

Anti B-Cell Maturation Antigen CAR T-cell and Antibody Drug Conjugate Therapy for Heavily Pre-Treated Relapsed and Refractory Multiple Myeloma

Evidence Report

April 5, 2021

Prepared for



This evidence report was updated on April 5, 2021 with corrections that resulted in small changes in the Long-Term Cost-Effectiveness section. No results were changed.

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Sei Lee served as the lead author for the report and wrote the background, other benefits, and contextual considerations sections. Molly Beinfeld led the systematic review and wrote the clinical effectiveness section of the report in collaboration with Noemi Fluetsch and Belén Herce-Hagiwara. Brett McQueen, Eric Gutierrez, and Sue Kwon developed the cost-effectiveness model and authored the corresponding sections of the report. Melanie Whittington provided methods guidance for the cost-effectiveness modeling effort and conducted the potential budget impact analysis. Daniel Ollendorf and Steven Pearson provided methodologic guidance on the clinical and economic evaluations. We would also like to thank Monica Frederick and Liis Shea for their contributions to this report.

About ICER

The Institute for Clinical and Economic Review (ICER) is an independent non-profit research organization that evaluates medical evidence and convenes public deliberative bodies to help stakeholders interpret and apply evidence to improve patient outcomes and control costs. Through all its work, ICER seeks to help create a future in which collaborative efforts to move evidence into action provide the foundation for a more effective, efficient, and just health care system. More information about ICER is available at http://www.icer.org.

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For drug topics, in addition to receiving recommendations <u>from the public</u>, ICER scans publicly available information and also benefits from a collaboration with <u>IPD Analytics</u>, an independent organization that performs analyses of the emerging drug pipeline for a diverse group of industry stakeholders, including payers, pharmaceutical manufacturers, providers, and wholesalers. IPD provides a tailored report on the drug pipeline on a courtesy basis to ICER but does not prioritize topics for specific ICER assessments.

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The Midwest Comparative Effectiveness Public Advisory Council (Midwest CEPAC) – a core program of ICER – provides a public venue in which the evidence on the effectiveness and value of health care services can be discussed with the input of all stakeholders. Midwest CEPAC seeks to help patients, clinicians, insurers, and policymakers interpret and use evidence to improve the quality and value of health care.

The Midwest CEPAC is an independent committee of medical evidence experts from across the Midwest, with a mix of practicing clinicians, methodologists, and leaders in patient engagement and advocacy. All Council members meet strict conflict of interest guidelines and are convened to discuss the evidence summarized in ICER reports and vote on the comparative clinical effectiveness and value of medical interventions. More information about Midwest CEPAC is available at https://icer.org/who-we-are/people/independent-appraisal-committees/midwest-comparative-effectiveness-public-advisory-council-m-cepac/.

The findings contained within this report are current as of the date of publication. Readers should be aware that new evidence may emerge following the publication of this report that could potentially influence the results. ICER may revisit its analyses in a formal update to this report in the future.

The economic models used in ICER reports are intended to compare the clinical outcomes, expected costs, and costeffectiveness of different care pathways for broad groups of patients. Model results therefore represent average findings across patients and should not be presumed to represent the clinical or cost outcomes for any specific patient. In addition, data inputs to ICER models often come from clinical trials; patients in these trials may differ in real-world practice settings. In the development of this report, ICER's researchers consulted with several clinical experts, patients, manufacturers, and other stakeholders. The following experts provided input that helped guide the ICER team as we shaped our scope and report. It is possible that expert reviewers may not have had the opportunity to review all portions of this report. None of these individuals is responsible for the final contents of this report, nor should it be assumed that they support any part of it. The report should be viewed as attributable solely to the ICER team and its affiliated researchers.

For a complete list of stakeholders from whom we requested input, please visit: http://icerorg.wpengine.com/wp-content/uploads/2020/10/ICER_MM_Key_Stakeholders_092720.pdf

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List of Acronyms and Abbreviations Used in this Report

95% Confidence Interval
Antibody drug conjugate
Adverse event
Agency for Healthcare Research and Quality
Academic in confidence
B-cell maturation antigen
Bendamustine + prednisone
Chimeric antigen receptor
Complete response
Calcium, renal, anemia, and bone
Cytokine release syndrome
Discontinuation
Dexamethasone + cyclophosphamide + etoposide + cisplatin
Duration of response
Extramedullary disease
European Organization for Research and Treatment of Cancer
Elotuzumab + pomalidomide + dexamethasone
Hazard ratio
Health-related quality of life
Immunomodulatory drug
Interquartile range
Idiopathic subglottic stenosis
Intention to treat
Carfilzomib + cyclophosphamide + dexamethasone
Kilogram
Carfilzomib + pomalidomide + dexamethasone
Milligram
Multiple Myeloma
Minimal residual disease
Number
Total number
Not applicable
Not estimable
National Institute for Health and Care Excellence
Not reported
Neurotoxicity
Odds ratio
Overall response rate
Overall survival
Ocular Surface Disease Index
Pomalidomide + cyclophosphamide + dexamethasone
Progression free survival
Proteasome inhibitor
Quality of Life Questionnaire C30
Relapsed or refractory multiple myeloma
Survey, Epidemiology, and End Results Program
Triple-class refractory multiple myeloma
Time to next treatment

Executive Summary

Multiple myeloma (MM) is a hematologic cancer of plasma cells, currently estimated to afflict approximately 150,000 Americans. The mainstays of current MM treatment include immunomodulatory agents, proteasome inhibitors and anti-CD38 monoclonal antibodies.¹ Most patients eventually relapse; these patients with relapsed or refractory multiple myeloma (RRMM) often cycle through different combinations of agents. When a patient's disease is no longer responsive to agents in each of the three classes, the disease is referred to as "triple-class refractory" MM (TCRMM).²

ICER reviewed three new treatments targeting the B-cell maturation antigen (BCMA) for heavily pre-treated patients with RRMM who have cycled through numerous previous lines of therapy. Belantamab mafodotin blmf (Blenrep[®], GlaxoSmithKline) is an antibody drug conjugate, with a monoclonal antibody to BCMA linked to a cytotoxic drug. Belantamab was studied in patients with heavily pre-treated (6-7 previous lines of therapy) TCRMM (majority quad- and penta-refractory, usually defined as refractory to 4 or 5 agents across all 3 drug classes outlined above). Idecabtagene vicleucel ("ide-cel", Abecma[®], Bristol Myers Squibb and bluebird bio) and ciltacabtagene autoleucel ("cilta-cel", Janssen and Legend biotech) are chimeric antigen receptor (CAR) T-cell therapies, involving engineering a patient's own T cells to target BCMA, and were studied in patients who were mostly TCRMM (majority triple- or quad-refractory patients).

Patients spoke about the burden of symptoms from both MM and its available treatments. Common symptoms of disease include fatigue, which can be overwhelming, and bony pain. Symptoms of the current treatments vary by medication, but frequently mentioned bothersome side effects include neuropathy as well as insomnia and psychosis from dexamethasone. Patients also noted substantial financial burden with annual out-of-pocket costs exceeding \$10,000 leading one patient to remark that one had to be a "mathematician" to navigate the costs of being a myeloma patient.

Response rates and survival statistics are presented in Table ES1. The CAR T-cell therapies (ide-cel and cilta-cel) appear to be superior to currently available treatment regimens for TCRMM, as estimated from the recent MAMMOTH observational study. In contrast, belantamab appears to be equivalent or slightly superior to the most relevant comparative set from MAMMOTH.

