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# *In vitro* evolution and whole genome analysis to study chemotherapy drug resistance in haploid human cells

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# ABBREVIATIONS

AF, allele frequency; CNV, Copy Number Variation; NGS, Next Generation Sequencing, WES, Whole Exome Sequencing; WGS, Whole Genome Sequencing; CML, Chronic Myelogenous Leukemia; IVIEWGA, *In Vitro* Evolution and Whole Genome Analysis; SNV, Single Nucleotide Variant; CNV, Copy Number Variation; TF, transcription factor; DOX, Doxorubicin; GEM, Gemcitabine; ETP, Etoposide; PTX, Paclitaxel; TPT, Topotecan; AML, Acute Myeloid Leukemia; TKIs, Tyrosine Kinase Inhibitors; MDR, Multi-Drug Resistance; gDNA, genomic DNA.

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# ABSTRACT

Background: In vitro evolution and whole genome analysis has proven to be a powerful method for studying the mechanism of action of small molecules in many haploid microbes but has generally not been applied to human cell lines in part because their diploid state complicates the identification of variants that confer drug resistance. To determine if haploid human cell could be used in MOA studies, we evolved resistance to five different anticancer drugs (doxorubicin, gemcitabine, etoposide, topotecan, and paclitaxel) using a near-haploid cell line (HAP1) and then analyzed the genomes of the drug resistant clones, developing a bioinformatic pipeline that involved filtering for high frequency alleles predicted to change protein sequence, or alleles which appeared in the same gene for multiple independent selections with the same compound. Applying the filter to sequences from 28 drug resistant clones identified a set of 21 genes which was strongly enriched for known resistance genes or known drug targets (TOP1, TOP2A, DCK, WDR33, SLCO3A1). In addition, some lines carried structural variants that encompassed additional known resistance genes (ABCB1, WWOX and RRM1). Gene expression knockdown and knockout experiments of 10 validation targets showed a high degree of specificity and accuracy in our calls and demonstrates that the same drug resistance mechanisms found in diverse clinical samples can be evolved, discovered and studied in an isogenic background.

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# INTRODUCTION

In human cells, methods for discovering genes that play a role in drug resistance or which encode drug targets, especially for poorly characterized compounds, such as natural products, are limited. Genome-wide CRISPR-*Cas9* knockdown experiments[1-3] in the presence of a drug are useful to broadly implicate relevant genes, but cannot readily reveal critical gain-of-function, single nucleotide alleles, such as imatinib-resistance conferring mutations in BCR-Abl. Discovering common alleles in whole genome sequences of tumors from cohorts of patients that have relapsed after drug treatment requires very large datasets and is complicated by patient heterogeneity. Furthermore, such studies also cannot be used on experimental therapies.

Work in other organisms has shown that *in vitro* evolution and whole genome analysis (IVIEWGA) is a powerful method to discover both a comprehensive set of drug resistance alleles, as well as the targets of compounds with unknown mechanisms of action[4, 5]. In this method, clonal or near clonal organisms are isolated and then clones are subjected to increasing levels of a drug that inhibits growth. After selection, the organism is cloned again. The genomes of resistant clones are then compared to the sensitive parent clone using next generation sequencing (NGS) methods. In organisms such as *Saccharomyces cerevisiae*[6], *Plasmodium falciparum*[4, 5], Mycobacteria[7], Trypanosomes[6], and *Acinetobacter baumannii*[8] this method has been used to comprehensively discover resistance conferring variants. Surprisingly, the data shows that typically only a small number of *de novo* variants are detected after evolution. If multiple selections are performed on independent clones, the same resistance gene will appear repeatedly, although often appearing with different alleles, providing a high level of statistical confidence that the allele has not arisen by chance.

Many of the organisms on which IVIEWGA has been used with success have both haploid and diploid phases of their lifecycle, which means that selections can be performed in a haploid stage. Selecting for resistant clones in a haploid organism greatly simplifies analysis as a driver resistance allele will approach 100% frequency. In addition, for loss of function alleles, only one mutation is needed (versus both copies). Although metazoans are all diploid, haploid human cells lines are nevertheless available: HAP1, is a human chronic mylogenous leukemia (CML)-derived cell line that is completely haploid except for a 30 megabase fragment of chromosome 15 [9].

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HAP1 has been used for genetic studies because mutated phenotypes are immediately exposed[10-15].

Using five different anticancer drugs (Doxorubicin, Gemcitabine, Etoposide, Topotecan, and Paclitaxel) as examples, we show that *in vitro* evolution in HAP1 cells can be used to identify the molecular basis of drug resistance in human-derived cells. Through our unbiased analysis of evolved clones, we detect a limited number of genes that acquire SNVs or CNVs after prolonged, sublethal exposure to our selected xenobiotics. We further demonstrate the power of the approach by using shRNAs and CRISPR-*Cas9* to downregulate or reintroduce selected alleles and demonstrate that this confers resistance or sensitivity to the drug which elicited the evolved genomic change. Our work has implications for clinical intervention strategies to prevent the emergence of drug resistance and tumor recurrence through gene mutations acquired through DNA damage from chemotherapeutics or natural variants which exist and persist from the heterogenous tumor cell environment.

# RESULTS

#### Selection of compounds for resistance studies

To identify xenobiotics with the best efficacy against HAP1 cells we first measured ATP levels (CellTiterGlo) treating HAP1 cells with serial dilutions of 16 different drug for 48 hours. Five drugs showed EC<sub>50</sub> values between 5 to 340 nM (**Fig. 1A-B, Table S1**). These included doxorubicin (DOX, EC<sub>50</sub> = 46.05 ± 4.6 nM), also known as adriamycin, an anthracycline antibiotic that works by inhibiting topoisomerase II alpha (*TOP2A*)[16, 17]; gencitabine (GEM, EC<sub>50</sub> = 8.7 ± 0.7 nM), a synthetic pyrimidine nucleoside prodrug that is used against a variety of hematopoietic malignancies[18-20]; etoposide (ETP, EC<sub>50</sub> = 338.6 ± 39.7 nM), a semisynthetic derivative of podophyllotoxin that also targets *TOP2A* and prevents re-ligation of the double-stranded DNA[21]; paclitaxel (PTX, EC<sub>50</sub> = 17.5 ± 4.0 nM) also known as taxol, an effective anticancer agent that targets tubulin, perturbing the cytoskeleton and causing M phase cell-cycle arrest[22], and topotecan (TPT, EC<sub>50</sub> = 5.6 ± 0.1 nM), a semisynthetic water-soluble derivative of camptothecin that inhibits topoisomerase I (*TOP1*)[23]. Our HAP1 EC<sub>50</sub> values were similar to those previously reported for other CML cell lines (<u>www.cancerrxgene.org [24, 25]</u>) with the with the exception of etoposide, which appeared more effective in HAP1 cells (EC<sub>50</sub> = 338.6 ± 39.7

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nM) relative to other CML cell lines (> 1  $\mu$ M in BV-173, KU812, EM-2, MEG-01, JURL-MK1, KCL-22, RPMI-8866, LAMA-84, K-562).

#### Evolution of resistance is readily achieved for all compounds.

Our next objective was to create drug resistant lines. Although we have had difficulty creating resistant lines for some drugs in some species ("irresistibles"[26]), there is precedent for resistance to the drugs included here[27-29]. To reduce the possibility of selecting mutations that were present in a nonhomogenous population of HAP1 cells and to facilitate later genomic analysis, we first cloned the cells. This was accomplished by diluting cells to an average density of ~0.5 cells per well in a poly-L-lysine treated 96-well plate (**Fig. 1C**) and then picking clones from wells that contained single colonies. Selections were initiated with different parent clones for the different drug-specific replicates (**Fig.1C**, **Fig. S1**).

To create drug resistant clones, cells were grown in tissue culture dishes (reaching 60-80% semi-confluence) in the presence of sublethal concentrations of each drug using one of two different methods. Most cell lines (DOX, GEM, TPT and PTX resistant clones) were subjected to a lethal concentration ( $\sim$ 3-5 × EC<sub>50</sub> value), killing more than 95% of the cells. Then, treatment was removed until cells reached semi-confluence again (doubling every 22 hours[30]) whereupon drug at ~ the EC<sub>95</sub> value was reapplied. Alternatively, for ETP-resistant clones a step-wise selection method was used whereby cells were repeatedly exposed to a concentration that killed around 50% of the cell population. Drug concentration was increased by 5-10% every 5 days while keeping the growth rate at 50% of untreated culture. Although others have used mutagenesis [31], we have found that this can increase the rate of background mutations, which would complicate an already difficult analysis. Because mutations will arise randomly during long term cell culture, we attempted at least three independent selections for each drug, in each case starting with an identical parental isolate (Fig. 1C). In a few cases, independent selections could not be achieved and dependent clones with a shared lineage (DOX-R4a and DOX-R4b; PTX-R2a and PTX-R2b; TPT-R4a, TPT-R4b and TPT-R4c) were collected. Resistance emerged after several months depending on the drug and the method used (7-30 weeks approximately, 49-210 generations) (Fig. S1).

