**Management of SARS-CoV-2 infection: recommendations of the Polish Association of Epidemiologists and Infectiologists as of April 26, 2021**

Robert Flisiak1, Andrzej Horban2,3, Jerzy Jaroszewicz4, Dorota Kozielewicz5, Agnieszka Mastalerz-Migas6, Radosław Owczuk7, Miłosz Parczewski8, Małgorzata Pawłowska5, Anna Piekarska9, Krzysztof Simon10, Krzysztof Tomasiewicz11, Dorota Zarębska-Michaluk12

1. Department of Infectious Diseases and Hepatology, Medical University of Białystok, Białystok, Poland
2. Department of Infectious Diseases for Adults, Medical University of Warsaw, Warsaw, Poland
3. Hospital for Infectious Diseases in Warsaw, Warsaw, Poland
4. Department of Infectious Diseases and Hepatology, Medical University of Silesia, Katowice, Poland
5. Department of Infectious Diseases and Hepatology, Faculty of Medicine, Ludwik Rydygier Collegium Medicum in Bydgoszcz of the Nicolaus Copernicus University in Toruń Bydgoszcz, Poland
6. Department of Family Medicine, Wroclaw Medical University, Wrocław, Poland,
7. Department of Anesthesiology and Intensive Care Unit, Medical University of Gdańsk, Poland,
8. Department of Infectious, Tropical Diseases and Acquired Immunodeficiency, Pomeranian Medical University, Szczecin, Poland
9. Department of Infectious Diseases and Hepatology, Medical University of Łódź, Łódź, Poland,
10. Department of Infectious Diseases and Hepatology, Medical University of Wrocław, Wrocław, Poland
11. Department of Infectious Diseases and Hepatology, Medical University of Lublin, Lublin, Poland
12. Department of Infectious Diseases, Jan Kochanowski University, Kielce, Poland

**Short title: Recommendations of management in SARS-CoV-2 infections, PTEiLChZ, v. 26-04-2021**

**Corresponding author:** Prof. Robert Flisiak,MD, PhD, Department of Infectious Diseases and Hepatology, Medical University in Białystok, 15-540 Białystok, ul. Żurawia 14, tel: +48 85 7416921, robert.flisiak1@gmail.com

**Conflict of interest:** none declared

**Introduction**

In early 2020 the world found out about the novel beta coronavirus which in December 2019 caused an epidemic of pneumonia in Wuhan, China. The virus was officially named severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) and the disease received the name of coronavirus disease 2019 (COVID-19) [1]. The outbreak has spread rapidly in many countries all over the world and on March 11, 2020 the World Health Organization (WHO) announced a pandemic of COVID-19. In the light of the increasing number of affected patients globally, there was an urgent need to establish the management of this emerging disease. In Poland, the first recommendations on the diagnosis and treatment of COVID-19 were issued by the Polish Association of Epidemiologists and Infectiologists on March 31, 2020 [2]. Those practical guidelines were created based on the previous experience with two known human pathogenic coronaviruses, SARS-CoV-1 and Middle East respiratory syndrome coronavirus (MERS-CoV), due to similarities and newly available data on SARS-CoV-2. The pathogenesis, transmission routes, methods of protecting against the infection, diagnostic procedures including molecular and serological methods, the definition of a suspected, probable and confirmed case of COVID-19 along with clinical symptoms of the disease were described. However, the most attention has been paid to specific management in the different clinical presentations of COVID-19, including asymptomatic or mild, the stable stage with respiratory and/or systemic symptoms, the unstable stage of respiratory failure, and the critical condition with acute respiratory distress syndrome (ARDS). The recommendations have been updated twice due to new data becoming available, especially in the field of pharmacological treatment. The most important change in the first Annex from June 2020 was the recommendation to use remdesivir (RDV) at an earlier stage of COVID-19, as a consequence of the Emergency Use Authorization issued by the United States Food and Drug Administration (FDA) and European Medicines Agency (EMA) [3, 4]. The second amendment from October 2020 solidified the recommendation to apply RDV at the stage of viral replication and tocilizumab and/or glucocorticosteroids at the unstable stage of cytokine storm based on the results from clinical trials and real-world experience studies [5, 6, 7, 8]. Moreover, several drugs were delisted due to evidence of therapeutic ineffectiveness [9]. After one year of the pandemic, despite global efforts of medical staff, we still do not have control of the disease. New hopes were raised with the development of vaccines; however, to date, the vaccination coverage added to the number of people who have experienced SARS-CoV-2 infection is insufficient to achieve herd immunity. In the face of emerging data and growing clinical experience, new recommendations on the management of COVID-19 are needed.

**Etiology**

Coronaviruses include a wide variety of viruses that cause disease in humans and animals. So far, seven species of coronaviruses pathogenic for humans, mainly for children, are known. In 2002 and 2012, two new species that are highly pathogenic to humans were identified – currently known as SARS-CoV-1 and MERS-CoV. In December 2019 in Wuhan, China, several cases of pneumonia were described, and in January 2020 a new type of betacoronavirus was confirmed as an agent. The genome of the new coronavirus named SARS-CoV-2 showed 79% similarity with SARS-CoV-1 and 50% with MERS-CoV. Like all coronaviruses, it has a positive RNA strand. The full genome sequencing and phylogenetic analysis were done at the beginning of 2020.

**Emerging variants**

Novel emerging variants of SARS CoV-2 are of significant concern due to changes in the binding affinity between the viral spike (S) protein and the ACE-2 receptor on the host cells, which may change the infectivity and disease course [10]. It should be noted that viral replicase introduced errors allow for the emergence of random mutations in the viral genome. Emergence of strains with higher replicative capacity and infectivity is a natural phenomenon which provides advantage to the novel strains with increased transmissibility [11]. Variants with increased infectivity and most likely pathogenicity are becoming the dominant circulating strains and are characterized by numerous point mutations within the spike region. So far several such lineages have been identified, the key ones being the B.1.1.7 lineage first detected in the United Kingdom, B.1.351 originating from South Africa and the P.1 lineage from Brazil [12, 13]. These variants of concern are likely the cause of the epidemic waves observed in multiple countries, and the likelihood of further spread is assessed as high, which may lead to an increase in the hospitalization rates and pressure on the health systems [14]. Also, vaccine responses, infectivity periods and mortality have differed depending on the variant [15]. Molecular surveillance on the emergence and transmission of the novel variants may be achieved by scaling up the sequencing capacity and systematic population sampling. Knowledge on the genomic variation may provide the basis for understanding of the spatio-temporal transmission patterns, vaccine efficacy and differences in the clinical course of the disease.

