In the USA and other western nations, respiratory syncytial virus is one of the most commonly encountered respiratory viruses among patients who have been diagnosed with a hematologic malignancy or who have undergone a stem cell transplant. Multiple studies have been performed to evaluate the complications associated with respiratory syncytial virus infections. Other studies have evaluated therapeutic agents and strategies in which these agents can be used. There have also been numerous reports of outbreaks in bone marrow transplant units and oncology wards, where infection control measures have been invaluable in controlling the spread of disease. However, despite these novel approaches, respiratory syncytial virus continues to be potentially fatal in immunocompromised populations. In this review, we discuss the incidence of respiratory syncytial viral infections, risk factors associated with progression from upper respiratory tract infection to lower respiratory tract infection, other complications and outcomes (including mortality), management strategies, and prevention strategies in patients with a hematologic malignancy and in hematopoietic cell transplant recipients.

Respiratory syncytial virus in hematopoietic cell transplant recipients and patients with hematologic malignancies

Fareed Khawaja and Roy F. Chemaly
Department of Infectious Diseases, Infection Control and Employee Health, Division of Internal Medicine, The University of Texas MD Anderson Cancer Center, Houston, TX, USA

ABSTRACT

Introduction

Community respiratory viruses are a common cause of respiratory infections. These viruses are perhaps best known for their seasonal variation. The outcomes of these infections vary on the basis of the patient population, with adverse outcomes having been described in hematopoietic cell transplant (HCT) recipients and patients with a hematologic malignancy (HM). One of the most common community respiratory viruses that may lead to the death of HCT recipients and HM patients is respiratory syncytial virus (RSV), whose incidence is second only to that of influenza according to prior reports; other viruses include parainfluenza virus, metapneumovirus, adenovirus, rhinovirus, and bocavirus.

Different strategies have been used for the management of RSV infections in immunocompromised patients, and HCT recipients in particular, including ribavirin in its different formulations, intravenous immunoglobulins (IVIG), RSV immunoglobulins, and RSV monoclonal antibodies. In addition, effective measures have been used to curtail outbreaks of RSV infection in numerous bone marrow transplantation units and oncology wards, with some success; however, despite some advances over the past two decades in early detection and management of RSV infections in immunocompromised patients, the outcomes related to these infections remain poor.

In this review, we summarize the published data on RSV infections in adult HCT recipients and HM patients, focusing on recent findings. We highlight the incidence of RSV infections, risk factors associated with progression from upper respiratory tract infection (URTI) to lower respiratory tract infection (LRTI), other complications and outcomes (including mortality), management strategies including new agents under investigation, and prevention strategies.

Incidence of respiratory syncytial virus Infection

RSV has been considered a major cause of respiratory viral infection in HM and HCT patients since the 1980s. In the USA alone, RSV accounts for approximately
30% to 37% of respiratory viral infections in this population, with approximately 19% to 36% of these infections resulting in LRTI. Determination of the true incidence of RSV infection among HCT recipients has been challenging, with a reported wide range of 5% to 49%. As many studies included symptomatic patients diagnosed with different laboratory assays such as RSV antigen detection, identification of RSV by direct fluorescent antibody, viral cultures, and molecular assays in recent years (Table 1). In addition, the wide range of reported incidences of RSV infections could be due to the increased awareness of this virus over the years and its impact on immunocompromised patients. On the other hand, lower incidences of RSV infections were reported in studies in which molecular assays such as multiplex polymerase chain reaction were used to detect RSV, with the reported range in these cases being 6-30%. This lower incidence could be explained by the increased rate of diagnosing other respiratory viruses, such as coronavirus, rhinovirus, and parainfluenza, with the use of molecular assays.

The proportion of RSV infections causing LRTI in HCT recipients ranges from 30% to 60% with higher rates reported in earlier years (in the early 1990s), whereas in more recent studies, since year 2000, lower rates of LRTI (24-44%) have been observed. This could possibly be explained by the increased use of ribavirin at earlier stages of RSV infection to prevent the infection from progressing to the lower respiratory tract, as well as the use of molecular assays for the detection not only of RSV but also other respiratory viruses. Recent studies from the Fred Hutchinson Cancer Research Center on respiratory viral infections in HCT patients applied specific definitions to better delineate the types of LRTI associated with respiratory viral infections. Patients with proven or probable LRTI were defined as patients with microbiological detection of respiratory viruses in the lower respiratory tract with or without radiological evidence of disease in the lungs, respectively. Patients with possible LRTI were defined as patients with microbiological detection of respiratory viruses in the upper respiratory tract only and with radiological evidence of disease in the lungs. In one of the studies, the proportions of patients with proven/probable or possible RSV LRTI were 48% and 52%, respectively. Interestingly, the authors showed that patients with proven or probable LRTI had a higher need for supplemental oxygen use and for mechanical ventilation compared to those with possible LRTI. These definitions were also applied in studies on coronavirus, parainfluenza virus, and rhinovirus.