Once resistance was observed in the batch culture, we isolated clones from batch drugselected cultures and the drug sensitivity of the clone was measured and compared to the isogenic parent clones (**Fig. 1D**). We observed an  $EC_{50}$  fold shift between 3.3 and 411.7 (**Table S2**) in

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paired comparisons. To demonstrate that the drug resistance phenotype was stable, drug pressure was removed for 8 weeks (approximately 56 generations) and clones were retested for sensitivity. We observed no changes in the  $EC_{50}$  values, indicating that resistance phenotypes were not due to transient adaptation.

# Identification of putative resistance variants using next-generation sequencing

We next performed whole genome and exome paired-end read sequencing on the 35 cell lines (both drug-resistant clones and their matched drug-sensitive parent clones). Our IVIEWGA studies in *Plasmodium*[5], have shown that stable drug resistance is typically conferred by SNVs in coding regions and thus exome sequencing was an efficient mechanism to find causal variants. However, gene amplifications, which contribute to 1/3 of drug resistance events in *Plasmodium*[5], are more accurately detected with WGS because exact chromosomal recombination sites, which may fall in intergenic regions, can be reconstructed from WGS data. Because of falling costs over the course of the project, more samples (N=21) were ultimately whole genome sequenced than whole exome sequenced (N=14).

Sequencing quality was high for all samples: alignment showed that, on average, 99.9% of 700 million WGS (40 million WES) reads mapped to the hg19 reference genome with 86% of the bases covered by 20 or more reads (**Table S3**). By comparing sequences of evolved clones to their respective parental clones, we discovered a total of 41,259 SNVs (**Table S4**), of which 26,625 were unique (**Table S5**, Methods). The majority of variants in all cell lines was non-coding (**Table S4**, **S5**) and were evenly distributed with respect to chromosome length (**Fig. S2**). Of the 26,625 mutations almost all (26,468) were present at allele frequencies (AF) of less than 85% relative to their parent clone and would thus not be expected to be driver mutations, given that the parents were cloned (to the best of our ability) before selections were initiated. The five drugs varied in the number of mutations, with TPT having the highest overall numbers (**Table 1**).

We next developed a pipeline (**Fig. S3A**, Methods) to filter the 26,625 "called" mutations (**Table S6**) to a final list of potential variants conferring drug resistance (**Table 1**). Our previous analyses in other species suggested that variants presented in coding regions are more likely to contribute to drug resistance even though this could exclude the variants associated with certain transcription factor (TF) binding sites. Therefore, our strategy focused on mutations that were in

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exonic regions and were drug-specific (Fig. S3A). We further considered only mutations likely to have a functional impact at the protein level (missense, nonsense, frameshift, start or stop gain or loss) which further reduced the number to an average of 35 and 23 nonsynonymous mutations for WGS and WES, respectively (Fig. S3A). Reasoning that resistance driver mutations (e.g. those actually causing resistance) would be present in 100% of the haploid cells in the sequenced culture, we selected only the variants with high allele frequency (AF > 0.85, as determined by sequencing read count). The top 2.5% of highest AF mutations corresponded to an AF > 0.85 (Fig S3B). At this cutoff, the majority of cell lines harbored a candidate resistance mutation. While selecting a cutoff represents a tradeoff with potentially missed relevant mutations, the full list of mutations is provided in the supplement (Table S6). We did not note any strong correlation between read depth and allelic fraction in our study (R2 = 0.06; Fig. S3C) and all of the final mutations selected for further analysis had a read depth > 10 reads, with the majority supported by over 20 (Fig. S3D). Although some with AF <0.85 could confer a beneficial advantage to the cell, most are likely to be random mutations that arose during long term culture. Finally, based on our experience with microbes whereupon genes with multiple, predicted independent amino acid changes (not expected by chance (4 genes, STARD9, CYP1B1, SLCO3A1 and DCK)) are often found for the same drug, we added these genes to our final list of 21 candidates (Table S7).

# Somatic Copy number variations (CNVs)

We next searched for CNVs (both amplifications and deletions) in our WGS and WES data using Control-FreeC[32]. Overall patterns of broad and focal alterations across the drugs and conditions varied (**Fig. S4A**, **Table S8**). Using a corrected p-value of less than 0.05, we identified 93 total amplification and 108 deletion events, with most appearing in the TPT-resistant samples (123) (**Table S8**). The CNVs had an average size of 8.5 Mbp (stdev 19 Mbp), ranged from 15,000 bp to 152 Mbp (**Fig. S4A**) and covered ~3% of the genome, on average. More CNVs were called in WES samples because of sequencing gaps—even for WGS samples, some CNVs were separated by short distances and were nearly contiguous (**Fig. S4A**). It is likely that some CNVs were also missed in the WES data. The number of events was proportional to chromosome size, with the exception of the Y chromosome, for which there were ~4 × more events (47) per unit length. Some CNV calls were supported by paired end red data, for example, the one near WWOX (**Fig. S4B**, **C**).

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# Doxorubicin resistance is associated with mutations in *TOP2A* and a solute carrier transporter

To evaluate the approach, we next considered the set of SNVs and CNVs for each drug. For DOX, we had six available selections from two different starting clones (WT-1 (WGS) and WT-5 (WES)) that were analyzed by WGS (DOX-R1, DOX-R2, DOX-R3) and by WES (DOX-R4a, DOX-R4b and DOX-R5) (Fig. 2A). High allele frequency missense mutations were found in only 11 genes (Table S7). Of note, DOX-R2 and DOX-R3 carried mutations in TOP2A at frequencies of 0.89 and 0.87, respectively. This is the known target of DOX[21, 33] and is known to play a role in drug resistance[33-35]. The amino acid mutation, Pro803Thr (Fig. S5), sits within the principal DNA-binding locus, the DNA-gate, a region conserved in type II topoisomerases (TOP2A and TOP2B). It is also adjacent to the catalytic tyrosine (Tyr805), responsible for nucleophilic attack on DNA[36]. While one explanation is that Pro803Thr creates steric hindrance and blocks DOX access to the site, a more likely explanation is that the mutation is a loss-offunction mutation, especially as knockdown of TOP2A activity has previously been shown to confer DOX resistance in a Eµ-Myc mouse lymphoma system[37]. To reproduce these results in our HAP1 human cells, TOP2A was downregulated using a shRNA pool containing constructs encoding target-specific shRNA hairpins for TOP2A. Western blots further showed the expected down regulation of protein levels (Fig. 2B) and an EC<sub>50</sub> analysis of the wildtype and the knockdown line revealed a 4.25-fold increase in DOX resistance compared to the isogenic parent (Fig. 2C, D).

We also found missense mutations present in 100% of the reads for several other attractive but less well characterized genes; *SLC13A4* (Gln165His, DOX-R4b), and *SPG7* (Lys593Asn, DOX-R5), as well as one uncharacterized gene (AC091801.1, His13Asn, DOX-R4a) in the three different clones that were subjected to WES and were derived from WT-5. *SLC13A4* is a solute carrier transport family member and members of this general solute carrier family have appeared in selections conducted in microbes (e.g. the UDP-galactose transporter and the AcetylCoA transporter[38]) and are also associated with cancer drug resistance[39]. The Gln165His mutation is located in the disordered region of the protein. To validate *SLC13A4* we performed a gene knockdown using a shRNA pool that targeted *SLC13A4*, containing three expression constructs each encoding target-specific 19-25 nucleotide shRNA hairpins. Protein expression levels of the

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knockdown line were measured by western blot followed by a dose-response assay to compare its  $EC_{50}$  value with the wildtype line (**Fig. 2E**). The 4 × increase in resistance suggests that *SLC13A4* contributes to resistance, although it may not account completely for the level of resistance of the sequenced clones, which ranged from 4 to 11 × (**Fig. 2F, G**).

#### Gemcitabine resistance is conferred by changes in DCK and RRM1 activity

Six selections were performed with GEM (starting from two different isogenic parents; WT-2 (WGS) and WT-3 (WES)). Among those, three GEM-resistant clones subjected to WGS (GEM-R1, GEM-R2 and GEM-R3) showed an average  $EC_{50}$  shift of 300 to 400-fold (**Fig. 3A**, **Table S2**), and the clones showed no change in HAP1 sensitivity to other drugs (**Fig. 3B**). As there were no candidate alleles with AF > 0.85, we looked for genes that acquired mutations in multiple selections, identifying deoxycytidine kinase (*DCK*) as likely important for resistance. Interestingly, across cell lines several distinct mutations were found in *DCK*, with varying effects (missense and frameshift) across several different positions (**Table 2**). In particular, the missense substitution Ser129Tyr, present in GEM-R1 and GEM-R3, not only alters the amino-acid size and charge also falls at the end of exon 3, within the ATP-binding pocket of a phosphorylation site, making it a strong causal candidate for GEM drug resistance (**Fig. S6**). GEM only becomes pharmacologically active if it is phosphorylated and the first phosphorylation is catalyzed by DCK. A shRNA knockdown of DCK was performed and confirmed by western blot analysis (**Fig. 3C**). Downregulation of the gene resulted in a 36.5-fold increase in the EC<sub>50</sub> value compared to both the isogenic parent line and the shRNA negative control (**Fig. 3D, E; Table 2**).

