Growth rate inhibition metrics correct for confounders in measuring sensitivity to cancer drugs

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Drug sensitivity and resistance are conventionally quantified by IC50 or Emax values, but these metrics are highly sensitive to the number of divisions taking place over the course of a response assay. The dependency of IC50 and Emax on division rate creates artefactual correlations between genotype and drug sensitivity, while obscuring valuable biological insights and interfering with biomarker discovery. We derive alternative small molecule drug-response metrics that are insensitive to division number. These are based on estimation of the magnitude of drug-induced growth rate inhibition (GR) using endpoint or time-course assays. We show that GR50 and GRmax are superior to conventional metrics for assessing the effects of small molecule drugs in dividing cells. Moreover, adopting GR metrics requires only modest changes in experimental protocols.

We expect GR metrics to improve the study of cell signaling and growth using small molecules and biologics and to facilitate the discovery of drug-response biomarkers and the identification of drugs effective against specific patient-derived tumor cells.

The quantification of drug response is fundamental to the discovery of therapeutic molecules, the investigation of their mechanisms of action1–3, and the study of signal transduction, cell division, and other biological processes using chemical biology approaches4,5. In the case of anticancer drugs, cells are typically exposed to drugs over a range of concentrations, and the number of viable cells (or surrogates, such as ATP level assayed using CellTiter-Glo, CTG) is measured several days later. Data comprising cell counts in the presence of drug divided by counts for untreated controls are fitted to a sigmoidal curve to compute (i) the concentration of drug at which the cell count is half the control (IC50), (ii) the fraction of viable cells at the highest drug concentration (Emax), and (iii) the area under the dose–response curve (AUC)6,7. Dose–response and genomic datasets are often combined to discover drug–response biomarkers1–3,8,9, but it has recently been found that large-scale drug–response data vary from one study to the next10 for reasons that remain poorly understood11–13.

We show here that, for dividing cells, traditional drug–response metrics such as IC50 suffer from a fundamental flaw when they are estimated from cell counts made at the end of the experiment (the standard approach): if control cells undergo different numbers of divisions during the course of an assay because of natural differences in proliferation rate, variation in growth conditions, or changes in the duration of an experiment, IC50, Emax, and AUC values will vary dramatically, independently of any changes in the underlying biology. Thus, biomarkers that predict sensitivity under one (potentially arbitrary) set of assay conditions may not predict sensitivity under slightly different conditions. We therefore propose a new method for parameterizing drug response, the normalized growth rate inhibition (GR), which is based on the comparison of growth rates in the presence and absence of drug. Parameterization of GR data yields GR50, GRmax, GRAOC and hGR (Hill slope), values that are largely independent of cell division rate and assay duration (we use ‘area over the curve’; GRAOC, rather than AUC for reasons discussed in Online Methods). GR metrics can be determined with modest changes in experimental procedures, and we propose that these metrics replace IC50 and Emax values in assessing cellular response to drugs, RNAi, and other perturbations in which control cells divide over the course of the assay.

RESULTS

Definition of normalized growth rate inhibition (GR)

We used computer simulation to model the drug response of three idealized cell lines with identical sensitivity to a cytostatic drug (i.e., a drug that arrests but does not kill cells) and different division times (Td = 1.8, 2.4, or 3.9 d). These division times correspond to the lower quartile, median, and upper quartile for breast cancer cell lines3 and are similar to the division times of NCI-60 cells14. In the slowly dividing cell line (Td = 3.9 d), the total number of cells did not double in an assay typically run over 3 days, and thus Emax was ≥0.5, and IC50 was undefined. In the case of the two faster-growing cell lines, IC50 and Emax values fell as division rate increased (Fig. 1a) because cell number (or CTG value) was normalized to a drug-naïve control in which cell number increased as division time fell (compare curves across panels of Fig. 1a).

We can compensate for the confounding effects of division rate on drug-response measurements by computing the GR value at time t in the presence of drug at concentration c:

$$\text{GR}(c, t) = 2^{k(c,t)/k(0)} - 1$$

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where $k(c,t)$ is the growth rate of drug-treated cells and $k(0)$ is the growth rate of untreated control cells (Fig. 1b). The GR value is simply the ratio between growth rates under treated and untreated conditions normalized to a single cell division. The sign of the GR value relates directly to response phenotype: it lies between 0 and 1 in the case of partial growth inhibition, it equals 0 in the case of complete cytostasis, and it lies between 0 and −1 in the case of cell death. Given GR values for a range of drug concentrations, GR50 is the concentration at which GR(c) = 0.5, GRmax is the maximal measured GR value, and $h_{GR}$ is the slope of the sigmoidal fit; $GR_{AOC}$ is calculated by integrating the GR curve over a range of concentrations (Online Methods).

In practice, GR values can be estimated from endpoint measurement of cell number in treated and untreated samples, given the initial cell number (Fig. 1c; this is related to the procedure for GI50 determination, see Supplementary Note). Alternatively, the doubling time for untreated cells can be measured under the same conditions in parallel experiments and used in place of the initial cell number (Online Methods). A time-dependent GR value can be evaluated given cell count measurements at two or more time points. Time-dependent GR values capture adaptive responses, varying kinetics of drug–target interaction, drug efflux, etc. (Fig. 1d). Introducing time as a variable makes it possible to relate drug-induced changes in cell states to dynamic measures of drug response at a molecular level (equations for all calculations are provided in Online Methods with links to scripts).

