Immunomodulators Acting on Covid-19: Actual Knowledge and Perspectives

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Abstract
Since the outbreak of SARS-CoV-2 infection (Covid-19), healthcare professionals worldwide have been trying to find disease management and control alternatives to encourage immunotherapies. Immunotherapy is an efficient therapeutic option used against comparable viral contaminations such as MERS-CoV and SARS-CoV. The aim of the current study is to assess the existing knowledge associated with SARS-CoV-2 immunotherapy. Information available in published articles and their quality highlights the importance of following strict scientific rules for clinical outcomes. Thus, these studies have shown enough data to confirm that immunomodulation is the main topic investigated in research about Covid-19 therapy. Therefore, it is possible saying that immunotherapy is certainly the appropriate option against this virus.

Keywords: COVID-19, Immunotherapy, Immunomodulator, Antibody, Plasma, Immunoglobulins.

Introduction
Covid-19 appears to be quite different from other SARS-CoV strains. Approximately 85% of contaminated individuals present mild-to-moderate symptoms, whereas approximately 10% of them present severe symptoms, approximately 5% present severe symptoms and half of them end up dying. Interestingly, contamination by Covid-19 in children and adolescents is milder than in adult patients; the most significant death rate is associated with elderly and adult individuals with comorbidities such as cardiopathic diseases, diabetes mellitus, among others. If we think about a good explanation for these findings, it is possible hypothesizing that Covid-19 and its interactions with the hosts’ immune network can help elucidating the route of Covid-19 infections [1-5]. Healthy adult individuals present undamaged innate immunity, as well as enhanced humoral and cell-mediated immunity. These conditions inhibit infection progression and enable patients to recover within few weeks, since SARS-CoV-2 does not reach their alveoli, differently from critically-ill patients. Elderly patients with, or without, co-morbidities and compromised immune system do not have the same innate and adaptive humoral immune response as healthy patients; thus, SARS-CoV-2 often remains mild until it reaches the alveoli, since the blood stream is protected by alveolar lymphocytes and macrophages. The immune system becomes more active in the alveoli because they...
are the last site for the virus to attack blood circulation. Epithelial cells in airways operate as immune effector molecules and release cytokines, chemokines, among other signaling proteins that play a key role in cell-mediated immunity. Thus, epithelial cells start eradicating infected cells, along with the emergence of systemic cytokines. This process leads to significant damage in lung tissues; therefore, alveoli get full of inflammatory exudates. After some time, patients experience severe hypoxia, as well as lung, liver and kidney damage, because the innate and humoral immunity in their alveoli triggers cell-mediated immunity to set up an extremely dangerous attack at the site. The phenomenon observed in adult and elderly individuals does not explain why children follow a different immunopathogenic path. Abdulamir and Hafidh have suggested that children's immune system is somehow lesser capable of attacking alveoli and lung tissues [6].

Assumingly, children’s immune system does not have previous memory of coronavirus infection, likely because SARS-CoV-2 shares similar proteins to those found in the common human coronavirus. This is a challenging immunopathogenesis topic that should be thoroughly investigated.

Based on the literature and on data shown in Table 1, almost all immunotherapy assays conducted with 2019-nCoV patients so far incorporate treatments comprising plasma, immunoglobulin, thymosin, tocilizumab, sarilumab, immunoglobulin Fc domain, CR3022, convalescent plasma transfusion, systemic anticoagulation, heparin, statin and interferon. IFN-α2b, Interferon IFN-β1b, BCG and OncoTherad have been used to treat SARS-CoV and they can become a promising alternatives against 2019-nCoV. Using immunotherapy against this virus could also be an adequate option [7-10].