The three WT-3 derived GEM-resistant clones (GEM-R4, GEM-R5 and GEM-R6) subjected to WES were not as resistant as those used in WGS ( $\sim 6 \times \text{versus} \sim 400 \times$ , Fig. 3F, Table S2). Our work in other species with well characterized compounds suggests this is not surprising and that even single nucleotide changes in the same gene can yield different levels of resistance. For example, repeated selections with dihydroorotate dehydrogenase (DHOD) inhibitors in a mouse model and *in vitro* culture gave rise to 13 different point mutations in parasite DHODH with levels of resistance ranging from 2- to  $\sim$ 400-fold [40]. No high AF SNVs were evident in these lines and *DCK* exons were not mutated. On the other hand, the three WES clones contained 20 CNVs that could play a role in drug resistance. Most CNVs were not shared between lines but GEM-R4, GEM-R5 and GEM-R6 all bore overlapping CNVs of varying sizes on chromosome 11,

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with all three lines bearing 3-4 copies (p value = 1.38e-37 to 2.05e-142) (**Fig. S4**). The chromosome 11 CNV was only found in GEM resistant lines and not in any of the other evolved lines (in contrast to CNVs on chromosome 1 or 16, for example). While it is difficult to determine which of the 140 genes in the smallest interval contribute to resistance, a known resistance mediator or target of GEM, ribonucleotide reductase (*RRM1*), was found within the amplified region. RRM1 is the largest catalytic subunit of ribonucleotide reductase, a key enzyme catalyzing the transformation of ribonucleotide diphosphates to deoxyribonucleoside diphosphates that are required for DNA synthesis and repair, and GEM is known to inhibit DNA polymerase by inhibiting *RRM1*[41]. Furthermore, overexpression of RRM1 is associated with poorer prognosis after gemcitabine treatment in non-small cell lung cancer[42] and in bladder cancer[43].

Western blot analysis of the evolved lines showed that the amplification was indeed associated with increased protein levels (**Fig. 3G**). As an additional validation, we performed a single shRNA knockdown of *RRM1* to reduce protein expression (**Fig. 3H**), followed by a doseresponse assay comparing  $EC_{50}$  values of both wildtype HAP1 and *RRM1* knockdown lines, which showed that downregulation of *RRM1* made HAP1 cells 31-fold more sensitive to GEM than their isogenic parent (**Fig. 3I, J**). As expected *RRM1* downregulation had no effect on HAP1 sensitivity to other drugs (**Fig. S7**).

# Etoposide resistance is modulated by levels of WDR33

We created three independent ETP resistant clones, all of which were subjected to WES, and compared them to one isogenic parent clone (WT-3) (23, 13 and 9-fold increased resistance respectively (**Fig. S8A, Table S2**). A single gene, *WDR33* (ETP-R3), carried a SNV (Pro622Thr) with a 100% allele frequency. This gene encodes for a member of the *WD* repeat protein family and is one of the six subunits of a multiprotein complex called *CPSF* (cleavage and polyadenylation specific factor)[44] involved in cell cycle progression, signal transduction, apoptosis and gene regulation. Disruption of *WDR33* can lead to slowed DNA replication forks[45], which could potentially explain why its disruption protects against topoisomerase inhibitors that block DNA unwinding. Lines in which *WDR33* was knocked down via shRNA acquired an EC<sub>50</sub> value 3.4 times greater than its parental line or the scrambled control (**Fig. S8B-D; Table 2**), despite an incomplete disruption of the gene by shRNA silencing.

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No clear candidate SNVs were evident for ETP-R1 and ETP-R2, which did not carry the *WDR33* mutation (**Table S6, Table S7**). All ETP lines carried multiple CNVs, however, including a large shared amplification on chromosome 15 (ETP-R1 and R3). Approximately 120 protein coding genes are found in this region, including *BUB1B*, the BUB1 mitotic checkpoint serine/threonine kinase B, *BMF*, a BCL-modifying factor, as well as the *RAD51* recombinase, whose overexpression has been previously shown to confer ETP resistance[46]. Overexpression of *RAD51* activity sensitizes cells to a variety of cancer drugs, including etoposide, doxorubicin and topotecan[47]. Notably, all ETP resistant lines were also cross-resistant to PTX, TPT and DOX, providing support for this general mechanism of resistance.

# Paclitaxel resistance is mediated by transporters SLCO3A1 and ABCB1

Seven different paclitaxel lines were created with different resistance levels (PTX-R1, R2a, R2b and R3, ~10 × to PTX-R4, R5, R6, 50X) (**Table S2**). The first four (**Fig. 4A**) were subjected to WGS and the latter three to WES. SNV analysis yielded no candidate genes (frameshift, indels, and missense mutations with an allele frequency >0.85). From genes with an allele frequency of less than 0.85, *SLCO3A1*, encoding another solute carrier transporter, was notable in that multiple missense alleles were identified (Ile587Asn, Ala263Thr). This class of transporter is known to play a role in the import of drugs as well as hormones such as prostaglandin[48]. Gene knockdown experiments showed that clones with loss of SLCO3A1 (**Fig. 4B**) resulted in HAP1 cells that were ~8 times more resistant than their isogenic parents to PTX (**Fig. 4C, D**).

Despite the lack of obvious coding SNVs, PTX-R1, R2a, R2b and R3 had a combined number of 47 CNVs, while PTX-R4, R5 and R6 had 10 (the fact that more CNVs were found in WGS samples may reflect the ease with which CNVs are called with WGS versus WES data). Potentially significant genes with CNVs were *ABCB1* (MDR1) and *ABCB4* (*MDR3*) (**Fig. 4E**) on chromosome 7 (PTX-R2a, R2b). *ABCB1* amplifications are associated with clinical resistance to PTX[49]. PTX-R4 and R5 showed structural variants on chromosome 1, and PTX-R4 show an amplification event on chromosome 17 that encompassed a variety of ABC transporters (*ABCA5*, *6*, *8*, *9*, *10*). No compelling candidate genes were found in CNVs for PTX-R6. On the other hand, inspection with IGV showed that read coverage was poor and that CNVs might not have been detected with WES data.

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To confirm the importance of ABC transporters in PTX resistance, clones were treated with both PTX and verapamil, a calcium channel-blocker which can reverse ABC-transporter mediated resistance [50, 51]. We observed a complete reversal of resistance in PTX lines (**Fig. 4F**). In contrast, we observed no reversal of resistance in GEM lines (**Fig. 4G**), suggesting the resistance role of ABC-transporters is PTX-specific.

# Topotecan resistance is associated with complex alterations in *TOP1*, deletion of *WWOX* and SNVs in cytochrome p450s (*CYP1B1*).

The six TPT samples were derived from four independent selection events (TPT-R4a-c are clones from the same selection with levels of resistance ranging from  $10-20 \times$ ; **Table S2**) and all six clones were subjected to WGS together with their parent clones (WT-6 and WT-7)

For TPT-R4a-c lines (**Fig. 5A**), 268 alleles were present with AF > 0.85, but of these, only six were coding mutations and the rest were intergenic. Three of the six coding mutations were frameshift mutations (His81) with AF = 1 in *TOP1* (**Fig. 5B, S6A**), the known target of topotecan[23]. The His81 frameshift mutation, which introduces a premature stop codon, was confirmed by examining the read alignments (**Fig. S9A**) and by the absence of the full-length protein using N-terminal antibodies (**Fig. 5C**). Because there were also complex structural variants in the region (**Fig 5D, S9B**) we also sequenced the 5' cDNA through the His81 frameshift for all three lines and as well as the parent line and confirmed the two-base deletion in the mutant as well as homozygosity in TPT-R4a-c evolved lines. We also observed a decrease in mRNA expression with TPT-R4a-b showing a statistically significant decrease in TOP1 mRNA expression of Top1 produces resistance to camptothecin, a close analog of topotecan[37]. Interestingly, of the 22 TOP1 frameshift or nonsense mutations in the COSMIC tumor database, 6 were located within a 30 amino acid span (of 765 total) that includes His81 (exon 4), suggesting likely clinical relevance[52]. The probability of this distribution by chance is  $9.65 \times 10^{-5}$ .

No clear coding SNVs with a high allele frequency were obvious in TPT-R1, R2 and R3 but we noted multiple SNVs (Asp217Glu from TPT-R4a,b,c and Val432Leu from TPT-R1) in *CYP1B1*, which encodes a cytochrome p450 isoform. Overexpression of CYP1B1 has previously been associated with TPT resistance[50]. TPT resistant lines (TPT-R1, R2 and R3 (**Fig. 5E**) also showed large chromosomal abnormalities at *WWOX* (**Fig. 5F**) with a clear deletion of the *WWOX* 

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gene region (chr16:78,569,166-78,792,736, exon7 and 8). WWOX bears a well-known fragile site (FRA16D) and encodes a putative oxidoreductase. The complete absence of WWOX protein was confirmed by Western in TPT-R1, 2 and 3 (**Fig. 5G**). Interestingly, lower levels of WWOX were also observed in TPT-R4a-c, which could be a consequence of other *cis* or *trans* variants in this cell line and might also contribute to this level of resistance. Knockdown of *WWOX* by shRNA resulted in marked resistance to TPT (**Fig. 5H, I**). WWOX acts as a tumor suppressor and plays a role in apoptosis. Its disruption may prevent TPT-induced apoptosis, promoting cell survival in the presence of TPT[53]. WWOX disruption also resulted in resistance to PTX (**Fig. 5J**), and as reported by others who examined WWOX-transfected epithelial ovarian cancer cells [54].

