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Weathering the storm: COVID-19 infection in patients with hematological malignancies^{*}

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Abstract: The coronavirus disease 2019 (COVID-19) is an emerging infectious disease caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). Within a matter of months, this highly contagious novel virus has led to a global outbreak and is still spreading rapidly across continents. In patients with COVID-19, underlying chronic diseases and comorbidities are associated with dismal treatment outcomes. Owing to their immunosuppressive status, patients with hematological malignancies (HMs) are at an increased risk of infection and have a worse prognosis than patients without HMs. Accordingly, intensive attention should be paid to this cohort. In this review, we summarize and analyze specific clinical manifestations for patients with coexisting COVID-19 and HMs. Furthermore, we briefly describe customized management strategies and interventions for this susceptible cohort. This review is intended to guide clinical practice.

Key words: Coronavirus disease 2019 (COVID-19); Hematological malignancies; Immunosuppressive status; Management strategies; Treatment regimen

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1 Introduction

The coronavirus disease 2019 (COVID-19) has spread worldwide in the past ten months. By Oct. 9, 2020, according to the World Health Organization (2020), **CLC number:** R73; R511

a total of 36361054 cumulative confirmed cases and 1056186 deaths had been documented globally in 216 countries and territories across six continents. Recent epidemiological study has revealed that the highly infectious causal pathogen, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), is commonly transmitted through respiratory droplets and close contact (Gu et al., 2020). It has also been detected in the sputum, blood, stool (Gu et al., 2020), urine (Jones et al., 2020), and even tears (Xia et al., 2020). At the molecular level, SARS-CoV-2 can enter host cells via interacting with angiotensin-converting enzyme 2 (ACE2) receptors located in various organs (Hamming et al., 2004; Zhou P et al., 2020), and therefore, could induce multiple-organ manifestations

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of infection (Gavriatopoulou et al., 2020). The severity and clinical outcomes of COVID-19 vary among individuals. Up to now, an advanced age, high sequential organ failure assessment score, high D-dimer level on admission, and preexisting chronic diseases have been recognized as risk factors for dismal outcomes (Wang XH et al., 2020; Zhou F et al., 2020). In particular, patients with cancer have a remarkably elevated risk of COVID-19 and poorer outcomes than those in non-cancer cohorts, which can be explained by the systemic immunosuppressive state caused by the malignancies and anti-cancer therapies (Liang et al., 2020).

Hematological malignancies (HMs) are a group of heterogeneous blood diseases, including leukemias, lymphomas, and multiple myelomas (MMs) (Li et al., 2018). According to the most recent statistics, both leukemia and non-Hodgkin lymphoma are among the top ten cancers in terms of incidence and death rates (Siegel et al., 2019). As cancer progresses, malignant cells proliferate extensively and inhibit normal hematopoiesis. The suppression of leukopoiesis confers increased susceptibility to several infections, such as bacterial, fungal, and viral infections (Busca, 2012). Though HMs remain challenging, the survival period and quality of life of patients with HMs have improved remarkably as a result of the introduction of chemotherapy, targeted therapy, monoclonal antibodies, and lately cell-based immunotherapy. However, these anti-cancer therapies might also expose patients to a higher risk of infection (Byrne et al., 2017).

Amid the COVID-19 outbreak, clinicians need to optimize the clinical management of HM patients in terms of treatment regimens, preventive measures for infection, clinical trials of novel therapies, and long-term follow-up. Recently, detailed case reports and cohort studies of COVID-19 in patients with preexisting HMs have become increasingly available in databases like PubMed, Web of Science, and medRxiv. Given the unfavorable outcomes of this susceptible group, strict vigilance is necessary to recognize and differentiate clinical manifestations, radiologic patterns, and laboratory findings. Moreover, the timely detection of COVID-19 and positive interventions are not only necessary to improve the survival rate of patients with HMs, but also to limit transmission among patients and caregivers. In this review, we outline recent reports of patients with coexisting COVID-19 and HMs. Also, we summarize the specific

clinical manifestations. Based on recent guidelines and clinical experience, we have compiled a set of management strategies for HM patients and interventional strategies for patients with HMs and COVID-19.

2 Regional reports of SARS-CoV-2 infections in patients with HMs

Owing to the characteristics of COVID-19 in HM cohorts, increasing numbers of case reports, large-cohort studies, and scattered clinical experiences have recently been presented. In this section, we summarize details of recent cases (Table 1) and provide an overview of several large-cohort studies.

In Wuhan, China, 128 hospitalized patients with HMs and 226 caregivers from two centers were included in a cohort study (He et al., 2020). The calculated incidences of SARS-CoV-2 infection in HM and caregiver cohorts were 10% and 7%, respectively. Though no significant difference was found in terms of prevalence, subjects with HMs tended to develop severe COVID-19. Based on the data of 530 individuals with chronic myeloid leukemia (CML), another local study group in Hubei Province reported a SARS-CoV-2 infection prevalence of 0.9% in patients and 0.1% in healthy individuals (Baumann et al., 2020). Note that advanced phase CML could enhance the risk of COVID-19, even if patients attained a complete cytogenetic response or significant molecular response.

Meanwhile, in the UK, Cook et al. (2020) conducted a real-world assessment, focusing on the clinical impact of COVID-19 in individuals with MMs. The risk of death was found to be significantly associated with male gender, advancing age, and other comorbidities. The authors also noted that patients with relapsed or refractory MM might be at the highest risk of dismal outcomes once infected. Moreover, multicenter retrospective research in China that focused on the same topic reported that HM patients have a poorer prognosis than solid tumor patients (Yang et al., 2020).

The European Myeloma Network (Terpos et al., 2020b) and Malard and Mohty (2020) published detailed management strategies for MM during the COVID-19 pandemic. In France, Malard et al. (2020) studied consecutive adult patients with hematologic disorders at the Hematology Department of the Saint-Antoine

