

GANDALF: GENERATIVE ATTENTION BASED DATA AUGMENTATION AND PREDICTIVE MODELING FRAMEWORK FOR PERSONALIZED CANCER TREATMENT

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ABSTRACT

Effective treatment of cancer is a major challenge faced by healthcare providers, due to the highly individualized nature of patient responses to treatment. This is caused by the heterogeneity seen in cancer-causing alterations (*mutations*) across patient genomes. Limited availability of response data in patients makes it difficult to train personalized treatment recommendation models on mutations from clinical genomic sequencing reports. Prior methods tackle this by utilising larger, labelled pre-clinical laboratory datasets (‘cell lines’), via transfer learning. These methods augment patient data by learning a shared, domain-invariant representation, between the cell line and patient domains, which is then used to train a downstream drug response prediction (DRP) model. This approach augments data in the shared space but fails to model patient-specific characteristics, which have a strong influence on their drug response. We propose a novel generative attention-based data augmentation and predictive modeling framework, GANDALF, to tackle this crucial shortcoming of prior methods. GANDALF not only augments patient genomic data directly, but also accounts for its domain-specific characteristics. GANDALF outperforms state-of-the-art DRP models on publicly available patient datasets and emerges as the front-runner amongst SOTA cancer DRP models.

1 INTRODUCTION

Cancer, a leading cause of deaths worldwide (Dattani et al., 2023), imposes a significant burden on global healthcare systems (Lopes, 2023). It is caused due to the presence of alterations (*mutations*) in the human genome, resulting in uncontrolled replication of cancer cells. Cancer patients exhibit a great deal of heterogeneity in their genomic mutation profiles, even when they have the same cancer type. This heterogeneity causes patients, of the same cancer type, to respond differently to the same treatment (Liao et al., 2023), making cancer treatment challenging (Wahida et al., 2023). Treatment, today, is largely guideline-based and prescribes drugs based on the cancer type (Planchard et al., 2018; Conroy et al., 2023; Morris et al., 2023). This approach fails to account for heterogeneity in patient mutations, and its impact on treatment outcomes. Precision oncology Sosinsky et al. (2024); Collins & Varmus (2015) is gradually shifting focus from a “one-size-fits-all” approach to more personalized treatment strategies.

To aid precision oncology, cancer patients undergo genomic sequencing as part of clinical diagnostics (Colomer et al., 2023). Clinical sequencing panels (Milbury et al., 2022; Wei et al., 2022) identify the set of mutations present in specific sections of the human genome (called *genes*), which have a known association with cancer. Cancer patients can exhibit a varying number of mutations in each of these genes (Saito et al., 2021). These mutations interact with each other and the drug in complex ways to determine patient response to treatment (Liu et al., 2022). While clinical trials have identified drugs that target specific mutations, these studies have largely been restricted to single mutations (Brachova et al., 2013; Randic et al., 2023). Conducting large scale clinical trials for all possible combinations of mutations in ~ 20000 genes of the human genome is practically intractable, thereby limiting their ability to identify the right treatment when a patient exhibits multiple mutations.

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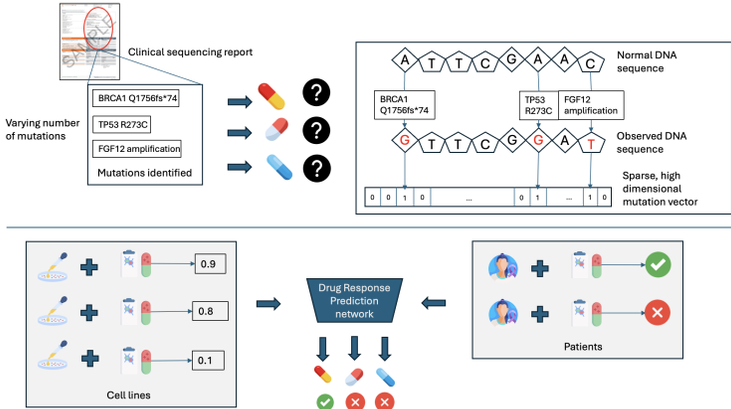


Figure 1: Overview of clinical challenge in cancer drug response prediction.

Machine learning (ML) approaches provide a promising avenue to predict patient response y_p to drugs d_p , based on the set of mutations X_p in their genomic profiles. However, guideline-based treatment in clinics prescribe only a small subset of drugs from all drugs approved for clinical use, thereby limiting the availability of labelled patient data (X_p, d_p, y_p) . The resulting scarcity poses a significant challenge in training supervised ML models to predict drug response in patients. Prior methods in Drug Response Prediction (DRP) literature have tackled this using data from a related domain called “cell lines”. Cell lines (Ghandi et al., 2019) are cancer cells extracted from patients, which are then cloned under controlled laboratory settings. Each clone X_c is administered a different drug d_c , and the corresponding response $y_c(X_c, d_c)$ is measured for various drug concentrations. Since these cells are studied outside the human body, it is possible to obtain y_c for a large set of drugs \mathcal{D} , resulting in abundant labelled data.

However, models trained only on (X_c, d_c, y_c) do not work well on patients (Mourragui et al., 2019; 2021; Sharifi-Noghabi et al., 2020). This is attributed to the inherent differences between patients and cell lines. As cell lines are studied outside the human body in the absence of blood vessels and the immune system (called *tumor microenvironment*), these cells can acquire mutations differently compared to patients, i.e. $P(X_c) \neq P(X_p)$. In addition, $y_c \in [0, 1]$ depends on drug concentration and number of surviving cells (called Area Under the Dose Response Curve, AUDRC), while $y_p \in \{0, 1\}$ indicates good or bad response (called Response Evaluation Criteria in Solid Tumors, RECIST, based on change in tumor volume), i.e. $domain(y_c) \neq domain(y_p)$, as shown in Figure 1.

Prior DRP methods (Jayagopal et al., 2024; 2023; Kim et al., 2024; He et al., 2022) have addressed these differences by learning shared domain-invariant representations Z_s between X_c and X_p , which are then used to train a downstream drug response prediction network f . Transforming X_c to Z_s increases samples in the shared space and allows f to use the larger (Z_s, d_c, y_c) in training, thereby tackling the data scarcity issue. However, Z_s does not capture patient-specific characteristics in X_p , which can influence y_p (Liao et al., 2023; Zhai & Liu, 2024). To capture this, we need to augment X_p directly. Prior DRP methods, except WISER, neglect this. WISER (Shubham et al., 2024) performs data augmentation by pseudolabelling unlabelled patient profiles $X_{p(u)}$ using (X_c, d_c, y_c) and trains f by combining (X_c, d_c, y_c) and pseudolabelled $X_{p(u)}$. However, while combining the two datasets, WISER assumes $domain(y_c) = domain(y_p)$, and does not account for $P(X_{p(u)}) \neq P(X_c)$. We tackle these issues using **GANDALF**, a *Generative Attention based Data Augmentation and predictive modeLing Framework*. GANDALF augments X_p directly, by generating more “patient-like” samples X_{aug} leveraging available X_c . It also generates their response labels y_{aug} to drugs $d_{aug} \in \mathcal{D}$. Unlike WISER, it explicitly models $domain(y_c) \neq domain(y_p)$ and $P(X_p) \neq P(X_c)$.

Data augmentation strategies are known to improve prediction performance in various fields of ML, like computer vision (Khosla & Saini, 2020) and natural language processing (Shorten & Khoshgoftaar, 2019). This is usually achieved through data transformations where identifying the label of the transformed data is relatively easy, e.g., a rotated image of a dog retains the label ‘dog’ after transformation. However, it is difficult to find such ‘label-invariant’ transformations for genomic data (Lacan et al., 2023). Although genomic data can be augmented by interpolation of available samples or sampling new data points from a known distribution, assigning labels to these samples

is difficult. Data points, which may be “close” together in the representation space, can still exhibit different responses to drugs. If patients are represented by binary vectors (each element corresponding to a gene, 1 indicating presence of mutations in a gene and 0 the absence), a perturbation is equivalent to addition or removal of a mutation. This perturbation can impact the functioning of the cells and the response to treatment (Hale et al., 2024). Identifying the response associated with each perturbation is difficult due to scarcity of labelled data, making data augmentation strategies challenging in DRP.

Though conclusively identifying labels for all possible perturbations is still an open problem, GANDALF takes a step towards leveraging data augmentation in DRP, by utilising available labelled data from cell lines. It generates X_{aug} by transforming X_c and assigns y_{aug} for generated $(X_{aug}, d_{aug}), d_{aug} \in \mathcal{D}$ by leveraging labelled information from (X_c, d_c, y_c) . We use attention mechanisms to ensure that X_{aug} retains information from X_c . $(X_{aug}, d_{aug}, y_{aug})$ is then used with (X_p, d_p, y_p) to train a downstream DRP classifier. Our paper makes the following contributions:

- We are the first to tackle, through a novel data augmentation approach, the challenging problem of limited labels for sparse patient genomic data, in cancer drug response prediction.
- We propose GANDALF, a generative, semi-supervised, attention-based data augmentation framework which uses labelled samples from the related cell line domain to generate labelled patient data.
- GANDALF performs data augmentation through a novel synthesis of denoising diffusion probabilistic models, transformers and multi-task learning.
- GANDALF demonstrates an improvement of upto 10.96% over SOTA DRP methods, in predicting patient response to drugs, on key benchmark datasets comprising real patient samples with responses to clinically approved anti-cancer drugs. GANDALF also outperforms baseline genomic data augmentation and pseudo-labeling strategies by 21% and 2.5% respectively.

2 RELATED WORK

2.1 DRUG RESPONSE PREDICTION MODELS

Prior DRP models perform transfer learning between the source domain (cell lines) and target domain (patients). These methods can be inductive, transductive or unsupervised (Pan & Yang, 2009), based on their use of labelled patient data. Inductive methods, like AITL (Sharifi-Noghabi et al., 2020), drug2tme (Zhai & Liu, 2024) and TCRP (Ma et al., 2021) use both labeled cell line and patient samples. They may either use multi-task learning approaches or few shot learning to capture the differences in label distribution across the two domains. Transductive methods like TUGDA (Peres da Silva et al., 2021), WISER (Shubham et al., 2024), PANCDR (Kim et al., 2024) use labeled cell line and unlabeled patient samples. The unjustified assumption is that the response label does not change across the domains. To this end, most papers convert the continuous valued cell line response to discrete categories as seen in patients, using arbitrary thresholds. Few methods, like CODE-AE (He et al., 2022), rely on unsupervised transfer learning using unlabeled cell line and patient datasets in pre-training. However, in most cases, the goal was to learn a shared representation space between the domains. The shared representation was then used to train a downstream DRP model. While the shared representation captures the similarities across the domains, this approach largely neglects the patient-specific characteristics, which is relevant for drug response prediction.

2.2 GENOMIC DATA AUGMENTATION

Genomic data augmentation is difficult due to lack of known label-invariant transforms (Lacan et al., 2023). Most existing methods augment transcriptomic data (Das & Shi, 2022; Chen et al., 2020), which is unavailable in a clinical setting. A few recent methods (Yu et al., 2024; Lee et al., 2023; Duncan et al., 2024; Lee et al., 2024) have augmented mutations, but they assume that the biological function and associated labels do not undergo changes during data transformation. Moreover, none of these methods focus on cancer drug response prediction as the downstream task, where it is known that even the addition or removal of a mutation can cause a change in drug response (Liao et al., 2023). Thus, patient mutation data augmentation for cancer drug response prediction is an open problem. GANDALF proposes a way forward, by using prior information available in labelled

cell lines to augment patient mutation data and to generate associated labels for DRP, rather than assuming label invariance.