Intervention	Study	Follow- Up Duration	As Treated ORR	ITT ORR	Median PFS or OS [*]	Toxicity
	CAR T Popu	lation (Triple	e- or Quad-	Refractory,	3+ prior lines of t	reatment)
lde-cel	KarMMa	13.3 months	73%	63%	As-treated PFS = 8.6 months	51% CRS Grade 2+
Cilta-cel	CARTITUDE- 1	12.4 months	97%	75%	As-treated PFS >12.4 months	44% CRS Grade 2+ 6% Treatment-related deaths
Usual Care	MAMMOTH	10.6 months		31%	PFS = 3.4 months	Variable
Belan	tamab Populat	ion (Triple-, 0	Quad- or Pe	enta- Refrac	tory MM, 4+ prior	lines of treatment)
Belantamab	DREAMM-2	13 months		32%	ITT OS = 13.8 months	18 – 46% Meaningful to moderate reversible visual decline (duration 22-33 days)
Usual Care	MAMMOTH subcohort ⁺	10.6 months		28%	Triple/quad OS = 9.2 months Penta OS = 5.6 months	Variable

Table ES1. Response Rates and Median PFS for Anti-BCMA Therapies

BCVA: Best Corrected Visual Acuity, CRS: cytokine release syndrome, ITT: intention-to-treat, ORR: overall response rate, OS: overall survival, PFS: progression free survival

* Ide-cel and cilta-cel PFS is as-treated. All other PFS and OS data are ITT

⁺ MAMMOTH comparator subcohort was defined by weighting the MAMMOTH triple/quad- and penta- refractory cohort proportions to the DREAMM-2 triple/quad- and penta- refractory proportions

Toxicities were common with both CAR T-cell therapies and belantamab. For CAR T-cell therapies, Grade 2+ cytokine release syndrome (usually requiring hospitalization) occurred in 51% of patients who received ide-cel and 44% of patients who received cilta-cel. In addition, 6% of patients who received cilta-cel died of treatment-related complications. For belantamab, 18-46% experienced meaningful to moderate decline in vision lasting 22-33 days.

Table ES2. ICER Evidence Ratings for Anti-BCMA Therapies

Treatment	Comparator	Evidence Rating			
Triple- or Q	Triple- or Quad- Refractory MM (3+ prior lines of treatment)				
lde-cel	Usual Care	B+			
Cilta-cel	Usual Care	B+			
lde-cel	Cilta-cel	1			
Triple-, Quad- or Penta- Refractory MM (4+ prior lines of treatment)					
Belantamab	Usual Care	P/I*			

MM: multiple myeloma.*Compared to current treatments, belantamab appears to be comparable to slightly superior. There is a small but nonzero likelihood of slight net harm. Current evidence does not support belantamab being substantially superior to current treatments.

ICER also performed cost-effectiveness modeling and analyses of the new therapies. The base-case findings from our analysis suggest that CAR-T therapies provide clinical benefit in terms of gains in both quality-adjusted and overall survival over current treatment options for triple- or quadrefractory MM patients exposed to three or more lines of therapy. The incremental costeffectiveness ratios for ide-cel versus the triple- or quad-refractory MM comparator market basket were approximately \$319,000 per QALY gained, \$250,000 per LY gained, \$280,000 per evLYG gained, and \$35,000 per additional PFS month gained. Threshold pricing suggests ide-cel would meet the \$100,000 per QALY threshold at a price of around \$200,000 or a >50% discount from the current list price. Cilta-cel would meet this threshold at a price of around \$300,000, but this finding is preliminary and an optimistic estimate given the extremely limited evidence currently available. Base-case findings for belantamab suggest current list pricing is within commonly cited costeffectiveness thresholds when compared to a triple-, quad-, or penta-refractory MM market basket. However, given uncertainties with the PFS-OS relationship and other parameters in the belantamab model, updated data should be generated and incorporated into future modeling analyses. Small changes in any of the key drivers changed belantamab model findings to a significant extent. Key drivers across all model findings included comparator market basket prices, progression-free survival for the active interventions, and utility of PFS (on or off treatment).

Several potential benefits and contextual considerations not fully captured in the economic modeling include the limited treatment options for patients with TCRMM. Since anti-BCMA treatments represent a novel mechanism of action, these treatments may provide efficacy for patients who currently have few alternatives. However, CAR-T therapies are complex and high-cost with significant side effects. Treatments with these characteristics have been underutilized by disadvantaged populations, suggesting that disparities may be worsened.

1. Background

Multiple Myeloma (MM) is a hematologic cancer of plasma cells.³ Uncontrolled proliferation of plasma cells can lead to a variety of clinical presentations, including:

- Bone pain and fractures due to lytic lesions from plasma cell proliferation in the marrow;
- Increased total or monoclonal protein, which can have direct toxic effects on the kidney, resulting in worsening renal function;
- Hypercalcemia;
- Anemia, due in part to plasma cells suppressing other hematopoietic cell lines and kidney disease.

MM is most often diagnosed through a bone marrow biopsy showing ≥10% plasma cells.⁴

MM is a relatively rare cancer, with an annual incidence of approximately 7 in 100,000 Americans. It is estimated that 32,270 new cases of MM were diagnosed in 2020 and 150,000 Americans are currently living with MM.³ It is primarily a disease of older adults, with a median age at diagnosis of 69. African-Americans appear to be at approximately twice the risk of white Americans, while Asian-Americans appear to be at lower risk.³ The rates of MM have been stable without evidence of increasing incidence over six decades.⁵ The direct medical costs of MM are substantial. A recent analysis of commercial and Medicare claims found that average costs exceeded \$250,000 over a 21-month period, and that 60% of these costs were medication-related.⁶

The last 15 years have seen a proliferation of new, approved therapies for MM, resulting in substantial improvements in survival.¹ In 2000, data from the Survey, Epidemiology, and End Results Program (SEER) suggested that 36% of MM patients achieved 5-year survival while in 2017, SEER models indicated that 56% of patients with MM will survive 5 years.³

Unfortunately, currently-approved therapies are not curative for most patients with MM. While modern combination treatments and autologous stem cell transplant can often lead to effective control with decreased signs and symptoms of MM, over time, most patients will relapse, showing signs and symptoms of renewed, active disease. Patients whose disease does not respond to treatment, or initially respond but are no longer responding to line of treatment are considered refractory. These patients with relapsed or refractory multiple myeloma (RRMM) often cycle through different combinations of agents, which may increase both their clinical and economic burden. Patients with MM whose disease has progressed through three common classes of anti-myeloma medications (monoclonal antibodies such as daratumumab or isatuximab; immunomodulatory drugs or IMiD's such as thalidomide, lenalidomide or pomalidomide; and proteasome inhibitors or PI's such as bortezomib, carfilzomib or ixazomib) are termed "triple class refractory" (TCR) MM.

Currently, there is no widely accepted preferred ordering of lines of therapy for patients with TCRMM. General principles that guide treatment choice include previous treatments, how patient's disease responded to these previous treatments, comorbidities, and risk stratification. One major consideration is incorporating as many new agents as possible (medications to which the patient has not been previously exposed) into each new line of treatment.⁷ This often results in regimens incorporating newer agents in one of the three major classes of anti-myeloma medications (such as pomalidomide or carfilzomib) as well as agents in other classes such elotuzumab or alkylator based treatments. Even with these treatments, patients with TCRMM unfortunately have limited survival, with overall survival <1 year.⁸ These patients with TCRMM represent the population that may potentially benefit from the three medications in this review. For our review, we focused on agents commonly used in the TCR population; thus, some agents, such as Selinexor, were not included as a component of usual care (and comparator to new treatments) due to low rates of use in these patients.

Three new treatments, idecabtagene vicleucel ("ide-cel", Abecma®, Bristol Myers Squibb and bluebird bio), ciltacabtagene autoleucel ("cilta-cel", Janssen and Legend Biotech) and belantamab mafodotin-blmf (Blenrep®, GlaxoSmithKline, referred to as belantamab for the remainder of the report) are proposed as the focus for this review. All three treatments target the B-cell maturation antigen (BCMA), which is overexpressed on plasma cells, but appears to be minimally expressed on other cells. In addition, BCMA appears to be essential for the survival of long-lived plasma cells, making BCMA an attractive therapeutic target.⁹ Blenrep is an antibody-drug conjugate, with a monoclonal antibody specific for BCMA that is linked to a cytotoxic drug. Belantamab is given as an intravenous infusion every 3 weeks. Belantamab received FDA approval in August 2020 for adult patients with relapsed or refractory MM who have received 4 prior lines of therapy including an anti-CD38 monoclonal antibody, a PI and an IMiD.

Ide-cel and cilta-cel are chimeric antigen receptor T (CAR-T) cell therapies, requiring a patient's own T lymphocytes to be obtained via leukapheresis and transduced in the lab with a gene to encode an anti-BCMA antibody. Ide-cel uses a mouse-derived CAR with a single BCMA recognition domain. Cilta-cel uses a camelid CAR with 2 BCMA recognition domains, which theoretically may strengthen the interaction between the CAR and target cells. These genetically modified CAR-T cells are expanded and then infused back into the patient intravenously. Ide-cel received FDA approval in March 2021 for the treatment of adult patients with relapsed or refractory multiple myeloma who previously received four prior lines of treatment, including an IMiD, a PI and an anti-CD38 monoclonal antibody.¹⁰ The biologic license application for cilta-cel was submitted December 21, 2020.

