Implementation of a hospital-at-home (HAH) unit for hematological patients during the COVID-19 pandemic: safety and feasibility

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Abstract

Background “Hospital-at-home” (HAH) programs have been shown to optimize resource utilization, shorten hospitalization and prevent nosocomial infection.

Methods We retrospectively analysed data regarding implementation of an HAH unit for caring patients with hematological malignancies in our center, during the COVID-19 pandemic.

Results Between January and November 2020, 105 patients were treated in the HAH unit for a total of 204 episodes. Nine patients with multiple myeloma (MM) received autologous HSCT (auto-HSCT). Three patients with acute myeloid leukemia (AML) received consolidation therapy, 32 patients underwent clinical and analytical monitoring, 20 were transplant recipients early discharged (5 auto-HSCT and 15 allo-HSCT) and 2 had received CART cells therapy. Azacitidine, bortezomib and carfilzomib were administered at home to 54 patients with AML, myelodysplastic syndrome (MDS) or MM. A median of 17 (IQR 13–19) days of admission per patient and a total of 239 visits to the Hematology day-care hospital were avoided. Overall, 28 patients (14% of all episodes) needed admission to the hospital, 4 of them due to COVID-19.

Conclusions Implementation of a Hematology HAH unit was feasible and safe, and provided thorough advanced care to a high-risk population. Advanced care-at-home strategies can be crucial during times of COVID-19 to minimize treatment interruptions and reduce the risk of cross-infections.

Keywords Hospital-at-home · Hematology · COVID-19 · QOL

Introduction

On December 2019, several cases of pneumonia with unknown etiology occurred in Wuhan, Hubei, China [1–3]. The pathogen was later identified as a novel coronavirus, named SARS-CoV-2 by the World Health Organization (WHO). The disease spread rapidly throughout China and then all over the world. On March 11, 2020, the WHO declared the coronavirus-19 disease (COVID-19), a global pandemic [4].

COVID-19 fatality can be as high as 15% in elderly population and patients with comorbidities [5]. Among them, patients with cancer are more susceptible to infection and have inferior outcomes than individuals without cancer [6–8]. Furthermore, patients with hematological malignancies and hematopoietic stem cell transplant (HSCT) recipients, are a potential high-risk population due to the immune
impairment from their underlying disease, the use of immunosuppressive treatments, and the need of frequent hospital visits for clinical monitoring, treatment delivery and admission. Several studies have described high mortality rates of nearly 30% in this specific population [9–18].

The city of Madrid in Spain, has been at the epicenter of the COVID-19 pandemic with approximately 370,000 diagnosed cases and more than 19,000 deaths (with confirmed or suspected COVID-19) as of early December 2020 [19]. The rapid development of the outbreak resulted in an enormous pressure to the medical institutions and the healthcare system was dramatically overloaded by the end of March 2020. These circumstances caused cross-infections among patients and medical workers and also limited the care of chronically ill individuals and patients with cancer, including hematologic patients. Therefore, novel strategies were needed to allow an appropriate care and treatment delivery for these patients, and at the same time reduce the risk of exposure avoiding hospital admissions and day hospital visits, considering that medical centers became one of the main sources of infection.

Advanced care at home or “hospital-at-home” (HAH) programs have significantly spread out in the last years. These strategies have demonstrated to improve patients satisfaction, avoid unnecessary hospital admissions and reduce healthcare costs [20–23]. Furthermore, hematologic patients have been successfully treated with outpatient approaches worldwide, including Spanish centers [24]. However, data regarding the use of these programs during COVID-19 are scarce.

The objective of this study is to describe the implementation of an HAH unit for clinical and analytical monitoring, treatment delivery and specific at-home treatment programs, for patients with hematological malignancies and HSCT recipients, during the COVID-19 pandemic in our area.

Patients and methods

We retrospectively analysed the implementation of an HAH unit for caring patients with hematological malignancies in a tertiary hospital located in the city of Madrid.

In January 2020, an HAH unit was developed in our center as a multidisciplinary unit, including Hematology physicians and nurses, pharmacists, nutritionists, and social workers.

The main objectives of the HAH unit were as follows: to improve the perception of the therapeutic process and quality of life, to optimize resources, to decrease hospitalization length, and to potentially prevent nosocomial infections.