It is known that coronavirus infection induces increased levels of T-helper 1 (Th1) cytokine, interferon (IFN)-gamma, inflammatory cytokines such as interleukins IL-1, IL-6 and IL-12, and connected cytokines and chemokines such as IL-8, chemokine (C-C motif) ligand 2 (CCL2 protein, also known as monocyte chemotactant protein-1 or MCP-1) and C-X-C motif chemokine 10 (CXCL10 protein, named as interferon-γ, induced protein 10 or IP-10) [11,12]. The release of these cytokines moderated by these inflammatory and Th1 cytokines activate monocytes/macrophages and neutrophils account for immunopathological outcomes observed for this contamination type.

These hyper-inflammatory immune responses lead to increased Covid-19 severity and, consequently, to high mortality rates. Thus, it is possible saying that hyper-inflammatory process inhibition is an unequivocal drug therapy alternative [11,13].

The current review addresses key medications suggested for Covid-19 treatment, based on immunological modulation, which have been reported in the literature, so far. Topics such as immunosuppression and vaccines were not addressed in the current manuscript.

### Data collection

Inclusion criteria were all immunomodulators in uses on Covid-19 (Plasma therapy, Immunoglobulin, Thymosin, Tocilizumab, Sarilumab, Immunoglobulin Fc domain, CR3022 (SARS-CoV), Anakinra, Convalescent plasma transfusion, Systemic anticoagulation, Heparin, Statin, Interferon, IFN-α2b, Interferon. IFN-β1b, BCG, BCG, Oncotherad). The data collection was done in the period of 2003 to 2020 with emphasis mainly in 2020 by several authors on this topic. Besides these data, several clinical trials were discussed and still in progress.

### Antibodies

Covid-19 antibodies are active subjects; among them one finds tocilizumab and sarilumab, which are the most studied ones; besides, they have recorded the most relevant results. The literature reports several clinical trials on different kinds of immunomodulators [14]; some of them will be herein addressed. A study conducted with 21 Chinese patients reported that tocilizumab has eliminated lung lesion opacity in 91% of cases, reduced oxygen intake in 75% of patients and ventilation was not necessary in approximately 5% of cases. High C-reactive protein levels significantly decreased in 84% of patients, whereas lymphocytes got back to normal levels in 53% of cases [15]. Another trial conducted with tocilizumab in China was lately approved by the National Health Commission of the People's Republic of China to be used in patients suffering with acute SARS-CoV-2 pulmonary implications [16]. Accordingly, sarilumab - which is an anti-human IL-6 receptor monoclonal antibody presenting activity similar to that of tocilizumab - started being used to treat rheumatoid arthritis. Sarilumab is capable of blocking IL-6, just like tocilizumab does; thus, it may have positive effects on Covid-19 patients presenting acute expressions of the infection and high IL6 levels. A phase II-III clinical trial was recently implemented in USA and in five European Countries [17].

Although anakinra is an immunosuppressor used against IL-1 (anti-IL-1), it is another medicinal alternative to treat patients with acute Covid-19. It is a recombinant and superficially transformed type of human interleukin 1 receptor antagonist protein (IL-1Ra). This protein is naturally liberated by monocytes and tissue macrophages that bind to IL-R and regulate its action. IL-1 blockade leads to inflammatory response suppression [18]. Based on a phase-III randomized controlled trial, anakinra was capable of increasing the survival rate of patients with sepsis without increased adverse effects [19].

Ten clinical trials focused on investigating the effects of anakinra application in Covid-19 patients have started in May 2020 and remain in progress [14]. As previously mentioned, anakinra is capable of inhibiting pro-inflammatory cytokines such as interleukin (IL)-1α/ IL-1β; data have shown its effectiveness in treating macrophage activation syndrome generated by several immunological circumstances, as well as in small assays conducted with Covid-19 patients [20,21].

