Second-line treatment for acute graft-versus-host disease with mesenchymal stromal cells: A decision model

Frederick W. Thielen1 | Hedwig M. Blommestein1,2 | Liesbeth E.M. Oosten3 | Friso G. Calkoen4 | Arjan C. Lankester4 | Jaap J. Zwaginga3,5 | Katarina Le Blanc6 | Alba Redondo7 | Fermin Sánchez-Guijo7 | Mattia Algeri8 | Franco Locatelli8 | Wim E. Fibbe3 | Carin A. Uyl-de Groot1

1Erasmus School of Health Policy and Management, Erasmus University, Rotterdam, The Netherlands
2Comprehensive Cancer Organisation, Utrecht, The Netherlands
3Department of Immunohematology and Blood Transfusion, Leiden University Medical Center, Leiden, The Netherlands
4Department of Paediatrics, Leiden University Medical Center, Leiden, The Netherlands
5Center for Clinical Transfusion Research, Sanquin Research, Leiden, The Netherlands
6Department of Laboratory Medicine, Karolinska Institutet, Stockholm, Sweden
7IBSAL-Hospital Universitario de Salamanca, Salamanca, Spain
8Ospedale Pediatrico Bambino Gesù, Rome, Italy

Correspondence
Frederick W. Thielen, Erasmus School of Health Policy and Management, Erasmus University, Rotterdam, The Netherlands. Email: thielen@eshpm.eur.nl

Funding information
This study received funding from the European Union’s Horizon 2020 research and innovation programme under grant agreement number 643580.

Abstract

Objective: No standard second-line treatment exists for acute graft-versus-host disease steroid-refractory (SR-aGvHD), and long-term outcomes remain poor. Mesenchymal stromal cells (MSCs) have been evaluated as treatment, but no disease model (DM) exists that integrates and extrapolates currently available evidence. The aim of this study was to develop such a DM to describe the natural history of SR-aGvHD and to predict long-term outcomes.

Method: The DM was developed in collaboration with experts in haematology-oncology. Subsequently, a model simulation was run. Input parameters for transition and survival estimates were informed by published data of clinical trials on MSC treatment for SR-aGvHD. Parametric distributions were used to estimate long-term survival rates after MSCs.

Results: The newly developed DM is a cohort model that consists of eight health states. For the model simulation, we obtained data on 327 patients from 14 published phase II trials. Due to limited evidence, DM structure was simplified and several assumptions had to be made. Median overall survival was 3.2 years for complete response and 0.5 years for no complete response.

Conclusion: The DM provides a comprehensive overview on the second-line treatment pathway for aGvHD and enables long-term predictions that can be used to perform a cost-effectiveness analysis comparing any treatment for SR-aGvHD.

KEYWORDS
aGvHD, decision model, MSC, paediatric haematology, socioeconomics and ethics, stem cell biology, supportive care, transplantation

1 | INTRODUCTION

Despite decades of research, acute graft-versus-host disease (aGvHD) is still one of the leading causes of death after haematopoietic allogeneic stem cell transplantation (HSCT) for both paediatric and adult patients.1,2 Immunosuppressive therapy with systemic corticosteroids is the first-line treatment option. However, latest available studies estimate that about 50%-70% of the patients...
do not respond to this therapy.\textsuperscript{3–5} Currently, there is no standard second-line treatment available and outcomes in terms of morbidity and mortality remain poor.\textsuperscript{5–9} Since 2004, mesenchymal stromal cells (MSCs) are increasingly studied in phase II clinical trials as a therapeutic option, demonstrating positive treatment effects for steroid-refractory (SR)-aGvHD.\textsuperscript{10,11} However, most of these trials are single-arm studies or case studies or include a limited amount of patients (<50 participants), making reliable and meaningful conclusions challenging. Ideally, the effectiveness of MSC should be tested through phase III randomised controlled trials (RCTs), but these trials are difficult to perform due to regulatory and patient population-related issues.\textsuperscript{11–13} In 2010, results of one RCT (Prochymal; Osiris Therapeutics, Inc. Columbia, Maryland, United States) were released in the form of an unreviewed abstract.\textsuperscript{34} Further, a multicentre RCT is currently being conducted by the European Union Horizon 2020–funded research consortium RETHRIM (ReGeneration ThrougH Im munomodulation)\textsuperscript{15} (ie, the HOVON 113 MSC trial\textsuperscript{16}). The latter study aims to determine the efficacy and cost-effectiveness of MSC as part of a second-line treatment for aGvHD.