It was initially created for specific programs: at-home autologous HSCT for patients with multiple myeloma (MM), at-home consolidation treatment for patients with acute leukemia and intravenous treatment delivery. Since March 2020, when COVID-19 cases rapidly increased in our center, the program expanded to other areas as part of our Hematology department strategy to ensure safe and high quality care for hematological patients during the outbreak.

Patients were included in the program if their residences were within a 30-min drive from our hospital and a caregiver was available at home 24 h a day. All patients gave informed consent before inclusion in the HAH unit.

Concerning the program flowchart, the hematologist prescribed the pharmacotherapeutic treatment in the hospital’s electronic prescription program. Subsequently, the pharmacist checked the prescription. All hazardous medications or those whose stability conditions required it were prepared centrally in laminar flow cabinets in the pharmacy service.

Patients were visited in their homes by nurses for clinical monitoring, blood samples collection, line care and intravenous drugs administration, among other activities. The frequency of visits was determined by clinical criteria and treatment needs. Additionally, patients received regular phone calls from nurses and physicians and 24 h on-call medical coverage was ensured. An inpatient bed in the Hematology ward was always available for these patients in case they would need urgent evaluation to receive rapid care and avoid visits to the emergency room.

After the COVID-19 outbreak, prevention measures were adopted to avoid cross-infection between patients and nurses. Social distancing was recommended and patients and caregivers were advised to stay at home as much as possible. Patients were instructed to immediately communicate any suspected COVID-19 symptoms (fever, cough, or dyspnea), in which case visits were discontinued and patients were evaluated at the hospital. On the other hand, nurses were required to be asymptomatic the day of the visit, and nasopharyngeal swabs were collected during the peak of the outbreak to diagnose asymptomatic cases as well. Nurses wore FFP2 or FFP3 facemasks and hand hygiene was intensified.

At-home autologous HSCT for patients with multiple myeloma

Patients were admitted to the HSCT unit to receive the conditioning regimen with melphalan (140 or 200 mg/m², according to local protocols). Cryotherapy was used to prevent mucositis.

Antimicrobial prophylaxis was performed with oral levofloxacin 500 mg qd, fluconazole 400 mg qd and acyclovir 800 mg bid from day 0. Additionally, intravenous ceftriaxone 1 g qd was administered from day +4 until the first day of febrile neutropenia or until achieving an ANC of 1 x 10^9/L.
Engraftment syndrome primary prophylaxis with oral steroids was administered, according to previous reports [25, 26]. Use of G-CSF was avoided.

Patients were discharged the day after stem cells were infused in case of clinical and analytical stability, were re-admitted in case of severe refractory mucositis, persistent febrile neutropenia or due to physician decision and finally left the at-home program when stable engraftment was achieved and major complications were solved.

At-home consolidation treatment for patients with acute leukemia

Patients were admitted to the Hematology ward to receive high/intermediate dose cytarabine (HDAC/iDAC) and were discharged the day after the cycle was administered providing clinical and analytical stability.

Antimicrobial prophylaxis was performed with oral levofloxacin 500 mg qd and acyclovir 800 mg bid from day 0. Additionally, intravenous ceftriaxone 1 g qd was administered from day +4 until the first day of febrile neutropenia or until achieving an ANC of 1 × 10^9/L. Besides, antifungal prophylaxis with posaconazole was added in this high-risk population.

Patients were re-admitted in case of persistent febrile neutropenia or due to physician decision, and left the program when aplasia and major complications were solved.

Other programs

Clinical and analytical monitoring and electrolyte replacement were performed at home to patients with higher risk of infection, including patients recently treated with autologous or allogeneic HSCT and chimeric antigen receptor (CAR) T cells therapy.

Intravenous antimicrobial agents (including foscarnet) were administered at home either for infections treatment (including febrile neutropenia and bloodstream infections) or prophylaxis.

Venetoclax dose ramp-up was performed at home to patients with chronic lymphocytic leukemia (CLL) and acute myeloid leukemia (AML) with high risk for tumour lysis syndrome (TLS), with close analytical monitoring and intravenous hydration.

Azacitidine was administered at home to patients with AML and myelodysplastic syndrome (MDS).

Bortezomib and carfilzomib were administered at home to patients with MM.

Definitions

At-home episode was defined as any time a patient was included in the HAH unit for a specific program previously described until patient was discharged. One patient could have more than one episode, depending on the treatment delivered.