3 METHOD

3.1 PROBLEM FORMULATION

Given a patient genomic mutation profile X_p and drug d_k , the goal in drug response prediction (DRP) is to classify whether the patient would respond well (label $y_p = 1$) or not (label $y_p = 0$), i.e. to learn a classifier $f_{d_k}(X_p) : \mathcal{R} \rightarrow \{0, 1\}$. Let \mathcal{M} denote the set of all possible mutations found in set of sequenced genes \mathcal{G} and \mathcal{A} denote the set of possible alterations in \mathcal{G} . Each mutation $m_l \in \mathcal{M}$ can be separated out into a gene component $g_l \in \mathcal{G}$ and alteration $a_l \in \mathcal{A}$. Let \mathcal{D} denote the set of chemotherapy drugs. Two related, albeit different datasets are available to perform the DRP task - labelled pre-clinical cell line data and clinical patient data. Cell line genomic data $X_c \subset \mathcal{P}(\mathcal{M})$ and labelled patient genomic data $X_p \subset \mathcal{P}(\mathcal{M})$, where $\mathcal{P}(\cdot)$ denotes the power set of \mathcal{M} . Let $\mathcal{N}_c = |X_c|$ and $\mathcal{N}_p = |X_p|$ denote the number of unique mutation profiles in each dataset. $y_{p(jk)} \in \{0, 1\}$ is a binary RECIST response associated with patient-drug pair (x_{pj}, d_k) , while $y_{c(jk)} \in [0, 1]$ is the real-valued AUDRC response for cell line-drug pair (x_{cj}, d_k) . To illustrate, a patient mutation profile $x_{p(1)} = \{m_5 = (g_2, a_{10}), m_7 = (g_{100}, a_8)\}$ has a response $y_{p(13)} = 1$ for drug d_3 . The goal is to predict the response $y_{p(jk)}$ for a new patient-drug pair (x_{pj}, d_k) . To achieve this, we perform patient data augmentation, i.e. generate $(X_{aug}, d_{aug}, y_{aug})$ using (X_c, d_c, y_c) and (X_p, d_p, y_p) . d_c and d_p denote the set of drugs available in labelled cell line and patient datasets, and $d_{aug} \subseteq \mathcal{D}$. In general, $|d_c| > |d_p|$, $d_c \subseteq \mathcal{D}$ and $d_p \subset \mathcal{D}$, as obtaining drug responses in cell lines for a wide range of drugs is easier than in patients. The real and generated labelled patient data $(X_{aug}, d_{aug}, y_{aug}) \cup (X_p, d_p, y_p)$ can then be used to train a downstream DRP classifier f . Please note that $*$ can denote c or p in subsequent sections, to denote cell lines and patients respectively.

3.2 METHOD OVERVIEW

We propose a *Generative Attention based Data Augmentation and predictive modelLing Framework* - GANDALF, to tackle the labelled patient data scarcity issue via data augmentation. The complete algorithm is available in Algorithm 1. GANDALF generates new patient-like samples from cell lines and assigns them labels in 5 steps - (1) pretraining diffusion models to learn representations of X_c and X_p , (2) generating new patient-like samples X_{aug} from X_c , (3) training a multi-task learning network using (X_c, d_c, y_c) and (X_p, d_p, y_p) , (4) assigning pseudolabels y_{aug} for $(X_{aug}, d_{aug}) \forall d_{aug} \in \mathcal{D}$ and selection of confident samples $(X_s, d_s, y_s) \subseteq (X_{aug}, d_{aug}, y_{aug})$ and (5) training DRP classifier f on $(X_s \cup X'_p, d_s \cup d_p, y_s \cup y_p)$.

The goal is to learn $g(\cdot) : X_{aug} = g(X_c) \sim P(X_p)$, which accounts for patient-specific characteristics. The intuition behind the transformation process is: if we decompose each domain into domain-invariant Z_s and domain-specific Z_p (for patients) and Z_c (for cell lines) representations (Lee & Pavlovic, 2021), to transform $X_c \rightarrow X_p$, we introduce Z_p over Z_s obtained from X_c . We can then augment (X_p, d_p, y_p) using $(X_{aug}, d_{aug}, y_{aug})$, $d_{aug} \in \mathcal{D}$, where y_{aug} can be generated by pseudolabelling (Lee et al., 2013; Kage et al., 2024). Our pseudolabelling approach assumes that y_c and y_p share certain characteristics, while differing in others.

3.2.1 STEP 1: PRETRAINING DIFFUSION MODELS

In this step, we learn Z_s, Z_p and Z_c representations. We assume $Z_s \sim \mathcal{N}(0, I)$, which can be modelled using denoising diffusion probabilistic model (DDPM) encoders (Ho et al., 2020). The DDPM decoders learn to remove the domain-specific noise, to reconstruct X . Transforming $X_c \rightarrow X_p$ would then involve the use of the patient DDPM decoder on Z_s . We train two DDPM models (TD_p and TD_c), one per domain, such that they share a common Z_s . In addition, we use the pretrained transformer encoder (T_e) from (Jayagopal et al., 2024), with padding, to model varying number of mutations. We use domain alignment losses (Sun et al., 2016) to align Z_s and KL-divergence loss to ensure $X_{aug} \sim P(X_p)$. We use cross-attention to ensure X_{aug} retains information from X_c .

T_e takes as input $\{m_l; m_l \in \mathcal{M}\}$. Each m_l has two parts - the gene part $g_l \in \mathcal{G}$ and the alteration part $a_l \in \mathcal{A}$. g_l and a_l are tokenized separately, padded and concatenated to generate a per-sample vector. In the embedding step, each a_l is embedded following the variant annotation procedure in (Jayagopal et al., 2024), to obtain a 23-dimensional embedding. This consists of a 17 dimensional binary vector from Annovar (Wang et al., 2010), a 3-dimensional binary vector each from GPD (Li et al., 2020) and ClinVar (Landrum et al., 2018). The embedding for each a_l is passed through a

Algorithm 1 GANDALF training

Require: Mutation profiles X_c, X_p , drugs \mathcal{D} , cell line-drug labels y_c , patient-drug labels y_p , time steps t , pre-trained transformer encoder T_e , DDPM networks TD_* , VAEs V_* , pre-train epochs e_p , pseudolabel generation epochs e_s , upper and lower thresholds t_u and t_l and DRP training epochs e_d .

- 1: **Step 1: Pretraining diffusion models**
- 2: Obtain transformer embedded samples $Z_{t*} = T_e(X_*) \in \mathcal{R}^{N_* \times k}$
- 3: Pre-train domain specific VAEs using Eq. 1 and 2
- 4: **for** e in range(e_p) **do**
- 5: Extract output from the tranformer-VAE encoder network $E = V_{*(e)}(T_e(\cdot))$
- 6: $Z_{v*} = S(\mu_*, \sigma_*) (S(\cdot) = \mu_* + \sigma_* \epsilon$, where $\epsilon \sim \mathcal{N}(0, 1)$, $\mu_*, \sigma_* = E_*(X_*)$)
- 7: $Z_{v*} = TD_{*(e)}(Z_{v*})$
- 8: $\mathcal{X}'_* = \text{denoise}(Z_{v*}, t, TD_{*(d)}(Z_{v*}))$
- 9: $\bar{Z}_{t*} = V_{*(d)}(\mathcal{X}'_*)$
- 10: $Z_{Att} = \text{softmax}(\frac{Z_{vpt} Z_{vct}^T}{\sqrt{t}}) Z_{vct}$
- 11: $\hat{Z}_{vpa} = \text{denoise}(Z_{Att}, t, TD_{p(d)}(Z_{Att}))$ using Eq. 5
- 12: Minimise loss L_{PRE} until convergence.
- 13: **end for**
- 14: **Step 2: Generating new patient-like samples**
- 15: $Z_{vct} = TD_{c(e)}(V_{c(e)}(Z_{tc}))$
- 16: $X_{aug} = \text{denoise}(Z_{vct}, t, \epsilon_{p\theta}); \epsilon_{p\theta} = TD_{p(d)}(Z_{vct})$
- 17: **Step 3: Training multi-task learning network**
- 18: **for** e in range(e_s) **do**
- 19: Obtain cell line and patient embeddings $Z_{v*} = S(E_*(X_*))$
- 20: Obtain drug embeddings $Z_{d*} = g_d(d_*)$
- 21: For each sample, drug pair concatenate the embeddings to get $O_{*d} = Z_{v*} || Z_{d*}$
- 22: Obtain AUDRC and RECIST predictions: $\hat{y}_c = g_a(O_{cd}); \hat{y}_p = g_r(O_{pd})$
- 23: Minimise L_{MTL} till convergence.
- 24: **end for**
- 25: **Step 4: Assigning pseudolabels and selection of confident samples**
- 26: $y_{aug} = g_r(X_{aug} || g_d(d_{aug}))$ for $d_{aug} \in \mathcal{D}$.
- 27: Set y_{bin} as 1 if $y_{aug} \geq t_u$, 0 if $y_{aug} < t_l$ and -1 otherwise.
- 28: Select confident tuples (non-abstained tuples) (X_s, d_s, y_s) , i.e. where $y_{bin} \neq -1$.
- 29: Combine (X_s, d_s, y_s) with $(\mathcal{X}'_p, d_p, y_p)$ to form $(X_{comb}, d_{comb}, y_{comb})$
- 30: **Step 5: Training drug response prediction classifier**
- 31: **for** e in range(e_d) **do**
- 32: $y_{comb} = f(X_{comb} || d_{comb})$
- 33: Minimise loss L_{BCE} in Eq. 10 till convergence.
- 34: **end for**

linear layer and concatenated with the corresponding g_l embedding (obtained by one hot encoding), before being fed into T_e . The resulting output is mean-aggregated to obtain sample embedding $Z_{t*} = T_e(X_*) \in \mathcal{R}^{N_* \times k}$, where k denotes the maximum sequence length. k is set based on maximum number of alterations in the training data, and all sequences are padded to match k . T_e was trained to predict the progression-free survival (PFS) for (X_p, d_p) . PFS is indicative of the time after treatment that a cancer patient survives without the cancer progressing. For further details, please refer to (Jayagopal et al., 2024).

To ease training (Rombach et al., 2022), we reduce the dimensionality of Z_{t*} from $k \rightarrow l, l < k$ using variational autoencoders (VAEs) (Kingma & Welling, 2013). We use 2 VAEs - V_c and V_p for cell line and patient domains respectively. These VAEs take as input $Z_{t*} \in \mathcal{R}^{N_* \times k}$ and estimate the mean $\mu_c, \mu_p \in \mathcal{R}^{N_* \times l}$ and standard deviation $\sigma_c, \sigma_p \in \mathcal{R}^{N_* \times l}$ of each domain. Samples generated using the estimated μ and σ are used to train TD_* . The VAEs are pretrained on each domain, to minimise reconstruction mean square error and KL divergence loss as in Eq. 1 and 2. The VAE pretraining loss is $L_{VAE} = L_R + L_{KLD}$.

$$L_R = \frac{1}{N_*} \sum_{N_*} (\hat{Z}_{t*} - Z_{t*})^2 \quad (1)$$

$$L_{KLD} = -(0.5/N_*) \sum_{N_*} (1 + \log(\sigma_*(Z_{t*})^2) - \mu_*(Z_{t*})^2 - \sigma_*(Z_{t*})^2) \quad (2)$$

where N_* denotes number of mutation profiles (N_c or N_p), \hat{Z}_{t*} is the reconstructed VAE output. Pretrained T_e attached to the encoder layers of the pretrained V_c and V_p , are henceforth referred to as encoder networks E_c and E_p ; $\mu_*, \sigma_* = E_*(X_*)$. Parameters of T_e are frozen for training.