In the absence of reliable and conclusive RCT data, several options exist to aggregate and synthesise currently available evidence of MSCs as a treatment option for SR-aGvHD. These include meta-analyses, observational databases, the aggregation of expert opinion or decision analytic modelling.\textsuperscript{17} Thus far, four reviews that include currently available trials testing MSC as treatment for SR-aGvHD have been published,\textsuperscript{11–13,18} of which two are meta-analyses.\textsuperscript{11,18} However, none of the reviews combine available patient-level data (PLD) of the phase II trials to predict and model long-term outcomes of MSC treatment.

The aim of this study was to develop a disease model (DM) to describe the natural history of SR-aGvHD progression and its treatment pathways. The DM can be used to predict long-term outcomes and cost-effectiveness of current (eg, MSC treatment) and future treatment options for SR-aGvHD. To test the practicability of the model, we aimed to gather and implement PLD of a second-line treatment option. Ultimately, our model may facilitate clinical decision-making under conditions of uncertainty.\textsuperscript{19–21} When costs are added, this model can be a valuable tool for reimbursement decision-makers.\textsuperscript{22}

2 MATERIALS AND METHODS

2.1 Part 1: designing and structuring the model

2.1.1 Model characteristics

The aim of the DM was 2-fold. First, it needed to represent the natural history of SR-aGvHD and its treatment pathways in a simplified manner. Therefore, the DM focuses on the SR-aGvHD only until patients either progress, relapse, develop chronic GvHD or die. Second, it needed to be easily adaptable to a cost-effectiveness model at a later stage. Therefore, the DM was built based on clinical expertise, previously published literature\textsuperscript{3,5,10,23,24} and the RETHRIM protocol\textsuperscript{15}. According to the ISPOR recommendations for good modelling practice,\textsuperscript{23} we consulted clinical experts to ascertain that the model represents the disease process and addresses the decision problem of determining which one second-line treatment option is (cost-)effective when compared to another therapy option. We employed a convenience sample to include the clinical experts from the RETHRIM consortium. To be part of RETHRIM, consortium members needed to prove extensive research and treatment experience in the field of HSCT and work at a HSCT specialised treatment centre. All experts involved in this research thus have various professional backgrounds (eg, internal medicine, haematology, oncology or transfusion medicine) and originate from five EU member states (Germany, Italy, The Netherlands, Spain and Sweden). A European perspective on the disease and treatment pathway(s) of SR-aGvHD was hence ensured.

2.1.2 Involvement of experts

Experts were consulted via email, during several telephone conferences and consortium meetings. This process was iterative until a final model version was regarded to fully represent the natural disease and treatment pathway. In addition to the several consultations, the experts were asked to give written feedback on an earlier model version by means of a semistructured questionnaire. Choosing the appropriate model type (eg, decision tree, Markov process) was also based on the ISPOR recommendations for good modelling practice.\textsuperscript{25}

2.2 Part 2: model simulation

2.2.1 Model input parameters

As MSCs are widely studied in numerous phase II clinical trials and case studies since 2004,\textsuperscript{10,13} we selected this treatment option to perform a model simulation. For the model input parameters, we identified relevant studies testing MSCs for the treatment of SR-aGvHD from the recent reviews of Chen et al\textsuperscript{18}, Munneke et al\textsuperscript{12} and Hashmi et al\textsuperscript{11} Additional PLD were obtained for studies whose principal investigator was a member of the RETHRIM consortium.

Data were extracted using a prespecified extraction form aiming at capturing all available data that describe the disease and treatment pathway (see Appendix S1). The clinical effectiveness of MSC treatment was obtained from PLD of the reported first response evaluation or reconstructed from the published Kaplan-Meier curves. Studies not reporting these data were excluded.

To extrapolate the survival data beyond the observed time horizon, we followed the 2013 updated NICE Decision Support Unit recommendations.\textsuperscript{26} Accordingly, considered parametric survival models for the DM included exponential, Weibull, Gompertz, log-logistic, lognormal and generalised gamma distributions. Parametric models were evaluated through visual inspection, Akaike’s information criterion (AIC) and Bayesian information criterion (BIC) tests as well as clinical validity according to the expert group.
2.2.2 | Simulation

We ran a base-case simulation with a hypothetical patient cohort comprising 100 patients with aGvHD grades II-IV. Survival was modelled on a lifetime horizon, whereas it was assumed that patients do not exceed the age of 99 years. Health outcomes of the simulation were expressed in life-years (LYs). As future health effects are valued lower than immediate effects, we adjusted future health outcomes (LYs) to "present values" according to the Dutch guideline for economic evaluations in health care.

All statistical analyses were conducted in RStudio (version 1.0.143, R version 3.4.1). Comprehensive R Archive Network (CRAN) packages used included survival, flexsurv, survminer and plyr.