**Pathogenesis**

SARS-CoV-2 spreads through respiratory droplet transmission, contaminated objects and surfaces and enters host cells via the functional cell receptor angiotensin converting enzyme II (ACE-2), with the support of transmembrane serine protease 2 (TMPRSS2) and integrins. The S proteins of SARS-CoV-2 bind to the ACE-2 receptor, localized *inter alia* on alveolar epithelial type 2 (AT2) cells. ACE-2 receptors are moreover highly expressed on the tubular epithelium of the upper esophagus, absorptive enterocytes of the ileum and colon, in the kidney, heart, pancreas, and, what is pathogenically important, in arterial and venous endothelial cells and arterial smooth muscle cells [16]. Immune responses induced by SARS-CoV-2 are biphasic [16, 17]. During the incubation period and early stages of the infection a specific adaptive immune response can eliminate the infection and blocks progression to more severe stages of this disease. It has been confirmed that susceptibility to various infectious diseases is associated with specific HLA haplotypes. In patients with impaired immune responses (e.g. older people with underlying diseases, young persons with impaired or dysregulated B-cell and T-cell immune responses) the virus will rapidly replicate with massive destruction of all cells with high ACE2 expression, e.g. alveolar epithelial type 2 (AT2) in lungs. This induces innate inflammation in the lungs, mediated by proinflammatory granulocytes and macrophages. Activated macrophages release over 100 cytokines (e.g. IL-1, IL-6, and TNFα) and chemokines (CXCL10 and CCL2) into the bloodstream (cytokine release storm-CRS) [18]. These events, endothelial dysfunction (endotheliitis) and procoagulant state lead to the life-threatening acute respiratory distress syndrome (ARDS) and impaired function of other organs [19].

**Clinical picture of the disease**

The median incubation period for COVID-19 is 5.1 days, and 97.5% of all infections develop within 11.5 days [20]. Infection with SARS-CoV-2 has a broad clinical spectrum from asymptomatic, oligosymptomatic to moderate or severe disease with multiorgan failure. The rate of asymptomatic infections ranges from 27 to 40%. Overall, approximately 90% of infections are uncomplicated, oligosymptomatic or with moderate symptoms not leading to hospitalization [21, 22]. Older age and co-morbidities, such as hypertension, chronic cardiac, pulmonary or kidney disease, diabetes, obesity as well as immunosuppression, and cancer are risk factors for severe disease and death [23].

Stage 1 of the disease includes asymptomatic and mildly symptomatic patients with SpO2≥94% on room air not requiring hospitalization. The symptoms of COVID-19 present at illness onset vary and most of them are non-specific, but mainly include: fever, chills, cough, shortness of breath or difficulty breathing, fatigue, muscle or body aches, headache, sore throat, runny nose, and conjunctivitis. Some patients also experience gastrointestinal symptoms, such as anorexia, nausea, vomiting diarrhea, and abdominal pain. New loss of smell (anosmia) or taste (ageusia) may be the sole presenting symptom, especially among women and younger or middle-aged patients [24]. Atypical symptoms can precede the respiratory symptoms of COVID-19 in elderly adults. Diarrhea with dehydration, fluctuating temperature with hypothermia, delirium and a fall are the most common. Since SARS-CoV-2 infection induces severe fatigue, falls can be common in older adults with little or no risk of falls [25].

In stage 2 patients present clinical and radiological signs of mild to moderate interstitial pneumonia with SpO2<94% on room air. In some patients fever, dry cough, shortness of breath, fatigue and other extrapulmonary symptoms are still present.

Stage 3 is a severe form of disease with respiratory failure (dyspnea, respiratory frequency over 30/min, SpO2<90% on room air, and/or lung infiltrates more than 50% of the lung field within 24–48 hours) and cytokine storm syndrome [26]. A dangerous phenomenon is the development of respiratory failure without subjective perception of dyspnea (“silent hypoxemia”). In these cases, hypocapnia caused by compensatory hyperventilation is an accompanying finding [27]. Skeletal muscle damage associated with increased CK levels and neurological manifestations involving both central and peripheral nervous systems are common among patients with severe infection [28] Acute cerebrovascular disease (ischemic stroke, intracerebral hemorrhage, deep cerebral venous thrombosis), encephalitis, Guillain-Barre syndrome, vision impairment, dizziness, impaired consciousness, ataxia, and seizure are reported [29]. Cardiovascular manifestations of COVID-19 (myocardial ischemia, myocarditis, cardiomyopathy, ventricular arrhythmias, and hemodynamic instability) are observed but at lower frequency than in critically ill patients [30]. The typical course of severe disease includes the appearance of overt dyspnea 6 days after the onset of first symptoms, hospitalization after a further 8 days and the need for tracheal intubation 10 days after hospitalization [31].

Stage 4 is a critical condition, which develops in around 5% of patients, with hypoxemic respiratory failure, septic shock, and/or multiple organ dysfunction (inter alia acute kidney injury, liver dysfunction, bleeding and coagulation dysfunction). In the ICU, venous and arterial thromboembolic events occur in 31% to 59% of patients [25]. Death is caused by progression to acute respiratory distress syndrome (ARDS) and multiorgan failure.

**Laboratory diagnostics**

Principles of molecular diagnostics

Currently, laboratory confirmation of SARS CoV-2 infection is based on molecular diagnostics: either detection of the viral nucleic acid or antigens from clinical samples. Typically, the highest sensitivity of molecular assays is observed with the initial symptoms of the disease, with genetic material often also detectable at the end of the pre-symptomatic phase of the infection [32].

Nucleic acid amplification testing (NAAT)

Since the beginning of the SARS CoV-2 epidemics nucleic acid amplification techniques allowing for the detection of genetic material of this virus have been rapidly introduced [33]. Polymerase chain amplification based techniques have been supplemented with loop mediated isothermal amplification (LAMA) and transcription mediated amplification (TMA) assays. NAAT tests, currently a gold standard of diagnostics, are designed to detect different genes of SARS CoV-2 including ORF, E, N, S RdRp and other regions. Molecular testing should be performed by a quality-controlled laboratory provider with the test targeting at least two (preferably three) regions of the viral genome. Detected genetic material is not a proof of infectivity, as it may be detectable after several weeks from onset of infection. Pooling of specimens may be useful for population diagnostics but should not be performed in the hospital setting for clinical purposes.

Sequencing of the SARS CoV-2 genome

To investigate molecular evolution of the virus genomic sequencing techniques may be implemented. This allows for the molecular surveillance of the emergence of mutations affecting transmissibility and pathogenicity of the virus, principally within the spike gene coding region and identification of newly emerging variants with pandemic potential.

