GenUI: Interactive and Extensible

² Open Source Software Platform for

³ De Novo Molecular Generation

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19 Abstract

20 Computer-aided *de novo* drug design holds promise to significantly accelerate the drug 21 discovery process and bring down its costs. Thanks to this outlook, the field has thrived in the 22 past few years and has seen a surge of new method development due to the proliferation of

generative deep neural networks. However, the widespread adoption of new de novo drug 23 24 design techniques has been slow in fields like medicinal chemistry or chemical biology. Such 25 development is not surprising since in order to successfully integrate de novo drug design in 26 existing processes and pipelines, a close collaboration between diverse groups of 27 experimental and theoretical scientists needs to be established. Therefore, to accelerate the 28 adoption of both modern and traditional de novo molecular generators, we developed GenUI 29 (Generator User Interface), a software platform that makes it possible to integrate molecular 30 generators within a feature-rich graphical user interface that is easy to use by experts of 31 varying backgrounds. GenUI is implemented as a web service and its interfaces offer tools for 32 data preprocessing, model building, molecule generation, and interactive chemical space 33 visualization. Moreover, the platform is easy to extend with customizable frontend React.js 34 components and backend Python extensions. GenUI is open source and a recently developed 35 de novo molecular generator, DrugEx, was integrated as a proof of principle. In this work, we 36 present the architecture and implementation details of the GenUI platform and discuss how it 37 can facilitate collaboration in the disparate communities interested in *de novo* drug design and molecule generation. 38

39 Keywords

40 graphical user interface, de novo drug design, molecule generation, deep learning, web

41 application

43 Introduction

44 Due to significant technological advances in the past decades, the body of knowledge on the 45 effects and roles of small molecules in living organisms has grown tremendously [1, 2]. At present, we assume the number of entries across all databases to be in the range of hundreds 46 47 of millions or billions (108-109) [3-5] and a large portion of this data has also accumulated in 48 public databases such as ChEMBL [6, 7] or PubChem BioAssay [1]. Still, these numbers are rather small in comparison to 10³³, a recently reported estimation of the size of the drug like 49 chemical space [8]. However, it should be noted that numerous studies in the past reported 50 51 numbers both bigger and smaller depending on the definition used [8-11]. In addition, 52 considering that only 1-2 measured biological activities per compound are available [12], the 53 characterization of known compounds also needs to be expanded.

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55 For a long time *de novo* molecular design algorithms for systematic and rational exploration 56 of chemical space [13-15] and quantitative structure-activity relationship (QSAR) modeling [16] 57 have been considered as tools that could broaden our horizons with less experimental costs and without the need to exhaustively evaluate as many as 10³³ possible drug-like compounds 58 59 to find the few of interest. The relevance of QSAR modeling and *de novo* molecular design for 60 drug discovery has been discussed many times [13-21], but these approaches can be just as 61 useful in the areas of chemical biology that require new tool compounds and chemical probes 62 that might not be constrained to drug-like molecules only [22].

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Thanks to the constant growth of bioactivity databases and widespread utilization of graphical processing units (GPUs) the application of powerful data-driven approaches based on deep neural networks (DNNs) has grown substantially. DNNs found many use cases in molecular virtual screening and *de novo* compound generation (**Figure 1**) [19]. This rapidly evolving class of algorithms has been influencing modern drug discovery by building more accurate QSAR models [12, 23], creating better molecular representations [24-26], predicting 3D

- protein structure with impressive accuracy [27] or achieving other promising results in many
 medicinal and clinical applications [3, 12, 17, 21, 28-30].
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Figure 1 Schematic view of a typical cheminformatics workflow involving a DNN. First, a data set of compound structures and their measured activities on the desired target molecule (most often a protein) is compiled and encoded to suitable representation. Second, the encoded data is used as input of the neural network in forward mapping. A large number of architectures can be used with recurrent neural networks (RNNs) and convolutional neural networks (CNNs) as the most popular examples. Finally, the neural network is trained by backpropagating the error of a suitable loss function to adjust the activations inside the network so that the loss is minimized. Depending on the architecture, the network is trained either as a bioactivity predictor (e.g. a QSAR model) or as a molecular generator.

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In the field of *de novo* drug design, the most attractive feature of DNNs is their ability to probabilistically generate compound structures [13, 31]. DNNs are able to take non-trivial structure-activity patterns into account, thereby increasing the potential for scaffold hopping and the diversity of designed molecules [32, 33]. A large number of generators based on DNNs were developed recently demonstrating the ability of various network architectures to generate compounds of given properties (biological activity included) [13, 31, 34-37].

91 Even though deep learning has been dominating *de novo* drug design in the recent years, it

should be noted that the field also has a long history of evolutionary heuristic methods such

93 as genetic algorithms on the forefront [20]. These traditional methods are still being 94 investigated and developed [38-43] and it is yet to be established how they compare to the 95 novel approaches based on deep learning [13]. Due to the simpler nature of these traditional 96 approaches non-obvious relationships can be easily missed, which may affect the quality of 97 the suggested chemical structures. However, simplicity can also be an advantage since 98 interpretation of simpler methods is easier. This is especially problematic for deep learning 99 models that can have more than thousands of parameters [44]. Moreover, a simpler method 100 requires less training data [38] without sacrificing chemical space coverage [45].

