High dimensional, tabular deep learning with an auxiliary knowledge graph

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Abstract

Machine learning models exhibit strong performance on datasets with abundant labeled samples. However, for tabular datasets with extremely high d-dimensional features but limited n samples (i.e. $d \gg n$), machine learning models struggle to achieve strong performance due to the risk of overfitting. Here, our key insight is that there is often abundant, auxiliary domain information describing input features which can be structured as a heterogeneous knowledge graph (KG). We propose PLATO, a method that achieves strong performance on tabular data with $d \gg n$ by using an auxiliary KG describing input features to regularize a multilayer perceptron (MLP). In PLATO, each input feature corresponds to a node in the auxiliary KG. In the MLP's first layer, each input feature also corresponds to a weight vector. PLATO is based on the inductive bias that two input features corresponding to similar nodes in the auxiliary KG should have similar weight vectors in the MLP's first layer. PLATO captures this inductive bias by inferring the weight vector for each input feature from its corresponding node in the KG via a trainable message-passing function. Across 6 $d \gg n$ datasets, PLATO outperforms 13 state-of-the-art baselines by up to 10.19%.

1 Introduction

Machine learning models have reached state-of-the-art performance in domains with abundant labeled data like computer vision [76, 10] and natural language processing [70, 12, 52]. However, for tabular datasets in which the number d of features vastly exceeds the number n of samples, machine learning models struggle to achieve strong performance [24, 41]. Crucially, many tabular datasets from scientific domains [21, 30, 79, 17, 16, 33] have high-dimensional features but limited labeled samples due to the high time and labor costs of experiments. For these and other tabular datasets with $d \gg n$, the performance of machine learning models is currently limited.

The key challenge for machine learning models when $d \gg n$ is the risk of overfitting. Indeed, deep models can have a large number of trainable weights, yet training is limited by the comparatively small number of labeled samples. As a result, tabular deep learning approaches so far have focused on data-rich regimes with far more samples than features $(n \gg d)$ [19, 18, 59]. In the low-data regime with far more features than samples $(d \gg n)$, the dominant approaches for single tabular datasets are still statistical methods [24]. These statistical methods reduce the dimensionality of the input space [1, 41, 66, 67], select features [64, 8, 14, 46], impose regularization penalties on parameter magnitudes [45], or use ensembles of weak tree-based models [15, 7, 34, 42, 51].

Here, we present a novel problem setting and framework that enables tabular deep learning when $d \gg n$ (Figure 1). Our key insight is that there is often abundant, auxiliary domain information

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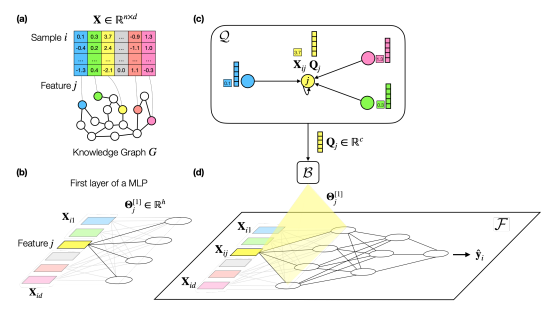


Figure 1: PLATO is a method that uses auxiliary domain information describing input features to regularize a multilayer perceptron (MLP) and achieve strong performance on tabular data with $d \gg n$. (a) In PLATO, each input feature j corresponds to a node in an auxiliary KG of domain information. (b) In the first layer of a MLP with h hidden units, each input feature j corresponds to a vector of weights $\Theta_j^{[1]} \in \mathbb{R}^h$ such that the weight vectors of all d features compose the weight matrix $\Theta^{[1]} \in \mathbb{R}^{d \times h}$. PLATO is based on the inductive bias that, if two input features j and k correspond to similar nodes in the auxiliary KG, they should have similar weight vectors $\Theta_j^{[1]}$ and $\Theta_k^{[1]}$ in the MLP. (c,d) PLATO captures this inductive bias by inferring the weight vector for each input feature j from its corresponding node in the KG. A trainable message-passing function Q creates a low-dimensional embedding $\mathbf{Q}_j \in \mathbb{R}^c$ for each input feature j. A neural network \mathcal{B} that is shared across all input features then infers the weight vector $\Theta_j^{[1]}$ corresponding to input feature j from \mathbf{Q}_j . Input features with similar embeddings produce similar weight vectors, regularizing the MLP.

describing input features which can be structured as a heterogeneous knowledge graph (KG). We propose a novel problem setting in which each input feature of a tabular dataset corresponds to a node in an auxiliary KG (Figure 1a). To represent diverse domain information describing the input features, the KG contains feature and non-feature nodes as well as multiple node and edge types. For example, consider a tabular medical dataset in which each row is a cancer patient, each column is a gene, and each value is the amount of a gene in the patient's tumor. For this tabular dataset, there exists an auxiliary KG with each gene (*i.e.* input feature) as a node. Each gene node has edges to other gene nodes (*i.e.* other feature nodes) with diverse edge types like "activates" or "inhibits." Each gene node also has edges to other nodes (*i.e.* non-feature nodes) representing the gene's function in the body like "heart rate". Finally, the function nodes (*i.e.* non-feature nodes) have edges to each other representing their anatomical relationships like "heart rate"-"part of"-"cardiac system". Note that the KG does *not* capture the relationships between *input data samples* but instead captures the relationships between *input features* and other domain information.

Within our novel problem setting, we propose PLATO, a method that enables deep learning for tabular data with $d \gg n$ by using an auxiliary KG describing input features (Figure 1). PLATO achieves strong performance by using the auxiliary KG to regularize a multilayer perceptron (MLP). In PLATO, each input feature corresponds to a node in the auxiliary KG (Figure 1a). In the first layer of the MLP, each input feature also corresponds to a weight vector such that the weight vectors of all features collectively compose the weight matrix (Figure 1b). PLATO is based on the inductive bias that two input features which correspond to similar nodes in the KG should have similar weight vectors in the first layer of the MLP. PLATO captures this inductive bias by inferring the weight vector for a feature from its corresponding node in the auxiliary KG with a trainable message-passing function (Figure 1c,d). Inferring the weights in the MLP's first layer also leads to a drastic reduction in the number of trainable weights, since most weights in a MLP are usually in the first layer when $d \gg n$.

We exhibit PLATO's performance on 6 $d \gg n$ tabular datasets with 13 state-of-the-art baselines spanning dimensionality reduction, feature selection, statistical models, graph regularization, weightinference, and tabular deep learning. Following a rigorous evaluation protocol from the tabular deep learning literature [19, 18], PLATO outperforms the prior state-of-the-art on all 6 datasets by up to 10.19%. Ablation studies demonstrate the importance of PLATO's trainable message-passing, the importance of non-feature nodes in the KG, and PLATO's robustness to missing edges in the KG. Ultimately, PLATO enables deep learning for tabular data with $d \gg n$ by using an auxiliary KG describing the input features.

2 Related Work

Tabular deep learning methods. In contrast to PLATO's setting, tabular deep learning methods have primarily been developed for settings with far more samples than features (*i.e.* $n \gg d$). Indeed, recent tabular deep learning benchmarks ignore datasets with a large number of features and a small number of samples [19, 18, 59]. In the $n \gg d$ setting, various categories of deep tabular models exist. First, decision tree models like NODE [50] make decision trees differentiable to enable gradient-based optimization [25, 36, 80]. Second, multilayer perceptrons (MLPs) apply sequential, non-linear transformations to input features [32, 31]. Third, tabular transformer architectures use an attention mechanism to select and learn interactions among features. Examples include TabNet [3], TabTransformer [29], FT-Transformer [18], TabPFN [26], SAINT [61], Non-Parametric Transformers [37], and AutoInt [62]. Finally, although PLATO focuses on single tabular datasets, transfer learning architectures can learn across multiple tabular datasets [39, 84, 73]. Ultimately, we compare PLATO to several benchmarked, state-of-the-art models for single, tabular datasets [19, 18, 59].

