CLINICAL ADVANCEMENT FORECASTING *

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Related Sciences

ABSTRACT

Choosing which drug targets to pursue for a given disease is one of the most impactful decisions made in the global development of new medicines. This study examines the extent to which the outcomes of clinical trials can be predicted based on a small set of longitudinal (temporally labeled) evidence and properties of drug targets and diseases. We demonstrate a novel statistical learning framework utilizing both constrained and unconstrained linear and tree models, that can predict clinical advancement for the top 2% of target-disease pairs 1.5-2x more effectively than Open Targets composite scores as well as common measures of genetic support that have already been established – and observed in this study – to confer a 2x higher likelihood of success (a 3-4x cumulative improvement). Utilizing a subset of our biomedical evidence base, non-negative linear models can produce simple weighting schemes across various types of human, animal, and cell model genomic, transcriptomic, proteomic, and clinical evidence to identify target-disease pairs with high probabilities of advancing beyond phase 2 trials, and uncover a range of previously undeveloped target-disease pairs poised for clinical success. In this study we further explore: i) how longitudinal treatment of evidence relates to leakage and reverse causality in biomedical research and how temporalized evidence can mitigate common forms of potential biases and inflation ii) the relative impact of different type of features on our predictions; and iii) an analysis of the space of currently undeveloped, tractable targets predicted with these methods to have the highest likelihood of clinical success. To ease reproduction and deployment, no data is used outside of Open Targets and the described methods require no expert knowledge, and can support expansion of lines of evidence to further improve performance.

1 Introduction

It has been well established that drugs with human genetic evidence linking their respective targets to indications in clinical trials are more likely to succeed [1, 2, 3, 4, 5, 6, 7, 8, 9], and to a lesser extent, that the same may also be true for single-cell transcriptomic evidence [10]. This information has been used to devise target and target-disease ranking algorithms based primarily on a synthesis of multiple genetic signals alone [11, 12, 13]. It is also possible to expand the breadth of this genetic support to more targets and diseases based on knowledge graphs, protein interactions and/or disease ontologies [14, 15, 16, 17, 18]. To our knowledge, all such expansion methods identify a larger space of opportunities at the expense of expected success rates. This is not a focus of this work as we aim, instead, to establish a framework for identifying target-disease (TD) pairs with the very highest possible likelihood of success. We accomplish this by integrating human clinical, genetic/genomic, transcriptomic and proteomic data as well as cell/animal model evidence, pathway information and basic literature metrics from Open Targets [12] in statistical models that are trained on past clinical outcomes, evaluated primarily on their performance at upper extremes of predictions and then deployed on undeveloped TD pairs to establish an evidence-based ranking pipeline. This method can be contrasted with far more

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integrative methods that rely on neural and/or graph models over extensive knowledge graphs [19, 20, 21, 22], which are more complex and difficult to interpret. We believe a desirable middle ground between these approaches and those that aim to combine many orthogonal indicators of success through expert knowledge in heuristic systems [23, 12] would: 1) permit inclusion of many types of evidence, spanning hundreds or even thousands of putative orthogonal, causal predictors, 2) be highly interpretable, 3) support expert judgement where necessary, and 4) not require manual ranking/weighting schemes. This is the basis for our internal ranking platform, and such operational advantages have made it possible to reach or exceed the inclusivity of highly integrative methods without sacrificing interpretability, modularity and ease of deployment.

A substantial challenge inherent to building such a system is the need to account for the longitudinal nature of knowledge discovery in biomedicine. This is vital because any method that optimizes for likely *future* clinical success based on *historical* clinical outcomes may easily be biased by the non-random nature with which evidence can be absent otherwise. We use longitudinal evidence, i.e. evidence for which timing of its emergence can be determined, where appropriate when training and evaluating our methods before ultimately applying them to present-day evidence that is not temporalized. We discuss motivations, prior research and our own analysis on how important this problem is for each type of evidence in Section 2.7.

While the use of longitudinal evidence is very rare in studies on target prioritization algorithms, it is not at nearly as rare in studies that attempt to predict clinical trial outcomes using a variety of biological evidence sources, clinical statistics and other organizational or behavioral factors [24, 25, 26]. The need for this is often clear in that setting where the inclusion of predictors like historical success rates for targets or diseases, trial sponsor track records, eventual patient enrollment, etc. constitute clear information leaks otherwise. This is discussed in some depth in [24] which notes several studies that do not account for this problem before drawing a clear distinction between quasi-prospective and prospective problem formulations. The difference between the two is that the former reconstructs timelines for predictors and outcomes based on recorded event dates while the latter relies on frozen predictions that are never evaluated until sufficient time has elapsed for more outcomes to occur. Nomenclature for these formulations is conflicting though, where this definition of a quasi-prospective formulation is deemed entirely prospective in some cases, e.g. [27]. To avoid any potential confusion, our formulation in this study is quasi-prospective. The distinct advantages and disadvantages of each formulation provide a complementary risk/reward trade-off that we capitalize on in our internal platform, and we assert that the distinction between them is important. Interestingly, the existence of Open Targets snapshots dating back to at least 2016 would enable the extension of this work to a prospective formulation as well. This is discussed more in Section 4.

The preceding works discussed so far can largely be categorized as either 1) target and target-disease prioritization methods evaluated based on how well they correlate with observed clinical trial success and 2) clinical trial outcome prediction models. Both are measured against the same outcomes and an important distinction between them lies in how the prioritization methods are **not** directly optimized for those outcomes while the outcome prediction methods are. In this study, we attempt to bridge these methodologies by predicting clinical trial advancement for target-disease pairs based solely on information that could be present well in advance of any drug program or individual trial. We then calibrate these predictions to determine what thresholds are necessary to match the observed success rates from benchmarks for genetic support like OMIM [28], ClinVar [29] and GWAS. Finally, we examine how many present-day target-disease pairs are undeveloped (i.e. have never been in clinical trials), have a tractable target and are likely to see success rates matching or exceeding those benchmarks.

2 Results

In order to model clinical advancement for target-disease pairs, we first define "advancement" as progression beyond any particular trial phase across all drugs associated with any one TD pair as indicated by the presence of a later-stage trial. All results to follow consider only advancement beyond phase 2 due to limitations described in Section 4. This binary outcome is then predicted based on a list of features shown in Table 1. Information for each of these features is only used when it was published before the year **prior** to the first phase 2 trial observed, with an exception for genetic

evidence discussed in Section 2.7. A training dataset is then formed by including only TD pairs where this first phase 2 year is between 1990 and 2015. The evaluation dataset then consists of all TD pairs entering phase 2 between 2016 and 2022, with a 2 year offset from the present year (2024) to allow enough time for some trials to complete. While the average phase 2 trial duration may be as low as 2 years [30], other estimates would suggest half of them take longer than 2.9 years [31]. This means a substantial fraction of outcomes are censored, that this is an important parameter to test sensitivity to and that time itself is likely to be a crucial covariate in this formulation. The distribution of these outcomes, the number of associated targets/diseases and a variety of other statistics on this dataset are presented in Supplementary Figure 8.