Statistical analysis

Quantitative variables were expressed as median and range. Qualitative variables were expressed as frequency and percentage. Analysis of normality was performed with the Kolmogorov–Smirnov test. Categorical data and proportions were analysed using chi-squared test or Fisher’s exact test as required. Statistical analysis was performed using the IBM SPSS Statistic program version 25. Follow-up of patients was updated on December 2020.

Results

One hundred five patients were included in the HAH unit between January and November 2020, with a total of 204 at-home episodes (Table 1). Fifty (47%) were female with a median age of 63 years (range 20–92).

Nine at-home autologous HSCT were performed in eight patients with MM (1 patient with high-risk MM received a tandem transplant). Six (75%) were female with a median age of 60 years (range 43–66). All patients were discharged the day after infusion, except one with persistent vomiting, later discharged on day +5. Median CD34(+) cells infused was 4.71 × 10^6/kg (range 4.32–11.7). Four patients developed grade 1 oral mucositis and 2 developed oral candidiasis. One patient needed in-hospital evaluation due to tachycardia, was diagnosed with atrial fibrillation, treated with oral drugs and discharged. Engraftment was achieved in a median time of 17 days (range 14–29) after transplant. Febrile neutropenia and engraftment syndrome rate was 0% and none of the patients were readmitted or diagnosed with SARS-CoV-2 infection. Median duration of the episodes was 18 days (range 13–20).

Three patients were included in the at-home consolidation treatment program for AML: 2 to receive HDAC/iDAC cycle and 1 with high-risk acute promyelocytic leukemia (APL) to administer consolidation chemotherapy according to PETHEMA LPA-2017 protocol. They were discharged from the Hematology ward the day after the cycle was fully administered. All were readmitted: 1 with persistent febrile neutropenia in spite of intravenous antimicrobials and 2 due to SARS-CoV-2 related pneumonia. One patient developed COVID-19 symptoms 48 h after discharge from the Hematology ward and the second patient needed several evaluations in the hospital during admission in the HAH unit for transfusions. HAH unit nurses did not developed SARS-CoV-2 symptoms simultaneously to these patients. Thus, SARS-CoV-2 cross-transmission between these two patients...
and HAH unit members was discarded. Median duration of an episode was 11 days (range 3–13).

Clinical and analytical monitoring or electrolyte replacement was performed at home in 32 patients (43 episodes). Five of these patients were autologous recipients who were early discharged (before achieving engraftment) from HSCT unit. Fifteen were allogeneic HSCT recipients and 2 had received CAR-T cells therapy. Eight patients were admitted to hospital during treatment delivery: 2 with febrile neutropenia, 1 with symptomatic progression disease, 1 with CMV infection and 1 with post-traumatic central nervous system (CNS) hemorrhage. None of these patients were diagnosed with SARS-CoV-2 infection.

Four patients, 3 diagnosed with AML and 1 with CLL, with high risk for TLS, started treatment with venetoclax, and dose ramp-up was performed at home. Prophylactic hydration was administered and blood samples were collected to early diagnose TLS. Two patients developed analytical TLS resolved with hydration. None of these patients were admitted to hospital or diagnosed with SARS-CoV-2 infection.

Azacitidine was administered at home to 22 patients (83 episodes) with AML or MDS. Four patients needed admission during treatment delivery: 2 with febrile neutropenia, 1 with symptomatic disease progression and 1 with CNS hemorrhage. None of them were diagnosed with SARS-CoV-2 infection.

Bortezomib was administered at home to 22 patients (34 episodes) with MM. One patient was admitted to hospital during treatment delivery due to acute limb ischemia and 1 patient was diagnosed with SARS-CoV-2 related pneumonia, at the same time than 2 other members of his family. HAH unit nurses did not develop SARS-CoV-2 symptoms simultaneously to these patients, and in consequence cross-transmission with HAH unit members was discarded.

Carfilzomib was administered at home to 8 patients (12 episodes) with MM. Only 1 scheduled treatment was not administered due to arterial hypertension, and no other incidences related to infusion were reported. One patient was admitted to hospital during treatment delivery due to fever and rash secondary to lenalidomide. None of the patients were diagnosed with SARS-CoV-2 infection.

Overall, more than 1000 visits were made to patient’s homes by nurses from the HAH unit; as a consequence, a median of 17 (IQR 13–19) days of admission per patient to the Hematology ward or HSCT unit and a total 239 visits to the Hematology day-care hospital were avoided. This translated to a significant decrease of bed occupancy in the Hematology ward and HSCT unit during the peak of the pandemic (Figs. 1, 2).