3 | RESULTS

3.1 | Part 1: designing and structuring the model

3.1.1 | Model characteristics

We opted for a cohort-based Markov model, which represents the most relevant responses and outcomes to second-line treatment for aGvHD in corresponding health states. A Markov model consists of mutually exclusive health states that are associated with different outcomes and costs and provides an efficient structure to simulate a group of patients over time. Patients can change from one health state to another (ie, transit) at specified time intervals (ie, cycles). Outcomes and costs are calculated for the entire time horizon of the model, taking into account the distribution of patients amongst the states at each cycle. Outcomes may entail clinical effectiveness outcomes (eg, response to treatment) as well as health-related quality of life (HRQoL) measures.

The DM comprises eight health states: (a) treatment for SR-aGvHD II-IV, (b) response to treatment (complete or partial response), (c) sustained response, (d) treatment failure (stable or progressive disease), (e) relapse/persistent aGvHD, (f) third-line aGvHD therapy, (g) relapse or adverse events of haematologic disease requiring reinitiation or intensification of immunosuppression and (h) death (see Figure 1 for the model diagram). These states were primarily based on the HOVON113 MSC treatment protocol and expert opinion (see section "Involvement of experts"). Four response categories are defined: complete response (CR) is defined as the absence of all signs and symptoms of aGvHD; partial response (PR) is the improvement of aGvHD by at least one grade; stable disease (SD) is no change in aGvHD; and progressive disease (PD) is the worsening of aGvHD by at least one grade. The cycle duration was set at 28 days according to the recommendation by Martin and colleagues on the standardised time period to evaluate aGvHD response. In addition, we applied a half-cycle correction for the calculation of the model outcomes.

The model starts with treatment for SR-aGvHD (a) at time 0 ($T_0$). Within 28 days after treatment ($T_{28}$), patients either respond [ie, transit to the response to treatment health state (b)] or have a treatment failure (d) or die (h). Responders (b) can have a sustained

![Figure 1](image)

* $T = \text{time in days post-MSC transfusion; } AE = \text{adverse events of GvHD treatment; } * = \text{requiring reinitiation or intensification of immunosuppression}.*
response (c), relapse or have a persistent aGvHD grade that requires the reinitiation or intensification of immunosuppression (e), or enter third-line therapy (f).

Patients with a treatment failure to the initial MSC treatment (d) will directly receive third-line treatment (f). Adverse events (AEs) of treatment are not defined as a separate health state, but are possible within health states (b), and (d)–(f). At any time and from any health state, patients can transit to relapse or experience of an adverse event of the underlining haematological disease (g), or death (h). Death is an absorbing health state, meaning that once entered, patients remain in this health state.

In the model, all health states are defined as mutually exclusive although in clinical practice patients may sometimes fit the criteria of multiple health states. In these cases, patients are assigned to the health state with the largest impact on outcomes and costs. For example, patients with treatment failure [health state (d)] may experience a relapse of the disease [health state (g)]. These patients are assigned to the relapse health state (g) although they also fit the criteria for treatment failure. The increasing numbering of the model states indicates the expected increasing impact on both outcomes and costs, according to expert opinion.

3.1.2 | Involvement of experts

The team of clinical experts consisted of eleven members from five different countries (Germany, Italy, the Netherlands, Spain and Sweden). A preliminary model based on the HOVON113 MSC treatment protocol was presented to all consortium members at the third RETHRIM consortium meeting. Because the structure of the model was deemed incomplete, alternative treatment pathways and additional health states were discussed in detail. This led to a refinement of the existing model states and to the addition of the following health states: relapse or adverse events of haematological disease, and adverse events. The new version of the model was then presented and discussed at the fourth RETHRIM consortium meeting. To allow for written feedback, experts were asked to fill out a semistructured questionnaire at the fifth consortium meeting. Subsequently, the model was amended and a final version was approved by all experts.

3.2 | Part 2: Model simulation

3.2.1 | Model input parameters

We detected 18 studies that reported on phase II trials and collection of case studies4,33–49, of which 9 reported PLD33–38,43,49 (see Appendix S2). Unpublished4,42 and additional data46 were requested and received for three studies. Two studies could be integrated by reconstructing the published Kaplan-Meier curves.44,48 Three studies were excluded because they did not publish survival data,47,50 or the proportion of patients in the response categories.41 Patients reported in the study by Ringdén et al45 were not included as they had been already included in the data presented by Le Blanc and colleagues.4 From Luchini et al49, only four patients with aGvHD stages II-IV were included.