	Reference	Chen et al., 2020	Zhao et al., 2020	Wu et al., 2020	eni et al., 2020	Farmer et al., 2020	n et al., 2020	ted as Baumann, et al., 2020 -19
	Ref				n Sien 2(-E	200 Baur
	Comment	 Patients with agranulocytosis tend to develop multiple infections or secondary viral infections and have dismal outcomes; SARS-CoV-2 might exist for an extended period in HM patients; Repetitive SARS-CoV-2 nucleic acid tests are needed to reduce false-negative rates and to detect the recurrence of infections. 	 Pediatric patients might bear longer durations of diagnosis than adults; Prolonged duration of cough might be caused by a low viral load. 	 Early symptoms of HM patients were uncharacteristic of the typical presentations of COVID-19; Mild COVID-19 might exacerbate the underlying hematologic diseases. 	Highly immunocompromised children on Sieni et al., anticancer therapy might have a 2020 favorable outcome under proper interventions.	 Both APL and COVID-19 are associated with coagulopathy, which raised diagnostic and therapeutic challenges; Prophylactic corticosteroids to prevent DS should be reappraised in the context of COVID-19. 	 Laboratory findings are not reliable in HM patients; HM patients, as well as the glucocorticoids-therapy receivers, might have a longer incubation period. 	 The prevalence of CLL was defined as 0.95% (4/420); Despite coexisting with various comorbidity, the course of COVID-19 was mild and controllable.
V 11D-17 Cases	Clinical outcome	Unstable and low SaO ₂ ; Referral to ICU	Recovery	Recovery	Recovery	Ϋ́	Remarkably improved	Recovery (4/4)
us ut reput teu co	Treatment regimen (for COVID-19 and HM)	CTX, Ara-C, up-leveled antibiotics, oxygen therapy, Arbidol, ribavirin, IFN, IVIG, etc.	Discontinued chemotherapy, IFN, sulfamethoxazole, others	Lopinavir/ritonavir, antibiotics	Anti-microbial and antifungal therapy, HCQ, lopinavir/ ritonavir	ATRA+ATO, enoxaparin, supportive management for COVID-19	Oral chlorambucil (reduced dose), IFN, IVIG, methylprednisolone	Lopinavir/ritonavir, hydroxychloroquine, azithromycin, others
CHILICAL CHARACTERISTICS AND INTEL VENTIONS OF LEPOT LEU COVID-17 CASES WITH TIME	Radiologic finding	Lung inflammation and consolidation (initially); Floc-density shadow with GGOs (after antibiotic therapies)	Dense texture in both sides of the lungs	Initial stage: normal; Mild infiltration in the Progressive stage: right lung hemocytopenia, PCT↑, BNP↑	Bilateral increased reticular marking	Extensive, predominantly peripheral, consolidative changes	GGOs and pleural fluid	Bilateral or single-lung infiltration
	Laboratory finding	WBCJ, ANCJ, ALCJ, HGBJ, PLTJ, IL-6f, IL-10f; Nucleic acid detection of SARS-CoV-2 ever turned to negative and soon back to positive	WBCJ, ALCJ, AST1, ALT1, CRP1; Initial negative results of RT-PCR	Initial stage: normal; Progressive stage: hemocytopenia, PCT↑, BNP↑	WBCL, HbJ, PLTL, CRP↑	WBCJ, ANCJ, ALCJ, PLTJ, PT↑, D-dimer↑, ferritin↑, LDH↑, CRP↑	WBC1, lymphocyte percentage1	Ferritin† (3/4), lymphocytopenia (2/4), D-dimer† (1/4)
I and I	Symptom and incubation period	Intractable fever, paroxysmal dry cough	Cough (for nearly 33 d), fever	Initial stage: productive cough, fatigue; fever, shortness of breath	Persistent fever	Fever, cough, sweats	Moderate fever, productive cough, dyspnea; 25 d	Fever (4/4), dry cough (2/4), diarrhea (1/4)
	Patient demographics (gender, age, place)	Male, 8 years, China	Male, 3 years and 10 months, China	Female, 62 years, China	Female, 13 years, Italy	Male, 36 years, UK Fever, cough, sweats	Male, 39 years, China	Male, >70 years, Spain
	Type of HM	T-cell ALL	B-cell ALL	ALL	AML	APL	CLL (progressed) and NHL	CLL (4 cases)

Table 1 Clinical characteristics and interventions of reported COVID-19 cases with HMs

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Type of HM	Patient demographics	Symptom and incubation period	Laboratory finding	Radiologic finding	Treatment regimen (for COVID-19 and	Clinical outcome	Comment	Reference
CML (accelerated	(gender, age, place) Male, 47 years, China	Cough, fever, dvsnnea.	Lymphocytopenia	No sign of pneumonia	HM) NA	Cured	Radiological findings of COVID-19 could be negative in CML.	Wang DY et al.
phase)		pharyngalgia						2020
CML (chronic phase)	Female, 89 years, China	Cough, dyspnea (oxygen therapy ineffective)	Lymphocytopenia, ALT and AST [↑] , Cr [↑] , troponin- positive	Typical pneumonia patterns	NA	Succumbed after a 3-d hospitalization	 Elderly patients with comorbidities have a poor prognosis and high mortality; 2. The degree of lymphopenia is related to the severity of the disease. 	Wang DY et al., 2020
CML (chronic phase)	Pregnant female, 26 years, Italy	Fever	Testing positive on the swab	NA	Paracetamol, antibiotics	Recovery	Chronic phase CML patients on TKl are not at higher risk of developing COVID-19.	Abruzzese et al., 2020
MM	Male, 60 years, China	Aggravating chest tightness without fever or cough	Aggravating chest ALCJ, CRP1, IL-6 [†] tightness without fever or cough	Bilateral, multiple GGOs and pneumatocele located in both subpleural spaces	Thalidomide, moxifloxacin IV, methylprednisolone, tocilizumab, Arbidol tablets	Recovery	 Symptoms of COVID-19 might be atypical with coexisting comorbidities; IL-6 receptor inhibitor might be effective in alleviating cytokine storm in COVID-19. 	Zhang et al., 2020 n
H	Female, 22 years, Ireland	Cough, fever, sore Lymphocytopenia, throat, chills, CRP1, LDH1, rigors PLT↓ (transiently	Lymphocytopenia, CRP1, LDH1, PLT1 (transiently)	Infiltrates in the lower zones bilaterally	Pembrolizumab, lopinavir/ritonavir, antibiotics, HCQ and azithromycin, corticosteroids	Better	 PDI-inhibitor-induced pneumonitis should be carefully differentiated in the context of COVID-19; Amid COVID-19, dual pathology for lung injury, CRS should be noted when using a PD1 inhibitor. 	O'Kelly et al., 2020
NHL	Male, 78 years, Italy	Persistent fever, cough, dyspnea	RT-PCR on pharyngeal swab was twice tested negative for COVID-19 and turrned to positive later	PET-CT: multiple FDG-avid lymphadenopathies and non-FDG-avid peripheral rounded GGO in the right upper lobe	Tocilizumab, CPAP ventilation	Worsen	 False or delayed results of RT-PCR should be taken into consideration; Co-infection with other pathogens might influence the accuracy of RT-PCR. 	Zanoni et al., 2020
Mantle cell lymphoma	France France	Asthenia, abdominal pain, dry cough	Leukocytosis, LDH1	Early CT: no sign of pulmonary infection but multiple adenomegalies PET/CT: FDG-avid bilateral ground glass and subpleural curvilinear opacities	NA	NA	The PET/CT features of COVID-19 were Playe et al., in accordance with recent publications. 2020	e Playe et al., 2020
ALC, absolute leukemia; Ara- chronic myelo cyclophosphan HMS, hematolo NHL, non-Hoć RT-PCR, real-t	ALC, absolute lymphocyte count; ALL, acute lymphoblastic leukemia; Ara-C, cytarabine; AST, aspartate aminotransfera chronic myeloid leukemia; COVID-19, coronavirus disease cyclophosphamide; DS, differentiation syndrome; FDG, fluo HMS, hematological malignancies; ICU, intensive care unit; IF NHL, non-Hodgkin lymphoma; PET/CT, positron emission RT-PCR, real-time polymerase chain reaction; SaO ₂ , oxygen	LL, acute lymphobl aspartate aminotrans aspartate aminotrans flow syndrome; FDG, U, intensive care uni I/CT, positron emiss reaction; SaO ₂ , oxy	astic leukemia; ALT, ferase; ATO, arsenic sease 2019; CPAP, o fluorodeoxyglucose; (ft, IFN, interferon; IL, iion tomography/comp gen saturation; SARS.	alanine aminotransferas trioxide; ATRA, all-tra ontinuous positive airw 3GO, ground-glass opac interleukin; IVIG, intravé puted tomography; PLT -CoV-2, severe acute res	e; AML, acute myeloi ns retinoic acid; BNP, ay pressure; Cr, creat ity; Hb, hemoglobin; I anous immunoglobulin; , platelet; PT, prothroi piratory syndrome cor	d leukemia; AN B-type natriure ine; CRP, C-re HCQ, hydroxycl (LDH, lactate d mbin time; PCT onavirus 2; TKI	ALC, absolute lymphocyte count; ALL, acute lymphoblastic leukemia; ALT, alamine aminotransferase; AML, acute myeloid leukemia; ANC, absolute neutrophil count; APL, acute promyelocytic leukemia; Ara-C, cytarabine; AST, aspartate aminotransferase; ATO, arsenic trioxide; ATRA, all-trans retinoic acid; BNP, B-type natrinuetic peptide; CLL, chronic lymphocytic leukemia; CML, chronic myeloid leukemia; COVID-19, coronavirus disease 2019; CPAP, continuous positive airway pressure; Cr, creatine; CRP, C-reactive protein; CRS, cytokine release syndrome; CTX, eyclophosphamide; DS, differentiation syndrome; FDG, fluorodeoxyglucose; GGO, ground-glass opacity; Hb, hemoglobin; HCQ, hydroxychloroquine; HGB, hemoglobin; HL, Hodgkin lymphoma; HMs, hematological malignancies; ICU, intensive care unit; IFN, interferon; IL, interleukin; IVIG, intravenous immunoglobulin; LDH, lactate dehydrogenase; MM, multiple myeloma; NA, not available; NHL, non-Hodgkin lymphoma; PET/CT, positron emission tomography/computed tomography; PLT, platelet; PT, prothrombin time; PCT, procalcitonin; PDI, programmed cell death protein I; RPCR, real-time polymerase chain reaction; SaO ₂ , oxygen saturation; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2; TKI, tyrosine kinase inhibitor; WBC, white blood cell	promyelocytic ukemia; CML, ndrome; CTX, kin lymphoma; v, not available; eath protein 1; ood cell

