor type associations that are
   currently used in clinical decision-making.
        We derived our clinical associations from a new project conducted by the Variant
75
   Interpretation for Cancer Consortium (VICC; https://cancervariants.org/), a Driver
   Project for the Global Alliance for Genomics and Health (GA4GH) (The Global Alliance
   for Genomics and Health, 2016). The VICC has curated annotations of known cancer
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variants at varying levels of evidence from multiple resources (A. H. Wagner et al., 2018):
the Precision Medicine Knowledgebase (L. Huang et al., 2016), MolecularMatch
(https://www.molecularmatch.com/index.html), OncoKB (Chakravarty et al., 2017),
Jackson Labs Clinical Knowledgebase (S. E. Patterson et al., 2016), Clinical
Interpretations of Variance in Cancers (CIViC) (Griffith et al., 2017), and the Cancer

Genome Interpreter (CGI) (Tamborero et al., 2018). These harmonized variants are hosted on an ElasticSearch (Kibana v6.0) platform called Genotype to Phenotype (G2P, 85 https://search.cancervariants.org/#*) that allows users to query and filter aggregated 86 evidence from the various databases listed above as well as the GA4GH beacon service 87 (http://beacon-network.org/#/), which yields access to genetic mutation data from over 88 200 datasets as of 2016 (The Global Alliance for Genomics and Health, 2016). For drug screen datasets, we focus on three well-known, large-scale drug projects: the CCLE and GDSC, which were mentioned above, and the Center for the Science of Therapeutics at the Broad Institute's Cancer Therapeutics Response Portal (CTRP) dataset (Basu et al., 2013; M. G. Rees et al., 2016; B. Seashore-Ludlow et al., 2015). We also analyzed the results from a large-scale CRISPR/Cas9 screen conducted by the Broad Institute's Cancer Dependency Map project (DepMap) using the Avana knockout library (Doench et al., 2016; Meyers et al., 2017), which will address whether gene essentiality screens via CRISPR yield results comparable to functional drug screens.

98 Results

99 Comparison of G2P drug-gene associations and CCLE, CTRP, and GDSC

G2P designed a set of standards for stratifying their database's cancer variant 100 interpretations, ranging from preclinical data at the low-evidence end (level D) to clinically 101 actionable interpretations at the high-evidence end (level A). To test only these clinically 102 actionable associations, we filtered the G2P dataset for only level-A G2P interpretations, of which there were 1,296 (see Methods). All compounds screened within the GDSC, CTRP, and CCLE datasets are considered "small molecules" and are respectively linked to 105 Compound (CID) numbers used for indexing in the PubChem database 106 (https://pubchem.ncbi.nlm.nih.gov/). As a result, non-small molecules, which include 107 proteins and biologics (i.e. monoclonal antibodies), that had level-A G2P evidence could 108

not be included in our comparisons. Thus, G2P interpretations for which the compound did not have a matching CID code were excluded from further analysis as were interpretations that did not have an associated compound, resulting in removal of 402 interpretations (Table 1). It should be noted that cisplatin, a small molecule, was excluded due to its being indexed with a ChEMBLdb identifier (https://www.ebi.ac.uk/chembl/), which we did not manually re-index to a CID to avoid accidental mischaracterization.

After filtering, 894 level-A G2P interpretations were carried forward in subsequent 115 analyses, constituting 57 unique drugs and 34 unique genes in 88 distinct level-A 116 combinations (Figure 1A). The number of level-A interpretations per drug ranged from 1 117 to 176 (M = 15.68, SD = 34.63), 1 to 264 (M = 26.24, SD = 61.58) per gene, and for each 118 level-A drug-gene combination, the range was 1 to 103 (M = 10.14, SD = 19.80). These 119 unique drugs, genes, and combinations are hereafter referred to as G2P drugs, G2P genes, 120 and G2P associations. Filtering our screening datasets by these clinical criteria yields 121 panels of potentially clinically relevant omic features. Of the 57 unique G2P drugs, 37 of 122 them were screened in at least one of the drug screen datasets, 17 drugs were screened in at 123 least two of the three datasets, and only 3 were screened in all three datasets. The 124 remaining 20 un-screened G2P drugs suggests an incompleteness of these drug screens, though these projects frequently release new screening data that may include these drugs in future releases. 127