# Some evolved mutations are associated with more drug resistance in human cancer cell lines

We further evaluated the association of mutations in our identified genes with drug resistance in cancer cell lines, reasoning that if resistance genes were already mutated before drug exposure, the cell lines would be more resistant. The set of genes we identified had mutations in multiple cancer cell lines [55, 56]. Considering only matched alteration types, we identified cell lines with mutations in the same set of resistance genes (Table S9). We grouped cell lines according to whether or not they had a drug-specific resistance mutation of matched type (SNV or CNV) in any resistance gene found in the HAP1 study and compared areas under the dose-response curve (AUC) between groups [24] (Fig. S11A). All five drugs trended toward higher dose response AUCs when resistance genes were already mutated indicating that they are more resistant, with differences in doxorubicin, gemcitabine and paclitaxel all showing significance after multiple test correction. This comparison is complicated by the fact that not all missense mutations are necessarily functional, thus some cell lines may be included in the mutated category that do not actually have altered protein function. Additionally, mutations and karyotypic abnormalities affecting other genes could also contribute to resistance in each cell line, thus cell lines lacking mutations in certain genes can still be resistant. We therefore attempted to further compare cell lines carrying mutations based on the predicted functional consequences of mutations in the context of specific gene-drug pairs. For some individual pairs, predicted loss of function (LOF) variants tended to have higher dose-response AUCs than variants predicted to have weaker effects on protein activity, in particular, SPG7 and SLC13A4 for doxorubicin, WDR33 for etoposide, and

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CYP1B1 for topotecan (Fig. S11B). Only CYP1B1 met statistical significance at a 0.1 Type 1 error rate.

# DISCUSSION and CONCLUSIONS

Here, we show for the first time that *in vitro* evolution and whole genome analysis (IVIEWGA) can readily lead to the identification of drug resistance mechanisms in human cells. Our results show *in vitro* resistance acquisition and provide a framework for the determination of chemotherapy resistance alleles that may arise in patients.

Our work using IVIEWGA in pathogens (see [55] for a review) guided our pipeline development: We focused on protein coding alterations that arose in association with a single treatment condition, that were nonsynonymous, occurring repeatedly and were high allele frequency. We also removed alleles for genes that are known to mutate frequently, like odorant receptors. Overall, our results are similar to what we have observed in eukaryotic pathogens with a mix of CNVs and single nucleotide variants giving rise to resistance.

Because of the substantially greater costs associated with WGS, here we did evaluate both WES and WGS sequencing methods. Despite a higher likelihood of discovering all changes by WGS, the disadvantage of WGS is cost and computational time. While human WES data can be analyzed on a laptop, human WGS data files are large and difficult to handle, computationally. It has been estimated, considering computational time, that a human genome costs upwards of \$25,000 to fully sequence [56].

The biggest disadvantage of using WES is that CNVs will be harder to call. This is partly for statistical reasons with many reads that support CNV calls located outside of coding regions in WGS samples. In addition, if one sequences over the exact location of the recombination event (or the start or end of the deletion) one can obtain additional support for location calls via split read analysis of paired-end libraries. In addition, one can extract the sequence of the short read and reconstruct the exact recombination breakpoint, as shown in **Fig. S3**. This would not be feasible with whole exome sequencing. Recently it was shown that CNV detection tools perform poorly on WES cancer genome samples. Comparative analysis showed a low consensus in CNV calling tools with moderate sensitivity ( $\sim$ 50% -  $\sim$ 80%), fair specificity ( $\sim$ 70% -  $\sim$ 94%) and poor FDRs ( $\sim$ 27% -  $\sim$ 60%). Also, using simulated data these authors observed that increasing the

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coverage more than 10× in exonic regions did not improve the detection power [57]. Of course, detecting CNVs is likely to be more challenging in diploid genomes, than haploid genomes. In support of this, we were able to identify and validate the RRM1 amplification event in GEM-R4, 5 and 6, which were only subjected to WES. In addition, in yeast, it appears CNVs are much less important than SNVs in driving drug resistance as well: In a more comprehensive *in vitro* evolution study in yeast with 80 different and 355 whole genome sequences we observed only 24 CNVs, including apparent aneuploidy (11 times, occurring in 10 clones) and small, intrachromosomal amplifications (13 times, occurring in 13 clones) in our set of 355 whole genome sequences [58].

A lesser disadvantage of WES, is the rare possibility that resistance is conferred by an intergenic mutation, which would be missed by WES data. Our work in other organisms has shown that almost all resistance conferring SNVs or small indels are nonsynonymous changes that would be detected by both WES and WGS. In the aforementioned yeast study, 271 mutations of the 1405 detected mutations in the 355 evolved lines were intergenic. Of these, only five were directly upstream or downstream of one of the 137 genes that were repeated identified in the study. In contrast to coding mutations, most intergenic mutations lacked any statistical support suggesting relevance and were likely to be background mutations [58]. Despite the lower probability that intergenic or other noncoding mutations may have functional effect, we recognize that there are examples from the literature where intergenic mutations have contributed to drug resistance. Noncoding RNAs such as *EGRF-AS1* and activating cis elements such as enhancers have previously been implicated in evasion of drug response[59-61]. The intergenic mutations with high allele frequency are present in our provided datasets and provide opportunity for reanalysis or for querying by those interested in a specific noncoding RNA or enhancer. It is feasible that even synonymous mutations could confer resistance if they altered the rate of protein folding.

A limitation of our HAP1 study, as presented, and is contrast to our work in other species, is that despite some level of repetition, we seldom achieved strong statistical confidence by just performing selections and sequencing. This may not be unexpected. Evolution is, unfortunately, a relatively stochastic process even when working with the exact same starting clone. In the yeast study [58] we only obtained the same allelic change in the same critical drug resistance gene a few times despite >3 repetitions per each of the 80 compounds. For example, two independent

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selections with hectochlorin both yielded an Arg116Lys in ACT1, the target of hectochlorin [58]. Similarly, a Leu671Phe change in YRM1 was observed 5 times for 4 different compounds.

Another disadvantage of using human cells is the challenge of validation of SNVs; we were not able to engineer any SNVs into HAP1 cells to demonstrate their importance. On the other hand, with the statistical confidence that comes from identifying the same gene repeatedly, CRISPR-*Cas9* validation becomes less important. In the same yeast study described above, YRM1, a gene encoding a transcription factor involved in drug resistance in yeast was independently identified 52 times with 27 different alleles. The likelihood of 355 selections yielding the same gene by chance is roughly  $3.53 \times 10^{-116}$ . This enrichment analysis becomes an attractive method for teasing apart driver and passenger mutations and may become possible with more repetitions despite the larger genome size of HAP1 cells. However, performing enough repetitions to achieve statistical confidence would require substantial resources with WGS. even with a thousand-dollar human genome. WES is thus likely to be more useful.

While HAP1 cells may not be considered a perfect model for human cancer biology, for the purposes of target identification, they are likely very useful. As with pathogens, our use of well-studied drugs, largely uncovered genes that were mostly already well known to confer resistance such as RRM1 [62], [41], DCK [63, 64], TOP2A [37]. and TOP1[37] in a variety of different cancer cell lines. Although it was initially argued that the *in vitro* evolution system might be artificial, in malaria parasites it has been used to discover or rediscover most, if not all (to our knowledge), clinically relevant drug resistance genes including the chloroquine resistance transporter[5], the artemisinin resistance gene, *Pfkelch13*[65], and well-known ABC transporters[5].

Despite questions about how much they mimic human cells, the value of using haploid cell lines is evident from our allele frequency data. If our lines had been diploid and we would have needed to consider allele frequency data of up to 0.4. There are 205 missense mutations with an AF of >0.4, making pinpointing the causative allele much more difficult without candidate genes or without many repetitions. Although *in vitro* evolution has been used repeatedly for discovering the mechanism of action of completely uncharacterized compound in malaria parasites (reviewed in [55]), there are fewer examples of *in vitro* evolution being used for de novo target discovery in diploid eukaryotic pathogens. Although there are some examples in trypanosomes [66-68], some hypothesis about the mechanism of action was already present before evolution studies were

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attempted. Despite this, low allele frequency data should not necessarily be discarded. There have been examples from haploid Plasmodium where a resistance-conferring allele was located within an amplification region and thus showed an AF < 0.5.

Although the HAP1 cells could be considered unnatural, it is likely that similar evolution experiments in other types of human cells will largely give the same genes. This is because conservation of drug targets and drug resistance mechanisms across phyla is often observed, although a given compound or inhibitor may show differences in selectivity and specificity. Resistance to topotecan/camptothecan in yeast is also provided by mutations in Top1[69]. Recent IVIEWGA studies in yeast also identified Top1 as the target in yeast [58]. Evolution studies with cladosporin in yeast and plasmodium both give the same resistance mechanism for cladosporin, lysyl tRNA synthetase [70].