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One of the open questions for both traditional and deep learning molecular generators is also how they should be benchmarked, compared and interpreted [40]. Therefore, benchmarking studies of *de novo* drug design approaches are also the subject of ongoing research [46-48] and much needed to ensure that these methods have conclusive real impact on new ligand discovery [49, 50]. However, the ultimate test of a *de novo* drug design method should always be prospective application in real projects with experimental validation of the generated molecules.

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110 Although *de novo* molecular design algorithms have been in development for multiple 111 decades [51] and experimentally validated active compounds have been proposed [18, 52-112 59], these success stories are still far away from the envisaged performance of the 'robot 113 scientist' [60-62]. Successful development of a completely automated and sufficiently accurate 114 process has been elusive and hindered mostly by the computational expense and poor 115 synthetic availability of the generated compounds [18]. Despite increasing efforts to automate 116 the scientific process of decision making [18, 63-65], human insight and manual labor are still 117 necessary to further refine the compounds generated by *de novo* molecular design algorithms. 118 In particular, human intervention is of utmost importance in the process of compound scoring 119 in which best candidates are prioritized for synthesis and experimental validation [18, 65].

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121 Though many *in silico* compound generation and optimization tools are available for free [66]. 122 it is still an exception that these approaches are routinely used. The vast majority of methods 123 described in the literature serve only as a proof of concept. Hence, they lack a proper graphical 124 user interface (GUI) through which non-experts could easily access the algorithms and 125 analyze their inputs and outputs in a convenient way. Even if such a GUI exists, it is often 126 simplistic and intended to be used only with one particular method [41, 43, 67, 68]. Lack of 127 easy to explain and auditable information systems is a factor leading to some level of 128 disconnection between medicinal and computational chemists [69], which can hinder tighter 129 collaboration that can stand in the way of effective utilization of many promising de novo drug 130 design methods. Many molecular generators would also benefit from a comprehensive and 131 easy to use application programming interface (API) that would enable easier integration with 132 existing computational infrastructures. Recently a tool called Flame was presented that offers 133 many of the aforementioned features in the field of predictive QSAR modeling [70], but while 134 there are closed-source solutions like BRADSHAW [71] or Chemistry42 [72] to the best of our 135 knowledge there is no such solution in the realm of open source software for *de novo* drug 136 design. However, there has been effort to develop interactive databases of generated 137 structures as evidenced by the most recent example, cheML.io [73].

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139 In this work we present the development of GenUI, a software framework that provides 140 a general-purpose GUI for molecular generators and enables easy integration of such 141 algorithms with existing drug discovery pipelines as well. The GenUI framework integrates 142 solutions for import, generation, storage and retrieval of compounds, visualization of the 143 created molecular data sets and basic utilities for QSAR modeling. All features can be easily 144 accessed through the web-based GUI or REST API to ensure that both human users and 145 automated processes can interact with the application easily. Integration of new molecular 146 generators and other features is facilitated by a Python API and GUI customization is possible 147 via custom components implemented with the React.js JavaScript library. To demonstrate the 148 features of the GenUI framework, our recently published molecular generator DrugEx [74] was

integrated within the GenUI ecosystem. The source code of the GenUI platform is distributed
under the MIT open-source license [75-77] and several Docker [78-80] images are also
available online for quick deployment [81].

152 Implementation

153 Software Architecture

- 154 User interaction with GenUI happens through the frontend web client which issues REST API
- 155 calls to the backend, which comprises five services (Figure 2). However, advanced users may
- 156 also implement clients and automated processes that use the REST API directly.

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Figure 2 Schematic depiction of the GenUI platform. On the frontend (A), users interact with the web-based GUI to access the backend server services (B). All actions and data exchange are facilitated through REST API calls so that any automated process can also interact with GenUI. The backend application comprises five REST API services each of which has access to the data storage and task queue subsystems. The services can issue computationally intensive and long-running asynchronous tasks to backend workers to ensure sufficient responsiveness and scalability. In the current implementation, tasks can be submitted to two queues: (1) the default CPU queue, which handles all tasks by default, or (2) the GPU queue, intended for tasks that can be accelerated by the use of GPUs.

169 The five backend services form the core parts of GenUI and can be described as follows:

- "Projects" service handles user account management, authorization, and workflows. It
 is used to log users in and organize their work into projects.
- "Compounds" service manages the compound database including deposition,
 standardization, and retrieval of molecules and the associated data (i.e. bioactivities,
 physicochemical properties, or chemical identifiers).
- 175 3. "QSAR Models" service facilitates the training and use of QSAR models. They can be
 176 used to predict biological activities of the generated compounds, but they are also
 177 integral to training of many molecular generators.
- 4. "Generators" service is responsible for the integration of *de novo* molecular generators.
 It is meant to be used to set up and train generative algorithms whether they are based
 on traditional approaches or deep learning.
- 181 5. "Maps" service enables the creation of 2D chemical space visualizations and
 182 integration of dimensionality reduction algorithms.
- 183
- 184 In the following sections, the design and implementation of each part of the GenUI platform185 will be described in more detail.

186 Frontend

187 Graphical User Interface (GUI)

The GUI is implemented as a JavaScript application built on top of the React.js [82] web framework. The majority of graphical components is provided by the Vibe Dashboard opensource project [83], but the original collection of Vibe components was considerably expanded with custom components to fetch, send, and display data exchanged with the GenUI backend 192 REST API. In addition, frameworks Plotly.js [84], Charts.js [85] and ChemSpace.js [86] are
193 used to provide helpful interactive figures.

