The urgent need for pan-antiviral agents: from multitarget discovery to multiscale design

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“multitarget drug discovery is a more complete paradigm than its one-drug-to-one target counterpart because finding multitarget inhibitors will also reveal information about target preferences and selectivity, thus providing insights for the design of selective inhibitors.”

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Viral infections continue to plague mankind. In the last 20 years the world has witnessed the increased emergence of different viral outbreaks: SARS in 2002, influenza A (H1N1) in 2009, the Middle East respiratory syndrome in 2012, Western/Eastern equine encephalitis in 2013 and Ebola in 2014. All these viral diseases are lethal and highly transmissible through travelers [1]. For most of these viruses, there are no treatments able to eradicate them. Recently, SARS-CoV-2, responsible for the COVID-19 pandemic has emerged, leaving a trail of more than 32.5 million infected people and more than 989,000 deaths as of 26 September 2020 [2]; the great morbidity and mortality associated with SARS-CoV-2/COVID-19 are yet expected to increase. The period between these outbreaks has shortened, suggesting that more viral epidemics will be seen in the years to come. Consequently, there is an urgent need to discover or design pan-antiviral agents (also known as broad-spectrum antivirals). In this article we highlight the different aspects that point to the development of pan-antiviral agents and the most promising in silico methodologies that can expedite this process. Emphasis is placed on finding potential treatments against SARS-CoV-2/COVID-19 from the perspective of discovering pan-antivirals which could also assist with the elimination of other viral pathogens.

The decadence of the one-target-one-drug paradigm

The vast majority of the approaches at both computational and experimental levels have focused on discovering drugs that inhibit specific targets. However, the many success stories in developing drugs to treat diseases barely compensate for the remarkably high expenditure of time (at least 12 years) and financial resources (around US$3 billion per new drug approved) [3]. Thus, the one-drug-to-one-target paradigm in drug discovery is characterized by many failures and great uncertainty. The most recent example demonstrating the drawbacks of designing highly specific monotargeted drugs is the disappointing result obtained in preliminary clinical trials by attempting to repurpose the drug combination lopinavir/ritonavir to treat SARS-CoV-2 [4]. These are well-known anti-HIV drugs acting as protease inhibitors, which were expected to have in vivo efficacy against SARS-CoV-2 by inhibiting the viral protein 3CLpro. The negative result of this study suggests two possible options: either antivirals should eradicate SARS-CoV-2 by inhibiting targets related to other processes, such as entry or replication, or they should act as multitarget inhibitors, disrupting several pathways necessary for virus survival and/or infectivity. Regarding the first option, DrugBank [5] reports 42 of the most promising anti-SARS-CoV2 experimental treat-
Drug repurposing of FDA-approved drugs as pan-antiviral agents using advanced computational models

In the race to find potential treatments against SARS-CoV-2/COVID-19 and other viruses, in silico models could serve as tools to identify FDA-approved drugs that can behave as pan-antivirals. Popular techniques relying on structure-based approaches (e.g., molecular docking) will always be useful but are computationally demanding if applied to the search for pan-antivirals via multitarget inhibition against diverse viral proteins. Therefore we will focus on 2D ligand-based approaches, which can be performed faster and without the diminution of accuracy. We would like to highlight three popular/important approaches that can be used to repurpose FDA-approved drugs as pan-antiviral agents. Each of them either directly or indirectly benefits from the fact that some proteins remain conserved across different viral species. Thus even if there is little information about an emerging pathogen, inhibiting different proteins of already known viruses (or the viruses themselves) could provide a starting point for inhibiting the new virus.

The first approach focuses on models combining perturbation theory and machine learning (PTML) [8,9] which, through the application of Box–Jenkins operators, can integrate in vitro screening data containing multiple viral proteins from different viruses. This enables the screening of FDA-approved drugs, and those predicted as versatile inhibitors of the viral proteins can be experimentally validated as potential treatments against SARS-CoV-2 and other viral pathogens. In the second approach, PTML models are applied to virtually screen drugs against different viruses in inhibition/replication assays. Again, the drugs predicted as pan-antivirals will be prioritized in biological testing against SARS-CoV-2. We would like to emphasize that, in antiviral research, PTML models have found successful applications in modeling anti-HIV activity through multitarget inhibition [10], linking in vitro activity with epidemiological outcomes [11] and virtually screening antiretroviral compounds [12]. Therefore PTML models could be easily adapted and built for the two aforementioned cases.

The third approach involves the development of alignment-free multitarget (AFMT) models [13]. These computational tools are constructed by calculating molecular descriptors from both the 2D structures of the ligands/inhibitors and the proteins’ amino acid sequences. The advantage of the AFMT models over their PTML counterparts is that the former can predict the inhibitory potency of large libraries of (either virtual or already synthesized) chemicals against proteins other than those used to develop the AFMT models [13,14]. AFMT models can therefore screen FDA-approved drugs against SARS-CoV-2 proteins (including the spike protein), whose sequences can be retrieved from public sources such as UniProt [15] or Protein Data Bank [16]; the drugs predicted as multitarget inhibitors of the SARS-CoV-2 proteins can then be fast-tracked for experimental confirmation.

Designing new molecular entities as potent & safe pan-antivirals

Considering the accumulated experience, the multifactorial nature of viral infections and the limitations of current antiviral therapies, new molecular entities (NMEs) could be computationally designed as pan-antivirals. We are referring to a chemical able to inhibit multiple viruses that are dissimilar in terms of their genomes, replication, transmission, mutability and infection mechanisms. Such a pan-antiviral agent will offer a better chance of eliminating SARS-CoV-2 and other viral pathogens. To reduce both the response time against a viral pandemic and financial resources, the chosen computational approach must integrate both in vitro and in vivo assays while focusing
on designing a small library of 5–20 chemicals that virtually exhibit potent pan-antiviral activity, low toxicity and adequate pharmacokinetic properties.

To date, the most suitable approach capable of designing an NME is multiscale de novo drug design [17], which involves four steps. First, a multitasking model for quantitative structure–biological effect relationships (mtk-QSBER) is built [18]. Then the most common molecular fragments present in the dataset used to create the model are extracted and their quantitative contributions to the pan-antiviral activity and the safety profiles are calculated; fragments with positive contributions will be used as building blocks to design NMEs. Third, the molecular descriptors present in the mtk-QSBER model are interpreted, providing information regarding the physicochemical properties and structural features that a molecule should possess to simultaneously display pan-antiviral activity, reduced toxicity and desirable pharmacokinetic profiles. Last, NMEs are generated by assembling fragments with positive contributions according to the interpretations of the molecular descriptors. In the context of antiviral research, multiscale de novo drug design has been successfully applied to the virtual generation of anti-hepatitis C agents [19] and anti-HIV compounds [20]. This approach represents the fastest and most economical solution to discover potent and safe pan-antivirals while preventing promiscuous (off-target) interactions potentially associated with side effects.

Conclusion

Viral infections represent a constant threat to humanity. The latest viral outbreaks have derived from RNA-based viruses, but it must not be taken for granted that DNA-based viruses are unable to cause pandemics. Developing pan-antiviral agents is perhaps the most efficient way to tackle all the emerging and re-emerging viral pathogens. The search for such therapeutic agents should be performed with the aim of finding chemicals that will disrupt dissimilar mechanisms of action, stopping the biomolecular machinery of viral infection and/or replication across multiple viral species. Computational approaches are and will be essential in accelerating the search for pan-antiviral agents, either by repurposing drugs with multitarget inhibition patterns via the PTML and AFMT models, or by using multiscale de novo drug design to rationally generate NMEs with the desired properties.

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