Our studies were not meant to study the process of evolution. Within the field of laboratory-based evolution, there are two broad areas of study. The first are those that fall under the heading of "experimental evolution" and which try to mimic evolution in natural conditions. Here, growth rates are often recorded and experimental conditions may be varied in a controlled manner (carbon sources, temperature, etc). Such studies include long term studies of E. coli or other bacteria(reviewed in [71]) and have also been performed with small molecules [8], [72, 73], primarily with known mechanism of action. Alternatively, there are also studies in which evolution has been used as a tool to discover targets and resistance genes for therapeutic purposes [7, 74] [55]. In many cases [74, 75], although not in all cases the term "in vitro evolution" is used instead of "experimental evolution." Based on our results here, resistance readily emerges in HAP1 cells but more work will need to be done to determine if this is because of the compounds that were used. Here we used in vitro evolution (versus experimental evolution) to select for mutant lines that could withstand treatment with the selected drugs. Although it may be possible to use HAP1 cells for experimental evolution, at present sequencing costs are so high that whole genome studies with whole genome analysis are not practical but this may change in the future. Questions that might be investigated include the fitness of different mutations, reproducibility of the process, impact of the starting clone, carbon sources or growth rate and whether one resistance mechanism predominates or if a variety are found.

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Finally, it is important to keep in mind that the compounds examined here are not modern cancer therapies and while still used clinically, they are imperfect. Newer molecules include bortezomib, a small molecule proteasome inhibitor, imatinib, a small molecule tyrosine kinase inhibitor or seliciclib, small molecule cyclin-dependent kinase inhibitor or even small molecule cancer immunotherapies. We anticipate mutations in the drug's targets will be identified sometimes, as is observed in microbes. In fact, unbiased IVIEWGA studies with bortezomid in P. falciparum have demonstrated mutations in the proteosome subunit, Pf20S  $\beta$ 5, [76] confer resistance, and similar resistance-conferring mutations have been discovered after using in vitro evolution in human cells, although whole genome sequencing was not performed and the mutations were identified using a candidate gene approach [77]. On the other hand, targeted therapies are less likely to work against HAP1 cells, as shown here for imatinib (Fig. S1), most likely because HAP1 cells do not harbor the appropriate sensitizing mutations (e.g. the BRC-Abl for imatinib or BRAF/EGFR mutations for vemurafenib, gefitinib or erlotinib, respectively [78]). Alternatively, the HAP1 cells may be intrinsically resistant because they harbor other resistance conferring mutations. Nevertheless, if they can be used or engineered to sensitivity, predicting resistance mechanisms for new drugs in clinical development, as well as for new drug combinations and may lead to better classes of drugs for chemotherapy.

# MATERIALS AND METHODS

### Compounds

All chemotherapeutic agents used in this study were obtained from Sigma-Aldrich, dissolved in DMSO at 10mM concentration and stored at -20°C.

# **Cell cultures**

The human chronic myelogenous leukemia cell line, HAP1, was purchased as authenticated at passage 7 from Horizon Discovery and cultured in tissue culture dishes (Genesee Scientific, Cat# 25-202) as a monolayer at 37°C in a humidified atmosphere with 5% CO<sub>2</sub> using Iscove's Modified Dulbecco's Medium (IMDM) (Life Technologies, CA) supplemented with 10% fetal bovine serum (FBS), 0.29mg/mL L-glutamine, 25mM HEPES, 100U/mL Penicillin and 100 $\mu$ g/mL Streptomycin (1% pencillin/streptomycin). Monoclonal and polyclonal stocks were made and stored in IMDM + 10% DMSO in liquid nitrogen.

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#### In vitro evolution of resistant HAP1 clones.

Prior to selection, an aliquot of the parental line was stocked as a reference for subsequent whole genome sequencing analysis. Three independent clones of HAP1 cells were cultured in tissue culture dishes exposed to increasing sublethal concentrations of each chemotherapeutic agent at a starting concentration previously determined by the EC<sub>50</sub> value for around 7-30 weeks depending on the drug, its speed of action and the method used as two methods were considered: high-pressure intermittent selection method and a step-wise selection method. For high pressure selection, cells were treated at a concentration  $3-10 \times EC_{50}$  value until more than 95% of the cells died. Then treatment was removed and cells were allowed to recover. After reaching around 60% semi-confluence, treatment was reinstalled and  $EC_{50}$  value monitored. For step-wise selection method, drug concentration used was at the  $EC_{50}$  which implied reduced growth rate of approximately 50% and drug pressure was increased in intervals of around 5-10% keeping growth inhibition around 50%. Once the  $EC_{50}$  values of the resistant lines were at least 5 times greater than the one used as control, cells were again cloned by limiting dilution and further reconfirmed for drug resistance and subsequent DNA extraction for whole genome sequencing analysis.

# Dose-response assay by EC<sub>50</sub> determination and bioluminescence quantification

Drug sensitivity and cell viability were assessed by a bioluminescence measurement as follows: twenty-four hours prior to addition of the drugs,  $2 \times 10^4$  cells/well for every replicate were seeded in a 96-well plate. Once attached, media was removed and 10 different concentrations of drug were added in serial dilutions 1:3 with a starting concentration of  $10\mu$ M or one of which the EC<sub>50</sub> value of the replicates fell within an intermediate drug concentration. When drug-resistant lines were co-treated in combination with verapamil, a fixed concentration of verapamil ( $10\mu$ M) was added to every concentration of the drug. After a 48-hour incubation period at 37°C and 5% CO<sub>2</sub> with the drug, cells were treated with CellTiterGlo (Promega) reagent (diluted 1:2 with deionized water) for quantification of HAP1 cell viability. Immediately after addition of the luminescence reagent, luminescence was measured using the Synergy HT Microplate Reader Siafrtd (BioTek). The data was normalized to 100% cell survival and 100% cell death and EC<sub>50</sub> values were obtained using the average normalized luminescence intensity of 8 wells per concentration and a non-linear variable slope four-parameter regression curve fitting model in

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Prism 8 (GraphPad Software Inc.). Unless otherwise noted, dose response experiments consisted of 4-8 technical replicates and 3 biological replicates.

#### Isolation of total DNA from drug resistant lines

Genomic DNA (gDNA) was extracted from drug-specific resistant cell lines together with their isogenic parental lines using the DNeasy<sup>®</sup> Blood & Tissue Kit (Qiagen) following the manufacturer's instructions. Samples were assessed for quantity with the Qubit<sup>™</sup> dsDNA BR Assay Kit (Life Technologies, Carlsbad, CA, USA). All samples (>2.0µg, >50ng/µL, >20µL) were prepared for quality control by testing gDNA degradation or potential contamination using agarose gel electrophoresis (1% Agarose, TAE, ~100 Voltage). Then gDNA concentration was again measured using the Qubit<sup>®</sup> DNA Assay Kit with the Qubit<sup>®</sup> 2.0 Fluorometer (Life Technologies). Finally, fragment distribution of the gDNA library was measured using the DNA 1000 Assay Kit with the Agilent Bioanalyzer 2100 system (Agilent Technologies, Santa Clara, CA, USA). DNA libraries were sequenced with 150 base pair (bp) paired single end reads on an Illumina HiSeq 4000 (PE150).

# **Genome Sequencing and Data Analysis**

The quality of the raw FASTQ files checked with FastQC was (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/). Whole genome sequencing (WGS) reads were mapped to GRCh37 (hg19) using BWA (v.0.7.17), specifically with the hs37d5 reference genome from 1000 Genomes project (Phase II). Whole exome sequencing (WES) samples were captured using Agilent SureSelect Human All Exon V6 (58 M), and the reads were also mapped to GRCh37 using BWA (v.0.7.17) with the same reference genome as WGS. Duplicate reads were removed using Picard (v.1.94); paired resistant and parent (WT) BAM files were used as input for The Genome Analysis Toolkit (GATK, v3.8-1). Local realignment and base quality recalibration were performed using default parameters. Somatic single nucleotide variants (SNVs) and small insertion and deletion (indels) were called using GATK MuTect2 following the state-of-the-art GATK Best Practices pipeline (https://ccbr.github.io/Pipeliner/Tools/MuTect2.html). In this project, the input to MuTect2 consisted of alignments for the parent and resistant clone in order. to call mutations with statistically significant differences in read support in the setting of resistance. Only the variants with PASS status, suggesting confident somatic mutations, were considered for further analysis.

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Variant allelic fraction was determined as the fraction of reads supporting the variant allele relative to total read depth at the variant position. Minimum callable depth was set to 10 and base quality score threshold was set to 18, following the default from MuTect2. All sequences have been deposited in SRA BioProject PRJNA603390.