2.1 Features

The features used throughout this study consist of 27 target-disease pair predictors, 5 target-specific predictors and 1 disease-specific predictor. These are listed in Table 1. The target and disease specific features are chosen carefully such that they are either capable of being associated with years in which events supporting them occurred or result from large-scale, unbiased methods that do not favor well-studied or drugged targets/diseases. Examples of this include target-specific tissue expression specificity scores computed from Human Protein Atlas [32] and LOEUF [33] scores from gnomAD. Simply put, our dataset combines scores from Open Targets for target-disease evidence and a select subset of target prioritisation [34] fields with almost no modifications, other than to add target and disease specific indicators of maximum trial phases reached and two extra genetic association features (described in Section 5).

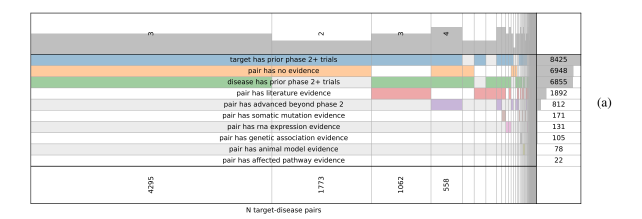
The combination of target, disease and target-disease features creates sparsity patterns that are important to understand before interpreting models built from it. Figure 1 demonstrates this sparsity by showing that of all the TD pairs entering phase 2 trials for the first time between 2016 and 2022 in our evaluation dataset (N=9010), less than 2% of them ever have evidence directly linking them other than literature co-mentions, which exist for 21% of those pairs. These TD pairs do, however, very frequently have prior clinical evidence for their associated targets and diseases. Specifically, 8,425 (94%) have a target and 6,855 (76%) have a disease that had already been in phase 2 or later trials previously. This is consistent with herding effects observed in recent drug development pipelines [35] over the same time period (2016-2022) and underscores the prognostic value such information may have as it is becoming more and more common and clearly confers lower clinical risk for new drug programs. We also observe that, on top of the clear theoretical, causal relationship between prior target and disease human clinical validation and the likely success of programs for new combinations of such targets and diseases, these features have strong, univariate predictive effects in the evaluation dataset of our study. This is illustrated in Supplementary Figure 10, which shows the relative risk of advancement for these features capturing the highest clinical stage previously reached by a target or disease.

Taken together, the paucity of TD pair evidence and the abundance of prior target or disease clinical validation should be considered carefully when interpreting predictive performance. TD pairs predicted to be highly likely to advance clinically despite a lack of target-disease-specific evidence are entirely plausible, but the value of such a prediction is dependent upon the application. We choose to minimize the influence of these cases in our study by focusing on rankings within therapeutic areas that do not extend beyond the number of TD pairs with direct evidence of some kind, as discussed in Section 5. This choice is also reflected in our primary performance metrics, as discussed in Section 2.3.

2.2 Models

We train a variety of models including constrained and unconstrained linear and tree models. The constrained variants of these models force effects of all features to increase monotonically, i.e. all effects are constrained to be non-negative. This is possible with no underlying feature transformations because all scores in Open Targets are constructed such that higher scores are presumed to be advantageous.

We also apply these models to our evaluation dataset using several feature ablations in order to assess the value of groups of related features. We refer to a "core" feature set consisting of all features listed in Table 1 except for the sole feature capturing the time since a target-disease pair first entered phase 2 trials (i.e. target_disease__time__transition).



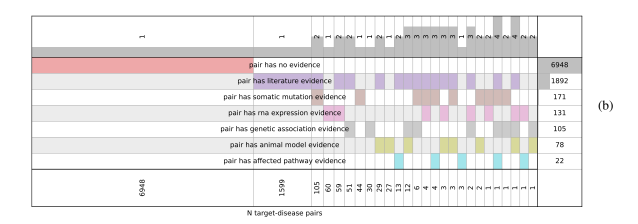


Figure 1: **Evaluation dataset feature presence**. (a) Presence and co-occurrence of various feature groupings or individual features indicated by overlapping sets of TD pairs, where the size of each bar is proportional to the number of TD pairs with a given combination of features and the sort ordering is determined by the number of TD pairs associated with any one feature (show in the right margin). The top margin shows the number of overlapping features for a combination of features and the bottom margin shows the number of TD pairs that are a part of that same combination. (b) Same as (a) without truly proportional bar sizes that are not scaled down beyond a limit necessary to fit all margin labels and combinatorial groupings. Features that are not specific to TD pairs are also omitted for brevity, along with the outcome feature used in this study and denoted here with the label "pair has advanced beyond phase 2".

Combinations of learning algorithms for the models and the feature sets to which they are applied are referred to using the following convention:

- RDG: Constrained, L2-regularized linear regressor (a.k.a. "Ridge regressor") fit with all core features
- **RDG-T**: **RDG** fit with all features instead of only core features, where the only difference is the inclusion of time since phase 2 transition for a TD pair
- RDG-X: RDG fit without human clinical and genetic evidence features
- GBM: Constrained, gradient-boosted machine with fit with all core features
- GBM-T: GBM fit with all features instead of only core features
- OTS: Open Targets composite score

The omitted human clinical and genetic evidence features for the RDG-X model are all of those in Table 1 with the midfix "clinical" or "genetic association". We omit these features specifically because they are known or expected to

comprise good predictors of human clinical success, so their exclusion examines the extent to which only literature and animal model evidence along with target-specific properties accomplish this task.

In order to compare these models to an Open Targets composite score (OTS), we use an equally weighted sum of all scores from individual sources except for those assigned lower weights in [36]. Scores from these sources are multiplied by the corresponding weight before being summed and only the TD-specific features of Table 1 are used. Neither the target/disease specific features nor the time since phase 2 transition feature are included in this calculation.

2.3 Metrics

The primary performance metric used in this study is relative risk (RR). This metric is commonly used to assess univariate measures of genetic support [1, 2, 3] and can be more intuitively understood, in the context of this study, as the probability that a TD pair among the top N TD pairs as ranked by a particular method will advance beyond phase 2 trials divided by that same probability of advancement among TD pairs with a rank greater than N. This provides a means to compare multivariate, model-based methods to univariate methods on a common scale. More specifically, any RR metric reported for a model among top N rankings is defined as:

$$\frac{P(advancement|rank >= N)}{P(advancement|rank < N)}$$
(1)

and RR metrics reported for univariate methods based on Open Targets scores for a single type of evidence, where not stated otherwise, are defined as:

$$\frac{P(advancement|score > 0)}{P(advancement|score = 0)}.$$
(2)

The use of such a metric is essential for properly assessing performance in this forecasting problem. While we also report more common measures of classifier performance like Receiver Operating Characteristic (ROC) and Average Precision (PR), neither of these adequately capture behavior in the upper extremes of rankings due the sparsity with which TD pair evidence is present for pairs that have ever entered phase 2 trials. This sparsity is further exacerbated in this study by the constraint that most of that evidence must have existed *before* such trials began. For more details on the extent of this sparsity, see Section 2.1. The figure presented there, Figure 1, also demonstrates that a substantial fraction (78%) of TD pairs that ultimately advance beyond phase 2 trials have targets and/or diseases with prior clinical validation despite no direct evidence linking the pairs themselves, which means a comprehensive measure of classifier performance (e.g. ROC) is far more likely to reflect the extent to which disease-only historical, clinical information or target-only information – including other attributes like conservation, essentiality and tissue expression – can predict clinical success. Again, this is not our primary focus as we want to evaluate the maximum achievable performance in this forecasting problem, and we assert that this is best accomplished when one or more lines of target-disease-specific evidence are present.