2. Patient and Caregiver Perspectives

ICER engaged with patients with MM (including those treated with anti-BCMA medications), representatives from advocacy organizations, and clinical experts to understand the patient perspective of living with MM. We also spoke with two patient advocacy groups who helped us identify patients who could speak to their experiences. We spoke with five patients over five calls. Finally, we also conducted a focus group with four patients, where we heard in depth about their lived experiences. Additional details, including the semi-structured interview guide and questions, are available in the <u>Supplement.</u> Additional details of the patient perspective from a survey of myeloma patients from the Cancer Support Community is available in the <u>Supplement section B.</u>

Patients spoke about the importance of **quality of life** beyond survival. One patient noted, "I don't want to lie in my bed. I want to meet with friends, go places." In addition, patients mentioned the negative impact of being continually tethered to the health care system. Another patient mentioned, "it's a burden to wake up early, go to the hospital, wait (there's always a delay), then get infused, and not get back home [until after dark]." A second patient summarized, "Visiting the doctor every week or two gets old." Thus, patients reported wanting low side effect treatments that would not require frequent returns to the clinic or hospital.

One of the most frequent side effects that patients mentioned was **fatigue** and **weakness**. Seventy percent of patients in the Cancer Support Community's (CSC) Myeloma registry reported fatigue within the past week. While some spoke of fatigue as a symptom of poorly controlled MM, for others, it was clearly a side effect of treatment. One patient mentioned, "The fatigue is bad—I find it more on POM [pomalidomide]. I have to take a break from cutting veggies." Others noted, "The weakness is the worst thing...it interferes with your ability to exercise and take care of yourself."

Patients also mentioned the impact their disease had on their loved ones and **caregivers**. One patient mentioned, "My wife was greatly impacted. I couldn't do the grocery shopping anymore and I had to sleep in an office chair because of the pain." One patient mentioned that the irritability caused by dexamethasone led to "a temporary estrangement with my spouse because of my short temper." Thus, MM and its treatments have profound effects on families and caregivers as well as the patients.

Several patients reported tremendous **financial strain** due to MM treatments. One patient stated, "My drugs were about \$250,000 a year. That first year I went into debt and had to refinance my home. I was 3 years from paying off my house and I had to start over." Another patient noted, "on top of being filled with cancer, you have to deal with all of these bills."

Clinicians noted, "We still see cases where patients decline to take their drugs because the outof-pocket expenses are so high that they'd have to choose between meds and food/housing." Data from the <u>Cancer Support Community's Multiple Myeloma Specialty Registry</u> indicate that nearly two-thirds of MM patients are concerned about the cost of their cancer care and 42% are often or always upset about the cost of their myeloma care.

Patients and clinicians mentioned that **African American** MM patients face additional barriers to effective treatment. One patient noted, "there's a mistrust of medical community [in the African American community] and is a real thing, and not as much awareness and understanding of MM. [African Americans] are wary in participating in trials and refuse stem cell transplant when its offered." A patient advocacy group mentioned, "navigating newer therapies can be like a maze...African Americans and patients with lower socioeconomic status are more like to get lost in that maze." These comments suggest that newer treatments must proactively engage with historically disadvantaged populations; otherwise, these treatments, because of their complexity, may worsen disparities.

We spoke with 2 patients who had **received CAR T-cell therapies** as part of a clinical trial. Both patients described the infusion and subsequent hospitalization as relatively easy ("a piece of cake") but long and monotonous. One patient described CAR-T therapies as "very liberating", since his doctors did not feel like he needed maintenance medications after CAR-T therapies. Thus, the frequency of doctors' visits and laboratory tests have decreased, and he noted, "if it wasn't for COVID [I could] travel to New York or Italy." A second CAR-T therapy patient had a different experience, since he required continued maintenance medications after CAR-T therapy with belantamab, but did not receive any replies.

We spoke with several patients who were **considering CAR T-cell therapies.** One patient stated, "I'm afraid of CAR-T." When asked what she was afraid of, she talked about how intensive it sounded and that it "only lasts for about a year." A second patient was more interested in CAR-T therapy, stating that he is currently responding to treatment, but that he would consider it for a future line of therapy. Even a patient who had undergone CAR-T therapy stated that he did so because he felt he had no other options. All patients spoke about the importance of having CAR-T therapy as an option saying, "At some point, we're all going to need this because all combinations eventually seem to stop working." CAR-T therapies appear to be primarily coordinated in larger cancer centers; this was a key consideration for patients, since it would require more frequent, longer drives to doctors' appointments.

Our conversations with patients informed our review by reinforcing the importance of specific symptoms including fatigue and weakness. In addition, patient perspectives helped focus our

review on the side effects of both current treatments as well as the side effects of the new interventions. Finally, all these issues reinforced the importance of considering health-related quality of life as a primary outcome for our review.

3. Comparative Clinical Effectiveness

3.1. Methods Overview

Procedures for the systematic review assessing the evidence on ide-cel, cilta-cel, and belantamab for heavily pre-treated relapsed and refractory multiple myeloma are described in the Detailed Methods section of the <u>Report Supplement</u>.

Scope of Review

This review compares the clinical effectiveness of ide-cel and cilta-cel for the treatment of adults with TCRMM (majority with triple- or quad-refractory disease) who have received at least three prior lines of therapy, as well as belantamab for adults with triple-, quad- or penta-refractory multiple myeloma who have received at least four prior lines of therapy in comparison with usual care (i.e., commonly used regimens for those exposed to \geq 3 and \geq 4 prior lines of therapy respectively). The primary patient-important outcomes included OS, PFS, ORR, and health-related quality of life (HRQoL). The full scope of the review is detailed in the Data Sources and Searches section of the <u>Report Supplement</u>.

Evidence Base

The clinical evidence is summarized qualitatively for each intervention separately because the key trials were all single arm studies, so quantitative comparisons were not possible. Details of key studies are described below and summarized in Table 3.1.

Ide-cel

A total of 12 references relating to two single-arm (one Phase I, one Phase II), open label trials of ide-cel met our inclusion criteria. Data from both trials were obtained from publications, conference abstracts, press releases, and information provided by the manufacturers (Table 3.1). In this report, we will report on the Phase II trial (KarMMa), but additional details of both trials are included in the Additional Clinical Evidence section of the <u>Report Supplement</u>.

<u>KarMMa</u>

The KarMMa trial is an ongoing Phase II multi-center, open-label, single-arm trial being conducted at 24 locations worldwide, including North America, Europe, and Japan.^{11,12} The trial screened 158 adults who had previously been exposed to an immunomodulatory drug (IMiD), a proteasome inhibitor (PI), an anti-CD38 antibody, and were refractory to the last prior therapy and enrolled 149 patients.¹³ 140 patients underwent leukapheresis and most (88%) received bridging therapy during the manufacturing process, before lymphodepletion with fludarabine and cyclophosphamide five

days prior to infusion (a total of 128 patients, or 86% of the enrolled population, were analyzed)¹². Retreatment with ide-cel was allowed among those who had a recurrence. The primary outcome was ORR. Secondary outcomes were complete response (CR), safety, duration of response (DoR), PFS, OS, pharmacokinetics, and HRQoL (Table 3.1).

Cilta-cel

A total of 10 references relating to two open-label, single arm trials (one Phase I and one Phase Ib/II) of cilta-cel met our inclusion criteria. At the time of this report, only results from selected sites from the Phase I trial (LEGEND-2) had been published.¹⁴ Data from both trials were therefore obtained from a combination of publications, conference abstracts, and press releases (Table 3.1). In this report, we focus attention on the Phase Ib/II trial (CARTITUDE-1), but additional details of both trials are included in the Additional Clinical Evidence section of the <u>Report Supplement</u>.