#### **Small-Variant Annotation for SNVs and Indels**

Somatic variants were annotated using snpEff (v 4.3q)[79]. The annotation was performed with parameters including (1) canonical transcripts and (2) protein coding to enable identification of different functional classes of variant and their impact on protein coding genes (Table 1 showing finalized and consolidated annotations; Table S4 shows the raw annotation from snpEff and consolidated classification used in Table 1; Table S7 shows all the SNVs with their raw annotations). The snpEff sequence ontology designation was used in the filtering steps to classify variants generally as noncoding or coding (Table S4).

# **Identification of Drug Specific Genes**

First, we excluded all variants in non-coding regions. Second, we excluded all nonfunctional variants, retaining only variants with a snpEff definition of HIGH or MODERATE impact (missense, stop lost, stop gain, and structural interaction variants). Finally, we selected only the variants with high allele frequency (AF > 0.85) and genes with multiple independent amino acid changes found in the same drug as the final list of candidates. The potential candidate variants were evaluated through Integrative Genomics Viewer (IGV)[80] to ensure coverage and allele fractions of the mutation positions. The top genes for each drug were included in Table 2 and Table S8.

#### Somatic Copy Number Variations (CNVs) Analysis

Copy number regions for WGS and WES were called by ControlFreeC<sup>47</sup> using the default settings for WGS and WES data. Background parental clone samples for each drug served as the control. Recurrent CNV regions were defined as regions observed in more than 1 sample, but not in all of clones from the tested drugs (as these are more likely to indicate potential sequencing artifacts).

# Gene knockdowns

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shRNAs targeting *TOP2A* (Cat# sc-36695-V), *DCK* (Cat# sc-60509-V), *SLCO3A1* (Cat# sc-62713-V), *SLC13A4* (Cat# sc-89760-V), *KLF-1* (Cat# sc-37831-V), *WWOX* (Cat# sc-44193-V), *WDR33* (Cat# sc-94735-V) and the non-coding control (Cat# sc-108080) were obtained in pLKO.1-Puro from Santa Cruz Biotechnology. *RRM1* (clone ID NM\_001033.2-476s1c1) and *CYP1B1* (clone ID NM\_000104.2-1176s1c1) were obtained in pLKO.1-Puro-CMV-tGFP from Sigma Aldrich.

Gene expression was knocked down using either a shRNA pool (Santa Cruz Biotechnology) containing between three and five expression constructs each encoding target-specific 19-25 shRNAs or a single shRNA (Sigma Aldrich). HAP1 cells were plated at 120,000 cells per well (~40% confluency) in a 24-well plate 24 hours prior to viral transduction. On the day of transduction, complete media was replaced with serum-free media and 7µg/mL Polybrene<sup>®</sup> (Cat# sc-134220) and virus was added to the cells at a multiplicity of infection of 0.5 and cells were incubated overnight at 37°C. The following day, media was replaced with complete media without Polybrene and cells were incubated at 37°C overnight. Cells were then split 1:3 and incubated for 24 hours more and finally stable clones expressing the shRNA were selected using complete media with 2µg/mL puromycin. After 7 days of selection with puromycin, knockdown efficiency was confirmed by western blot. Cells transduced with shRNAs containing fluorescent tags, were trypsinized (TrypLE<sup>™</sup> Express; Cat# 12605-010, Gibco) after puromycin selection, washed twice with DPBS (1X) (Gibco) and sorted by flow cytometry.

# **Knockout of USP47**

*USP47* was knocked out (Cat# HSPD0000092816) using a single plasmid CRISPR-*Cas9* system, using as lentivirus backbone the LV01 U6-gRNA:ef1a-puro-2A-Cas9-2A-tGFP targeting *USP47* (Sigma Aldrich). The target sequence (5'-3') was CAATGGGGGCTTCTACTAGG. Transduction was as described above. HAP1 cells were plated at 120,000 cells per well (~40% confluency) in a 24-well plate 24 hours prior to viral transduction. On the day of transduction, complete media was replaced with serum-free media and  $7\mu$ g/mL Polybrene<sup>®</sup> (Cat# sc-134220), virus was added to the cells at a multiplicity of infection of 0.5 and cells were incubated overnight at 37°C. The following day, media was replaced with complete media without Polybrene and cells were incubated at 37°C overnight. Cells were then split 1:3 for 24 hours more and stable clones expressing the CRISPR-*Cas9* sequence were selected using complete media with 2 $\mu$ g/mL

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puromycin. After 14 days of selection with puromycin and propagation as required, cells were trypsinized (TrypLE<sup>TM</sup> Express; Cat# 12605-010, Gibco), washed twice with DPBS (1X) (Gibco) and sorted by flow cytometry using the GFP fluorochrome which is expressed with Cas9. GFP positive cells were plated at an average density of 0.5 cells per well in a 96-well plate (previously treated with poly-L-Lysine (Sigma #P4707-50ml) to improve cell adhesion) in presence of  $2\mu g/mL$  puromycin (limiting dilution cloning). Cell growth was monitored via microscopy during 25 days to select those wells which were observed to contain single colonies and *USP47* knockout was confirmed in those monoclonal HAP1 cell lines first via PCR and then reconfirmed by western blot using the *USP47* rabbit polyclonal antibody (Abcam, Cat# ab97835).

#### Immunoblotting

HAP1 cells (at least  $5 \times 10^6$ ) were trypsinized, washed twice with cold  $1 \times$  DPBS and then lysed in 500µL Pierce<sup>TM</sup> RIPA Buffer (Thermo Scientific) containing 1:100 protease inhibitor (Halt<sup>TM</sup> Protease & Phosphatase Inhibitor Cocktail, Thermo Scientific) and 1:100 0.5M EDTA Solution (Thermo Scientific). Total protein concentration was measured using the DC Protein Assay (Bio-Rad). Equal amounts of proteins were resolved by SDS-PAGE and transferred to nitrocellulose membranes (Bio-Rad #1704271), blocked in PBS with 5% (w/v) Blotting-Grade Blocker (Bio-Rad #170-6404) and 0.1% (v/v) Tween20 for 1h and probed. As secondary antibodies, HRP-linked anti-mouse or anti-rabbit (Cell Signaling Technology) were used and the HRP signal was visualized with SuperSignal<sup>®</sup>West Pico Chemiluminescent Substrate (Thermo Scientific #34080) using Syngene G-Box imager. Protein enrichment was calculated relative to *vinculin*,  $\gamma$ -tubulin or  $\beta$ -actin. Primary antibodies are listed below. Full size western blots are shown in **Fig. S12**.

# Antibodies

TOP2A (Sigma #SAB4502997), USP47 (Abcam #ab97835), WDR33 (Abcam #ab72115), DCK (Abcam #ab151966),  $\beta$ -actin (Cell Signaling #3700S),  $\gamma$ -tubulin (Cell Signaling #4285S), Vinculin (Invitrogen #700062), SLC13A4/SUT-1 (Abcam #ab236619), WWOX (Abcam #ab137726), EKLF/KLF-1 (Abcam #175372), SLCO3A1/OATP-A (Santa Cruz #sc-365007), TOP1 (Proteintech #20705-1-AP), CRISPR-Cas9 (Sigma #SAB4200701), RRM1 (Abcam #ab133690), CYP1B1 (Abcam #ab137562), SPG7 (Sigma #SAB1406470 and Abcam #ab96213), goat anti-mouse (Invitrogen #G21040), goat anti-rabbit (Invitrogen #G21234).

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# RNA isolation, RT-PCR analysis and sequencing of TOP1 (His81)

TPT-resistant cells and TPT-WT ( $1 \times 10^6$  cells) were dissociated from plates by the addition of 2mL of TrypLE (Cat #12605-010, Gibco), washed and total RNA was isolated and purified using a Qiagen RNeasy® Mini Kit (Cat #74104, Qiagen) according to manufacturer's instructions. cDNA was synthesized from 1µg of total RNA using the Superscript<sup>™</sup> First-Strand Synthesis System for RT-PCR Kit (Invitrogen #11904-018) and random hexamers. The primers used to amplify the region containing His81 were FWD: GATCGAGAACACCGGCAC and REV: TCAGCATCATCCTCATCTCGAG. DNA from PCR product was extracted, using the QIAquick® Gel Extraction Kit (Qiagen #28706) following the manufacturer's instruction, measured using the Qubit<sup>®</sup> DNA Assay Kit with the Qubit<sup>®</sup> 2.0 Fluorometer (Life Technologies), and sequenced. The cDNA was sent to Eton Biosciences for Sanger sequencing. Quantification of TOP1 expression was performed using PerfeCTa<sup>®</sup> Sybr Green Fast Mix (Quanta #95072-250) the following primers: FWD: CGAATCATGCCCGAGGATATAA; **REV**: CCAGGAAACCAGCCAAGTAA, following the manufacturer's instruction.