Data on RSV infections in HM patients are scarce. Some studies have reported the rate of RSV infections in these patients. In an early study, the incidence of RSV was 31% among all symptomatic HM patients, including HCT recipients. A large proportion of these infections were diagnosed as LRTI (56%). Similar studies reported a range of 3% to 37%, but the sample sizes were relative-

**Table 1.** Incidence rates of respiratory syncytial virus infections and lower respiratory tract infections among symptomatic hematopoietic cell transplant recipients.

| Date of publication | Country     | Study type                | Detection methods                        | Percentage of allogeneic HCT | Total number of RSV cases (%) | Percentage of LRTI | Authors        |
|---------------------|-------------|---------------------------|------------------------------------------|-------------------------------|-------------------------------|-------------------|---------------|
| 1991                | USA         | Retrospective observational| RSV antigen and viral culture            | 73%                           | 3 (5%)                        | 33%               | Garcia et al. |
| 1996                | USA         | Retrospective observational| RSV antigen, viral culture and pathology| 52%                           | 33 (49%)                      | 61%               | Whimbey et al.|
| 2001                | Multiple European nations | Prospective observational   | RSV antigen and viral culture            | 90%                           | 20 (10%)                      | 60%               | Ljungman et al.|
| 2002                | England     | Retrospective observational| RSV antigen and viral culture            | 100%                          | 13 (37%)                      | 46%               | Chakrabarti et al.|
| 2002                | USA         | Retrospective observational| RSV DFA, RSV antigen and viral culture  | 89%                           | 48 (9%)                       | 52%               | Small et al. |
| 2003                | Brazil      | Prospective observational  | RSV antigen                             | 72%                           | 27 (40%)                      | 56%               | Machado et al.|
| 2003                | USA         | Prospective observational  | Viral culture and RSV PCR               | 45%                           | 11 (30%)                      | 36%               | Roghmann et al.|
| 2003                | England     | Retrospective observational| RSV antigen and viral culture            | 83%                           | 6 (21%)                       | 33%               | Hassan et al. |
| 2005                | Spain       | Prospective observational  | RSV antigen and viral culture            | 54%                           | 19 (11%)                      | 37%               | Martino et al.|
| 2009                | Sweden      | Retrospective observational| RSV antigen and viral culture            | 100%                          | 32 (12%)                      | 44%               | Averissyan et al.|
| 2013                | Brazil      | Retrospective observational| RSV PCR                                 | 79%                           | 14 (14%)                      | NA                | Moreira et al.|
| 2014                | Italy       | Prospective observational  | RSV PCR                                 | 95%                           | 21 (8%)                       | 11%               | Mikulska et al.|

HCT: hematopoietic cell transplant; RSV: respiratory syncytial virus; LRTI: lower respiratory tract infection; DFA: direct fluorescent antibody detection; PCR: polymerase chain reaction; NA: not available.
ly small. In a recent study, 181 HM patients with RSV infections were described over 13 years. Of these, 65% and 35% presented with URTI and LRTI, respectively. Among the HM patients with URTI, 13% progressed to develop a LRTI (75% were patients with leukemia, 27% with multiple myeloma, and none with lymphoma). In a recent study from our institution focusing on RSV LRTI in HM patients who had or had not undergone HCT, we found that most HM patients who had not undergone HCT were defined as having possible RSV LRTI as bronchoscopically had not been performed in most of these patients at the time of diagnosis.

Few data are available with regards to RSV infections in pediatric HCT recipients. The estimated incidence of RSV infections among this population is 3% to 7%, with 22%–37% developing LRTI.

**Risk factors for the progression of upper to lower respiratory tract infection**

The most significant complication of RSV infection in HM patients and HCT recipients is progression to LRTI, which is associated with a higher mortality rate. Many risk factors for progression have been identified in the hopes that target populations that could benefit from early therapy could be identified. These risk factors primarily consist of host factors, as previous studies on RSV severity.”

**Table 2. Risk factors for progression to RSV lower respiratory tract infections among hematopoietic cell transplant recipients and patients with hematologic malignancies.**

| Risk factors for progression to RSV LRTI | HCT recipients | Patients with hematologic malignancies |
|----------------------------------------|----------------|---------------------------------------|
| Neutropenia ≤ 500 cells/mL              | Neutropenia ≤ 500 cells/mL |
| Lymphopenia ≤ 200 cells/mL             | Lymphopenia ≤ 200 cells/mL |
| Age ≥ 40 years                         | Age ≥ 40 years |
| HCT within 1 year of infection         | HCT within 1 year of infection |
| Myeloablative chemotherapy within 1 year of infection | Myeloablative chemotherapy within 1 year of infection |
| Presence of GvHD                        | Presence of GvHD |
| Use of steroids within 2 weeks of infection | Use of steroids within 2 weeks of infection |
| History of smoking                     | History of smoking |
| Nosocomial infection                   | Nosocomial infection |
| Hypoxia                                | Hypoxia |
| Matched unrelated donor/mismatched donor status | Matched unrelated donor/mismatched donor status |
| Prior autologous HCT                   | Prior autologous HCT |
| Cord blood as transplant cell source   | Cord blood as transplant cell source |

There are currently no predictive scoring index for the progression of respiratory viruses in patients with leukemia, lymphoma, or multiple myeloma. On the other hand, there are well-described risk factors that are associated with progression to LRTI in HM patients; these include lymphopenia and neutropenia, which are generally defined as <200 lymphocytes/mL and <500 neutrophils/mL, respectively. In multiple retrospective analyses of HCT recipients, lymphocytopenia and neutropenia at the time of the diagnosis of RSV were independent predictors of progression to LRTI.