Using cell-line-specific point mutation annotations from the CCLE, we compared z-score-transformed area under the dose-response curve (AUC) distributions between mutant and wildtype cell lines for all G2P genes for each G2P drug tested in the CCLE, CTRP, and GDSC drug screens (Figures 1B and 2A). At an unadjusted $\alpha = 0.05$, Wilcoxon rank-sum tests revealed that in CCLE, the AUC distributions of mutant and wildtype groups for 18 out of 170 (10.59%) tested drug-gene combinations were significantly different, 135 out of 918 (14.71%) were significantly different for CTRP, and 89 out of 850 (10.47%) for GDSC.

Controlling for a false discovery rate (FDR) < 5%, selumetinib-BRAF (p < .001) was the only significant drug-gene combination tested in CCLE ($p_k < 8^{-4}$). CTRP ($p_k < 7^{-4}$) yielded 918 significant combinations, one of which was selumetinib-BRAF.

We observe little concordance between the number of interpretations for each G2P association and the significance of mutation status on AUC for said G2P association, suggesting that mutation status alone is not a robust predictor of clinically actionable genetic targets. This led us to examine the correlations between AUC and gene expression and AUC and copy number, two other genomic features that are known to associate with cancer and drug response.

We computed Spearman correlation coefficients (r_s) between AUC of each drug-gene association and gene expression (RPKM) and copy number (log₂ ratio) of the gene in the association

Do small-molecule screens in cancer cell lines recapitulate clinical drug-gene associations?

Cut this analysis since we are comparing distributions of p-values?

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We compared the distribution of level-A G2P associations (Figure 1A) and the 151 associated p-values from the mutation status comparisons (Figure 1B) with a null 152 distribution of p-values derived from all pairwise combinations of G2P genes and the G2P 153 drugs tested in each dataset, less any of the 88 distinct combinations with G2P 154 interpretations, i.e. the non-gray squares in the right heatmap of Figure 1A (Figure 3A). To formally test concordance between clinically actionable variants and drug screen results, we conducted an independent, two-group t-test to compare the distributions of p-values of 157 drug-gene combinations that either were or were not represented with level-A evidence in 158 G2P, which indicated that the distributions of p-values did not significantly differ for 159 CCLE [t(9) = 0.25, p = 0.81], CTRP [t(51) = 0.85, p = 0.40], nor GDSC [t(47) = -0.56, p]160

161 = 0.58] (Table 2). This further supports that mutation status is not a robust genomic
162 feature for predicting therapeutic responses.

Similar tests were conducted for the distributions of Pearson correlation coefficients 163 (r) between the number of level-A G2P associations and gene expression (RPKM) and copy 164 number (log2 ratio) of the gene in the association (Figures 3B-C). Example r derivations 165 are shown in Figures 2B-C. Absolute values of r were used to simplify extreme values in 166 both directions for testing. The difference in the CCLE |r| distributions for both gene 167 expression [t(8) = 1.43, p = 0.19] and copy number [t(8) = 2.48, p = 0.23] were 168 non-significant, which is likely explained by the small sample size since only five G2P drugs 169 were screened in the CCLE dataset. For both the CTRP and GDSC screens, |r| values for 170 gene expression correlations were significantly greater for the level-A G2P associations than 171 the null distribution [tCTRP(49) = 3.79, pCTRP < 0.001; tGDSC(44) = 2.97, pGDSC = 172 0.005]. The |r| distributions using copy number were also significantly different for CTRP 173 [t(49) = 2.48, p = 0.017] but not for GDSC [t(44) = 1.95, p = 0.058]. 174