To compare GR dose–response curves to conventional curves, we created synthetic data for cells that had $T_d$ ranging from 1 to 4 d and that were exposed to a drug that was partially cytostatic, fully cytostatic, or cytotoxic (models are described in Online Methods). For all three drugs, IC50 and Fmax values were strongly correlated with division time and assay duration, but this was not true of corresponding GR metrics (Fig. 1e,f, Supplementary Fig. 1a.b, and Online Methods). In the case of drugs that kill cells rapidly and independently of cell cycle state, GRmax still varies with growth rate, and time-dependent GR values are preferable (Supplementary Fig. 2 and Online Methods). AUC combines IC50 or Fmax data and is less sensitive to experimental noise1,6,11, but it suffers from the same dependence on cell division time; $GR_{AOC}$ corrects for this (Supplementary Fig. 1c).

**GR metrics are robust to experimental variability** To study how changing cell division affects IC50 values, we expressed BRAFV600E in hTERT-immortalized retinal pigment epithelial (RPE) cells and then exposed them to etoposide, a topoisomerase II inhibitor that has a cytostatic effect in RPE cells (and whose mechanism of action is independent of BRAF). Oncogene overexpression is known to slow the growth of non-transformed cells15,16, and expression of BRAFV600E in RPE cells under the control of a doxycycline-inducible promoter was observed to increase division time three-fold as the doxycycline concentration increased from 0 to 60 nM. Under these conditions the estimated IC50 value for etoposide increased 100-fold and Fmax increased from 0.25 to 0.6 (Fig. 2a and Supplementary Fig. 3a). In contrast, GR50 and GRmax values varied only slightly under these conditions because the effects of etoposide per cell division were unchanged.

In a second experiment, we measured etoposide sensitivity in MCF 10A cells grown in serum-free medium supplemented with different concentrations of epidermal growth factor (EGF).
The division time of MCF 10A cells varied from >10 d at 0.02 ng ml⁻¹ EGF to <2 d at 1 ng ml⁻¹ EGF (Fig. 2c). Across this range, IC₅₀ and Eₘₐₓ for etoposide were substantially more variable than GR₅₀ or GR₉₅ values (Fig. 2b and Supplementary Fig. 3b). Thus, potentially arbitrary and unintended differences in cell culture conditions can lead to large variation in IC₅₀ and Eₘₐₓ values that is unlikely to have a biological basis.

To test the GR approach in a typical small-scale drug sensitivity screen, we exposed MCF 10A and BT-20 cells (expressing H2B-mCherry to facilitate automated cell counting) to five different drugs chosen for diverse mechanisms of action. We measured cell number approximately every 8 h over 3 d using an automated microscope. Estimated IC₅₀ and Eₘₐₓ values converged only after three divisions (~60 h), whereas GR metrics stabilized by the first division (<20 h for MCF 10A and ~36 h for BT-20; Fig. 2c and Supplementary Fig. 3c). This confirms that GR metrics are substantially less dependent on assay duration than are IC₅₀ and Eₘₐₓ. In the case of very slow and uneven growth (by primary human tumor cells, for example), the stabilization of GR values within one cell division is likely to be a real advantage in obtaining reliable estimates of drug sensitivity.

When grown in 3D culture, MCF 10A cells are known to adapt to inhibitors of PI3K and mTOR (such as omipalisib), becoming less sensitive over time¹⁷. Endpoint measures of drug sensitivity do not report on such adaptive responses. However, when we monitored spheroids by live-cell imaging, we observed that the time-dependent GR and GR₅₀ values were lowest ~20 h after omipalisib addition, and they increased ~10-fold by day 4 (Fig. 2d); endpoint GR₅₀ values lay midway in this range (Fig. 2d). Thus, time-dependent GR data directly capture the decreasing effectiveness of omipalisib in MCF 10A spheroids, enabling detection and further analysis of adaptive mechanisms.

**Analysis of high-throughput data using GR metrics**

The majority of large-scale drug-response datasets published to date neither report cell division rates nor allow one to estimate them. An exception is a study by Heiser et al.³, which recorded cell numbers for breast cancer cell lines before and after exposure to a panel of anticancer drugs for 3 days. For many drugs in this dataset, IC₅₀ correlates with division rate⁶ (e.g., for cell cycle inhibitors, regression coefficient of −0.54, Spearman’s P-value < 10⁻⁶⁶, N = 2,956;
Figure 3 | Evaluation of GR metrics in a high-throughput dataset. (a,b) Fitted dose–response curves for (a) relative cell count and (b) GR values for paclitaxel in breast cancer cell lines (3-day assay data from Heiser et al.3). Red denotes cytotoxic response, and blue denotes cytostatic response; darker curves denote cell lines with fewer divisions. Marginal distributions (below and to the right of the dose–response plots) show the relationship between estimated IC50 or GR50 and Emax or GRmax values and number of divisions for that line over 3 d; p-values are Spearman’s correlation coefficients. (c) Number of divisions over 3 d for cell lines in the dataset grouped by subtype: HER2-amp (HER2amp), triple-negative breast cancer (TNBC), hormone receptor positive (HR+), and nonmalignant (NM). (d,e) Distribution of IC50 (green) and GR50 (purple) for all drugs in Heiser et al.3 (d) or for ErbB2 inhibitors only (e) grouped by clinical subtype. P-values were derived from a rank-sum test for IC50 or GR50 values for NM cell (d) or for HER2amp cell lines (e) versus all other subtypes.

Supplementary Fig. 4a). However, we could reproduce this correlation by using an idealized model that does not assume any biological connection between drug sensitivity and division rate and by repeatedly simulating drug responses using random parameters (Supplementary Fig. 4b and Online Methods). This finding suggests that the correlation between drug sensitivity and division rate found in experimental data is spurious; also, the correlation was absent when drug response was measured using GR50 (Spearman’s P-value = 0.31, Supplementary Data 1), suggesting that it is an artifact of the way IC50 is calculated.