In total, we extracted data from 327 patients from 14 studies to estimate the proportion of individuals in the different health states and the proportion of patients changing between these states (ie, transition probabilities).31 Age and sex were only reported for 177 and 152 patients, respectively, which made the originally planned subgroup analysis for age and sex impossible. All patients had aGvHD grades II–IV prior to MSC treatment; for 204 cases, the exact grade was known (18.6% aGvHD II, 45.6% aGvHD III and 35.8% aGvHD IV). Response categories in the underlying studies were defined heterogeneously. Whereas one study did not provide any definition for the response categories employed,37 only complete response (CR = the complete resolution of all aGvHD symptoms) was unanimously defined in all other studies. Therefore, all response categories other than CR were grouped as “no complete response” (nCR).

Due to the lack of sufficiently detailed observational data, we were forced to use a simplified version of our model simulation. In this version, we could only model the first response to treatment (at day +28) and long-term survival of patients after the first response evaluation to MSCs. The health states sustained response,4 relapse or persistent aGvHD,3 third-line therapy, chronic GvHD,7 relapse or adverse events of haematological disease,8 and adverse events in general could not be modelled.

3.2.2 | Transition probabilities for the first two model cycles

Reported mean and median time of first evaluation was 26.6 and 28 days, respectively (range = 2-58). To integrate all available observations into the model, we assumed that all responses were evaluated within the first 28 days after MSC treatment. Based on this, we calculated a transition matrix showing the probability of response to MSC after day 28 (see Table 1).

Table 1 shows that 43.4% patients with aGvHD II-IV had a CR at first response evaluation whereas 43.7% had nCR, and 12.8% died within the first 28 days.

3.2.3 | Long-term survival estimation

To extrapolate survival estimates beyond study observations, survival data, available for 235 patients (115 CR, 120 nCR), were used. For the excluded cases, the last time of follow-up or time to death was not reported. Patients who died before day 28 were already included in the 28-day transition rates to death. Median survival times were reached at approximately five years (1819 days) for complete responders and at approximately four months (115 days) for nCR.

Kaplan-Meier curves and fitted parametric models for survival are depicted in Figure 2 for both response categories. AIC and BIC values are presented in Appendix S3. From a statistical point of view, the fit of the parametric models to the empirical data was similar. Nevertheless, the extrapolation after observation was different. For instance, the extrapolation according to
the generalised gamma function predicted that 40% of the CR patients survived more than 25 years after MSC treatment. This was not deemed plausible from a clinical perspective by the experts. Based on clinical expertise, the lognormal distribution would present the best balance between the statistical tests of fit and visual inspection for both CR and nCR.

### 3.3 Results of the model simulation

At the 28-day treatment evaluation, of the 100 simulated cases with aGvHD II-IV, approximately 43 and 44 observations had a CR and nCR, respectively. The median survival time modelled for CR was 3.2 and 0.5 years for nCR. Overall median survival for all patients irrespective of their response category was 9.6 months. Average per-person life-years were estimated at 0.32 years (ie, 3.8 months).

In Figure 3, modelling results are plotted on the empirical Kaplan-Meier curves for CR and nCR. Table 2 depicts the estimated overall survival probabilities for different years after the first response evaluation.

### 4 DISCUSSION

In this study, we present a DM for the second-line treatment of aGvHD. In line with current recommendations for good modelling practice, we involved an international group of clinical experts to develop this DM and used evidence available from the literature. Consequently, we ensured that the model sufficiently represents current clinical practice.

The DM can serve manifold purposes. Updated with clinical evidence, it can be used by clinicians and researchers to estimate long-term health outcomes of different treatment alternatives. When costs are added to the clinical evidence, for example, from RETHRIM, economic evaluations can be performed to inform reimbursement decision-makers on the implementation of treatment alternatives.

#### TABLE 1 Transition rates from pre- to post-MSC treatment

| From   | To     | CR    | nCR   | Death |
|--------|--------|-------|-------|-------|
| aGvHD II-IV | 43.4%  | 43.7% | 12.8% |

CR: complete response; nCR: no complete response.
To run a first model simulation, we searched for available evidence on both costs and clinical evidence on treatment alternatives for aGvHD. We found that MSCs for the treatment of SR-aGvHD are widely studied. Therefore, we were able to integrate more than a decade of empirical data into our DM. Nevertheless, mainly due to the absence of randomised phase III studies, the number of patients included and the restricted follow-up periods, we faced several challenges in integrating the collected information. Consequently, we had made a number of assumptions.

First, MSC products and their administration varied between the studies. With the exception of the study by Fang et al. where MSCs were derived from human adipose tissue, all other studies used bone marrow–derived products. In addition, the number of MSC infusions (between 1 and 21 infusions) as well as the dosage of infused MSCs (between 0.6 and 20 × 10^6 cells per kg body weight) varied across the studies. In this regard, we assumed that type and administration of MSC products had no effect on transition probabilities or mortality rates. Whereas this made the integration of the study results possible, the data did not allow for further stratification to test potential MSC derivation–related or dose–related effects on the response rates.