### Immunoglobulins/Heparin

Preliminary studies have shown the effectiveness of
<table>
<thead>
<tr>
<th>Treatment by</th>
<th>Type of immunology (vaccine and...)</th>
<th>Accompanied by another treatment</th>
<th>Number of patient treated</th>
<th>Outcomes, results and relationship</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plasma therapy</td>
<td>Polyclonal antibody immune response</td>
<td>Standard treatment</td>
<td>300 patients in COVID-19 clinical trial</td>
<td>Clinical improvement</td>
<td>Zhang et al., [73]</td>
</tr>
<tr>
<td>Immunoglobulin</td>
<td>----</td>
<td>Standard treatment</td>
<td>80 patients in COVID-19 clinical trial</td>
<td>Clinical improvement</td>
<td>Zhang et al., [73]</td>
</tr>
<tr>
<td>Thymosin</td>
<td>Polypeptide hormone for maturation of T cells</td>
<td>Camrelizumab (anti–PD-1 immune checkpoint inhibitor), conventional treatment</td>
<td>120 patients in COVID-19 clinical trial</td>
<td>Lung injury score</td>
<td>Zhang et al., [73]</td>
</tr>
<tr>
<td>Tocilizumab</td>
<td>Monoclonal antibody against the IL-6 receptor (IL-6R)</td>
<td>-----</td>
<td>188 patients in COVID-19 clinical trial</td>
<td>Cure rate</td>
<td>Zhang et al., [73]</td>
</tr>
<tr>
<td>Tocilizumab (Actemra)</td>
<td>Monoclonal antibody against IL-6 inhibition - reduction in cytokine storm</td>
<td>-----</td>
<td>21 patients In COVID-19 clinical trial</td>
<td>Cure rate</td>
<td>Xu et al., [15]</td>
</tr>
<tr>
<td>Tocilizumab</td>
<td>Monoclonal antibody against IL-6 inhibition - reduction in cytokine storm</td>
<td>Conventional therapy</td>
<td>188 patients in Covid-19 clinical trial</td>
<td>Cure rate</td>
<td>CCTR [16]</td>
</tr>
<tr>
<td>Sarilumab</td>
<td>Monoclonal antibody against IL-6R</td>
<td>-----</td>
<td>300 patients Trials: Italy, Spain, Germany, France, Japan, Canada and Russia</td>
<td>Eur. Pharm. Review Report [17],</td>
<td></td>
</tr>
<tr>
<td>Immunoglobulin Fc domain</td>
<td>ACE2 immunoadhesin</td>
<td>-----</td>
<td>-----</td>
<td>Untested</td>
<td>Kruse [71]</td>
</tr>
<tr>
<td>CR3022 (SARS-CoV)</td>
<td>Monoclonal antibody (cross reactive antibody)</td>
<td>Alone or combination with other neutralizing antibody (e.g. m396, CR3014)</td>
<td>-----</td>
<td>Untested</td>
<td>Tian et al., [72]</td>
</tr>
<tr>
<td>Anakinra</td>
<td>Recombinant human nonglycosylated interleukin-1 (IL-1) receptor antagonist</td>
<td>Conventional therapy (subcutaneous or intravenous)</td>
<td>Different numbers of patients in France, Greece, USA, Belgium and UK</td>
<td>-----</td>
<td>Alijotas-Reig et al., [14]</td>
</tr>
<tr>
<td>Convalescent plasma transfusion</td>
<td>Convalescent plasma transfusion from cure patients rich containing specific antibody (IgG)</td>
<td>Associated to mechanical ventilation, interferon-alpha-1b and favipiravird</td>
<td>5 critical patients 2 remaining hospitalized. 3 discharged home</td>
<td>Cure rate</td>
<td>Shen et al., [24]</td>
</tr>
<tr>
<td>Systemic anticoagulation</td>
<td>Not specified</td>
<td>Associated to mechanical ventilation</td>
<td>786 patients</td>
<td>Cure rate</td>
<td>Paranjpe et al., [33]</td>
</tr>
<tr>
<td>Heparin</td>
<td>Not specified</td>
<td>-----</td>
<td>2075 patients, several hospitals in Spain</td>
<td>Cure rate</td>
<td>Ayerbe et al., [32]</td>
</tr>
<tr>
<td>Heparin</td>
<td>Low molecular weight heparin</td>
<td>-----</td>
<td>99 patients</td>
<td>Cure rate</td>
<td>Tang et al., [28]</td>
</tr>
<tr>
<td>Statin</td>
<td>One of them: Atorvastatin, fluvastatin, pravastatin, rosuvastatin, simvastatin</td>
<td>-----</td>
<td>154 patients</td>
<td>Cure rate</td>
<td>Dashti-Khavidaki and Khalili, [36]</td>
</tr>
<tr>
<td>Interferon IFN-α2b</td>
<td>Presence and absence of Arbidol</td>
<td>-----</td>
<td>77 patients</td>
<td>Cure rate</td>
<td>Zhou et al., [43]</td>
</tr>
<tr>
<td>Interferon IFN-β1b</td>
<td>Associated to lopinavir–ritonavir, and ribavirin, compared with single-drug lopinavir–ritonavir</td>
<td>-----</td>
<td>127 patients</td>
<td>Cure rate</td>
<td>Hung et al., [42]</td>
</tr>
<tr>
<td>Interferons</td>
<td>Boosts the antiviral immune response in the early phase of the disease</td>
<td>-----</td>
<td>A total of 227 patients (4 trials)</td>
<td>Cure rate</td>
<td>Saber-Ayad et al., [2]</td>
</tr>
<tr>
<td>BCG</td>
<td>Several clinical trials are in progress</td>
<td>-----</td>
<td>A total of 17378 patients</td>
<td>Cure care</td>
<td>Soliman et al.; [55]</td>
</tr>
<tr>
<td>OncoTherad</td>
<td>Associated to bladder cancer treatment</td>
<td>Patients refractory to BCG treatment</td>
<td>5 patients</td>
<td>Cure care of COVID-19 infections</td>
<td>Delafiori et al.; MS-Brazil, [10,70]</td>
</tr>
</tbody>
</table>