In a cohort of 237 allogeneic HCT recipients with RSV infection, the hazard ratio (HR) of experiencing progression from RSV URTI to LRTI was 4.1 for an absolute neutrophil count of <500 cells/mL and 2.6 for an absolute lymphocyte count of <200 cells/mL, making them the most predictive factors for progression. Among leukemia patients, post-induction chemotherapy neutropenia and leukopenia have also been shown to increase the risk of progression to LRTI. In a recent retrospective study, neutropenia and lymphopenia were not independently associated with progression of disease in HM patients, but the combination of the two factors was associated with a higher risk of progression. However, time from last chemotherapy was not shown to play a role in progression to LRTI in HM patients. On the other hand, lack of ribavirin therapy was associated with progression of dis-

**Table 3. Outcome data stratified by respiratory syncytial virus-Immunodeficiency Scoring Index from three different cohorts.**

| ISI risk stratification | Shah et al. 2014 | Wang et al. 2016 | Damaj et al. 2015 |
|------------------------|------------------|-----------------|------------------|
|                        | Low n=69         | Medium n=147    | High n=21        |
|                        | Low n=17         | Medium n=20     | High n=7         |
|                        | Low n=11         | Medium n=28     | High n=6         |

| Mortality rate, %      | 0                | 3                | 29               |
|                       | 0                | 20               | 14               |
|                       | 0                | 4                | 50               |

| Rate of progression to LRTI, % | 7                | 15               | 42               |
|                               | 6                | 15               | 29               |
|                               | NA               | NA               | NA               |

ISI: Immunodeficiency Scoring Index; LRTI: lower respiratory tract infection.
ease in HCT recipients and HM patients with RSV infections. The protective benefits of ribavirin are discussed further in the treatment section.

**Long-term complications of respiratory syncytial virus infection**

Certain long-term complications after RSV infection have been described in HCT recipients, with the most widely described long-term complication being a reduction in pulmonary function. Delayed engraftment after RSV infection is less commonly described, with an uncertain association in small numbers of patients. In fact, previous studies have shown that a decrease in FEV, of 10% or greater is associated with the development of bronchiolitis obliterans. However, the relationship between respiratory viruses (RSV in particular) and changes in FEV, or other markers of pulmonary function is not fully understood.

There are limited data on HCT recipients and pulmonary function (including changes in FEV, and oxygen diffusion capacity) after an RSV infection. Avestisyan et al. showed that, compared to a control group, HCT recipients with RSV infections were more likely to develop mild or marked changes in vital capacity or diffusion capacity during pulmonary function tests. Subsequently, Seo et al. described significant decreases in patients’ diffusion capacity 3 months after RSV infection, which persisted for a year. There were some indications that FEV, and total lung capacity were also affected, but the sample size was too small to draw definite conclusions. Another study compared the effects of different viruses on pulmonary function in HCT recipients. RSV and parainfluenza were associated with FEV, decreases of at least 10%. It was postulated that subclinical shedding of these viruses may augment airway inflammation, leading to airway restriction. In comparison, lung transplant recipients who are infected with respiratory viruses are at increased risk of developing bronchiolitis obliterans, with an associated mortality of up to 29%. Similarly, HCT recipients with prior respiratory viral infections were more likely to develop bronchiolitis obliterans or changes in FEV, with a higher mortality rate than that of patients without bronchiolitis obliterans (HR: 2.7).

Delayed or failed engraftment of stem cells during or after acute RSV infection has been reported; however, it is an uncommon complication with a total of only seven patients having been described in a few case series. This association was first described in 1999 by McCarthy et al., who noted that four patients with graft failure had had an RSV infection in the pre-engraftment period, with no other identified infections. Furthermore, during an RSV outbreak in a hematologic unit in Australia, delayed neutrophil and platelet engraftment occurred in two autologous HCT recipients and graft failure occurred in one allogeneic HCT recipient with an RSV infection. On the other hand, a study by Waghmare et al. showed no significant changes in lymphocyte count dynamics in HCT recipients who experienced progression to RSV LRMI compared to patients who did not. Overall, there are very limited data supporting that RSV infection per se leads to graft failure or contributes to a delay in engraftment.