In summary, 28 patients (14% of all episodes) needed admission to the hospital while being treated by the HAH unit, 4 of them because of SARS-CoV-2 related pneumonia (Table 2). None of the personnel from the HAH unit developed SARS-CoV-2 infection while being actively caring patients at home.

### Table 1 Characteristics of patients and at-home episodes

| Patients, n   | 105 |
| Episdes, n    | 204 |
| Median age, years (range) | 63 (20–92) |
| Female, n (%)  | 50 (47%) |
| Diagnosis, n (%) |
| MM    | 35 (33%) |
| AML   | 27 (26%) |
| NHL   | 16 (15%) |
| MDS   | 15 (14%) |
| ALL   | 5 (5%)  |
| Others | 7 (7%)  |

**Program, episodes (%)**

**Admission avoidance programs**

At home autologous HSCT 9 (5%)

At home consolidation in acute leukemia 3 (1.5%)

Monitoring and electrolyte replacement (early discharge) 43 (21%)

Antimicrobial drugs delivery 16 (8%)

Venetoclax dose ramp-up 4 (2%)

**Day-hospital avoidance programs**

Azacitidine 83 (40%)

Bortezomib 34 (16.5%)

Carfilzomib 12 (6%)

**Re-admissions, n (%)**

SARS-CoV-2 related pneumonia 4 (2%)

Other causes 24 (12%)

Febrile neutropenia 7

Progression of underlying disease 4

Other infections 8

Other 5

*ALL acute lymphoid leukemia, AML acute myeloid leukemia, HSCT hematopoietic stem cell transplantation, HL Hodgkin lymphoma, MM multiple myeloma, MDS Myelodysplastic syndrome, NHL non-Hodgkin lymphoma*
Discussion

In this study, we introduce our experience in the implementation of a Hematology HAH unit during the COVID-19 pandemic for caring patients with hematological malignancies and HSCT recipients.

Since March 2020 when the WHO declared the coronavirus-19 disease (COVID-19) a global pandemic, Spain, and particularly the city of Madrid, has been at the
epicenter of the outbreak [15, 27]. During the months of March and April, healthcare system was dramatically overloaded, and hospitals became high risk places for cross-transmission. Social distancing was recommended to help reduce the infection rates. In this adverse context, maintaining appropriate patients care and particularly in cancer patients, became a challenge [28, 29]. Consequently, new strategies to reduce admissions and hospital visits, especially in tertiary centers, and without jeopardizing patients care and safety, needed to be implemented.

Professional care at-home services have significantly increased worldwide lately. HAH programs improve patients quality of life and reduce healthcare costs [22, 23, 30]. Moreover, outpatient approaches avoid unnecessary admissions and day hospital visits. Additionally, previous studies have shown that outpatient strategies can be safe and effective in hematological patients, including HSCT recipients [24, 31–34]. Besides, given a choice between care in institution or at home, most people would prefer to stay at their own home [35]. In this direction, HAH services appear to be an interesting option during COVID-19 pandemic to keep social distancing and reduce hospital visits, particularly for high-risk population [36], and especially during high peaks.

In our centre, the development of a Hematology HAH unit concurred with COVID-19 pandemic emergency. Patients with diverse hematological conditions and varied treatments, were safely and effectively cared in their houses. More than 1000 visits were made to patient’s homes by nurses, allowing high risk patients to continue with their treatments and reducing the exposure to healthcare establishments.

In adult population, the risk for severe COVID-19 increases with age, with older adults at highest risk specially if other comorbidities are associated [5, 37]. Besides, patients with hematological malignancies and HSCT recipients are thought to be at higher risk due to underlying disease, use of immunosuppressive treatments and the need for regular visits to hospitals for different reasons, including treatment delivery and admission, with many reports showing a worse outcome in this particular population and supporting measures to minimize the risk of infection in these patients [9–15].

Without an available vaccine or consistent effective treatments, shielding has been extensively recommended, mainly to high-risk population. Besides, as previously exposed, healthcare system overload has caused difficulty in patient’s care. Thus, there has been concern regarding treatment delay in patients with cancer and its impact on underlying disease outcome [8]. The use of an HAH program can overcome this issue, allowing patients to keep social distancing and to continue or even start new treatments for malignancies.