Table 1: Main characteristics of included studies: Immunotherapy potentials (except vaccine) on Covid-19
intravenous immunoglobulin (IVIg) applications in the therapy of patients with acute inflammatory discomfort associated with influenza and SARS-CoV infections. A study reported the effectiveness of high IVIg immunoglobulin doses in 3 Covid-19 patients [22]. However, it is necessary conducting further studies with a larger number of Covid-19 patients in order to validate this clinical outcome. Thus, a randomized controlled clinical trial based on IVIg immunoglobulin application in patients with acute SARS-CoV-2 infection was implemented (Clinical Trial.gov: NCT 04261426) [14].

Alijotas-Reig et al. have pointed out two potentially unfavorable effects of acute IVIg use that may have harmful survival effect on patients with acute Covid-19, namely: 1) [14]. Adverse effect presenting high death rate, called TRALI (transfusion (immunoglobulin)-related severe lung injury), which comprises severe respiratory distress syndrome within few perfusion hours [23]; 2) and thrombotic events associated with IVIg therapy, whose estimated incidence reaches up to 17% [23]. The aforementioned authors suggested that, due to the dubious effectiveness of IVIg therapy in patients with SARS-CoV and to their likely risk of developing acute lung injury and thrombosis, IVIg should be carefully applied to Covid-19 patients as alternative therapy. Besides, according to comments by Alijotas-Reig et al. [14] about preliminary reports on convalescent plasma/hyperimmune immunoglobulin therapy, this treatment was capable of decreasing patient mortality rates, mainly when it was applied at early stages of the disease, right after symptoms onset. However, the true effectiveness of this therapy remains controversial; thus, the aforementioned authors have suggested that this treatment should be assessed through well-designed clinical trials [24].