**Mortality and associated risk factors**

High mortality rates have been reported in HM patients and HCT recipients with RSV infection. RSV-attributable mortality rates in HCT recipients vary between 0% in outbreak situations, in which some patients received reduced-intensity conditioning regimens, and 43% in other circumstances. When HCT recipients develop RSV LRMI, the mortality rate can range from 21% to 83%. Of note, when HCT recipients with RSV infections were classified into those with possible RSV LRMI (only radiological evidence of chest abnormalities and negative or no bronchoscopy data) or proven RSV LRMI (RSV detected in the lower respiratory tract), the mortality rate increased from 0% to 26%, respectively. Multiple other risk factors for mortality from RSV have been identified, most of which are host-related, including neutropenia and lymphopenia, time from transplant to infection, cell source, older age, steroid exposure, graft-versus-host disease, hypoxia, and the use of myeloablative chemotherapy.

The RSV-IsI has been validated to predict mortality risk in HCT recipients and HM patients. On the derivative cohort of allogeneic HCT recipients with RSV infections, the predicted mortality for patients with high RSV-IsI was 29% and 50% in one of the validation cohorts. Interestingly, some studies showed that ribavirin may have a protective effect in HCT recipients and HM patients. This is discussed further in the treatment section.

Delaying transplant in patients diagnosed with RSV or other respiratory viruses prior to HCT was shown to improve survival. Campbell et al. reviewed 116 patients who had pre-transplant respiratory viral infections and found that, regardless of the virus, they had a higher 100-day mortality rate than did those without infections. At our institution, HCT is delayed for approximately 2 weeks when patients are diagnosed with RSV infections prior to transplantation.

Among HCT recipients with RSV infections, mortality was considerably lower in pediatric patients than in adults. Mortality varied from 0 to 5% among all pediatric HCT recipients with RSV infections and is higher in patients with LRMI (up to 15%).

The mortality rate amongst HM patients with RSV can be as high as 18%. The mortality rate in a small study conducted in the mid-1990s was 80% among leukemia patients with RSV LRMI who had recently undergone myeloablative chemotherapy. More recent studies of HM patients with RSV LRMI found mortality rates of 8% to 17%. This better outlook could be explained in part by the improvement in supportive care over the years and the use of ribavirin for the treatment of RSV in this population of patients.

These host factors reflect the patients’ immune status and its ability to curtail the impact of RSV infections. The severity and stage of the infection (URTMI vs. LRMI) or the virulence of the RSV may also affect mortality. In HM patients with proven LRMI, the 30-day mortality rate was reported to be 36% compared to 14% in patients with possible LRMI. In a recent retrospective analysis, patients with HM, including HCT recipients, in whom respiratory viral infections were detected in the intensive care unit.
had a higher intensive care unit mortality rate, with the association being strongest when influenza, parainfluenza or RSV was detected.10

Treatment options for respiratory syncytial virus infection in patients with hematologic malignancies and in hematopoietic cell transplant recipients

The current treatments for RSV infections in immunocompromised adult patients are ribavirin, in different formulations [although it has not been approved by the Food and Drug Administration (FDA) for this purpose], and immune-modulators, such as conventional IVIG or RSV monoclonal antibodies (palivizumab).

Ribavirin

Ribavirin is a nucleoside analog that is active against a broad spectrum of RNA viruses. It acts through intercalation into the RNA virus, enhancing its mutation rate. Ribavirin is available in aerosolized, oral, and intravenous formulations. Aerosolized ribavirin was approved by the FDA for the treatment of RSV LRTI in hospitalized infants and young children in 1985 and is the most studied formulation in HCT recipients.4,5,34,45,46,49,53,56,71-77 The conventional dosing regimen is 6 g delivered over 18 h through a small particle aerosol generator, with patients in a scavenger tent to decrease environmental contamination and exposure to healthcare workers (ribavirin is teratogenic). In an alternative regimen, the same total dose of ribavirin is given but is divided into three doses per day (2 g over 3 h, 3 times a day). This intermittent regimen was shown to be equivalent to the conventional continuous regimen in an adaptive randomized trial.78

Most of the studies on the use of ribavirin for RSV infections in HCT recipients and HM patients are retrospective in nature, lacking a comparison or control group.26,45,73,74,79,80 Yet cumulative evidence, albeit not from clinical trials, suggests a better outcome when ribavirin is used in HCT recipients with early disease or URTI vs. LRTI.4,46,71,73-75,81 The observed hepatic toxicity was probably due to the coadministration of interferon therapy.92 In a retrospective analysis comparing outcomes of HCT recipients with RSV infections who received either aerosolized or oral ribavirin, only one of the 38 patients who received oral ribavirin developed hemolytic anemia and lactic acidosis. The latter was thought to be due to severe gastrointestinal graft-versus-host disease. In a recent retrospective analysis comparing outcomes of HCT recipients with RSV infections who received either aerosolized or oral ribavirin, two of 29 (6.9%) patients on oral ribavirin developed new-onset grade 3 or more anemia at day 14 compared to two of 41 (4.9%) patients who received the aerosolized formulation. These studies demonstrate that both aerosolized and oral ribavirin have similar safety profiles.
Intravenous ribavirin has been used to treat RSV infection in HCT recipients. The most common regimen used is a single 33 mg/kg loading dose, followed by 16 mg/kg four times a day for 4 days with a maintenance dose of 8 mg/kg three times a day until symptoms resolve. The intravenous formulation of ribavirin is not readily available in the USA due to the lack of FDA approval; however, it can be acquired through the FDA for emergency or compassionate use.