 $d \gg n$ methods. For PLATO's setting in which $d \gg n$, various tabular machine learning approaches exist [24]. First, dimensionality reduction techniques like PCA [1] aim to reduce the dimensionality of the input data while preserving as much of the the variance in the data as possible [41, 66, 67]. Second, feature selection approaches select a parsimonious set of features, leading to a smaller feature space. Feature selection approaches include LASSO [64] and its variants [8, 14, 46]. For feature selection with deep learning, Stochastic Gates [77] are among the best performing of many variants [4, 43]. Finally, tree-based models like XGBoost learn ensembles of weak decision trees models to make an overall prediction [15, 7, 34, 51].

Weight inference. Using one network to infer the weights of another has been studied extensively [11, 58, 6]. For example, [22] infers the weights in all layers of a sequential model (*i.e.* RNN, LSTM) by using information about the weights' structure. Diet Networks [54] infer weights by hand-crafting prior information about the input features or using random projections. By contrast, PLATO infers the weights in a MLP from prior information describing the input features in an auxiliary KG. PLATO's weight inference uniquely captures the inductive bias that two input features corresponding to similar nodes in a KG should have similar corresponding weight vectors in the first layer of a MLP (Figure 1).

Graph regularization. Graph regularization approaches regularize the weights of a linear model based on a simple graph between input features. The graph is typically constructed from the tabular data based on covariance relationships. Approaches then add a regularization penalty to the loss function which forces the weights of the linear model to vary smoothly over the corresponding feature nodes in the graph. State-of-the-art methods include GraphNet [20] and Network-Constrained LASSO [40] which are based on a Laplacian regularization [60, 2] as well as Network LASSO [23] which generalizes the Group LASSO [83] to a network setting. PLATO differs from graph regularization approaches in two key ways. First, PLATO's KG includes both feature and non-feature nodes and multiple edge types, thereby modeling diverse, prior domain information that is missing in graph regularization approaches. Second, PLATO infers the weights of a deep non-linear model (*i.e.* a MLP) rather than adding a regularization penalty to a loss, representing a distinct regularization mechanism.

Knowledge graph methods. Existing KG approaches are designed for tasks directly on the graph like link prediction or node classification [71, 65, 72, 78, 13]. By contrast, PLATO does not make any predictions on the KG. Instead, PLATO makes predictions on a separate, tabular dataset by using the KG as a prior. Graph classification methods also do not apply (Appendix B).

3 Plato

PLATO is a machine learning method for tabular datasets with $d \gg n$ and an auxiliary knowledge graph (KG) with input features as nodes (Section 3.1). PLATO's key insight is that there often exists abundant domain information describing input features which can be structured as an auxiliary KG *G* (Figure 1a). PLATO uses the auxiliary KG to regularize a multilayer perceptron (MLP) and achieve strong performance on tabular data when $d \gg n$.

3.1 Problem setting

Consider a tabular dataset $\mathbf{X} \in \mathbb{R}^{n \times d}$ with labels $\mathbf{y} \in \mathbb{R}^n$ and far more d features than n samples such that $d \gg n$. The goal is to train a model \mathcal{F} to predict labels $\hat{\mathbf{y}}$ from the input \mathbf{X} . PLATO assumes the existence of an auxiliary knowledge graph G = (V, E) with |V| nodes and |E| edges such that each input feature j corresponds to a node in G. Formally, $\forall j \in \{1, \ldots, d\}$, $\exists v \in V$ s.t. $j \mapsto v$, as shown in Figure 1a. G also contains additional nodes which represent broader knowledge describing the domain. The edges in G are (head node, relation type, tail node) triplets.

3.2 PLATO's inductive bias

In PLATO, each input feature j corresponds to a node in the auxiliary KG (Figure 1a). In the first layer of a MLP with h hidden units, each input feature j also corresponds to a weight vector $\Theta_j^{[1]} \in \mathbb{R}^h$ such that the weight vectors of all features collectively compose the weight matrix $\Theta^{[1]} \in \mathbb{R}^{d \times h}$ (Figure 1b). PLATO is based on the inductive bias that two input features j and k which correspond to similar nodes in the KG should have similar weight vectors $\Theta_j^{[1]}$ and $\Theta_k^{[1]}$ in the first layer of the MLP. PLATO captures this inductive bias by inferring the weight vector for a feature from its corresponding node in the auxiliary KG with a trainable message-passing function (Figure 1c,d).

3.3 PLATO overview

PLATO has four key steps. First, PLATO uses a self-supervised objective on the auxiliary KG to pretrain an embedding for each input feature (Section 3.4). Second, PLATO updates each feature embedding with a trainable message-passing function that is trained on the supervised loss objective for the tabular data (Section 3.5, Figure 1c). Third, PLATO infers the weights in the first layer of the MLP from the feature embeddings with a small neural network that is shared across input features (Section 3.6, Figure 1d). Finally, the MLP predicts the label for the input sample.

3.4 Pretraining feature embeddings with self-supervision on the knowledge graph

First, PLATO learns general prior information about each input feature j from the auxiliary KG G. PLATO represents the general prior information about each input feature j as a low-dimensional embedding $\mathbf{M}_j \in \mathbb{R}^c$. Since each input feature j corresponds to a node in G, PLATO can learn \mathbf{M}_j by learning an embedding for the corresponding feature node in G. Any self-supervised node embedding method on G can be used within PLATO's framework.

Formal notation. Formally, PLATO uses self-supervision on G to pretrain an embedding for each input feature according to

$$\mathbf{M} = \mathcal{H}(G). \tag{1}$$

 $\mathbf{M} \in \mathbb{R}^{d \times c}$ is the matrix of all feature embeddings. \mathcal{H} is a self-supervised node embedding method. We refer to Eq. (1) as pretraining since only the auxiliary KG G is used but the tabular data \mathbf{X} , \mathbf{y} is ignored. After pretraining, the feature embeddings \mathbf{M} are fixed.

For \mathcal{H} , we choose ComplEx as it is a prominent and highly scalable KG node embedding method [65]. ComplEx uses a self-supervised objective which learns an embedding for each node in G by classifying whether a proposed edge exists in G. ComplEx's proposed edges include both feature nodes and other nodes in G, thereby integrating prior information about the input features and the broader domain. We also test KG embedding methods DistMult [78] and TransE [72] in Appendix C.

3.5 Updating feature embeddings with a message-passing function trained on tabular data

PLATO next updates each feature embedding with a trainable message-passing function that is trained on the supervised loss for the tabular data (Figure 1c). During message-passing, PLATO updates the embedding of each input feature to be a weighted aggregation of it's neighbors' embeddings.

Formal notation. Formally, PLATO uses a message-passing function Q on the KG to update each pre-trained feature embedding $\mathbf{M}_j \in \mathbb{R}^c$ to feature embedding $\mathbf{Q}_j \in \mathbb{R}^c$ according to

$$\mathbf{Q} = \mathcal{Q}(\mathbf{M}, G, \mathbf{X}_i; \mathbf{\Pi}). \tag{2}$$

As input, the message-passing function considers the pre-trained feature embeddings \mathbf{M} , the knowledge graph G, and the sample value \mathbf{X}_i . Q uses an attention mechanism which considers the sample value \mathbf{X}_i . The only trainable weights in Q are in the attention mechanism and are $\mathbf{\Pi}$.