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Hospital, beginning on Mar. 9, 2020. After nearly one month, 25 patients with hemopathies had confirmed COVID-19. In this cohort, fever (22/25), cough (19/25), and shortness of breath (19/25) were the most common complaints at diagnosis. Complementary examinations were also typical, for example, lymphopenia (23/25) and bilateral ground-glass opacities (25/25). The HM cohort was at a higher risk of developing severe COVID-19 with acute respiratory distress syndrome. Within this cohort, patients with MM accounted for a high proportion, emphasizing the impact of steroidinvolved chemotherapy.

Foà et al. (2020) collected data on adult patients with acute lymphoblastic leukemia (ALL) from 40 hematological centers in Italy during the COVID-19 outbreak. Notably, in the Philadelphia chromosomepositive (ph+) cohort, only one individual had verified COVID-19, and one was symptomatic but tested negative. The low infection rate supported the effectiveness of the tyrosine kinase inhibitor (TKI)-based "soft" approach for the treatment of ph+ ALL. Moreover, the effectiveness of this regimen has been identified in GIMEMA (Gruppo Italiano Malattie Ematologiche dell'Adulto) nationwide trials. More importantly, amid the COVID-19 pandemic, this regimen has reduced the hospitalization days and enabled home-based treatment.

Subsequently, in Barcelona, Spain, 804 chronic lymphocytic leukemia (CLL) patients were registered, 420 of whom maintained follow-up. Of this CLL cohort, only four patients (0.95%) were diagnosed with symptomatic COVID-19. The details of the four infected individuals with CLL are summarized in Table 1. These published case reports and cohort studies were performed at different research centers, with differences in various factors, such as medical level and treatment planning. Thus, an overview of their experiences could provide insight into optimal choices for patient care.

3 Specific clinical manifestations, radiologic patterns, and laboratory findings in patients with COVID-19 and HMs

3.1 Clinical manifestations

In 2002 and 2003, most patients with SARS had a fever (99%–100%). In contrast, fever (43.8% on admission, 88.7% during hospitalization) and nonproductive cough (67.8%) are not specific symptoms for COVID-19 (Guan et al., 2020; Han et al., 2020). Those asymptomatic SARS-CoV-2 carriers might contribute to the acceleration of COVID-19 transmission. Based on recent literature, patients with HMs are more likely to be asymptomatic carriers presenting with atypical and delayed COVID-19 symptoms. As shown in Table 1, some COVID-19 patients with HMs might experience extended incubation periods (Jin et al., 2020). Patients with HMs, along with COVID-19, might be non-febrile on admission or during the entire period of illness (Zhang et al., 2020). Moreover, mild initial symptoms (e.g., a slight cough, dyspnea, or chest tightness) might be followed by rapid deterioration. Worse still, overlapping clinical manifestations of the infection and tumor would hinder diagnosis, as discussed in detail later.

3.2 Radiologic findings

In most patients with COVID-19, the initial radiologic findings show multiple patchy shadows and interstitial changes in the lungs before the appearance of ground-glass opacities (Guan et al., 2020). In HM cohorts, radiologic imaging findings might not be significantly typical or recognizable. Because of their low immunity, lung imaging of patients might initially show mixed lesions of bacterial, viral, or fungal infections. Signs of viral infection would take time to emerge after the administration of antimicrobial and antifungal medications. In immunocompromised patients, the critical viral infection could lead to "extensive white lung," which is caused mainly by the respiratory syncytial virus, cytomegalovirus, or SARS-CoV-2, individually or collectively. In clinical practice, we also discovered that patients with COVID-19 would have radiographic images and patterns of disease progression based on chest imaging (X-ray and computed tomography (CT) scan) similar to those of patients with cytomegalovirus-related pneumonia with diffuse large B-cell lymphoma (Fig. 1). Therefore, besides radiologic imaging, a definite diagnosis would be required with nasopharyngeal swabs or bronchoalveolar lavage samples, testing for COVID-19 and other pathogens.

3.3 Laboratory tests

The characteristic laboratory features of COVID-19 include lymphocytopenia, leukopenia, and thrombocytopenia (Guan et al., 2020). However, in patients

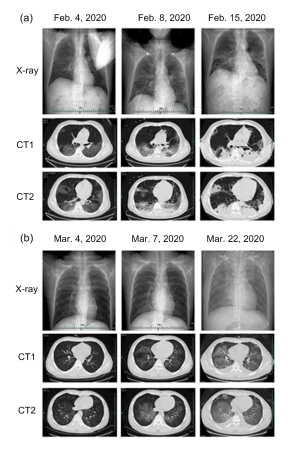


Fig. 1 Comparisons of chest imaging (X-ray and CT scans) of COVID-19 and CMV pneumonia

These two types of pneumonia (COVID-19 vs. CMV) showed a somewhat similar progression in the initial stage, progressive stage, and consolidation stage/diffuse infiltration stage of infection. Initially, both presented with typical GGOs. (a) Chest imaging of COVID-19 pneumonia. A 51-year-old male was confirmed with COVID-19 and received CT scans on 4, 8, and 15 February, 2020. (b) Chest imaging of CMV pneumonia. A 26-year-old male with diffuse large B cell lymphoma (non-GCB, IVB stage) was diagnosed with CMV pneumonia during chemotherapy. He received CT scans on 4, 7, 22 March, 2020. COVID-19, coronavirus disease 2019; CMV, cytomegalovirus; CT, computed tomography; GCB, germinal center B-cell-like lymphoma; GGOs, ground-glass opacities

with HMs, laboratory results are not as reliable as those for COVID-19 patients without such malignancies. Hematological neoplasms and treatment regimens might influence the blood system to variable degrees. For example, leukemia cells could boost leukocyte counts to abnormally high levels, whereas other malignant cells could impair the function of hematopoiesis and leukopoiesis. Moreover, the unexplained phenomenon of "COVID-induced lymphocytosis (CIL)," with remarkably elevated levels of leukocytes and lymphocytes, was observed in patients with CLL (Paneesha et al., 2020). Additionally, highdose chemotherapeutic agents and lymphocyte-targeted immunotherapy would influence normal hematopoiesis and deplete lymphocytes non-selectively. For example, rituximab, an anti-CD20 monoclonal antibody, and chimeric antigen receptor (CAR) T-cell therapy were designed to effectively recognize and lyse malignant and non-malignant B-cells (Freeman and Sehn, 2018; June and Sadelain, 2018).