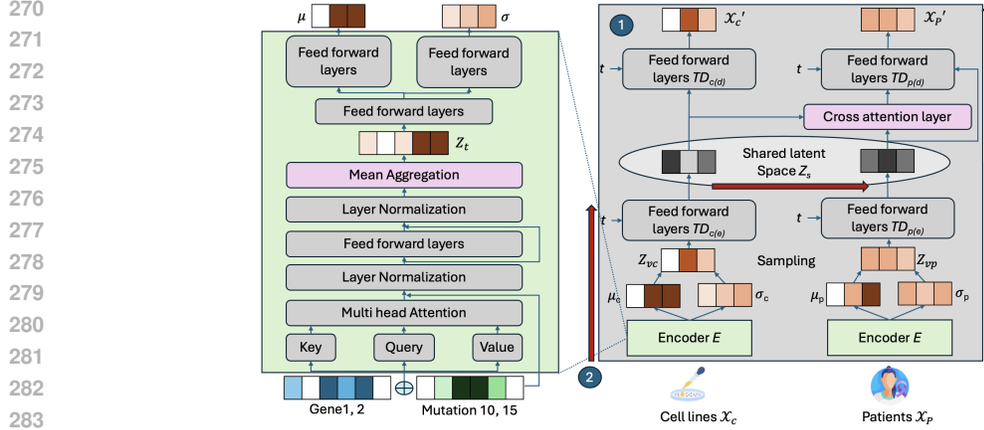


Figure 2: GANDALF architecture used for pretraining domain-specific diffusion models and to generate new patient-like samples using available cell line data. Circled numbers in blue indicate steps from Algorithm 1.

The sampled output from E_* , $Z_{v*} = S(\mu_*, \sigma_*)$ ($S(\cdot) = \mu_* + \sigma_* \epsilon$ denotes VAE sampling, where $\epsilon \sim \mathcal{N}(0, I)$) is fed into TD_c and TD_p , with encoder $TD_{*(e)}$ and decoder $TD_{*(d)}$. Since Z_{v*} is a vector, we used feed forward linear layers in TD_* (Kotelnikov et al., 2023). To learn Z_s , we perform domain alignment, using CORAL loss (Sun et al., 2016). CORAL loss minimises the co-variance between the latent spaces, as in Eq. 4. Although in theory, DDPM encoders should yield isotropic Gaussians as $T \rightarrow \infty$, the use of CORAL loss enforces that the two domains share Z_s , when \hat{T} is finite. TD_c and TD_p are trained jointly with the CORAL loss using $L_{ALIGN} = L_{DDPM} + L_{CORAL}$, as in Eq. 3 and 4.

$$L_{DDPM} = E_{(Z_{vc}, \epsilon_c, t)} [\epsilon_c - \epsilon_{c\theta}(Z_{vct}, t)]^2 + E_{(Z_{vp}, \epsilon_p, t)} [\epsilon_p - \epsilon_{p\theta}(Z_{vpt}, t)]^2 \quad (3)$$

$$L_{CORAL} = \sum_l \sum_l |C(Z_{vct}) - C(Z_{vpt})|^2; C(Z) = \frac{1}{n} \sum_n (Z_i - \bar{Z}_i)(Z_i - \bar{Z}_i)^T \quad (4)$$

ϵ_c and ϵ_p are ground truth noise distributions added to X_c and X_p . $Z_{vct} = TD_{c(e)}(Z_{vc})$ and $Z_{vpt} = TD_{p(e)}(Z_{vp})$ are the noisy representations after t timesteps through $TD_{*(e)}$. $\epsilon_{c\theta}$ and $\epsilon_{p\theta}$ are estimated by $TD_{*(d)}$. \bar{Z} denotes mean. $Z_{v* t}$ is denoised using $\epsilon_{*\theta}$ (Eq. 5) to obtain \mathcal{X}'_c and \mathcal{X}'_p . These are passed through VAE decoders to obtain $\hat{Z}_{tc} = V_{c(d)}(\mathcal{X}'_c)$ and $\hat{Z}_{tp} = V_{p(d)}(\mathcal{X}'_p)$. β_t in Eq. 5 is the variance schedule (Nichol & Dhariwal, 2021) of ϵ_c and ϵ_p at diffusion step time t .

$$\mathcal{X}'_c = \text{denoise}(Z_{vct}, t, \epsilon_{c\theta}); \mathcal{X}'_p = \text{denoise}(Z_{vpt}, t, \epsilon_{p\theta})$$

$$\text{where } \text{denoise}(X_t, t, \epsilon) = \frac{1}{\sqrt{\hat{\alpha}_t}} (X_t - \sqrt{1 - \hat{\alpha}_t} \epsilon); \hat{\alpha}_t = \prod_{i=1}^t (\alpha_i); \alpha_t = 1 - \beta_t \quad (5)$$

To ensure that X_{aug} preserves information from X_c , we use cross-attention Rombach et al. (2022). Given, Z_{vct} and Z_{vpt} , we obtain $Z_{Att} = \text{softmax}(\frac{Z_{vpt} Z_{vct}^T}{\sqrt{l}}) Z_{vct}$. Z_{Att} pays attention to Z_{vct} . Z_{Att} is passed through $TD_{p(d)}$ and denoised using $\epsilon_{p\theta}$ to obtain \hat{Z}_{vpa} . A KL divergence loss L_{KLDA} is also calculated between the distributions of Z_{vp} and \hat{Z}_{vpa} to ensure eventual adherence to $P(X_p)$, as in Equation 6. Additional mean square error terms L_{MSE} between Z_{t*} and \hat{Z}_{t*} and KL divergence terms L_{KLDV} for Z_{v*} are calculated as in Equation 7.

$$L_{KLDA} = 0.5 \sum_{N_*} (-1 + \log(\sigma(\hat{Z}_{vpa})^2) - \log(\sigma(Z_{vp})^2) + \exp(\log(\sigma(Z_{vp})^2) - \log(\sigma(\hat{Z}_{vpa})^2))) + (\mu(Z_{vp}) - \mu(\hat{Z}_{vpa}))^2 \exp(-\log(\sigma(\hat{Z}_{vpa})^2)) \quad (6)$$

$$L_{MSE} = \frac{1}{N_*} \sum_{N_*} (Z_{t*} - \hat{Z}_{t*})^2 \quad (7)$$

$$L_{KLDV} = -(0.5/N_*) \sum_{N_*} (1 + \log(\sigma_*(Z_{v*})^2) - \mu_*(Z_{v*})^2 - \sigma_*(Z_{v*})^2)$$

The overall training loss is $L_{PRE} = L_{ALIGN} + L_{KLDA} + L_{KLDV} + L_{MSE}$. Architecture details are available in Figure 2. The training is done in an unsupervised manner and does not require labeled data.

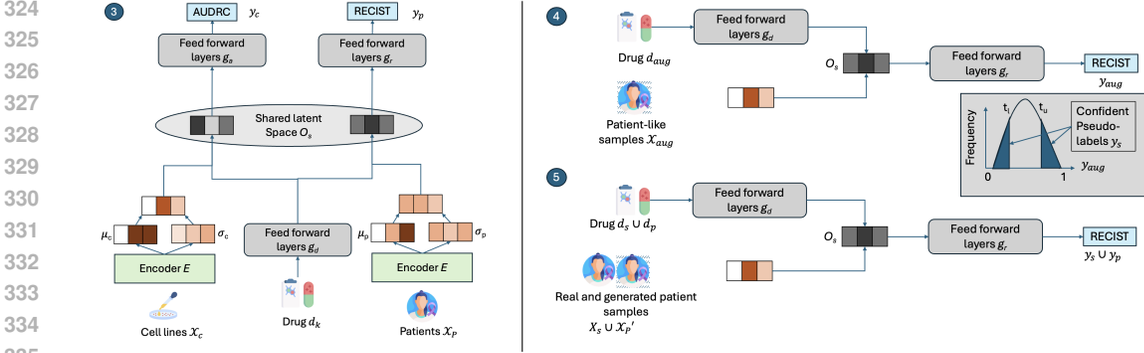


Figure 3: GANDALF architecture for multi-task training (left), pseudolabel generation and selection of confident samples (right, top) and training downstream DRP model (right, bottom). Circled numbers in blue indicate steps from Algorithm 1.

3.2.2 STEP 2: GENERATING NEW PATIENT-LIKE SAMPLES

To generate X_{aug} , we run inference on the trained model using X_c . X_c is first passed through T_e , followed by $V_{c(e)}$, to get Z_{vct} . This is then passed through $TD_{c(e)}$ to get Z_{vct} . This is analogous to removing Z_c from the input samples. As the latent spaces of the DDPMs are already aligned, Z_{vct} can be denoised using $TD_{p(d)}$ to obtain X_{aug} . This step corresponds to introducing Z_p to Z_s . The red arrows in Figure 2, indicates the generation of X_{aug} from X_c .

3.2.3 STEP 3: TRAINING MULTI-TASK LEARNING NETWORK

In this step, the goal is to train a network to assign $y_{aug} \forall (X_{aug}, d_{aug}), d_{aug} \in \mathcal{D}$. A naive approach would involve training a classifier \hat{f} on (X_p, d_p, y_p) and using it to predict y_{aug} . However, $d_p \subset d_{aug}$, since only a small subset of drugs are provided to patients as per clinical guidelines. This implies that $P(X_p, d_p, y_p)$ learnt by \hat{f} may not fully model $P(X_p \cup X_{aug}, d_p \cup d_{aug}, y_p \cup y_{aug})$. During inference, \hat{f} may encounter drugs outside of the training set, yielding noisy y_{aug} . A similar constraint exists in using weak supervision methods (Ratner et al., 2017; Zhang et al., 2022) to assign pseudo-labels. Further, \hat{f} can be prone to overfitting, given the small size of (X_p, d_p, y_p) .

