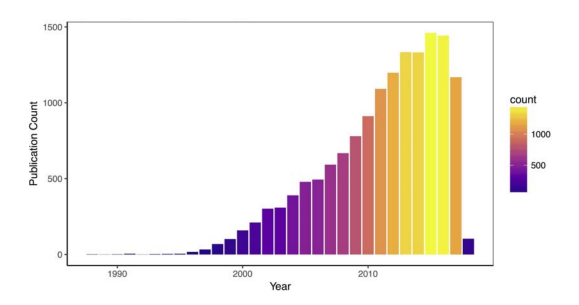
1 2	Title
- 3 4	Probing the chemical-biological relationship space with the Drug Target Explorer
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12	
13	Abstract
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15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31	Modern phenotypic high-throughput screens (HTS) present several challenges including identifying the target(s) that mediate the effect seen in the screen, characterizing 'hits' with a polypharmacologic target profile, and contextualizing screen data within the large potential space of drugs and biological screening model combinations. To address these challenges, we developed an interactive web application that enables exploration of the chemical-biological interaction space. Compound-target interaction data from public resources were quantified for over 280,000 molecules. Each molecule was annotated with a name and chemical structure, and every target was annotated with gene identifiers. The Drug-Target Explorer allows users to query molecules within this database of experimentally-derived and curated compound-target interactions and identify structurally similar molecules. It also enables network-based visualizations of the compound-target interaction space, and incorporates comparisons to publicly-available <i>in vitro</i> HTS datasets. Users can also identify compounds given one or more targets of interest. The Drug Target Explorer is a multifunctional platform for exploring chemical space as it relates to biological targets, and may be useful at several steps along the drug development pipeline including target discovery, structure-activity relationship, and lead compound identification studies.
 32 33 34 35 36 37 38 39 40 41 42 43 44 	Keywords Drug targets, polypharmacology, webapp, phenotypic drug screen, compound-target network

45 Introduction

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47 In the modern drug discovery and development process, high-throughput screens (HTS) 48 of drugs have become a common and important step in the identification of novel treatments for 49 disease. In the past decade, studies describing or citing high throughput drug screening are 50 increasingly prevalent, topping 1000 per year for the past 5 years (Figure 1) and span many 51 disease domains such as cancer, neurodegenerative disease, and cardiopulmonary diseases. 52 These screens are often phenotypic in nature whereby a large panel of compounds of known, 53 presumed known, and/or unknown mechanisms of action are tested in a biological model of 54 interest and generate phenotypic readouts such as apoptosis or proliferation. While these types 55 of screens facilitate the rapid identification of biologically active drugs or chemical probes, they 56 also present several challenges.



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58 **Figure 1 - High throughput drug screening is an increasingly common experimental** 59 **approach.** Yearly count of Pubmed-indexed publications that appear with the search term "high 60 throughput drug screening." Search performed on January 30, 2018.

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62 One prevailing challenge is the identification of the specific biological mechanisms within a cell that determine the response in a screen. The search for novel drugs constantly pushes 63 64 the pharmaceutical researchers to include novel chemical sets in phenotypic screens, with the 65 caveat that the underlying mechanism of action (MoA) of a particular compound cannot usually 66 be gleaned from the phenotypic screens. (1) Most of the time, identifying the MoA requires additional experimentation, particularly if the molecule represents a novel or understudied 67 68 chemical entity. Another challenge is that the polypharmacologic nature of many small 69 molecules can make it difficult to interpret HTS results as a given drug may affect multiple 70 targets with a range of efficacy. This, in turn, presents the difficulty of consolidating multiple 71 targets into a unified biological mechanism or set of mechanisms leading to poorly annotated 72 targets, misunderstood MoAs (2), and unknown or ambiguous off-targets with potential deadly side effects (3,4). A final challenge is that identification of related molecules and their targets is
not always straightforward; in the context of HTS analysis, structurally and functionally related
molecules that are not contained in a screening library might be useful to explore.