194

The GUI reflects the structure of the GenUI backend services (**Figure 2** and **Figure 3**). Each backend service (Projects, Compounds, QSAR, Generators, and Maps) is represented as a separate item in the navigation menu on the left side of the interface (**Figure 3**a). Upon clicking a menu item, the corresponding page opens rendering a grid of cards (**Figure 3**b) that represent the objects corresponding to the selected backend service. Various actions related to the particular service can be performed from the action menu in the top right of the interface (**Figure 3**c).

202 Projects

The "Projects" interface serves as a simple way to organize user workflows. For example, a project can encapsulate a workflow for the generation of novel ligands for one protein target (**Figure 3**). Each project contains imported compounds, QSAR models, molecular generators and chemical space maps. The number of projects per user is not limited and they can be deleted or created as needed.

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🔽 GenUl	< Projects	c) Action Menu Actions - A
Drojects		
S A2A Receptor	COX 1/2 Created: 3/22/2021 - 4:12:37 PM Last Update: 3/29/2021 - 7:30:05 PM	A2A Receptor Created: 7/28/2020 - 4:52:21 PM Last Update: 3/10/2021 - 2:12:15 PM
🗘 Compounds	Short description of the project	An example project where we use DrugEx to find new ligands for the Adenosine A2A receptor.
∽ QSAR Models	Onon Deleta	Onen Delate
Ø Generators ✓	open belete	open Delete
DrugEx	Create New Project	
🕅 Maps 🗸 🗸	Name	
Creator	New Project Name Description	
Explorer	Short description of the project	
a) Navigation Menu		
	Create & Open	
	· · · · · · · · · · · · · · · · · · ·	
		b) Card Grid
	Copyright \otimes 2019 Nice Dash, 2019-2020 Martin Šícho. All rights reserved.	

Figure 3 A screenshot showing part of the GenUI web GUI. In the figure, the GUI is in a state where the "A2A Receptor" project is already open so the menu on the left can be used to access its data. The GUI consists of three main parts: a) navigation menu, b) card grid and c) action menu. The navigation menu is used to browse data associated with various GenUI services ("Projects" in this case). If a link is clicked in the navigation menu, the data of the selected service is displayed as a grid of interactive cards. Each card allows the users to manage particular data items (a project in this case). The action menu in the top right is also updated depending on the service selected in the navigation menu and performs actions not related to a particular data item. In this case, the action menu was used to bring up the project creation form on the bottom left of the card grid.

- 218 Compounds
- Each project may contain any number of compound sets (Figure 4). Each set of compounds
- 220 can have a different purpose in the project and come from a different source. Therefore, the
- 221 contents of each card on the card grid depend on the type of compound set the card represents.
- 222 Compounds can be generated by generators, but also imported from SDF files, CSV files or
- 223 obtained directly from the ChEMBL database [6, 7]. New import filters can be easily added by
- 224 extending the Python backend and customizing the components of the React API accordingly
- 225 (see Python API and JavaScript API). For each compound in the compound set the interface
- can display its 2D representation (Figure 4), molecular identifiers (i.e. SMILES, InChI, and
- 227 InChIKey), reported and predicted activities (Figure 4) and physicochemical properties (i.e.
- 228 molecular weight, number of heavy atoms, number of aromatic rings, hydrogen bond donors,
- 229 hydrogen bond acceptors, logP and topological polar surface area).
- 230

🔽 GenUl	CHEMBL251							
Projects	Info Compounds Activities	Edit						
😂 A2A Receptor	Molecules in CHEMBL251	*						
O Compounds		Info Activities	Proper	ties				
小 QSAR Models	14							
Ø Generators ✓		All Ki F	(i_pChEMBL	Active	Probability			
DrugEx		TYPE	VALUE	UNITS	RELATION	ASSAY	TARGET	SOURCE
₩ Maps ✓ Creator		Ki	23.00	nM	-	CHEMBL644656	CHEMBL251	CHEMBL Activities (importe
Explorer		Ki_pChEMBL	7.64		-	CHEMBL644656	CHEMBL251	CHEMBL Activities (importe
		Active Probability	0.99					A2A Activ Predictio
		<						>

Figure 4 A screenshot showing part of the "Compounds" GUI. In this page, users can import data sets from various sources. A card representing an already imported data set from the ChEMBL database [7] is shown. The position and size of each displayed card can be modified by either dragging the card (reposition) or adjusting the bottom right corner (size change). The card shown is currently expanded over two rows of the card grid (Figure 3b) in order to accommodate the displayed data better. The "Activities" tab in the compound overview shows summary of the biological activity data associated with the compound. The activities are grouped by type and aside from experimentally determined activities the interface also displays activity predictions of available QSAR models. For example, in the view shown the "Active Probability" activity type is used to denote the output probability from a classification QSAR model. Each activity value also contains information about its origin (the "Source" column) so that it can be tracked back to its source.