The message passing network Q. Let $\mathbf{Q}_{j}^{[r]}$ be the embedding of input feature j after round $r \in \{1, ..., R\}$ of message passing. For each input feature j, Q first initializes the updated feature embedding to the pretrained feature embedding.

$$\mathbf{Q}_j^{[0]} = \mathbf{M}_j. \tag{2a}$$

Q then conducts R rounds of message passing. In each round of message passing, the feature embedding $\mathbf{Q}_{j}^{[r]}$ is updated from the feature embedding of each neighbor k in the prior round $\mathbf{Q}_{k}^{[r-1]}$ and its own feature embedding in the prior round $\mathbf{Q}_{j}^{[r-1]}$. The "message" being passed is the embedding of each feature from the prior round.

$$\mathbf{Q}_{j}^{[r]} = \sigma \left[\underbrace{\beta(\sum_{k \in N_{j}} \alpha_{ijk} \mathbf{Q}_{k}^{[r-1]})}_{k \in N_{j}} + \underbrace{(1-\beta) \mathbf{Q}_{j}^{[r-1]}}_{\text{Weighted message from self}} \right].$$
(2b)

 σ is an optional nonlinearity. N_j are the neighbors of feature node j in G.

During message-passing, Q uses two scalar values $\beta \in \mathbb{R}$ and $\alpha_{ijk} \in \mathbb{R}$ to control the weights of messages. First, Q uses hyperparameter $\beta \in \mathbb{R}$ to control the weight of the messages aggregated from the feature node's neighbors vs. from the feature node itself. Second, Q calculates an attention coefficient $\alpha_{ijk} \in \mathbb{R}$ to allow distinct nodes in the same neighborhood to have distinct weights. The coefficient α_{ijk} specifies the weight of the message between feature j and neighbor k for sample i.

After R rounds of message-passing, the updated feature embeddings Q_i are set.

$$\mathbf{Q}_j = \mathbf{Q}_j^{[R]}.\tag{2c}$$

The attention coefficient. PLATO's attention coefficient α_{ijk} is inspired by [68] in which node attributes are used to calculate the weight of a message between neighboring nodes. For a sample *i* in PLATO, the node attributes for features *j* and *k* are their sample values $\mathbf{X}_{ij} \in \mathbb{R}$ and $\mathbf{X}_{ik} \in \mathbb{R}$. PLATO thus uses the sample values \mathbf{X}_{ij} and \mathbf{X}_{ik} to calculate the attention coefficient. The attention coefficient e_{ijk} indicates the importance of node *j* to node *k* for sample *i*.

$$e_{ijk} = \mathcal{A}(\mathbf{X}_{ij}, \mathbf{X}_{ik}; \mathbf{\Pi}).$$
(2d)

 \mathcal{A} is a shallow neural network parameterized by Π that is shared across samples and features. The number of trainable weights in Π is small since the input of \mathcal{A} is \mathbb{R}^2 and the output of \mathcal{A} is a scalar \mathbb{R} .

To make the attention coefficients comparable across different nodes, PLATO normalizes the attention coefficients with a softmax function across the neighbors N_j of node j.

$$\alpha_{ijk} = \operatorname{softmax}_k(e_{ijk}) = \frac{\exp(e_{ijk})}{\sum_{t \in N_j} \exp(e_{ijt})}.$$
(2e)

Algorithm 1: The PLATO Algorithm.

Input: A data sample $\mathbf{X}_i \in \mathbb{R}^d$, a knowledge graph G containing each input feature in \mathbf{X} as a node, a matrix of input feature embeddings $\mathbf{M} \in \mathbb{R}^{d \times c}$ pre-trained over G. Output: A predicted label $\hat{\mathbf{y}}_i \in \mathbb{R}$.

- 1 Use a trainable message-passing function Q to update the pre-trained feature embeddings: $\mathbf{Q} = Q(\mathbf{M}, G, \mathbf{X}_i; \mathbf{\Pi}), \mathbf{Q}_i \in \mathbb{R}^c, \mathbf{Q} \in \mathbb{R}^{d \times c}$
- 2 Infer the weight vector in the first layer of a MLP that corresponds to an input feature j with a neural network \mathcal{B} :

 $\hat{\mathbf{\Theta}}_{j}^{[1]} = \mathcal{B}(\mathbf{Q}_{j}|\mathbf{X}_{i};\mathbf{\Phi}), \, \hat{\mathbf{\Theta}}_{j}^{[1]} \in \mathbb{R}^{h}$

- 4 Concatenate the first layer inferred weights with the trainable weights in the rest of the MLP layers:

$$\hat{\mathbf{\Theta}} = \{ \hat{\mathbf{\Theta}}^{[1]} | \mathbf{X}_i \} \cup \{ \mathbf{\Theta}^{[2]}, \dots, \mathbf{\Theta}^{[L]} \}.$$

5 Predict the label with a MLP \mathcal{F} that is parameterized by $\hat{\Theta}$

 $\hat{\mathbf{y}}_i = \mathcal{F}(\mathbf{X}_i; \hat{\mathbf{\Theta}} | \mathbf{X}_i), \hat{\mathbf{y}}_i \in \mathbb{R}$ Trainable weights: $\mathbf{\Pi}, \mathbf{\Phi}, \mathbf{\Theta}^{[2]}, \dots, \mathbf{\Theta}^{[L]}$.

3.6 Inferring the first layer of weights in \mathcal{F} from the updated feature embeddings

Finally, PLATO infers the weights in the first layer of a MLP \mathcal{F} from the updated feature embeddings (Figure 1d). In the first layer of a MLP with h hidden units, each input feature j corresponds to a weight vector $\Theta_j^{[1]} \in \mathbb{R}^h$ (Figure 1b). The weight matrix in the first layer of the MLP, $\Theta^{[1]} \in \mathbb{R}^{d \times h}$, is simply the concatenation of d weight vectors, one corresponding to each input feature. For each input feature j, PLATO infers the weight vector $\hat{\Theta}_j^{[1]} \in \mathbb{R}^h$ from the feature embedding $\mathbf{Q}_j \in \mathbb{R}^c$ by using a shallow neural network shared across input features. Input features with similar feature embeddings will produce similar weight vectors. Thus, PLATO captures the inductive bias that input features corresponding to similar nodes in the KG should have similar corresponding weight vectors in the MLP's first layer.

Formal notation. PLATO infers the weight vector associated with each input feature j in the first layer of \mathcal{F} with

$$\hat{\boldsymbol{\Theta}}_{j}^{[1]} = \mathcal{B}(\mathbf{Q}_{j} | \mathbf{X}_{i}; \boldsymbol{\Phi}).$$
(3)

 \mathcal{B} is a shallow neural network with trainable weights Φ . \mathbf{Q}_j is the updated feature embedding of j which is conditioned on the specific input sample \mathbf{X}_i since the input sample is used as an input in its calculation (Section 3.5, Equation 2). Φ are the weights of \mathcal{B} . \mathcal{B} and its weights Φ are shared across each feature $j \in \{1, \ldots, d\}$.

PLATO drastically reduces the number of trainable weights compared to a standard MLP. The sharing of \mathcal{B} and Φ across all input features drastically reduces the number of trainable weights compared to a standard MLP. For a high-dimensional tabular dataset (*i.e.* $d \gg n$), a standard MLP \mathcal{T} with h hidden units has a large number of trainable weights in the first layer since $\Theta^{[1]} \in \mathbb{R}^{d \times h}$. A standard MLP \mathcal{T} must learn all dh of these trainable weights by backpropagation. By contrast, \mathcal{B} uses a shared set of trainable weights Φ to infer $\hat{\Theta}_j$ from \mathbf{Q}_j for every $j \in \{1, \ldots, d\}$. The number of trainable weights in Φ is small compared to dh since \mathcal{B} need only transform every $\mathbf{Q}_j \in \mathbb{R}^c$ to $\hat{\Theta}^{[1]} \in \mathbb{R}^h$. Thus, $|\Phi| = ch$ (assuming \mathcal{B} is a single layer neural network). c, the dimensionality of the feature embedding, is much less than d the number of input features. As a result, $|\Phi| = ch \ll dh$ and PLATO drastically reduces the number of trainable weights in the first layer of a MLP.