# **GDSC** analysis methods

Mutations and copy number data for cancer cell lines were obtained from the DepMap 2021 quarter 1 release via the DepMap portal (https://depmap.org/portal/download/) on 02/01/2021. Copy number alterations in genes were determined by filtering for a  $log_2(copy number + 1)$  greater in absolute value than 1.5. Cell lines were first grouped according to whether they had a mutation or copy number alteration that matched any of those found in Table 2. EC<sub>50</sub> and dose-response area under the curve (AUC) data were obtained from the GDSC 8.3 Release (June 2020). Dose-response AUC distributions for doxorubicin, etoposide, gemcitabine, paclitaxel, and topotecan were compared between the cell lines with or without a mutation using the Mann-Whitney U test (**Fig. S11A**). P-values were corrected for multiple testing using the Benjamini-Hochberg (BH) method[81]. All cell lines with SNVs in the genes listed in Table 2 were then grouped based on functional predictions by the Variant Effect Scoring Tool (VEST4.0)[82]. Cell lines with a mutation that had a VEST score > 0.8 were labeled as "Likely LOF" cell lines, whereas cell lines with mutations that scored <= 0.8 or that were silent were labeled as "No Likely LOF Mutation" cell lines. As most predicted functional mutations result in loss of function, we assumed this was the likely consequence, though it is possible that some high scoring mutations could in

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fact be gain of function. Dose-response AUC distributions for these groups were then compared for each gene using the Mann-Whitney U test (Fig. S11B) and p-values corrected by the BH procedure.

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Author contributions: JCJ performed selections, validation experiments and wrote the manuscript. MD, AK performed all computational analyses, and assembled figures and tables and wrote the manuscript. GF assisted with CRISPR-*Cas9* experiments. KC performed RNA extraction and RTqPCR experiments. AK performed sequence analysis. HC provided advice and management and wrote the manuscript. EAW performed data analysis, provided advice, obtained funding and wrote the manuscript. TI obtained funding and provided advice.

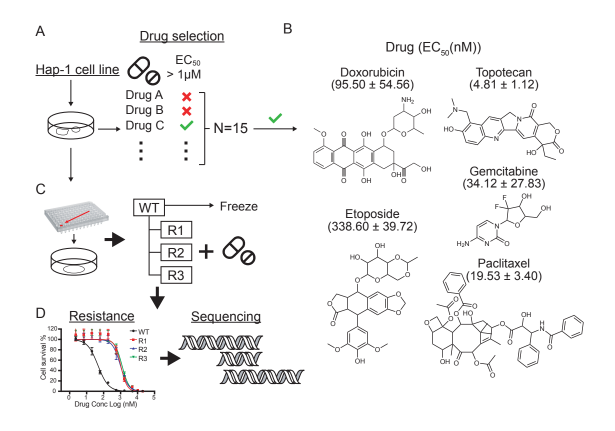
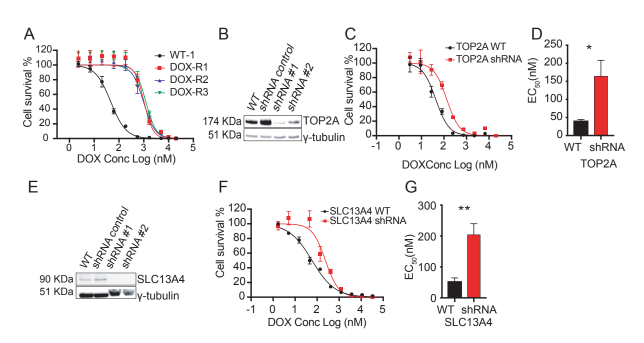
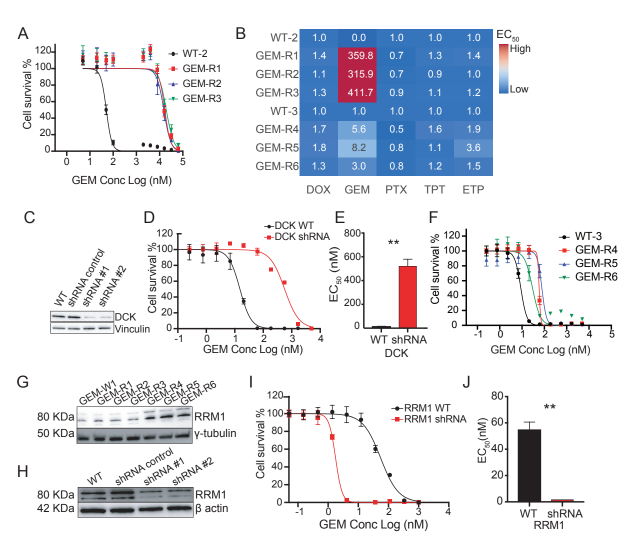


Fig. 1. Experimental workflow. A. Chemotherapy drug evaluation.  $EC_{50}$  dose response assays were performed on 15 different chemotherapeutic agents (Table S1). Only drugs to which HAP1 cells were sensitive ( $EC_{50}$  value below 1µM) were considered for IVIEWGA. B. Chemical structures of the chemotherapy agents ultimately used for IVIEWGA.  $EC_{50}$  values are presented as the mean ± s.e.m., for n=3 biological replicates and n=8 with technical replicates per concentration point. C. Clone selection. To ensure a homogenous genetic background limiting dilution cloning was used to isolate individual cells prior to drug selection. For each drug three independent selections were performed. Resistance was confirmed using dose-response assays D. Drug resistance was achieved in 7 to 30 weeks approximately (49 and 210 generations). The parental cell line and the drug resistant lines were then sequenced.

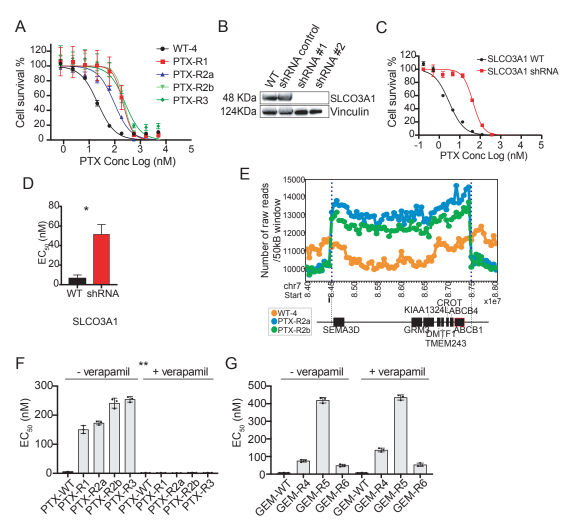


**Fig. 2. A.** DOX EC<sub>50</sub> curves for DOX evolved clones using 8 technical replicates for each concentration **B.** Western blot confirming that shRNA gene depletion downregulates TOP2A protein level. **C.** EC<sub>50</sub> curves of the WT and shRNA (shRNA #1) knockdown cell lines for *TOP2A*. **D.** Barplot of the WT and shRNA knockdown cell line (shRNA#1) for *TOP2A*. **E.** Western blot confirming protein downregulation of SLC13A4 by shRNA. **F.** EC<sub>50</sub> curves of the WT and shRNA knockdown cell lines for *SLC13A4*. **G.** Barplot of the WT and shRNA knockdown cell lines for *SLC13A4*. Data is represented by mean  $\pm$  s.e.m. with n=2 biological replicates and n=8 technical replicates (*TOP2A*) or n=3 biological replicates and n=4 technical replicates (*SLC13A4*) for every concentration point. \* = p value < 0.05, \*\* = p < 0.01. p values determined by two-tailed *t* test.

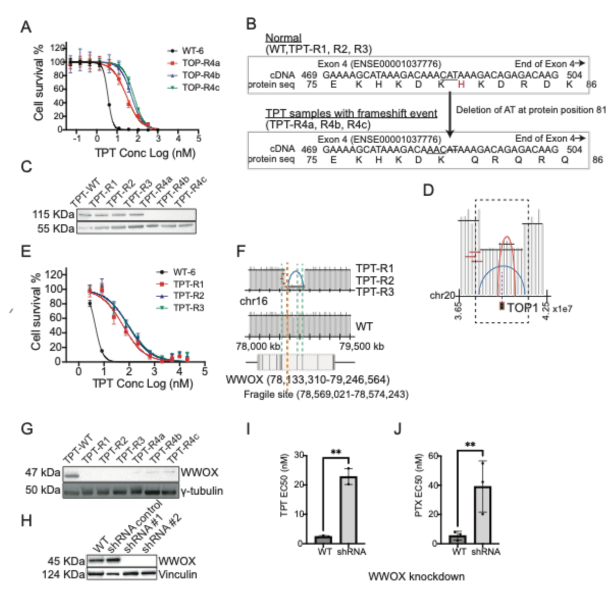




**Fig. 3. A.** GEM EC<sub>50</sub> for curves first set of GEM evolved lines using n=8 technical replicates per concentration point **B.** EC<sub>50</sub> ratio matrix showing absence of multidrug resistance pathways of GEM resistant lines. **C.** Western blot confirming that shRNA gene depletion downregulates protein levels for *DCK*. **D.** EC<sub>50</sub> curves of the WT and shRNA knockdown cell lines for *DCK*. n=8 with individual technical replicates overlaid for every concentration point. **E.** Barplot of the WT and shRNA knockdown cell lines for *DCK*. **F.** GEM EC<sub>50</sub> curves for second set of GEM evolved lines. **G.** Western blot for *RRM1* across all GEM samples showing overexpression pattern of *RRM1* in GEM-R4-6 resistant clones. *γ*-tubulin is used as a loading control. **H.** Western blot confirming that shRNA gene depletion downregulates protein levels for *RRM1*.  $\beta$ -actin is used as a loading control. **I.** EC<sub>50</sub> curves of the WT and shRNA knockdown cell lines for *RRM1*. All data is represented by mean ± s.e.m. with n=3 with individual biological replicates overlaid. \*\* = p value < 0.01. p values determined by two-tailed *t* test.