Previous studies have described an association between COVID-19 cases and recent hospital visits or admission in general population and specifically in patients with cancer and hematological malignancies, including HSCT recipients [11, 13, 38]. Furthermore, the number of infected healthcare workers has been proportionally high during the pandemic, showing that healthcare centres are high-risk

| Table 2 Characteristics and outcome of patients diagnosed with SARS-CoV-2 infection during admission in the HAH unit |
|----------------|----------------|----------------|----------------|----------------|
| Patient  | 1    | 2     | 3     | 4     |
| Sex      | Female | Male  | Female | Female  |
| Age (years) | 47    | 73    | 40    | 79    |
| Diagnosis | AML   | AML   | Hodgkin lymphoma | MM |
| HAH program | AML consolidation treatment | AML consolidation treatment | Clinical monitoring after allo-HSCT | At-home Bortezomib |
| Admission in HAH unit (date) | 26-03-2020 | 24-03-2020 | 20-03-2020 | 07-09-2020 |
| Days in HAH unit | 3    | 10    | 18    | 18    |
| Number of visits from HAH unit | 1    | 6     | 6     | 4     |
| COVID-19 diagnosis (date) | 29-03-2020 | 03-04-2020 | 07-04-2020 | 25-09-2020 |
| Symptoms and signs | Fever, cough and rhinorhoea | Fever and cough | Fever and cough | Fever, cough and dyspnea |
| Pneumonia | Yes | Yes | Yes | Yes |
| Treatment | Lopinavir/ritonavir + HQ/GCs/tocilizumab | HQ/GCs/tocilizumab | HQ/GCs/tocilizumab/Anakinra/HI Plasma | GCs/Rimdesivir/Tocilizumab |
| ICU admission | Yes | No | Yes | No |
| Resolution | Yes | Yes | Yes | Yes |

*allo-HSCT allogeneic hematopoietic stem cell transplantation, AML acute myeloid leukemia, COVID-19 coronavirus-19 disease, GCs glucocorticoids, HAH hospital at home, HI hyperimmune, HQ hydroxychloroquine, ICU intensive care unit, MM multiple myeloma*
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places for transmission [39–42]. In this sense, avoiding visits to the hospital might have an impact in reducing the probability of infection.

In all, only four patients were diagnosed with COVID-19 while receiving any treatment from our HAH unit, in spite of being an elderly and immunocompromised population. Although concerns about the safety of delivering treatments at home during high incidence of SARS-CoV-2 transmission in the community can be posed, taking appropriate measures such as intensifying barrier measures and regular PCR collection to healthcare, can overcome the risk of cross-transmission. In this sense, HAH strategies seem to reduce exposure to SARS-CoV-2 and might decrease the risk of infection of high-risk patients. Of note, none of the cases acquired the infection from HAH personnel. Nevertheless, this is a retrospective analysis, and prospective studies should be conducted to test this hypothesis.

Although cost-effectiveness analysis was not performed in this study, healthcare cost must have been reduced in consequence to previous reports [22, 23, 30], since a median of 17 days of hospitalization per patient and a total of 239 visits to the Hematology day-care hospital were avoided.

On the other hand, implementing these programs requires a remarkable effort from different areas and precisely coordinated actions between physicians, nurses, pharmacists, caregivers and patients, among others. However, we believe this effort is well worth it considering the extremely positive feedback from staff, caregivers and specially from patients [43]. Furthermore, spared hospitalization can be used for more complex procedures and decrease waiting lists.

Overall, implementation of a hematology HAH unit was feasible, safe and provided a thorough advanced care to a high-risk population. Admissions and visits to day hospital were significantly reduced and SARS-CoV-2 infection rate was low and probably not related with cross-transmission between patients and nurses from the unit.

Hence, we believe the use of advanced care at-home strategies can be crucial during times of COVID-19 to minimize treatment interruptions and reduce the risk of cross-infections.

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Author contributions Conception and design: IGC, MK; provision of study materials or patients: all authors; collection and assembly of data: IGC, MK; data analysis and interpretation: all authors; manuscript writing: IGC, MK and final approval of manuscript: all authors.

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Availability of data and materials The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

Code availability Not applicable.

Declarations

Conflict of interests The authors declare that they have no competing interests.

Ethics approval and consent to participate The study was approved by the institutional review board and all patients gave written informed consent for us to collect their clinical information before transplantation or initiation of treatment for research purposes according to the local ethics policy and the Declaration of Helsinki.

Consent for publication All patients provided written informed consent for publication.

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