3.7 The PLATO algorithm

PLATO is outlined in Algorithm 1.

Dataset		MNSCLC	СМ	PDAC	BRCA	CRC	СН
# of fe	eatures d	15,390	13,183	12,932	12,693	18,206	19,902
# of sa	amples n	295	286	321	476	562	924
0	l/n	52.2	46.1	40.3	28.2	22.6	19.7
Classic Stat ML Ridge		0.153±0.000	$0.390{\scriptstyle\pm0.000}$	$0.344{\scriptstyle\pm0.000}$	$\underline{0.538{\scriptstyle\pm0.000}}$	$0.376{\scriptstyle \pm 0.000}$	$0.546{\scriptstyle\pm0.000}$
Dim. Reduct.	PCA	0.156±0.113	0.070 ± 0.000	$0.232{\scriptstyle\pm0.121}$	$0.452{\scriptstyle\pm0.000}$	$0.193{\scriptstyle \pm 0.163}$	$0.237{\scriptstyle\pm0.232}$
Feat. Select.	LASSO STG	$ \begin{vmatrix} 0.168 \pm 0.000 \\ 0.132 \pm 0.130 \end{vmatrix} $	$\frac{0.431{\pm}0.000}{0.366{\pm}0.043}$	$\begin{array}{c} 0.346 {\pm} 0.000 \\ 0.258 {\pm} 0.055 \end{array}$	$\begin{array}{c} 0.470 {\pm} 0.000 \\ 0.485 {\pm} 0.037 \end{array}$	$\frac{0.400{\pm}0.000}{0.301{\pm}0.010}$	$\begin{array}{c} 0.547 {\pm} 0.000 \\ 0.262 {\pm} 0.076 \end{array}$
Decision Tree	XGBoost	-0.02±0.000	$0.225{\scriptstyle\pm0.000}$	$\underline{0.363{\scriptstyle\pm0.000}}$	$0.347{\scriptstyle\pm0.000}$	$0.354{\scriptstyle\pm0.000}$	$\underline{0.728{\scriptstyle\pm0.000}}$
Graph Reg.	GraphNet NC LASSO Network LASSO	$ \begin{vmatrix} 0.169 \pm 0.030 \\ 0.210 \pm 0.014 \\ 0.212 \pm 0.046 \end{vmatrix} $	$\begin{array}{c} 0.277 {\pm} 0.099 \\ 0.339 {\pm} 0.044 \\ 0.243 {\pm} 0.058 \end{array}$	$\begin{array}{c} 0.249 {\pm} 0.018 \\ 0.327 {\pm} 0.053 \\ 0.136 {\pm} 0.027 \end{array}$	$\begin{array}{c} 0.350 {\pm} 0.069 \\ 0.458 {\pm} 0.083 \\ 0.348 {\pm} 0.033 \end{array}$	$\begin{array}{c} 0.125{\scriptstyle\pm0.061}\\ 0.220{\scriptstyle\pm0.030}\\ 0.171{\scriptstyle\pm0.040}\end{array}$	$\begin{array}{c} 0.646 {\pm} 0.051 \\ 0.415 {\pm} 0.083 \\ 0.212 {\pm} 0.091 \end{array}$
Param. Infer.	Diet	-0.04±0.205	$0.054{\scriptstyle\pm0.149}$	$0.309{\scriptstyle\pm0.096}$	$0.213{\scriptstyle \pm 0.036}$	$0.087{\scriptstyle\pm0.112}$	$0.148{\scriptstyle\pm0.008}$
Tabular DL	MLP NODE TabTransformer TabNet	$ \begin{vmatrix} 0.128 \pm 0.126 \\ 0.003 \pm 0.000 \\ \underline{0.265 \pm 0.000} \\ 0.085 \pm 0.028 \end{vmatrix} $	$\begin{array}{c} 0.322{\pm}0.043\\ 0.150{\pm}0.000\\ 0.072{\pm}0.000\\ 0.010{\pm}0.068\end{array}$	$\begin{array}{c} 0.289 {\pm} 0.047 \\ 0.190 {\pm} 0.000 \\ 0.029 {\pm} 0.000 \\ 0.088 {\pm} 0.037 \end{array}$	$\begin{array}{c} 0.240 {\pm} 0.067 \\ 0.512 {\pm} 0.000 \\ 0.202 {\pm} 0.000 \\ 0.055 {\pm} 0.037 \end{array}$	$\begin{array}{c} 0.355 {\pm} 0.022 \\ 0.344 {\pm} 0.000 \\ 0.238 {\pm} 0.000 \\ 0.018 {\pm} 0.016 \end{array}$	$\begin{array}{c} 0.044 {\pm} 0.039 \\ 0.181 {\pm} 0.000 \\ 0.020 {\pm} 0.000 \\ 0.039 {\pm} 0.026 \end{array}$
Ours	PLATO	0.272±0.130	$0.435{\scriptstyle \pm 0.022}$	$0.400{\scriptstyle\pm0.021}$	$0.583{\scriptstyle \pm 0.019}$	$0.401{\scriptstyle \pm 0.019}$	$0.770{\scriptstyle\pm0.003}$

Table 1: **PLATO outperforms statistical and deep baselines when** $d \gg n$. For every dataset, the best overall model is in **bold** and the second best model is <u>underlined</u>.

Table 2: PLATO's performance depends on updat-
Table 3: PLATO's performance depends on
ing feature embeddings with a trainable message- both feature nodes in G and other nodes
passing (MP) function.Comparison of the sector of the s

Weight Infer. $\mathcal B$ Input	Feature Info.	Trainable MP	PearsonR	Auxiliary KG	Feature Info.	Broader Info.	PearsonR
Updated feat. embed. \mathbf{Q}	1	1	0.583±0.019	Full KG	1	 Image: A set of the set of the	0.583±0.019
General feat. embed ${f M}$	~	×	0.522 ± 0.030	Feature-only KG	 Image: A second s	×	$0.539{\scriptstyle \pm 0.038}$
None	×	×	0.240±0.067	No KG	×	×	$0.240{\scriptstyle \pm 0.067}$

4 Experiments

We evaluate PLATO against 13 baselines on 10 tabular datasets (6 with $d \gg n$, 4 with $d \sim n$).

Datasets. We use 6 tabular $d \gg n$ datasets, $4 d \sim n$ datasets [16, 17, 30, 79], and a KG from prior studies [44, 35, 38, 56, 63, 74, 75] (Appendix G, H). The KG contains 108,447 nodes, 3,066,156 edges, and 99 relation types. All datasets include features which map to a subset of knowledge graph nodes. Code, data, and the KG are available at https://github.com/snap-stanford/plato.

Baselines. We compare PLATO to 13 state-of-the art statistical and deep baselines. We consider regularization with Ridge Regression [45], dimensionality reduction with PCA [1] followed by linear regression, feature selection with LASSO [64], deep feature selection with Stochastic Gates [77], and gradient boosted decision trees with XGBoost [7]. We consider tabular deep learning with a standard MLP, self-attention-based methods with TabTransformer [29] and TabNet [3], differentiable decision trees with NODE [50], and weight inference with Diet Networks [54]. We also attempted FT-Transformer [18], but it experienced out of memory issues on all datasets due to the large number of features. Finally, we consider graph regularization methods which also have access to the knowledge graph including GraphNet [20], NC LASSO [40], and Network LASSO [23] (Appendix E).

Fair Comparison of PLATO with Baselines. To ensure a fair comparison with baselines, we follow evaluation protocols in recent tabular benchmarks [19, 18]. We conduct a random search with 500 configurations of every model (including PLATO) on every dataset across a broad range of hyperparameters (Appendix A). We split data with a 60/20/20 training, validation, test split. All results are computed across 3 data splits and 3 runs of each model in each data split. We report the mean and standard deviation of the Pearson correlation (PearsonR) between y and \hat{y} across runs and splits on the test set. Each model is run on a GeForce RTX 2080 TI GPU.