In this step we alleviate overfitting concerns using larger data (X_c, d_c, y_c) , in a multi-task learning (MTL) setup, with additional regularizing loss terms. Moreover, $d_c \simeq \mathcal{D}$, which allows the network to learn from drugs $\notin d_p$. We also capture the shared traits between y_c and y_p by projecting labelled (X_c, d_c) and (X_p, d_p) into a shared latent space O_s , and capture the differences, via two separate prediction heads - a classification head $\hat{y}_p = g_r(X_p, d_p) \in \{0, 1\}$ and a regression head $\hat{y}_c = g_a(X_c, d_c) \in [0, 1]$. O_s is learnt by aligning the latent representations, using CORAL loss (Sun et al., 2016), as in Equation 8. X_c and X_p are first passed through the pretrained encoder network $E(\cdot)$ to obtain μ_c, μ_p, σ_c and σ_p . Sampling S is applied as before to obtain Z_{vc} and Z_{vp} . Z_{vc} and Z_{vp} are concatenated with drug embeddings obtained from a feedforward multi-layer perceptron (MLP) $Z_{d*} = g_d(d_*) \in \mathcal{R}^{N_* \times l}$. The resulting concatenated representations $O_{cd} = Z_{vc} || Z_{dc} \in \mathcal{R}^{N_c \times 2l}$ and $O_{pd} = Z_{vp} || Z_{dp} \in \mathcal{R}^{N_p \times 2l}$ where $||$ denotes concatenation, $N_c = |(X_c, d_c, y_c)|$ and $N_p = |(X_p, d_p, y_p)|$ denote number of labelled sample, drug pairs ($N_p < N_c$).

$$L_{CORALO} = \Sigma_{2l} \Sigma_{2l} \|C(O_{cd}) - C(O_{pd})\|^2; C(Z) = \frac{1}{n} \Sigma_n (Z_i - \bar{Z}_i)(Z_i - \bar{Z}_i)^T \quad (8)$$

O_{cd} is passed through a feed-forward MLP g_a to predict AUDRC values $\hat{y}_c = g_a(O_{cd})$. O_{pd} is passed through another feed forward MLP g_r to predict RECIST values $\hat{y}_p = g_r(O_{pd})$. The entire network is trained to minimise $L_{MTL} = L_{BCE} + L_{MSE} + L_{CORALO}$ as in Equation 9, where $\sigma(x) = \frac{1}{1+e^{-x}}$. MTL architecture is shown in Figure 3(left).

$$L_{BCE} = -\frac{1}{N_p} \Sigma_{N_p} [y_p \log(\sigma(\hat{y}_p)) + (1 - y_p) \log(1 - \sigma(\hat{y}_p))]; L_{MSE} = \frac{1}{N_c} \Sigma_{N_c} (y_c - \hat{y}_c)^2 \quad (9)$$

3.2.4 STEP 4: ASSIGNING PSEUDOLABELS AND SELECTION OF CONFIDENT SAMPLES

To obtain y_{aug} , we first generate all possible $N_c \times |\mathcal{D}|$ pairs $(X_{aug}, d_{aug}), d_{aug} \in \mathcal{D}$. We pass the drug representation d_{aug} through g_d . We concatenate the resulting drug embedding $g_d(d_{aug})$ with

378 X_{aug} . This is then passed through g_r and $\sigma(\cdot)$ to get $y_{aug} \in [0, 1]$, as shown in Figure 3(right, top).
 379 $(X_{aug}, d_{aug}, y_{aug})$ may however be noisy due to incorrect predictions from g_r . Prior work on subset
 380 selection (Lang et al., 2022) has identified that choosing a subset of more confident pseudolabelled
 381 samples is more effective than using the complete pseudolabelled dataset. We use y_{aug} , to select
 382 this subset. y_{bin} is generated by binning y_{aug} into 3 groups, using an upper and lower threshold
 383 t_u and t_l . $y_{bin} = 1$, if $y_{aug} \geq t_u$; $y_{bin} = 0$, if $y_{aug} < t_l$ and $y_{bin} = -1$ otherwise (abstained
 384 samples). Only $N_s < (N_c \times |D|)$ high confidence (non-abstained) samples ($y_{bin} \neq -1$) are used
 385 for the downstream DRP classifier training.

3.2.5 STEP 5: TRAINING DRUG RESPONSE PREDICTION CLASSIFIER

386 The non-abstained, high confidence generated “patient”-drug pairs after pseudo labeling
 387 $((X_s, d_s, y_s)$ of size N_s) are combined with N_p (X'_p, d_p, y_p) pairs to train a drug response pre-
 388 dicting feed forward neural network f (Figure 3, right, bottom). f is trained to minimise BCE loss
 389 in Eq. 10.

$$391 L_{BCE} = -\frac{1}{N_p + N_s} \sum_{N_p + N_s} [y_i \log(\sigma(\hat{y}_i)) + (1 - y_i) \log(1 - \sigma(\hat{y}_i))] \quad (10)$$

392 GANDALF offers several advantages. The use of VAEs and DDPMs makes the model generative
 393 in nature. While generation in DDPMs usually involves sampling from $\mathcal{N}(0, I)$ and denoising, here
 394 the sampling incorporates prior knowledge from X_c . This also enables the use of (X_c, d_c, y_c) in
 395 generating pseudo-labels for X_{aug} . When $N_s > 0$, it reduces chances of overfitting.
 396

397 4 EXPERIMENTS AND RESULTS

398 4.1 DATASETS

399 We used publicly available cell line and patient datasets, for all our experiments. Cell line mutation
 400 profiles were obtained from the Cancer Cell Line Encyclopedia (CCLE) DepMap (v23Q4) (Ghandi
 401 et al., 2019; Barretina et al., 2012). AUDRC responses were obtained from the GDSCv2 (Iorio et al.,
 402 2016; Yang et al., 2012). Patient mutation profiles and associated response labels for drugs were col-
 403 lected from The Cancer Genome Atlas (TCGA) (Weinstein et al., 2013), CbioPortal (CBIO) (Hard-
 404 ing et al., 2019; Nixon et al., 2019; de Bruijn et al., 2023; Gao et al., 2013; Cerami et al., 2012) and
 405 UC San Diego Moores Cancer Center (Moores) (Schwaederle et al., 2016). Patient response, mea-
 406 sured via RECIST were coalesced into binary labels (1: positive response; 0: negative) (Peres da
 407 Silva et al., 2021). Drugs were encoded using 2048 dimensional binary Morgan fingerprints (Mor-
 408 gan, 1965). We exclude samples on multiple drug regimen and retain only patients given a single
 409 drug at a time. This results in 1197 CCLE samples, 541 TCGA, 44 Moores and 84 CBIO patient
 410 samples with documented response labels for 211 drugs in cell lines and 56 drugs across patients. We
 411 restrict our analysis to the 324 genes found in a popular clinical sequencing panel, FoundationOne
 412 CDx (Milbury et al., 2022) and removed samples without mutations in these genes. We also removed
 413 samples with responses to drugs without a Morgan fingerprint. For the transformer pretraining, we
 414 used 71 non-small cell lung cancer and 71 colorectal cancer samples from GENIE (Choudhury et al.,
 415 2023; Garcia et al., 2023), with a documented progression-free survival. We had a total of 156441
 416 train, 17371 validation and 21589 test cell line, drug pairs. We also had 488/488/487 train, 53/54/56
 417 validation and 115/114/113 test patient, drug pairs over 3 folds (folds 0/1/2 respectively) (details in
 418 Appendix Section A.1).

419 4.2 COMPARISON WITH CANCER DRUG RESPONSE PREDICTION METHODS

420 We compared GANDALF against 4 recent state-of-the-art (SOTA) methods which take sample,
 421 drug pairs as model inputs, namely, DruID (Jayagopal et al., 2023), PREDICT-AI (Jayagopal et al.,
 422 2024), drug2me (Zhai & Liu, 2024) and PANCDR (Kim et al., 2024). We also compared GAN-
 423 DALF against CODE-AE (He et al., 2022) and WISER (Shubham et al., 2024), which train separ-
 424 ate models per drug. We report performance metrics on 5 drugs, with samples available in all 3
 425 test folds, namely Cisplatin (Cis), Paclitaxel (Pac), 5-Fluorouracil (Flu), Gemcitabine (Gem) and
 426 Temozolomide (Tem). We do drug-specific model tuning in GANDALF, by only augmenting with
 427 sample, drug pairs for the drug considered. For CODE-AE and WISER, we train separate models
 428 per drug. Apart from GANDALF, only PREDICT-AI could handle varying length inputs. For all
 429 other methods, we converted the mutation profiles into fixed length input vectors of 7776 dimen-
 430 sions, following the pre-processing in (Jayagopal et al., 2023). Validation set correlation between
 431 predicted and actual response was used for early stopping and hyper-parameter selection. As shown
 in Table 1, GANDALF achieves the best AUROC in Flu, Gem, Pac and Tem and second-best in Cis.
 GANDALF achieves the best AUPRC score in Flu, Gem and Pac, and second-best in Cis.

Table 1: Performance comparison across SOTA drug response prediction methods. Best performing results are highlighted in bold, while the second best performing results are underlined.

AUROC (Mean \pm Standard deviation)					
Method	Cis	Flu	Gem	Pac	Tem
GANDALF	<u>0.6343 \pm 0.0306</u>	0.7309 \pm 0.0664	0.6188 \pm 0.0674	0.7728 \pm 0.1253	0.6451 \pm 0.0776
DruID	0.6764 \pm 0.1447	0.6071 \pm 0.1988	<u>0.5092 \pm 0.1005</u>	0.5119 \pm 0.2324	0.6194 \pm 0.0420
PANCDR	0.6278 \pm 0.0308	0.4762 \pm 0.1798	0.4429 \pm 0.2268	0.4236 \pm 0.4168	<u>0.6436 \pm 0.2310</u>
PREDICT-AI	0.5072 \pm 0.0331	0.3869 \pm 0.0372	0.5046 \pm 0.1181	<u>0.6815 \pm 0.1786</u>	0.5350 \pm 0.0606
drug2tme	0.5243 \pm 0.1301	<u>0.7167 \pm 0.1957</u>	0.4568 \pm 0.0857	0.3194 \pm 0.3127	0.5951 \pm 0.2541
WISER	0.4622 \pm 0.1685	0.6095 \pm 0.193	0.4305 \pm 0.0867	0.3641 \pm 0.2522	0.5297 \pm 0.0738
CODE-AE	0.6322 \pm 0.1872	0.5381 \pm 0.1606	0.5085 \pm 0.0503	0.3611 \pm 0.3155	0.4332 \pm 0.3123
AUPRC (Mean \pm Standard deviation)					
Method	Cis	Flu	Gem	Pac	Tem
GANDALF	<u>0.9093 \pm 0.0355</u>	0.8483 \pm 0.0933	0.5874 \pm 0.175	0.9558 \pm 0.024	0.2535 \pm 0.1108
DruID	0.9176 \pm 0.0671	0.7588 \pm 0.1484	0.4515 \pm 0.1297	<u>0.8897 \pm 0.0223</u>	0.3014 \pm 0.1039
PANCDR	0.9018 \pm 0.0324	0.6951 \pm 0.1530	0.4562 \pm 0.2270	0.8561 \pm 0.1019	<u>0.3049 \pm 0.2653</u>
PREDICT-AI	0.8622 \pm 0.0189	0.5885 \pm 0.0581	0.3873 \pm 0.0489	0.8687 \pm 0.1090	0.1373 \pm 0.0050
drug2tme	0.8754 \pm 0.0523	<u>0.8092 \pm 0.1722</u>	<u>0.4826 \pm 0.0947</u>	0.7824 \pm 0.1023	0.3058 \pm 0.1327
WISER	0.8454 \pm 0.0685	0.7505 \pm 0.0657	0.3901 \pm 0.0885	0.7724 \pm 0.1585	0.1762 \pm 0.0243
CODE-AE	0.9059 \pm 0.0521	0.6665 \pm 0.1435	0.4735 \pm 0.0701	0.8208 \pm 0.0574	0.1756 \pm 0.0929

Table 2: Contribution of various components (ablation) in GANDALF, comparisons with other augmentation and pseudolabeling strategies.