76 Multiple tools and databases have attempted to address various aspects of the 77 challenges outlined above (see Table 1). These tools allow the user to explore known 78 polypharmacology of small molecules. Many also allow users to explore compound-target 79 relationships by querying either by molecule or by target: DGIdb, DT-Web, BindingDB, 80 Polypharmacology Browser, STITCH, and SuperTarget allow users to identify MoAs/targets of a 81 given compound by evaluating a guery drug (5–10), while DT-Web, BindingDB, 82 Polypharmacology Browser, and STITCH allow users to search by chemical similarity using any 83 guery molecule (Table 1). Probe Miner, alternatively, is designed primarily to handle target-84 based queries (11). All tools listed in Table 1 allow users to identify molecules with known 85 polypharmacology, but only two, STITCH and SuperTarget, provide the ability to summarize 86 these targets into biological pathways/mechanisms using a gene list enrichment approach 87 (9.10). The final challenge - identifying structurally or functionally related molecules - is 88 addressed by DT-Web, BindingDB, Polypharmacology Browser, and STITCH (6-9).

89 While several of the tools listed address one or more of these challenges, there are 90 some gaps (Table 1). For example, ChEMBLSpace does not have a web interface and therefore requires installation on a compatible system before use (12). In addition, not all of these tools 91 92 are open-source (STITCH, SuperTarget, and BindingDB). An easy to modify open-source 93 application could enable users to create features that are helpful for their specific analyses. 94 While most tools allow both drug-based and target-based queries, none appear to facilitate 95 queries for molecules that affect several targets, which may be useful for users who want to 96 leverage polypharmacology by employing drugs that inhibit multiple biological mechanisms. 97 While multiple targets can be queried at one time in STITCH, it is not straightforward to identify 98 single molecules that affect all guery targets. In addition, DGIdb and ChEMBLSpace cannot be 99 used to explore similar chemical space to the query molecule. These two, plus SuperTarget, 100 also cannot be queried using molecules that are not in the database; a feature that might help 101 users with novel preclinical candidate drugs. With the exception of DT-Web and STITCH, these 102 tools do not allow visualization of drug-target networks, which may help users address the 103 challenge of identifying structurally or functionally related drugs. No tools other than STITCH 104 perform gene list enrichment, which may help users interpret the biological MoAs of 105 polypharmacologic molecules.

106 To address these gaps, we developed the Drug-Target Explorer. Specifically, the Drug-107 Target Explorer enables the user to (1) look up targets for individual molecules and groups of 108 molecules, (2) explore networks of targets and drugs, (3) perform gene list enrichment of targets 109 to assess target pathways of compounds, (4) compare query molecules to cancer cell line 110 screening datasets, and (5) discover bioactive molecules using a query target and exploration of 111 these networks. We anticipate that the users will include biologists and chemists involved in 112 drug discovery who are interested in performing hypothesis generation of human targets for 113 novel molecules, identifying off-targets for bioactive small molecules of interest, and exploring of 114 the polypharmacologic nature of small molecules.

	Drug-Target Explorer	Probe Miner	DGIdb v3.0	DT-Web	BindingDB	Polypharmacology Browser	STITCH	ChEMBLSpace	SuperTarget
Web app?	х	х	х	Х	х	х	Х		х
Open-source software?	х		х	X - underlying R package only	unknown	х		х	
Search by targets to find drugs?	х	х	х	х	х	X - only by PDB-listed ligands	х	х	х
Search by drugs to find targets?	х		х	х	х	х	х		х
Identification of molecules that are associated with multiple query targets?	х			unknown					
Drug structure input?	х			х	х	х	х		
Drug name/ID input?	х		х	х	х	х	х		х
Visualize drug-target networks?	х			X, with user provided drug-target networks	not currently functioning		х		
Identify chemically similar drugs?				X, with user provided drug-target networks	х	х	х		х
Allows queries using molecules not in database?	х			X, with user provided drug-target networks	х	х	х		
Target organism?		human	human	human	human and others	human and others	human and others	unknown	human and othe
Target space?	3.6k	2.2k	6.1k	3.8k	>7k	4.6k	9.6mil	unknown	>6k
Chemical space?	280k	400k	10k	4.4k	>642k	870k	500k	unknown	>196k
Quantitative interactions?	х	х		unknown	х	х	х	х	х
Qualitative interaction?	х	х	х	х			unknown		х
Explore polypharmacology?	х	х	х	Х	х	х	х	х	х
Polypharmacologic target enrichment?	х						х		х
Comparison of query molecule to HTS drug response datasets?	х								
Target prediction?				х	х	х			
Database access?	Open	Open	Open	Open	Open	Open	Full database requires license	unknown	unknown
Last known update	2018	2018	2018	2018	2018	2016	2016	2015	2012