**Adjunctive therapies**

Many immunomodulators have been used as adjunctive therapy for RSV infections in adult HM patients and HCT recipients. There have been no randomized control trials comparing the benefits of adding IVIG to ribavirin therapy. Many retrospective analyses have reported variable results for this combination. Shah et al. suggested a minimal benefit in HCT recipients with RSV infections; all 51 HCT patients who were treated with combination therapy survived, and only one of 61 patients treated with ribavirin alone died. This was not evaluated in multivariate analysis. Another study showed no clinical benefit in HCT recipients who received weekly or high-dose IVIG as adjunctive therapy to ribavirin as monotherapy for RSV infections in adult HCT recipients. In a recent retrospective study of a mixed population of adult HCT recipients and patients with HM, those treated with palivizumab as treatment in HCT recipients with RSV is very limited. Studies showed no survival benefit when comparing palivizumab as adjunctive therapy to ribavirin as monotherapy for RSV infections in adult HCT recipients. In a recent retrospective study of a mixed population of adult HCT recipients and patients with HM, those treated with palivizumab as treatment in HCT recipients with RSV is very limited. Studies showed no survival benefit when comparing palivizumab as adjunctive therapy to ribavirin as monotherapy for RSV infections in adult HCT recipients. In a recent retrospective study of a mixed population of adult HCT recipients and patients with HM, those treated with palivizumab as treatment in HCT recipients with RSV is very limited. Studies showed no survival benefit when comparing palivizumab as adjunctive therapy to ribavirin as monotherapy for RSV infections in adult HCT recipients.

**Investigational agents**

As previously discussed, there are limited therapeutic options for RSV infections and new therapies are desperately needed. Presatovir is an oral RSV fusion inhibitor with selective anti-RSV activity in vitro. In a phase I, first-in-human, single- and multiple-ascending dose study, presatovir had an excellent safety profile, despite having an extended half-life. The oral bioavailability was also

---

*If patient is unable to take oral ribavirin, it can be substituted with the aerosolized formulation.
good, regardless of prandial state. 19 Although clinical trials on RSV infections in transplant recipients have been challenging due to low recruitment, 72 two phase II trials on presatovir in HCT recipients with RSV infections were recently completed 73,100 (ClinicalTrials.gov number NCT02254408). To overcome potential low recruitment, 189 HCT recipients with RSV infections limited to the upper respiratory tract were recruited from a total of 43 centers in nine countries over 2.5 years. 73,100 Preliminary data showed that presatovir was not effective at reducing nasal RSV viral load over time or at reducing the incidence of lower respiratory tract complications. However, in an exploratory analysis, presatovir did reduce the rate of progression to lower respiratory tract complications in patients with lymphopenia relative to the rate in placebo-treated patients. Furthermore, presatovir was not effective at reducing nasal RSV viral load, supplemental oxygen use or all-cause mortality in another phase II trial in HCT recipients with RSV LRTI. 73,100 Several lessons were learned from the two trials. HCT recipients with one or more risk factors (i.e. lymphopenia, neutropenia, or infected within a year from transplant) for poorer outcomes from RSV infections may benefit the most from an effective antiviral therapy. On the other hand, time from symptom onset and time before the development of lower respiratory tract complications are probably critical factors in determining the effectiveness of fusion inhibitors.

The development of another novel agent, ALX-0171, was recently stopped and the drug may not enter clinical trials on RSV infections in HCT recipients. This inhaled agent is a trivalent nanobody that inhibits RSV replication by binding the F-protein on the surface of the virus and thereby neutralizes RSV by blocking virus uptake into cells. A recent phase 1 trial in infants with RSV infections was prematurely terminated due to lack of efficacy (ClinicalTrials.gov identifier: NCT03418574).

Prevention of respiratory syncytial virus infection

The prevention of RSV in HM patients and HCT recipients is limited to infection control measures and interventions. There are many reports of outbreaks in these populations of patients 13,18,101,102 and the emphasis has been on multifaceted interventions, including compliance with hand hygiene, contact precautions with gowns and the use of gloves, screening visitors and healthcare workers for respiratory symptoms and restricting visitors and healthcare workers if they are symptomatic, grouping RSV-infected patients together, and sometimes screening asymptomatic patients in the same treatment areas. These interventions have been shown to be effective in mixed adult and pediatric patients. 19,103,104 At our institution, we place all patients diagnosed with respiratory viral infections on contact and droplet precautions. 103

Interestingly, the use of chemoprophylaxis with palivizumab was described during an outbreak 23 in patients who were at high risk of acquiring RSV infections in a bone marrow transplant ward. Sixteen asymptomatic patients were given one prophylactic dose of palivizumab during the outbreak, and none developed an RSV infection. These results warrant further investigation into the use of palivizumab in an outbreak; however, no definite recommendation can be made at this time. Finally, an RSV vaccine for children or adults is not available at present. Multiple trials in children and healthy adults are in progress (www.ClinicalTrials.gov). Whether immunization for RSV would be beneficial in immunocompromised patients is uncertain.