Similarly to SARS-CoV patients, Covid-19 patients in acute conditions can develop disseminated intravascular coagulation, arterial thrombosis and pulmonary thromboembolism (PTE), which is one of the reasons why different diagnostic alternatives were suggested for Covid-19 patients with clinical suspicion of PTE [25]. Besides, 50 Covid-19 patients subjected to autopsy presented microthrombosis and, in some cases, thrombus involving wide pulmonary artery regions. Histological analyses have shown that alveolar and interstitial inflammation spread towards juxtaposed pulmonary vasculature, as well as the normal circulatory fibrinogen amounts and local fibrinolysis with high D-dimer (fibrin degradation product) formation, small protein fragment in the blood) at the initial stage of Covid-19 pneumonia, were not associated with severe onset of macrophage activation syndrome/hemophagocytic syndrome [26]. This hyper-inflammatory intra-pulmonary infection may influence patients’ trend to develop acute regional vascular dysfunction, together with microthrombosis and hemorrhage. This process leads to lung centric pulmonary intravascular coagulopathy that resembles disseminated intravascular coagulation [27].

Some patients presented significantly abnormal coagulation function, which suggests that the initial intravenous immunoglobulin application and the low molecular weight heparin anticoagulation treatment were very relevant. This treatment is recommended for Covid-19 patients who present elevated D-Dimer levels [23]. Tang et al. [28] found the progress of clot activation markers that have reduced the rate of death by Covid-19 when heparin was administered. Besides, heparin has pharmacological effects other than the antithrombotic ones. It also has anti-inflammatory and immune-modulating properties. Low-molecular weight heparin (LMWH) enables the survival of human endothelial cells undergoing apoptosis in response to TNF-α (tumor necrosis factor-alpha) binds to antiphospholipid antibodies and stops complement pathway activation [29,30]. Interestingly, fondaparinux (saccharide with antithrombotic activity) is often used in heparin-allergy cases and presents excellent results. All these outcomes are associated with viral infections (e.g., SARS-CoV, SARS-CoV-2) and end up presenting antiphospholipid antibodies with potential pathogenic effects [31]. It is possible saying that the heparin-based anticoagulant therapy helps avoiding thrombosis and simultaneously down-regulates pro-inflammatory pathways [14].

A clinical trial comprising 2,075 Covid-19 patients subjected to heparin therapy, as well as to others drugs, was conducted in 17 hospitals in Spain; results have shown that 301 patients died, 1,447 were discharged from hospital, 201 remained hospitalized and 126 were transferred to other hospitals (these patients were excluded from the study) - heparin was used in 1,734 patients after this study. Finally, heparin was associated with reduced mortality rates when the model was adjusted based on patients’ age and sex [32].

A clinical trial was conducted with 99 Covid-19 patients treated with heparin (mainly with low molecular-weight heparin) for a week or longer. There was no difference in 28-day mortality between heparin users and non-users. However, the 28-day mortality of heparin in patients with SIC (sepsis-induced coagulopathy) was lower than that of non-users; score ≥4 or D-dimer >6-fold the upper normal limit. Then, anticoagulant therapies, mostly the ones based on low molecular-weight heparin, are associated with better prognosis in acute Covid-19 patients who meet SIC criteria or present significantly high D-dimer levels [28].

Another clinical trial run in a hospital reported low mortality rate among 786 Covid-19 patients treated with anticoagulation drugs (no anticoagulant name was mentioned) [33].

Actually, several clinical trials are currently recruiting patients for heparin therapy (NCT04401293-USA; NCT04359277-France; NCT04366960-Italy; NCT04362085-Switzerland; NCT04345848-Switzerland; NCT04367831-USA; NCT 04373 707-France; NCT04366960-Italy; NCT04362085-Canada; NCT04359277-USA).