Paclitaxel, a taxane microtubule inhibitor widely used in chemotherapy, is one drug that exhibits a strong negative correlation between sensitivity and division rate as well as substantial variation (100-fold) in IC50 across cell lines (Fig. 3a). This relationship has previously been described and is thought to arise because paclitaxel acts primarily on mitotic cells, and the faster a cell line divides, the more likely it is to be in mitosis18,19. However, reanalysis of data in Heiser et al.3 shows that GR50 values for paclitaxel actually span a narrow range centered around 10 nM (Fig. 3b and Supplementary Fig. 4c), close to the estimated affinity of paclitaxel for assembled microtubules in vitro20. In contrast, GRmax values for paclitaxel vary considerably across cell lines and subtypes. In HER2-amp (HER2amp) and triple-negative breast cancer (TNBC) lines GRmax values are negative, which is indicative of a cytotoxic response, whereas hormone receptor-positive (HR+) and nonmalignant lines generally exhibit cytostatic responses (Supplementary Fig. 4c). This is consistent with data showing that HER2amp and TNBC (basal-like) human tumors are more taxane-sensitive than tumors in other types of breast cancer21. We propose that future attempts to find biomarkers predictive of paclitaxel response focus on variation in GRmax rather than variation in IC5022,23.

The average division rate of breast cancer cell lines in culture differs with clinical subtype: among cells studied by Heiser et al.3, HR+ and HER2amp subtypes divided most slowly (median Td = 3.2 d), TNBC cells divided faster (median Td = 2.3 d), and nonmalignant cells divided still faster (median Td = 1.8 d, Fig. 3c). As measured by IC50 values, nonmalignant cells were, on average, more sensitive to anticancer drugs than tumor cells, whereas GR50 values showed that the mean and range of drug sensitivity was similar (Fig. 3d). Focusing on HER2amp lines, IC50 values for inhibitors of EGFR and ErbB2 were similar across breast cancer subtypes, even though HER2amp human tumors are preferentially sensitive to such drugs and ErbB2 inhibitors are frontline therapy for this disease24,25. GR50 data for HER2amp cell lines in Heiser et al. show that this subtype of breast cancer is ~10-fold more sensitive than other subtypes to EGFR/ErbB2 inhibitors in vitro (Fig. 3e; P = 1.3 × 10−4). The failure of IC50 values to show the preferential sensitivity of HER2amp cell lines to EGFR/ErbB2 inhibitors arises because their relatively slow growth rate is a hidden confounder in IC50 calculation. From these data we conclude that artefactual dependency of IC50 on cell division rate creates associations where none exist and also obscures meaningful associations between genotype and drug sensitivity.

Supplementary Fig. 4c) and number of divisions for that line over 3 d; p-values are Spearman’s correlation coefficients. (c) Number of divisions over 3 d for cell lines in the dataset grouped by subtype: HER2-amp (HER2amp), triple-negative breast cancer (TNBC), hormone receptor positive (HR+), and nonmalignant (NM). (d,e) Distribution of IC50 (green) and GR50 (purple) for all drugs in Heiser et al.3 (d) or for ErbB2 inhibitors only (e) grouped by clinical subtype. P-values were derived from a rank-sum test for IC50 or GR50 values for NM cell (d) or for HER2amp cell lines (e) versus all other subtypes.

Figure 4 | Plating density affects division rate and drug sensitivity. (a) Spearman’s correlation between estimated values for IC50, Emax, GR50, and GRmax and seeding number for the six breast cancer cell lines shown. Data derive from an experiment in which cells were plated at a range of six densities and treated with 11 drugs having diverse mechanisms of action (Online Methods and Supplementary Data 2). Significance: *P < 0.05, **P < 0.01, ***P < 0.001. (b) Relationship between seeding number and the number of divisions in 3 d. Error bars are the s.e.m. of five to nine biological replicates.
**Effects of cell density on drug sensitivity**

GR metrics allow us to quantify how drug sensitivity changes in the face of variables that affect division rate. One such variable is seeding density: increasing density has widely been reported to promote drug resistance. To investigate this, we cultured six breast cell lines representative of different subtypes at six seeding densities over a 32-fold range. 24 h after plating, cells were exposed to 11 drugs with diverse mechanisms of action. Growth rates, IC$_{50}$, $E_{\text{max}}$, and GR metrics were estimated by imaging and counting fixed cells at the time of drug addition and 72 h after treatment (Supplementary Data 2). Overall, IC$_{50}$ and $E_{\text{max}}$ correlated positively with seeding density for MCF 10A, MDA-MB-231, and Hs 578T cells, and they correlated negatively for BT-20 and SK-BR-3 cells ($p < 0.01$, Fig. 4a). In the first three cell lines, division rate decreased with density, presumably because of contact inhibition, depletion of essential medium components, and other effects (Fig. 4b).$^{30–32}$ In BT-20 and SK-BR-3 cells, division rate increased with density, presumably because of conditioning of the growth medium.$^{31,33}$ Correlations between density and IC$_{50}$ and $E_{\text{max}}$ were weakest for MCF7 cells, which grew equally well across all densities. Thus, the effect of density on number of divisions, and consequently on IC$_{50}$ and $E_{\text{max}}$, varied dramatically among cell lines (Supplementary Fig. 5a). However, only a small number of cell line–drug pairs exhibited a statistically significant association between plating density and GR$_{50}$ or GR$_{\text{max}}$ values. We wondered whether these were situations in which the biology of drug response was altered by density.