Reasonable alternative choices for this primary metric include those more common in information extraction literature or other machine learning studies with a focus on ranking rather than classification, such as mean reciprocal rank (MRR), precision at k (P@k) and normalized discounted cumulative gain (NDCG) [37, 38]. Precision at k is the most similar among these to relative risk at k since it is equivalent to the numerator in the relative risk calculation. We use relative risk instead because it is more intuitive than most ranking metrics, has well established analytical solutions for confidence intervals [39] and is consistent with prior work in this field.

Lastly, we emphasize that the interpretation of "risk" for the relative risk metric is to be inverted in this context. A higher "risk" in this study actually corresponds to a greater probability of success. The name "Relative Success" is used for this metric instead in [3] even though it has the same underlying definition. We choose not to use this label because we also present generic performance measures like ROC and AP, thereby prioritizing consistency with a domain-independent nomenclature.

2.4 Performance

2.4.1 Open Targets comparison

Figure 2 demonstrates how well our primary model in this study, RDG, ranks TD pairs by comparison to a composite score from Open Targets, OTS. This comparison highlights relative risk (RR) as our primary performance indicator along with secondary measures of performance like Receiver Operating Characteristic (ROC) and Average Precision (PR), as discussed more in Section 2.3. The third ranking method presented in Figure 2, "RDG-T", differs from the RDG model only in that it uses time since the phase 2 transition as a predictive factor in addition to all others. We observe that the use of this information greatly improves standard performance metrics like receiver operating characteristic (ROC) and average precision (AP), however it adds little to no value in rankings beyond a level where substantial relative risk increases can be observed. In other words, it constitutes an effective but coarse mechanism for ranking TD pairs while lacking the high precision of other factors like genetic support. More implications of this and opportunities it may imply are discussed in Section 4. As a more practical concern, we refrain from focusing on RDG-T, or the similar GBM-T model, because neither is readily applicable to undeveloped TD pairs for which the time since phase 2 transition is not available. They do, however, present a useful performance ceiling towards which future work might build.

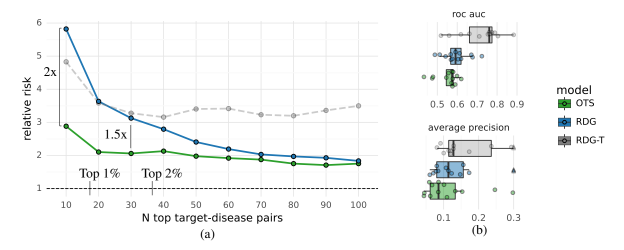


Figure 2: **Performance compared to Open Targets composite scores**. (a) Equally-weighted average relative risk estimates across 13 therapeutic areas, by number of top rankings and 3 methods: RDG (ours), RDG-T (ours) and OTS (Open Targets composite scores). (b) Receiver operating characteristic (ROC) and average precision scores across the same 13 therapeutic areas with no limit on the number of rankings. See Supplementary Figure 11 for raw data underlying (a). Annotations indicate RR multiples between RDG and OTS for the top 10 and top 30 TD pairs as well as how many TD pairs comprise the top 1% (N=18) and top 2% (N=36) of rankings on average across therapeutic areas. All results shown are from the evaluation dataset.

We also note that Figure 2 presents average RR estimates drawn across a subset of therapeutic areas, and the criteria used to select them is described more in Section 5. A full list of therapeutic areas meeting these criteria can be seen in Supplementary Figure 11 along with the RR estimates summarized in Figure 2. Furthermore, a comparison of the distribution of these estimates by model is presented in Supplementary Figure 12 along with the statistical significance of their differences.

2.4.2 Genetic benchmark comparison

In order to establish baseline levels of success and coverage across TD pairs, we examine ranking performance in comparison to well established, univariate indicators of genetic support in Figure 3. This figure presents OMIM and

GWAS baselines, in the parlance of [2], [1] and [3], as well as an intermediate baseline from the European Variation Archive (EVA) [40] containing evidence predominantly from ClinVar [29].

One key objective of this study is to determine if any model, e.g. RDG, can sort TD pairs with genetic support such that at least some portion of that sorted list has a likelihood of advancement that consistently exceeding what is expected from any one source of genetic support alone. We find that this goal is met and exceeded by the RDG model, which actually identifies more TD pairs than those that have either EVA or GWAS support alone at an expected rate of advancement exceeding that of the single source (respectively). This does not appear to be the case with the OMIM baseline, however the lack of examples in our evaluation dataset with OMIM support makes any determination difficult. See Section 2.6 for more on how these benchmarks are employed to contextualize opportunities among undeveloped TD pairs and Supplementary Figure 14 for top predictions from the RDG model. This latter, supplementary figure further emphasizes our focus on prioritizing opportunities beyond those with genetic support and provides examples of TD pairs with multiple lines of evidence.

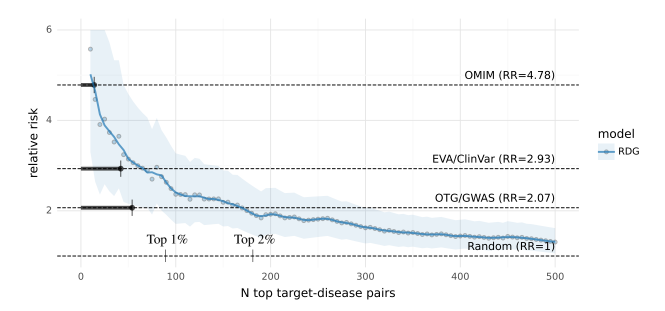


Figure 3: **Performance compared to genetic support benchmarks**. The RR estimates for each benchmark annotated on the right are shown as horizontal dotted lines and are calculated across all TD pairs (N=9010). The number of pairs with support from each benchmark is represented by the bars extending horizontally (to the right) from the y-axis. The RDG RR point estimate is shown in blue and bounds around that estimate correspond to a Katz 90% confidence interval. Annotations on the x-axis indicate how many TD pairs comprise the top 1% (N=90) and top 2% (N=180) of rankings. All results shown are from the evaluation dataset.

We also note that Supplementary Figure 9 shows confidence intervals for each of the genetic benchmarks of Figure 3 in isolation, as well as all other target-disease-specific evidence sources, in addition to confidence intervals for the RDG model at various top ranking cutoffs. Similar comparisons for target-specific and disease-specific features can be seen in Supplementary Figure 10. These findings suggest that 1) genetic support for TD pairs is highly predictive but rare, 2) human clinical support for targets and diseases in isolation is also predictive while being more common and 3) constraint and expression specificity of targets exhibit modest but significant effects. While only human clinical evidence appears to have a prognostic value rivaling that of genetic support when considered on a univariate basis, the combined influence of non-genetic, non-clinical evidence is examined in Section 2.4.3 where a model using only this information, RDG-X, still outperforms Open Targets composite scores by all measures.