Antigen tests

Detection of viral antigens allows for rapid (15-30 minutes) SARS CoV-2 diagnosis, improving efficacy of testing and detection of infectious cases and reducing the cost of molecular diagnostics. Sensitivity of the antigenic tests is lower comparted to the NAAT, and antigen based testing remains limited to the initial period of infection. Recommended (ECDC) diagnostic sensitivity of the antigen assays is ≥90% with specificity of ≥97% [34]. This testing is intended for symptomatic cases within the first 5 to 7 days after the onset of the disease. In asymptomatic cases a negative result of the antigen test following the exposure does not rule out the possibility of infection, but a positive antigen test should be regarded as confirmation of active infection. A positive antigenic test in immunocompromised individuals may be indicative of prolonged viral shedding and infectivity [35].

Serological testing

Serological tests reflect the antibody responses following exposure and may reflect long-term immunity against reinfection. However, protective antibody titers have not yet been established [33]. Data indicate that following an infection neutralizing antibodies in protective titers are present for 3-6 months, but it is unknown whether neutralizing function will vary for the emerging viral variants. Due to the serological window of 10-14 days tests detecting antibodies are not useful for the diagnosis of active SARS CoV-2 infection, with clinical utility as confirmation of infection in cases with clinically suspected COVID-19 and negative molecular test results. Serology may also be used for population surveillance studies, vaccine response analyses, to identify candidates for plasma donation, and in the diagnostic workup of postinflammatory syndromes. To detect antibody responses following vaccination serological assays detecting IgG antibodies against the S1 subunit of the spike protein should be used. Rapid immunochromatographic tests are of low sensitivity and specificity and should not be used in practice.

**Therapy**

In mildly symptomatic patients, only antipyretic, antitussive drugs and inhaled budesonide, are recommended. Vitamin D3 supplementation and low-molecular-weight heparin in prophylactic doses should be determined individually. In moderate or severe disease primary treatment is remdesivir, tocilizumab, dexamethasone or methylprednisolone. As a supportive therapy, low-molecular-weight heparins, rehydration, antibiotics (if necessary) and symptomatic medications are recommended. The vast majority of hospitalized patients require oxygen supplementation, and the flow depends on the demand. In patients with critically severe disease with respiratory failure, it is recommended to use high-flow oxygen therapy, mechanical ventilation or ECMO. So the treatment regimen depends on the stage of the disease and is detailed below.

* Stage 1

In the vast majority of patients (80%–90%), SARS‐CoV‐2 infection develops with no or mild symptoms and such individuals do not require hospitalization [9]. However, in some cases, this clinical form may represent stage 1 of the disease, preceding the fully symptomatic stage 2 (Table 1). Stage 1 patients followed up by a primary care physician usually require no treatment, and monitoring of their clinical status is a sufficient management option [9]. All adult patients staying at home should be registered in the Home Medical Care (Pulsocare) system with monitoring of oxygen saturation as measured by a pulse oximeter (SpO2), which should be at least 94%. All patients followed by the Pulsocare system who report measurements are looked after by consultants and physicians who will assess indications for hospitalization in the event of the occurrence of alarm values of measurements and may call the Medical Rescue Team. However, baseline oxygen saturation in patients with chronic respiratory diseases can be decreased, which does not necessarily mean that SARS‐CoV‐2-induced respiratory failure is progressing. If symptoms arise, patients may require antipyretic agents (non-steroidal anti-inflammatory drugs or paracetamol is most effective). It is recommended to supplement vitamin D3 in each patient, in accordance with the recommendations for the Polish population [9]. There is evidence indicating a risk of a more severe course of the disease in patients with vitamin D3 deficiency, with a low risk associated with the use of this agent [36]. It is recommended to use antitussive drugs in patients with severe cough (making it difficult to talk and sleep). In severe cases, the use of agents containing codeine can be considered, although its abuse and misuse may cause respiratory depression. In adult patients with symptomatic COVID-19, it is recommended to administer budesonide in a daily dose of 2 x 800 µg by inhalation [37]. In patients who are chronically bedridden and those with risk factors for deep vein thrombosis and/or pulmonary embolism, a prophylactic dose of low-molecular-weight heparin is recommended [38, 39, 40]. However, there is insufficient evidence to support or not recommend the routine use of antiplatelet drugs in COVID-19 [41]. Of note, patients with stage 1 disease should not be treated with glucocorticosteroids. These drugs have been shown to be ineffective in patients who do not require oxygen therapy. Furthermore, using glucocorticosteroids too early may increase the viral replication and, hence, worsen patient prognosis [8].

Treatment with antibiotics or anti‐influenza drugs is not recommended in SARS‐CoV‐2 infection, unless it is required because of another medical condition. Home oxygen therapy is not recommended in the acute phase of the disease due to the risk of a sudden, life-threatening deterioration. The need to use oxygen therapy in the treatment of COVID-19 is an absolute indication for hospitalization.

* Stage 2

Patients with increasing dyspnea and SpO2 <94% need oxygen therapy, which requires hospitalization. As part of the thromboembolic prevention, all hospitalized patients should receive low-molecular-weight heparin, usually in prophylactic doses, which can be increased in justified cases. If symptoms have appeared within 7 days, the patient requires oxygen therapy, and if SpO2 is below 94%, antiviral therapy is indicated. After 7 days of the disease, antiviral therapy becomes useless due to the disappearance of replication of the virus. The only currently approved antiviral drug with proven efficacy against SARS-CoV-2 in adults and children over 12 years of age is remdesivir, which should be administered by intravenous infusion at a dose of 200 mg on the first day, and 100 mg on the following four days [5, 6]. Extending the treatment period beyond 5 days does not improve the effectiveness of the therapy [42]. The main contraindication to the use of remdesivir is renal failure with GFR <30 ml/min. Remdesivir should be discontinued if alanine aminotransferase levels exceed 5 times the upper limit of normal [43]. In the case of lack of availability of remdesivir or contraindications to its use, the use of convalescent plasma may be considered. However, its effectiveness has not yet been clearly confirmed [44]. It is assumed that the therapeutic effect can be expected only when plasma with a high antibody titer (at least 1: 500) is used. In the event of no clinical improvement despite the use of antiviral therapy, dexamethasone in a daily dose of 4-8 mg may be considered [8]. However, it should not be used in the first week of illness due to the risk of an increase in viral replication. Antibiotics can be considered in case of bacterial superinfection.