In MCF 10A cells, six drugs were associated with significant variation in GR$_{50}$ or GR$_{\text{max}}$ (or both) across seeding densities; RNA-seq revealed that gene expression in these cells also varied with cell density, with significant enrichment for genes involved in catabolic processes, cellular respiration, and wound responses.$^{30–32,34}$ (Supplementary Fig. 6a). When the data was analyzed by principal-component analysis, the first principal component (which captured 32% of variance) was strongly correlated with the number of cells at the time of collection (Spearman’s $p = 0.98$) and less so with time in culture ($p = 0.57$), suggesting that the cell density at the time of assay and not the history of the culture was the primary determinant of transcriptional state (Supplementary Fig. 6b). Follow-up studies with methotrexate showed that the ratio between cell number and medium volume, and not cell density per se, was the key variable for sensitivity to this drug (this was also true of oligomycin, an inhibitor of ATP synthase; Fig. 5a and Supplementary Fig. 7); since cells grew in a constant volume of medium, GR$_{50}$ was therefore time dependent. Density- (and time-) dependent variation in GR$_{50}$ were also observed for linsitinib, an IGF1R inhibitor currently in Phase II clinical trials (Fig. 5b). Variation in GR$_{50}$ with time and density was reduced by cotreatment with the metalloprotease inhibitor btmastat.

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**Figure 5** Time-dependent GR metrics reveal diverse mechanisms of drug sensitivity and resistance. Evaluation of three selected drugs for which seeding density had a significant effect on GR values in MCF 10A cells based on data summarized in Figure 4. (a) Influence of ratio of cell number to medium volume for two drugs related to metabolism. Time-dependent GR$_{50}$ values for MCF 10A cells at different seeding numbers following exposure to 0.1 µM methotrexate (left). Time-dependent GR$_{50}$ values for MCF 10A cells for methotrexate or oligomycin (middle); values are capped at 30 µM for illustration purposes. GR dose–response curves at 3 d for MCF 10A treated with methotrexate with a constant number of seeded cells but in different volumes of medium (right). Data shown derive from one of two biological replicates. (b) Medium conditioning and adaptive resistance to linsitinib, an IGF1R inhibitor. GR dose–response curves at 3 d for MCF 10A treated with linsitinib at different cell densities (left). Error bars are the s.e.m. of three biological replicates. Time-dependent GR$_{50}$ values for MCF 10A at different seeding numbers following exposure to 20 µM linsitinib in the absence (middle) or presence (right) of 10 µM batimastat, a matrix metalloprotease inhibitor. Data shown are from one of three biological replicates. (c) Paclitaxel-induced cytostasis at low cell density and apoptosis at high cell density. GR values for MCF 10A cells seeded at different densities and treated with paclitaxel for 3 d (left). Error bars are the s.e.m. of three biological replicates. Fraction of MCF 10A cells positive for cleaved caspase-3 at different time points and cell densities following exposure to 0.03 µM paclitaxel (middle). Time-dependent GR$_{\text{max}}$ values for paclitaxel in MCF 10A cells seeded at different densities (right). Data shown are from one of two biological replicates.
batimastat, suggesting a role for autocrine conditioning of the microenvironment in drug response.

In the case of paclitaxel, GR_{50} values remained at ~5–10 nM across plating densities, but GR_{max} was strongly density dependent, varying from ~0 (cytostatic) at low cell densities to negative values (cytotoxic) at higher densities (Fig. 5c, left). Time-dependent GR_{max} reached its greatest negative value at 24 h, concomitant with an increase in the fraction of taxol-treated cells that contain cleaved caspase-3, confirming that negative GR_{max} values corresponded to elevated apoptosis (Fig. 5c, middle and right). Though its molecular basis is unknown, this effect may be one reason for the discrepancy between cell-killing dynamics in low-density culture and in high-density xenograft tumors.

**DISCUSSION**

Accurate measurement of drug sensitivity and resistance is the cornerstone of cancer biology, pharmacology, and of many fundamental studies on cell signaling and cell division. In this paper we demonstrate theoretically and experimentally that variation in division rate seriously confounds existing drug–response metrics. We also show that division rate varies with cell type, medium composition, and seeding density, often in unpredictable ways. Cell division rate slows down in some cell lines as density increases, while it speeds up in others. Such variation can change apparent IC_{50} 100-fold or more and therefore introduce artificial correlations in data, obscure the true effects of drug action, and introduce unknown complications into biomarker discovery. As an alternative, we propose GR metrics that are computed by comparing growth rates in the presence and absence of drug. GR_{50} and GR_{max} are robust to variation in cell division rate and should replace IC_{50} and E_{max} in studies in which control cells divide (including the study of drugs, gene depletion or overexpression, and variation in the extracellular environment). GR_{50} quantifies the potency of a drug on a per-division basis, ensuring that fast- and slow-growing cells with similar biochemical responses to drug are scored equivalently. GR_{max} captures the maximal drug effect on growth rate and differs from E_{max} in that it falls between 1 and ~1, where negative values denote cell death, 0 denotes cytostasis, and positive values denote partial inhibition. GR_{AOC} and h_{GR} values can also be calculated from GR curves; the former is often the most robust metric in the face of experimental noise, and the latter quantifies an important relationship between dose and response that is often neglected.

In the simplest version of our method, GR metrics are computed after measuring cell number or a surrogate (e.g., CTG value) before and after exposure of a culture to varying concentrations of drug or other perturbation for a fixed time (Fig. 1c). When cell division rates under similar culture conditions are known from previous data, only the final cell number is needed (although we believe that before and after data on cell number are a valuable control to ensure data quality). Time-course data make it possible to compute time-dependent GR values and to quantify phenomena such as delayed response, drug adaptation, or variation in the kinetics of drug–target interaction (Fig. 1d). To facilitate the use of GR metrics by others we provide MATLAB and python routines and an online GR calculator (http://www.grcalculator.org/). Moreover, the data in this paper, including the GR_{50} and GR_{max} values for the Heiser et al. dataset, are available online (http://lincs.hms.harvard.edu/).