Second, there were not enough data available to estimate transitions between response categories after a first response evaluation. We had assumed survival can be predicted based on the initial response category at 28 days post-MSC transfusion. However, in clinical practice response categories may change in any direction after the first evaluation. Of the included studies, only Prasad et al. assessed response to MSCs in paediatric patients more than once. In this study, five of twelve patients further improved after day 32 to a complete remission. Future studies measuring response on several time points after MSC treatment could inform our model on subsequent changes in response categories. Whereas this enables a better estimate for subgroups, the effect on the average survival of the entire population is most likely negligible.

Third, it needs to be noted that the simulation did not consider the underlying haematological disease, nor the patients’ age or sex. This choice was made as not all studies did report on these variables and any further stratification would have resulted in a reduction in the population on which the different estimates are based. However, we acknowledge that the underlying haematological disease and age and sex can be important determinants for long-term survival. Detailed reporting on patient characteristics and their diagnosis may help to enable further analyses for these subgroups.

Our modelling results, however, did show the expected longer survival after MSC transfusion for patients that achieved CR at first response evaluation, when compared to patients with nCR. These estimates are in line with previous findings highlighting the importance of complete response for long-term survival.

To our knowledge, there are no long-term survival results published for SR-aGvHD patients treated with MSCs other than the two reports that are based on studies included in our study. Therefore, we attempted to compare our results to survival estimates of studies that tested other second-line treatment options for aGvHD. To find suitable studies, we consulted the most recent NHS England clinical commissioning policy on the treatments for GvHD following HSCT. This guideline is based on an extensive and updated review of the literature and concludes that there is sufficient evidence only for extracorporeal photopheresis (ECP) to be routinely commissioned for the treatment of aGvHD. Therefore, we focussed on a comparison with ECP. For other treatment options such as infliximab, etanercept, inolimomab, alemtuzumab, pento‐statin or MSCs, the reviewers found that there was no sufficient evidence available to propose the routine commissioning for aGvHD.

Regarding survival of aGvHD patients treated with ECP, the largest series was published by Greinix et al. in a phase II prospective study. Every week, 59 patients with steroid-refractory or steroid-dependent GvHD received two consecutive ECP treatments. CR and nCR was defined as in our study. The reported median survival was below 6 months after HSCT for patients with nCR, confirming our median survival estimates for nCR to MSC of approximately 6 months. In the study of Greinix et al., median survival for complete responders was never reached during the reported follow-up period of 9 years after HSCT. For this study, this implies that the probability of surviving nine years after HSCT would be approximately 59% for patients with a CR. With an estimate of approximately 27% at ten years post-MSCs, our estimates for patients with a CR are significantly lower. This may be explained by differences in patient population before treatment (ie, Greinix et al. included a higher number of patients with aGvHD grade II (61%)). In addition, the study by Greinix and colleagues was based on a limited amount of complete responders (n = 41).

Although we were able to demonstrate longer survival for patients with CR when compared to nCR, the relatively high mortality rate of SR-aGvHD, even with MSCs, can still be regarded as unacceptably high. And although there were numerous studies reporting on MSCs as a treatment alternative for SR-aGvHD, none of them included patient-reported outcomes (PROs) such as HRQoL. Updating our results with HOVON113 findings, including HRQoL measures, might improve the current survival estimates and show favourable treatment outcomes in terms of quality of life for MSCs when compared to placebo. However, until these study results are presented, this remains subject

| Response category | Years after the first response evaluation |
|-------------------|------------------------------------------|
|                   | 1  | 2  | 5  | 10 | 20 | 50 | 80 | 99 |
| CR                | 73.1 | 59.6 | 40.1 | 26.7 | 16.0 | 6.8 | 4.1 | 0  |
| nCR               | 25.7 | 11.4 | 2.6  | 0.6  | 0.1  | 0   | 0   | 0  |
| All               | 43.0 | 30.9 | 18.6 | 11.9 | 7.0  | 3.0 | 1.8 | 0  |
5 | CONCLUSION

The designed DM provides a comprehensive overview on the second-line treatment pathways for aGVHD in general. The model simulation with data from previously published studies on MSCs as a second-line treatment option for aGVHD presented outcomes matching the literature as well as clinical expectations. This demonstrates the practicability and usefulness of the model. However, to date, only insufficiently detailed data are available to fully model all health states and to perform a cost-effectiveness analysis. The yet restricted model simulation would therefore benefit from additional data, preferably from a phase III RCT. The integration of effectiveness results together with health-related quality of life measures (eg, from the EQ-5D questionnaires) and different cost components derived from the RETHRIM trial could overcome this limitation and enable a full cost-utility analysis.