2.4.3 Model comparison

Figure 4 presents average performance across therapeutic areas for select combinations of learning algorithm, constraint type and feature group described in Section 2.2. Several key findings illustrated in this figure are:

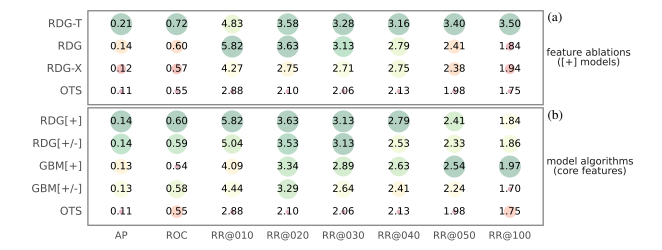


Figure 4: **Performance across model algorithms and feature ablation groups**. Average precision (AP) and receiver operating characteristic (ROC) scores with relative risk (RR) at ranking cutoffs denoted by RR@N . (a) Performance across constrained RDG models using different groups of features as described in Section 2.2 (b) Performance across constrained ([+]) and unconstrained ([+/-]) linear and gradient-boosted models using the core feature set.

- 1. The RDG-T model achieves far higher ROC and AP scores through the use of the time since transition feature, which indicates the number of years a TD pair has been underdevelopment after having reached phase 2.
- 2. The RDG model, however, matches or exceeds RDG-T in performance among top TD pairs
- 3. The RDG-X model, using no human clinical or genetic evidence linked to a disease, outperforms the Open Targets composite score and nearly matches the performance of the RDG model beyond top rankings
- 4. Linear models outperform gradient-boosting models by nearly all measures
- 5. Constrained linear models outperform unconstrained linear models by nearly all measures

We conclude from these results that constrained linear models are an optimal choice for this problem due both to their greater performance and interpretability. This interpretability is illustrated more in Section 2.5 and owed much to the effort Open Targets has already undertaken to construct evidence scores such that they can be assumed to have a monotonically increasing effect on the likelihood that a causal relationship exists between a target and a disease.

2.5 Effects

The coefficients learned by the RDG model, and the average effects they have across the evaluation dataset, are shown in Figure 5. This model most highly prioritizes genetic signals that have the greatest coverage, i.e. associations from GWAS studies through the ot_genetics_portal feature and associations from any curated clinical genetics source, i.e. EVA, Orphanet, UniProt, Genomics England, ClinGen and gene2phenotype, via the curated feature. Notably, literature and target/disease specific clinical features also have substantial effects, followed by indicators of animal evidence and target genetic constraint / expression specificity. Any features not shown were deflated to have no effect, which is possible in this model due to the non-negativity constraint. One such feature worth emphasizing is transcriptomic

evidence from Expression Atlas. We found this somewhat surprising, but it is supported by arguments against transcript over/under expression as an indicator of genes that influence disease rather than the other way around [41].

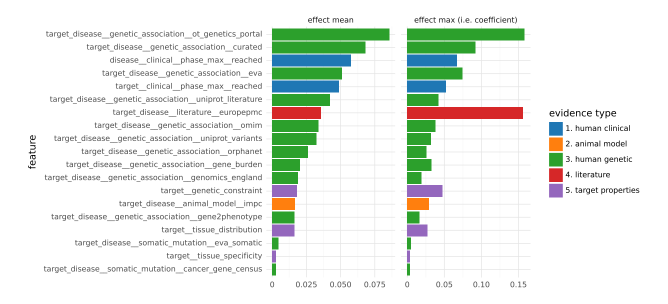


Figure 5: **RDG model feature effects**. The **effect max** values are equivalent to RDG model coefficients for the corresponding feature while the **effect mean** values indicate average values of the product between the coefficient and a particular feature value, when that feature is present.

It is worth noting that the discordance between the coefficients and the average feature effects of Figure 5 arises from both the frequency with which features exist and the distribution of their underlying scores. Scores for many clinical genetics features (e.g. OMIM, Genomics England, UniProt) are very frequently absent or close to 1. By comparison, scores for literature associations are typically far lower, even when limited only to cases where they exist, with a median value of 0.12 (mean=.23) in the evaluation data. This is why the europepmo feature has a relatively large associated coefficient, but a much smaller average effect across predictions.

2.6 Opportunities

A common method for identifying druggable opportunities within a specific disease context involves first ranking TD pairs according to some prioritization methodology followed by filtering or reprioritizing those ranks based on knowledge of target tractability [42, 43, 13]. We use a similar approach to identify tractable targets associated with TD pairs that have yet to enter clinical trials. To aid in interpreting this approach, we also draw on the results of Figure 3. The data in this figure suggests thresholds for the RDG model that align to expected rates of advancement compared to several genetic support benchmarks. These thresholds are used to bucket undeveloped TD pairs before further bucketing them based on levels of tractability. The tractability buckets in Supplementary Table 2 provide HIGH, MED, and LOW confidence ratings for each type of tractability evidence based on the priorities suggested in [44].

Figure 6 shows the distribution of TD pair counts for select buckets across therapeutic areas as well as across the current maximum phase reached for any one pair. We find that there are \sim 2,400 small-molecule-enabled, \sim 1,400 antibody-enabled, and 14 PROTAC-enabled TD pairs with a probability of advancement that is nearly 3x other TD pairs based on the EVA threshold RR=2.93 in Figure 3. Top antibody-enabled pairs are shown in Supplementary Figure 15 along with their corresponding genetic and clinical support.

	stage [threshold=EVA]					threshold [stage=NONE]			tractability [stage=NONE, confidence=HIGH, threshold=EVA]				
	ALL	NONE	Phase 1	Phase 2	Phase 3	Phase 4	EVA	ОМІМ	OTG	АВ	ос	PR	SM
therapeutic area													
ALL	9724	8560	165	367	274	358	8560	679	20821	1456	655	14	2408
genetic, familial or congenital disease	4867	4638	32	73	58	66	4638	402	7641	399	271	12	993
cancer or benign tumor	2063	1531	122	209	125	76	1531	96	6506	430	124	0	639
nervous system disease	2093	1923	10	54	49	57	1923	166	3710	153	74	3	446
musculoskeletal or connective tissue disease	1597	1469	15	46	35	32	1469	158	2863	206	129	0	353
gastrointestinal disease	1149	995	22	48	30	54	995	82	3265	257	67	0	334
immune system disease	1294	1114	30	69	36	45	1114	107	2762	339	69	4	290
nutritional or metabolic disease	1491	1409	1	16	16	49	1409	132	2316	94	29	0	310
endocrine system disease	1071	933	17	42	26	53	933	99	2685	184	80	4	287
cardiovascular disease	956	834	7	25	32	58	834	81	1908	126	70	0	315
psychiatric disorder	820	735	2	22	16	45	735	73	1442	59	18	2	168
disorder of visual system	838	823	1	8	3	3	823	87	1289	40	24	0	131
integumentary system disease	733	646	14	35	22	16	646	51	1502	115	40	0	164
hematologic disease	702	589	23	41	31	18	589	45	1394	158	73	1	202
respiratory or thoracic disease	590	472	8	52	33	25	472	23	1552	177	45	0	134
urinary system disease	550	490	5	19	19	17	490	42	1164	94	47	0	121
reproductive system or breast disease	534	452	9	30	18	25	452	33	1240	69	22	6	128
phenotype	353	307	3	10	10	23	307	13	921	74	54	0	142
pancreas disease	326	278	7	9	7	25	278	37	770	65	24	0	96
measurement	7	7	0	0	0	0	7	1	69	1	2	0	3
disorder of ear	5	5	0	0	0	0	5	0	21	1	0	0	1

Figure 6: **Present-day target-disease pair counts by stage, likelihood of advancement and tractability**. The **stage** panel contains counts by maximum trial phase reached, the **threshold** panel contains counts of pairs with a RDG model score exceeding that of the associated benchmark for only undeveloped pairs, and the **tractability** panel shows pair frequencies among undeveloped pairs exceeding the EVA threshold that also have a HIGH tractability rating as defined in Supplementary Table 2. This corresponds to targets that have all been in clinical development already, except for the **OC** modality in which case it indicates that a target has been approved.