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Table 1 - Summary of selected features/uses of databases and applications for exploring
 molecule-target relationships and their overlapping features with the Drug-Target
 Database. Related tools include Probe Miner (11), DGldb (5), DT-Web (6), BindingDB (7),
 Polypharmacology Browser (8), STITCH (9), ChEMBLSpace (12), and SuperTarget (10).

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123 Results

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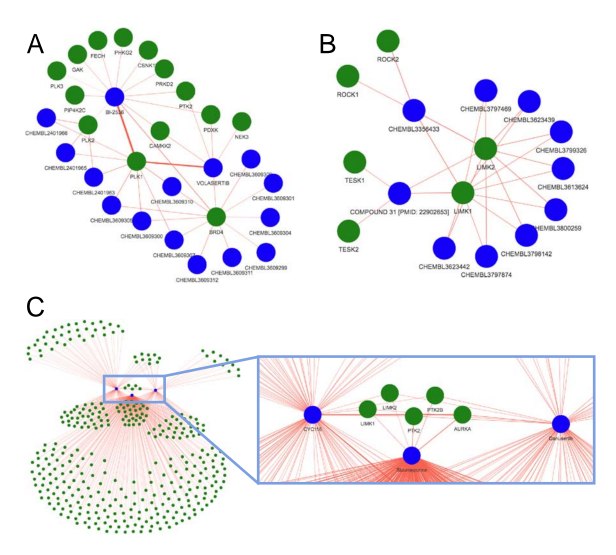
The Drug Target Explorer was designed to facilitate the following use-cases: hypothesis generation of targets for newly-discovered molecules, identification of off-targets for bioactive research molecules, and exploration of the polypharmacologic nature of many drugs. Below, we include vignettes highlighting how the Drug-Target Explorer can facilitate analysis in these areas.

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Identifying potential off-target effects of novel molecules

133 To highlight the use of this app to find potential off-targets of a novel molecule, we 134 queried the Drug-Target Explorer for C21, a recently-published Polo like kinase (PLK) inhibitor 135 that is not captured in our database (13). This molecule inhibits Plk2 and Plk1 in the low nM 136 range, and Plk3 in the low uM range (13). Using a Tanimoto similarity of 0.65 or greater, we 137 identified 14 molecules (Figure 2A, Supplemental Table 1). PLK1, PLK2, and PLK3 are known 138 targets of several of these molecules, such as BI 2536 and volasertib. Curiously, CAMKK, 139 BRD4, PDXK, and PTK2 are also targeted by molecules in this chemical set, with pChEMBL 140 values >6-8. A plausible hypothesis could be that these targets are affected by this family of 141 molecules, including the query molecule, in the 10-1000 nM range, which would indicate that 142 further research is needed to determine the selectivity of C21 or other structurally related 143 molecules.

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147 Figure 2 - Molecule-target networks highlight targets within chemical families. (A) 148 Using the novel Plk inhibitor C21 as a query with a Tanimoto cutoff of 0.65 (SMILES: 149 CCNC(=0)C1=CC2=C(C=C1)N(C=C2)C1=NC=C2N(C)C(=0)[C@@H](CC)N(C3CCCC3)C2=N1), we identify 14 related molecules (blue vertices), and observe several targets (green vertices) common to multiple 150 151 members of this family, including PLK1, PLK2, BRD4, CAMKK2, PTK2, and PDXK. (B) A gene-152 based query for two targets (green vertices), LIMK1 and LIMK2, identifies 10 molecules (blue 153 vertices), as well as other targets affected by these molecules. (C) A query for multiple targets relevant to tumors caused by neurofibromatosis type 2 identifies three promiscuous molecules 154 155 that have associations with these targets.