Experiment	Method	AUROC (mean \pm std)	AUPRC (mean \pm std)
Ablation	GANDALF	0.8409 \pm 0.0437	0.778 \pm 0.0255
	<i>W/O MTL</i>	0.753 \pm 0.1637	0.6448 \pm 0.1604
	<i>W/O cross-attention</i>	0.752 \pm 0.165	0.6443 \pm 0.1636
	<i>W/O transformer</i>	0.6007 \pm 0.08	0.5632 \pm 0.1101
Augmentation	<i>W perturbation</i>	0.6306 \pm 0.0255	0.5967 \pm 0.0611
	<i>W/O aug</i>	0.6052 \pm 0.0219	0.5784 \pm 0.0394
Pseudolabeling	<i>W majority vote</i>	0.8153 \pm 0.0541	0.756 \pm 0.0827

4.3 ABLATION STUDY

Next, we performed an ablation study to empirically verify the importance of each component in the architecture. We successively removed each component and measured the overall AUROC and AUPRC performance across all the drugs in the test set. The key components of GANDALF are the MTL network for pseudolabeling, cross-attention in pretraining DDPMs and use of transformers to model varying length inputs. We first removed the cell line head in the MTL network (*W/O MTL*). Next, we removed the cross-attention KL divergence loss L_{KLDA} (*W/O cross-attention*). We then removed the use of pretrained transformer (*W/O transformer*) in the input to the network and instead used the 7776 dimensional input used by other SOTA methods. The full model with all components shows the best performance in terms of both AUROC and AUPRC, highlighting the importance of each component in the overall performance (Table 2, Ablation). We also analyse test performance sensitivity to increased volume of pseudolabelled data; details in Appendix Section A.2. A low to moderate volume of high confidence samples is better than large volume of low confidence samples.

4.4 COMPARISON WITH OTHER AUGMENTATION STRATEGIES

There are no known label-invariant mutation data augmentation approaches for cancer DRP (refer Section 2.2 for details). As a baseline, we compare GANDALF against a naive data augmentation approach (Lee et al., 2023), where we perturb the 7776 dimensional inputs, using samples from $\mathcal{N}(0, I)$. This is done once per patient, drug pair (*W perturbation*) in the training data, and the associated label is assumed to remain the same as in the original sample, resulting in a dataset of size $2N_p$. In addition, we also compare GANDALF against a vanilla feed-forward MLP (*W/O aug*), trained using only (X_p, d_p, y_p) . We compare the learning curves (Appendix Figure 6) and test performance metrics (Table 2, Augmentation). In both cases, we fix training epochs. In all folds, no augmentation and Gaussian perturbation strategies result in overfitting, where the validation loss show an increase while the training loss remains low. This is consistent with the fact that smaller datasets can result in overfitting. The test performance metrics for these methods is lower than that of

GANDALF. The slight improvement due to perturbation indicates the benefit of data augmentation in improving overall performance.

4.5 COMPARISON WITH MAJORITY VOTE BASED PSEUDOLABELING

We compared MTL based pseudolabeling strategy against another pseudolabeling strategy similar to Dong-DongChen & WeiGao (2018). The augmented data $(X_{aug}, d_{aug}, y_{aug})$ is passed through 3 separate feed-forward networks, trained on (X_p, d_p, y_p) . The pseudolabels generated by each network is aggregated by majority voting (Lang et al., 2022). As before, non-abstained samples are used to train the downstream DRP model, along with (X_p, d_p, y_p) . The results comparing GANDALF against this approach (*W majority vote*) are shown in Table 2, Pseudolabeling. While the majority voting strategy does perform well, GANDALF outperforms it in overall AUROC and AUPRC. This may be potentially due to the use of the larger cell line labelled data, with more drugs, as opposed to the smaller labelled patient dataset.

5 CONCLUSIONS AND DISCUSSION

In this paper, we propose GANDALF, a generative patient data augmentation framework, to tackle the challenge of training a cancer DRP model with limited labelled data. Unlike prior DRP methods that augment data in the shared space between patients and cell lines, we utilise the larger labelled cell line dataset to generate more patient-like samples as well as their pseudo-labels. GANDALF outperforms SOTA DRP methods, and also shows improved performance when compared to baseline genomic data augmentation and pseudo labeling approaches. GANDALF has a large number of parameters and sub-modules, each of which needs pretraining, increasing overall training time. Learning the underlying data distributions is limited by available labelled cell lines and patients.

There are several future directions to explore, which may improve GANDALF further. In this paper, we have only considered labelled patient profiles for training, although the pretraining stage supports unlabelled data. Future work can evaluate the use of unlabelled patient profiles in all steps of training. We examined the quality of the generated samples by comparing the distributions against the original patient data. More extensive studies to examine the biological significance of the generated samples and their fidelity can shed light on the patterns captured by the model. Generative strategies, which can incorporate known biological information on co-occurring mutations, can also be explored in the future. Overall, GANDALF sets the stage for using generative techniques in the field of cancer DRP research, and emphasises the importance of capturing patient domain-specific characteristics for improving downstream prediction performance.

6 REPRODUCIBILITY

Our code and data are made publicly available at <https://anonymous.4open.science/r/GANDALF>.

REFERENCES

- Jordi Barretina, Giordano Caponigro, Nicolas Stransky, Kavitha Venkatesan, Adam A Margolin, Sungjoon Kim, Christopher J Wilson, Joseph Lehár, Gregory V Kryukov, Dmitriy Sonkin, et al. The cancer cell line encyclopedia enables predictive modelling of anticancer drug sensitivity. *Nature*, 483(7391):603–607, 2012.
- Pavla Brachova, Kristina W Thiel, and Kimberly K Leslie. The consequence of oncomorphic tp53 mutations in ovarian cancer. *International journal of molecular sciences*, 14(9):19257–19275, 2013.
- Ethan Cerami, Jianjiong Gao, Ugur Dogrusoz, Benjamin E Gross, Selcuk Onur Sumer, Bülent Arman Aksoy, Anders Jacobsen, Caitlin J Byrne, Michael L Heuer, Erik Larsson, et al. The cbio cancer genomics portal: an open platform for exploring multidimensional cancer genomics data. *Cancer discovery*, 2(5):401–404, 2012.
- Junjie Chen, Mohammad Erfan Mowlaei, and Xinghua Shi. Population-scale genomic data augmentation based on conditional generative adversarial networks. In *Proceedings of the 11th ACM International Conference on Bioinformatics, Computational Biology and Health Informatics*, pp. 1–6, 2020.

- 540 Noura J Choudhury, Jessica A Lavery, Samantha Brown, Ino de Bruijn, Justin Jee, Thinh Ngoc Tran,
541 Hira Rizvi, Kathryn C Arbour, Karissa Whiting, Ronglai Shen, et al. The genie bpc nslc cohort:
542 A real-world repository integrating standardized clinical and genomic data for 1,846 patients with
543 non-small cell lung cancer. *Clinical Cancer Research*, 29(17):3418–3428, 2023.
- 544 Francis S Collins and Harold Varmus. A new initiative on precision medicine. *New England journal*
545 *of medicine*, 372(9):793–795, 2015.
- 547 Ramon Colomer, Jesús Miranda, Nuria Romero-Laorden, Javier Hornedo, Lucía González-Cortijo,
548 Silvana Mouron, Maria J Bueno, Rebeca Mondéjar, and Miguel Quintela-Fandino. Usefulness
549 and real-world outcomes of next generation sequencing testing in patients with cancer: an obser-
550 vational study on the impact of selection based on clinical judgement. *EClinicalMedicine*, 60,
551 2023.
- 552 T Conroy, P Pfeiffer, V Vilgrain, Angela Lamarca, T Seufferlein, EM O’Reilly, T Hackert, T Golan,
553 G Prager, K Haustermans, et al. Pancreatic cancer: Esmo clinical practice guideline for diagnosis,
554 treatment and follow-up. *Annals of oncology*, 34(11):987–1002, 2023.
- 555 Supratim Das and Xinghua Shi. Offspring gan augments biased human genomic data. In *Proceed-*
556 *ings of the 13th ACM International Conference on Bioinformatics, Computational Biology and*
557 *Health Informatics*, pp. 1–10, 2022.
- 559 Saloni Dattani, Fiona Spooner, Hannah Ritchie, and Max Roser. Causes of death. *Our World in*
560 *Data*, 2023. <https://ourworldindata.org/causes-of-death>.
- 561 Ino de Bruijn, Ritika Kundra, Brooke Mastrogiacomo, Thinh Ngoc Tran, Luke Sikina, Tali Mazor,
562 Xiang Li, Angelica Ochoa, Gaofei Zhao, Bryan Lai, et al. Analysis and visualization of longitu-
563 dinal genomic and clinical data from the aacr project genie biopharma collaborative in cbiportal.
564 *Cancer research*, 83(23):3861–3867, 2023.
- 566 W Dong-DongChen and ZH WeiGao. Tri-net for semi-supervised deep learning. In *Proceedings of*
567 *twenty-seventh international joint conference on artificial intelligence*, pp. 2014–2020, 2018.
- 568 Andrew G Duncan, Jennifer A Mitchell, and Alan M Moses. Improving the performance of super-
569 vised deep learning for regulatory genomics using phylogenetic augmentation. *Bioinformatics*,
570 40(4):btae190, 2024.
- 571 Jianjiong Gao, Bülent Arman Aksoy, Ugur Dogrusoz, Gideon Dresdner, Benjamin Gross, S Onur
572 Sumer, Yichao Sun, Anders Jacobsen, Rileen Sinha, Erik Larsson, et al. Integrative analysis of
573 complex cancer genomics and clinical profiles using the cbiportal. *Science signaling*, 6(269):
574 p11–p11, 2013.
- 576 Enrique Sanz Garcia, Eric Chen, Marios Giannakis, Gregory J Riely, Jeremy L Warner, Michele L
577 LeNoue-Newton, Jessica Weiss, Katrina Hueniken, Kenneth L Kehl, Deborah Schrag, et al. Ge-
578 nomic characteristics and clinical outcomes of early onset colorectal cancer (eocr): Findings
579 from aacr project genie biopharma collaborative registry. *Cancer Research*, 83(7_Supplement):
580 1177–1177, 2023.
- 581 Mahmoud Ghandi, Franklin W Huang, Judit Jané-Valbuena, Gregory V Kryukov, Christopher C
582 Lo, E Robert McDonald III, Jordi Barretina, Ellen T Gelfand, Craig M Bielski, Haoxin Li, et al.
583 Next-generation characterization of the cancer cell line encyclopedia. *Nature*, 569(7757):503–
584 508, 2019.
- 585 Joseph J Hale, Takeshi Matsui, Ilan Goldstein, Martin N Mullis, Kevin R Roy, Christopher Ne Ville,
586 Darach Miller, Charley Wang, Trevor Reynolds, Lars M Steinmetz, et al. Genome-scale analysis
587 of interactions between genetic perturbations and natural variation. *Nature Communications*, 15
588 (1):4234, 2024.
- 589 James J Harding, Subhiksha Nandakumar, Joshua Armenia, Danny N Khalil, Melanie Albano,
590 Michele Ly, Jinru Shia, Jaclyn F Hechtman, Ritika Kundra, Imane El Dika, et al. Prospective
591 genotyping of hepatocellular carcinoma: clinical implications of next-generation sequencing for
592 matching patients to targeted and immune therapies. *Clinical Cancer Research*, 25(7):2116–2126,
593 2019.