In all three datasets for both level-A G2P associations and null distribution 175 associations, there are more significant |r| values when using gene expression as the 176 genomic feature than when using copy number (Table 2), which corresponds with previous 177 work showing that gene expression is an informative molecular feature when assessing drug 178 sensitivity (Jang et al., 2014; Sirota et al., 2011). Additionally, in all cases, there are more 179 significant r values when using copy number than there are significant p-values from the 180 mutation status comparisons. Though observational, these results suggest an order of 181 predictive robustness for these three genomic features when used in isolation: Gene expression > copy number > mutation status. Additionally, across all three genomic features, CTRP consistently identified a higher percentage of its possible level-A G2P 184 associations than GDSC, and similarly, GDSC identified a higher percentage than CCLE. 185 In the CCLE-GDSC debate, inconsistency between the datasets was attributed to the 186 groups using two very different assays for generating AUC measurements though it was 187

undecided which was more accurate (Haibe-Kains et al., 2013; Jang et al., 2014; Weinstein & Lorenzi, 2013). Our results provide very preliminary evidence that GDSC is more accurate than CCLE in regards to reproducing clinical associations. However, CTRP may be even more accurate than both GDSC and CCLE.

Do these pancancer drug-gene associations penetrate to the lineage-specific level?

- What do these correlations with mutation status, gene expression, and copy number look like when you constrict to a specific lineage?
 - When you restrict, do the correlations become more distinct?

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- Are the lineage-specific results approximately the same as the pancancer results?
- Looking at a histogram of p-values, if the histogram is flat, then nothing is significant (note from meeting on 11/16)

In the 894 level-A G2P interpretations, there was a wide variation in the specificity of 200 cancer description, ranging from "Waldenström macroglobulinemia" and "hyper 201 eosinophilic advanced syndrome" on the more detailed end to "cancer" on the broad end. 202 Similarly, the drug screen datasets had a wide range in lineage specificity, e.g. CCLE, 203 GTRP, and GDSC all had lineages labeled "leukemia" and "T-cell childhood acute 204 lymphocytic leukemia" and the CRISPR dataset had cell lines labeled "Epstein-Barr 205 virus-related Burkitt lymphoma" and "lymphoma." To more easily conduct comparative 206 analyses in a cancer-specific manner, we developed a more general lineage grouping method 207 derived from the Human Disease Ontology identity codes (DOIDs, 208 http://disease-ontology.org/) (Schriml et al., 2019) assigned to each cell line, yielding 27 209 unique lineage groupings. This custom lineage grouping was included in our harmonized 210 cancer cell line database (see Methods). 211

In the level-A G2P interpretations, there were 102 unique drug-gene-lineage

combinations (DGLs), representing seven of our curated cancer lineages: cancer, lung 213 cancer, breast cancer, leukemia, thyroid cancer, skin cancer, and ovarian cancer. CCLE had 214 data for 9 DGLs, CTRP had data for 61, and GDSC had data for 53, yielding a total of 73 215 unique DGLs across all three datasets. Only 7 DGLs were screened in all three datasets, 216 Prior to correction for an FDR of 5%, no DGLs were identified in the CCLE screen 217 when using mutation status as a correlate (n = 7, 2 DGLs) were not analyzed due to lack of 218 mutation annotations in UGT1A1 in "cancer" lineages), but CTRP (n = 48) and GDSC (n=39) identified 6 and 5, respectively, in skin cancer, lung cancer, and cancer. After Benjamini-Hochberg FDR correction, none of the datasets yielded significant DGL 221 interactions. 222

Are CRISPR/Cas9 gene essentiality results comparable to those of functional drug screens?

The CRISPR screen from the Broad Institute reports gene dependency using CERES scores (Meyers et al., 2017), which are generated from sgRNA depletion scores and eliminate bias arising from the effect of copy number variation on Cas9 DNA cleavage. The lower the CERES score, the higher the likelihood that the gene is essential in the associated cell line. Scores are scaled per cell line such that a score of 0 is the median effect of nonessential genes and -1 is the median effect of common core essential genes. Of the 34 G2P genes, only *G6PD* was not targeted by an sgRNA in the CRISPR dataset.

