Repurposing Cardio-Metabolic Drugs to Fight Covid19

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Since November 2019, healthcare systems around the world had to face the consequences of the pandemic caused by the Sars-CoV-2 virus. This virus is a positive-sense single-stranded RNA virus classified as coronavirus. Two previous epidemics caused by coronaviruses developed in 2000s, but had been confined to specific regions of the world. On the contrary, the disease associated with Sars-Cov-2 infection, known as Covid19, due to its very high contagiousness, has taken on the measure of a pandemic. To date, more than two hundred million cases are confirmed, with over four million deaths globally [1].

Covid19 mostly manifests with interstitial pneumonia, even though it can affect heart and central nervous system as well. Although the pathophysiology of the clinical manifestations of Covid19 is not yet completely known, it has been shown that the entry of Sars-CoV-2 into cells is mediated by angiotensin-converting enzyme 2 (ACE2), expressed also in pulmonary epithelium [2]. After that, symptoms and the severity of Covid19 depend on the virus replication but also on the individual's immune response, often establishing an immune/inflammatory storm leading to tissue damage that can have severe prognosis [2, 3].

In parallel with the progressive knowledge about the mechanisms of Sars-CoV-2 infection and the pathophysiology of Covid19, is not yet completely known, it has been shown that the entry of Sars-CoV-2 into cells is mediated by angiotensin-converting enzyme 2 (ACE2), expressed also in pulmonary epithelium [2]. After that, symptoms and the severity of Covid19 depend on the virus replication but also on the individual's immune response, often establishing an immune/inflammatory storm leading to tissue damage that can have severe prognosis [2, 3].

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The first is that a single drug interacts with multiple targets, so that a drug already known might have new targets of action [10, 11]. The second is that targets associated with a disease might be involved in many biological processes [12, 13] so that a known target might have a new indication. Therefore, it has been considered that 75% of known drugs could be repositioned for various diseases [9].

Moreover, another study by Fehr et al. [24] revealed that SARS-CoV-2 infects human cells by using the host’s translation pathway to produce twenty-nine viral proteins. These twenty-nine proteins bind to multiple human proteins to set up the molecular processes necessary to develop the infection. Among these protein, Gorden et al. [25] using affinity purification mass spectrometry released the twenty-six proteins that the SARS-CoV-2 uses to infect the human body and identified more than three hundred human proteins involved in these viral proteins binds. More than sixty of these proteins appear to be druggable human proteins with already existing drugs. Therefore, the identification of interactions between host and virus proteins might provide significant insights to find drug targets for drug development against SARS-CoV-2.

Among the new approaches for drug repurposing, Mendelian randomization (MR) analyses have also been used. MR analysis focusing on actionable genes, defined as genes that encode proteins targets of drugs already existent, could therefore contribute to repurpose drugs useful in the treatment of Covid19 [21].

The first is that a single drug interacts with multiple targets, so that a drug already known might have new targets of action [10, 11]. The second is that targets associated with a disease might be involved in many biological processes [12, 13] so that a known target might have a new indication. Therefore, it has been considered that 75% of known drugs could be repositioned for various diseases [9]. Therefore, during the first year of the pandemic, several drugs were repurposed including those drugs acting through virus related targets such as RNA genome (i.e. remdesivir); drugs acting through polypeptide packing (i.e. lopinavir and ritonavir); drugs acting through host targets such as antiviral immunity (i.e. interferons), drugs targeting the virus uptake pathways (i.e. Chloroquine) and drugs acting on host proinflammatory cytokines (i.e. Tocilizumab) [14]. Few of these drugs have been also tested in the Solidarity clinical trial for Covid19 [15], which is a multinational phase III-IV clinical trial organized by the World Health Organization (WHO) and partners to compare four untested treatments for hospitalized people with severe Covid19 illness.