Furthermore, severe COVID-19 could be characterized by a hyper-inflammatory status with a significant elevation of serum cytokines and chemokines (Mehta et al., 2020). High serum procalcitonin, ferritin, C-reactive protein, lactate dehydrogenase, interleukin-6 (IL-6), and D-dimer levels are associated with a dismal prognosis for COVID-19, prompting timely intervention to improve clinical outcomes (Lagunas-Rangel, 2020; Terpos et al., 2020a).

Undeniably, microbial pathogen detection and serological tests are the most accurate methods to diagnose COVID-19, even in patients with HMs. Using real-time polymerase chain reaction (RT-PCR) and next-generation sequencing, it is possible to identify SARS-CoV-2 nucleic acids from various specimen types like nasopharyngeal swabs, sputum samples, and blood samples. With respect to serological tests, immunoglobulin M (IgM) specific to SARS-CoV-2 can be detected nearly 3 d after infection, and IgG levels increase four-fold during the recovery phase. However, for subjects with negative RT-PCR results, typical CT patterns could be found in more than 70% of suspected individuals (Ai et al., 2020). Therefore, it is better to increase the frequency of examination or combine nucleic acid tests with serological tests (or imaging results) to improve the COVID-19 detection rate in patients with HMs.

4 Clinical diagnostic criteria and differential diagnoses of patients with COVID-19 and HMs

During the COVID-19 pandemic, clinical diagnostic criteria are essential to differentiate between infected and non-infected cohorts efficiently. In line with the certainty of diagnosis, these criteria divide patients into three categories, specifically "surveillance cases," "clinically COVID-19 suspected cases," and "confirmed COVID-19 cases" (Table 2).

In clinical practice, the efficient identification of "suspected cases" is an urgent and challenging issue, especially in HM cohorts, due to diverse manifestations with various confounding factors. Besides SARS-CoV-2, pneumonia in patients with HM could be induced by other viruses, fungal pathogens, or even medications. Table 3 clearly describes the similarities and unique features of various types of pneumonia in patients with HMs. According to the Chinese guidelines for COVID-19, suspected cases can be ruled out if an individual tests negative twice in the SARS-CoV-2 nucleic acid test and achieves negative serological results 7 d after disease onset. As a side note, samples for nucleic acid detection should be collected using a throat swab or nasal swab (at longer than a 1-d interval).

Another significant issue is the partial overlap between the common complaints of patients with HMs and patients with COVID-19. Fever, as a systemic warning symptom of COVID-19, is commonly observed in the hematology department. The causative factors, including infection, cancer, and treatment, should be dissected with meticulous attention (Table 4). Organ-specific manifestations of COVID-19 also partly resemble the symptoms of HMs. In a large retrospective study, the neurological complication of COVID-19 was observed in over 36.4% of infected patients, and 24.8% of infected patients presented symptoms from the central nervous system (CNS) (Gavriatopoulou et al., 2020; Mao et al., 2020). The most common CNS symptoms were dizziness and headache, which overlapped with the CNS involvement of leukemia or myeloma in patients with HMs. Kidney involvement, a typical complication of MM, is also common in SARS-CoV-2-infected individuals. For instance, a large prospective study, which involved both moderate and severe COVID-19 cases, demonstrated that 43.9% presented with proteinuria and 26.7% with hematuria on admission (Boulware et al., 2020; Gavriatopoulou

Table 2 Stratified diagnosis criteria of SARS-CoV-2 infection in patients with HMs

Case category	Criteria for diagnosis
Surveillance ^a	 The patient presents with systemic or respiratory signs and symptoms, such as fever, fatigue, nausea, vomiting, and diarrhea. The patient had a place context with plinically prepared on early firmed COVID 10 second.
	2. The patient had a close contact with clinically suspected or confirmed COVID-19 cases.
Suspected ^b	 History of exposure: The patient had a living history or travel history in and around areas with high COVID-19 prevalence stayed in the community with suspected COVID-19 cases.
	• The patient had close contact with individuals who had been to areas with high COVID-19 prevalence the community with suspected COVID-19 symptoms.
	• The patient was from one of the COVID-19 clusters.
	• The patient had a contact history with confirmed COVID-19 patients.
	2. Clinical manifestations:
	• The patient presents with fever or respiratory symptoms like non-productive cough, dyspnea, and cher tightness.
	• The imaging test of the patient shows pneumonia features with multiple spot shadows and interstitial changes. It could progress into multiple infiltration shadows on bilateral lung and pulmonary consolidation in severe cases; however, pleural effusion is rare.
	The total leukocyte count, granulocyte count, and (or) lymphocyte count could be reduced.When there are no definite or convincing etiologies to explain the clinical manifestations thoroughly.
Confirmed ^c	1. Positive nucleic acid detection of the SARS-CoV-2 from the samples (respiratory tract secretion, blood etc.) by RT-PCR;
	 Sequencing of the SARS-CoV-2 viral genomes from samples of the respiratory tract or blood; Both IgM and IgG serological results of SARS-CoV-2 test positive.

viduals with a confirmed history of exposure, accompanied by two of the clinical manifestations. ^cCases that are compatible with each of the diagnostic items can be diagnosed as a confirmed case of COVID-19. COVID-19, coronavirus disease 2019; HMs, hematological malignancies; Ig, immunoglobulin; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2; RT-PCR, real-time polymerase chain reaction. Stratified diagnosis criteria refer to Liang et al. (2020), China National Health Commission (2020), and Subspecialty Group of Hematology and Oncology, Society of Pediatrics of Hubei (2020)