2.7 Inflation

Like most studies of this kind, we assume a "closed-world" [19] over the space of target-disease pairs and any evidence between them. This means that we do not differentiate between evidence that an association for any one pair truly does **not** exist (or is too weak to be relevant under the omnigenic model [45]), and the lack of any attempt to find that evidence in the first place. This also means that our estimate of the prognostic value for any one evidence source is subject to historical trends in biomedical research and the myriad ways that this research can be biased towards particular targets and diseases. We avoid attempting to comprehensively survey these biases in favor of offering an illustrative list of specific examples that are relevant in this study:

- 1. Mendelian randomization research is biased towards cardiovascular diseases as they have a disproportionate number of known, modifiable exposures [46]
- 2. Putative protein interactions that do not result from genome-scale or otherwise unbiased assays result in an overrepresentation of successful drug targets in resources like STRING [47], thereby inflating the success of network expansion methods over these databases to identify such targets [16].
- 3. Transcript expression studies run in late-stage clinical trials for a single indication, e.g. [48] linking SLE to IFN genes, are a degenerate indicator of advancement beyond earlier stage trials when the timing of this evidence is not accounted for.
- 4. Targets tested against more indications in clinical trials enrich for failures because the marginal cost of testing more indications decreases, but the evidence for these indications is often weaker [14].

5. Herding effects in pharma R&D pipelines around particular drug targets are becoming increasingly clear over time [35] and generate an excess of clinical evidence for those targets.

We also note that the skew in basic drug target research towards those that already have rich annotations and well characterized molecular function [49] as well as the disproportionate representation of particular target families in pharma R&D pipelines [50, 51] and the fact that literature is well known to be biased away from negative results in general [52] are all problematic.

While it is not possible to address all of these issues, we emphasize that there is a clear pattern across the examples in the list above in that they require **past** clinical successes and/or failures to arise in the first place. This suggests that accounting for when evidence first emerged would limit the extent of these problems. We do so in this study based solely on publication dates associated with any one piece of information linking target-disease pairs. This also offers a novel opportunity to attempt to quantify what kind of evidence suffers most from these biases. Figure 7 presents results for this based on a relative risk statistic defined as:

$$\frac{P(A|B)}{P(A|\neg B)}\tag{3}$$

where:

- A is the event that evidence for a TD pair arises after its first early-stage (phase 1 or 2) trial rather than before
- B is the event that a TD pair advances into late-stage trials (phase 3 or 4)

We refer to this as "inflation risk" so as not to confuse it with the relative risk statistic used in all other contexts, and it can be more simply described as the fraction of TD pairs for which evidence arises **after** the beginning of an ultimately successful early-stage trial divided by that same fraction for TD pairs that do not advance to late-stage trials. The intuition for this statistic is that it will be higher if successful trials lead to the generation of evidence of a particular type, and it should be 1 in cases where the emergence of evidence is independent of clinical success. We also measure this potential lack of independence through the more commonly used Fisher's exact test, e.g. [53], and both are presented in Figure 7.

We find that evidence from Reactome is the worst offender by this metric, implying that it often only arises for TD pairs after a certain level of clinical success has been attained. We also find that long-running aggregators/curators of published research often focused on individual diseases/phenotypes, like Expression Atlas, IMPC, CGS and Cancer Biomarkers exhibit this form of inflation as well.

Sources of genetic evidence appear to be much less inflated, or have too little data to reach significance. This is to be expected for GWAS evidence arising from genome-wide, phenome-wide biobank consortia, however much of historical GWAS evidence is not phenome-wide. More context on how much this is likely to matter comes from [54] in which it was estimated that as few as 6% of 500 FDA-approved targets for non-cancer drugs arose from programs highly motivated by pre-existing genetic support and that "the remaining 94% were probably identified using conventional pharmacology, biochemistry or molecular biology approaches". We then speculate that if the initiation of new drug programs was not historically motivated highly by the existence of genetic support, then the incentives for pursing new genetic evidence based on clinical and commercial success are likely to be minimized. This, in conjunction with existing precedent [1, 2, 3, 4, 5, 7] and our inflation results, ultimately led us to the use of genetic evidence without temporalization. In other words, we do not treat genetic evidence as longitudinal features like all others associated with TD pairs. A breakdown of which features are treated in which manner is provided in Table 1.

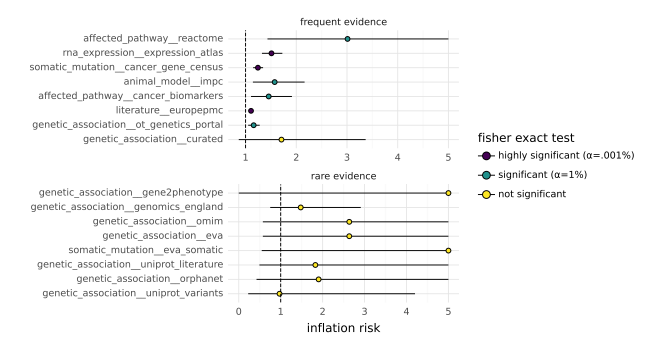


Figure 7: Clinical success drives evidence discovery.

3 Conclusion

We have demonstrated that simple machine learning methods applied to longitudinal biomedical evidence from many sources can be used to predict clinical outcomes for combinations of drug targets and diseases, without knowledge of molecular properties or trial design details. We have also shown that these methods are more precise in the extremes of their predictions than composite, heuristic scores like those from Open Targets. They also outperform such baselines by more comprehensive, traditional measures of classifier performance; however, we find this less compelling and easier to accomplish than improving performance among the upper tail of the opportunities implied by the very highest predictions. This framework would also support the addition of new lines of evidence over time well as it is designed to automatically determine the relevance of any new information without intervention. Lastly, we find that the space of present-day, undeveloped targets within a disease context that both exceed baseline levels of tractability and have a high predicted likelihood of clinical advancement is substantial. It is likely to grow quickly as well since the breadth of much of the underlying evidence is expanding rapidly [55, 56, 57].

4 Discussion

An important question, that remains difficult to answer, is how much target prioritization methods like the one explored in this study can improve the efficiency of drug discovery. One 2018 estimate suggests that improving success rates from phase 1 to approval by 1.4x (from 9.6% to 13.8%) would reduce the median R&D cost for a **single drug** by \$480 million USD [58]. If a 2x increase in success rates from phase 1 to approval could be expected when pursuing only genetically validated TD pairs [1], then this would increase to a \$686 million cost savings per drug. Pushing that even further to 3x or 4x increase by incorporating more lines evidence, as we have done here, would suggest a per-drug cost savings of \$1.03 billion and \$1.37 billion, respectively. These savings would only be realized for drugs tested against targets and diseases with sufficient supporting evidence, however, we show in Section 2.6 there are more than 8,000 such opportunities (i.e. untested, present-day TD pairs) where at least a 3x improvement in success rates could be expected.

Another way to quantify the impact that employing superior target prioritization strategies could have is to consider what would happen if this was done across the entire pharmaceutical development industry. Our internal research shows that \$130B (92%) of global pharma R&D costs (\$150B) are expended on failed assets each year. This assumes an 8% success rate that if improved by 3x to 24%, would lead to a total cost savings of approximately \$22B annually ...