Conclusions

RSV is a common respiratory viral infection in adult HM patients and HCT recipients. Its incidence is affected by its seasonality and geography and the diagnostic method used. The mortality rate associated with RSV can be high in severely immunocompromised patients with LRTI. The use of an ISI is helpful to stratify HCT recipients into risk categories; a similar scoring system is needed for HM patients. Ribavirin, with or without IVIG, may mitigate the impact of RSV infections in this population of patients. The further development of new antiviral agents or other treatment modalities is of the utmost importance to have a greater impact on rates of progression to LTRI and mortality in adult HM patients and HCT recipients.

References

1. Wu X, Wang Q, Wang M, et al. Incidence of respiratory viral infections detected by PCR and real-time PCR in adult patients with community-acquired pneumonia: a meta-analysis. Respira-tion. 2015;90(4):343-352.

2. Ruskanen O, Lahti E, Jennings LC, Murdoch DR. Viral pneumonia. Lancet. 2011;377(9773):1264-1275.

3. Jain S, Self WH, Wunderink RG, et al. Community-acquired pneumonia requiring hospitalization among U.S. Adults. N Engl J Med. 2015;373(5):415-427.

4. Chemaly RF, Ghosh S, Bodey GP, et al. Respiratory viral infections in adults with hematologic malignancies and human stem cell transplantation recipients: a retrospective study at a major cancer center. Medicine (Baltimore). 2006;85(5):278-287.

5. Ljungman P, Ward KN, Crooks BN, et al. Respiratory virus infections after stem cell transplantation: a prospective study from the Infectious Diseases Working Party of the European Group for Blood and Marrow Transplantation. Bone Marrow Transplant. 2001;28(5):479-484.

6. Chemaly RF, Shah DP, Bodey GP. Management of respiratory viral infections in hematopoietic cell transplant recipients and patients with hematologic malignancies. Clin Infect Dis. 2014;59(Suppl 5):S344-S351.

7. Atilla E, Sahin D, Atilla PA, et al. Upper respiratory viral infections in patients with hematologic malignancies after allogeneic hematopoietic stem cell transplantation: a single-center experience. Open Forum Infect Dis. 2018;5(5):ofy077.

8. Mikulska M, Del Bono V, Gandolfo N, et al. Epidemiology of viral respiratory tract infections in an outpatient haematology facility. Ann Hematol. 2014;93(4):669-676.

9. Moreira LF, Watanabe AS, Carrao E, et al. A survey strategy for human respiratory syncytial virus detection among haematopoietic stem cell transplant patients: epidemiological and methodological analysis. Mem Inst Oswaldo Cruz. 2013;108(1):119-122.

10. Legoff J, Zuurman N, Lemiale V, et al. Clinical significance of upper airway virus detection in critically ill hematology patients. Am J Respir Crit Care Med. 2019;199(4):516-528.

11. Spahr Y, Tschudin-Sutter S, Baettig V, et al. Community-acquired respiratory paramyxovirus infection after allogeneic hematopoietic cell transplantation: a single-center experience. Open Forum Infect Dis. 2018;5(5):ofy077.

12. Wang L, Allen J, Dong C, et al. Respiratory virus infection after allogeneic hematopoietic stem cell transplant in a tropical center: predictive value of the immunodeficiency
Low mortality rates related to respiratory syncytial virus infection in allogeneic hematopoietic stem cell transplant recipients: viral RNA detection in serum samples. Clin Infect Dis. 2013;57(12):1731-1741.

Menaud C, Xie H, Seo S, et al. Mortality rates of human metapneumovirus and respiratory syncytial virus lower respiratory tract infections in hematopoietic cell transplantation recipients. Biol Blood Marrow Transplant. 2015;19(8):1220-1226.

Mendes ET, Ramos J, Peixoto D, et al. An outbreak of respiratory syncytial virus infection in hematopoietic stem cell transplantation patients: viral RNA detection in blood, antiviral treatment, and clinical outcomes. Clin Infect Dis. 2014;59(9):1396-1406.

Ljungman P. Respiratory virus infections in stem cell transplant patients: the European experience. Blood. Marrow Transplant. 2001;7(5):55-75.

Khanna N, Widmer AF, Decker M, et al. Respiratory syncytial virus infection in patients with hematological diseases: single-center study and review of the literature. Clin Infect Dis. 2008;46(3):402-412.