Statins

Statins trigger strong inhibition - via protein geranylgeranylation - of pro-inflammatory cytokine production (IL-6, -8, -10. and TNF-α) in synovial and endothelial mononuclear cells. Besides, they act as T-cell inhibitors (e.g., activation and/or proliferation) and trigger immunomodulatory activity.
Statins also preserve the gene expression of myeloid differentiation primary response 88 (MyD88) at normal levels during hypoxia and mitigate the activation of factor nuclear kappa B (NF-κB). These features explain the trend to use statins to treat Covid-19 infection [34]. Accordingly, in a murine model a high pathogenic infection, abnormal signaling coming from a lack of toll-like receptors (TLRs) adaptor TRIF or MyD88 causes severe respiratory distress syndrome (ARDS) as principal reason of mortality in SARS-CoV disease [35]. MyD88 operates downstream TLRs and results from SARS-CoV contamination. In addition, either the under-expression or over-expression of the MyD88 gene is associated with increased mortality rates after MERS-CoV infection. Thus, an NF-κB downstream TLRs-MyD88 pathway is triggered by coronavirus infections. This is an interesting finding, since NF-κB inhibition has improved lung infection and survival rates in murine models subjected to SARS-CoV infection. It may have happened due to counterbalancing activation of other innate immune factors. Although many US hospitals have incorporated statin to Covid-19 therapies applied to patients, some concerns about its use remain and caution must be taken at the time to adopt such an alternative [36].

Despite all these comments about statin use in Covid-19 patients, a clinical trial was conducted with 154 Covid-19 positive patients, who were treated with a combination of angiotensin-converting enzyme inhibitors (ACEI) (e.g., captopril)/angiotensin II receptor blockers (ARB) (e.g., losartan) and/or statin (e.g., simvastatin). Results have shown statistically significant association between statin intake and lack of symptoms in Covid-19 infection cases. This association remained statistically significant when patients were assessed based on age, sex, diabetes mellitus, hypertension and functional status. There was not statistically significant association between ACEi/ARB and asymptomatic status or severe clinical outcome. Based on these data, statin intake by elderly, fragile individuals can be associated with substantial beneficial effect on Covid-19-associated clinical symptoms [37].

**Interferon**

Data showing that interferon reacts to coronaviruses in a way different from that to other respiratory viruses is an important finding [38]. Trials assaying the application of type I or type III interferons are already in course. Type I interferon (IFNαβ) receptors are ubiquitously expressed, whereas type I interferons can lead to antiviral effects; however, they can also induce immune cell activation and increase tissue pathologies [39,40]. Moreover, SARS-CoV-2 did not efficiently induce types I, II, or III interferons in human lung tissues infected ex-vivo as it happened with 2003 SARS-CoV [41]. Thus, interferon therapy application to jump-start or enhance patients' antiviral response would be an obvious approach [42].

Interferon (IFN-α2b) application was investigated in a cohort of Covid-19 positive cases in China; 77 adult individuals infected with Covid-19 were treated with either nebulized IFN-α2b or arbidol (Umifenovir), or with the combination of both. All serial SARS-CoV-2 tests were conducted; IFN-α2b application in separate, or in association with arbidol, has effectively shortened the time the virus remained detectable in patients' upper respiratory tract and simultaneously diminished the length of high blood levels recorded for inflammatory markers such as IL-6 and C-reactive protein. Although it was a non-controlled exploratory study, results have suggested that IFN-α2b should be further investigated as therapy for Covid-19 patients [43].

A clinical trial was conducted with 127 patients treated with triple antiviral therapy based on interferon beta-1b (IFN-β1b), lopinavir/ritonavir and ribavirin (Clinical trial gov.:NCT04276688). The triple antiviral therapy was safe and better than the use of lopinavir/ritonavir alone, since it mitigated virus dissemination, relieved patients’ symptoms and enabled individuals with mild-to-moderate Covid-19 to be discharged from hospital. The interferon/lopinavir/ritonavir/ribavirin association has also diminished IL-6 levels. The clinical and antiviral efficiency of this association enabled shorter hospital stays and better infection control [42]. Other clinical trials associated with antiviral drugs are currently in progress (IFN-β1b: NCT04350684-Iran; NCT04350671-Iran; IFN-α: NCT04343976-USA) [2].