* Stage 3

Clinical deterioration usually occurring at the beginning of the 2nd week of the disease may be a sign of a cytokine storm commencing. Due to the increasing shortness of breath and the reduction of SpO2 below 90%, low-flow oxygen therapy ensuring the supply of oxygen up to a maximum of 15 l/min is no longer sufficient. Patients require high-flow oxygen therapy, reaching 60 l/min. In this phase of the disease, it is crucial to catch the onset of the cytokine storm. The presence of an increase in IL-6 concentration above 100 pg/ml justifies the administration of tocilizumab, an inhibitor of IL-6 receptors, which significantly reduces the risk of mechanical ventilation and death [7, 45]. Tocilizumab should be administered as an intravenous infusion over 1 hour as a single dose based on body weight up to 800 mg (see table 1 for dosing details). In the event of no effect, another infusion may be given after 8-12 hours. Tocilizumab should not be administered in patients with a neutrophil count below 2x109/l, a platelet count below 50x103/l or alanine aminotransferase levels exceeding 5 times the upper limit of normal [46]. At this stage of the disease, patients should still receive low-molecular-weight heparin and glucocorticosteroids, and antibiotics if necessary due to bacterial infections.

* Stage 4

Further deterioration of the patient's condition means the development of the stage in which acute respiratory distress syndrome (ARDS) occurs, requiring the use of high-flow oxygen therapy and in most such cases also tracheal intubation and mechanical ventilation, requiring the patient to be transferred to the intensive care unit. At this stage of the disease, 6-8 mg of dexamethasone is recommended, although higher doses of glucocorticoids are being studied. There are no indications for introduction or continuation of remdesivir therapy in intensive care unit patients. A lung-protective ventilation strategy should be applied – that means use of low tidal volumes, appropriate values of positive end-expiratory pressure (PEEP), adjustment of inspired oxygen fraction to partial pressure of oxygen in the blood (PaO2) and arterial oxygen saturation (SaO2) values. The use of ventilation in the prone position provides good results in some patients unresponsive to conventional methods of respiratory therapy [47]. Unfortunately, the mortality rate among COVID-19 patients requiring mechanical ventilation is as high as 67% [48]. The use of veno-venous extracorporeal membrane oxygenation (VV ECMO) is indicated only in selected individuals, and its use is limited to expert centers with appropriate experience and technical capabilities. It may be beneficial in patients with ARDS (moderate or severe) diagnosed prior to introduction of mechanical ventilation, ventilated no longer than 7 days, with severe disturbances in lung gas exchange despite optimal conventional ventilation and failed attempts of oxygenation improving methods (e.g. prone position, neuromuscular blockade). It is also used as a bridging therapy to lung transplantation. There are also numerous contraindications to this therapy [49].

**Treatments of unproven efficacy**

A number of medications have been considered for use in COVID-19 and a few as a prophylaxis, but only a small proportion have been proved in clinical randomized controlled trials (RCTs) and/or in observational studies. The following medications failed to have their efficacy and safety confirmed or trials are ongoing.

Antiviral medications

* Favipiravir is a potent inhibitor of many RNA viruses. Preliminary study results may suggest that the drug can shorten the time to recovery for patients with COVID-19, help achieve viral clearance early so as to positively impact disease transmission in the community, and reduce the health burden by shortening the length of stay at the hospital [50]. Disadvantages include severe side effects, such as anaphylactic shock and pneumonia.
* Anti-influenza medications and non-specific antivirals. A limited number of trials have evaluated oseltamivir, amantadine, rimantadine, zanamivir, and acyclovir. No significant benefit for repurposed antivirals was observed and oseltamivir may increase mortality [51].
* Ivermectin. Some clinicians concluded that this anti-parasitic drug could significantly reduce viral load and accelerate recovery in patients with mild and moderate cases [52]. The WHO and FDA advise against this anti-parasitic drug to treat COVID-19.
* Anti-HIV medications have been found ineffective in SARS-CoV-2 infection, but a number of trials are ongoing.
* Interferons. Subcutaneous injection of interferon β-1a showed no significant improvement in time to clinical response, but the overall mortality at 28 days was reduced.
* Intravenous immunoglobulin (IVIg) has been used as an adjuvant therapy or in a concentrated (hyperimmune) form. The potential utility of IVIg for the treatment of SARS-CoV-2 is unknown at this time.
* Monoclonal anti-SARS-CoV-2 antibodies. Bamlanivimab with etesevimab and the combination of casirivimab plus imdevimab are anti-severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) monoclonal antibodies [53, 54]. There are currently insufficient data to recommend either for or against their use.

Anti-inflammatory medication

* Data about use of non-steroidal anti-inflammatory drugs (NSAIDs) are controversial. There are reports that ibuprofen blocks the diffuse inflammation produced by SARS CoV-2 and thus might prevent COVID-19 complications [55]. However, NSAIDs do have fairly well-established side effects, so it is reasonable to use these medicines to treat the symptoms of mild COVID19, but to minimize the dose and duration of treatment where possible.
* Anticytokine medications – apart from tocilizumab, a number of immune antagonists have been studied in clinical trials. So far there is no confirmation of their efficacy in COVID-19, including anakinra, which failed to improve clinical outcomes in patients with mild-to-moderate COVID-19 [56].
* Further studies are needed to assess the suitability of early inhaled budesonide administration [56]

Unproven preventive – Mouthwash and nasal spray

Some recommend gargling with certain mouthwashes to kill viruses in the throat and temporarily reduce the risk of shedding viruses. Antiviral nasal sprays work by coating viral particles within the nose to block their entrance into cells within the body. There are no clinical studies proving that it is effective in preventing human-to-human transmission.

**Late sequelae of COVID-19**

Post-acute COVID-19 syndrome (PC19, long COVID) refers to persistence of symptoms or organ dysfunctions following the acute phase of COVID-19. Although the incidence, clinical picture and therapeutic recommendation are yet to be established, PC19 refers to abnormalities beyond 4 weeks after onset of COVID-19 [57, 58]. It can be divided into subacute within 4-12 weeks and chronic beyond 12 weeks. The prevalence of PC19 varies between 30% and 80% and depends on the methodology of the study [59]. The most prevalent complications are pulmonary (dyspnea, hypoxia, reduced diffusion capacity, persistent inflammatory lesions and/or fibrosis in computed tomography (CT)); hematologic (thromboembolic events, anemia); cardiovascular (palpitations, dyspnea, chest pain, arrhythmias, myocardial fibrosis/scarring); neuropsychiatric (chronic fatigue, myalgia, headache, olfactory/gustatory dysfunctions, anxiety, depression, sleep disturbances and posttraumatic stress disorder). Multiple laboratory abnormalities including reduced eGFR, increased liver function tests, and hyperglycemia were reported 3-6 months following the acute phase of COViD-19 [59]. Pediatric inflammatory multisystem syndrome (PIMS/MIS-C) is a separate post-COVID-19 condition in children and young adults with specific diagnostic criteria [60].