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Identifying off targets of existing molecules

This app may also be useful in identifying off-targets of existing molecules in a preclinical or exploratory research setting. In order to confidently interrogate the role of cellular targets, one must use compounds with specificity for those targets. A well-known example of a non-specific inhibitor is imatinib. This molecule, developed for use in the treatment of chronic myelogenous leukemia, was initially considered a selective inhibitor of Abl (14). More recently, several other targets have been identified for imatinib such as KIT, PDGFRA, and PDFGRB (15). Querying the Drug Target Explorer indicates that there is evidence for 61 targets of imatinib, several of which have pChEMBL values within a reasonable range of Abl, PDGFRA, and PDFGRB (Supplemental Table 2). These targets must all be considered when evaluating imatinib in human model systems.

169 A more recent example is the tool compound G-5555, a selective PAK1 inhibitor (16). 170 This compound has been used to demonstrate the role of PAK1 in cellular processes such as 171 invasion (17). A search of the Drug-Target Explorer database showed that this molecule not 172 only binds PAK1 (mean pChEMBL = 8.01), but there is qualitative evidence for effects on 173 PAK2/3, and guantitative evidence suggesting an effect on SIK2, MAP4K5, and PAK2 at similar 174 concentrations of G-5555 (mean pChEMBLs 8.05, 8, and 7.96 respectively, Table 2). G-5555 175 also may have an effect on STK family proteins (STK3, STK24, STK25, STK26) and LCK. 176 Therefore, any findings with G-5555 with regards to PAK1 inhibition must be validated with other 177 selective inhibitors or genetic approaches, as Jeannott and colleagues did (using other PAK 178 inhibitors such as FRAX597 and FRAX1036, as well as PAK1 silencing RNA), to confirm that 179 the effects observed are PAK1 specific (17).

180

Molecule Name	HGNC Symbol	Mean pChEMBL	n Quantitative	n Qualitative	KSI	Confidence
CHEMBL3770443	PAK1	8.01	3	1	0.1	11
CHEMBL3770443	PAK2	7.96	2	1	0.1	9.96
CHEMBL3770443	SIK2	8.05	1		0.1	9.05
CHEMBL3770443	MAP4K5	8	1		0.1	9
CHEMBL3770443	STK26	7.7	1		0.1	8.7
CHEMBL3770443	STK25	7.47	1		0.1	8.47
CHEMBL3770443	STK24	7.37	1		0.1	8.37
CHEMBL3770443	STK3	7.37	1		0.1	8.37
CHEMBL3770443	LCK	7.28	1		0.1	8.28
CHEMBL3770443	PAK3			1	0.1	1

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Table 2 – Targets of G-5555 found in the Drug-Target Explorer Database.

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Identifying polypharmacologically-targeted pathways and drugs with similar biological effects

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187 In order to provide biological context, this app allows the user to aggregate multiple 188 targets from compounds into functional categories. Using the previous example of G-5555, we 189 performed enrichment analysis on the list of targets to identify potential biological pathways and 190 MoAs that this molecule may disrupt. In doing so, we observed that G-5555 targets are enriched 191 in several Gene Ontology terms and KEGG Pathways like T-cell receptor signaling, Ras/MAPK 192 signaling, and Golgi-localized proteins (Supplemental Table 3). The app also allows the user to 193 compare the query molecule to drugs in the Cancer Cell Line/CTRP and GDSC/Sanger cell line 194 screening datasets. Specifically, the app identifies the most similar molecule available in these

195 datasets and uses that molecule as a reference to plot chemical similarity vs drug response 196 correlation.