CARTITUDE-1

CARTITUDE-1 is an ongoing Phase Ib/II single-arm trial of cilta-cel being conducted in 21 sites in the United States and Japan.¹⁵ The trial enrolled 126 adults with TCRMM who had progressive disease after at least three prior therapies (including a PI, an IMiD, and anti-CD38 antibody), who are double refractory to an IMiD and PI.¹⁶ A total of 113 patients underwent leukapheresis (90% of the enrolled population), 101 underwent lymphodepletion with fludarabine and cyclophosphamide five days prior to infusion (80%), and 97 patients (77% of the enrolled population) were included in the analysis. The primary outcomes were adverse events (AEs) and ORR. Secondary outcomes were OS, PFS, and minimal residual disease (MRD) (Table 3.1).

Belantamab

A total of eight references pertaining to one open-label Phase II clinical trial (DREAMM-2), one conference abstract relating to a pooled post-hoc analysis, and one conference abstract relating to an expanded access program met our inclusion criteria. At the time of this report, two published manuscripts were available for the DREAMM-2 trial,^{17,18} which were supplemented with conference abstracts and information provided by the manufacturer. Additional details are available in the <u>Supplement.</u>

DREAMM-2

DREAMM-2 is an ongoing Phase II, open-label, global, multicenter trial comparing the efficacy and safety of two doses of belantamab (2.5 mg/kg and 3.4 mg/kg) in adults with TCRMM.¹⁷ In total, 196 patients (97 in the 2.5 mg/arm and 99 in the 3.4 mg/kg arm) who had been treated with at least three prior lines of treatment and who were refractory to an IMiD, a PI, and refractory to and/or were not able to tolerate an anti-CD38 monoclonal antibody were enrolled in the trial.¹⁷ At the time of this report, information on the proportion of patients who are classified as penta-refractory was not publicly available (submitted as academic in confidence). In August 2020, the FDA granted accelerated approval to belantamab for the treatment of patients with RRMM with progressive disease after having been treated with four prior lines of therapy. For the purpose of this review, we only present data for the FDA approved dose of 2.5 mg/kg (N=97). The primary outcome assessed was ORR, and secondary outcomes assessed included DoR, time to response, PFS, OS, and safety. All patients who received at least one dose of belantamab were included in the evaluation of efficacy outcomes (ITT population, N=97).

See <u>Supplement</u> for detailed inclusion and exclusion criteria, and definitions of measurable disease and outcomes reported.

Intervention & Trial	Inclusion/Exclusion Criteria	Outcomes	Baseline Characteristics [‡]
lde-cel	<u>Inclusion:</u> — Received at least 2 cycles of ≥3	<u>Primary:</u> ORR	 Age, median (range): 61 (33-78)
<u>KarMMa^{13 Munshi, 2021, 1051}</u> Phase II, open-label single- arm	prior treatment regimens (incl. PI, IMiD, anti-CD38 antibody) and refractory to last regimen	<u>Secondary:</u> CR, OS, PFS, AEs,	 Prior lines of therapy, median (range): 6 (3-16) Triple-refractory: 84% Penta-refractory: 26%
N=149 ⁺	Exclusion: – Previous allogeneic SCT	HRQoL	– EMD: 39% – High-risk cytogenetics: 35%
Cilta-cel <u>CARTITUDE-1^{16,19}</u> Phase II, open-label single- arm	Inclusion: – Received ≥3 prior treatment regimens (incl. PI, IMiD, anti-CD38 antibody) or are double refractory to a PI and IMiD	<u>Primary:</u> AEs, ORR <u>Secondary:</u> OS, PFS, MRD	 Age, median (range): 61 (43-78) Prior lines of therapy, median (range): 6 (3-18) Triple-refractory: 88% Penta-refractory: 42%
N=126 ⁺	Exclusion: – Allogeneic SCT within 6 months or autologous SCT within 4 months		 – EMD: 13% – High-risk cytogenetics: 23%
Belantamab DREAMM-2 ¹⁷ Phase II, open-label, two-	<u>Inclusion</u> : — Received ≥3 previous lines of treatments — Refractory to IMiD and PI, and	Primary: ORR <u>Secondary</u> :	 Age, median (IQR): 65.0 (60.0-70.0) Prior lines of therapy, median (range): 7 (3-21)
arm N=97*	refractory/intolerant to an anti- CD38 therapy <u>Exclusion</u> :	DoR, time to response, PFS, AEs	 Triple-refractory: 100% Penta-refractory: EMD: 23%
	 Received allogeneic SCT Current corneal epithelial disease 		 High-risk cytogenetics: 42%

AEs: Adverse events, CR: Complete response, DoR: Duration of response, EMD: Extramedullary disease, HRQoL: Health-related quality of life, IQR: interquartile range, MRD: Minimal residual disease, N: total number, ORR: Overall response rate, OS: Overall survival, PFS: Progression free survival, SCT: stem cell transplant.

*2.5 mg/kg arm only.

+ Sample sizes are based on the intention-to-treat population

[‡] Baseline characteristics from KarMMa and CARTITUDE-1 are based on the as-treated population

Usual Care

Our systematic literature review did not reveal any prospective studies evaluating the effectiveness of usual care (defined as commonly-used combination regimens described in our research protocol (<u>https://osf.io/3dtr4/</u>) in triple-class refractory patients. The most relevant evidence to support the clinical effectiveness of usual care for triple-class refractory patients to compare with CAR T-cell treatments came from a retrospective observational study (Table 3.2).⁸

The MAMMOTH study was a multi-center US-based retrospective analysis of 275 multiple myeloma patients (data were collected between January 2017 and June 2018), of whom 218 were refractory to at least three lines of therapy (PI, IMiD, and anti-CD38 monoclonal antibody).⁸ 70 (25%) were "penta-refractory" (refractory to two PIs, two IMiDs, and an anti-CD38 monoclonal antibody). Primary outcomes were OS, PFS, and ORR. Two additional retrospective studies were identified and are described in the Additional Clinical Evidence Section of the <u>Report Supplement</u>.

Study	Inclusion/Exclusion Criteria	Outcomes	Patient Characteristics
MAMMOTH ⁸	Inclusion:	Primary:	– Age, median (range): 65 (27-90)
Retrospective chart review	 Refractory to at least 1 PI, 1 IMID, and 1 anti-CD38 	OS, PFS, ORR	 Prior lines of therapy, median (range): 4 (1-16)
N=275 (54% triple/ quad-refractory, 25% penta-refractory)	Exclusion: NR		 Triple- and quad-refractory: 75% Penta-refractory: 25% High-risk cytogenetics: 29%

Table 3.2. Overview of MAMMOTH Study

IMiD: Immunomodulatory drug, N: total number, NR: not reported, PI: Proteasome inhibitor, ORR: Overall response rate, OS: Overall survival, PFS: Progression free survival

Key Differences Across Studies

See the <u>Report Supplement Table D3.2</u> for details on baseline characteristics of the key trials of the interventions. Key differences are summarized below.

Although the study participants in KarMMa and CARTITUDE-1 were of similar ages (median age 61 in both) and had received a similar amount of pre-treatment (median of 6 prior lines of therapy), patients in the CARTITUDE-1 trial were more likely to be penta-refractory than patients in the KarMMa trial (42% vs 26%), but less likely to have extramedullary disease (EMD, the presence of plasma cells outside the bone marrow, a marker of more aggressive disease) (13% vs 39%), and high-risk cytogenetics (23.7% vs 35.2%).^{12,19} In KarMMa, 28 patients (22% of the treated population) were retreated with ide-cel after disease progression. At the time of this report, retreatment was not reported in CARTITUDE-1. In DREAMM-2, the patients undergoing treatment with belantamab were typically older (median age 65), had undergone more pre-treatment (median of seven prior lines of therapy), and were more likely to have high-risk cytogenetics (42.3%);¹⁷ the percentage of penta-refractory patients was provided to us as academic in confidence (

A key difference between the pivotal CAR-T therapy trials and DREAMM-2 was the approach for inclusion in the outcomes analysis. KarMMa and CARTITUDE-1 both only included patients who were infused in the analysis (86% and 77% of enrolled and 91% and 86% of leukapheresed patients, respectively) in an as-treated approach, whereas DREAMM-2 reports on the full intention-to-treat (ITT) population.