Traditionally, the repurposing of an existing agent, appealing due to its biological plausibility, takes much experimental work both in vitro and in vivo. Expertise in pharmacology, as well as expertise in translational drug development and clinical trials is needed to set up small-scale studies and then larger population-wide study are crucial to confirm the use of a given compound for a new indication [16]. Although large randomized trials have begun to identify drugs that can be repurposed to face Covid19 [17–21], many of them have been conducted only in hospitalized and severely ill patients mostly failing to show any efficacy. Therefore, to identify additional drugs that can be repurposed also for early management in Covid19, remains a priority.

Alongside with this “traditional” process based on pathophysiology and pharmacodynamics, systems available in the 21st century have made it possible to add other ways to identify drugs already on the market and potentially useful to treat Covid19. Broadly, there are three kinds of approaches that can be used in drug repositioning: biological experimental approaches (the traditional ones), computational approaches and mixed approaches that can be especially useful in emergency situations such as the Covid19 pandemic [14].

Many computational methods are now available to examine the relationships between chemical compounds, molecular targets, biological systems, therapeutic endpoints and diseases [22]. For example, it has been presented a new network-based approach for drug repositioning, called SAvReRUNNER (Searching off-lAbel dRUg aNd NEtwoRk), which predicts drug-disease associations by quantifying the interplay between the drug targets and the disease-specific proteins in the human interactome via a novel network-based similarity measure that detect associations between drugs and diseases locating in the same network neighborhoods. With this approach, many drugs possibly useful in Covid19 have been found including ACE-inhibitors (ACEi), monoclonal antibodies (e.g., anti-IFNγ, anti-TNFα, anti-IL12, anti-IL1β, anti-IL6), and thrombin inhibitors [23].
inhibitors, but they also play many pleiotropic effects. They can reduce vascular and endothelial inflammation, improve endothelial function also by ameliorating the vascular redox balance and inhibit platelet aggregation [33]. Moreover, they seem to have an immunomodulatory activity by modifying the release of cytokines during the cytokine storm and the expression of major histocompatibility complex class II increasing the production of protective Th-2 cytokines [34]. Moreover, statins can exert some direct antiviral activity by inhibiting cholesterol synthesis, which is involved in the viral cell cycle [35]. Indeed, the presence of cholesterol-rich subdomains on the plasma membrane of host cells, namely lipid rafts, is crucial for viral fusion and entry [36, 37]. In addiction, via epigenetic histone modifications, statins might upregulate the expression of ACE2 [38] which is known to be associated with reduced severity of acute respiratory distress syndrome (ARDS) [39].

Last year, Zhang et al. reported the first large observational study of statin use in Covid19 [40]. In a retrospective cohort 13,981 patients hospitalized with Covid19 in Hubei Province, China, among which 1219 received statins they examined the association between statin use and outcomes. The authors found that the risk for 28-day all-cause mortality was 5.2% in the statin group and 9.4% in the matched non-statin group (HR 0.58; 95% CI, 0.43–0.80; p = 0.001). Despite the promising results, it must be stated that there are important limitations in this study, such as its retrospective nature, the lack of data on pre-hospital statin use and concomitant therapies, suboptimal matching techniques between the two groups. To date, even though consistent evidence from prospective studies is not currently available, most of available data from retrospective studies support the beneficial effect of statins against SARS-CoV-2 infection in terms of inflammatory parameters, severity of clinical manifestations and mortality [41]. On the contrary Cariou et al. [42], using data from the large-scale multicenter CORONADO study carried out in France in diabetic patients hospitalized for Covid19, reported that routine statin treatment is significantly associated with increased mortality (based on 7-day and 28-day in-hospital death rates).

Recently, a total of eleven highly heterogeneous observational studies were included in a meta-analysis for a total of 3462 statin users and 10,560 non-users. Although no significant reductions in either in-hospital mortality or Covid19 severity were reported among statin users compared with non-users after univariate comparisons, such reductions were observed after adjusting for confounding factors [43].