Large-scale drug-response studies based on existing response metrics are discrepant for poorly understood reasons, raising concerns about the value of drug-response biomarkers. We speculate that comparison of datasets across centers (or even within a center) might be confounded by differences in plating density, growth medium, and other factors that affect cell division rate. It might be possible to correct for this in existing data by computing GR values post facto, but this will require recreating the original assay conditions and then measuring division rates.

Based on the results in this paper, we believe that use of GR metrics in lieu of traditional IC_{50}, E_{max}, or AUC values will improve our ability to identify genes and biological processes responsible for drug sensitivity and resistance. GR metrics decouple any effect that genotype or microenvironment have on division rate from their effect on drug sensitivity. Cell biology studies involving the modification of genes or the microenvironment often result in changes in cell division rate, leading to potentially spurious correlations with drug sensitivity (as illustrated here by oncogene overexpression and changes in EGF levels), and should also benefit from the use of GR metrics. By analogy with antimicrobial susceptibility testing for bacterial infections, it has recently been suggested that cancer therapy might be personalized by screening primary human tumor cells against panels of drugs. Such cells grow slowly and unevenly in culture, making division number a poorly controlled variable. Accounting for such differences using GR metrics should create drug–response data that are more reproducible and useful for optimizing patient therapy.

**METHODS**

Methods and any associated references are available in the online version of the paper.

**Accession codes.** Data are deposited in the Gene Expression Omnibus (GEO) database with accession number GSE80297.

**Note:** Any Supplementary Information and Source Data files are available in the online version of the paper.

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**AUTHOR CONTRIBUTIONS**

M.H., M.N., and P.K.S. conceived this study and wrote the paper. M.H. and M.C. performed the experiments; M.H. conceived GR metrics and performed the computational analyses.

**COMPETING FINANCIAL INTERESTS**

The authors declare no competing financial interests.

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ONLINE METHODS

Metrics of drug response. Determining relative cell counts. Cell counts in the presence of drug are normalized to DMSO-treated controls grown on the same plate under the same conditions. In the current study, relevant conditions include seeding density, the concentration of exogenous growth factors (e.g., EGF in the case of MCF 10A cells), and the concentration of a second drug such as doxycycline or batimastat. For each cell line, drug, and drug concentration, we define the relative cell count as $x(c)/x_{ctrl}$ where $x(c)$ is the count in the presence of drug at concentration $c$ and $x_{ctrl}$ is the 50%-trimmed mean of the count for control cells. Technical replicates are averaged to yield an average relative cell count. We typically collect data from three technical replicates (on three separate plates).

Calculating GR values using endpoint drug-response data. Normalized growth rate inhibition is calculated according to the formula:

$$\text{GR}(c) = \frac{\log_2(x(c)/x_0)}{\log_2(x_{ctrl}/x_0)} - 1 \quad \text{(Fig. 1c)},$$

where $x(c)$ and $x_{ctrl}$ are as described above, and $x_0$ is the 50%-trimmed mean of the cell count from a sample grown in parallel and measured just prior to drug exposure.

Alternatively, the untreated division time $T_d = \ln(2)/k(0)$ can be measured in independent experiments and used in place of the initial cell number ($x_0 = x_{ctrl} \times 2^{-7T_d}$):

$$\text{GR}(c) = \frac{1 + \log_2(x(c)/x_{ctrl})}{T/T_d} - 1,$$

where $T$ is the duration of the assay.

Calculating time-dependent GR values. GR values can be evaluated over a time interval $2 \times \Delta t$ around any time point $t$ based on the equation:

$$\text{GR}(c,t) = \frac{\log_2(x(c+tΔt)/x(c-tΔt))}{\log_2(x(0+tΔt)/x(0-tΔt))} - 1 \quad \text{(Fig. 1d)}.$$

The time-dependent GR values in the current paper were computed with $2 \times \Delta t = 12$ h to 18 h, which corresponds to about half a cell division time.

Curve fitting and estimating drug-response metrics. GR data are fitted to a sigmoidal curve as follows (Supplementary Fig. 8b):

$$\text{GR}(c) = \text{GR}_{inf} + \frac{1 - \text{GR}_{inf}}{1 + (c/\text{GEC}_{50})^{h_{GR}}},$$

where the fitted parameters are:

- $\text{GR}_{inf}$: the effect of the drug at infinite concentration ($\text{GR}_{inf} \equiv \text{GR}$ ($c \rightarrow \infty$)). $\text{GR}_{inf}$ lies between $-1$ and $1$; negative values correspond to cytotoxic responses (i.e., induction of cell death, Supplementary Fig. 8a), a value of $0$ corresponds to a fully cytostatic response (Supplementary Fig. 8b), and a positive value corresponds to partial growth inhibition (Supplementary Fig. 8c).
- $\text{GEC}_{50}$: the concentration at half-maximal effect. To avoid artefacts in curve fitting we constrain $\text{GEC}_{50}$ to be two orders of magnitude higher and lower than the experimentally tested concentration range (in practice, this is usually about $10^{-5}$ to $10^{3}$ μM).
- $h_{GR}$: the Hill coefficient of the fitted curve, which reflects how steep the dose–response curve is. In practice, we typically constrain $h_{GR}$ to a value between 0.1 and 5.
- $\text{GEC}_{50}$: the concentration at half-maximal effect. To avoid artefacts in curve fitting we constrain $\text{GEC}_{50}$ to be two orders of magnitude higher and lower than the experimentally tested concentration range (in practice, this is usually about $10^{-5}$ to $10^{3}$ μM).

