Thesis proposal An alpha version of the draft

Nil Adell Mill Winter 2020 Institute of Neuroinformatics

1 Introduction/Background

- 2 Drug discovery—the process by which a potential new medicine is identified—is a complex process that
- encompases the intersection of several fields (such as biology, statistics, chemistry or pharmacology). The
- 4 entire process is a long and costly endeavour, with a typical time-frame of 10 to 20 years till maket release and
- an estimated cost between 1 and 2 billion USD. With just a small quantity of the initially identified compounds
- 6 actually becoming an approved medicine. Many of these dropouts happening at the early stages of the entire
- 7 pipeline.
- 8 It exists, then, a need for better mechanisms for detecting better candidates. One of the most promising
- 9 directions is to improve the *in-silico* methods—computational simulations are relatively cheap and quick
- run that makes them an interesting solution. *In-silico* simulations then cover two main aspects: modelling
- the dynamics of the human body—such that any effect relevant to the durg or the disease will be captured
- by it—and methods to generate good candidates that are effective at exploring the vast space of possible
- 13 compounds.
- 14 Among the different computational approaches that have been used in the process of drug discovery deep
- learning (DL) has shown signs to be a potential game changer [Dargan et al., 2019]. DL has been able to
- capitalize on the exponential growth of data and the higher availability of computational resources. For examle,
- DL has had a remarkable success on computer vision (CV) and natural language processing (NLP), and has
- become the go-to solution for any problem in these two fields. It is, at the same time, penetrating into other
- fields, drug-discovery being one of them [Chen et al., 2018].
- 20 When we deal with this biological and molecular data, it exists a challenge on how to deal with the intrinsic
- 21 structure of the data. If we look at the case of deep learning for CV, where we deal with images, a key element
- 22 of any architecture for it's success was the use of convolutional layers—one will mostly observe convolutional
- 23 neural networks (CNNs) when analizying the state of the art in CV—which introduce a structural a prior based
- on the structure of the data. A similar case can be made for NLP. For that reason, there exists a strong signal
- of the structure of the data. A similar case can be made for 1421. For that reason, there exists a strong signa
- to look for models that can leverage the structural equivalent when in molecule or protein data, i.e. leverage
- 26 graph structures [Wu et al., 2019]. Sign [Sign? need to rewrite that] of that is the recent advancements in that
- direction [Sun et al., 2019].
- 28 Another of the big challenges is to unify all the aspects of drug-discovery and be able to incorporate all the
- 29 rellevant biological information when designing possible candidate molecules. An initial success story on
- that line is a recently paper [Zhavoronkov et al., 2019] where the authors describe a deep learning method by
- 31 which they are able to discover inhibitors of discoidin domain receptor 1 (DDR1)—a kinase implicated in
- 32 fibrosis—in just 21 days.
- 33 Those promising results, albeit encouraging, are just the tip of the iceberg. There is still a long way till a model
- 34 can satisfactorily capture the biological complexity of any arbitrary target and produce promising candidates.
- 35 On top of that, there is an added dimension, as such model should account for the variability from patient to
- patient and be able to generate a molecule that accommodates for all the genotipic and phenotipic variants, or
- generate different candidates for each of the genetic populations of interest. [need a ref here]
- 38 [I am not completely sure about this paragraph but I leave it here so I don't forget for now] Even more, in the
- 39 case of diseases like cancer, an heterogeneous population may appear within a single patient. So the same
- 40 variant effects arise inside a dynamic ecosystem, where a drug that just targets a subpopulation may lead to an

- evolutionary pressure complicating further the treatment outlook [reference paper of evolutionary perspective
- to cancerl. 42
- There is then a great need to develop models that can be conditioned based on a large set of biological 43
- [conditions?] and meaningfully account for this variations when generating a compound or/and evaluating a 44
- compunds effect when administered.
- In fact it is of interest to develop multi-scale models that capture system complexity at the different levels.
- For instance, a model that is able to learn protein-compound interactions—commonly known as the docking
- problem—while at the same time use this information to predict effects of the introduction of the compound 48
- on the larger protein-protein interaction (PPI) network.

Aim & Methods

- [Should I separate em in two different sections?] 51
- The aim of this thesis will be two fold. One the one side, analyze how the explicit use of graph convolutional
- neural networks (GCNNs) may open new oportunities when dealing with biological and checmical data. On 53
- the other side, explore how modelling the biology at different levels (e.g. molecular structure v.s. molecular 54
- interacton network [okay here I need to develop further about PPI, maybe mention NetBite (as Jannis referenced
- in the mail)]) may help with our understanding [of the biology? of compounds interaction?] and help generate
- better models. Furthermore, evaluate how these may be integrated toguether.
- This precise work will be focused around exploring all these concepts in the context of drug design for cancer 58
- [...] the work will be done in colaboration with the Computational Systems Biology group at IBM Research 59
- (Zurich). [...] The group is currently focused on individualised paediatric cure (iPC), so an end goal of this
- project is for the end results of it to help in that effor, for instance in contibuting to the ongoing research in 61
- neuroblastoma.
- As mentioned previously, the idea of using GCNNs is not a new one in the literature [Sun et al., 2019].
- My project will build upon those ideas presented in the literature, expand them and test their feasibility by
- implementing them into a wider framework for drug design [Born et al., 2019]. In that context two main
- areas of application appear. One of them would be to re-desing the drug conditional generator, for instance
- by reframing the vairational autoencoders, used for molecule generation, to architectures that operate over
- graphs [Simonovsky and Komodakis, 2018, Li et al., 2018a, Li et al., 2018b]. The second area would be to
- find better ways to asses the activity of these molecules, and in a wider context, assess their relevance as drug
- candidates. In the concrete case of the mentioned framework it is done by using a critic network proposed
- in [Manica et al., 2019]. This could be expanded on a set of different fronts: usign structural data instead
- of SMILES [Li et al., 2017, Do et al., 2019], by using GCNNs to cover a much wider network of genes
- [Oskooei et al., 2019, Wang et al., 2019], or by introducing particular scores (rewards) based in the interaction 73 of the compound to certain targets [Yingkai Gao et al., 2018, Zhavoronkov et al., 2019] or the combination of
- the compund with other drugs [Zitnik et al., 2018] —a common practice in patients with cancer. 75
- All these possible changes on the critic model would apply at different abstraction levels. That opens 76
- the door to seek for ways to integrate the representations learnt at those different stages [Ying et al., 2018,
- Ma and Zhang, 2019, Huang et al., 2019]. On top of that information extracted from here could be then
- leveraged on the drug generation part of the framework.

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