ACKNOWLEDGEMENTS

We gratefully acknowledge the valuable input from participants at the 2017 lowlands Health Economists’ Study Group (lolaHESG) conference in Rotterdam, The Netherlands. We also thank all RETHRIM members for their manifold ways of supporting this work, especially all members of the clinical expert group: Katarina Le Blanc, Wim Fibbe, Franco Locatelli, Dietger Niederwieser, Liesbeth Oosten, Fermín Sánchez-Guijo and Jaap Zwaginga. We also acknowledge Ida Duprez for her contribution to a thorough data analysis.

ORCID

Frederick W. Thielen  
http://orcid.org/0000-0002-0312-5891

REFERENCES

1. D’Souza A, Zhu XC. Uses and outcomes of hematopoietic cell transplantation (HCT): CIBMTR summary slides, 2016. https://www.cibmtr.org
2. Nassereddine S, Rafei H, Elbahesh E, Tabbara I. Acute graft-versus-host disease: a comprehensive review. Anticancer Res. 2017;37:1547-1555.
3. Garnett C, Aperley JF, Pavlu J. Treatment and management of graft-versus-host disease: improving response and survival. Ther Adv Hematol. 2013;4:366-378.
4. Le Blanc K, Frassoni F, Ball L, et al. Mesenchymal stem cells for treatment of steroid-resistant, severe, acute graft-versus-host disease: a phase II study. The Lancet. 2008;371:1579-1586.
5. Deeg HJ. How I treat refractory acute GVHD. Blood. 2007;109:4119-4126.
6. Martin PJ, Rizzo JD, Wingard JR, et al. First- and second-line systemic treatment of acute graft-versus-host disease: recommendations of the american society of blood and marrow transplantation. Biol Blood Marrow Transplant. 2012;18:1150-1163. https://doi.org/10.1016/j.bbmt.2012.04.005.
7. Kim SS. Treatment options in steroid-refractory acute graft-versus-host disease following hematopoietic stem cell transplantation. Ann Pharmacother. 2007;41:1436-1444.
8. Westin JR, Saliba RM, De Lima M, et al. Steroid-refractory acute GVHD: predictors and outcomes. Adv Hematol. 2011;2011:1–8.
9. NHS England. (NHS England, 2017). Clinical Commissioning Policy: Treatments for Graft versus Host Disease (GvHD) following Haematopoietic Stem Cell Transplantation. https://www.england.nhs.uk/wp-content/uploads/2017/03/gvhd-hematopoietic-stem-cell.pdf. Accessed January 24, 2018.
10. Le Blanc K, Rasmussen I, Sundberg B, et al. Treatment of severe acute graft-versus-host disease with third party haploidentical mesenchymal stem cells. The Lancet. 2004;363:1439-1441.
11. Hashmi S, Ahmed M, Murad MH, et al. Survival after mesenchymal stromal cell therapy in steroid-refractory acute graft-versus-host disease: systematic review and meta-analysis. Lancet Haematol. 2016;3:e45-e52.
12. Munneke JM, Spruit MJ, Cornelissen AS, vanHoeven V, Voermans C, Hazenberg MD. The potential of mesenchymal stromal cells as treatment for severe steroid-refractory acute graft-versus-host disease: a critical review of the literature. Transplantation. 2016;100(11): 2309-2314.
13. Amorin B, Alegreppi AP, Valim V, et al. Mesenchymal stem cell therapy and acute graft-versus-host disease: a review. Human cell. 2014;27:137-150.
14. Martin P, et al. Prochymal improves response rates in patients with steroid-refractory acute graft versus host disease (SR-GVHD) involving the liver and gut: results of a randomized, placebo-controlled, multicenter phase III trial in GVHD. Biol Blood Marrow Transplant. 2010;16:5169-5170.
15. RETHRIM, www.rethrim.eu (2017).
16. HOVON. HOVON 113 MSC, https://www.hovon.nl/studies/studies-per-ziektebeeld/sct.html?action=showstudie&studie_xmendid=97&categorie_xmlid=11 (n.d.).
17. Gale R. Markov model of CLL transplants. Bone Marrow Transplant. 2012;47:1145.
18. Chen X, Wang C, Yin J, et al. Efficacy of mesenchymal stem cell therapy for steroid-refractory acute graft-versus-host disease following allogeneic hematopoietic stem cell transplantation: a systematic review and meta-analysis. Plos One. 2015;10:e0136991. https://doi.org/10.1371/journal.pone.0136991.
19. Goeree R, Diaby V. Introduction to health economics and decision-making: is economics relevant for the frontline clinician? Best Pract Res Clin Gastroenterol. 2013;27:831-844. https://doi.org/10.1016/j.bgpg.2013.08.016.
20. Beck JR, Pauker SG. The Markov process in medical prognosis. Med Decis Making. 1983;3:419-458.
21. Sonnenberg FA, Beck JR. Markov models in medical decision making: is economics relevant for the frontline clinician? Med Decis Making. 1993;13:322-338.
22. Drummond M, Sculpher MJ, Claxton K, Torrance GW. Methods for the Economic Evaluation of Health Care Programmes. Oxford, UK: Oxford University Press, 2015.
23. Digan FL, Clark A, Amrolia P, et al. Diagnosis and management of acute graft-versus-host disease. Br J Haematol. 2012;158:30-45.
24. Glucksberg H, Storb R, Fefer A, et al. Clinical manifestations of graft-versus-host disease in human recipients of marrow from HL-A-matched sibling donors. Transplantation. 1974;18:295-304.
25. Roberts M, Russell LB, Paltiel AD, et al. Conceptualizing a model of survival analysis for economic evaluations alongside clinical trials—extrapolation with patient-level data (2011). http://nicedsu.org.uk/wp-content/uploads/2016/03/NICE-DSU-TSD-Survival-analysis-updates-March-2013.v2.pdf. Accessed January 24, 2018.