The study populations in MAMMOTH were of similar age (median 60 years) to those in KarMMa and CARTITUDE-1 (median 61 years), but appeared to have received less pre-treatment overall (median of four prior lines of therapy versus six or seven).⁸ For this reason, we report the outcomes for triple and quad-penta-refractory patients separately. Furthermore, the exclusion criteria for MAMMOTH were not reported, making it difficult to interpret differences in study populations relative to the key studies of the interventions. See the <u>Report Supplement</u> Table D3.19 for details on baseline characteristics of the studies of usual care.

3.2. Results

Clinical Benefits

The primary outcomes that are used in the economic model are PFS and OS as defined in the clinical trials. The key clinical benefits of ide-cel are described first, followed by cilta-cel and belantamab. Additional outcomes are described in the <u>Report Supplement</u>.

Ide-cel

In the KarMMa trial, with a median follow-up time of 13.3 months (range 0.2-21.2 months) the astreated median PFS across all target CAR-T therapy doses was 8.8 months, and the as-treated median OS was 19.4 months.¹² The as-treated median PFS varied by dose, with the highest dose (450x10⁶ CAR-T cells) achieving the longest median PFS (12.1 months). The as-treated ORR was 73% (94 out of 128 infused patients) and the as-treated stringent complete or complete response rate (sCR or CR) was 33% (42 out of 128) across all doses. The reported outcomes from KarMMa likely represents an optimistic estimate of the results since they are based on patients who received infusion of CAR-T cells, excluding patients who did not receive the therapy due to death prior to infusion, disease progression, or AEs. When calculated on an ITT (that is, including all enrolled patients, including those who were enrolled, leukapheresed but not infused), however, ORR was 63% (94 out of 149 enrolled patients) and sCR or CR was 28% (42 out of 149) (Table 3.3).¹⁶ 33 patients (26% of the treated population) had MRD-negative status.

Importantly, 28 patients (22% of the treated population) were retreated with ide-cel after disease progression, a statistic that was not available until publication of the trial results. Limited data were available on the characteristics of these patients and their associated outcomes; however, PFS in these patients was generally poor, with a median of one month following retreatment.¹²

In the KarMMa trial, HRQoL was assessed using the EQ-5D, EORTC QLQ-C30, and MY20 scales (details on these cancer-specific instruments are available at https://qol.eortc.org/) prior to induction and at day one and nine months post-infusion with ide-cel. Physical functioning, fatigue, pain, and global health sub-scales all improved at nine months compared to baseline (Table 3.4); however data on only 59 of 111 (53%) patients assessed at day one were available at nine months.²⁰ More details on patient-reported outcomes for ide-cel are available in Table D3.10 of the Report Supplement.

Cilta-cel

At the time of this report, as-treated median PFS and OS were not reached with a median of 12.4 months of follow-up (range 1.5-24.9 months) in the CARTITUDE-1 trial.¹⁹ Using an as-treated approach, ORR was 97% (94 out of 97 infused patients) and sCR was 67% (65 out of 97 infused patients had an sCR). However, in an ITT analysis, using the overall enrolled population, ORR was 75% (94 out of 126 enrolled patients) and sCR was 52% (65 out of 126) (Table 3.3).¹⁶ 53 patients (55% of the treated population) had MRD negative status. No data were available on whether any patients receiving cilta-cel required retreatment. More details on outcomes data for cilta-cel are available in Table D3.4 of the <u>Report Supplement.</u>

In CARTITUDE-1 the EORTC QLQ-C30 was administered prior to induction and at various time points post-treatment. At the time of the report, only data on fatigue and pain sub-domains were available. At 184 days, both fatigue and pain scores improved relative to baseline (Table 3.4), however data were available for only 30 out of the 68 (44%) patients who were assessed at baseline.²¹ More details on patient-reported outcomes for cilta-cel are available in Table D3.10 of the <u>Report Supplement</u>.

Belantamab

At the 13-month follow-up (data cut-off date: January 14 2020), patients treated with 2.5 mg/kg belantamab had a median PFS of 2.8 months and median OS of 13.7 months.²² Please refer to the <u>Report Supplement</u> for OS rates at three, six, nine, and 12 months. PFS rates at three to 12 months were submitted as academic in confidence (**Section 2010**). Thirty-one out of 97 participants (32%) achieved an overall response, with five and two patients achieving CR and sCR, respectively.

In DREAMM-2 HRQoL was assessed by means of the EORTC-QLQ-C30, EORTC-QLQ-MY20, and Ocular Surface Disease Index (OSDI) scales. Between weeks 0 to 25, patients' HRQoL was stable across all domains (disease symptoms, pain, fatigue, role functional, physical functioning, global health status, social functioning, and future perspective). For the 25% of patients who responded to treatment and remained on treatment beyond 25 weeks, there was a trend toward improvement in some of these HRQoL domains.²³ However, this trend toward improvement is only applicable to the small minority of patients who remain on therapy beyond 25 weeks.

Additional data on HRQoL data for belantamab can be found in Table D3.10 of the <u>Report</u> <u>Supplement</u>.

Intervention	Trial (N)	Median Follow-Up Duration	As-treated PFS, Median Months (95% CI)	As-treated OS, Median Months (95% CI)	ITT ORR, n (%); [95% Cl]
Ide-cel	KarMMa ^{12,13} (N=149) [‡]	13.3 months [§]	8.8 (5.6, 11.6) [§]	19.4 (18.2, NE) [§]	94 (63.0); [NR]
Cilta-cel	CARTITUDE- 1 ^{16,19} (N=126) [‡]	12.4 months [§]	Not reached at 12.4 months [§]	Not reached at 12.4 months [§]	94 (75.0); [NR]
Belantamab	DREAMM-2 ¹⁷ (N=97 [*])	13 months	2.8 (1.6, 3.6) ‡	13.7 (9.9, not reached) [‡]	31 (32.0); [97.5%Cl: 21.7, 43.6]

 Table 3.3. Key Trial Results of Ide-cel, Cilta-cel, and Belantamab

95%CI: 95% confidence interval, 97.5%CI: 97.5% confidence interval, NE: not estimable, NR: not reported, n: number, ORR: Overall response rate, OS: Overall survival, PFS: progression free survival

*2.5 mg/kg arm only

+ Xi'an site only

‡ Intention-to-treat

§ Median Follow-up duration, PFS and OS for KarMMa and CARTITUDE-1 are based on the as-treated population

Table 3.4. Change from Baseline in Health-Related Quality of Life (Selected EORTC QLQ-C30* Sub-
Domains)

Intervention (Trial)	Time from Baseline (N)	Physical Functioning, Mean (95% Cl)	Fatigue, Mean (95% CI)†	Pain, Mean (95% CI)†	Global Health, Mean (95% CI)
Ide-cel‡	9 months	13.2	-22.8	-23.8	15.4
(KarMMa) ²⁰	(N=59)	(7.9, 17.9)	(-29.1, -17.1)	(-30.2, -18.3)	(9.8, 20.9)
Cilta-cel‡	6 months	NR	-9.2	-8.9	NR
(CARTITUDE-	(N=30)		(-16.4, -2.0)	(-17.6, -0.3)	
1) ²¹	C m a m th a	0.1	2.0	2.0	4 7
Belantamab ‡	6 months	-0.1	3.6	2.6	-4.7
(DREAMM-	(N=19)	(-5.3, 5.3)	(-7.6, 14.6)	(-6.5, 11.4)	(-12.1, 2.8)
2) ^{23,24}					

95%CI: 95% confidence interval, N: total number, NR: not reported, ORR: Overall response rate.

*EORTC QLQ-C30: European Organization for Research and Treatment of Cancer Quality of Life Questionnaire for Cancer Patients

⁺Negative changes indicate a reduction in pain or fatigue

‡Mean changes from baseline have been digitized and should be interpreted with caution

Usual Care

We identified three retrospective observational studies to inform our comparison of usual care to the interventions.^{8,25,26} In this report, we present outcomes from the MAMMOTH study because outcomes are available across various lines of therapy as well as on an overall basis.⁸ The majority of patients in the MAMMOTH study were refractory to daratumumab (93%), lenalidomide (77%), bortezomib (68%), and pomalidomide (65%) and a minority were refractory to carfilzomib (47%), ixazomib (12%), and thalidomide (8%). Median PFS was 3.4 months in the overall population but was not separately reported in the triple-quad and penta-refractory populations. Median OS was 9.3 months in the overall population, 9.2 months in the triple-quad population, and 5.6 months in the penta-refractory population. Overall response was 29-31% (Table 3.5). Additional details on the outcomes from MAMMOTH and the additional retrospective studies are provided in Tables D3.15 and D3.16 in the Report Supplement.