Fraction of edges in KG	PearsonR	Model	Description	PearsonR
90%	$\begin{array}{c} 0.583 \pm 0.019 \\ 0.570 \pm 0.017 \\ 0.537 \pm 0.044 \end{array}$	Plato Plato-LR	MLP with first layer weights inferred Linear regression	$\begin{array}{c} 0.583 \pm 0.019 \\ 0.550 \pm 0.020 \end{array}$
	0.412 ± 0.011		with weights inferred	

Table 4: PLATO's performance with an incom-
plete knowledge graph.Table 5: PLATO's MLP layers $2, \ldots, L$ with
trainable weights are useful for performance.

Table 6: **PLATO's performance is competitive with baselines when** $d \sim n$ **.** For every dataset, the best overall model is in **bold** and the second best model is <u>underlined</u>.

Da	taset	ME	BC	SCLC	NSCLC
# of features d		19,902	18,261	18,437	18,308
# of sa	mples n	10,064	10,101	10,712	16,730
d	/n	2.0	1.8	1.7	1.1
Classic Stat ML	Ridge	0.566±0.008	$0.483{\scriptstyle\pm0.008}$	$0.604{\scriptstyle\pm0.057}$	$0.679{\scriptstyle \pm 0.008}$
Dim. Reduct.	n. Reduct. PCA		$0.233{\scriptstyle \pm 0.294}$	$0.284{\scriptstyle\pm0.274}$	$0.645{\scriptstyle\pm0.000}$
Feat. Select.	LASSO STG	$\begin{array}{ c c c c c c c c c c c c c c c c c c c$	$\begin{array}{c} 0.633 {\pm} 0.000 \\ 0.643 {\pm} 0.000 \end{array}$	$\begin{array}{c} 0.669 {\pm} 0.000 \\ 0.668 {\pm} 0.000 \end{array}$	$\begin{array}{c} 0.637 {\pm} 0.000 \\ 0.646 {\pm} 0.000 \end{array}$
Decision Tree XGBoost		0.875±0.000	$\underline{0.826{\scriptstyle\pm0.000}}$	$\underline{0.878{\scriptstyle\pm0.000}}$	$0.843{\scriptstyle \pm 0.000}$
Graph Reg.	GraphNet NC LASSO Network LASSO	$ \begin{vmatrix} 0.675 \pm 0.047 \\ 0.733 \pm 0.016 \\ 0.401 \pm 0.034 \end{vmatrix} $	$\begin{array}{c} 0.723 {\pm} 0.026 \\ 0.730 {\pm} 0.027 \\ 0.451 {\pm} 0.022 \end{array}$	$\begin{array}{c} 0.742 {\pm} 0.039 \\ 0.793 {\pm} 0.009 \\ 0.417 {\pm} 0.074 \end{array}$	$\begin{array}{c} 0.627 {\pm} 0.042 \\ 0.746 {\pm} 0.023 \\ 0.465 {\pm} 0.034 \end{array}$
Param. Infer.	Diet	0.105±0.000	$0.037{\scriptstyle\pm0.000}$	$\textbf{-0.050}{\scriptstyle \pm 0.000}$	0.002 ± 0.000
Tabular DL	MLP NODE TabTransformer TabNet	$ \begin{vmatrix} 0.487 \pm 0.131 \\ 0.870 \pm 0.000 \\ 0.305 \pm 0.028 \\ 0.667 \pm 0.002 \end{vmatrix} $	$\begin{array}{c} 0.508 \pm 0.061 \\ 0.420 \pm 0.169 \\ 0.010 \pm 0.000 \\ 0.624 \pm 0.001 \end{array}$	$\begin{array}{c} 0.537 {\pm} 0.061 \\ 0.801 {\pm} 0.102 \\ 0.288 {\pm} 0.203 \\ 0.657 {\pm} 0.004 \end{array}$	$\begin{array}{c} 0.573 {\pm} 0.005 \\ 0.487 {\pm} 0.197 \\ 0.503 {\pm} 0.187 \\ 0.647 {\pm} 0.000 \end{array}$
Ours	Plato	0.875±0.004	$0.844{\scriptstyle\pm0.003}$	$0.883{\scriptstyle \pm 0.002}$	$\underline{0.839{\scriptstyle\pm0.000}}$

4.1 Results

PLATO outperforms statistical and deep baselines when $d \gg n$. **PLATO outperforms all baselines** across all 6 datasets with $d \gg n$ (Table 1). PLATO achieves the largest improvement on the PDAC dataset, improving by 10.19% vs. XGBoost, the best baseline for PDAC (0.400 vs. 0.363). While PLATO achieves the strongest performance across all 6 datasets, the best performing baseline varies across datasets. Ridge Regression is the strongest baseline for BRCA, LASSO for CM and CRC, XGBoost for PDAC and CH, and TabTransformer for MNSCLC. The remaining baselines are not the strongest baseline for any dataset. We also find that the performance of a specific baseline depends largely on the dataset. TabTransformer, for example, is the best baseline for the MNSCLC dataset but the worst baseline for the CH dataset. The rank order of all models on all datasets is Appendix D.

PLATO's performance depends on updating feature embeddings with a trainable messagepassing function. PLATO infers the weights $\hat{\Theta}^{[1]}$ in the first layer of a MLP \mathcal{F} by using feature embeddings which contain prior information about the input features. PLATO first pretrains general feature embeddings $\mathbf{M} \in \mathbb{R}^{d \times c}$. PLATO then updates the feature embeddings to $\mathbf{Q} \in \mathbb{R}^{d \times c}$ with a trainable message-passing function. We test whether updating the feature embeddings based on the trainable message-passing function is necessary by evaluating PLATO's performance on the BRCA dataset in three configurations (Table 2). The default configuration uses the updated feature embeddings \mathbf{Q} generated by the message-passing function to infer $\hat{\Theta}^{[1]}$ according to $\hat{\Theta}_{j}^{[1]} = \mathcal{B}(\mathbf{M}_{j})$. The third configuration does not use feature embeddings and thus ablates to a standard MLP. Using general feature embeddings \mathbf{M} improves over not using feature embeddings at all (0.522 vs. 0.240). Using feature embeddings \mathbf{Q} that are generated by the trainable message-passing function further improves performance (0.583 vs. 0.522). Thus, updating the feature embeddings to \mathbf{Q} based on the trainable message-passing function is key to PLATO's performance. **PLATO's performance depends on both feature nodes and broader knowledge nodes in the auxiliary KG.** PLATO relies on an auxiliary KG *G* which contains information describing input features and the broader domain. Information describing input features is represented as feature nodes while information describing the broader domain is represented as other nodes in *G* (Methods 3.1). To test the relative importance of the feature information in *G* vs. the broader domain information, we measured the performance of PLATO on the BRCA dataset in two KG configurations: PLATO with the full KG (*i.e.* both the feature nodes and the broader domain nodes) and PLATO with a "feature-only KG" (*i.e.* an induced subgraph on only the feature nodes) (Table 3). We also compare to a "No KG" configuration in which PLATO does not have access to the KG. Without auxiliary information describing the input features or the broader domain, PLATO is ablated to a standard MLP.

We find that both the feature nodes and the broader knowledge nodes are important for PLATO's performance. Using the "feature-only KG" configuration of PLATO improves performance vs the "no KG" configuration (0.539 vs 0.240). Using the "full KG" configuration further improves performance vs the "feature-only KG" configuration (0.583 vs 0.539). PLATO's performance thus relies on both the feature information and the broader domain information in the KG.