			Viral pneumonia	monia				
Characteristics	COVID-19	Adenovirus pneumonia	CMV pneumonia	RSV pneumonia	Influenza virus pneumonia	HPIV pneumonia	- Fungal pneumonia	Drug-induced pneumonia
Causative factor	SARS-CoV-2	Adenovirus	CMV	RSV	Influenza virus	HPIV	Pneumocystis jirovecii	Medicine (e.g., MTX, RTX)
Epidemiology	Travel history to an area with high prevalence or contact history with infected patients	Year-round	Mostly after allo-HCT	Seasonal variation of incidence	Seasonal variation of incidence	Year-round	Common in immunodeficiency patients	
Symptoms and signs	Common symptoms: onset with symptoms of upper respiratory infection, such as fever, headache, sore throat, and nasal congestion. When it progresses to pneumonia, most develop a non-productive cough with persistent high fever, dyspnea, etc. Their manifestations vary from asymptomatic patterns to end-organ damage, such as ARDS. Common signs: diminished breath sounds with wheezing rales or crackles.	with symptoms When it progresse yspnea, etc. Theii S. ed breath sounds	of upper respin s to pneumonia r manifestation: with wheezing	atory infection, suc t, most develop a nc s vary from asympti- rales or crackles.	ch as fever, head on-productive cc omatic patterns	ache, sore throat, ugh with to end-organ	Fever, non-productive cough, dyspnea, etc.	Dyspnea, fever, cough
Laboratory findings	Most viral infection induces a reduction of leukocytes and lymphocytes, whereas a routine blood test is not always reliable and typical. Combination of viral nucleic acid detection and antibody detection could differentiate various infections.	ces a reduction of ical. leic acid detectior	Teukocytes and	l lymphocytes, whe detection could diff	rreas a routine bl	lood test is not s infections.	Special staining of the sample from BAL	
CT imaging	Diffuse bilateral ground-glass infiltrates small multifocal nodules, bronchial vascular thickening, and/or airspace consolidation.	glass infiltrates sn	nall multifocal	nodules, bronchial	vascular thicken	ing, and/or	Bilateral, patchy GGOs, and cystic lesions	Diffuse interstitial infiltrates
Distribution of lesions	Peripheral	Multifocal	Diffuse	Airway, multifocal	Airway, multifocal	Airway, multifocal		
660	*+	+++++	+ + + +	+	+	+	Common (not as common as in patients without HMs)	Common
Consolidation	Uncommon	+++	+	+	+	+	Uncommon	
Nodule	Rare	Centrilobular+	+	Centrilobular+++	+++++	Centrilobular++	Uncommon	
Pleural effusion	Uncommon	Common	Rare	Common	Uncommon	Uncommon	Rare	
References	Koo et al., 2018; Fontana and Strasfeld, 2019	and Strasfeld, 20	19				Cordonnier et al., 2017	Atzeni et al., 2013
[#] Involvement of lu ARDS, acute respi virus; MTX, methor	[#] Involvement of lung was graded by "+/++/++++": +, 10%-<25%; ++, 25%-<50%; +++, 50%-<75%; ++++, no less than 75%. Allo-HCT, allogeneic hematopoietic cell transplantation; ARDS, acute respiratory distress syndrome; BAL, bronchoalveolar lavage; CMV, cytomegalovirus; GGO, ground-glass opacity; HMS, hematological malignancies; HPIV, human parainfluenza virus; MTX, methotrexate; RSV, respiratory svncytial virus; RTX, rituximab; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2	-/++++": +, 10%< L, bronchoalveolar tial virus; RTX, ritt	25%; ++, 25%- lavage; CMV, c tximab; SARS-C	<50%; +++, 50%-<7 ytomegalovirus; GGC oV-2, severe acute res	75%; ++++, no le), ground-glass ol spiratory syndrom	ss than 75%. Allo-H pacity; HMs, hemato e coronavirus 2	10%-<25%; ++, 25%-<50%; +++, 50%-<75%; ++++, no less than 75%. Allo-HCT, allogeneic hematopoietic cell transplantation; Iveolar lavage; CMV, cytomegalovirus; GGO, ground-glass opacity; HMS, hematological malignancies; HPIV, human parainfluenza TX, rituximab; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2	cell transplantation; human parainfluenza

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Table 3 Differential diagnoses of several types of pneumonia in patients with HMs

Causative factor	Example
Infectious fever	Viral infections, including SARS-CoV-2, CMV, and RSV Fungal infections Bacterial infections Others, such as mycoplasma
Treatment-associated fever	Chemotherapeutics, such as high-dose MTX, L-ASP, and Ara-C aGVHD after HSCT CRS induced by CAR-T therapy Others
Oncological fever	NHL: body temperature >38 °C accompanied by weight loss or (and) night sweats HL: patients could present with unknown persistent fever; 1/6 patients could develop Pel Ebstein fever Others

Table 4 Principal causative factors of fever in patients with hematological cancers

aGVHD, acute graft versus host reaction; Ara-C, cytarabine; CAR-T, chimeric antigen receptor T cell; CMV, cytomegalovirus; CRS, cytokine release syndrome; HL, Hodgkin lymphoma; HSCT, hematopoietic stem cell transplantation; L-ASP, L-asparaginase; MTX, methotrexate; NHL, non-Hodgkin lymphoma; RSV, respiratory syncytial virus; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2

et al., 2020). Moreover, in acute promyelocytic leukemia (APL), petechia on the skin, gingival hemorrhaging, and other familiar manifestations of coagulation dysfunction might disguise COVID-19-induced coagulation abnormalities (Wang YD et al., 2020).

Governments' restrictive measures, public anxiety, and a shift in the primary focus of medical staff to SARS-CoV-2 infection amid the pandemic might lead to underdiagnosis of symptoms in HM patients, deferral of hematological laboratory and radiological tests, and postponement of other hematological exams. Most acute leukemias deteriorate rapidly with severe complications, making prompt diagnosis and treatment crucial. Therefore, telephone calls, telemedicine, or digital platforms should be used to screen for suspicious hematological symptoms (Molica et al., 2020). After a referral to a hospital, a comprehensive evaluation of SARS-CoV-2 and hematological disease should be performed simultaneously.

5 General management strategies for patients with HMs during the COVID-19 pandemic

The outbreak of COVID-19 has not only had a massive impact on patients with HMs, but also posed tremendous challenges for medical institutions and medical staff. Routine medical procedures and standards for the diagnosis, treatment, follow-up, and prevention of nosocomial infections need to be adjusted to account for COVID-19.

5.1 Considerations for hospital admission procedures

Patients with HMs are advised to follow different hospital admission procedures in view of the prevalence of COVID-19 in different areas of a hospital (Fig. 2). On admission, patients with HMs should be classified according to the risk stratification system for COVID-19 (Table 2). Those patients with a history of COVID-19 exposure or suspicious symptoms should be further investigated. Detailed contact history with exact locations, time, and individuals should be tracked and recorded. Also, blood tests, biochemical analysis, viral nucleic acid detection, viral antibody detection, and chest CT imaging are strongly recommended, to aid in decision-making regarding subsequent diagnostic processes and cancer regimens.

Furthermore, each medical team should carefully weigh the advantages and disadvantages of each inclusion for clinical trials. Clinical trials allow access to off-label drugs or combinations that can be highly beneficial but are otherwise unavailable (Malard and Mohty, 2020). Additionally, routine SARS-CoV-2 testing is also applicable in clinical trials before inclusion.

For highly suspected or confirmed COVID-19 cases, HM patients should be transferred to assigned hospitals for further treatment and isolation. Moreover, a multidisciplinary team (MDT) involving hematologists, infectious disease experts, and other related specialists is necessary to discuss and tailor individualized treatment plans. Throughout the entire medical process, medical staff should be well-protected with full personal protective equipment.

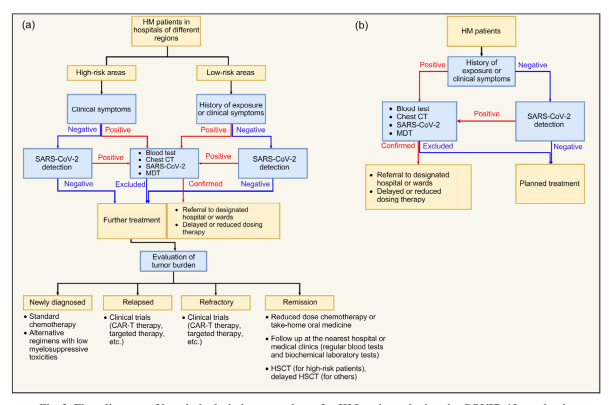


Fig. 2 Flow diagram of hospital admission procedures for HM patients during the COVID-19 pandemic (a) Admission procedures in areas with a high prevalence of COVID-19; (b) Admission procedures in areas with a low prevalence of COVID-19. CAR-T, chimeric antigen receptor T cell; COVID-19, coronavirus disease 2019; CT, computed tomography; HM, hematological malignancy; HSCT, hematopoietic stem cell transplantation; MDT, multidisciplinary team; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2

5.2 Considerations for hematological therapeutic regimens

Current options for the treatment of HMs include chemotherapy, immunotherapy, targeted therapy, hematopoietic stem cell transplantation (HSCT), radiotherapy, and cell-based therapy. All of these options would cause detrimental effects on hemopoietic functions, to some extent. Therefore, treatment needs to be adjusted during the COVID-19 outbreak.