If the fit of the curve is not significantly better than that of a flat curve (i.e., $\text{GR}(c) = \text{GR}_{inf}$) based on an F-test with cutoff of $P = 0.05$, the response is considered flat and the parameter $\text{GEC}_{50}$ is set to 0 (Supplementary Fig. 8d).

Inferred drug-response metrics. The $\text{GR}_{50}$ value is the concentration of drug at which $\text{GR}(c = \text{GR}_{50}) = 0.5$. If the value for $\text{GR}_{inf}$ is above 0.5, the $\text{GR}_{50}$ value is not defined and is therefore set to $+\infty$ (Supplementary Fig. 8c). By extension, other thresholds can be defined in a similar manner. For example, $\text{GR}_{100}$ corresponds to the concentration at which a drug is fully cytostatic: $\text{GR}(c = \text{GR}_{100}) = 0$.

The $\text{GR}_{max}$ is the maximum effect of the drug at the highest tested concentration, and it lies between $-1$ and $1$; a value of $0$ corresponds to a fully cytostatic response, and a negative value corresponds to a cytotoxic response. $\text{GR}_{max}$ can be estimated from the fitted curve or obtained directly from experimental data; we often do the latter.

For time course data, all metrics are evaluated at each time point individually.

Area under the curve and over the curve ($\text{GR}_{AOC}$). Another common metric for quantifying dose–response data is the area under the response curve ($\text{AUC}$), which is based on integrating the dose–response curve over the range of tested concentrations. In the case of GR curves, which can have negative values, it is more intuitive to use the area over the curve:

$$\text{GR}_{AOC} = \int 1 - \text{GR}(c) dc \equiv 1 - \sum c_i \text{GR}(c_i),$$

where $\text{GR}(c_i)$ are measured GR values at discrete concentrations $c_i$. $\text{GR}_{AOC}$ has the benefit that, in the case of no response, it has a value of $0$. The $\text{GR}_{AOC}$ can be normalized to the range of concentrations as, for example, $\text{GR}_{AOC}/\log_{10}(c_{max}/c_{min})$, where $c_{max}$ and $c_{min}$ are the highest and lowest tested concentrations. It is important to note that $\text{GR}_{AOC}$ values (like conventional $\text{AUC}$) should only be used to compare responses evaluated across the same drug concentration range.

The $\text{GR}_{AOC}$ value captures variation in potency and efficacy at the same time. The calculation of $\text{GR}_{AOC}$ at discrete (experimentally determined) concentrations has the advantage that it does not require curve fitting and is therefore free of fitting artifacts. This is especially useful for assays where fewer than five concentrations are measured, and curve fitting is unreliable. $\text{GR}_{AOC}$ values are also more robust to experimental noise than metrics derived from curve fitting; $\text{GR}_{max}$ values are particularly sensitive to outlier values when directly obtained from data.
Drug concentration range. The drug concentrations used for fitting drug-response curves need to span a sufficiently wide range and have sufficiently intermediate values in order to obtain reliable estimates for GEC50, kGR, and GRinf. Denser sampling provides more precise estimates, especially in the case of steep dose–response curves. Optimal design of dose–response curves has been discussed elsewhere. In practice, we suggest using nine doses spanning four orders of magnitude from 1 nM to 10 μM. This range can be shifted to lower concentrations for more potent drugs or to higher concentrations for less potent drugs with the caveat that GR_AOC values for different drugs should be compared only if evaluated over the same concentration range. We suggest discarding any GR50 value that is more than an order of magnitude above the highest tested concentration because values extrapolated from the fitted curves are more subject to fitting artefacts than interpolated values. Similarly, GR_inf value is not properly constrained if the GR(c) dose–response curve does not reach a plateau at the highest measured concentration. In such cases, GR_AOC is the most reliable metric.

Computing GR metrics. Source code for computing GR metrics is provided (Supplementary Software). To facilitate the computation of GR metrics we provide updated source code available under an open source software license and MATLAB and python scripts at https://github.com/sorgerlab/gr50_tools. We also provide an online calculator at http://www.grcalculator.org. This website contains a user guide, various tutorials and explanatory materials, and example datasets, including all of the data in the current manuscript.

Theoretical model of drug response. To simulate the effect of division time on GR and conventional drug-responses metrics under different assumptions about the degree of cytostasis or cell killing, we developed a theoretical model of drug response. To the first approximation, cell growth can be considered exponential, kL

where x is the cell count, k is the untreated growth rate (per day), c is the drug concentration, SM is the maximal inhibitory effect, S50 is the concentration at half-maximal effect of drug, and h is the Hill coefficient. The growth rate k corresponds to the division rate k0 as k = ln(2) × k0 = ln(2)/T_d, where T_d is the division time. SM can be larger than 1 to account for drugs inducing cell death at a specific phase of the cell cycle. The model can also be generalized to account for drugs that induce cell death independent of the cell cycle:

\[ \dot{x} = x \times k \left(1 - \frac{S_M \times c^h}{S_{50}^h + c^h}\right) - x \times \frac{k_L \times c^h}{L_{50}^h + c^h}, \]

where k_L is the maximal killing rate (per day) and L50 is the concentration of drug that produces half-maximal cell killing.