PLATO's performance with an incomplete knowledge graph. All KGs are incomplete since there is undiscovered knowledge. PLATO thus uses low-dimensional embeddings from KG embedding approaches [72, 78, 65] which are designed to account for missing information, enabling predictive performance even with missing edges. We conduct an ablation study to assess PLATO's robustness to missing edges in the KG. We randomly remove edges from the KG and measure PLATO's performance on the BRCA dataset. We observe that with only 50% of the KG's edges, PLATO still has 71% of the performance as PLATO with 100% of the KG's edges (0.412 vs. 0.583) (Table 4).

The importance of MLP layers $2, \ldots, L$, the layers with trainable weights, for PLATO. PLATO is a MLP in which the weights in the first layer are inferred from the knowledge graph (KG) but the weights in the remaining layers $2, \ldots, L$ are trained normally. We conduct an ablation study to determine whether MLP layers $2, \ldots, L$ are necessary for PLATO's performance or whether the first layer of inferred weights are sufficient. Note that a single layer of inferred weights in PLATO is equivalent to a linear regression in which the weights are inferred from the KG. We thus compare PLATO to PLATO-LR, a linear regression in which the weights are inferred from the KG (Table 5). PLATO 's standard configuration outperforms PLATO-LR on the BRCA dataset (0.583 vs. 0.550). Therefore, layers $2, \ldots, L$ of the MLP are important for PLATO's performance.

For datasets with $d \sim n$, PLATO is competitive with baselines. Finally, we test PLATO's performance for datasets with $d \sim n$. We test 4 datasets with $d \sim n$ ranging from $\frac{d}{n} = 1.1$ to 2.0 (Table 6). We find that on 4 datasets with $d \sim n$, PLATO is competitive with the best baseline, XGBoost, but does not improve performance substantially. PLATO's stronger performance for datasets with $d \sim n$ is justified. PLATO's key idea is to include auxiliary information describing the input features. Auxiliary information is likely to help performance the most in settings with the least labeled data (*i.e.* $d \gg n$). When $d \sim n$, auxiliary information is less helpful since the tabular dataset may already have enough information to train a strong predictive model. We further find that XGBoost is the strongest baseline for all datasets with $d \sim n$, in contrast to XGBoost's varied performance on the datasets with $d \gg n$ (Table 1).

5 Discussion

PLATO achieves strong performance on tabular data when $d \gg n$ by using an auxiliary KG describing input features to regularize a multilayer perceptron (MLP). Across 6 datasets, PLATO outperforms 13 state-of-the-art baselines by up to 10.19%. Ablations demonstrate the importance of PLATO's trainable message-passing function, of including nodes in the KG that don't represent input features but instead represent domain information, and of the layers in the MLP whose weights are trained directly rather than inferred. We also test PLATO's robustness to missing information in the KG. PLATO has several limitations. First, PLATO matches but does not improve the performance of baselines for high-dimensional datasets with more samples (*i.e.* $d \sim n$). Second, PLATO depends on the existence of an auxiliary KG of domain information. Overall, PLATO enables tabular deep learning when $d \gg n$ by using an auxiliary KG of domain information describing input features.

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A Evaluation protocol and hyperparameter ranges

To ensure a fair comparison with baselines, we follow evaluation protocols outlined in tabular benchmarks [19, 18]. We conduct a random search with 500 configurations of every model (including PLATO) on every dataset across a broad range of hyperparameters. We base the hyperparameter ranges on the ranges used in prior tabular learning benchmarks [19, 18] and the ranges mentioned in the original papers of the methods. Hyperparameter ranges for PLATO are given in Table 7. Hyperparameter ranges for baseline methods are given in Table 8.

Module in PLATO	Hyperparameter	Range
General	Learning rate Batch size L2	LogUniform(1e-4, 5e-3) [16, 32, 64] 0, LogUniform(1e-5, 1e-2)
$\operatorname{KG} \mathcal{H}$	Embedding dimension c Embedding model	200 ComplEx
Message Passing (MP) Q	# Rounds R β Hidden dimension in \mathcal{A}	2 LogUniform(1e-4, 1e-1) UniformInt(16, 512)
Weight Inference \mathcal{B}	# Layers Hidden dimension	UniformInt(2, 6) UniformInt(16, 512)
Layers $2, \ldots, L$ in MLP \mathcal{F}	# Layers L Hidden dimension	UniformInt(2, 6) UniformInt(16, 512)

Table 7: Hyperparameter ranges used for PLATO.

Model	Hyperparameter	Range
LASSO	L1	LogUniform(1E-4, 10)
Ridge	L2	LogUniform(1E-4, 10)
XGBoost	n-estimators Max depth Min weight Subsample Learning rate Col sample by level Col sample by tree Gamma Lambda Alpha Booster Early-stopping-rounds Iterations	UniformInt(1,2000) UniformInt(3, 10) LogUniform(1E-8,1E5) Uniform(0.5, 1) LogUniform(1E-5,1) Uniform(0.5, 1) Uniform(0.5, 1) 0, LogUniform(1E-8, 1E2) 0, LogUniform(1E-8, 1E2) "gbtree" 50 100
PCA	Number of PCA Components	UniformInt(2,1000)
STG	Hidden dimension Number of layers Activation Learning rate Sigma Lambda	UniformInt(10, 500) UniformInt(1, 5) [Tanh, Relu, Sigmoid] LogUniform(1e-4, 1e-1) Uniform(0.001, 2) LogUniform(1e-3, 10)
MLP	Number of layers Hidden dimension Dropout Learning rate L2	UniformInt(1, 8) UniformInt(1, 512) 0, Uniform([0,0.5]) LogUniform(1e-5, 1e-2) 0, LogUniform(1e-6, 1e-3)
TabNet	Decision Steps Layer size Relaxation factor Sparsity loss weight Decay rate Decay steps Learning rate Iterations	UniformInt(3, 10) 2, 4, 8, 16, 32, 64 Uniform[1, 2] LogUniform[1e-6, 1e-1] Uniform[0.4, 0.95] 100, 500, 2000 Uniform(1e-3, 1e-2) 100
TabTransformer	Embedding dimension Number of heads Number of attention blocks Attention dropout rate Add norm dropout Transformation activation L2 Learning rate FF dropout FF hidden multiplier Out FF activation Out FF dropout	4, 8, 16, 32, 64, 128 UniformInt(1, 10) UniformInt(1, 12) Uniform(0, 0.5) Uniform(0, 0.5) [Tanh, Relu, LeakyReLU] LogUniform(1e-6, 1e-1) LogUniform(1e-6, 1e-3) Uniform(0, 0.5) 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 [Tanh, Relu, LeakyReLU] Uniform(0, 0.5)
NODE	Learning rate Number of layers Number of trees Depth	LogUniform(1e-5, 1) UniformInt(1, 10) UniformInt(2, 2048) UniformInt(1, 10)

Diet Network

	Number of layers Hidden dimension Dropout Learning rate L2	UniformInt(1, 8) UniformInt(1, 512) 0, Uniform([0,0.5]) LogUniform(1e-5, 1e-2) 0, LogUniform(1e-6, 1e-3)				
GraphNet	Hidden dimension Learning rate λ L1 coefficient	UniformInt(1, 512) LogUniform(1e-5, 1e-2) 0, LogUniform(1e-5, 1e2) 0, LogUniform(1e-5, 1e2)				
NC Lasso	Hidden dimension Learning rate λ L1 coefficient	UniformInt(1, 512) LogUniform(1e-5, 1e-2) 0, LogUniform(1e-5, 1e2) 0, LogUniform(1e-5, 1e2)				
Network Lasso	Hidden dimension Learning rate λ L1 coefficient	UniformInt(1, 512) LogUniform(1e-5, 1e-2) 0, LogUniform(1e-5, 1e2) 0, LogUniform(1e-5, 1e2)				
Table 8: Hyperparameter range for all baselines.						