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Finding a drug for known targets

200 Finally, the tool allows users to perform a reverse search, i.e. identify molecules that 201 have an association with a query target or targets and assess the known selectivity of these 202 molecules. For example, Petrilli et. al. identified LIM domain kinases as targets of interest in 203 tumors caused by the genetic disease neurofibromatosis type 2 (NF2) (18). They found that 204 pharmacologic (LIMK1/2 inhibitor BMS-5) and genetic modulation of LIMK1 and LIMK2 caused 205 cell-cycle inhibition and reduced viability in merlin (Nf2) deficient Schwann cells (18). In the 206 context of follow-up and validation studies, it may be beneficial to use alternate molecules that 207 target LIMK1/2 at the same or greater potency than BMS-5. We used the Drug-Target Explorer 208 to find molecules that target LIMK1 and LIMK2 (Supplemental Table 4, Figure 2B). For example, 209 BMS-5 (CHEMBL2141887 in the Drug-Target Explorer) has mean pChEMBLs of 7.33 and 7.07 210 for LIMK1 and LIMK2 respectively. A good alternative to validate the effects of this molecule 211 might be CHEMBL3623442, a relatively structurally distinct small molecule (extended fingerprint 212 Tanimoto similarity of 0.433 to BMS-5 in this database), with pChEMBLs of 9 and 8.52 for 213 LIMK1 and LIMK2 respectively. Another interesting possibility is the identification of multiple 214 molecules with overlapping desired targets and non-overlapping off-targets to reduce off-target 215 effects, or to identify synergistic/additive single-target, multi-drug combinations as outlined by 216 Fitzgerald et al 2006 (19). Using the above scenario with LIMK1/2, it may be possible to use 217 structurally distinct molecules in combination or in sequence, like CHEMBL3356433 and 218 Compound 31 highlighted in Figure 2B, to reduce off-target effects or inhibit LIMK1/2 in an 219 additive or synergistic manner. The opposite approach could also be taken by finding a single 220 molecule that binds multiple desired targets. In the case of merlin-deficient cells, focal adhesion 221 kinases (FAKs) such as PTK2 (FAK2) and PTK2B, as well as Aurora kinase A (AURKA) have 222 been highlighted as potential targets of interest (18,20,21). Using the Drug-Target Explorer, we 223 can identify molecules that target LIMK1/2, PTK2/2B, and AURKA (Supplemental Table 5, 224 Figure 2C). Using this information, a rational hypothesis might be that CYC116 or danusertib 225 could be effective and selective for NF2-deficient tumor cells; to our knowledge, the use of these 226 molecules in this setting has yet not been explored.

227 228

229 Discussion

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In the present study, we demonstrate that the Drug-Target Explorer enables the user to look up targets for novel and known molecules such as C21, G-5555, and imatinib, as well as explore networks of these drugs and their targets. Users can perform target enrichment to consolidate multiple targets to into pathways, compare query molecules to screening datasets, and identify bioactive molecules given a query target.

236 Several future directions are envisioned for this application. The code and database has 237 been designed in such a way that any database with structural information and drug-gene target 238 information (qualitative associations, or quantitative associations that can be coerced to

239 pChEMBL values) can be harmonized and integrated into the database. Therefore, as new 240 datasets become available, such as the recently-published Drug-Target Commons (22), they 241 can be integrated and released. We also envision occasional errors being identified as the 242 database is explored and vetted by users and have included a feedback form for users to 243 suggest new data to integrate, as well as to highlight necessary corrections to the dataset. 244 Currently, the guery molecule to full database similarity calculation is computationally intensive. 245 One solution to speed up calculation times may be to implement a locality sensitive hashing 246 method in future versions of the database and web app, such as the method devised by Cao et 247 al 2010 (23). An additional planned feature for this app is the implementation of a bulk 248 annotation feature to allow users to annotate HTS data with targets and/or putative targets of 249 identical or structurally related molecules. Finally, the integration of a predictive framework for 250 identifying targets of query drugs based on drug and target feature data would enable users to 251 quantitatively predict targets of novel molecular entities rather than manually exploring 252 structurally similar molecules.