- 594 Di He, Qiao Liu, You Wu, and Lei Xie. A context-aware deconfounding autoencoder for robust
595 prediction of personalized clinical drug response from cell-line compound screening. *Nature*
596 *Machine Intelligence*, 4(10):879–892, 2022.
- 597 Jonathan Ho, Ajay Jain, and Pieter Abbeel. Denoising diffusion probabilistic models. *Advances in*
598 *neural information processing systems*, 33:6840–6851, 2020.
- 600 Francesco Iorio, Theo A Knijnenburg, Daniel J Vis, Graham R Bignell, Michael P Menden, Michael
601 Schubert, Nanne Aben, Emanuel Gonçalves, Syd Barthorpe, Howard Lightfoot, et al. A landscape
602 of pharmacogenomic interactions in cancer. *Cell*, 166(3):740–754, 2016.
- 603 Aishwarya Jayagopal, Robert J Walsh, Krishna Kumar Hariprasannan, Raguathan Mariappan, De-
604 babrata Mahapatra, Patrick William Jaynes, Diana Lim, David Shao Peng Tan, Tuan Zea Tan,
605 Jason J Pitt, et al. A multi-task domain-adapted model to predict chemotherapy response from
606 mutations in recurrently altered cancer genes. *medRxiv*, pp. 2023–11, 2023.
- 607 Aishwarya Jayagopal, Hansheng Xue, Ziyang He, Robert J Walsh, Krishna Kumar Hariprasannan,
608 David Shao Peng Tan, Tuan Zea Tan, Jason J Pitt, Anand D Jeyasekharan, and Vaibhav Rajan.
609 Personalised drug identifier for cancer treatment with transformers using auxiliary information. In
610 *Proceedings of the 30th ACM SIGKDD Conference on Knowledge Discovery and Data Mining*,
611 pp. 5138–5149, 2024.
- 612 Patrick Kage, Jay C Rothenberger, Pavlos Andreadis, and Dimitrios I Diochnos. A review of pseudo-
613 labeling for computer vision. *arXiv preprint arXiv:2408.07221*, 2024.
- 614 Cherry Khosla and Baljit Singh Saini. Enhancing performance of deep learning models with dif-
615 ferent data augmentation techniques: A survey. In *2020 International Conference on Intelligent*
616 *Engineering and Management (ICIEM)*, pp. 79–85. IEEE, 2020.
- 617 Juyeon Kim, Sung-Hye Park, and Hyunju Lee. Pancdr: precise medicine prediction using an adver-
618 sarial network for cancer drug response. *Briefings in Bioinformatics*, 25(2):bbae088, 2024.
- 619 Diederik P Kingma and Max Welling. Auto-encoding variational bayes. *arXiv preprint*
620 *arXiv:1312.6114*, 2013.
- 621 Akim Kotelnikov, Dmitry Baranchuk, Ivan Rubachev, and Artem Babenko. Tabddpm: Modelling
622 tabular data with diffusion models. In *International Conference on Machine Learning*, pp. 17564–
623 17579. PMLR, 2023.
- 624 Alice Lacan, Michèle Sebag, and Blaise Hanczar. Gan-based data augmentation for transcriptomics:
625 survey and comparative assessment. *Bioinformatics*, 39(Supplement_1):i111–i120, 2023.
- 626 Melissa J Landrum, Jennifer M Lee, Mark Benson, Garth R Brown, Chen Chao, Shanmuga Chi-
627 tipiralla, Baoshan Gu, Jennifer Hart, Douglas Hoffman, Wonhee Jang, et al. Clinvar: improving
628 access to variant interpretations and supporting evidence. *Nucleic acids research*, 46(D1):D1062–
629 D1067, 2018.
- 630 Hunter Lang, Aravindan Vijayaraghavan, and David Sontag. Training subset selection for weak
631 supervision. *Advances in Neural Information Processing Systems*, 35:16023–16036, 2022.
- 632 Dong-Hyun Lee et al. Pseudo-label: The simple and efficient semi-supervised learning method for
633 deep neural networks. In *Workshop on challenges in representation learning, ICML*, volume 3,
634 pp. 896. Atlanta, 2013.
- 635 Hyunjung Lee, Utku Ozbulak, Homin Park, Stephen Depuydt, Wesley De Neve, and Joris Vanker-
636 schaver. Assessing the reliability of point mutation as data augmentation for deep learning with
637 genomic data. *BMC bioinformatics*, 25(1):170, 2024.
- 638 Mihee Lee and Vladimir Pavlovic. Private-shared disentangled multimodal vae for learning of la-
639 tent representations. In *Proceedings of the IEEE/CVF conference on computer vision and pattern*
640 *recognition*, pp. 1692–1700, 2021.

- 648 Nicholas Keone Lee, Ziqi Tang, Shushan Toneyan, and Peter K Koo. Evoaug: improving gener-
649 alization and interpretability of genomic deep neural networks with evolution-inspired data aug-
650 mentations. *Genome Biology*, 24(1):105, 2023.
- 651
652 Ginny XH Li, Dan Munro, Damian Fermin, Christine Vogel, and Hyungwon Choi. A protein-
653 centric approach for exome variant aggregation enables sensitive association analysis with clinical
654 outcomes. *Human mutation*, 41(5):934–945, 2020.
- 655
656 Jinzhuang Liao, Xiaoying Li, Yu Gan, Shuangze Han, Pengfei Rong, Wei Wang, Wei Li, and
657 Li Zhou. Artificial intelligence assists precision medicine in cancer treatment. *Frontiers in oncol-*
658 *ogy*, 12:998222, 2023.
- 659
660 Ruishan Liu, Shemra Rizzo, Sarah Waliany, Marius Rene Garmhausen, Navdeep Pal, Zhi Huang,
661 Nayan Chaudhary, Lisa Wang, Chris Harbron, Joel Neal, et al. Systematic pan-cancer analysis of
662 mutation–treatment interactions using large real-world clinicogenomics data. *Nature Medicine*,
28(8):1656–1661, 2022.
- 663
664 Gilberto Lopes. The global economic cost of cancer—estimating it is just the first step! *JAMA*
665 *oncology*, 9(4):461–462, 2023.
- 666
667 Jianzhu Ma, Samson H Fong, Yunan Luo, Christopher J Bakkenist, John Paul Shen, Soufiane Mour-
668 ragui, Lodewyk FA Wessels, Marc Hafner, Roded Sharan, Jian Peng, et al. Few-shot learning
669 creates predictive models of drug response that translate from high-throughput screens to individ-
670 ual patients. *Nature Cancer*, 2(2):233–244, 2021.
- 671
672 Coren A Milbury, James Creeden, Wai-Ki Yip, David L Smith, Varun Pattani, Kristi Maxwell,
673 Bethany Sawchyn, Ole Gjoerup, Wei Meng, Joel Skoletsky, et al. Clinical and analytical valida-
674 tion of foundationone® cdx, a comprehensive genomic profiling assay for solid tumors. *PLoS*
675 *One*, 17(3):e0264138, 2022.
- 676
677 Harry L Morgan. The generation of a unique machine description for chemical structures—a tech-
678 nique developed at chemical abstracts service. *Journal of chemical documentation*, 5(2):107–113,
679 1965.
- 680
681 Van K Morris, Erin B Kennedy, Nancy N Baxter, Al B Benson III, Andrea Cercek, May Cho, Kristen
682 K Ciombor, Chiara Cremolini, Anjee Davis, Dustin A Deming, et al. Treatment of metastatic
683 colorectal cancer: Asco guideline. *Journal of Clinical Oncology*, 41(3):678–700, 2023.
- 684
685 Soufiane Mourragui, Marco Loog, Mark A Van De Wiel, Marcel JT Reinders, and Lodewyk FA
686 Wessels. Precise: a domain adaptation approach to transfer predictors of drug response from
687 pre-clinical models to tumors. *Bioinformatics*, 35(14):i510–i519, 2019.
- 688
689 Soufiane MC Mourragui, Marco Loog, Daniel J Vis, Kat Moore, Anna G Manjon, Mark A van de
690 Wiel, Marcel JT Reinders, and Lodewyk FA Wessels. Predicting patient response with models
691 trained on cell lines and patient-derived xenografts by nonlinear transfer learning. *Proceedings of*
692 *the National Academy of Sciences*, 118(49):e2106682118, 2021.
- 693
694 Alexander Quinn Nichol and Prafulla Dhariwal. Improved denoising diffusion probabilistic models.
695 In *International conference on machine learning*, pp. 8162–8171. PMLR, 2021.
- 696
697 Mellissa J Nixon, Luigi Formisano, Ingrid A Mayer, M Valeria Estrada, Paula I González-Ericsson,
698 Steven J Isakoff, Andrés Forero-Torres, Helen Won, Melinda E Sanders, David B Solit, et al.
699 Pik3ca and map3k1 alterations imply luminal a status and are associated with clinical benefit from
700 pan-pi3k inhibitor buparlisib and letrozole in er+ metastatic breast cancer. *NPJ Breast Cancer*, 5
701 (1):31, 2019.
- 702
703 Sinno Jialin Pan and Qiang Yang. A survey on transfer learning. *IEEE Transactions on knowledge*
704 *and data engineering*, 22(10):1345–1359, 2009.
- 705
706 Rafael Peres da Silva, Chayaporn Suphavilai, and Niranjana Nagarajan. Tugda: task uncertainty
707 guided domain adaptation for robust generalization of cancer drug response prediction from in
708 vitro to in vivo settings. *Bioinformatics*, 37(Supplement_1):i76–i83, 2021.