Integrating these equations for an assay of t days yields the cell count at concentration c:

\[ x(c, t) = x_0 \times \exp \left(t \times k \left(1 - \frac{S_M \times c^h}{S_{50}^h + c^h}\right) - t \times \frac{k_L \times c^h}{L_{50}^h + c^h}\right) \]

where x_0 = x(t = 0) is the cell number at the time of treatment. Thus, the relative cell count is:

\[ IC(c, t) = x(c, t)/x_{ctrl} = \exp \left(-t \times k \frac{S_M \times c^h}{S_{50}^h + c^h} - t \times \frac{k_L \times c^h}{L_{50}^h + c^h}\right), \]

and the normalized growth rate inhibition (GR value) is:

\[ GR(c) = \left( \frac{\log_2 x(c, t)}{\log_2 x_{ctrl}} \right) - 1 = 2^{\frac{x(c, t)}{x_{ctrl}}} \left(1 - \frac{S_M \times c^h}{S_{50}^h + c^h} \times \frac{k_L \times c^h}{L_{50}^h + c^h}\right) - 1. \]

This equation for GR(c) is independent of the length of the assay t and, thus, the metrics GR50, GRmax, GR_AUC, and h_GR are also independent of t. For cases where drug action is mainly related to the cell cycle (k_L = 0), GR values are also independent of the untreated growth rate k. As shown in Supplementary Figures 2 and 3a,b, the impact of k_L > 0 is minimal on GR50 and relatively small on GRmax. This is also illustrated by the analytical formula for GR_inf: GR_inf = 2^(1 - S_M - k_L/k) - 1. Note that the metrics derived from growth inhibition (GI) values used in Heiser et al, such as GI50, are more robust than traditional metrics, but they still depend on both division time and assay length (see Supplementary Note).

Parameters for simulations in figure panels. Model parameters used in the numerical simulations shown in the main and supplementary figures (Fig. 1b; Supplementary Figs. 1, 2, and 7) were as follows:

- Cytostatic: S_M = 1, S_{50} = 1.5, T_M = 0, h = 1.6
- Partial response: S_M = 0.65, S_{50} = 1.2, T_M = 0, h = 1.6
- Cytotoxic: S_M = 2.6, S_{50} = 2, T_M = 0, h = 1.6
- Partial toxic response: S_M = 0.45, S_{50} = 1.2, T_M = 0.05, T_{50} = 5, h = 1.6
- Mixed response: S_M = 0.65, S_{50} = 1.2, T_M = 0.1, T_{50} = 3, h = 1.6
- Complete cytotoxic: S_M = 2.6, S_{50} = 1.2, T_M = 1, T_{50} = 3, h = 1.6

For Supplementary Figure 3b, the parameters were randomly sampled within the following distribution:

- Division rate k: normal distribution around 0.9 divisions per day with s.d. of 0.46 and a lower bound of 0.14
- Hill coefficient h: uniform distribution between 1.5 and 2.5
- Maximum inhibition S_M: uniform distribution between 0 and 2
- Half inhibition concentration S_{50}: log-uniform distribution ranging from 0.31 to 10

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Experimental methods and data processing. Cell lines and tissue culture. MCF 10A, Hs 578T, MDA-MB-231, MCF7, SK-BR-3, and BT-20 were obtained from the ATCC and grown according to ATCC recommendations. For time-lapse experiments, MCF 10A and BT-20 cells were modified by inserting an H2B-mCherry expression cassette (gift of R. Benezra, Addgene plasmid # 20972) into the AAVS1 safe harbor genomic locus using CRISPR/Cas9. MCF 10A-H2B-mCherry cells were grown in the same manner as the parental strain, with the exception that traditional DMEM was replaced with FluoroBrite DMEM (Thermo Fisher Scientific) for imaging. The modified hTERT RPE-1 cells (gift from J. Chen) were created by inserting the full-length BRAF V600E expression cassette (Addgene plasmid # 22072) into the AAVS1 safe harbor genomic locus.

Evaluating drug sensitivity in MCF 10A spheroids over time. MCF10 A-H2B-mCherry cells were plated at 200 cells per well into ultralow attachment-coated, flat-bottom, 384-well plates (Corning) with the addition of 2.5% growth factor reduced basement membrane matrix Matrigel (Corning) to the growth medium. After 48 h, cells were treated with a dilution series of omipalisib using a D300 Digital Dispenser (Hewlett-Packard) and imaged in an IncuCyte ZOOM live-cell imager (Essen Bioscience) for an additional 96 h. Drug sensitivity was evaluated by computing GR values every 15 h over a 10–90 h time frame.

Evaluating drug sensitivity in breast cancer cells plated at different seeding densities. MCF 10A, Hs 578T, MDA-MB-231, MCF7, SK-BR-3, and BT-20 were plated at densities ranging from 150 to 5,000 cells per well in 384-well plates using the Multidrop Combi Reagent Dispenser (Thermo Scientific) and grown for 24 h. Cells were treated with a dilution series of the indicated drugs using a D300 Digital Dispenser (Hewlett-Packard). Cells were stained and fixed for analysis at the time of drug treatment and after 72 h of incubation with drug. For these experiments, we used the following drugs:

- Etoposide, topoisomerase inhibitor
- Linsitinib, IGF1R inhibitor
- Omipalisib/GSK2126458, panPI3K/mTOR inhibitor
- PLX4720, B-RAF inhibitor
- Tanespimycin/17-AAG, HSP90 inhibitor

Manipulating cell growth rate to determine effects on drug sensitivity. RPE-1 or MCF 10A-H2B-mCherry cells were plated in 384-well plates using the Multidrop Combi Reagent Dispenser (Thermo Scientific) at 250 and 500 cells per well, respectively. To modulate the growth rate in RPE-1 cells, we induced expression of the BRAFV600E oncogene by treating cells with indicated doses of doxycycline using a D300 Digital Dispenser (Hewlett-Packard). To modulate the growth rate in MCF 10A-H2B-mCherry we serum-starved cells twice with DMEM/F12 medium supplemented with 0.1% bovine serum albumin and 1% penicillin-streptomycin. Medium changes and cell washing were performed using an EL406 Microplate Washer Dispenser (BioTek). Cells were treated with indicated doses of human epidermal growth factor (EGF, Peprotech) using a D300 Digital Dispenser. After 24 h the cells were treated with a dilution series of etoposide using a D300 Digital Dispenser. RPE-1 cells were stained and fixed for analysis at the time of drug treatment and after 72 h. MCF 10A-H2B-mCherry cells were imaged in an IncuCyte ZOOM live-cell imager (Essen Bioscience) starting at the time of EGF treatment, and drug sensitivity was evaluated 72 h after drug addition.