Study	Population (N)	PFS, Median Months, (95% CI)	OS, Median Months (95% CI)	ORR, n (%)
MAMMOTH ⁸	Overall (N=275)	3.4 ⁺ (2.8-4.0)	9.3 ⁺ (8.1, 10.6)	85 (31.0) [*]
	Triple- and quad-	NR	9.2 (7.1, 11.2)	80 (29.0) [*]
	refractory (N=148)			
	Penta-refractory (N=70)	NR	5.6 (3.5, 7.8)	21 (30.0)*

95%CI: 95% confidence interval, N: total number, n: number, ORR: Overall response rate, OS: overall survival, PFS: progression-free survival.

* ORR calculated based on first line of therapy.

 \dagger PFS and OS on next line after T₀.

Harms

Harms of ide-cel, cilta-cel, and belantamab are presented based on the number of study participants who were actually infused/treated ("safety population").

Ide-cel

In the KarMMa trial, cytokine release syndrome (CRS) was the most commonly-reported AE, reported by 84% of patients (48% Grade 1; 31% Grade 2) and lasting a median of 5 days.¹² Of 107 patients who had CRS with or without neurotoxicity, 19 (17.8%) required intensive care unit admission.²⁷ CRS was most likely to be managed with tocilizumab (52%), followed by corticosteroids (15%). Risk of CRS appeared to be dose-related, reported by 96% of patients in the 450x10⁶ dose compared to 76% in the 300x10⁶ dose. Patients in the 450x10⁶ dose were also more likely to require tocilizumab to manage their CRS than patients in the 300x10⁶ dose (67% vs 43%). Older patients were also more likely to experience CRS; all 20 patients ≥70 reported CRS across all doses. Other important AEs included neurotoxicity (18%), thrombocytopenia (63%), and neutropenia (91%) (Table 3.6). Forty-four (34%) patients died during the study, with 27 (21%) due to progressive disease. Three deaths (2%) were treatment-related AEs within 8 weeks of infusion (CRS, pneumonia, gastrointestinal hemorrhage). An additional treatment-related death from pneumonia was reported within 6 months of infusion.¹² More information on harms of ide-cel are available in Table D3.11 of the <u>Report Supplement</u>.

Cilta-cel

In CARTITUDE-1, 95% of patients reported CRS, with most 95% experiencing low to moderate CRS (51% Grade 1 and 39% Grade 2).¹⁹ Median time to onset of CRS was seven days (range 1-12) and lasted a median of four days (range 1-97). CRS was most likely to be managed with tocilizumab (69%), followed by corticosteroids (22%) and anakinra (19%). Other important AEs included neurotoxicity (21%), thrombocytopenia (79%), and neutropenia (96%) (Table 3.6). A total of 14 deaths (14.4%) were reported during the study, five due to progressive disease, three due to AEs unrelated to treatment (pneumonia and other cancers), and six due to AEs related to treatment (sepsis, CRS, lung abscess, respiratory failure, neurotoxicity).¹⁹ More information on harms of ciltacel are available in Table D3.12 of the <u>Report Supplement.</u>

Belantamab

In DREAMM-2, AEs were reported by 97.9% of patients treated with belantamab 2.5 mg/g (N=95).²² The vast majority of AEs were considered to be related to the study treatment (88.4%). Three patients (3.2%) died during the study due to AEs (myocardial infarction (n=2), sepsis (n=1)), with one death being treatment related (sepsis).²² At 6.3 months of follow-up, over three-quarters of the enrolled participants had discontinued study treatment, mainly due to disease progression or death (60.8% and 32.6%, respectively).¹⁷ Nine patients (10%) discontinued study treatment due to AEs (one due to keratopathy and one due to blurred vision).²² AEs frequently led to dosing modifications, with over half (54%) experiencing dose delays and over a third (35%) requiring dose reductions. Keratopathy, defined as changes to the corneal epithelium, was reported by 72% of patients; however at 13 months follow-up, 77% had recovered from their first, and 48% from their last corneal event.^{17,18} Forty-six percent of patients experienced a moderate decline in vision (Grade 2 toxicity, Best Corrected Visual Acuity), which resolved in most patients at the end of study follow-up. A decline in vision in their better-seeing eye to 20/50 or worse was reported by 18% of patients. The majority, 82%, recovered (improved to 20/40 or better) by the end of the follow-up period. No patients reported permanent vision loss. More information on harms of belantamab are available in Table D3.13 of the Report Supplement.

Intervention	Trial (N)	Treatment- related SAEs	Important AEs	D/C due to AEs	Overall Mortality at Median Follow-Up Time
lde-cel	KarMMa ¹² \ (N=128)	3.1%	- CRS: 84% - NT: 18% - Thrombocytopenia: 63%	NR	34% at 13.3 months (range 0.2- 21.2)
Cilta-cel	CARTITUDE- 1 ¹⁹ (N=97)	NR	- CRS: 95% - NT: 21% - Thrombocytopenia: 79%	NR	14% at 12.4 months (range 1.5- 24.9)
Belantamab	DREAMM- 2 ^{17,22} (N=95) [†]	11.6%	 CRS: 0% NT: 0% Thrombocytopenia: 38% Moderate decline in vision in more affected eye (BVCA Grade 2+): 46% Meaningful decline in vision (worse than 20/50 in better eye): 18% 	10%	33% at 6.3 months (IQR 3.7-7.7)

Table 3.6. Key Harms of Ide-cel, Cilta-cel, and Belantamab

AE: adverse event, CRS: cytokine release syndrome, D/C: discontinuation, N: total number, NT: neurotoxicity, SAE: serious adverse event, BCVA: Best Corrected Visual Acuity, IQR: interquartile range

* Xi'an site only.

+ safety population (2.5 mg/kg).

Usual Care

The retrospective observational studies we selected to represent the effectiveness of usual care did not provide sufficient or consistent information on the harms of the treatment regimens. Therefore, we selected representative prospective trials of commonly used treatments for TCRMM (Elo-Pom-Dex: elotuzumab-pomalidomide-dexamethasone, Car-Cy-Dex: carfilzomibcyclophosphamide-dexamethasone, Ixa-Len-Dex: ixazomib-lenalidomide-dexamethasone).²⁸⁻³⁰ Serious AEs were common, reported by roughly half of the participants. The most commonly reported grade 3 or 4 AEs included neutropenia, anemia, infection, and thrombocytopenia. Discontinuation rates due to AEs varied from 14 to 18% (Table 3.7). Differences in harms between these regimens and that of the interventions should be interpreted with caution, however, as the trials were generally conducted in less heavily pre-treated populations.

Treatment	Trial (N)	Serious AEs	Important Grade 3/4 AEs	Deaths (all)	Discontinuation Due to AEs
Elo-Pom-Dex	ELOQUENT-3 ³⁰	53%	– Neutropenia: 13%	22%	18%
	(N=60)		– Anemia: 10%		
			– Infection: 13%		
Car-Cy-Dex	Bringhen 2014 ²⁹	NR	– Neutropenia: 20%	13%	14%
	(N=56)		– Anemia: 11%		
			– Infection: 5%		
Ixa-Len-Dex	TOURMALINE-	46.5%	– Neutropenia: 22%	4%	17%
	MM1 ²⁸		– Anemia: 9%		
	(N=361)		– Infection: <1%		

Table 3.7. Harms of Selected Commonly-Used Usual Care Regimens

AE: adverse events, Car: carfilzomib, Cy: cyclophosphamide, Dex: dexamethasone, Elo: elotuzumab, Ixa: Ixazomib, Len: lenalidomide, n: number, N: total number, Pom: pomalidomide.

Subgroup Analyses and Heterogeneity

Data on efficacy outcomes by subgroups of patients (such as by age, high risk cytogenetics, race/ethnicity, etc.) were not consistently reported. In the KarMMa trial, as-treated median PFS for ide-cel was 8.6 months for patients 65 years or older (n=45) and 10.2 months for 70 or older (n=20), compared to 8.8 months for the overall population.³¹ As-treated median PFS for the 50 patients in the KarMMa trial who had EMD was 7.9 months, and for the 45 patients with high cytogenetic risk was 8.2 months.³² At the time of this report, efficacy data by subgroups was not available for the CARTITUDE-1 trial.