Campbell AF, Chien JW, Kuypers J, et al. Respiratory virus pneumonia after hematopoietic cell transplantation (HCT): associations between respiratory syncytial virus in bronchoalveolar lavage samples, viral RNA detection in serum samples, and clinical outcomes. HCT. J Infect Dis. 2010;201(9):1404-1415.

Mazzulli T, Peret TC, McGeer A, et al. Molecular characterization of a nosocomial outbreak of human respiratory syncytial virus in an adult leukemia/lymphoma ward. J Infect Dis. 1999;179(6):1686-1692.

Shah DP, Ghanotoi SS, Ariza-Heredia EJ, et al. Immunodeficiency scoring index to predict poor outcomes in hematopoietic cell transplant recipients with RSV infections. Blood. 2014;123(21):3263-3268.

Kim YJ, Guthrie KA, Waghmare A, et al. Respiratory syncytial virus in hematopoietic cell transplant recipients: factors determining progression to lower respiratory tract disease. J Infect Dis. 2014;209(5):1195-1204.

Nichols WG, Gooley T, Boeckh M. Community-acquired respiratory syncytial virus and parainfluenza virus infections after hematopoietic stem cell transplantation: the Fred Hutchinson Cancer Research Center experience. Biol Blood Marrow Transplant. 2012;18(7):955-959.

Damlaj M, Ratoo G, Carin-Ceba R, et al. Corticosteroid use as adjunct therapy for respiratory syncytial virus infection in adult allogeneic stem cell transplant recipients. Transplant Infect Dis. 2016;18(2):216-225.
57. Whimbey E, Couch RB, Englund JA, et al. Respiratory syncytial virus pneumonia in hospitalized adult patients with leukemia. Clin Infect Dis. 1995;21(2):376-379.

58. Torres HA, Aguilar EA, Mattuzzi GN, et al. Characteristics and outcome of respiratory syncytial virus infection in patients with leukemia. Haematologica. 2007;92(9):1216-1223.

59. Sheshadri A, Shah D, Kneid J, et al. Pulmonary function decline after respiratory viral infection predicts mortality after hematopoietic stem cell transplantation. Am J Resp Crit Care Med. 2017;195:A2706.

60. Ebbert JO, Limper AH. Respiratory syncytial virus upper respiratory tract illnesses in adult blood and marrow transplantation recipients. Treatment with aerosolized ribavirin and intravenous immunoglobulin. Bone Marrow Transplant. 2000;25(7):751-755.

61. Fishbein CF, Freikas CS, Lease ED, et al. Symptomatic respiratory virus infection and chronic lung allograft dysfunction. Clin Infect Dis. 2016;62(5):513-519.

62. Abdallah A, Rowland KE, Schepetiuk SK, To LB, Bardy P. An outbreak of respiratory syncytial virus infection in a bone marrow transplant unit: effect on engraftment and outcome of pneumonia without specific antiviral treatment. Bone Marrow Transplant. 2003;32(2):195-203.

63. Chien JW, Martin PJ, Gooley TA, et al. Airflow obstruction after myeloablative hematopoietic stem cell transplantation. Am J Resp Crit Care Med. 2003;168(2):208-214.

64. Abedin S, Yank GA, Braun T, et al. Predictive value of bronchiolitis obliterans syndrome stage Op in chronic graft-versus-host disease of the lung. Biol Blood Marrow Transplant. 2015;21(6):1127-1131.

65. Khalilah F, Hashem RM, Chakinala MM, et al. Respiratory viral infections are a distinct risk for bronchiolitis obliterans syndrome and death. Am J Resp Crit Care Med. 2004;170(2):181-187.

66. Versluis AB, Rossen JW, van Ewijk B, Schuurman R, Bierings MB, Boelens JJ. Strong association between respiratory viral infection early after hematopoietic stem cell transplantation and the development of life-threatening acute and chronic allogeneic lung syndromes. Biol Blood Marrow Transplant. 2010;16(6):782-791.

67. Yu S, Li A, et al. Virus infection facilitates the development of severe pneumonia in transplant patients with hematologic malignancies. Oncotarget. 2016;7(35):59366-59406.

68. Ebbert JO, Limper AH. Respiratory syncytial virus pneumonia in immunocompromised adults: clinical features and outcome. Respir Res. 2005;7(2):265-269.

69. Peck AJ, Corey L, Boeckh M. Pretransplantation respiratory syncytial virus infection: impact of a strategy to delay transplantation. Clin Infect Dis. 2004;39(9):675-680.

70. Campbell AP, Guthrie KA, Englund JA, et al. Clinical outcomes associated with respiratory virus detection before allogeneic hematopoietic stem cell transplantation. Clin Infect Dis. 2015;61(2):192-202.

71. Boeckh M, Englund J, Li Y, et al. Randomized controlled multicenter trial of aerosolized ribavirin for respiratory syncytial virus pneumonia complicating transplant recipients. Clin Infect Dis. 2007;44(2):245-249.