Evaluating drug-response metrics in MCF 10A and BT-20 over time. MCF10 A-H2B-mCherry and BT-20-H2B-mCherry cells were plated at 1,250 and 2,500 cells per well, respectively, in 384-well plates using the Multidrop Combi Reagent Dispenser (Thermo Scientific) and grown for 24 h. Cells were treated with a dilution series of the indicated drugs using a D300 Digital Dispenser (Hewlett-Packard) and imaged after drug addition in an Operetta High-Content Imaging System (PerkinElmer) equipped with a live-cell chamber over a period of 72 h. For these experiments, we used the following drugs:

- Etoposide, topoisomerase inhibitor
- Linsitinib, IGF1R inhibitor
- Omipalisib/GSK2126458, panPI3K/mTOR inhibitor
- PLX4720, B-RAF inhibitor
- Tanespimycin/17-AAG, HSP90 inhibitor

Investigating density-dependent drug effects. MCF 10A-H2B-mCherry cells were plated at densities that ranged from 150 to 5,000 cells per well in 384-well plates using the Multidrop Combi Reagent Dispenser (Thermo Scientific) and grown for 24 h. Cells were treated with a dilution series of drugs using a D300 Digital Dispenser (Hewlett-Packard) and imaged after drug addition in an Operetta High-Content Imaging System (PerkinElmer) equipped with a live-cell chamber over a period of 72 h. In the case of methotrexate and oligomycin, 1,250 cells were plated in 20–120 µL of medium per well, treated with a dilution series of drug, and imaged for 72 h.
In the case of linsitinib, cells were treated with a dilution series of linsitinib either with or without 10 µM batimastat using a D300 Digital Dispenser and imaged in an IncuCyte ZOOM live-cell imager (Essen Bioscience) for an additional 72 h.

In the case of paclitaxel, cells were treated with a dilution series of paclitaxel and 200 nM of NucView 488 caspase-3 substrate (Biotium) using a D300 Digital Dispenser (Hewlett-Packard) and imaged after drug in an IncuCyte ZOOM live-cell imager (Essen Bioscience) for an additional 72 h.

For immunofluorescence experiments, cells were grown for 24 h and then treated with a dilution series of paclitaxel using a D300 Digital Dispenser (Hewlett-Packard) and incubated for 3, 6, 12, and 24 h. Cells were fixed for 30 min in 3% formaldehyde, permeabilized for 30 min in PBS with 0.3% Triton X-100 (Sigma-Aldrich), washed twice in PBS with 0.1% Tween 20 (Sigma-Aldrich; PBS-T), and blocked for 60 min with Odyssey blocking buffer (LI-COR Biosciences). Anti-active Caspase-3 antibody (BD Biosciences) was diluted 1:1,000 in Odyssey blocking buffer and incubated for 16 h at 4 °C. Cells were washed three times in PBS-T for 5 min and incubated with Alexa Fluor 488-conjugated goat anti-rabbit secondary antibody diluted 1:1,000 in Odyssey blocking buffer and incubated for 16 h at 4 °C. Cells were washed two times in PBS-T, washed once with PBS, stained for 30 min with whole cell stain (Thermo Fisher Scientific) and Hoechst (Thermo Fisher Scientific), and washed three times in PBS.

Fixed-cell endpoint assays. After drug treatment, cells were stained at the indicated timepoints with 2 µM Hoechst 33342 (Thermo Fisher Scientific) and LIVE/DEAD Far Red Dead Cell Stain (Thermo Fisher Scientific) for 30 min and fixed with 3% formaldehyde (Sigma-Aldrich) for 30 min. Fixed cells were imaged using an Operetta microscope and analyzed using the Columbus image data storage and analysis system (PerkinElmer).

Live-cell time course assays. Cells expressing H2B-mCherry were imaged at the indicated timepoints after drug treatment in an Operetta High-Content Imaging System (PerkinElmer) equipped with a live-cell chamber or an IncuCyte ZOOM live-cell imager. Dead cells were identified by counterstaining cells with YOYO-1 or TOTO-3 (Thermo Fisher Scientific), and apoptotic cells were identified with the NucView 488 caspase-3 substrate (Biotium). Live, dead, or apoptotic cell lines were identified using the Columbus image data storage and analysis system or the IncuCyte analysis software. Spheroid growth was estimated by measuring the sum of the spheroid area per well using IncuCyte analysis software.

mRNA analysis. mRNA analysis. Cells plated at different densities in 384-well plates were harvested at the indicated times, and RNA was extracted using the RNeasy mini kit (Qiagen). To ensure sufficient amounts of RNA, wells with a low number of cells were pooled. Libraries were prepared by the Broad Technology Labs (BTL), following the protocol for SCRB-Seq described in ref. 41. Transcripts were quantified by the BTL computational pipeline using Cuffquant v.2.2.1 (ref. 42). Analysis of the transcriptional data was performed in MATLAB using standard libraries and inhouse scripts. Gene set enrichment analysis was performed using the GSEA software v2.1.0 from the Broad Institute with the MSigDB v4.0 GO biological process set43.

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