Throughout the majority of this study, we focus primarily on performance among the top 2% of TD pairs. Reasons for this are introduced in Section 2.1 and Section 2.3. It is constructive to expand on this choice by comparing it to alternatives, and noting that it is quite likely that many other predictors could improve performance for the remaining 98%. We offer some evidence of this through the use of a feature indicating how long a TD pair has been in phase 2 trials, which improves wholistic measures of ranking performance like ROC by 12 points (see Figure 4). While time is a necessary but not sufficient condition for clinical advancement of a TD pair, given that a single trial can take years to complete, it also correlates with how many distinct trials, drugs, companies, sponsors, investigators, etc. attempt to validate any one pair. We posit that this is crucial because even if a druggable, mechanistic link exists between a target and a disease, many (or even most) trials testing that link will fail for reasons unrelated to efficacy or safety of a drug [4]. Some of these reasons include 1) commercial factors like a lack of funding, pipeline reprioritization and competitive density, 2) administrative or logistical factors like a lack of enrollment, retention problems, poor trial design, drug supply chain shortages, epidemics (COVID-19) and 3) regulatory factors like legislative shifts or extensive regulatory approval delays. Many of these factors are fundamentally difficult or impossible to predict. Epidemic outbreaks, labor shortages leading to supply chain problems and regulatory shifts are examples of exogenous shocks for which accurate predictors are unlikely to exist. However, factors like funding, competition, enrollment and certain aspects of trial design have measurable predictors such as preceding venture capital and private equity investments, patent filings and revenue for related drugs, disease severity and prevalence, and the existence of biomarkers or other enabling factors for better trial designs (respectively). As a more detailed example, patient recruitment failures are the most common reason trials fail [4] and recruitment statistics are highly predictive of eventual trial outcomes [26, 25]. It stands to reason then that predicting trial enrollment failures well in advance of when a trial is even conceived, i.e. for a target-disease pair, should be possible because recruitment in trials is determined, in part, by how prevalent a disease is, how debilitating it is, what patient demographics it inflicts, whether existing treatments exist for it and how successful recruitment for past trials (if any exist) has been – all of which are measurable. Signals like this could be used to further improve performance in the bottom 98% of TD pairs since they better capture causal factors in non-biological trial failures that time under development – the only non-biological feature in this study – does not.

Another notable focus of this study is on evidence for TD pairs that can be **directly** attributed to them. This is a departure from earlier research in this space like [1] that often includes measures of similarity between disease ontology terms to account for the fact that bridging disease nomenclatures used in clinical trial datasets with those used in databases maintaining genetic evidence is difficult. It is not uncommon for disease terms from either source to be sufficiently similar such that it is appropriate to consider them as equivalent. It can also be argued that similarity between disease terms offers an important dimension for expanding evidence in a biologically meaningful way, regardless of technical mapping issues. This is taken to further extremes in studies like [14] that propagates evidence using both measures of target similarity (from protein interaction networks) and measures of disease similarity (from ontologies and literature co-occurrence). We believe such expansion methods are very compatible with our approach by either using them to derive new predictors or to propagate predictions from models like these across networks or knowledge graphs.

- Discuss the possibility to do prospective evaluation with OT snapshots
- Cover limitations with OT concerning temporalization for both evidence and drug approvals, and why only transitions from phase 2 are relevant for this work
- Discuss the limitations of the tractability analysis and why they might not be tractable targets
- From [14]: "It is important to bear in mind therefore that what we are measuring when looking at historical trial outcomes is not an unbiased measure of any given gene's true disease associations, but rather a view on how useful a given evidence source or analytical method has been for choosing drug targets based on current

and historical drug discovery practices. Dramatic changes in these practices in the future could render some of our conclusions obsolete, though the fundamental observation that genetic association itself is retained in molecular networks will remain valid."

5 Methods

- Discuss why all target prioritisation data fields are not used due to the potential leakage they may impose (e.g. target families, GO annotations, etc.)
- Describe how therapeutic areas were selected based on having at least 100 TD pairs with target-disease-specific
 evidence of any kind and with explicit omissions: ("biological_process", "pregnancy or perinatal disease",
 "injury, poisoning or other complication", "pregnancy or perinatal disease", "medical procedure", "infectious
 disease", "animal disease")
- RDG models are fit using the LightGBM [59] algorithm Ridge implementation from scikit-learn and GBM models are fit using the
- Cite supervenn
- · Training/evaluation results are limited by temporalization while the present-day predictions are not
- OMIM is defined as EVA associations with publications
- The "genetic_association__curated" field is a union of all genetic association sources other than "gene_burden" and "ot_genetics_portal"
- Mention that the scores for a TD pairs in a year are the maximum score for that source, not the harmonic sum
- Mean imputation is used for target-specific features
- Mention specifics on RDG and GBM implementations, i.e. lightGBM and scikit-learn
- RDG is always trained on all features, but only applied to feature subsets where relevant

6 Appendix

6.1 Features



Figure 8: Training and evaluation dataset summary statistics.

Table 1: Features used in modeling and analysis

	feature	entity	kind
1	diseaseclinicalphase_maxreached	disease	temporal
2	targetclinicalphase_maxreached	target	temporal
3	targetgenetic_constraint	target	static
4	targetmouse_ko_score	target	static
5	targettissue_distribution	target	static
6	targettissue_specificity	target	static
7	target_diseaseaffected_pathwaycancer_biomarkers	target_disease	temporal
8	target_diseaseaffected_pathwaycrispr	target_disease	temporal
9	target_diseaseaffected_pathwaycrispr_screen	target_disease	temporal
10	target_diseaseaffected_pathwayprogeny	target_disease	temporal
11	target_diseaseaffected_pathwayreactome	target_disease	temporal
12	target_diseaseaffected_pathwayslapenrich	target_disease	temporal
13	target_diseaseaffected_pathwaysysbio	target_disease	temporal
14	target_diseaseanimal_modelimpc	target_disease	temporal
15	target_diseasegenetic_associationclingen	target_disease	static
16	target_diseasegenetic_associationcurated	target_disease	static
17	target_diseasegenetic_associationeva	target_disease	static
18	target_diseasegenetic_associationgene2phenotype	target_disease	static
19	target_diseasegenetic_associationgene_burden	target_disease	static
20	target_diseasegenetic_associationgenomics_england	target_disease	static
21	target_diseasegenetic_associationomim	target_disease	static
22	target_diseasegenetic_associationorphanet	target_disease	static
23	target_diseasegenetic_associationot_genetics_portal	target_disease	static
24	target_diseasegenetic_associationuniprot_literature	target_disease	static
25	target_diseasegenetic_associationuniprot_variants	target_disease	static
26	target_diseaseknown_drugchembl	target_disease	temporal
27	target_diseaseliteratureeuropepmc	target_disease	temporal
28	target_diseaseoutcomeadvanced	target_disease	temporal
29	target_diseaserna_expressionexpression_atlas	target_disease	temporal
30	target_diseasesomatic_mutationcancer_gene_census	target_disease	temporal
31	target_diseasesomatic_mutationeva_somatic	target_disease	temporal
32	target_diseasesomatic_mutationintogen	target_disease	temporal
33	target_diseasetimetransition	target_disease	temporal