**Immunotherapies: BCG (Bacillus Calmette-Guérin) and OncoTherad**

The recent Covid-19 outbreak has made the association between BCG vaccination and Covid-19 morbidity and mortality rates reemerge in several countries [44-53]. Some studies have shown significant association between BCG vaccination and Covid-19 incidence, and likely mortality, in countries counting on BCG vaccination policies [54,55].

According to Soliman et al, there is robust evidence about the association between BCG vaccination and prevention of severe respiratory infections. This statement meets information from data available in some systematic reviews; one of them incorporated a meta-analysis and a randomized controlled trial (RCTs), which evidenced a high confirmation level for prevention studies [55,56].

Clinical trials are currently in progress and recruiting participants to determine whether BCG vaccination helps protecting healthcare workers working in the front lines of the Covid-19 pandemic and in other pandemic situations in different countries (Clinical trial gov.: NCT04327206-Australia; NCT04328441-Netherlands; NCT04379336-South Africa; NCT04347876-Egypt; NCT 04414267-Greece; NCT04373291-Denmark; NCT04348370-USA).

As previously mentioned, patients’ immune response to SARS-CoV-2 pulmonary infection involves a series of cytokines (IL-1B, IFN-γ, IP10, MCP1 and TNF-α) identified at higher concentrations in the plasma of infected individuals [57]. Robust adaptive immune response to viral clearance through IFN-γ requires early type-1 IFN response. However, previous knowledge has shown that cytokine dysregulation is the main feature of early SARS-CoV infection; CoV-cell recognition and type-1 IFN response dynamics may determine patients’ pro- or anti-inflammatory immune response and injury severity [58,59]. Therefore, immunomodulators capable of controlling early cytokine
response, such as the recruitment of effector cells and viral clearance, may be a good strategy to be used against SARS-CoV-2.

Several studies have successfully defeated CoVs based on direct IFN administration to patients. The combination of type-I IFN and IFN-γ was capable of synergistically inhibiting virus replication in vitro [60]. Larkin et al. have indicated that the combination of IFN-α and IFN-γ in vitro enabled strong synergistic antiviral activity at much lower IFN doses than often necessary [61]. Reducing the IFN dose in combination therapies has the advantage of reducing unwanted adverse reactions in patients. Nagata et al. described the destructive effect of cytokine storm in adult mice after SARS-CoV infection [62]. Although intravenous TNF-α injection did not show beneficial effects, intraperitoneal IFN-γ injection has protective effect on SARS-CoV infection. Scagnolari et al. also reported the synergistic effects of IFN-γ and IFN-β on SARS-CoV-infected Vero cells. Thus, IFN induction can play a fundamental role in protecting the human body from CoV infections [63].

Accordingly, immunotherapy with OncoTherad stands out as a potential treatment for Covid-19. The research team of the Urogenital Carcinogenesis and Immunotherapy Laboratory (UNICAMP), produced an immunomodulator and registered as OncoTherad. OncoTherad works as Biological Response Modifier and contributes to local immune system activation in tumor microenvironments. Pre-clinical, clinical-veterinary and Phase I/II clinical studies conducted with humans have shown that immunotherapy with OncoTherad has activated patients’ innate immune system mediated by Toll-like receptors (TLRs) 2 and 4; this process resulted in increased interferon signaling pathway (TLR4, TRIF, IRF3, INF-γ), which is associated with the greater effectiveness of this nano-immunotherapy to treat non-muscle invasive bladder cancer (NMIBC) in comparison to standard treatments with BCG (Bacillus Calmette-Guérin). OncoTherad was capable of decreasing RANK/RANK-L system expression in animal models and, consequently, of preventing metastases formation and/or progression (Fávaro and Durán, 2017; Böckelmann et al., 2019; Durán et al., 2019; Fávaro et al., 2019ab; Fávaro and Durán, 2020) [7,64-67].