When we don't account for lineage, Wilcoxon rank-sum tests ($p_k < 5^{-3}$, FDR < 5%)
revealed that mutations in 2 genes significantly correlated with CERES score: KRAS (p < .001) and BRAF (p < .001).

Gene expression (RPKM) of 12 out of 33 (36.36%) G2P genes significantly correlated with CERES score, as did copy number (log2-ratio) for 12 out of 33 (36.36%) G2P genes.

For the purpose of comparison to the drug screen data, if we treat CRISPR

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knockdown like a small-molecule drug, then these results are similar to the results from CCLE, CTRP, and GDSC (Table 2).

• Lineage?

Are known copy number driven associations recapitulated in the drug and CRISPR screens?

If a gene's response to is copy number driven, we would expect copy number to significantly correlate with AUC and gene essentiality scores, as would gene expression is expression and copy number are associated.

In a lineage-agnostic context of the CRISPR screen, ERBB2 gene essentiality score had a significant negative correlation with ERBB2 gene expression $(p < 10^{-12}, r = -0.31)$ and ERBB2 copy number $(p < 10^{-15}, r = -0.34)$ (Figure ##2). Similarly, MET gene essentiality also correlated with both MET gene expression $(p < 10^{-12}, r = -0.31)$ and copy number $(p < 10^{-8}, r = -0.25)$. Notably, BRCA1 also significantly correlated with both gene expression (p < 0.01, r = 0.14). BRCA2 significantly correlated with all three of the genomic features tested: gene expression (p < 0.05, r = 0.10), copy number $(p < 10^{-7}, r = 0.23)$, and mutation status (p < 0.001).

Which genomic features drive KRAS, EGFR, and BRAF drug-gene associations?

Should we do this?

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257 Discussion

Level-A G2P interpretations correspond to clinical evidence that suggests efficacy of gene targets and/or drugs, and this work questions whether these relationships manifest

themselves in in vitro drug screens. We have demonstrated that on its own, mutation status is not a robust predictor of drug efficacy and that copy number and gene expression fare 261 better as predictors. However, for these three genomic features in all three datasets, only 262 11.1% (mutation status, 1 of 9, CCLE) to 47.9% (gene expression, 23 of 48, CTRP) (Table 263 2), of the level-A G2P drug-gene associations in the respective dataset were significant at 264 an uncorrected $\alpha = 0.05$. This suggests that while the drug screens successfully identify 265 some clinically relevant drug-gene associations, many associations are also missed, which 266 raises the question of the extent to which clinical researchers can rely on in vitro drug 267 screens when attempting to develop and select therapeutics for cancer patients. 268

These results also highlight the need to account for the complex differences inherent 269 between data generated in clinical scenarios and data captured in highly controlled, 270 artificial in vitro environments. Potential sources of variation and error include the fact 271 that cells growing in a laboratory as opposed to those, even of the same tissue type, 272 growing in a multicellular organism are exposed to differing sets of stressors and signals 273 that can significantly impact intracellular signaling pathways, irrespective of shared 274 genomic profiles. These differences can include, but are not be limited to, immunologic 275 reactions, both adaptive and intrinsic (e.g. cytokine signaling, inflammation), endocrine (e.g. stress hormones), and nervous system stimulation, all of which can have significant 277 downstream implications on cellular behavior in the context of therapeutic efficacy. Similarly, a laboratory environment and in vitro cell culture introduce abnormal growth conditions with respect to extracellular matrix composition, nutrient availability, and cell 280 density, all of which have the potential to alter cellular signaling and thus render a drug 281 ineffective during screening despite would-be in vivo activity. 282

In the era of precision medicine, the ultimate boon would be that the unique omics
signatures of a patient's individual cancer can be used to guide treatment.