7 Supplementary Material

7.1 Performance

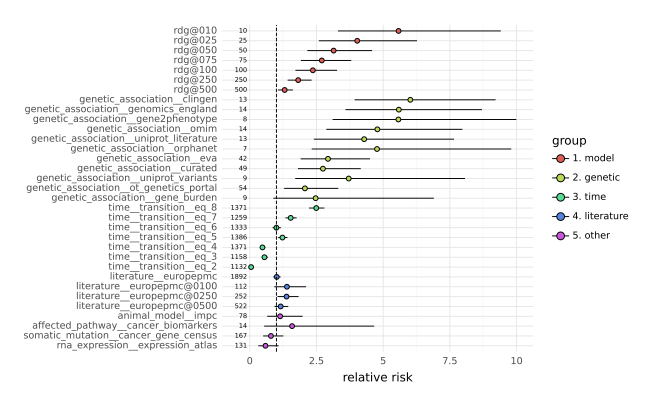


Figure 9: **Performance of individual features and predictive scores as measured by relative risk**. RDG model results denoted by rdg@N indicate performance for the N top TD pairs. The same convention is used for literature evidence and the time_transition_eq_X convention denotes RR estimates when the time since the phase 2 transition is equal to X years. The omim, eva, and ot_genetics_portal features correspond to the OMIM, EVA and OTG baselines of Figure 3, respectively. All other features are assessed based on their existence. The counts along the origin indicate how many TD pairs were used to compute the RR numerator.

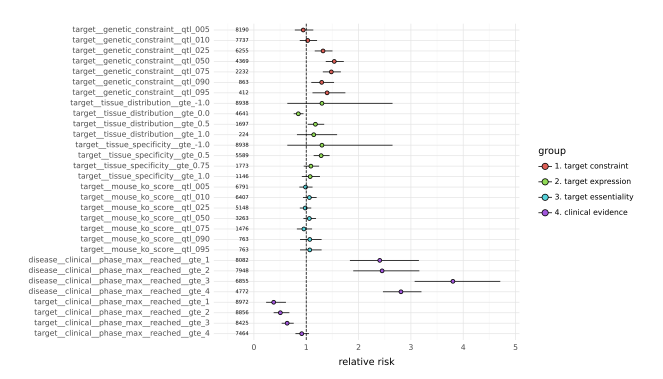


Figure 10: **Relative risk scores for target/disease features**. The features ending with qtl_Q denote binary indicators constructed from cases where the feature meets or exceeds quantile Q of its distribution. The features ending with gte_X denote indicators for when the feature meets or exceeds a specific value X.

		ОМІМ	EVA/ClinVar	OTG/GWAS	0TS@010	OTS@020	OTS@030	01S@050	OTS@100	RDG@010	RDG@020	RDG@030	RDG@050	RDG@100
therapeutic_area	n_pairs													
average		9.65	5.78	1.95	2.88	2.10	2.06	1.98	1.75	5.82	3.63	3.13	2.41	1.84
all	9010	4.78	2.93	2.07	6.70	3.91	3.35	2.69	2.13	5.58	3.91	3.73	3.14	2.37
cancer or benign tumor	4013		3.69	0.00	2.58	3.91	5.31	3.65	2.68	5.20	5.00	4.40	4.28	3.26
genetic, familial or congenital disease	2035	3.43	2.37	0.68	3.08	1.53	1.02	0.81	0.91	3.08	2.58	1.71	1.23	0.80
nervous system disease	1534	3.64	1.95	0.00	2.54	1.15	1.70	1.54	2.14	2.54	1.26	2.59	2.08	1.56
gastrointestinal disease	1260		0.00	3.48	5.68	4.33	2.68	3.17	2.82	8.72	5.90	3.90	4.34	2.44
immune system disease	1225	21.09	21.44	2.88	6.39	4.30	4.43	2.61	1.99	11.05	5.48	4.43	3.10	2.53
hematologic disease	1023	26.89	27.59	0.00	5.48	2.71	1.79	2.05	1.68	8.44	4.18	3.78	2.22	1.64
endocrine system disease	949	0.00	0.00	11.26	0.00	0.00	2.12	2.88	2.21	10.83	5.36	4.90	2.88	1.77
reproductive system or breast disease	948		0.00	0.00	0.00	0.00	0.00	1.68	1.72	8.79	4.35	3.95	2.99	2.51
musculoskeletal or connective tissue disease	757	8.82	6.67	2.68	4.74	3.62	2.73	2.48	2.18	6.02	5.00	4.76	3.09	2.80
respiratory or thoracic disease	644		0.00	0.00	4.37	2.74	1.80	1.97	1.52	3.22	2.15	2.19	1.51	1.83
cardiovascular disease	617	3.65	1.66	2.09	1.83	1.66	1.48	0.86	0.87	2.21	1.85	1.35	0.93	0.68
integumentary system disease	563			0.00	0.00	0.00	0.82	1.43	1.49	4.37	2.94	1.92	1.76	1.12
urinary system disease	561		3.99	2.29	0.79	1.40	0.92	0.61	0.59	1.19	1.19	0.78	0.86	0.93

Figure 11: **Relative risk scores by method, benchmark and therapeutic area**. The **average** therapeutic area indicates mean values across all others except for **all**, which is an ungrouped estimate across all diseases regardless of therapeutic area.

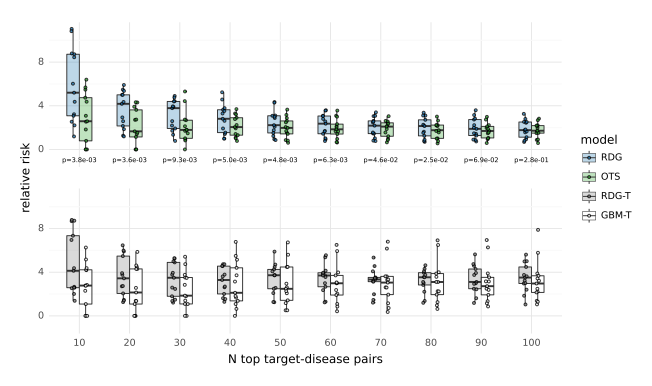


Figure 12: **Relative risk distributions across select therapeutic areas**. P-values are computed from a one-sided Wilcoxon signed-rank test with the alternative that the RDG model RR averages across therapeutic areas exceed OTS averages.

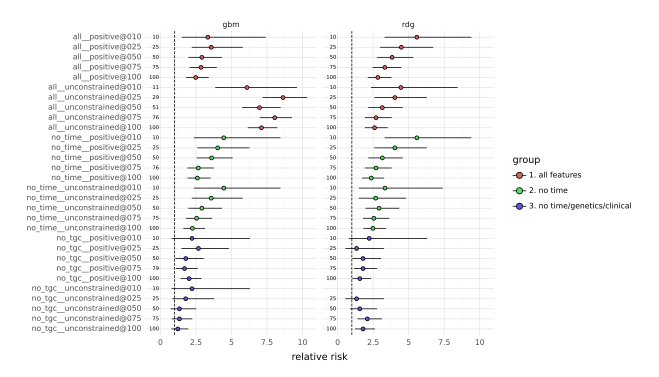


Figure 13: Performance by algorithm, constraint type and feature grouping.