B Graph classification approaches

Graph classification models are not appropriate for PLATO's setting. In graph classification models, every input sample is a graph with node attributes, and a model must make a prediction for that graph. The PLATO problem setting breaks key assumptions made by typical graph classification models. First, graph classification models assume that different samples correspond to different graphs [81, 28, 27]. However, in PLATO every sample corresponds to the exact same graph. There is a single background knowledge graph for all samples, and every sample has input features that correspond to the exact same nodes within the knowledge graph. Second, graph classification approaches typically assume that every node in an input graph has a node attribute [81, 28, 27]. However, in PLATO only a small subset of the nodes in the knowledge graph have measured feature values. Finally, graph classification approaches typically assume small graphs: the largest graph classification task in the Open Graph Benchmark has only 244 nodes [27]. However in PLATO, the knowledge graph contains 108,447 and the smallest dataset has 12,932 features corresponding to nodes.

C PLATO's performance across node embedding methods for pre-training the feature embeddings

We conduct an ablation study to assess how PLATO's performance depends on the node embedding method used to pre-train the feature embeddings (Methods 3.4). We test three shallow node embedding methods for knowledge graphs which are scalable and prominent: TransE [72], DistMult [78], and ComplEx [65]. We find that PLATO's performance is similar across TransE, DistMult, and ComplEx (Table 9). More generally, PLATO makes no assumption about what type of self-supervised node embedding method is used to pre-train the feature embeddings. The self-supervised embedding step is simply a module that pre-trains feature embeddings which are then passed to the message passing and weight inference modules of PLATO.

KG Node Embedding Method	PearsonR (Test) on BRCA Dataset
TransE DistMult ComplEx	$\begin{array}{c} 0.582 \pm 0.025 \\ 0.575 \pm 0.011 \\ 0.583 \pm 0.019 \end{array}$

Table 9: PLATO's performance is consistent across knowledge graph node embedding methods.

D Rank ordering of methods for datasets with $d \gg n$

In Table 10, we show the rank order performance of all models on all $d \gg n$ datasets. We find that PLATO exhibits consistent and strong performance while the performance of the baselines depends on the specific $d \gg n$ dataset. For example, TabTransformer is the second best performing of all models on the MNSCLC dataset but the worst performing of all models on the PDAC and CH datasets. Similarly, XGBoost is the second best performing of all models on PDAC but only the tenth best performing of all models on BRCA. The baselines with the most stable performance are LASSO and Ridge Regression which rank consistently between the second and eighth best of all models.

Table 10: For datasets with $d \gg n$, PLATO exhibits consistent and strong performance. By contrast, the performance of the baselines varies with each dataset. For every dataset, the rank order of performance from Table 1 is shown. The best overall model is in **bold** and the second best model is underlined.

Da	Dataset		СМ	PDAC	BRCA	CRC	СН
# of features d	/ # of samples n	52.2	46.1	40.3	28.2	22.6	19.7
Classic Stat ML	Ridge	8	3	4	<u>2</u>	3	5
Dim. Reduct.	PCA	7	12	10	7	10	8
Feat. Select.	LASSO STG	6 9	$\frac{2}{4}$	3 8	5 4	$\frac{2}{7}$	4 7
Decision Tree	XGBoost	14	9	<u>2</u>	10	5	2
Graph Reg.	GraphNet NC LASSO Network LASSO	5 4 3	7 5 8	9 5 12	8 6 9	12 9 11	3 6 9
Param. Infer.	Diet	13	13	6	12	13	11
Tabular DL	MLP NODE TabTransformer TabNet	$\begin{vmatrix} 10\\12\\2\\11 \end{vmatrix}$	6 10 11 14	7 11 14 13	11 3 13 14	4 6 8 14	12 10 14 13
Ours	PLATO	1	1	1	1	1	1

Table 11: For datasets with $d \sim n$, PLATO is competitive with baselines. For XGBoost is consistently the strongest baseline. For every dataset, the rank order of performance from Table 11 is shown. The best overall model is in **bold** and the second best model is underlined.

Da	Dataset			SCLC	NSCLC
# of features d	# of features d / # of samples n			1.7	1.1
Classic Stat ML	Ridge	9	9	9	4
Dim. Reduct.	PCA	13	12	13	7
Feat. Select.	LASSO	7.5	6	6	8
i cut. Sciect.	STG	5	5	7	6
Decision Tree XGBoost		1.5	<u>2</u>	<u>2</u>	1
	GraphNet	6	4	5	9
Graph Reg.	NC LASSO	4	3	4	3
	Network LASSO	11	10	11	13
Param. Infer. Diet		14	13	14	14
	MLP	10	8	10	10
Tabular DL	NODE	3	11	3	12
	TabTransformer	12	14	12	11
	TabNet	7.5	7	8	5
Ours PLATO		1.5	1	1	2

E Graph regularization baselines

We test the state-of-the-art graph regularization baselines GraphNet [20], Network-Constrained LASSO [40], and Network LASSO [23]. The graph regularization baselines can only consider a homogeneous graph with only features as nodes and a single edge type. For the graph regularization baselines, we thus induce a subgraph between feature nodes from the knowledge graph and collapse all edge types between feature nodes into a single edge type. In this context, GraphNet, Network-constrained LASSO, and Network LASSO correspond to a LASSO model with a mean-squared error loss and a graph regularization penalty. Let λ be the graph regularization coefficient, j and k be two input features, let E be the set of edges in the graph, let $\Theta \in \mathbb{R}^d$ be the weights of the linear regression for d input features, and let D_j be the degree of feature node j. The graph regularization penalty for GraphNet is $\lambda \sum_{j,k \in E} (\Theta_j - \Theta_k)^2$, the penalty for Network-constrained LASSO is $\sum_{j,k \in E} |\Theta_j - \Theta_k|$.

F Number of trainable weights in PLATO vs. a multilayer perceptron

Table 12: **PLATO drastically reduces the number of trainable weights compared to a multilayer perceptron (MLP) across all of the datasets.** The number of trainable weights in the best model from the hyperparameter sweep is shown for each dataset.

Model	MNSCLC	СМ	PDAC	BRCA	CRC	CH ME	BC	SCLC	NSCLC
MLP	429665	416961	820097	425217	200529	589761 586945	296113	298929	594209
Plato	17154	42498	32066	17154	28386	61890 17154	28386	32066	17154

G Dataset Details

We use 6 datasets with $d \gg n$ and 4 datasets with $d \sim n$ [16, 17, 30, 79]. In all datasets, a machine learning model must predict the response of a cell line or a mouse tumor model to a drug. As input, the model considers a tabular dataset. In the tabular dataset, every row corresponds to a specific cell line or mouse tumor model. Every column corresponds to a gene name. Every value corresponds to the amount of that gene in the cell line or in the mouse tumor (*i.e.* gene expression). In practice, the number of genes is large for all tasks and the number of cell lines or mouse tumor models is comparatively small (*i.e.* $d \gg n$). For every row, the model also takes as input a fixed feature vector corresponding to the drug (*i.e.* a 200-dimensional ComplEx [65] embedding of the drug node in the knowledge graph). The output label is the response of the cell line (*i.e.* ln-ic50) [30, 79, 17] or mouse tumor model (*i.e.* minimum average percent tumor growth "min-avg-pct-tumor-growth") to the drug [16]. Data is available at https://github.com/snap-stanford/plato.

Gene expression datasets were pre-processed following a standard process in [48]. Briefly, gene expression values underwent TMM normalization and log transformation (*i.e.* log(x + 1)). Values were made to have zero mean and unit standard deviation. Dataset abbreviations are breast carcinoma (BC) [30, 79, 17], breast carcinoma (BRCA) [16], chondrosarcoma (CH) [30, 79, 17], colorectal cancer (CRC) [16], cutaneous melanoma (CM) [16], melanoma (ME) [30, 79, 17], non-small cell lung carcinoma (MNSCLC) [16], non-small cell lung carcinoma (NSCLC) [30, 79, 17], pancreatic ductal carcinoma (PDAC) [16], and small cell lung carcinoma (SCLC) [30, 79, 17].