27. Martin PJ, Counts GW, Appelbaum FR. Life expectancy in patients surviving more than 5 years after hematopoietic cell transplantation. J Clin Oncol. 2010;28:1011-1016.

28. Zorginstituut Nederland. Guideline for economic evaluations in healthcare (2016). https://www.zorginstituutnederland.nl/binaries/content/documents/znl-wwv/documenten/publicaties/publicaties-in-engels/2016/1606-guideline-for-economic-evaluations-in-healthcare/Guideline+for+economic+evaluations+in+healthcare.pdf

29. Severens JL, Milne RJ. Discounting health outcomes in economic evaluation: the ongoing debate. Value Health. 2004;7:397-401.

30. Briggs AH, Claxton K, Sculpher MJ. An introduction to Markov modelling for economic evaluation. Oxford United Kingdom: Oxford University Press. Handbooks in Health Economic. E. 2006.

31. Martin PJ, Bachier CR, Klingemann HG, et al. Endpoints for clinical trials testing treatment of acute graft‐versus‐host disease: a consensus document. Biol Blood Marrow Transplant. 2009;15:777. https://doi.org/10.1016/j.bbmt.2009.03.012.

32. Fang B, Song Y, Liao L, Zhang Y, Zhao R. Favorable Response to Umbilical Cord-Derived Mesenchymal Stem Cells in Steroid-Refractory Acute Graft-Versus-Host Disease. Transplantation Proceedings. 2007;39:10, 3358–3362 (Elsevier). https://doi.org/10.1016/j.transproceed.2007.08.103.

33. Von Bonin M, Stözel F, Goedecke A, et al. Treatment of refractory acute GVHD with third-party MSC expanded in platelet lysate-containing medium. Bone Marrow Transplantation. 2009;43:245-251.

34. Müller I, Kordwich S, Holzwarth C, et al. Application of multipotent mesenchymal stromal cells in pediatric patients following allogeneic stem cell transplantation. Blood Cells Mol Dis. 2008;40:25-32.

35. Prasad VK, Lucas KG, Kleiner GL, et al. Efficacy and safety of ex vivo cultured adult human mesenchymal stem cells (Prochymal™) in pediatric patients with severe refractory acute graft-versus-host disease in a compassionate use study. Biol Blood Marrow Transplant. 2011;17:534-541.

36. Pérez-Simon JA, Lopez-Villar O, Andreu EJ, et al. Mesenchymal stem cells expanded in vitro with human serum for the treatment of acute and chronic graft-versus-host disease: results of a phase I/II clinical trial. Haematologica. 2011;96:1072-1076.

37. Herrmann R, Sturm M, Shaw K, et al. Mesenchymal stromal cell therapy for steroid-refractory acute and chronic graft versus host disease: a phase 1 study. Int J Hematol. 2012;95:182-188.

38. Chen G, Yang T, Tian H, et al. Safety and efficacy of mesenchymal stromal cells (Prochymal™) in patients with severe acute graft-versus-host disease after allogeneic hematopoietic stem cell transplantation. Blood. 2010;15:777.

39. Jacobsohn DA, Hallick J, Anders V, et al. Infliximab for steroid-refractory acute GVHD: a case series. Am J Hematol. 2003;74:119-124.

40. Remmerber M, Ringdén O. Treatment of severe acute graft-versus-host disease with mesenchymal stromal cells: a comparison with non-MSC treated patients. Int J Hematol. 2012;96:822-824. https://doi.org/10.1007/s12185-012-1218-3.