**Fig. 4. A.** PTX EC<sub>50</sub> curves for evolved lines with n=8 technical replicates (values are in Table S2) **B**. Western blot confirming shRNAs downregulate protein levels for *SLCO3A1*. **C.** EC<sub>50</sub> curves of the WT and shRNA knock-down cell lines. **D**. Barplot of the WT and shRNA knockdown cell lines for *SLCO3A1*. **E**. Raw copy number profile for the amplification event containing protein coding genes including ABC transporters (*ABCB1/ABCB4*) PTX cell lines. The amplification region (chr7:84,500,000-87,300,000) had a higher number of raw reads (labeled with blue dash lines) with default window size of 50K bp. Genes associated with the CNV event are depicted by black boxes underneath according to their position and sizes. *ABCB1* is highlighted with red outline. **F**. Barplot of EC<sub>50</sub> of the PTX treated cell lines with and without verapamil and verapamil alone showing sensitization in presence of verapamil as ABC inhibitor (n=4 technical replicates). **G**. Barplot of EC<sub>50</sub> of the GEM cell lines  $\pm$  verapamil showing no EC<sub>50</sub> shift for GEM cell lines when co-treated with verapamil. Unless otherwise noted, all data is represented by mean  $\pm$  s.e.m. with n=3 with individual biological replicates overlaid. \*\* = p value < 0.01. p values determined by two-tailed *t* test.



**Fig. 5. A.** TPT EC<sub>50</sub> curves for TOP-R4a,b and c. **B.** Transcript sequence (cDNA) and protein sequence for TOP1 transcript for normal (top) and TOP 4a, b and c. The figure shows only part of the cDNA (position 469-504) and protein sequence (position 75-86) for TOP1 at the affected exon (Exon 4, ENSE00001037776). The frameshift deletion of nucleotides 'CAT' to 'C' observed in TPT samples (TPT-R4a, R4b, and R4c) is predicted to give a frameshift at amino acid 81 (His, red highlight in normal). Amino acids affected by the frameshift deletion are highlighted in red. **C.** Western blot TOP1 protein depletion in evolved lines. **D.** Schematic showing complex read depth patterns around *TOP1*. *E.* TPT EC<sub>50</sub> curves for evolved TOP-R1, R2 and R3. **I.** Schematic of chr16 reads around *WWOX* for TPT-R1, R2, and R3 compared to the WT chromosome 16 parental cell line. Blue arch represents a deleted region. *WWOX* below shows the exonic (black lines) and intronic (white box) regions of the gene. The start of the deletion event is also close to a known fragile site (orange dashed line). **G.** Western blot showing WWOX protein levels in TPT resistant

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clones. **H.** Western blot shows downregulation of protein levels for *WWOX* in shRNA samples compared to WT and scrambled control. Barplot of the WT and shRNA knockdown cell lines for *WWOX* showing EC<sub>50</sub> values for TPT (**I**) and PTX (**J**). Data is represented by mean  $\pm$  s.e.m. with n=3 biological replicates and 4-8 technical replicates. \*\* = p < 0.05. p values determined by ratio, paired two-tailed *t* test.

# TABLES

			WGS				WES			
	DOX	GEM	PTX	TPT	TPT	DOX	ETP	GEM	PTX	
	(n=3)	(n=3)	(n=4)	(n=3)	(n=3)	(n=3)	(n=3)	(n=3)	(n=3)	
Indels										
Disruptive inframe ins.	0.00	0.00	0.00	0.00	1.00	1.00	0.00	0.00	0.00	
Frameshift	0.00	1.00	1.33	2.33	0.00	1.00	1.00	1.00	1.00	
Frameshift plus stop-gained	0.00	0.00	0.00	0.33	0.00	0.00	1.00	0.00	0.00	
Inframe insertion	0.00	0.00	1.00	0.33	0.00	0.00	1.00	0.00	0.00	
Intergenic	27.67	43.00	26.75	24.67	47.67	2.00	3.50	1.00	1.00	
Intragenic	10.00	5.67	9.00	9.00	14.33	1.00	2.00	1.00	1.00	
Intron	12.00	20.33	16.25	15.67	32.33	1.00	1.00	4.00	1.50	
Splice region plus intron	0.00	0.00	0.00	103.3	0.00	0.00	0.00	1.00	0.00	
			S	NVs						
Disruptive inframe del.	0.00	1.00	1.00	0.67	0.00	1.00	0.00	1.00	1.33	
Frameshift	1.00	2.00	1.00	22.33	2.00	3.00	1.67	1.00	3.00	
Inframe deletion	0.00	0.00	1.00	0.00	0.00	0.00	2.00	0.00	0.00	
Intergenic	898.33	1303.3	1416.6	258.6 7	2635.7	25.00	19.00	17.00	21.33	
Intragenic	272.00	403.33	389.25	77.67	834.67	15.33	12.33	10.33	11.33	
Intron	448.67	701.33	764.00	128.3	1358.33	28.00	27.00	33.00	22.67	
Missense	16.00	14.00	12.75	7.00	34.33	15.00	19.67	21.67	15.67	
Others	1.00	1.00	0.00	0.00	1.33	1.00	1.00	1.50	1.00	
Splice region plus intron	1.00	1.67	2.00	30.00	3.33	1.67	1.50	1.00	1.33	
Start lost	0.00	0.00	0.00	0.00	0.00	0.00	0.00	1.00	1.00	
Stop gained	0.00	2.50	1.00	0.00	1.67	1.50	2.00	2.67	0.00	
Stop lost	0.00	1.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	
Synonymous	3.00	7.33	4.00	6.33	7.67	6.67	5.33	7.67	5.33	

**Table 1. Summary of average number of mutations.** Number of selections performed with the drug is given by n. SNVs and Indels were grouped according to snpEff sequence ontology annotations (Methods, Table S4), and detailed counts per clone can be found in Table S5.

2	2
3	5

Drug	Sample	Gene	Amino Acid Change	Туре	AF	EC <sub>50</sub>	EC <sub>50</sub> (nM)
3	1		0	<i></i>		(nM)	KD/KO/ĆI
						ŴŤ	
DOX	R1, R2, R3	TOP2A	Pro803Thr	MS	0.89, 0.87	$38.6\pm4.3$	$164.3\pm43.9$
	R4b	SLC13A4	Gly165His	MS	1	$52.9 \pm$	$204.3\pm35.7$
						11.6	
	R5	SPG7	Lys593Asn	MS	1	NE*	
PTX	R1, R2a, R3	WWOX	16q23.1	CNV	-	$5.8\pm2.5$	$42.9\pm11.7$
	R2a, R2b	ABCB1	7q21.12	CNV	-	$252\pm38$	$1.3\pm0.1$
						$218 \pm 14$	$1.3 \pm 0.1$
	R2b, R6	SLCO3A1	Ile587Asn (R2b),	MS		$6.6\pm3.3$	$51.5\pm9.9$
			Ala263Thr (R6)				
GEM	R1, R2, R3	DCK	Ser129Tyr (R1, R2),	MS,		$14.3 \pm 1.7$	$521.7 \pm 58.3$
			Asn80fs (R1. R3);	FS			
			Asn113fs (R2);				
			Thr184fs (R3)				
	R4, R5, R6	RRM1	11p15.4	CNV	-	$54.9\pm5.8$	$1.8 \pm 0.1$
ETP	R2, R3	WNT3A	1q42.13 (R2)	CNV	-	ND	-
	R3	WDR33	P622T	MS	1	$241.5 \pm$	$821.6 \pm$
						31.0	226.9
ТРТ	R1, R4a,	CYP1B1	Val432Leu;	MS	0.13, 0.40,	$6.3\pm0.2$	$13.3\pm0.3$
	R4b, R4c		Asp217Glu (R4a,b,c)		0.43, 0.42		
	R1, R2, R3	WWOX	16q23.1	CNV	-	$2.4\pm0.3$	$22.8\pm2.7$
	R4a, R4b,	TOP1	His81fs; 20q12	FS;	1	ND	-
	R4c			CNV			
	R4a, R4b,	USP47	Arg408*	Stop	0.38, 0.57,	$3.05 \pm 0.2$	$1.07 \pm 0.07$
	R4c				0.58		

**Table 2.** Validation (knockdown) results for selected genes. CNV, copy-number variant. MS, missense. FS, frameshift variant, KO/KD/CI, knockout, knockdown, chemical inhibition (verapamil, ABCB1). ND: No data: gene knockdowns were attempted but could not be achieved. NE: Not expressed (protein not detected by Western blot, preventing validation).  $EC_{50}$  WT and  $EC_{50}$  KO/KD/CI are from matched pairs for the given drug and represent the mean  $\pm$  s.e.m. (n=3 biological replicates).

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