- 242 QSAR Models
- 243 All QSAR models trained or imported in the given project are available from the "QSAR Models"

page (Figure 5, Figure 6). Each QSAR model is represented by a card with several tabs. The

- ²⁴⁵ "Info" tab contains model metadata, as well as a serialized model file to download (Figure 5).
- 246 The "Performance" tab lists various performance measures of the QSAR model obtained by
- cross-validation or on an independent hold out test set (Figure 6). The validation procedure
- can be adjusted by the user during model creation (Figure 5). Making predictions with the
- 249 model is possible under the "Predictions" tab. Each QSAR model can be used to make
- 250 predictions for any compound set listed on the "Compounds" page and the calculated
- 251 predictions will then become visible in that interface as well (Figure 4).
- 252

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n Projects										
S A2A Receptor	QSAR Training Set and You have to choose a	Activity Endpoin	t ou can choose any	compound set in the	^		i			
Compounds	current project.	СНЕ	MBL251 -			Info Perform	ance	Predictions		
 ✔ QSAR Models Ø Generators 	Choose the desired a table below. The cho as the output variabl	activity endpoint sen activity type e for the resulti	: by clicking the con from the given ac ng model.	rresponding row in the tivity set will be used		Description A QSAR model pro Tasks <mark>3 Comple</mark>	dicting	the Ki_PCHEMBL	. (-log(Ki)) value.	
DrugEx	ACTIVITY TYPE	DATA POINTS	MOLECULES	ACTIVITY SET		Model Data				
Creator				CHEMBL251		ITEM			VALUE	
Evolorer	Ki	6129	5135	Activities (imported)		Compound Set			CHEMBL251	
explorer	Ki_pChEMBL	5142	4257	CHEMBL251 Activities (imported)		Predictions Activ	Ki_pChEMBL	1BL		
				CHEMBL251		Training Settings				
	Inhibition	2227	2087	Activities (imported)		PARAMETER		VALUE		
				CHEMRI 251	~	Algorithm		RandomForest	t	
	Selected endpoint: Ki_ Activities (imported)	pChEMBL from C	HEMBL251	Choose Model Parameters		Delete				

Figure 5 A screenshot showing part of the "QSAR Models" GUI. The card on the left side of the screen shows how training data is chosen for a new model while the card on the right shows metadata about an already trained model.

🔽 GenUl	< QSAR Models		Create New 👻 🔺
Projects			
😂 A2A Receptor	A2A Ki_pChEMBL Regressor Created: 8/12/2020 - 11:49:18 AM Last Update: 8/25/2020 - 9:39:22 AM		A2A Activity Classifier Created: 728/2020 - 542:59 PM Last Update: 8725/2020 - 9:36:15 AM
	Info Performance Predictions		Info Performance Predictions
∽ QSAR Models	Independent Validation Set	a) Regression Metrics	Independent Validation Set b) Classification Metrics
	MSE 0.3382		ROC Curve (Independent Test Set) Baseline Independent Set
DrugEx	R2 0.7708		
🕅 Maps 🗸 🗸	Cross-Validation		2 07 8 06 8 05 9 04
	Metrics Summary		9 03 9 02 0.1
Explorer	MSE	R2	
	MIN 0.5746	0.2855	Metrics Summary
	MAX 0.9980	0.6571	MCC 0.7004
	AVG 0.7761	0.4301	ROC (AUC) 0.9381
	SD 0.1557	0.1170	·
			Delete
	Delete		

Figure 6 Performance evaluation view for a) regression and b) classification QSAR model. In a) the mean-squared error (MSE) and the coefficient of determination (R2) are used as validation metrics. In b) the performance is measured on a hold out independent validation test set with the Matthews correlation coefficient (MCC) and the area under the receiver operating characteristic (ROC) curve (AUC). The ROC curve itself is also displayed above the metrics.

262 New QSAR models are submitted for training with a creation card (Figure 5) that helps users 263 choose model hyperparameters and a suitable training strategy (i.e. the characteristics of the 264 independent hold out validation set, the number of cross-validation folds or the choice of 265 validation metrics). The "Info" tab of a trained model contains important metadata as well as 266 a hyperlink to export the model and save it as a reusable Python object. This import/export 267 feature enables users to archive and share their work, enhancing the reusability and 268 reproducibility of the developed models [87]. The "Performance" tab can be used to observe 269 model performance data according to the chosen validation scheme (Figure 6). This 270 information is different depending on the chosen model type (regression vs. classification, 271 Figure 6a vs. Figure 6b) and the parameters used (i.e. the choice of validation metrics). 272 Additional performance measures and machine learning algorithms can be integrated with the 273 backend Python API. Creation of such extensions does not even require editing of the GUI for 274 many standard algorithms (see Python API).

275 Generators

Under the "Generators" menu item, the users find a list of individual generators implemented in the GenUI framework (**Figure 7**). Currently, only the DrugEx generator [74] is available, but other generators can be added easily by extending the Python backend and customizing the existing React components. In fact, the GUI for DrugEx is based on the same React components as the "QSAR Models" view.

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283 Figure 7 A screenshot showing part of the "DrugEx" GUI with a model creation card with a) DrugEx training 284 285 parameters and b) performance overview of a trained DrugEx network. In a) the fields to define the compound set for the process of fine-tuning the 'parent' recurrent neural network trained on the ZINC data set [74] are shown. In 286 addition, the form provides fields to set the number of learning epochs, training batch size, frequency of 287 performance monitoring and size of the validation set. In b) the "Performance" tab tracks model performance. It 288 289 shows values of the loss function on the training set and validation set (top) and the SMILES error rate (bottom) at each step of the training process. The performance view is updated according to the chosen monitoring frequency 290 in real time as the model is being trained. Each model also has the "Info" tab which holds the same information as 291 for QSAR models.