41. Ho VT, Zahrin D, Hochberg E, et al. Safety and efficacy of defludeluxin difltix in patients with steroid-refractory acute graft-versus-host disease after allogeneic hematopoietic stem cell transplantation. Blood. 2004;104:1224-1226.

42. Ball LM, Bernardo ME, Roelofs H, et al. Multiple infusions of mesenchymal stromal cells induce sustained remission in children with steroid-refractory, grade III-IV acute graft-versus-host disease. Br J Haematol. 2013;163:501-509.

43. Introna M, Lucchini G, Dander E, et al. Treatment of graft versus host disease with mesenchymal stromal cells: a phase I study on 40 adult and pediatric patients. Biol Blood Marrow Transplant. 2014;20:375-381.

44. Kurtzberg J, Prokop S, Teira P, et al. Allogeneic human mesenchymal stem cell therapy (remestemcel-L, Prochymal) as a rescue agent for severe refractory acute graft-versus-host disease in pediatric patients. Biol Blood Marrow Transplant. 2014;20:229-235.

45. Ringdén O, Uzunel M, Rasmusson I, et al. Mesenchymal stem cells for treatment of therapy-resistant graft-versus-host disease. Transplantation. 2006;81:1390-1397.

46. Sánchez-Guijo F, Caballero-Velázquez T, López-Villar O, et al. Sequential third-party mesenchymal stromal cell therapy for refractory acute graft-versus-host disease. Biol Blood Marrow Transplant. 2014;20:1580-1585.

47. Zhao K, Lou R, Huang F, et al. Immunomodulation effects of mesenchymal stromal cells on acute graft-versus-host disease after hematopoietic stem cell transplantation. Biol Blood Marrow Transplant. 2015;21:97-104.

48. Te Boome L, Mansilla C, van der Wagen LE, et al. Biomarker profiling of steroid-resistant acute GVHD in patients after infusion of mesenchymal stromal cells. Leukemia. 2015.

49. Luchini G, Gaipa G, Perseghin P, et al. Platelet-lysate-expanded mesenchymal stromal cells as a salvage therapy for severe resistant graft-versus-host disease in a pediatric population. Biol Blood Marrow Transplant. 2010;16:1293-1301.

50. Chen GH, Yang T, Tian H, et al. [Clinical study of umbilical cord-derived mesenchymal stem cells for treatment of nineteen patients with steroid-resistant severe acute graft-versus-host disease]. Zhonghua xue ye xue za zhi. 2012;33:303-306.

51. Martin PJ, Schoch G, Fisher L, et al. A retrospective analysis of therapy for acute graft-versus-host disease: secondary treatment. Blood. 1991;77:1821-1828.

52. Weisdorf D, Haake R, Blazar B, et al. Treatment of moderate/severe acute graft-versus-host disease after allogeneic bone marrow transplantation: an analysis of clinical risk features and outcome. Blood. 1990;75:1024-1030.

53. Martin PJ, Schoch G, Fisher L, et al. A retrospective analysis of therapy for acute graft-versus-host disease: initial treatment. Blood. 1990;76:1464-1472.

54. Greinix HT, Knobler RM, Worel N, et al. The effect of intensified extracorporeal photochemotherapy on long-term survival in patients with severe acute graft-versus-host disease. Haematologica. 2006;91:405-408.

55. Ho VT, Zahrieh D, Hochberg E, et al. Safety and efficacy of defludeluxin difltix in patients with steroid-refractory acute graft-versus-host disease after allogeneic hematopoietic stem cell transplantation. Blood. 2004;104:1224-1226.

56. Jacobsohn DA, Hallick J, Anders V, et al. Infliximab for steroid-refractory acute GVHD: a case series. Am J Hematol. 2003;74:119-124.

57. Remmerber M, Ringdén O. Treatment of severe acute graft-versus-host disease with mesenchymal stromal cells: a comparison with non-MSC treated patients. Int J Hematol. 2012;96:822-824. https://doi.org/10.1007/s12185-012-1218-3.

58. von Bahr L, Sundberg B, Lönnies L, et al. Long-Term Complications, Immunologic Effects, and Role of Passage for Outcome in Mesenchymal Stem Cell Therapy. Biol Blood Marrow Transplant. 2012;18:557-564. https://doi.org/10.1016/j.bbmt.2011.07.023.

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

How to cite this article: Thielen FW, Blommestein HM, Oosten LEM, et al. Second-line treatment for acute graft-versus-host disease with mesenchymal stromal cells: A decision model. Eur J Haematol. 2018;101:676-683. https://doi.org/10.1111/ejh.13158