- 702 D Planchard, ST Popat, K Kerr, S Novello, EF Smit, Corinne Faivre-Finn, TS Mok, M Reck,
703 PE Van Schil, MD Hellmann, et al. Metastatic non-small cell lung cancer: Esmo clinical practice
704 guidelines for diagnosis, treatment and follow-up. *Annals of Oncology*, 29:iv192–iv237, 2018.
705
- 706 Tijana Randic, Stefano Magni, Demetra Philippidou, Christiane Margue, Kamil Grzyb, Jasmin Re-
707 nate Preis, Joanna Patrycja Wroblewska, Petr V Nazarov, Michel Mittelbronn, Katrin BM
708 Frauenknecht, et al. Single-cell transcriptomics of nras-mutated melanoma transitioning to drug
709 resistance reveals p2rx7 as an indicator of early drug response. *Cell Reports*, 42(7), 2023.
- 710 Alexander Ratner, Stephen H Bach, Henry Ehrenberg, Jason Fries, Sen Wu, and Christopher Ré.
711 Snorkel: Rapid training data creation with weak supervision. In *Proceedings of the VLDB en-
712 dowment. International conference on very large data bases*, volume 11, pp. 269. NIH Public
713 Access, 2017.
- 714 Robin Rombach, Andreas Blattmann, Dominik Lorenz, Patrick Esser, and Björn Ommer. High-
715 resolution image synthesis with latent diffusion models. In *Proceedings of the IEEE/CVF confer-
716 ence on computer vision and pattern recognition*, pp. 10684–10695, 2022.
- 717 Yuki Saito, Junji Koya, and Keisuke Kataoka. Multiple mutations within individual oncogenes.
718 *Cancer science*, 112(2):483–489, 2021.
- 720 Maria Schwaederle, Barbara A Parker, Richard B Schwab, Gregory A Daniels, David E Piccioni,
721 Santosh Kesari, Teresa L Helsten, Lyudmila A Bazhenova, Julio Romero, Paul T Fanta, et al.
722 Precision oncology: The uc san diego moores cancer center predict experience. *Molecular cancer
723 therapeutics*, 15(4):743–752, 2016.
- 724 Hossein Sharifi-Noghabi, Shuman Peng, Olga Zolotareva, Colin C Collins, and Martin Ester. Aitl:
725 adversarial inductive transfer learning with input and output space adaptation for pharmaco-
726 genomics. *Bioinformatics*, 36(Supplement_1):i380–i388, 2020.
- 727
728 Hossein Sharifi-Noghabi, Parsa Alamzadeh Harjandi, Olga Zolotareva, Colin C Collins, and Martin
729 Ester. Out-of-distribution generalization from labelled and unlabelled gene expression data for
730 drug response prediction. *Nature Machine Intelligence*, 3(11):962–972, 2021.
- 731 Connor Shorten and Taghi M Khoshgoftaar. A survey on image data augmentation for deep learning.
732 *Journal of big data*, 6(1):1–48, 2019.
- 733
734 Kumar Shubham, Aishwarya Jayagopal, Syed Mohammed Danish, Prathosh AP, and Vaibhav Ra-
735 jan. Wiser: Weak supervision and supervised representation learning to improve drug response
736 prediction in cancer. *arXiv preprint arXiv:2405.04078*, 2024.
- 737 Alona Sosinsky, John Ambrose, William Cross, Clare Turnbull, Shirley Henderson, Louise Jones,
738 Angela Hamblin, Prabhu Arumugam, Georgia Chan, Daniel Chubb, et al. Insights for precision
739 oncology from the integration of genomic and clinical data of 13,880 tumors from the 100,000
740 genomes cancer programme. *Nature Medicine*, 30(1):279–289, 2024.
- 741 Baochen Sun, Jiashi Feng, and Kate Saenko. Return of frustratingly easy domain adaptation. In
742 *Proceedings of the AAAI conference on artificial intelligence*, volume 30, 2016.
- 743
744 Adam Wahida, Lars Buschhorn, Stefan Fröhling, Philipp J Jost, Andreas Schneeweiss, Peter Lichter,
745 and Razelle Kurzrock. The coming decade in precision oncology: six riddles. *Nature Reviews
746 Cancer*, 23(1):43–54, 2023.
- 747 Kai Wang, Mingyao Li, and Hakon Hakonarson. Annovar: functional annotation of genetic variants
748 from high-throughput sequencing data. *Nucleic acids research*, 38(16):e164–e164, 2010.
- 749
750 Bo Wei, John Kang, Miho Kibukawa, Gladys Arreaza, Maureen Maguire, Lei Chen, Ping Qiu, Lixin
751 Lang, Deepti Aurora-Garg, Razvan Cristescu, et al. Evaluation of the trusight oncology 500 assay
752 for routine clinical testing of tumor mutational burden and clinical utility for predicting response
753 to pembrolizumab. *The Journal of Molecular Diagnostics*, 24(6):600–608, 2022.
- 754 John N Weinstein, Eric A Collisson, Gordon B Mills, Kenna R Shaw, Brad A Ozenberger, Kyle
755 Ellrott, Ilya Shmulevich, Chris Sander, and Joshua M Stuart. The cancer genome atlas pan-cancer
analysis project. *Nature genetics*, 45(10):1113–1120, 2013.

756 Wanjuan Yang, Jorge Soares, Patricia Greninger, Elena J Edelman, Howard Lightfoot, Simon
757 Forbes, Nidhi Bindal, Dave Beare, James A Smith, I Richard Thompson, et al. Genomics of
758 drug sensitivity in cancer (gdsc): a resource for therapeutic biomarker discovery in cancer cells.
759 *Nucleic acids research*, 41(D1):D955–D961, 2012.

760 Yiyang Yu, Shivani Muthukumar, and Peter K Koo. Evoaug-tf extending evolution-inspired data
761 augmentations for genomic deep learning to tensorflow. *Bioinformatics*, 40(3):btae092, 2024.

762 Jia Zhai and Hui Liu. Cross-domain feature disentanglement for interpretable modeling of tumor
763 microenvironment impact on drug response. *IEEE Journal of Biomedical and Health Informatics*,
764 2024.

765 Jieyu Zhang, Cheng-Yu Hsieh, Yue Yu, Chao Zhang, and Alexander Ratner. A survey on program-
766 matic weak supervision. *arXiv preprint arXiv:2202.05433*, 2022.

767 A APPENDIX

768 A.1 EXPERIMENT SETTINGS

769 A.1.1 DRUG SELECTION CRITERIA

770 The patient dataset we used had 56 drugs. For each of the 56 drugs in patients, we first consider those
771 with at least 20 labelled patient samples (He et al., 2022) - this reduced labelled data to 12 drugs.
772 For each drug, we divided the samples into groups based on cancer type and data source. Each
773 group with > 20 samples was divided into 2:1 ratio in 3-fold label based stratified cross-validation.
774 For some groups, no test samples were available. We excluded these to get 7 drugs. These drugs
775 were used in the ablation studies in Table 2, to report overall performance metrics. We removed
776 drugs which had < 3 positive samples as it would cause issues in CV, where one fold may have test
777 samples with only a single label - this resulted in the five drugs shown in Table 1.

778 A.1.2 TRAIN-TEST SPLIT

779 RECIST labels in patients were initially coalesced into 2 groups - Complete and Partial Response
780 as label 1 (good response), Stable and Progressive Disease as label 0 (bad response). The labelled
781 patient samples obtained were grouped based on the drug, cancer type and source of dataset (TCGA,
782 Moores, CBIO). Each group with ≥ 20 samples was divided into 3-fold cross validation train-test
783 splits, stratified by label. Groups with < 20 samples were only used for training. The train and test
784 labelled samples across all groups and folds were combined to form 3 train-test folds respectively.
785 Each of the 3 train folds were further divided in a 90:10 ratio to obtain a train-validation split. Cell
786 line data was also grouped in a similar fashion and divided into a single train-validation and test
787 fold. The training and evaluation in all cases use sample, drug pairs where the sample could be from
788 either domain. We had a total of 156441 train, 17371 validation and 21589 test cell line, drug pairs.
789 We also had 488/488/487 train, 53/54/56 validation and 115/114/113 test patient, drug pairs over the
790 3 folds. We run inference on test patient, drug pairs, and report the average AUROC and AUPRC
791 metrics across 3 test folds.

792 A.2 SENSITIVITY TO VOLUME OF PSEUDOLABELLED DATA

793 We examined the sensitivity of the overall model performance to increasing the quantity of pseu-
794 dolabelled data. We change the amount of pseudolabelled data by varying the upper and lower
795 thresholds t_u and t_l . Increasing t_l and decreasing t_u is equivalent to adding more pseudolabelled
796 samples. We varied t_l from 0.1 to 0.4, t_u from 0.5 to 0.9, in increments of 0.1. In all cases, only
797 a single parameter was changed while all others were left constant. Figure 4 indicates that a lower
798 value of t_l shows better performance. This may result in fewer non-abstained samples with label 0,
799 and improve performance in the samples selected for the downstream DRP task. A higher t_u in gen-
800 eral improves performance with 0.8 yielding the best. If t_u is too low or t_l is too high, it may admit
801 more low-confidence samples, y_{aug} being closer to 0.5. If t_u is too high, it may drastically reduce
802 the number of positive labels available for downstream DRP training, also reducing performance.
803 Table 3 indicates the number of pseudolabelled samples added in each case.

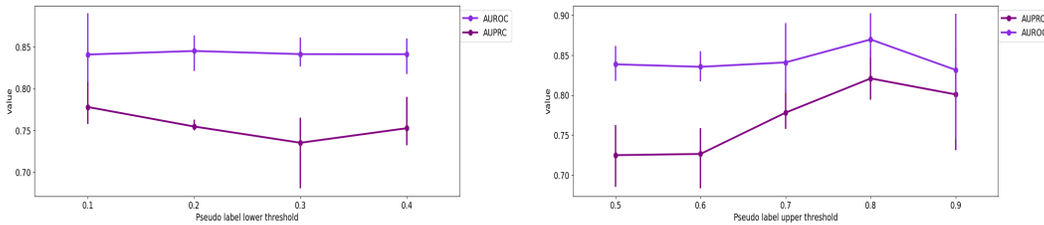


Figure 4: Sensitivity tests on value of pseudo label lower (left) and upper (right) thresholds.

Lower threshold values	Fold 0 psuedolabelled responders / non-responders	Fold 1 psuedolabelled responders / non-responders	Fold 2 psuedolabelled responders / non-responders
0.1	3830/60101	874/15668	241/7157
0.2	3830/192454	874/125098	241/81011
0.3	3830/355849	874/323572	241/274803
0.4	3830/481589	874/479348	241/462177
Upper threshold values	Fold 0 psuedolabelled responders / non-responders	Fold 1 psuedolabelled responders / non-responders	Fold 2 psuedolabelled responders / non-responders
0.5	29599/60101	25932/15668	25554/7157
0.6	9568/60101	6336/15668	4023/7157
0.7	3830/60101	874/15668	241/7157
0.8	1578/60101	27/15668	0/7157
0.9	500/60101	0/15668	0/7157

Table 3: Number of pseudolabelled samples used in sensitivity test of thresholds.

A.3 SENSITIVITY TO DIFFERENT AMOUNTS OF TRAINING DATA

We conducted two experiments to evaluate the effect of varying amounts of real and synthetic patient data.

A.3.1 EFFECT OF VARYING AMOUNTS OF PSEUDOLABELLED DATA

We retain all the real train patient data and randomly sample 25%, 50%, 75% and 100% of the generated pseudolabelled data, and use this in training the DRP model. 0% setting indicates no augmented data in the DRP training. Results are shown in the Table 4 below. 0% does the worst, without any augmentation. Best AUROC is at 50% addition of pseudolabelled data, best AUPRC at 25% pseudolabelled data. Across 25-100% settings, the difference in performance is not statistically significant. For the case of 0% vs any other level of augmentation, differences are statistically significant, indicating that adding psuedolabelled data improves performance. To answer the question of how much pseudolabelled is helpful we will need further studies on possibly larger datasets.

% of pseudolabelled data	Average AUROC over 3 folds	Average AUPRC over 3 folds	Number of pseudo labelled patient data (fold 0)	Number of real labelled patient data (fold 0)
0%	0.5263 \pm 0.0195	0.5229 \pm 0.0249	0	488
25%	0.8584 \pm 0.0361	0.7838 \pm 0.0564	15983	488
50%	0.8613 \pm 0.0279	0.7796 \pm 0.0437	31966	488
75%	0.8577 \pm 0.0269	0.7677 \pm 0.0354	47948	488
100%	0.8409 \pm 0.0437	0.778 \pm 0.0255	63931	488

Table 4: Performance comparison for varying quantities of pseudolabelled data

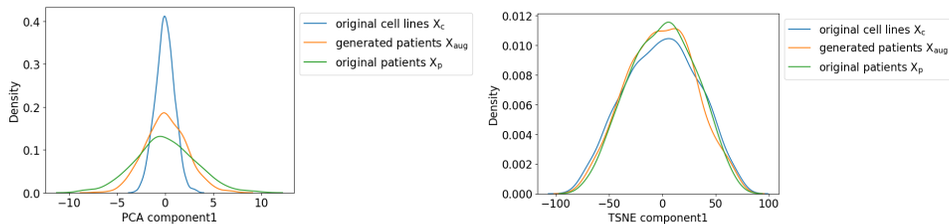


Figure 5: Kernel Density Estimation plot comparing the distribution of first PCA component (left) and first TSNE component (right) across original cell line, real patient and generated patient data

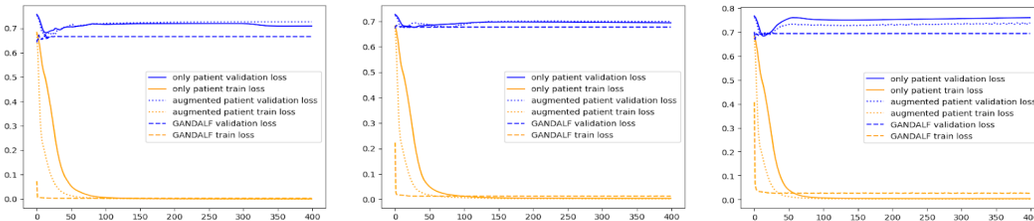


Figure 6: Learning curves on (left to right) 3 cross-validation folds, with orange line indicating train loss and blue indicating validation loss. Dotted lines indicate augmentation with Gaussian perturbation, solid lines indicate no augmentation, dashed lines indicate GANDALF augmentation.