High-throughput, in vitro screens of targeted cancer agents against cell lines with known
omics profiles are beneficial for testing hypotheses concerning the mechanisms of action for

these agents and they allow for scalability and systematic screening. However, 287 extrapolating the potential downstream effects of inhibiting a major growth pathway 288 (e.g. MAPK, IP-DAG) to predict the clinical prognosis and progression of a multicellular 289 tumor mass growing in a complex environment, compounded with the cross-reacting effects 290 of tumor genomic heterogeneity, make it a questionable statement that one can safely 291 predict clinical consequences from suppressing a single pathway in a model system, as 292 evidenced by our findings of fewer than half of the known clinical associations in the three 293 drug screen datasets. The benefits of these cancer cell line models, in contrast, lie in their 294 ability to assess the big-picture effects of these agents. In order to fully understand 295 therapeutic drug response, a higher degree of granularity is needed. Additional effort to 296 combine more cell line information such as gene expression, epigenetic profiles, and 297 proteomic data with mutational profiles is essential to improving the efficacy and validity of in vitro cell line screens. Further benefit could be derived from conducting these screens in environments that are more in line with the clinical scenarios we are trying to predict. This would include things such a 3-dimensional cell culture, coculturing with stromal cells 301 to mimic the tumor microenvironment, as well as other modalities that can attempt to 302 better replicate in vivo environments. 303

As a next step to improve the informational granularity of the analysis presented
here, our subsequent goal will be to assess the extent to which these drug-gene associations
are specific to individual cancer cell line lineages, which are available and annotated in the
data sets analyzed within this report. By evaluating how many clinically actionable
associations are identified in these large-scale functional screens, we can begin to address
best practices for translating discoveries from these tumor models into clinical trials.

310 Methods

G2P cancer variants

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Cancer variants with the highest level of evidence (i.e. level A) in the Genotype to 312 Phenotype (G2P) database were retrieved from the Variant Interpretation for Cancer 313 Consortium (VICC) portal (https://search.cancervariants.org/#*) (A. H. Wagner et al., 314 2018) using a customized JSON-query script. All JSON queries were passed to the 315 available application program interface (API) where a request was made for all drug-gene 316 associations with level-A evidence via the G2P evidence label, 317 i.e. association.evidence label. From manual inspection using G2P's front-end Kibana 318 interface, it was known that, at the time of the last query, 27 November 2018, there were 319 1,297 known level-A associations. Due to the limited request processing capabilities of the 320 G2P JSON API, queries were batched into packets of 10 data points, for a total of 130 321 requests made in succession. Each returned request was processed as JSON object and 322 searched to identify any existing key:value pairs for the following variables: evidence level, 323 mutation, gene, chromosome, start, end, ref, alt, direction, phenotype description, 324 phenotype family, phenotype ID, drug, drug ID, feature names, and sequence ID. For any evidence point where a given key:value pair was not found or unavailable, a value of -1 was assigned. One of the 1,297 level-A entries was irretrievable, yielding a final dataset of 1,296 level-A G2P interpretations, which were stored in a pandas data table and exported in CSV format. The VICC's methods for the harvesting and harmonizing of cancer variants is 329 available in a GitHub repository from Oregon Health & Science University 330 (https://github.com/ohsu-comp-bio/g2p-aggregator). 331

DepMap data retrieval and processing

CRISPR/Cas9, CCLE mutation calls, and CCLE copy number data were all retrieved from the Broad Institute's Cancer Dependency Map (DepMap) Public 18Q4 release via the