Preliminary results of the Polish prospective study SILCOV-19 (the Silesian Complications of COVID-19 Database) show that after a median of 90 days of COVID-19 onset the most prevalent symptoms are: fatigue (~50%), reduced lung diffusion capacity (~30%), ongoing inflammatory lesions in HRCT (35% of hospitalized, 10% of non-hospitalized patients), anxiety (18-24%), sleep disturbances (19-34%), depression (5-15%) and bradyarrhythmias (13-16%). The most prevalent laboratory abnormalities are anemia (48%) and neutropenia (28%), NT-proBNP>125 pg/mL (26% hospitalized, 9% non-hospitalized), increased D-dimers (20% hospitalized, 9% non-hospitalized), and hyperbilirubinemia (~10%). [61]

Clinical guidelines for PC19 lack strong evidence and over 40 prospective trials are underway. There is no evidence for routine screening in asymptomatic subjects. Symptom assessment should be performed 4-6 weeks and 12 weeks after discharge in patients with moderate to severe COVID-19. Diagnostics should be symptom oriented. The objective methods for pulmonary evaluations are the Borg Dyspnoea Scale, home pulse oximetry, 6-minute walk test (6MWT), chest X-ray, pulmonary function tests and, if recommended by a specialist, chest CT/HRCT (usually not earlier that 3 months after COVID-19). For cardiac symptoms: 24-hour Holter monitoring and echocardiogram, in selected cases cardiac MRI or CT coronary angiogram. Some authors recommend routine screening for anxiety, depression and sleep disorders. The need for laboratory testing depends on abnormalities during hospitalization. It is important to follow patients with acute kidney or liver injury, a thromboembolic event, chronic cardiovascular, pulmonary, liver, or metabolic disorders. Currently, there is no recommendation for routine screening for coagulation following COVID-19.

In the post-COVID19 convalescence, rest, relaxation and pulmonary rehabilitation with breathing techniques play an important role. There is no evidence for routine thromboprophylaxis although high-risk patients may require prophylactic anticoagulation for up to 30 days after discharge (Table 2) [62]. Direct oral anticoagulants and low-molecular-weight heparins are preferred over vitamin K antagonists. The role of antiplatelet agents has not been established. The therapy of sequelae should be based on general recommendations. It is underlined that prompt initiation of anxiolytic and antidepressants improves prognosis. The benefits of corticosteroids or antifibrotic agents in post-COVID inflammatory lung disease in not confirmed. It is advisable to include subjects after moderate-severe COVID-19 in prospective studies evaluating PC19.

**Vaccination**

The Emergency Use Authorizations of the first COVID-19 vaccine was issued by the European Medicines Agency (EMA) in December 2020 [64, 65]. Currently available vaccines have been constructed using mRNA, vector and recombination technologies. Regardless of the technology used, all EMA-approved COVID-19 vaccines meet stringent efficacy and safety criteria when used in accordance with the summary of product characteristics (SmPC) [66, 67]. Recommendations for management of the most common doubts regarding vaccination against SARS-CoV-2 are presented below.

* The priority in vaccination against SARS-CoV-2 infections after vaccination of medical workers should be given to people over the age of 60, in order from the age of over 80. Introducing any groups before the end of vaccination of seniors is unacceptable; such action would be unjustified from the point of view of public health, inhumane and would increase the number of deaths, contributing to the extension of the duration of the epidemic with the consequence of prolonging paralysis of the health care system. People with cancer and diabetes under the age of 60, as well as parents of premature babies, should be considered as another group. After completion of vaccinations in the above‑mentioned groups, patients with other chronic diseases should be included. Only after vaccinating people over 60 years of age and chronically ill is it justified to take into account the social factors that determine the priority of vaccination.
* People who develop COVID-19 after the first dose of the vaccine may receive a second dose according to the schedule and time frame permitted by the vaccine SmPC. If this is not possible due to persistent disease symptoms, consideration should be given to restarting vaccination at least 6 months after the onset of disease symptoms, or after specific anti-spike antibodies have disappeared.
* There are no contraindications for vaccination after COVID-19. However, the persistence of the specific immune response following SARS-CoV-2 infection is unknown. Based on the current state of knowledge, it is possible to postpone the vaccination of people who have had COVID-19 up to 6 months. However, it should be emphasized that due to the emerging new research results, the recommended time interval between disease and vaccination will probably be extended.
* A contraindication to vaccination against SARS-CoV-2 is hypersensitivity to the active substance or to any of the excipients contained in the vaccine, or a history of any anaphylactic reaction in the past. It is permissible to vaccinate such a person with full protection against shock in hospital conditions, after prior notification of possible risks and obtaining written informed consent for vaccination.
* Chronic diseases, including cancer, or treatments, including immunosuppressive, are not contraindications to vaccination against SARS-CoV-2. However, vaccination is not recommended in the course of febrile diseases and in the period of exacerbation of chronic diseases. In people who have undergone organ or bone marrow transplantation, it is recommended not to vaccinate for 6 months after the procedure. The optimal timing for vaccination in patients scheduled for or undergoing immunosuppressive/biological therapy should be checked with appropriate specialist guidelines. Due to the risk of extravasation and bleeding, particular care should be taken when administering the vaccine to patients with severe coagulation disorders; it is advisable to compensate for any deficiencies before vaccination and apply pressure at the injection site after injection.
* According to CDC pregnant females with COVID-19 might be at increased risk of adverse pregnancy outcomes, such as preterm birth, compared with pregnant women without COVID-19 [68]. Before deciding whether to vaccinate, pregnant women must be informed about the lack of studies on the effects of vaccination against SARS-CoV-2 on fetal development. After considering the risk of infection, which can be dangerous in pregnancy, they should provide written informed consent to be vaccinated when deciding to immunize.
* Lactation is not a contraindication to vaccination against SARS-CoV-2. Due to the lack of research in this group, the decision to vaccinate should be made individually by the breastfeeding woman and confirmed by written informed consent to undergo vaccination.
* Vaccination against SARS-CoV-2 can be carried out with a 4-week interval from vaccines containing live microorganisms. In other cases the interval should be at least 2 weeks in order to differentiate possible post-vaccination reactions and avoid the risk of a reduced specific immune response to one of the vaccines.