72. Field K, Slavin MA, Seymour JF. Severe respiratory syncytial virus pneumonia complicating Budasphamide and cyclophosphamide treatment of chronic lymphocytic leukemia. Eur J Haematol. 2002;69(3):54-57.

73. Ghosh S, Champlin RE, Englund J, et al. Respiratory syncytial virus upper respiratory tract illnesses in adult blood and marrow transplantation recipients. Treatment with aerosolized ribavirin and intravenous immunoglobulin. Bone Marrow Transplant. 2000;25(7):751-755.

74. McColl MD, Conser RB, Brennan J, Chopra R. Respiratory syncytial virus infection in adult BMT recipients: effective therapy with short duration nebulised ribavirin. Bone Marrow Transplant. 1998;21(4):423-425.

75. McCoy E, Kiely JM, Jaffar AU, Wynd MA, Sebti R, Munk GB. Treatment of respiratory syncytial virus infection in adult patients with hematologic malignancies based on an institution-specific guideline. Transpl Infect Dis. 2011;13(2):117-121.

76. Makrelid E, Ljungman P, Ekelof-Andstrom E, et al. Ribavirin therapy in bone marrow transplant recipients with viral respiratory tract infection. Bone Marrow Transplant. 1997;19(9):905-908.

77. Vieira R, Torres HA, Munsell ME, et al. Successful systemic high-dose ribavirin treatment of respiratory syncytial virus and adenovirus isolates from the respiratory tract of healthy children. J Infect Dis. 1995;172:77-88.

78. Schuurman R, Bierings MB, Boelens JJ. Respiratory infecions in hematopoietic stem cell transplantation: a pilot study. Transpl Infect Dis. 2015;21(6):1127-1131.

79. Lewinsohn DM, Bowden RA, Mattson D, et al. An adaptive randomized trial of an intermittent dosing schedule of aerosolized ribavirin in patients with cancer and respiratory syncytial virus infection. J Infect Dis. 2012;206(9):1567-1571.

80. Scharf L, Ljungman P, Ekelof-Andstrom E, et al. Ribavirin treatment in bone marrow transplantation recipients with respiratory syncytial virus infection. Clin Infect Dis. 2016;62(3):313-319.

81. Rhind SC, O'Sullivan KM, de Heer ME, et al. Ribavirin aerosol for the treatment of respiratory syncytial virus in allogeneic hematopoietic stem cell transplantation recipients. Clin Infect Dis. 2018 Sep 8. [Epub ahead of print].

82. German P, Xin Y, Chien JW, et al. Phase 1 and 2b, randomized, double-blind, placebo-controlled trial of presatovir (GS-5806) for the treatment of respiratory syncytial virus (RSV) infections in allogeneic hematopoietic stem cell transplantation recipients. J Infect Dis. 2011;117(4):1087-1102.

83. Scharf L, Ljungman P, Ekelof-Andstrom E, et al. Ribavirin treatment in bone marrow transplantation recipients with respiratory syncytial virus infection. J Infect Dis. 2016;18(4):634-636.

84. Quintero A, Soto J, et al. Respiratory syncytial virus infection in hematopoietic stem cell transplantation recipients with aerosolized ribavirin and the humanized monoclonal antibody palivizumab: a single centre experience. Br J Haematol. 2009;146(5):574-576.

85. Scharf L, Ljungman P, Ekelof-Andstrom E, et al. Ribavirin treatment in bone marrow transplantation recipients with respiratory syncytial virus infection. J Infect Dis. 2016;18(4):634-636.
Chemaly RF, Sanjeet S, Bergeron A, et al. A phase 2b, randomized, double-blind, placebo-controlled trial of presatovir (GS-5806), an oral fusion inhibitor for the treatment of respiratory syncytial virus (RSV) upper respiratory tract infection (URTI) in hematopoietic-cell transplant (HCT) recipients. BMT Tandem Meeting, Salt Lake City, UT, USA; February 21-25, 2018. Presentation LBA 6.

Jalal H, Bibby DF, Bennett J, et al. Molecular investigations of an outbreak of parainfluenza virus type 3 and respiratory syncytial virus infections in a hematology unit. J Clin Microbiol. 2007;45(6):1690-1696.

Lehners N, Schnitzler P, Geis S, et al. Risk factors and containment of respiratory syncytial virus outbreak in a hematology and transplant unit. Bone Marrow Transplant. 2013;48(12):1548-1553.

Gala CL, Hall CB, Schnabel KC, et al. The use of eye-nose goggles to control nosocomial respiratory syncytial virus infection. JAMA. 1986;256(19):2706-2708.

Krasinski K, LaCouture R, Holzman RS, Waihe E, Bonk S, Hanna B. Screening for respiratory syncytial virus and assignment to a cohort at admission to reduce nosocomial transmission. J Pediatr. 1990;116(6):894-898.

Ariza-Heredia EJ, Chemaly RF. Update on infection control practices in cancer hospitals. CA Cancer J Clin. 2018;68(5):540-555.