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DepMap data portal (https://depmap.org/portal/download/) (Broad DepMap, 2018).
   Gene expression data was retrieved from the DepMap Public 18Q3 release (Broad
336
   DepMap, 2018) from the same data portal. For cell lines with no CCLE annotation, the
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   DepMap group drew raw copy number and mutation data from whole exome sequencing
338
   data produced by the Wellcome Trust Sanger Institute [Catalogue Of Somatic Mutations
339
   In Cancer (COSMIC, https://cancer.sanger.ac.uk/cell lines) (S. Bamford et al., 2004; S.
340
   A. Forbes et al., 2017); European Genome-phenome Archive (Lappalainen et al., 2015),
341
   accession number EGAD00001001039 and processed the data following the CCLE
   pipelines to ensure consistency.
343
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CRISPR/Cas9 screen. Gene effect scores for 17,634 genes in 517 cell lines were 344 inferred from a CRISPR/Cas9 (clustered regularly interspaced short palindromic 345 repeats/CRISPR-associated 9) screen using the Broad Institute's Avana knockout library 346 (Doench et al., 2016). The Broad Institute's data processing and screening methods are 347 available from the figshare record (Broad DepMap, 2018) and the original publication of 348 CERES, the algorithm that computes inferred gene dependency scores (Meyers et al., 349 2017). The DepMap releases new data quarterly, with the current data set being the 18Q4 350 release. This release has 175 more cell lines than the original data release that was published with CERES (dataset: gene effect.csv, accessed 15 November 2018). 352

To preclude variation in genomic feature calls across the datasets, Mutation calls. 353 we used identical annotations from CCLE for all cell lines screened in the three datasets. 354 The CCLE provided mutation annotations in 19,280 genes across 1,596 cell lines (dataset: 355 depmap 18Q4 mutation calls.csv, accessed 14 November 2018). CCLE called substitutions using MuTect (Cibulskis et al., 2013) and annotated variants using Oncotator (Ramos et al., 2015) and indels using Indelocator 358 (https://software.broadinstitute.org/cancer/cga/indelocator). We filtered mutation calls 359 for point mutations, defined as single-nucleotide insertions, deletions, and substitutions, 360 regardless of result, i.e. frameshift, missense, or nonsense mutation. For this analysis, cell 361

lines harboring non-silent point mutations were considered "mutant" for the gene in
question. All other mutations and genes without annotation in the Mutation Annotation
Format (MAF) file were considered "wildtype." "Mutant" and "wildtype" calls were
binarily encoded per-gene/per-cell line, regardless of quantity of harbored mutations per
gene in a given cell line.

Copy number data. The CCLE generated genomic copy number (CN) data using
the Affymetrix Genome-Wide Human SNP Array 6.0 and GenePattern pipeline (The
Cancer Genome Atlas Research Network, 2008) and normalized segmented CN log2-ratios
for 23,299 genes across 1,098 cell lines using circular binary segmentation (Olshen,
Venkatraman, Lucito, & Wigler, 2004) (dataset: public_18Q4_gene_cn.csv, accessed 15
November 2018).

Gene expression data. The CCLE reports gene expression in RPKM (reads per kilobase per million mapped reads) for 54,356 genes across 1,156 cell lines, generated on the GeneChip Human Genome U133 Plus 2.0 Array (dataset:

CCLE DepMap 18q3 RNAseq RPKM 20180718.gct, accessed 18 July 2018).

Drug screen dataset retrieval and harmonization

The Cancer Therapeutics Response Portal (CTRP) drug screen dataset (v2) (Basu et 378 al., 2013; M. G. Rees et al., 2016; B. Seashore-Ludlow et al., 2015) was retrieved from the 379 National Cancer Institute's Cancer Target Discovery and Development (CTD2) Network's 380 data portal (https://ocg.cancer.gov/programs/ctd2/data-portal, accessed 6 June 2018). 381 The Genomics of Drug Sensitivity in Cancer (GDSC) drug screen dataset (Garnett et al., 2012; Yang et al., 2012) was retrieved from the data portal at (https://www.cancerrxgene.org/downloads, accessed 19 June 2018). The Cancer Cell Line Encyclopedia (CCLE) drug screen dataset (J. Barretina et al., 2012; T. C. C. L. E. 385 Consortium & Consortium, 2015) was retrieved from the Broad Institute's data portal 386 (https://portals.broadinstitute.org/ccle, accessed 20 June 2018). 387