**References**

1. Ciotti M, Angeletti S, Minieri M, et al. COVID-19 Outbreak: An Overview. Chemotherapy 2019; 64: 215-223.
2. Flisiak R, Horban A, Jaroszewicz J, et al. Management of SARS-CoV-2 infection: recommendations of the Polish Association of Epidemiologists and Infectiologists as of March 31, 2020. Pol Arch Intern Med. 2020; 130: 352-357
3. Flisiak R, Horban A, Jaroszewicz J, et al. Management of SARS-CoV-2 infection: recommendations of the Polish Association of Epidemiologists and Infectiologists. Annex no. 1 as of June 8, 2020. Pol Arch Intern Med. 2020; 130: 557-558.
4. Ye M, Fu D, Ren Y, et al. Treatment with convalescent plasma for COVID-19 patients in Wuhan, China. J Med Virol. 2020; 92: 1890-1901.
5. Beigel JH, Tomashek KM, Dodd LE, et al. Remdesivir for the Treatment of Covid-19 - Final Report. N Engl J Med. 2020; 383: 1813-1826.
6. Flisiak R, Zarębska-Michaluk D, Berkan-Kawińska A, et al. Remdesivir-based therapy improved the recovery of patients with COVID-19 in the multicenter, real-world SARSTer study. Pol Arch Intern Med. 2021; 131: 103-110.
7. Tomasiewicz K, Piekarska A, Stempkowska-Rejek J, et al. Tocilizumab for patients with severe COVID-19: a retrospective, multi-center study. Expert Rev Anti Infect Ther. 2021; 19: 93-100.
8. Horby P, Lim WS, Emberson JR, et al. Dexamethasone in Hospitalized Patients with Covid-19. N Engl J Med. 2021; 384: 693-704.
9. Flisiak R, Parczewski M, Horban A, et al. Management of SARS-CoV-2 infection: recommendations of the Polish Association of Epidemiologists and Infectiologists. Annex no. 2 as of October 13, 2020. Pol Arch Intern Med. 2020; 130: 915-918.
10. Volz E, Mishra S, Chand M, et al. Transmission of SARS-CoV-2 Lineage B. 1.1. 7 in England: Insights from linking epidemiological and genetic data. medRxiv 2020.12.30.20249034
11. European Centre for Disease Prevention and Control (ECDC). Sequencing of SARS-CoV-2. 23 December 2020, https://www.ecdc.europa.eu/sites/default/files/documents/sequencing-of-SARS-CoV-2.pdf. Accessed April 2, 2021.
12. Rambaut A, Loman N, Pybus O, et al. Preliminary genomic characterisation of an emergent SARS-CoV-2 lineage in the UK defined by a novel set of spike mutations. https://virological.org/t/preliminary-genomic-characterisation-of-an-emergent-sars-cov-2-lineage-in-the-uk-defined-by-a-novel-set-of-spike-mutations/563. Accessed April 2, 2021.
13. Rambaut A, Holmes EC, O’Toole Á, et al. A dynamic nomenclature proposal for SARS-CoV-2 lineages to assist genomic epidemiology. Nat Microbiol. 2020; 5: 1403–1407.
14. Davies NG, Barnard RC, Jarvis CI, et al. Estimated transmissibility and severity of novel SARS-CoV-2 Variant of Concern 202012/01 in England. medRxiv 2020.12.24.20248822
15. Davies NG, Jarvis CI, CMMID COVID-19 Working Group. et al. Increased mortality in community-tested cases of SARS-CoV-2 lineage B.1.1.7. Nature. 2021. https://doi.org/10.1038/s41586-021-03426-1 . Accessed April 6, 2021.
16. Hoffmann M, Kleine-Weber H, Schroeder S, et al. SARS-CoV-2 cell entry depends on ACE2 and TMPRSS2 and is blocked by a clinically proven protease inhibitor. Cell. 2020; 181: 271-280.
17. Attaway AH, Scheraga RG, Bhimraj A, et al. Severe covid-19 pneumonia: pathogenesis and clinical management. BMJ. 2021; 372: n436.
18. Trougakos IP, Stamatelopoulos K, Terpos E, et al. Insights to SARS-CoV-2 life cycle, pathophysiology, and rationalized treatments that target COVID-19 clinical complications. J Biomed Sci. 2021; 28: 9.
19. Becker RC. COVID-19 update: Covid-19-associated coagulopathy. J Thromb Thrombolysis. 2020; 50: 54-67.
20. Lauer SA, Grantz KH, Bi Q, et al. The incubation period of coronavirus disease 2019 (COVID-19) from publicly reported confirmed cases: estimation and application. Ann Intern Med. 2020; 172: 577-582.
21. Pollán M, Pérez-Gómez B, Pastor-Barriuso R, et al. Prevalence of SARS-CoV-2 in Spain (ENE-COVID): a nationwide, population-based seroepidemiological study. Lancet. 2020; 396: 535-544.
22. Lavezzo E, Franchin E, Ciavarella C, et al. Suppression of a SARS-CoV-2 outbreak in the Italian municipality of Vo’. Nature. 2020; 584: 425-429.
23. Gupta S, Hayek SS, Wang W, et al. Factors associated with death in critically ill patients with coronavirus disease 2019 in the US. JAMA Intern Med. 2020; 180: 1-12.
24. Wiersinga WJ, Rhodes A, Cheng AC, et al. Pathophysiology, Transmission, Diagnosis, and Treatment of Coronavirus Disease 2019 (COVID-19): A Review. JAMA. 2020; 324: 782-793.
25. Blain H, Rolland Y, Benetos A, et al. Atypical clinical presentation of COVID‑19 infection in residents of a long‑term care facility. Eur Geriatr Med. 2020; 11: 1085-1088.
26. Umakanthan S, Sahu P, Ranade AV, et al. Origin, transmission, diagnosis and management of coronavirus disease 2019 (COVID-19). Postgrad Med J. 2020; 96: 753-758.
27. Xie J, Tong Z, Guan X, et al. Critical care crisis and some recommendations during the COVID-19 epidemic in China. Intens Care Med. 2020; 46: 837-840.
28. Mao L, Jin H, Wang M, et al. Neurological manifestation of hospitalized patients with coronavirus disease 2019 in Wuhan, China. JAMA Neurol. 2020;77: 683-690.
29. Helms J, Kremer S, Merdji H, et al. Neurologic features in severe SARS-CoV-2 infection. N Engl J Med. 2020; 382: 2268-2270.
30. Long B, Brady WJ, Koyfman A, et al. Cardiovascular complications in COVID-19. Am J Emerg Med. 2020; 38: 1504-1507.
31. Bouadma L, Lescure FX, Lucet JC, et al. Severe SARS-CoV-2 infections: practical considerations and management strategy for intensivists. Intens Care Med. 2020; 46: 579–582.
32. Kucirka LM, Lauer SA, Laeyendecker O, et al. Variation in false-negative rate of reverse transcriptase polymerase chain reaction-based SARS-CoV-2 tests by time since exposure. Ann Intern Med. 2020; 173: 262-267.
33. World Health Organization. Laboratory testing for coronavirus disease (‎‎COVID-19)‎‎ in suspected human cases: interim guidance. https://apps.who.int/iris/handle/10665/331501. Accessed April 2, 2021.
34. European Centre for Disease Prevention and Control. Options for the use of rapid antigen tests for COVID-19 in the EU/EEA and the UK. https://www.ecdc.europa.eu/en/publications-data/options-use-rapid-antigen-tests-covid-19-eueea-and-uk. Accessed April 2, 2021.
35. World Health Organization. Antigen-detection in the diagnosis of SARS CoV-2 infectios using rapid immunoassays. Interim guidance 11.09.2020. https://www.who.int/emergencies/diseases/novel-coronavirus-2019/serology-in-the-context-of-covid-19. Accessed April 2, 2021.