The Drug-Target Explorer enables users to explore known molecule-human target relationships as they relate to chemical similarity rapidly and with minimal effort. We anticipate that users such as biologists and chemists using chemical probes or studying preclinical therapeutics will find this tool useful in several areas. Specifically, this tool may aid drug discovery efforts by accelerating hypothesis generation, simplifying the transition from phenotypic HTS results to mechanistic studies, and streamlining the identification of candidate molecules that target a protein or mechanism of interest.

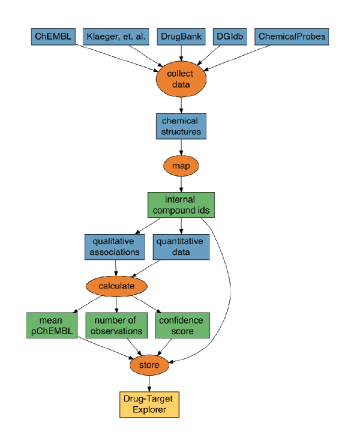
261 Methods

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263 To build the database of known compound-target interactions, we aggregated five data 264 sources containing qualitative and quantitative interactions (Figure 3). We considered qualitative 265 interactions to be curated compound-target associations with no associated numeric value. 266 Quantitative interactions were defined as compound-target information with a numeric value 267 indicating potency of compound-target binding or functional changes. Qualitative compound-268 target associations were retrieved from the DrugBank 5.0.11 XML database, the DGIdb v3.0.1 269 interactions.tsv file, and ChemicalProbes.org (acc. Jan 17 2018) (5,24,25). pChEMBL, IC50, 270 C50, EC50, AC50, Ki, Kd, and potency values for Homo sapiens targets were retrieved from the 271 ChEMBL v23 MySQL database (26). Kd values were also obtained from Klaeger et al 2017, in 272 which the authors determined the Kd of 244 kinase inhibitors against 343 kinases (27). For all 273 quantitative and qualitative data sources, compound structural information (SMILES) was 274 retrieved when available. When not available, it was batch annotated using the Pubchem 275 Identifier Exchange Service, or, in some cases, manually annotated via PubChem and 276 ChemSpider search (28,29).

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Figure 3 - Process for developing the Drug-Target Explorer. Molecule-target and chemical structure data were collected from public sources. In the case of DGIdb, chemical structures were assigned using the PubChem Chemical Identifier Exchange or manually assigned using ChemSpider and PubChem. Chemical structures were converted to circular fingerprints and the databases were mapped to internal Drug-Target Explorer identifiers. Qualitative and quantitative data were summarized by calculating several summary statistics, and these data were stored together with the internal identifiers to form the Drug-Target Explorer database.

287 To consolidate data for "identical" molecules within and across multiple databases, the 288 functional connectivity fingerprint (FCFP6)-like 'circular' fingerprint for each SMILES was 289 calculated using the R interface (rcdk) to the Java Chemical Development Kit (CDK) (30-32). 290 The package was modified to use the latest version of the CDK (2.1.1), which enables 291 perception of chiral centers, enabling differentiation between isomeric molecules. Each unique circular fingerprint and all external IDs and SMILES associated with that fingerprint were then 292 293 assigned an internal identifier, so that groups of molecules with identical fingerprints were 294 assigned to the same internal ID. The internal molecular IDs were then mapped to each 295 database to permit their aggregation. All datasets were combined and summaries were 296 generated for each compound-target comparison using functions from the R 'tidyverse' (33).

The summary metrics described in Table 3 were calculated. One of these metrics, pChEMBL, is used to convey the efficacy of a given molecule. It is calculated from one of

several semi-comparable values in the ChEMBL database, and is defined as the negative log 10 molar of the IC50, XC50, EC50, AC50, Ki, Kd, or potency (26). For example, a pChEMBL value of 7 would indicate that there is a measurable effect on a given target in the presence of 100 nM of molecule. To harmonize the data from Klaeger et al with ChEMBL data, the Kd values were converted to pChEMBLs. The mean pChEMBL was calculated for every moleculetarget combination, as well as the number of quantitative and qualitative associations found in the source databases.