A.3.2 EFFECT OF VARYING AMOUNTS OF REAL PATIENT DATA

We randomly sample 25%, 50%, 75% and 100% of real labelled patient data. In each case we sample twice the number of real samples from the pseudolabelled data. 100% setting thus refers to 3 times the size of real labelled patient data. In general, as seen in Table 5, as more real labelled data is added performance improves, as generally expected.

A.4 HYPERPARAMETER SELECTION

For baseline models, we used the hyperparameter ranges defined in each of the papers. We did a hyperparameter sweep over these ranges using Bayesian Optimization for maximum of 15 runs, to determine the best hyperparameters for our dataset. For GANDALF, we mainly focused on the hyperparameters in the supervised training stages, key being the lower and upper thresholds and learning rate parameters for the DRP and MTL models. We varied the lower threshold between 0.1 to 0.5 and upper threshold from 0.5 to 0.9, with increments done based on quantiles calculated from predicted probability of response after MTL training. This was done for each drug separately. The hidden layers from the VAE were set to 64 dimensions based on our GPU memory restrictions and batch size was 512. For the transformer we used 64 dimensional embeddings, 4 heads and 8 encoder layers. For the cell line VAE, we used 1024, 128 and 64 hidden units and for patient VAE we used 512, 128 and 64 hidden units. Both VAEs used tanh activation. The DDPM uses linear layers with dimensions as the VAE representation size, and uses dropout and ReLU. The MTL network uses 2 linear layers each in embedding drugs, predicting RECIST and predicting AUDRC, with ReLU

% of real data (pseudolabelled data = 2 x real data)	Average AUROC over 3 folds	Average AUPRC over 3 folds	Number of pseudo labelled patient data (fold 0)	Number of real labelled patient data (fold 0)
25%	0.5326 ± 0.0152	0.5239 ± 0.0222	244	122
50%	0.581 ± 0.0216	0.5505 ± 0.0274	488	244
75%	0.6888 ± 0.0257	0.638 ± 0.0348	732	366
100%	0.7086 ± 0.0247	0.6533 ± 0.0374	976	488

Table 5: Performance comparison for varying quantities of real patient data

Table 6: Performance comparison across different ablation tests, where each test removes one component from GANDALF. Best performing results are highlighted in bold.

AUROC (Mean \pm Standard deviation)					
Method	Cis	Flu	Gem	Pac	Tem
GANDALF	0.6343 \pm 0.0306	0.7309 \pm 0.0664	0.6188 \pm 0.0674	0.7728 \pm 0.1253	0.6451 \pm 0.0776
<i>W/O MTL</i>	0.3409 \pm 0.219	0.5333 \pm 0.075	0.5587 \pm 0.1787	0.2758 \pm 0.1461	0.7513 \pm 0.0805
<i>W/O cross-attention</i>	0.6061 \pm 0.0475	0.7309 \pm 0.0834	0.6188 \pm 0.0674	0.7728 \pm 0.1253	0.6152 \pm 0.1074
<i>W/O transformer</i>	0.3735 \pm 0.1404	0.4143 \pm 0.1122	0.5718 \pm 0.0805	0.5625 \pm 0.3903	0.2106 \pm 0.0457
<i>W/O VAE</i>	Out of memory issues				
<i>W/O DDPM</i>	0.4849 \pm 0.0909	0.6929 \pm 0.1189	0.5162 \pm 0.1247	0.5208 \pm 0.4161	0.3138 \pm 0.0647
AUPRC (Mean \pm Standard deviation)					
Method	Cis	Flu	Gem	Pac	Tem
GANDALF	0.9093 \pm 0.0355	0.8483 \pm 0.0933	0.5874 \pm 0.175	0.9558 \pm 0.024	0.2535 \pm 0.1108
<i>W/O MTL</i>	0.8101 \pm 0.0793	0.7345 \pm 0.1	0.5697 \pm 0.0628	0.7582 \pm 0.1012	0.3215 \pm 0.1623
<i>W/O cross-attention</i>	0.9183 \pm 0.0255	0.8483 \pm 0.0967	0.5873 \pm 0.1753	0.9558 \pm 0.024	0.2068 \pm 0.0585
<i>W/O transformer</i>	0.8047 \pm 0.0417	0.6400 \pm 0.1231	0.4760 \pm 0.0865	0.8478 \pm 0.1374	0.0993 \pm 0.0195
<i>W/O VAE</i>	Out of memory issues				
<i>W/O DDPM</i>	0.8696 \pm 0.0098	0.8241 \pm 0.0741	0.4932 \pm 0.1867	0.8273 \pm 0.1636	0.1150 \pm 0.0263

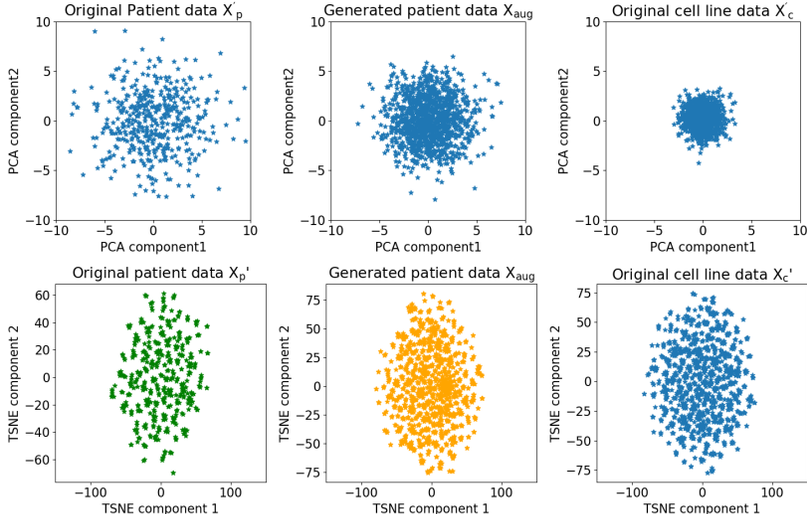


Figure 7: Comparison of distribution (left to right) across real and generated data, using PCA (top) and TSNE (bottom) methods.

activation. Max epochs were set to 500 for pretraining, 100 for the MTL and DRP training. Early stopping was done using patient validation set pearson correlation as in (Sharifi-Noghabi et al., 2021).

A.5 ADDITIONAL ABLATION EXPERIMENTS

To understand the contribution of each individual component, we added additional ablation studies where each test removes just one component of the architecture. We also performed drug specific tuning in each case. Table 6 shows the results of each ablation test. Apart from the ablation tests in Table 2, we also added two more tests *W/O VAE* and *W/O DDPM*. In *W/O VAE*, we attempt to directly feed the output of the transformer encoder layer to the domain-specific DDPMs, bypassing the VAEs. In *W/O DDPM*, we replace the DDPMs with two domain-specific VAEs. The data augmentation is done by passing the cell lines through the cell line VAE encoder and the patient VAE decoder. The pseudolabelling and downstream DRP training remains the same as GANDALF in both cases. The use of transformers and MTL appear to contribute the most to the model performance.

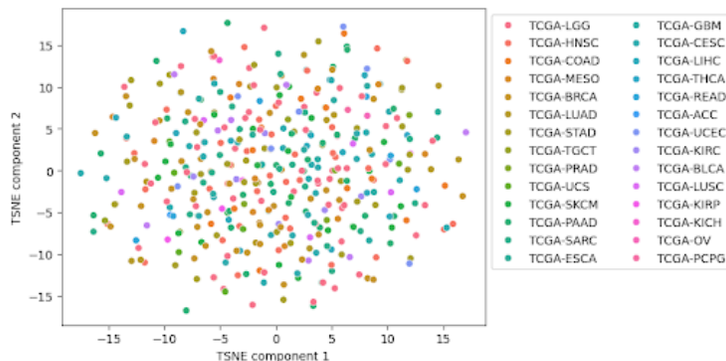


Figure 8: TSNE plots of first two components of the patient data in the representation space, color coded based on TCGA cancer types.

A.6 COMPARISON OF DISTRIBUTION

We examine the distribution of the X_{aug} with respect to the real distributions of \mathcal{X}'_c and \mathcal{X}'_p . We expect X_{aug} to be closer to the patient distribution than the original cell lines, while also retaining information from the original cell line data. Each dataset is further subjected to principal component analysis (principal components (PCs) from \mathcal{X}'_p) to obtain lower dimensional representations for easier visualization. Figure 7 (top, right) shows that original cell line data had lower variance compared to the real patient data (Figure 7, top left). However, the generated patient data (Figure 7, top middle) is closer to the real patient data in terms of the variance of data points. This indicates that the generated data captures patient-specific heterogeneity. A similar trend is seen in the density plots of the first PC in Appendix Figure 5. Quantitatively, we also examine the Kolmogorov–Smirnov (KS) test between the PCs of the 3 distributions. KS distance statistic between generated patient data and real patient data over 3 folds is 0.0694 ± 0.0071 , while the same between original cell line data and patients is 0.2524 ± 0.0022 . The PCs of the augmented data is closer to that of the real patient data, when compared to the distance between the PCs of the original cell line and patient data. This indicates that the augmented data starts resembling the patient data while retaining information from the original cell line data.

A.7 CHECKING FOR BATCH EFFECTS IN THE REPRESENTATION SPACE

Our patient data comes from three different sources - TCGA, CBIO and Moore’s. To ensure that these representations do not inadvertently capture batch effects, we perform a TSNE based visualization, where the patient latent representations are colored based on the cancer type (as coded in TCGA). For Moore’s and CBIO datasets, we identified the corresponding category in TCGA. Figure 8 shows the TSNE plot for the first two components, after embedding the patient data into the representation space. The lack of well defined boundaries across cancer types (indicated by various colors) suggest that there is no batch effect across the mutation datasets.

A.8 PERFORMANCE ACROSS CANCER TYPES

During the train-test split, we split the data based on cancer type and drug. Then we divided each group into 2:1 ratio if more than 20 samples were present per group. The train data thus contained all available cancer types. The evaluation was on a limited set of cancer types - 'TCGA-BRCA', 'TCGA-CESC', 'TCGA-HNSC', 'TCGA-STAD', 'TCGA-PAAD', 'TCGA-LGG'. Performance per cancer type from existing test splits - we calculated the metrics over the available test splits by grouping based on cancer type. Table 7 shows the results.

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Cancer type	AUROC over 3 folds	AUPRC over 3 folds
TCGA-BRCA	0.8947 ± 0.0368	0.8720 ± 0.0712
TCGA-CESC	0.3197 ± 0.1403	0.7704 ± 0.0716
TCGA-HNSC	0.7652 ± 0.137	0.9788 ± 0.0137
TCGA-STAD	0.7119 ± 0.1318	0.8253 ± 0.1222
TCGA-PAAD	0.6620 ± 0.0765	0.6239 ± 0.0516
TCGA-LGG	0.4309 ± 0.0563	0.1406 ± 0.0061

Table 7: Comparison of performance across various cancer types.