292 Like QSAR models, DrugEx networks can also be serialized and saved as files. For example,

a cheminformatics researcher can build a DrugEx model outside of the GenUI ecosystem (i.e.

using a script published with the original paper [74]) and provide the created model files to

another researcher who can import and use the model from the GenUI web-based GUI.

296 Therefore, it is easy to share work and accommodate various groups of users in this way.

- 297 Maps
- 298 Interactive visualization of chemical space is available under the "Maps" menu item. The menu
- separates the creation of the chemical space visualization, the "Creator" page (Figure 8), and
- 300 its exploration, the "Explorer" page (**Figure 9**).

🔽 GenUl		<	Map Creator					Create New
Projects								
😂 A2A Receptor			Create New TSNE Model Model Name		Î	A2A ChEMBL Tested + Created: 3/10/2021 – 10:20: Last Update: 3/10/2021 – 10	DrugEx Suggestee 14 AM 0:20:14 AM	d
			New TSNE Model			Info		
			Write more about thi	s model if needed		Tasks 1 Complet	ted	
						Training Settin	gs	
DrugEx			Compound Sets	A2A Ligands from DrugEx	^	PARAMETER	VALUE	
🕅 Maps				CHEMBL203		Algorithm	TSNE	
Creator			Training Paramete	CHEMBL251	×	Parameters	n_iter=500;per early_exaggera	rplexity=30;early_exaggeration=12; ation_iter=250;exaggeration=0
Explorer			Mode			Mode	map	
			map		~	Training Set	CHEMBL251;A	2A Ligands from DrugEx
			Descriptor Sets	accriptor sets to use in the calculations		Descriptor Sets	MORGANFP	
			MORGANFP	scriptor sets to use in the calculations.	^	Validation Sett No validation data	ings available for this	s model.
						Model File		Auxiliary Files
		TSNE Parameters				TSNE_main_ma	iin.joblib.gz	chemspaceJSON_aux.json
			n_iter	500	۲			

Figure 8 The "Creator" interface of GenUI "Maps" page. On the left a form to create a new t-SNE [88] mapping of
 two sets of compounds using Morgan fingerprints is shown while information about an existing map can be seen
 on the right.





In this particular configuration, the shapes and colors of the points indicate the compound set to which the compounds belong to. The color scheme of points can be changed with the menu in the top left corner of the plot. It is possible to color points by biological activities, physicochemical properties and other data associated with the compounds. The same can also be done with the size of the points. The points drawn in the map are interactive and hovering over a point shows a box with information about the compound inside and on the right side of the map. Groups of points can also be selected by drawing a rectangle over them in which case a list of selected compounds is shown in the "Selection List" tab (Figure 10) and their bioactivity data is summarized under the "Selection Activities" tab (Figure 11).

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The "Creator" page is implemented as a grid of cards each of which represents an embedding of chemical compounds in 2D space (**Figure 8**). Implicitly, the GenUI platform enables t-SNE [88] embedding (provided by openTSNE [89]). However, new projection methods can be easily added to the backend through the GenUI Python API with no need to modify the GUI (see Python API) [90].

322

323 The purpose of the "Explorer" page is to interactively visualize chemical space embedding 324 prepared in the "Creator" (Figure 9). In the created visualization the users can explore 325 compound bioactivities, physicochemical properties, and other measurements for various 326 representations and parts of chemical space. Thanks to ChemSpace.js [86] up to 5 327 dimensions can be shown in the map at the same time: X and Y coordinates, point color, point 328 size and point shape. The map can be zoomed in by drawing a rectangle over a group of 329 points. Such points form a selection and their detailed information is then displayed under the 330 "Selected List" (Figure 10) and "Selected Activities" tabs (Figure 11).

🔽 GenUl	< Map Explorer								Actions 👻 📝
Projects	A2A ChEMBL Tested + DrugEx Suggested Select	tion List Selection Activiti	ies						
😂 A2A Receptor									
	CHEMBL251								
	1	Info Activities	Properties						
	° / -)	Ali Ki Ki_pC	hembl A	Active Proba	bility				
🛱 Maps 🗸 🗸		Ki	24.00	DM	RELATION	ASSAY	CHEMBI 251	CHEMBI 251 Activities (importe	d)
		Ki_pChEMBL	7.62		-	CHEMBL1018640	CHEMBL251	CHEMBL251 Activities (importe	d)
Explorer	<u>^</u>	Active Probability	0.96					A2A Activity Prediction	
		Info Activities	Properties						
		All Ki Ki_pC	hembl A	Active Proba	bility				
		TYPE	VALUE	UNITS	RELATION	ASSAY	TARGET	SOURCE	
		кі	1561.00	nM	-	CHEMBL1018640	CHEMBL251	CHEMBL251 Activities (imported)	
		KI_pChEMBL	5.81	-	-	CHEMBL1018640	CHEMBL251	CHEMBL251 Activities (imported)	
		Active	0.21					A2A Activity Prediction	

Figure 10 View of the "Selected List" tab of the "Explorer" page. The tab shows the selected molecules in the map as a list which is the same as the one used in the "Compounds" view (**Figure 4**). For easier navigation, the compounds are also grouped by the compound set they belong to and the view for each set can be accessed by switching tabs above the displayed list (only one compound set, CHEMBL251, is present in this case).