36. Castillo ME, Costa LME, Barrios JMV, et al. Effect of calcifediol treatment and best available therapy versusbest available therapy on intensive care unit admission and mortality among patients hospitalized for COVID-19: A pilot randomized clinical study. J Steroid Biochem Mol Biol. 2020; 203: 105751.
37. Ramakrishnan S, Nicolau Jr DV, Langford B, et al. Inhaled budesonide in the treatment of early COVID-19 (STOIC): a phase 2, open-label, randomised controlled trial. Lancet Respir Med. 2021. doi.org/10.1016/S2213-2600(21)00160-0. Online ahead of print.
38. Shi C, Wang C, Wang H, et al. Clinical observations of low molecular weight heparin in relieving inflammation in COVID-19 patients: A retrospective cohort study. medRxiv 2020.03.28.20046144.
39. Tang N, Bai H, Chen X, et al. Anticoagulant treatment is associated with decreased mortality in severe coronavirus disease 2019 patients with coagulopathy. J Thromb Haemost. 2020; 18: 1094-1099.
40. Ayerbe L, Risco C, Ayis S. The association between treatment with heparin and survival in patients with Covid-19. J Thromb Thrombolysis. 2020; 50: 298-301.
41. Salah HM, Mehta JL. Meta-Analysis of the Effect of Aspirin on Mortality in COVID-19. Am J Cardiol. 2021; 142: 158–159.
42. Goldman JD, Lye DCB, Hui DS, et al. Remdesivir for 5 or 10 days in patients with severe Covid‑19. N Engl J Med. 2020; 383: 1827-1837.
43. Veklury – summary of product characteristics. https://www.ema.europa.eu/en/documents/product‑information/veklury‑epar‑product‑information\_pl.pdf. Accessed March 31, 2021.
44. Moniuszko-Malinowska A, Czupryna P, Zarębska-Michaluk D, et al. Convalescent plasma transfusion for the treatment of COVID-19-experience from Poland: a multicenter Study. J Clin Med. 2020; 10: E28.
45. Flisiak R, Jaroszewicz J, Rogalska M, et al. Tocilizumab improves the prognosis of COVID-19 in patients with high IL-6. J Clin Med. 2021; 10, 1583.
46. RoActemra – summary of product characteristics. https://www.ema.europa.eu/en/documents/product‑information/roactemra‑epar‑product‑information\_pl.pdf. Accessed March 31, 2021.
47. Wujtewicz MA, Dylczyk-Sommer A, Aszkiełowicz A, et al. COVID-19 – what should anaethesiologists and intensivists know about it? Anaesthesiol Intensive Ther. 2020; 52: 34-41.
48. Flisiak R. Mortality due to COVID-19, SARSTer database. http://www.pteilchz.org.pl/wp-content/uploads/2021/01/%C5%9Bmiertelno%C5%9B%C4%87-w-Polsce-26-01-2021.pdf. Accessed March 31, 2021.
49. Badulak J, Antonini MV, Stead CM, et al. CMO for COVID-19: Updated 2021 Guidelines from the Extracorporeal Life Support Organization (ELSO), ASAIO J. 2021. doi:10.1097/MAT.0000000000001422. Online ahead of print.
50. Dabbous HM, Abd-Elsalam S, El-Sayed MH. et al. Efficacy of favipiravir in COVID-19 treatment: a multi-center randomized study. Arch Virol. 2021; 166: 949–954
51. Ravikirti, Roy R, Pattadar C, et al. Ivermectin as a potential treatment for mild to moderate COVID-19–A double blind randomized placebo-controlled trial. medRxiv 2021.01.05.21249310
52. Mancilla-Galindo J, García-Méndez JÓ, Márquez-Sánchez J, et al. All-cause mortality among patients treated with repurposed antivirals and antibiotics for COVID-19 in Mexico City: a real-world observational study. EXCLI Journal. 2021; 20: 199–222.
53. Gottlieb RL, Nirula A, Chen P, et al. Effect of Bamlanivimab as Monotherapy or in Combination With Etesevimab on Viral Load in Patients With Mild to Moderate COVID-19: A Randomized Clinical Trial. JAMA. 2021; 325: 632-644.
54. Lundgren JD, Grund B, Barkauskas CE, et al. A neutralizing monoclonal antibody for hospitalized patients with COVID-19. N Engl J Med. 2020: 384: 905-914.
55. Chen JS, Madel Alfajaro M, Chow RD, et al. Non-steroidal anti-inflammatory drugs dampen the cytokine and antibody response to SARS-CoV-2 infection. J Virology. 2021; 95: e00014-21
56. Khan FA, Stewart I, Fabbri L, et al. Systematic review and meta-analysis of anakinra,sarilumab, siltuximab and tocilizumab for COVID-19. Thorax. 2021. doi: 10.1136/thoraxjnl-2020-215266. Online ahead of print.
57. Greenhalgh T, Knight M, A'Court C, et al. Management of post-acute covid-19 in primary care. BMJ. 2020; 370: m3026.
58. National Institute of Health. COVID-19 Treatment Guidelines. Clinical Spectrum of SARS-CoV-2 Infection. https://www.covid19treatmentguidelines.nih.gov/overview/clinical-spectrum/ Accessed Mar 31 2021.
59. Nalbandian A, Sehgal K, Gupta A, et al. Post-acute COVID-19 syndrome. Nat Med. 2021. doi: 10.1038/s41591-021-01283-z. Online ahead of print.
60. Okarska-Napierała M, Ludwikowska K, Książyk J, et al. Approach to a child with paediatric inflammatory multisystem syndrome with COVID-19. Przegl Pediatr 2020; 49: 1-9.
61. Jaroszewicz J. Gąsior M. Kompleksowa opieka nad chorym z zespołem Post-COVID-19 (PC19). I-medica Warszawa 2021, ISBN: 978-83-959922-1-6.
62. Spyropoulos AC, Levy JH, Ageno W, et al. Subcommittee on Perioperative, Critical Care Thrombosis, Haemostasis of the Scientific, Standardization Committee of the International Society on Thrombosis and Haemostasis. Scientific and Standardization Committee communication: Clinical guidance on the diagnosis, prevention, and treatment of venous thromboembolism in hospitalized patients with COVID-19. J Thromb Haemost. 2020; 18: 1859-1865.
63. Spyropoulos AC, Anderson FA Jr, Fitzgerald G, et al. Predictive and associative models to identify hospitalized medical patients at risk for VTE. Chest. 2011; 140: 706-714.
64. Polack FP, Thomas SJ, Kitchin N et al. Safety and Efficacy of the BNT162b2 mRNA Covid-19 Vaccine. N Engl J Med. 2020; 383: 2603-2615.
65. Comirnaty – summary of product characteristics. https://www.ema.europa.eu/en/documents/product-information/comirnaty-epar-product-information\_pl.pdf. Accessed March 31, 2021.
66. Rzymski P, Borkowski L, Drąg M, et al. The Strategies to Support the COVID-19 Vaccination with Evidence-Based Communication and Tackling Misinformation. Vaccines (Basel) 2021; 9: 109.
67. Borkowski L, Drąg M, Fal AM, et al. Vaccinations against COVID-19. Innovative technologies and efficiency. https://naukaprzeciwpandemii.pl/en/. Accessed March 31, 2021.
68. Centers for Disease Control and Prevention. Information about COVID-19 vaccines for people who are pregnant or breastfeeding. https://www.cdc.gov/coronavirus/2019-ncov/vaccines/recommendations/pregnancy.html. Accessed April 12, 2021.