7.2 Predictions

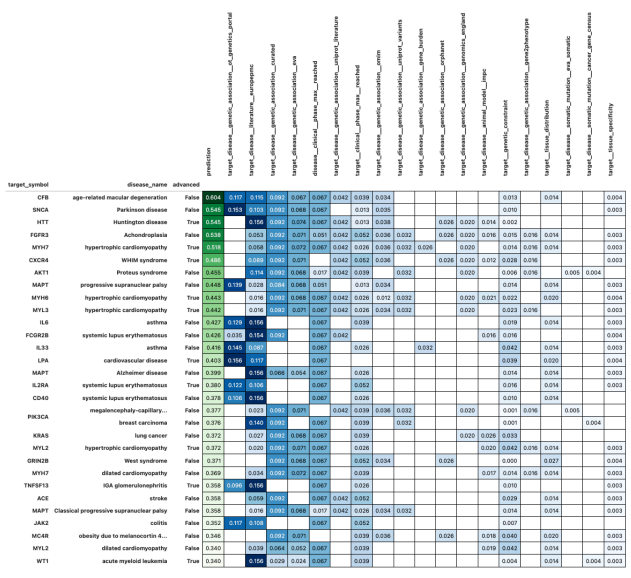


Figure 14: **Top RDG model evaluation dataset predictions**. Feature contributions are shown as the product of their underlying values and the RDG coefficients. The **advanced** field indicates whether the associated TD pair advanced beyond phase 2 as of 2024.

7.3 Opportunities

Table 2: Tractability bucket assignments

	evidence	modality	
1	Phase 1 Clinical	OC	LOW
2	Advanced Clinical	OC	MED
3	Approved Drug	OC	HIGH
4	GO CC med conf	AB	LOW
5	Human Protein Atlas loc	AB	LOW
6	UniProt SigP or TMHMM	AB	LOW
7	UniProt loc med conf	AB	LOW
8	GO CC high conf	AB	MED
9	UniProt loc high conf	AB	MED
10	Advanced Clinical	AB	HIGH
11	Approved Drug	AB	HIGH
12	Phase 1 Clinical	AB	HIGH
13	Database Ubiquitination	PR	LOW
14	Half-life Data	PR	LOW
15	Small Molecule Binder	PR	LOW
16	Literature	PR	MED
17	UniProt Ubiquitination	PR	MED
18	Advanced Clinical	PR	HIGH
19	Phase 1 Clinical	PR	HIGH
20	Druggable Family	SM	LOW
21	High-Quality Pocket	SM	LOW
22	Med-Quality Pocket	SM	LOW
23	High-Quality Ligand	SM	MED
24	Structure with Ligand	SM	MED
25	Advanced Clinical	SM	HIGH
26	Approved Drug	SM	HIGH
27	Phase 1 Clinical	SM	HIGH

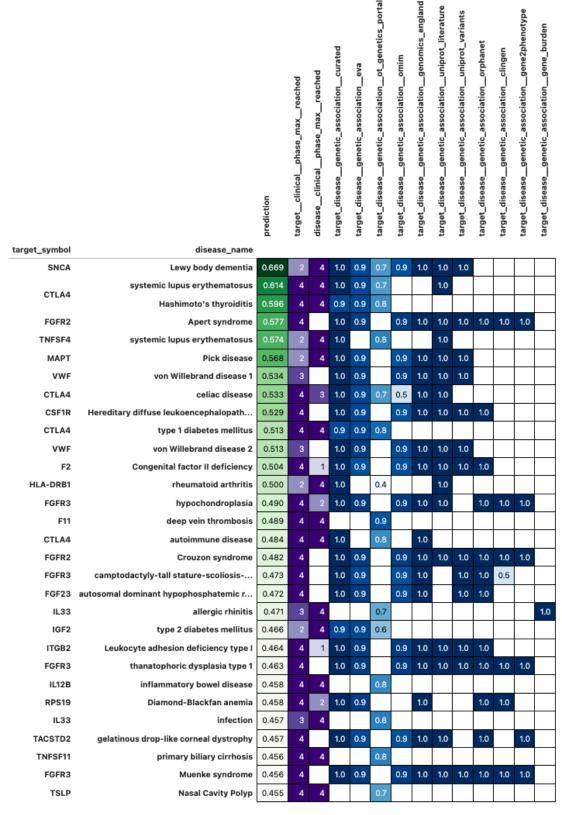


Figure 15: **Top ranked undeveloped, tractable target-disease pairs**. The highest scoring TD pairs per the RDG model that have not entered clinical trials despite having a target that has been in trials of an antibody-based drug. All values shown other than **prediction** are raw feature values, unweighted by RDG coefficients.

7.4 Sensitivity

In order to validate the stability of our findings in Section 2.4, we repeat this analysis across 18 different configurations listed in Supplementary Table 3. This includes 3 separate versions of Open Targets, 3 choices for the year defining the split between training and evaluation data and 2 choices for the length of the minimum advancement window (in years).

We find that the mean RR values from the RDG model consistently exceed the OTS model in all configurations among the very highest rankings (N=10) and also exceed the OTS model in all configurations except for 1 for N between 20 and 60. This data is shown in Supplementary Figure 16. The significance of these differences drops notably after N=40, which can be seen in the distribution of p-values from a Wilcoxon signed-rank test shown in Supplementary Figure 17.

Table 3: Configurations for sensitivity analysis

	open_targets_version	max_training_year	min_time_to_advancement_years
1	23.09	2017	4
2	23.12	2017	2
3	23.09	2015	2
4	23.12	2015	2
5	23.09	2017	2
6	23.06	2015	4
7	23.06	2017	2
8	23.06	2013	2
9	23.06	2015	2
10	23.06	2017	4
11	23.12	2013	4
12	23.09	2015	4
13	23.09	2013	2
14	23.12	2013	2
15	23.09	2013	4
16	23.12	2017	4
17	23.12	2015	4
18	23.06	2013	4

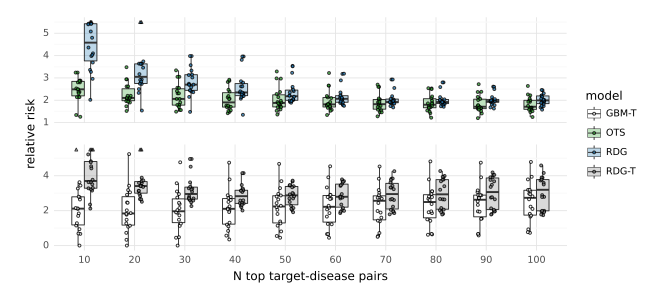


Figure 16: **Relative risk distributions across configurations in sensitivity analysis**. The distribution of the mean RR values displayed for a single configuration in Figure 2 is shown here across 18 configurations.

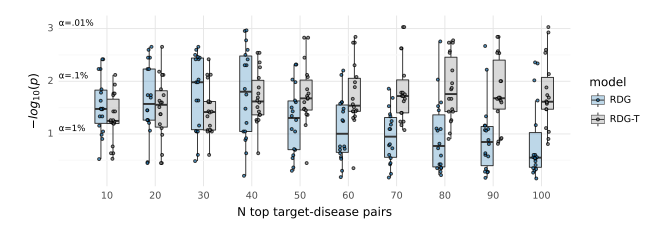


Figure 17: **P-value distributions across configurations in sensitivity analysis**. The distribution of the p-values displayed for a single configuration in Supplementary Figure 12 is shown here across 18 configurations.

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