🔽 GenUl		AZA C	hEMBL Tested + Dr	ugEx Suggested	Selection List	Selection Activities				
Projects		1/1	Ki ochTMDI	Inhibition	Active Drobability					
😂 A2A Receptor		KI	KI_PCITEWIDE	minipition	Active Probability					
	_				Distribu	tions of Ki_pChEMBL				
								CHEMBL251		
	~		9.5					A2A Ligands from DrugEx	F	1
									Sel 1	
🕅 Maps	~		9							
			8.5							
Explorer										
			8							
			7.5						Properties	
									PROPERTY	VALUE
						6 8691 A	A Ligands		AMW	457.505
									NUMHEAVYATOMS	34
									NUMAROMATICRINGS	4
			. :	Ψ/					HBA	6
			°						HBD	2
									LOGP	7.1131
			2.5						TPSA	76.39

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Figure 11 View of the "Selection Activities" tab of the "Explorer" page. In this view, violin plots representing distributions of activities in the set of selected compounds are displayed. Each violin plot corresponds to one compound set and one activity type. The violin plots are also interactive and hovering over points updates the compound structure and its physicochemical properties are displayed on the right.

341 JavaScript API

- 342 Two main considerations in the development of GenUI are reusability and extensibility.
- 343 Therefore, the frontend GUI comprises a large library of over 50 React components that are

encapsulated in a standalone package (Figure 12). The package is organized into
subpackages that follow the structure and hierarchy of design elements in the GenUI interface.
In the following sections, we use the two most important groups of the React API components
as case studies to illustrate how the frontend GUI can be extended. The presented
components are "Model Components", used to add new trainable models, and the "REST API
Components", used to fetch and send data between the frontend and the GenUI REST API
services.



351

Figure 12 Schematic depiction of the GenUI React library which contains customized styles, utility functions and the components used in the GenUI web client. The "components" subpackage organizes the components into groups related to the structure of the GenUI interface. For example, components filed under the "models" subpackage are used in the creation of the "QSAR Models" (Figure 5), "DrugEx" (Figure 7) and "Maps" (Figure 8) interfaces while components under the "compounds" subpackage are used to implement the "Compounds" view (Figure 4). General purpose components (i.e. the card grid or the card tab widget) are in the root of the "components" subpackage.

359 Model Components

Much of the functionality of the GenUI platform is based on trained models. The "QSAR Models", "DrugEx" and "Maps" pages all borrow from the same library of reusable GenUI React components (**Figure 12**). At the core of the "models" component library (**Figure 12**) is the *ModelsPage* component (**Figure 13**). *ModelsPage* manages the layout and data displayed in model cards. When the users select to build a new model, the *ModelsPage* component is also responsible to show a card with the model creation form. The information that the *ModelsPage* displays can be customized through various React properties (**Figure 13**) that 367 represent either data (data properties) or other components (component properties). Such an 368 encapsulation approach and top-down data flow is one of the main strengths of the React 369 framework. This design is very robust since it fosters appropriate separation of concerns by 370 their encapsulation inside more and more specialized components. This makes the code easy 371 to reuse and maintain.



372

Figure 13 A simplified illustration of the high-level components in the GenUI React API for rendering model cards. The main *ModelsPage* component has two kinds of attributes (called "properties" in React): a) *data properties* and b) *component properties*. The values of data properties are used to display model data while the values of component properties are used as child components and injected into the GUI at appropriate places. If no component property is specified, default components are used as children instead (i.e. *ModelCard* and *NewModelCard*). The child components can accept data and component properties as well from their parent (i.e. *ModelsPage*). This creates a hierarchy of reusable components that can be easily assembled and configured to accommodate the different needs of each model view in a standardized and consistent manner.

381 REST API Components

382 Because the GUI often needs to fetch data from the backend server, several React 383 components were defined for that purpose. In order to use them, one just needs to provide 384 the required REST API URLs as React component properties. For example, the 385 ComponentWithResources component configured with the '/maps/algorithms/' URL will get all 386 available embedding methods as JSON and converts the result to a JavaScript object. Many 387 components can also periodically update the fetched data, which is useful for tracking information in real time. For paginated data there is also the ApiResourcePaginator 388 389 component that only fetches a new page if a given event is fired (i.e. user presses a button).

This makes it convenient to create GUIs for larger data sets. In addition, user credentials arealso handled automatically.

392

Many more specialized components are also available to fetch specific information. For example, the *TaskAwareComponent* tracks URLs associated with background asynchronous tasks and it regularly passes information about completed, running, or failed tasks to its child components. However, other specialized components exist that automatically fetch and format pictures of molecules, bioactivities, physicochemical properties or create, update and delete objects in the UI and the server [76].

399 Backend

400 The backend services are the core of the GenUI platform and the GenUI Python API provides 401 a convenient way to write backend extensions (i.e. new molecular generators, compound 402 import filters, machine learning algorithms for QSAR modeling, and dimensionality reduction 403 methods for chemical space maps). All five backend services (Figure 2) are implemented with 404 the Django web framework [91] and Django REST Framework [92]. For data storage, a freely 405 available Docker [80] image developed by Informatics Matters Ltd. [93] is used. The Docker 406 image contains an instance of the PostgreSQL database system with integrated database 407 cartridge from the RDKit cheminformatics framework [94]. The integration of RDKit with the 408 Django web framework is handled with the Django RDKit library [95]. All compounds imported 409 in the database are automatically standardized with the current version of the ChEMBL 410 structure curation pipeline [96].