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Metric	Unit	Meaning
mean IC50/AC50/EC50/C50/Potency/Ki/Kd	nM	mean of values obtained from quantitative datasets; available in database but not app
mean pChEMBL	-log10(nM)	mean -log10(nM) of all semi-comparable quantitative values
n_qualitative	count	number of qualitative associations identified
n_quantitative	count	number of quantitative associations identified
known selectivity score	N/A	1 divided by the number of known targets, lower is less selective
confidence score	N/A	mean pChEMBL, multiplied by n_quantitative, plus n_qualitative

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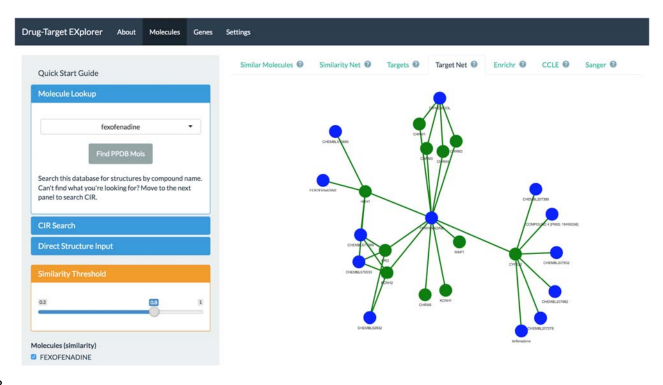
308Table 3 - Drug-target association metrics summarized in the Drug-Target Explorer309database.

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We calculated a known selectivity score for each molecule, which we defined as 1 divided by the total number of targets for that molecule (lower values correspond to lower molecule selectivity), and a confidence score for each molecule-target relationship, which we defined as the mean pChEMBL multiplied by the number of quantitative measurements, in addition to the number of qualitative annotations. A larger confidence score indicates greater confidence in this relationship; this confidence is weighted by the potency to give increased preference to high-potency compound-target interactions.

318 This resulted in a database containing 3645 human targets (represented by HUGO gene 319 symbols), ~280,000 small molecules, and ~623,500 molecule-target relationships summarized 320 from ~598,000 quantitative associations and ~25,000 qualitative associations. Finally, this 321 database as well as fingerprints and chemical aliases for each molecule were saved as R binary 322 files and stored on Synapse. All of the data, as well as snapshots of the source databases used 323 to build the Drug Target Explorer database (with the exception of DrugBank, which requires a 324 license to access) are accessible at <u>www.synapse.org/dtexplorer</u>. The Drug-Target Database is 325 licensed under CC BY-SA 4.0.

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Figure 4 - **Layout of the Drug-Target Explorer.** The "About" tab describes the apps functions and uses, the "Molecules" tab permits molecule-based searching, the "Genes" tab permits target queries, and the "Settings" tab allows the user to pick the fingerprinting method used.

339 We developed a Shiny application to permit exploration of the database (34,35). For 340 chemical queries, users can search for molecules in the database by one of three methods: 341 from a list of aliases obtained from the source databases, retrieving the chemical structure using 342 the 'webchem' interface to the Chemical Identifier Resolver, or by directly inputting the SMILES 343 string (36). A Tanimoto similarity threshold allows the user to narrow or widen the chemical 344 space of the results. After querying, the input molecule is converted to a fingerprint and it's 345 similarity calculated relative to all molecules in the database, using 'extended' fingerprints. The 346 user then can view the resulting set of molecules as well as the molecule-target relationships in 347 interactive tables and graphs (Figure 4). In addition, the user can remove or include molecules 348 on an a-la-carte basis, view the 2D structural representation of the input molecule, and perform target list enrichment analysis (37,38). Furthermore, the query molecule can be compared 349 350 against molecules in the CTRP and Sanger cancer cell line drug-screening datasets to identify 351 identical or similar structures in these datasets, and compare the relationship between chemical 352 structure and correlations in drug response.