H Knowledge Graph Details

We compile a general biomedical knowledge graph from prior studies [44, 35, 38, 56, 63, 74, 75] to use across all datasets. A schematic of the KG is in Supplementary Figure 2. A detailed breakdown of relation types is in Supplementary Table 13. The knowledge graph is available at https://github.com/snap-stanford/plato.

The knowledge graph contains 108,447 total nodes, including 7,975 drugs, 18,370 diseases, 11,447 phenotypes, 22,319 genes, 11,153 molecular functions, 28,748 biological processes, and 4,184 cellular components. Every gene and every drug in every dataset is present as a node in the knowledge graph. The knowledge graph also contains 3,066,156 edges with 99 distinct relation types. The remaining node types and their relationships serve as broader domain knowledge.

Edges between drug nodes and gene/protein nodes were derived from Drugbank [75], Gao [16], and the Genomics of Drug Sensitivity in Cancer [79, 30, 17]. Edges between diseases and genes/proteins were derived from DisGeNet [5]. Edges between diseases and phenotypes were derived from the Human Phenotype Ontology [35]. Edges between drugs and diseases were derived from the Multiscale Interactome [56]. Edges between drugs and side effects were derived from SIDER [38]. Edges between genes/proteins and other genes/proteins were derived from BioGRID [49], [55], the Database of Interacting Proteins [57], [44], [47], [53], [82], [69], and STRING [63]. Finally, edges from genes/proteins to molecular functions, biological processes, and cellular components as well as edges between molecular functions, biological processes, and cellular components were derived from the Gene Ontology [9].

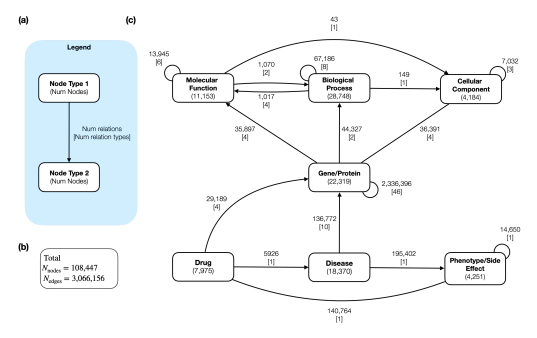


Figure 2: **Knowledge graph as a unified knowledge backbone.** We constructed a knowledge graph as a unified knowledge backbone across all 6 datasets. (a) Legend. For each node type, the number of nodes is given in parentheses. Between node types, the number of edges and the number of relation types are given. (b) Number of total nodes and edges across entire knowledge graph. (c) Visual schematic of knowledge graph across each node type.

Head type	Relation	Tail type	#
JI			edges
BiologicalProcess	EndsDuring	BiologicalProcess	1
BiologicalProcess	HappensDuring	BiologicalProcess	8
BiologicalProcess	HasPart	BiologicalProcess	229
BiologicalProcess	IsA	BiologicalProcess	53015
BiologicalProcess	NegativelyRegulates	BiologicalProcess	2768
BiologicalProcess	PartOf	BiologicalProcess	5193
BiologicalProcess	PositivelyRegulates	BiologicalProcess	2756
BiologicalProcess	Regulates	BiologicalProcess	3216
BiologicalProcess	OccursIn	CellularComponent	149
BiologicalProcess	HasPart	MolecularFunction	173
BiologicalProcess	NegativelyRegulates	MolecularFunction	269
BiologicalProcess	PositivelyRegulates	MolecularFunction	274
BiologicalProcess	Regulates	MolecularFunction	301
_	HasPart		179
CellularComponent CellularComponent	IsA	CellularComponent CellularComponent	4863
CellularComponent	PartOf	CellularComponent	1990
Disease	AlteredExpression	Gene	7157
Disease	Biomarker	Gene	107160
Disease	ChromosomalRearrangement	Gene	162
Disease	FusionGene	Gene	166
Disease	GeneticVariation	Gene	15076
Disease	GermlineCausalMutation	Gene	4677
Disease	ModifyingMutation	Gene	10
Disease	SomaticCausalMutation	Gene	130
Disease	SusceptibilityMutation	Gene	441
Disease	Therapeutic	Gene	1793
Disease	Has	Phenotype	195402
Drug	Treats	Disease	5926
Drug	Carries	Gene	866
Drug	Enzymes	Gene	5382
Drug	Targets	Gene	19817
Drug	Transports	Gene	3124
Drug	Has	Phenotype	140764
Gene	Associates	BiologicalProcess	43857
Gene	NotAssociates	BiologicalProcess	470
Gene	Associates	CellularComponent	35306
Gene	Colocalizes	CellularComponent	914
Gene	NotAssociates	CellularComponent	160
Gene	NotColocalizes	CellularComponent	11
Gene	Acetylation	Gene	9
Gene	Activation	Gene	58502
Gene	AdpRibosylation	Gene	2
Gene	Ampylation	Gene	5
	1.		18
Gene	Association	Gene	
Gene	Binary	Gene	56565 287641
Gene	Binding	Gene	287641
Gene	Catalysis	Gene	344801
Gene	Cleavage	Gene	22
	Complexes	Gene	62552
			52
Gene	CovalentBinding	Gene	
Gene Gene	CovalentBinding Deacetylation	Gene	8
Gene Gene Gene	CovalentBinding Deacetylation Demethylation	Gene Gene	8 6
Gene Gene Gene Gene Gene	CovalentBinding Deacetylation	Gene	8

Gene	DirectInteraction	Gene	290
Gene	DisulfideBond	Gene	5
Gene	DosageGrowthDefect	Gene	9
Gene	DosageLethality	Gene	112
Gene	DosageRescue	Gene	63
Gene	Enzymatic	Gene	2
Gene	Expression	Gene	188
Gene	GeneticInterference	Gene	32
Gene	Hydroxylation	Gene	26
Gene	Inhibition	Gene	201
Gene	Kinase	Gene	119
Gene	Literature	Gene	174
Gene	Metabolic	Gene	106
Gene	Methylation	Gene	25
Gene	NegativeGenetic	Gene	344
Gene	OxidoreductaseActivityElectronTransferAssay	Gene	2
Gene	PhenotypicEnhancement	Gene	209
Gene	PhenotypicSuppression	Gene	214
Gene	Phosphorylation	Gene	166
Gene	Phosphotransfer	Gene	1
Gene	PhysicalAssociation	Gene	824
Gene	PositiveGenetic	Gene	233
Gene	PostTranslationalModification	Gene	530
Gene	ProteinCleavage	Gene	48
Gene	PutativeSelfInteraction	Gene	3
Gene	Reaction	Gene	400
Gene	Regulation	Gene	265
Gene	Signaling	Gene	654
Gene	SyntheticGrowthDefect	Gene	407
Gene	SyntheticLethality	Gene	816
Gene	SyntheticRescue	Gene	91
Gene	Associates	MolecularFunction	35(
Gene	Contributes	MolecularFunction	596
Gene	NotAssociates	MolecularFunction	285
Gene	NotContributes	MolecularFunction	4
MolecularFunction	PartOf	BiologicalProcess	106
MolecularFunction	Regulates	BiologicalProcess	2
MolecularFunction	OccursIn	CellularComponent	43
MolecularFunction	HasPart	MolecularFunction	204
MolecularFunction	IsA	MolecularFunction	136
MolecularFunction	NegativelyRegulates	MolecularFunction	42
MolecularFunction	PartOf	MolecularFunction	11
MolecularFunction	PositivelyRegulates	MolecularFunction	27
MolecularFunction	Regulates	MolecularFunction	30
Phenotype	IsA	Phenotype	140