The most recent available meta-analysis [44] was limited to seven retrospective studies performed in Western countries with 2398 patients with 44.8% taking statins. Five out of seven studies evaluated patients with statin treatment before hospital admission while the others considered both treatments started before admission or during hospital stay. Covid19 patients taking statins had nearly 40% lower odds of incurring in the composite endpoint of severe illness or death (OR: 0.59; 95% CI: 0.35–0.99), with only one study [45] reporting an OR > 1 (i.e., OR: 1.60; 95% CI: 0.88–2.92).

Lately, it has been also proposed that fenofibrates, suppressing inflammation and apoptosis cascades through inactivation of NF-kB and stimulation of adenosine monophosphate-activated protein kinase (AMPK) signaling, might be useful in treating Covid19 [46].

Moreover, metformin, which has been firstly discovered in the search for antimarial agents and now has become one of the most prescribed antidiabetic drugs worldwide seems to have a new potential therapeutic application in Covid19 [47]. Indeed, retrospective studies [48] and meta-analyses reported significant reduction of infection-related mortality in patients affected by Covid19 and treated with metformin [49, 50]. Currently, the “COVIDOUT—Outpatient Treatment of COVID-19 with Metformin” (REGISTRATION: URL: https://www.clinicaltrials.gov NCT04510194) a phase II/III trial involving 750 participants aged 30–85 years is investigating whether metformin treatment in non-hospitalized adults with SARS-CoV-2 can (a) prevent hypoxia and emergency department utilization, (b) prevent disease progression in Covid19, and (c) improve viral load and C-reactive protein (CRP). Moreover, a phase II trial “Pilot Study into the Use of Metformin and Low Dose Naltrexone (LDN) for Patients with Coronavirus Disease 2019 (COVID-19)-Assessment of Short and Long Term Effects” (REGISTRATION: URL: https://www.clinicaltrials.gov; NCT04604678) involving 80 participants aged 30–70 years aims to study monthly the effect of a combination of metformin and LDN on hard and soft outcomes in Covid19 patients.

Furthermore, through an integrative bioinformatics approach including the search of the biomedical literature for high confidence DPP4-protein/gene associations followed by functional analysis using network analysis and pathway enrichment it has been found that DPP4 networks are highly enriched in viral processes required for viral entry and infection, and as a result, DPP4 was proposed as an important possible target for the treatment of Covid19. Since protein-chemical interaction networks identified strong interactions between DPP4 and sitagliptin, this latter has been postulated to be beneficial for the treatment of Covid19 disease, either as monotherapy or in combination with other therapies, especially for high CV risk patients [51].

Finally, since Covid19 is associated with the ARDS and higher activity of ACE2 leads to attenuation of ARDS and people taking angiotensin receptor blockers (ARBs) or ACEIs have higher levels of expression of ACE2, experimental and clinical studies have been performed to investigate the influence of ARBs or ACEIs on Covid19 [52]. Moreover, spironolactone, a potassium-sparing diuretic
drug antagonizing mineralocorticoid receptors, display favorable effects on ACE2 expression, preventing acute lung injuries, also in Covid19 [53, 54]. Therefore, several prospective randomized control trials investigating the protective effect of renin-angiotensin-aldosterone system-blockers in Covid19 are currently ongoing (REGISTRATION: URL: https://www.clinicaltrials.gov; Unique identifier: eg, NCT04493359, NCT04394117, NCT04591210, NCT04312009, NCT04351581, NCT04366050; and DRKS00021732, NCT04345887).

To date, the main advantage of drug repurposing lies in its economy. Indeed, the approaches used are faster and less expensive than the design and development of a drug from scratch. However, there are not many centers that can afford to perform a reliable computational approach and many of the drugs that have so far been identified with these approaches have not given the desired results.

Therefore, if on the one hand the strategy of using drugs already placed on the market to save time and resources appears fascinating, on the other hand it seems more likely that on a global level it will not be sufficient to change the fate of Covid19.

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Conflicts of interest Nothing to declare.

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