5.2.1 Chemotherapy

High-dose chemotherapy would severely impair the immune system via non-selective anti-cancer activity. Then, overcoming the sequelae of chemotherapy typically requires extensive medical equipment, medicine, or blood products. However, most medical equipment has been mobilized to emergency departments or COVID-19 wards during this critical period. Therefore, it is necessary to adjust conventional chemotherapy regimens for patients with HMs. The core ideas for converted regimens are as follows: (1) safe and effective oral chemotherapeutic drugs are preferable to reduce the duration of hospitalization; (2) therapeutic regimens should be simplified, with less frequent visits and shorter duration; (3) for patients maintaining remission, the dose of chemotherapy or the frequency of follow-up visits could be reduced; and (4) non-urgent chemotherapy or medical procedures should be deferred.

For newly diagnosed patients with a high tumor burden or multiple high-risk genetic aberrations, the risks and benefits should be carefully weighed because underdosing might not kill the malignant cells efficiently and may induce tolerance to therapy. In Tables 5–7, we briefly summarize the considerations for treatment regimens of common hematological cancers during the COVID-19 pandemic based on recommendations from the American Society of Hematology (ASH), the European Hematology Association (EHA), and other published guidelines. Before the administration of myelosuppressive treatment, it is vital to rule out COVID-19 infection.

	Table 5 Considerations for treatment of leukemia during the COVID-19 pandemic						
Leukemia	Induction treatment	Consolidation and maintenance therapy	R/R patient	Reference			
Acute leukemi	ia						
ALL	Ph-positive:1. TKI+minimal corticosteroid exposure2. TKI+low-dose chemotherapy	 Ph-positive: 1. Consolidation therapy is not suggested for CR patients 2. TKI+corticosteroids (for maintenance therapy) 	 B-ALL: 1. Inotuzumab 2. Blinatumomab or CD22 CAR-T (with enough medical resources) 	Gruber et al., 2009; Gavillet et al., 2020; Stock et al., 2020			
ALL	 Ph-negative: Standard chemotherapy Reduced dose for patients with high risk of complications Minimized corticosteroids to relieve symptoms G-CSF to facilitate the recovery of leukocyte count 	 Ph-negative: 1. Consolidation therapy is not suggested for CR patients 2. Allo-HSCT 3. Vincristine+prednisone+ 6-MP+MTX (for maintenance therapy) 	T-ALL:1. Standard nelarabine-based regimens2. Salvage chemotherapy and targeted therapies when necessary	Gavillet et al., 2020; NCCN, 2020; Stock et al., 2020			
AML	 APL: 1. Targeted oral drugs (e.g., ATO+ATRA), for low-risk patients; for high-risk pa- tients, regimens should in- clude a cytotoxic agent 2. Dexamethasone (to prevent DS) 	APL: ATO to achieve a low relapse rate	 Salvage intensive re-induction HSCT, aggressive chemotherapy or targeted therapies 	Raza et al., 2020; Tallman et al., 2020			
AML	 Non-APL: 1. "7+3" regimen 2. Preventive antimicrobial drugs+G-CSF 3. Targeted drugs with reduced doses 4. Adjusted blood supply standards 	 Non-APL: 1. High-dose cytarabine+ reduced-dose cytarabine (decreased to three cycles) 2. Oral BCL-2 inhibitor alone or combined with 5-azacitidine 		Gavillet et al., 2020; NCCN, 2020; Tallman et al., 2020			
Chronic leuker	mia						
CLL	 First-line treatment like ibrutin Monoclonal anti-CD20 antibo Avoid the combination of targ Venetoclax should be cautious quent hemogram monitoring 	dies and IVIG eted therapy		Shadman et al., 2020			
CML	 Newly diagnosed: first-line tre Accelerated phase: continuing For others, dose interruptions 	TKI if clinical response is po	ositive	Mauro et al., 2020			

 Table 5 Considerations for treatment of leukemia during the COVID-19 pandemic

ALL, acute lymphoblastic leukemia; Allo-HSCT, allogeneic hematopoietic stem cell transplantation; AML, acute myeloid leukemia; APL, acute promyelocytic leukemia; ATRA, all-trans retinoic acid; ATO, arsenic trioxide; B-ALL, B cell-ALL; CAR-T, chimeric antigen receptor T cell; CLL, chronic lymphoblastic leukemia; CML, chronic myeloid leukemia; COVID-19, coronavirus disease 2019; CR, complete remission; DS, differentiation syndrome; G-CSF, granulocyte-colony stimulating factor; IVIG, intravenous immunoglobulin; 6-MP, 6-mercapto-purine; MTX, methotrexate; Ph, Philadelphia chromosome; R/R, relapsed or refractory; T-ALL, T-cell ALL; TKI, tyrosine kinase inhibitors

5.2.2 Immunotherapy

Unlike chemotherapy, novel therapies, like monoclonal antibodies, bispecific T-cell receptor-engaging antibodies, and chimeric antigen receptor T cell (CAR-T) therapies, aim to precisely eradicate malignant cells by recognizing surface antigens on tumor cells in relapsed or refractory HMs. However, for CAR-T therapy, long-term B-cell aplasia would increase susceptibility to infections. Also, cytokine release syndrome, the most common toxicity associated with CAR-T therapy, would promote cytokine storms with COVID-19. Fortunately, Bachanova et al. (2020) proposed appropriate strategies and useful guidelines for CAR-T therapy during the COVID-19 pandemic.

Lymphoma	Induction treatment	Consolidation and maintenance therapy	R/R patient	Reference
HL	Early stage: Four cycles of ABVD with an interim of PET-CT (favorable disease) Advanced stage: 1. ABVD+PET/CT 2. AAVD+G-CSF	Conventional HL regimens+ prophylactic antibiotics	 Second-line regimens (e.g., gemcitabine-based regimens) Brentuximab vedotin or PD1 antibodies 	Advani et al., 2020b; ESMO, 2020b
NHL (mainly focused on DLBCL)	 High-risk: 1. CHOP-21 regimens 2. G-CSF to minimize neutropenia Medium-risk: 1. Non-curative treatment could be delayed 2. Avoid T-cell suppressive agents (e.g., bendamustine) 	High-risk: Lenalidomide+RTX (R2) or lenalidomide+ corticosteroids Others: 6-MP+MTX	 High-dose chemotherapy+ ASCT Second-line chemotherapy Lenalidomide-based therapy Polatuzamab+ bendamustine CAR-T cell therapy 	Advani et al., 2020a; ESMO, 2020a

Table 6 Considerations for treatment of lymphoma during the COVID-19 pandemic

AAVD, brentuximab vedotin, doxorubicin, vinblastine, and dacarbazine; ABVD, doxorubicin, bleomycin, vinblastine, and dacarbazine; ASCT, autologous stem cell transplantation; CAR-T, chimeric antigen receptor T cell; CHOP-21, three-weekly cyclophosphamide, doxorubicin, vincristine, and prednisolone; COVID-19, coronavirus disease 2019; DLBCL, diffuse large B-cell lymphoma; G-CSF, granulocyte-colony stimulating factor; HL, Hodgkin lymphoma; 6-MP, 6-mercapto-purine; MTX, methotrexate; NHL, non-Hodgkin lymphoma; PD1, programmed cell death protein 1; PET/CT, positron emission tomography/computed tomography; R/R, relapsed or refractory; RTX, rituximab