411

412 Because the backend services also handle processing of long-running and computationally 413 intensive tasks, the framework uses Celery distributed task queue [97] with Redis as 414 a message broker [98] to dispatch them to workers. Celery workers are processes running in 415 the background that consume tasks from the task queue and process them asynchronously.

416 Workers can either run on the same machine as the backend services or they can be 417 distributed over an infrastructure of computers (see Deployment).

418 Python API

419 The GenUI backend codebase [77] is divided into multiple Python packages that each 420 encapsulate a part of the GenUI Django project (Figure 14). Any package that resides in the 421 root directory is referred to as the root package. Root packages facilitate many of the REST 422 API endpoints (Figure 2), but they also contain reusable classes that are intended to be 423 extended by extensions (see Generic Views and Viewsets, for example). In the following 424 sections, some important features of the backend Python API are briefly highlighted. However, 425 a much more detailed description with code examples is available on the documentation page 426 of the project [90].



Figure 14 Schematic depiction of the GenUI backend Python code. The backend is formed by a single Django project which is designated by its *settings* package and the *urls* and *wsgi* modules. The GenUI code itself is divided into a number of *root packages*. Each root package has a predefined structure with the code of the package organized in its own modules and packages. Each root package of the GenUI framework also has the *extensions* subpackage, which is a collection of extension modules. GenUI extensions and packages can also define the *genuisetup* module, which is used to automatically configure the individual package or extension.

- 435 Extensions
- 436 Just like in the case of the GenUI React API, modularity and extensibility were also the main
- 437 concerns during the design of the GenUI backend services. Each of the aforementioned root
- 438 packages contains a Python package called *extensions* (Figure 14). The *extensions* package
- 439 can contain any number of Django applications or Python modules, which ensures that the
- 440 extending components of the GenUI framework are well-organized and loosely coupled.
- 441
- 442 Provided that GenUI extensions are structured a certain way they can take advantage of
- 443 automatic configuration and integration (see Automatic Code Discovery). Before the Django

444 project is deployed, GenUI applications and extensions are detected and configured with the 445 *genuisetup* command, which makes sure that the associated REST API endpoints are 446 exposed under the correct URLs. The *genuisetup* command is executed with the *manage.py* 447 script (a utility script provided by the Django library).

448 Automatic Code Discovery

The root packages of the GenUI backend library define many abstract and generic base classes to implement and reuse in extensions. These classes either implement the REST API or define code to be run on the worker nodes inside Celery tasks. Automatic code discovery uses several introspection functions and methods to find the derived classes of the base classes found in the root packages. By default, this is done when the *genuisetup* command is executed (see Extensions).

455

456 For example, if the derived class defines a new machine learning algorithm to be used in 457 QSAR modelling, automatic code discovery utilities make sure that the new algorithm appears 458 as a choice in the QSAR modelling REST API and that proper parameter values are collected 459 via the endpoint to create the model. Moreover, all changes also get automatically propagated to the web-based GUI because it uses the REST API to obtain algorithm choices for the model 460 461 creation form. Thus, no JavaScript code has to be written to integrate a new machine learning 462 algorithm. These mechanisms are also used when adding molecular generators, 463 dimensionality reduction methods, or molecular descriptors.

464 Generic Views and Viewsets

When developing REST API services with the Django REST Framework, a common practice is using generic views and sets of views (called viewsets). In Django applications, views are functions or classes that handle incoming HTTP requests. Viewsets are classes defined by the Django REST Framework that bring functionality of several views (such as creation, update or deletion of objects) into one single class. Generic views and viewsets are then

470 classes that usually do not stand on their own, but are designed to be further extended and471 customized.

472

473 The GenUI Python library embraces this philosophy and many REST API endpoints are 474 encapsulated in generic views or viewsets. This ensures that the functionality can be reused 475 and that no code needs to be written twice, as stated by the well-known DRY ("Don't Repeat 476 Yourself") principle [99]. An example of such a generic approach is the *ModelViewSet* class 477 that handles the endpoints for retrieval and training of machine learning models. This viewset 478 is used by the gsar and maps applications, but also by the DrugEx extension. All these 479 applications depend on some form of a machine learning model so they can take advantage 480 of this interface, which automatically checks the validity of user inputs and sends model 481 training jobs to the task queue.

482 Asynchronous Tasks

483 Many of the GenUI backend services take advantage of asynchronous tasks which are 484 functions executed in the background without blocking the main application. Moreover, tasks 485 do not even have to be executed on the same machine as the caller of the task, which allows 486 for a great deal of flexibility and scalability (see Deployment).

487

488 The Celery task queue [97] makes creating asynchronous tasks as easy as defining a Python 489 function [100]. In addition, some GenUI views already define their own tasks and no explicit 490 task definition is needed in the derived views of the extensions. For example, the compounds 491 root package defines a generic viewset that can be used to create and manage compound 492 sets. The import and creation of compounds belonging to a new compound set is handled by 493 implementing a separate initializer class, which is passed to the appropriate generic viewset 494 class [90]. The initialization of a compound set can take a long time or may fail and, thus, 495 should be executed asynchronously. Therefore, the viewset of the *compounds* application

automatically executes the methods of the initializer class asynchronously with the help of anavailable Celery worker.