In DREAMM-2, ITT median PFS for belantamab was 2.9 months for patients who had previously been unsuccessfully treated with three to six therapies and 2.2 months for those who had received seven lines of treatment or more.³³ ITT median PFS for patients with high cytogenetic risk factors was 2.1 months.³⁴ For those with mild to moderate renal impairment, ITT median PFS was 2.2 and 3.7 months, respectively.³⁵ At 6.3 months of follow up, 43.6% of patients aged 65 to < 75 years achieved an overall response, while only one patient (7.7%) in the age group 75 and above achieved an overall response. 31.6% of White, and 37.5% of Black patients achieved an overall response at 6.3 months of follow-up. ORR at 13 months of follow-up was submitted to ICER as academic in confidence (**Context**). Similar to other studies, substantially fewer Black patients were enrolled in this study compared to White patients (16 and 72 patients, respectively).¹⁷

Uncertainty and Controversies

Several important uncertainties remain in our evaluation of CAR T-cell therapies. First, 9% of the ide-cel patients who were leukapheresed did not receive treatment (14% for cilta-cel) and were not included in the published ORR and PFS estimates. Since manufacturing failures (i.e., inability to successfully encode the patients' T-cells) are now rare, most patients who were enrolled but not

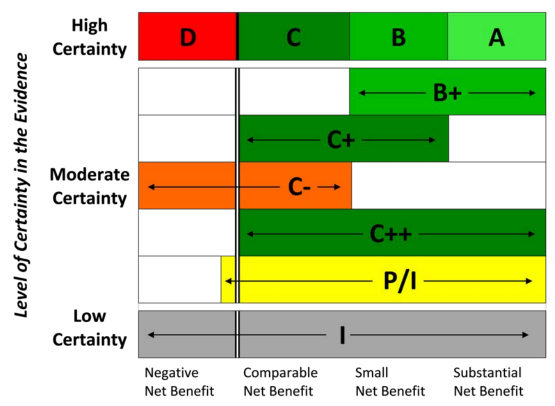
able to receive treatment likely had more severe or more aggressive disease. Thus, it is likely that accounting for these patients would diminish the benefits seen with ide-cel and cilta-cel; future studies should publish an ITT analysis incorporating these patients. Our calculated ITT analysis decreased the ORR from 73% to 63% for KarMMa and from 97% to 75% for CARTITUDE-1. Second, additional follow-up data are needed for ide-cel and cilta-cel to quantify the median PFS and OS; outcomes data are particularly limited for cilta-cel, as no comprehensive presentations or peer-reviewed publications are available, and the manufacturer did not respond to requests for additional data. Longer-term data may also allow us to definitively determine whether a small minority of patients are able to achieve a long-term, durable response, and to understand whether retreatment with CAR-T therapy might be necessary in others. Third, additional information is needed regarding the specific nature and potential causes of cilta-cel treatment-related deaths (6% in CARTITUDE-1). Fourth, while there is interest in utilizing CAR T-cell therapies earlier in the MM disease course, studies are needed to determine whether these therapies are superior to current therapies for first or second relapse of MM.

Uncertainties also remain in our evaluation of belantamab. First, additional studies should examine the median OS with belantamab. Across 22 RCTs in relapsed and refractory patients MM, a metaanalysis found that the ratio between PFS and OS was approximately 3.1³⁶; previous meta-analyses have generated comparable results.³⁷ However, in the pivotal DREAMM-2 study, the 2.5mg/kg arm had a median PFS of 2.8 months and a median OS of 13.7 months, for a 4.9 ratio. While it is possible that the OS/PFS ratio would differ substantially for belantamab compared to all other treatments for MM, further studies are needed to confirm or refute the current OS/PFS ratio seen for belantamab. Second, while belantamab was approved as a single-agent treatment, other MM treatments are most effective as doublet or triplet therapies. We await the results of ongoing studies combining belantamab with other treatments to determine whether belantamab would be helpful as a component of novel combination therapies. Lastly, more research is needed to determine a treatment approach best suited for the management of keratopathy, and ultimately to reduce the burden of ocular toxicities on patients, improve patient outcomes and reduce the need for dose adjustments or treatment discontinuation.¹⁸

3.3. Summary and Comment

An explanation of the ICER Evidence Rating Matrix (Figure 3.1) is provided in the Supplement.





Comparative Clinical Effectiveness

Comparative Net Health Benefit

A = "Superior" - High certainty of a substantial (moderate-large) net health benefit

B = "Incremental" - High certainty of a small net health benefit

C = "Comparable"- High certainty of a comparable net health benefit

D= "Negative"- High certainty of an inferior net health benefit

B+= "Incremental or Better" – Moderate certainty of a small or substantial net health benefit, with high certainty of at least a small net health benefit

C+ = "Comparable or Incremental" - Moderate certainty of a comparable or small net health benefit, with high certainty of at least a comparable net health benefit

C- = "Comparable or Inferior" – Moderate certainty that the net health benefit is either comparable or inferior with high certainty of at best a comparable net health benefit

C++ = "Comparable or Better" - Moderate certainty of a comparable, small, or substantial net health

benefit, with high certainty of at least a comparable net health benefit

P/I = "Promising but Inconclusive" - Moderate certainty of a small or substantial net health benefit, small likelihood of a negative net health benefit

I = "Insufficient" - Any situation in which the level of certainty in the evidence is low

Our systematic review of the evidence suggests that CAR T-cell therapies for patients with triple-or quad- refractory MM likely provides small to substantial net health benefits over current usual care. Benefits included longer survival as well as improved quality of life. Counterbalancing these benefits were the harms, including CRS, which is temporary but often requires hospitalization and intensive care unit level care.

Our systematic review of cilta-cel and ide-cel suggests that the evidence is insufficient to determine whether one agent is superior to the other. There are no studies comparing these agents directly, nor sufficient data to perform quantitative indirect comparisons. We conclude that belantamab is promising but inconclusive compared to usual care for patients with triple-, quad- and penta-refractory MM exposed to 4+ prior lines of treatment. The ORR and OS suggests a possible small net benefit. However, the frequency and severity of visual impairment and lack of demonstrated improvement in HRQoL suggests that any net benefits are likely to be modest. The current evidence precludes a substantial benefit; additional data is required to preclude small overall net harm.

Table 3.8. Evidence Ratings

Treatment	Comparator	Evidence Rating		
Triple- or quad- refractory MM (3+ prior lines of treatment)				
lde-cel	Usual Care	B+		
Cilta-cel	Usual Care	B+		
lde-cel	Cilta-cel	1		
Triple-, quad- or penta- refractory MM (4+ prior lines of treatment)				
Belantamab	Usual Care	P/I		

MM: multiple myeloma

4. Long-Term Cost-Effectiveness

4.1. Methods Overview

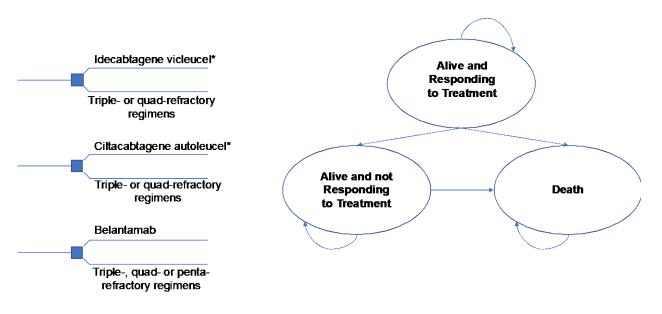
The primary aim of this section of the review was to assess the lifetime cost-effectiveness of ide-cel, cilta-cel, and belantamab as compared to relevant comparator treatments. We developed a *de novo* decision analytic model for this evaluation, informed by key clinical trials and prior relevant economic models.^{38,39}

Specific to CAR-T therapies, an initial decision tree was used to calculate the costs and consequences from treatment initiation (i.e., leukapheresis) to T-cell infusion. We note this differs somewhat from our reporting of "intent to treat" results based on all enrolled trial patients, but better fits the purpose of the model, which is to reflect costs and outcomes following the initiation of the CAR-T therapy process. The decision tree included patients who were eligible for CAR-T therapy and who had undergone leukapheresis. After initiating leukapheresis, patients could continue to receive the infusion of the engineered T-cells; discontinue (before infusion but after leukapheresis) because of disease progression, adverse events, or manufacturing failures; or die before receiving the infusion. Those who discontinued prior to T-cell infusion received the costs, benefits, and risks of the market basket of triple- or quad-refractory comparators.