Table 7 Considerations for treatment of multiple myeloma during the COVID-19 pandemic

Multiple myeloma	Induction treatment	Consolidation and maintenance therapy	R/R patient	Reference
Standard-risk	 Lenalidomide, bortezomib, and dexamethasone (RVd regimen) Oral ixazomib, lenalidomide, and dexamethasone (IRd regimen; 28-d cycle) CRd regimen; 28-d cycle DRd regimen for elderly patients CyBorD regiment: the only approved initial therapy 	 Reduced dose of dexamethasone with follow-up for those with good responses after 3–4 cycles Bortezomib on a weekly basis after 6–8 cycles 	 Salvage ASCT (for aggressive relapse or secondary PCL) Daratumumab- based treatment 	Al Saleh et al., 2020; Malard and Mohty, 2020; Rajkumar et al., 2020
	6. Clinic visits at an interval of 4 weeks7. Consider administering additional cycles of induction, and delaying transplant until the first relapse	3. If remission after 10–12 cycles, le- nalidomide		
High-risk	 Twenty-eight-day cycle of carfilzomib, lenalidomide, and dexamethasone (KRd regimen) High-dose melphalan and ASCT as first-line treatments Weekly outpatient clinic visits 	 RVd or Rd regimens Acquiring ASCT post-pandemic 		Al Saleh et al., 2020; Rajkumar et al., 2020

ASCT, autologous stem cell transplantation; COVID-19, coronavirus disease 2019; CRd, cyclophosphamide, lenalidomide, and dexamethasone; CyBorD, cyclophosphamide, bortezomib, and dexamethasone; DRd, daratumumab, lenalidomide, and dexamethasone; IRd, ixazomib, lenalidomide, and dexamethasone; KRd, carfilzomib, lenalidomide, and dexamethasone; PCL, plasma cell leukemia; Rd, lenalidomide and dexamethasone; R/R, relapsed or refractory; RVd, lenalidomide, bortezomib, and dexamethasone

5.2.3 Hematopoietic stem cell transplantation

HSCT is an established and indispensable therapy for helping advanced-stage patients to achieve longterm survival (Gruber et al., 2009). Before its administration, patients should be treated with high-dose chemotherapy, inducing a relatively long period of pancytopenia and increasing susceptibility to various infections. However, its implementation is challenging during the COVID-19 pandemic. Therefore, strict measures, such as the use of rooms with laminar flow hoods or positive pressure isolation wards, should be adopted for this cohort. Furthermore, non-urgent transplantation procedures for patients with stable diseases could be deferred. Moreover, COVID-19 screening should be conducted for both donors and recipients (Ljungman et al., 2020). To reduce the falsenegative rate, the combination of SARS-CoV-2 nucleic acid and antibody tests should be performed at least twice at an interval of at least 24 h prior to HSCT.

To minimize the influence of unpredictable factors, the cryopreservation of stem cells before the conditioning regimen and the preparation of a back-up donor is recommended by the European Society for Blood and Marrow Transplantation (Ljungman et al., 2020). If a transplant candidate is diagnosed with COVID-19, HSCT should be delayed for at least three months. For individuals who were in close contact with patients with COVID-19, HSCT should not be performed for at least 14 to 21 d (Ljungman et al., 2020).

5.2.4 Recommendations for regimen adjustments

5.2.4.1 Acute leukemia

In acute myeloid leukemia (AML), the regimens differ remarkably between APL and non-APL. For non-APL, following a negative COVID-19 test, intensive induction chemotherapy is recommended with a curative intent (Raza et al., 2020). Lower-intensity therapies such as a hypomethylating agent with venetoclax can be administered to people as outpatients to reduce exposure (DiNardo et al., 2019). Moreover, the consolidation with high-dose cytarabine should be offered to individuals with negative COVID-19 testing. The dose could be lowered (from 3.0 to 1.5 mg/m^2) to shorten the duration of myelosuppression, and the cycles could be reduced (Mayer et al., 1994; Burnett et al., 2013; Brissot et al., 2020). For APL, the regimen of "oral all-trans retinoic acid (ATRA) and arsenic trioxide" is more desirable for low-risk individuals, owing to the reduced duration of myelosuppression (Zhu and Huang, 2014). High-risk patients must still receive a regimen that includes a cytotoxic agent. In ALL, glucocorticoids should be prescribed in a recommended dose, given the paucity of information on their potential adverse role in SARS-CoV-2 infections. Meanwhile, bacterial and fungal infections and the inherent risk of thrombotic complications should be prevented in advance (Brissot et al., 2020).

5.2.4.2 Chronic leukemia

Among patients with CLL, a "watch and wait" policy has been widely adopted during this epidemic. Notably, given the dismal outcome of infection, selfisolating should be strictly performed to limit a patient's exposure to viral infection (Paneesha et al., 2020). Because of the potential of Bruton tyrosine kinase (BTK) inhibitors to disturb pulmonary inflammation and immune response in COVID-19, they could be reserved for the treatment of CLL and other indications, such as mantle cell lymphoma and Waldenstrom macroglobulinemia (Scarfò et al., 2020).

As for CML, the absolute risk of contracting a severe SARS-CoV-2 infection was lower than 1% (Ector et al., 2020; Li et al., 2020). Leukemia-associated factors, including no complete hematologic response and subjects in advanced phase CML, were found to be the risk factors of infection, whereas the use of TKI was not (Li et al., 2020). Therefore, it was safe to continue their TKI.

5.2.4.3 Lymphoma

Diffuse large B-cell lymphoma is the most common aggressive non-Hodgkin lymphoma. For newly diagnosed patients, R-CHOP (rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisone), as the standard regimen, is recommended to be performed immediately. Moreover, the cycle of R-CHOP could be reduced to four cycles if the positron emission tomography (PET) is negative at the end of treatment (di Ciaccio et al., 2020).

5.2.4.4 Multiple myeloma

As for MM, high-dose melphalan and autologous HSCT (ASCT) remain the standard care. At the same time, hemopoietic recovery takes a long time after this therapy. Therefore, it was recommended to perform induction regimens of up to six cycles to postpone the ASCT. In the standard-risk group, transplant could be delayed until the first relapse because no significant effect on overall survival rate has been observed in this adjusted regimen. In the high-risk group, ASCT should still be regarded as first-line therapy, and SARS-CoV-2 screening should be highlighted throughout the therapy (Malard and Mohty, 2020). Referring to the current guidelines, Table 7 summarizes induction, consolidation, and maintenance therapies, and regimens for relapsed or refractory MM.

5.3 Considerations for follow-up plans

Unfortunately, routine follow-up has been strongly influenced by traffic restrictions and travel bans during the COVID-19 pandemic. In this context, patients should seek medical care at the nearest hospital to ensure treatment in the shortest time and minimize opportunities for exposure to COVID-19. Furthermore, telemedicine could be introduced to enable communication between patients and doctors. Video conferencing platforms, such as Zoom, WeChat, and Skype, could be used for collaborations among different medical institutions. If infection or disease progression is suspected, patients could search for timely medical assistance.

Once the travel bans are lifted, pre-planned treatment and follow-up strategies could be resumed. However, the screening and testing of suspected COVID-19 cases should be continued. The precautions of proper handwashing methods, masking, so-cial distancing, and good personal hygiene should be disseminated and strictly practiced among vulnerable cohorts. Above all, the most effective method to prevent COVID-19 transmission is still likely to be vaccination, once it becomes available.