For target queries, users can input one or more query HUGO gene(s) and identify molecules that are reported to bind those targets, and view these data in an interactive table. Users can also view these drugs in an interactive graph format to view their association with the query target and their other targets. The Drug-Target Explorer is available at www.synapse.org/dtexplorer. The source code for the Drug-Target Explorer app is available at

358 https://github.com/Sage-Bionetworks/polypharmacology-db. The source code is licensed under 359 Apache 2.0. 360 361 362 Supplemental table legends: 363 364 Supplemental Table 1 – Targets of C21-like compounds in the Drug-Target Explorer 365 Database. 366 367 Supplemental Table 2 – Targets of imatinib in the Drug-Target Explorer Database. 368 369 Supplemental Table 3 – Target enrichment analysis of G-5555 highlights putative 370 mechanistic effects. G-5555 targets were enriched in multiple Gene Ontology terms and 371 KEGG pathways. 372 373 Supplemental Table 4 - Molecules targeting LIMK1/2. The database was queried for 374 molecules that may modulate LIMK1 and LIMK2; this analysis revealed a large set of putative 375 tool compounds. 376 377 Supplemental Table 5 – Identification of multi-kinase-targeting molecules for NF2. A query 378 of the database for molecules that target several kinases of interest in NF2 (AURKA, LIMK1/2, 379 PTK2/2B) identified 3 polypharmacologic compounds. 380 381 References 382 383 1. Wagner BK. The resurgence of phenotypic screening in drug discovery and development. 384 Expert Opin Drug Discov [Internet]. 2016 Feb 10 [cited 2018 Mar 13];11(2):121-5. 385 Available from: http://www.tandfonline.com/doi/full/10.1517/17460441.2016.1122589 386 2. Santos R, Ursu O, Gaulton A, Bento AP, Donadi RS, Bologa CG, et al. A comprehensive map of molecular drug targets. Nat Rev Drug Discov [Internet]. 2017 Jan 2 [cited 2018 387 388 Mar 26];16(1):19-34. Available from: http://www.nature.com/articles/nrd.2016.230 389 van Esbroeck ACM, Janssen APA, Cognetta AB, Ogasawara D, Shpak G, van der Kroeg 3. 390 M, et al. Activity-based protein profiling reveals off-target proteins of the FAAH inhibitor 391 BIA 10-2474. Science [Internet]. 2017 Jun 9 [cited 2018 Mar 26];356(6342):1084-7. 392 Available from: http://www.ncbi.nlm.nih.gov/pubmed/28596366 393 4. Roy M, Dumaine R, Brown AM. HERG, a primary human ventricular target of the 394 nonsedating antihistamine terfenadine. Circulation [Internet]. 1996 Aug 15 [cited 2018] 395 Mar 26];94(4):817–23. Available from: http://www.ncbi.nlm.nih.gov/pubmed/8772706 396 5. Cotto KC, Wagner AH, Feng Y-Y, Kiwala S, Coffman AC, Spies G, et al, DGldb 3.0: a 397 redesign and expansion of the drug-gene interaction database. Nucleic Acids Res 398 [Internet]. 2017 Nov 16 [cited 2018 Mar 20]; Available from: 399 http://academic.oup.com/nar/article/doi/10.1093/nar/gkx1143/4634012 400 Alaimo S, Bonnici V, Cancemi D, Ferro A, Giugno R, Pulvirenti A. DT-Web: a web-based 6. 401 application for drug-target interaction and drug combination prediction through domain-402 tuned network-based inference. BMC Syst Biol [Internet]. 2015 [cited 2018 Mar 403 20];9(Suppl 3):S4. Available from: http://www.ncbi.nlm.nih.gov/pubmed/26050742 404 7. Gilson MK, Liu T, Baitaluk M, Nicola G, Hwang L, Chong J. BindingDB in 2015: A public

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