Table 1

Recommended pharmacological management at different clinical stages of SARS‑CoV‑2 infection including primary and supportive treatment (modified from reference 9).

|  |  |  |
| --- | --- | --- |
| **Disease stage** | **Primary treatment** | **Supportive therapy** |
| **Stage 1:**  **asymptomatic or mildly symptomatic**   * SpO2 ≥94% * no hospitalization is necessary | * antipyretic drugs (paracetamol, ibuprofen, etc.), * rest, * oral hydration, * low-molecular-weight heparins in chronically bedridden patients, * antitussive drugs for severe cough, * **Budesonide,** 2 x 800µg daily by inhalation * systemic glucocorticosteroids are contraindicated, * antibiotics, anti-influenza drugs contraindicated, unless there is a bacterial co-infection or concomitant influenza, * oxygen saturation control – implementation of the Pulsocare remote alarm system (using pulse oximeters). | |
| **Stage 2:**  **fully symptomatic** (viral multiplication)   * SpO2 <94% * usually week 1 after disease onset * hospitalization is required | **Remdesivir** administered intravenously once daily for 5 days, with a loading dose of 200 mg on day 1, followed by a maintenance dose of 100 mg. If remdesivir is not available high antibody titer convalescent plasma can be considered. | * low-molecular-weight heparin at prophylactic or therapeutic doses * dexamethasone in patients receiving remdesivir and oxygen therapy, iv or po 4-8 mg daily; should not be used in the first week of the disease if remdesivir is not administered. * antibiotic therapy in case of secondary bacterial infection * symptomatic treatment * oxygen therapy * oral or intravenous hydration |
| **Stage 3:**  **respiratory failure** (cytokine storm)   * SpO2 <90% * usually week 2 after disease onset * hospitalization is required | **Tocilizumab** (in patients with IL-6 >100 pg/ml) is used in single dose of 800 mg iv if patient’s body weight (PBW) >90 kg; 600 mg iv if PBW >65 kg and ≤90 kg; 400 mg iv if PBW >40 kg and ≤65 kg and 8 mg/kg iv if PBW ≤40 kg. The second dose can be administered 8-24 hours after the first one, if the patient’s general condition has not improved; and/or  **Dexamethasone phosphate** administered intravenously at a daily dose of 6-8 mg\* for 7-10 days. | * low-molecular-weight heparin at prophylactic or therapeutic doses * antibiotic therapy in case of secondary bacterial infection * symptomatic treatment * low-/high-flow oxygen therapy * intravenous hydration |
| **Stage 4:**  **ARDS**   * unsuccessful pharmacotherapy to date * need for mechanical ventilation * ICU treatment is required | **Dexamethasone phosphate** administered intravenously at a daily dose of 6-8 mg\* for 7-10 days. If dexamethasone is not available, other glucocorticosteroids can be used at corresponding doses, ie.: hydrocortisone: 50 mg iv. 3 times daily, methylprednisone: 10 mg iv. 4 times daily, prednisone: 40 mg po. once daily; and/or  **Tocilizumab** combined with dexamethasone may be considered in mechanically ventilated individuals (should be started early, within the first day of treatment). | * high-flow oxygen therapy * non-invasive ventilation * mechanical ventilation * ECMO in selected cases * low-molecular-weight heparin in prophylactic or therapeutic doses depending on clinical scenario * empiric antibiotic therapy is strongly not recommended unless there are evident signs of secondary bacterial infection |

\* according to manufacturers, 6 or 8 mg/ml dexamethasone phosphate contained in the available injection solutions correspond respectively 4.95 and 6.6 mg/ml dexamethasone.

Abbreviations: ARDS, acute respiratory distress syndrome; ECMO, extracorporeal membrane oxygenation; ICU, intensive care unit; IL‑6, interleukin 6; SpO2, oxygen saturation as measured by pulse oximetry.

Table 2

The IMPROVE VTE risk-assessment model to guide prophylactic anticoagulation after discharge from hospital after the acute phase of COVID-19 [63]. A score of 2 or more suggests the need of prophylaxis.

|  |  |
| --- | --- |
| Risk factor | Points |
| Prior venous thromboembolism | 3 |
| Diagnosed thrombophilia | 2 |
| Current lower limb paralysis | 2 |
| Current cancer | 2 |
| Immobilized for at least 7 days | 1 |
| Stay in ICU or coronary care unit | 1 |
| More than 60 years old | 1 |