498 Deployment

499 Docker Images

Since the GenUI platform consists of several components with many dependencies and spans 500 501 multiple programming languages, it can be tedious to set up the whole project on a new system. 502 Docker makes deployment of larger projects like this easier by encapsulating different parts 503 of the deployment environment inside Docker images [78-80]. Docker images are simply 504 downloaded and deployed on the target system without the need to install any other tools 505 beside Docker. GenUI uses many official Docker images available on the Docker image 506 sharing platform Docker Hub [101]. The PostgreSQL database with built-in RDKit cartridge 507 [93], Redis [102] and the NGINX web server [103, 104] are all obtained by this standard 508 channel. In addition, we defined the following images to support the deployment of the GenUI 509 platform itself [81]:

- 510
- 511 1. *genui-main*: Used to deploy both the frontend web application and the backend 512 services.

513 2. *genui-worker*. Deploys a basic Celery worker without GPU support.

- *genui-gpuworker*. Deploys a Celery worker with GPU support. It is the same as the
 genui-worker, but it has the NVIDIA CUDA Toolkit already installed.
- 516

517 The tools to build these images are freely available [81]. Therefore, developers can create 518 images for extended versions of the GenUI that fit the needs of their organizations. In addition, 519 the separation of the main application (*genui-main*) from workers also allows distributed 520 deployment over multiple machines, which opens up the possibility to create a scalable 521 architecture that can quickly accommodate teams of varying sizes.

522 Future Directions

523 Although the GenUI framework already implements much of the functionality needed to 524 successfully integrate most molecular generators, there are still many aspects of the 525 framework that can be improved. For instance, it would be beneficial if more sources of 526 molecular structures and bioactivity information are integrated in the platform besides 527 ChEMBL (i.e PubChem [105], ZINC [106], DrugBank [107], BindingDB [108] or Probes and Drugs [109]). Currently, GenUI also lacks features to perform effective similarity and 528 529 substructure searches, which we see as a crucial next step to improve the appeal of the 530 platform to medicinal chemists. The current version of GenUI would also benefit from 531 extending the sets of descriptors, QSAR machine learning algorithms and chemical space 532 projections since the performance of different methods can vary across data sets. Finally, the 533 question of synthesizability of the generated structures should also be addressed and 534 a system for predicting chemical reactions and retrosynthetic pathways could also be very 535 useful to medicinal chemists if integrated in the GUI (i.e. by facilitating connection to a service 536 such as the IBM RXN [110] or PostEra Manifold [111]).

537

538 Even though it is hard to determine the requirements of every project where molecular 539 generators might be applied, many of the aforementioned features and improvements can be 540 readily implemented with the GenUI React components (see JavaScript API) and the Python 541 API (see Python API). In fact, the already presented extensions and the DrugEx interface are 542 useful case studies that can be used as templates for integration of many other 543 cheminformatics tools and de novo molecular generators. Therefore, we see GenUI as 544 a flexible and scalable framework that can be used by organizations to quickly integrate tools 545 and data the way it suits their needs the most. However, we would also like GenUI to become 546 a new useful way to share the progress in the development of novel de novo drug design 547 methods and other cheminformatics approaches in the public domain.

548 Conclusions

549 We implemented a full stack solution for integration of de novo molecular generation techniques in a multidisciplinary work environment. The proposed GenUI software platform 550 551 provides a GUI designed to be easily understood by experts outside the cheminformatics 552 domain, but it also offers a feature-rich REST API for programmatic access and 553 straightforward integration with automated processes. The presented solution also provides 554 extensive Python and JavaScript extension APIs for easy integration of new molecular 555 generators and other cheminformatics tools. We envision that the field of molecular generation 556 will likely expand in the future and that an open source software platform such as this one is 557 a crucial step towards more widespread adoption of novel algorithms in drug discovery and 558 related research. We also believe that GenUI can facilitate more engagement between 559 different groups of users and inspire new directions in the field of *de novo* drug design. 560

561 **Declarations**

562 Authors' Contributions

563 GvW suggested the original idea of developing a graphical user interface for a molecular 564 generator and supervised the study along with DS. MŠ extended the original idea and 565 developed all software presented in this work. XL is the author of DrugEx and helped with its 566 integration as a proof of concept. MŠ and XL also prepared the manuscript, which all authors 567 proofread and agreed on.

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573 Competing Interests

574 The authors declare that they have no competing interests.

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578 Availability of Data and Materials

579 The complete GenUI codebase and documentation is distributed under the MIT license and

- 580 located in three repositories publicly accessible on GitHub:
- <u>https://github.com/martin-sicho/genui</u> (backend Python code)
- <u>https://github.com/martin-sicho/genui-gui</u> (frontend React application)
- <u>https://github.com/martin-sicho/genui-docker</u> (Docker files and deployment scripts)
- 584 A reference application that was described in this manuscript can be deployed with Docker
- 585 images that were uploaded to Docker Hub: <u>https://hub.docker.com/u/sichom</u>. However, the

586 images can also be built with the available Docker files and scripts (archived at

- 587 <u>https://doi.org/10.5281/zenodo.4813625</u>). The reference web application uses the following
- 588 versions of the GenUI software:
- 0.0.0-alpha.1 for the frontend React application (archived at
- 590 <u>https://doi.org/10.5281/zenodo.4813608</u>)
- 0.0.0.alpha1 for the backend Python application (archived at
- 592 <u>https://doi.org/10.5281/zenodo.4813586</u>)

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