Cell line harmonization. In an effort to compare CCLE, CTRP, GDSC and 388 CRISPR results and include lineage specificity, it was necessary to consolidate cell lines 380 screened in each dataset. Given that unique identifiers can be used between studies, it was 390 necessary to standardize the identity of each cell line with a common source. To achieve 391 this, a freely available cell line database called Cellosaurus (v26.0, 14 May 2018) (Bairoch, 392 2018) was downloaded and merged with the Broad Institute's DepMap cell line data 393 (accessed 8 June 2018) to create a harmonized cancer cell line database containing 394 synonymous identifiers that enabled consolidation of all cell lines used in CCLE, CTRP, 395 GDSC and CRISPR into a common framework. Subsequent additions to this database 396 include the merging of DepMap IDs used in the 18Q3 and 18Q4 DepMap data releases. We 397 curated missing synonyms and identifiers, and we manually annotated the granularity of 398 lineage labeling for all cell lines analyzed in this study based on their Human Disease Ontology identity codes (DOIDs, http://disease-ontology.org/) (Schriml et al., 2019), resulting in 27 unique lineages. For ease of comparison between datasets, we labeled all cell lines with custom, randomized identifiers. 402

AUC harmonization. The CCLE reports cell line sensitivity to small molecules as
an activity area measured above the dose-response curve (ActArea), while both CTRP and
GDSC report small molecule activity as area under the dose-response curve (AUC). Since
ActArea is reported with a scale of 0 to 8, we converted CCLE's ActArea values to AUC via

$$AUC = 1 - \frac{ActArea}{8}$$

All AUC values across the three drug screen datasets were z-score transformed:

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$$z = \frac{AUC - \mu_{AUC}}{\sigma_{AUC}}$$

where AUC is the mean AUC across all drug-cell line combinations in the dataset and AUC is the standard deviation.

Computational

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For Figure 1, all data were imported and manipulated using pandas (v0.23.0)
411
   (McKinney, 2011) and plotted using seaborn (v0.90) (Waskom, 2012), which sat on top of
412
   matplotlib (v2.2.2) (Hunter, 2007) and was executed on Python (v2.7.15) (Python Software
413
   Foundation, https://www.python.org/) in an IPython (v5.5.0) (Pérez & Granger, 2007)
414
   kernel within a localhost Jupyter (v5.2.2) notebook (Kluyver et al., 2016). All other
415
   analyses were conducted in R (v3.5.0) (R Development Core Team, 2010,
416
   https://www.R-project.org/) in an RStudio environment (v1.1.447) (RStudio, 2012). All
417
   other plots were drawn using the ggplot2 (v3.0.0) R package (Wickham, 2016).
         Statistics.
                      Wilcoxon tests were conducted using the compare_means function in
419
   the ggpubr (v0.1.7.999) R package (Kassambara, 2018). compare means does
420
   Benjamini-Hochberg p-value adjustment using the p,adjust function from the stats base
421
   R package.
422
         For computing Spearman correlation coefficients. If there were 3 or fewer data points,
423
   then the correlation was not done and a value of NA was supplied.
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Finally, the APA guidelines require a note at the start of the reference section that explains
426
           what an asterisk means. This note can be added at the end of the document as
427
          follows.
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Table 1 ${\it Distribution~of~level-A~G2P~drugs~excluded~from~analyses}$

Compound	Compound identifier	Number of G2P interpretations	Proportion of excluded interpr
cetuximab	CHEMBL1201577	74	18.41
cisplatin	CHEMBL2068237	3	0.75
EGFR	SID160769799	1	0.25
ipilimumab	CHEMBL1789844	2	0.50
mAb	CHEMBL2109423	16	3.98
nivolumab	CHEMBL2108738	4	1.00
panitumumab	CHEMBL1201827	75	18.66
pembrolizumab	CHEMBL3137343	3	0.75
pertuzumab	CHEMBL2007641	1	0.25
trastuzumab	CHEMBL1201585	9	2.24
NA	NA	214	53.23

Note. mAb = monoclonal antibody; NA indicates G2P interpretation does not have an associated compo