6 Comprehensive intervention strategies for patients with COVID-19 and preexisting HMs

The coexistence of HMs and COVID-19 infection is rather complicated and challenging for medical experts. During comprehensive care, in severely infected cases, COVID-19 should be prioritized over HMs. Generally, patients with COVID-19 and HMs should be isolated, and comprehensive treatment strategies should be designed according to the severity of the disease. The core interventions include the clearance of viremia, relieving cytokine storms, and improving organ function.

Aiming to target the direct action of the virus on multiple organs, antivirals (e.g., lopinavir/ritonavir, remdesivir) and antibodies (e.g., convalescent plasma) are under evaluation for their safety and feasibility for managing COVID-19 (Wiersinga et al., 2020). Emerging evidence shows the favorable effect of convalescent plasma infusion on improving symptoms and biomedical indices, and proves that early administration with convalescent plasma is preferable before seroconversion for SARS-CoV-2. Convalescent plasma also shows an acceptable safety profile (Psaltopoulou et al., 2020). Most other medications did not show significant benefits for patients (Wiersinga et al., 2020). A cytokine storm initiated by SARS-CoV-2 would induce a series of adverse effects, such as a hypercoagulable inflammatory state, increasing the risk of secondary infections, and impairing humoral immunity (Chauhan et al., 2020; Jamilloux et al., 2020). Therefore, targeted immunomodulatory agents (e.g., tocilizumab, ruxolitinib) and anti-inflammatory agents (e.g., dexamethasone) have been used to curb cytokine storms. Thromboprophylaxis is necessary among COVID-19 patients. Low-molecular weight heparins are the firstline choice, and other antithrombotic treatments such as antithrombin could be integrated (Terpos et al., 2020a).

The monitoring of blood oxygen saturation and oxygen therapy should be appropriately and promptly adopted as hypoxemia, or cytokine release syndrome would contribute to the development of shock, tissue perfusion disorders, and even multiple organ failure. Oxygen saturation should be maintained at 93%–96% for patients without chronic pulmonary disease and at 88%–92% for patients with chronic type II respiratory failure.

Artificial liver support systems and blood purification efficiently reduce the incidence of shock, hypoxemia, and respiratory distress syndrome (Liang et al., 2020). For patients with HMs and COVID-19, the prophylactic use of antibiotics is necessary (quinolones, second- or third-generation cephalosporins, and β -lactamase inhibitor compounds). Alternatives, such as carbapenems, β -lactamase inhibitor compounds, linezolid, and vancomycin, can be used in critically ill patients.

Supportive treatment and nutritional support are crucial to maintaining the balance of the intracellular environment. A perturbation of the intestinal microecological balance in patients with COVID-19 can result in gastrointestinal symptoms, such as diarrhea, abdominal pain, and even secondary infection. Therefore, patients with severe COVID-19 infection would benefit from regular oral feeding and enteral nutrition.

With the improvement of fever, respiratory symptoms, and radiologic and laboratory results, patients with HMs could be discharged after approval of the MDT. Importantly, two consecutive nucleic acid tests from throat swab specimens should test negative, with a sampling interval of at least 24 h (Liang et al., 2020). After discharge, patients with COVID-19 and HMs must continue two weeks of home quarantine and isolation. Personal hygiene measures, such as handwashing and mask-wearing, should be emphasized. Re-examinations should include liver and kidney functions, blood tests, nucleic acid tests of sputum and stool samples, and a pulmonary function test or lung CT scan, according to the patient's condition. Remote management and telemedicine should be introduced for follow-ups (Balduzzi et al., 2020).

7 Conclusions

We have summarized atypical symptoms and features of patients with COVID-19 and HMs based on our broad experience and data, as well as the recently published literature. The prolonged incubation period, complicated laboratory patterns, and confounding radiologic findings might eventually lead to the misdiagnosis of COVID-19 in patients with HMs. Furthermore, we have described additional considerations related to the diagnosis, differential diagnosis, and management of patients with HMs based on our clinical experiences. Amid the COVID-19 pandemic, patients with HM should be managed individually based on the severity and type of disease. Hematological cancer treatments during this critical period should be adjusted on a case-by-case basis. For patients with HMs and confirmed COVID-19, timely supportive treatment, antiviral regimens, oxygen therapy, and prophylactic antibiotics should be administered. An overview of management strategies for patients with HMs during the COVID-19 pandemic is illustrated in Fig. 3.

We have also acknowledged the importance of home-based oral chemotherapy to reduce clinic visits. Moreover, a simplified chemotherapy regimen with higher therapeutic efficacy and lower toxicity for patients with HMs is indispensable. Lastly, hematology wards should be alert and prepared for emergencies and disease outbreaks. For example, procedures for hospital admission, mass screening, quarantine and isolation, and novel treatments for emergency conditions should be well-planned. Strengthening emergency preparedness, immediate responses, and timely recovery are critical to avoid future pandemics.

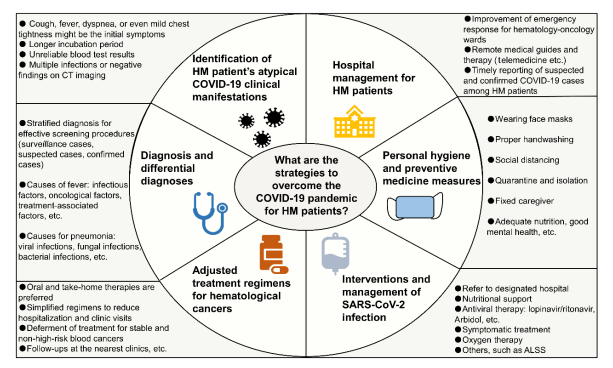


Fig. 3 Overview of management strategies for HM patients during the COVID-19 pandemic

ALSS, artificial liver support system; HM, hematological malignancy; COVID-19, coronavirus disease 2019; CT, computed tomography; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2

Contributors

He HUANG, Arnon NAGLER, and Yong-xian HU took the lead in writing the manuscript. Lin-qin WANG and Elaine TAN SU YIN wrote and edited the manuscript. Guo-qing WEI participated in shaping the tables and figures. All authors have read and approved the final manuscript and, therefore, have full access to all the data in the study and take responsibility for the integrity and security of the data.

Compliance with ethics guidelines

He HUANG, Arnon NAGLER, Yong-xian HU, Lin-qin WANG, Elaine TAN SU YIN, and Guo-qing WEI declare that they have no conflict of interest.

This article does not contain any studies with human or animal subjects performed by any of the authors.

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<u>中文概要</u>

- 题 目: 应对恶性血液肿瘤在新型冠状病毒肺炎疫情中的 挑战
- 概要:新型冠状病毒肺炎(COVID-19)是一种由严重急性呼吸综合征冠状病毒2(SARS-CoV-2)引起的新型传染病。SARS-CoV-2具有极强的传染性,因此在引发流行病后又使其迅速升级为全球大流行,至今仍在部分国家迅速传播。在COVID-19的感染者中,存在肿瘤等慢性病或伴有相关合并症的患者预后往往较差。同时,由于恶性血液肿瘤患者长期处于免疫抑制状态,往往感染风险更高且预后更差,所以十分值得我们关注。这篇综述归纳并分析了恶性血液肿瘤患者合并COVID-19后的临床表现。此外,归纳了疫情期间恶性肿瘤的管理策略和感染后的综合干预措施,旨在为临床提供指导。
- 关键词:新型冠状病毒肺炎(COVID-19);恶性血液肿瘤; 免疫抑制;管理策略;治疗方案