A prospective, monocentric clinical study (Hospital Municipal de Paulínia, São Paulo, Brazil), based on phase I/II single-arm immunotherapy with OncoTherad, was conducted with patients refractory to BCG treatment. Results have shown that this treatment promoted recurrence-free survival rate (complete response) of approximately 80% in the 24-month follow-up; this outcome shows that this immunotherapy was better than the standard treatment with BCG and/or intravesical chemotherapy, a fact that makes OncoTherad a beneficial therapeutic option to treat NMIBC. This assay has also shown significantly increased immunoreactivity to TLR4, TRIF, IRF-3, INF-γ and iNOS in primary lymphoid follicle regions in the bladder, as well as significantly low adverse effects [66,68].

Based on the action mechanism of the OncoTherad-based immunotherapy and on the role played by the interferon signaling pathway in controlling Covid-19 infection, it was possible identifying the potential action of this immunotherapy to control SARS-CoV-2 infection in 5 NMIBC patients (CAAE: 93619718.7.0000.5404; Brazilian Clinical Trial RBR-6sqwd2), who tested positive for Covid-19.

The most emblematic case among the herein evaluated ones was the case report about a 78-year-old Brazilian male patient with history of systemic arterial hypertension, anterior myocardium revascularization and former smoker, who was recruited for the OncoTherad clinical trial to attend BCG-refractory or relapsed high grade-NMIBC. After interrupting the treatment for a certain period-of-time due to an international trip, he arrived at the Hospital with Covid-19 symptoms. All analyses have shown advanced Covid-19 stage; the patient was treated with OncoTherad, in association with chemotherapy support and ventilation; he was discharged at the 10th hospitalization day [10]. Besides all these data, patient’s clinical improvement was featured by high free fatty acid, acylcarnitine and phosphatidylglycerol (PG) levels (based on deep leaning technique carried out through mass spectrometry), which reinforced previous findings about serum metabolome [69] in recovered SARSCoV patients.

The other 4 patients subjected to the clinical protocol based on OncoTherad application to NMIBC cases who tested positive for Covid-19 have shown similar behavior to that of the patient described in the aforementioned case report. The evolution and recovery of patients who received OncoTherad immunotherapy took shorter time (10 days) than that observed for patients subjected to the standard treatment. Thus, if one takes into consideration patients’ age and comorbidities, it is possible assuming that the immunotherapy with OncoTherad has played a protective role in stopping the evolution of Covid-19 infection to the most severe stages of it and in promoting patients’ fast recovery without the need of hospitalization in Intensive Care Units [10]. Actually, patients presenting these conditions have been recruited to receive OncoTherad therapy in a Clinical trial in progress at Hospital Municipal de Paulínia in São Paulo, Brazil [70-73].

Conclusions

Since the outbreak of Covid-19 in December 2019, different new and old pharmaceutical materials have been tested in a large number of clinical trials focused on assessing their effectiveness against SARS CoV-2 infection. The first strategy lied on using anti-rheumatic drugs, likely due to their immunomodulating or immunosuppressing properties; however, many aspects related to their applications should be taken into consideration. Zhong et al. [13] have pointed out aspects such as hyper-inflammation and viral replication (hyper-activation of immune response leads to tissue damage and organ failure), immunomodulation treatment timing (important and direct evidences deriving from randomized controlled trial are required to determine the appropriate treatment timing for patients with Covid-19) and pharmacokinetics of oral medications (administration route is an important factor).
Immunotherapy is an effective alternative treatment against Covid-19; the main methods mentioned in the current study were capable of enhancing clinical outcomes in Covid-19 infected patients. Several trials yet in progress will certainly help better understanding the potential action of immunomodulation treatments on Covid-19 infection associated with hyper-inflammation processes.

Acknowledgments

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