The cohort of patients were assigned to three mutually exclusive and exhaustive health states in a partitioned survival model (Figure 4.1). Health states included 1) alive and progression free or responding to therapy, 2) alive and not responding to therapy/subsequent relapse, and 3) dead from multiple myeloma-related complications or other causes. We accounted for on/off therapy through application of differential utilities within a health state. At the end of each cycle, patients in the alive and progression free or responding to therapy health states did not transition treatments. Those in the alive and not responding to therapy/subsequent relapse health state transitioned to a progressed state that included a market basket of subsequent therapies. Patients remained in the model until they died. Health state occupancy was derived using partitioned survival techniques that included the direct extrapolation of progression-free survival (PFS) and overall survival (OS) Kaplan-Meier curves. A detailed description of curve digitization is available in <u>Supplement Section E2</u>.

Model outcomes included total life years (LYs) gained, quality-adjusted life years (QALYs) gained, equal-value life years gained (evLYG), time progression free/responding to treatment, and total costs for each intervention over a lifetime time horizon discounted at 3% per annum. <u>Supplement Table E.3.1-E.3.3</u> present undiscounted results.

Figure 4.1. Model Structure



*Includes up-front decision tree to account for patient disposition from leukapheresis and through CAR-T infusion.

Target Population

The model focused on an intention-to-treat analysis, with a hypothetical cohort of heavily pretreated patients with MM beginning at age 60. The CAR-T trials' enrollment criteria required patients to have been treated with at least 3 previous lines of therapy. However, enrolled patients had received a median of 6 previous lines of therapy and were 84% – 88% TCRMM. For belantamab, 100% were at least triple class-refractory and enrolled patients had received a median of 7 previous lines of therapy. Cohort characteristics for each treatment group are described in <u>Supplemental Table E.1.2 and E.1.3</u>.

Treatment Strategies

Interventions

The list of interventions was developed with input from patient organizations, clinicians, manufacturers, and payers on which treatments to include. The full list of interventions is as follows:

- Idecabtagene vicleucel (Abecma[®], Bristol Myers Squibb, Bluebird bio, Inc.)
- Ciltacabtagene autoleucel (Janssen, Legend Biotech)
- Belantamab mafodotin-blmf (Blenrep[®], GlaxoSmithKline)

Comparator Treatments

Given the numerous available therapies used by clinicians at various lines of therapy, a market basket approach was used to compare to each intervention based on level of pretreatment using the MAMMOTH study and a recent conference proceeding that estimated the distribution of treatments by line of therapy.^{8,26} The market basket composition was approximated by both broad-therapy and specific-therapy estimations. PFS and OS curves were either directly informed by the MAMMOTH study or derived as described in the model structure section.⁸ Supplement Tables E.2.3 and E.2.4 provide dosing, administration schedules, and unit costs for each market basket of comparators.

4.2. Key Model Assumptions and Inputs

Our model includes several key assumptions described in Table 4.1.

Assumption	Rationale
For CAR-T therapies, patients received at least one full single course of therapy. A proportion of patients were re-treated and assigned the full costs associated with CAR-T, including infusion and other resource utilization.	A proportion of patients in the KarMMa trial were re- treated and we assume the cost of re-treatment would be incurred for the infusion and other resources during and after infusion. In the absence of data from CARTITUDE-1 on retreatment, we assumed the same percentage across CAR-Ts and tested this in a scenario analysis.
Subsequent treatments received after progression are uniform within each population/line of therapy.	Patients progressing but still alive are assumed to receive subsequent therapy consistent with the line of therapy by population.
Parametric curve functions were fit separately for each population/treatment and used to extrapolate the data over a lifetime horizon.	Given different populations and lines of treatment, an indirect treatment comparison with the same baseline comparator across populations/treatments was not feasible.
Recent observational evidence on patients using a mix of therapies was used to estimate PFS and OS of relevant comparators.	There is wide variation in therapies used by line of therapy; we used a population with a mix of the most recently used therapies to reflect survival under conditions of current practice in RRMM.
In cases with immature survival data, calibration methods were used to adjust the relationship between PFS and OS based on prior evidence in multiple myeloma.	In some cases, OS data were immature and to calculate health outcomes, we used calibration methods to adjust relationships between PFS and OS.
Patients who discontinue CAR-T therapy due to an AE, or manufacturing failure before receiving the T-cell infusion received comparator treatment benefits, risks, and costs; those who died were accounted for prior to CAR-T infusion.	Patients with MM often receive some level of therapy or intervention until death and therefore patients that discontinued received a market basket of subsequent therapies consistent between each arm of the model.

Table 4.1. Key Model Assumptions

The model included only grade 3/4 adverse events and specific toxicities as well as all grades of cytokine release syndrome.	Less severe adverse events are not expected to significantly impact patient health outcomes or costs, although there is evidence to suggest an impact of cytokine release syndrome and keratopathy on
	outcomes and cost across all grades.

95%CI: 95% confidence interval, CAR-T: chimeric antigen receptor T-cell, MM: multiple myeloma, OS: overall survival, PFS: progression-free survival

Model inputs were estimated from the clinical review, published literature, and information from expert stakeholders. Model inputs included PFS, OS, occurrence of adverse events, quality-of-life utility values, and health care costs. We note that data on PFS are extremely limited for cilta-cel; we used reported data for PFS from CARTITUDE-1 and used prior estimates of the relationship between PFS and OS to estimate the median OS. Probabilities, costs, and other inputs differed between treatments to reflect varying effectiveness between interventions. Health state utility values were consistent across interventions within the same disease, although different utilities were applied for patients in the progression-free state depending on whether they were on or off therapy. Key model inputs are described in Tables 4.2 and 4.3 for interventions in the triple- or quad-refractory population and for interventions in the triple-, quad-, or penta-refractory population, respectively.

Table 4.2. Key Model Inputs for Population with Triple- or Quad-Refractory MM (3+ prior lines of treatment)

Parameter	Ide-Cel	Cilta-Cel	CAR-T Comparator Market Basket
Progression-Free Survival, Median	8.8 Months	Not reached; 12 month % PFS used	3.4 Months
Overall Survival, Median	19.4 Months	NR	9.2 Months
Progression-Free on Therapy and Responding Utility	0.78		
Progression-Free Off Therapy and Responding Utility	0.82	0.82	N/A
Progressed Disease/Not Responding to Therapy Utility	0.71	·	·
Proportion of patients infused (CAR-T specific)	91%	86%	N/A
Proportion of patients re-treated (CAR-T specific)	22% of those initially infused	22% of those initially infused	N/A
Treatment Acquisition Price*	\$419,500	\$419,500 (assumption based on ide-cel list price)	\$24,829 per cycle applied until progression
Administration, Monitoring, and Adverse Event Management Costs Applied in First Model Cycle [†]	\$18,500 (CRS grade 1) - \$121,500 (CRS grade 4); other AE costs and monitoring included in decision tree	\$18,500 (CRS grade 1) - \$121,500 (CRS grade 4); other AE costs and monitoring included in decision tree	\$4,661
Other Management-Related Costs per Model Cycle	\$540		
Key Sources (see inputs section and supplement for all sources)	Delforge et al 2020 ⁴⁰ ;Hari et al, 2020 ²⁷ ;Munshi et al, 2021 ¹²	Delforge et al 2020 ⁴¹ ;Hari et al, 2020 ²⁷ ;Madduri et al, 2020 ⁴²	Gandhi et al, 2019 ⁴¹ ;Gandhi et al, 2019 ⁸ ;Mehra et al, 2020 ²⁶

CAR-T: chimeric antigen receptor T-cells, N/A: not applicable, NR: not reported.

*Comparator market basket price assumes 15% discount for oral therapies based on Federal Supply Schedule pricing

⁺ For ide-cel approximately 80% had a grade 1-4 CRS event, whereas for cilta-cel approximately 95% had a grade 1-4 CRS event

Table 4.3. Key Model Inputs for Population with Triple-, Quad-, or Penta- Refractory MM (4+ prior
lines of treatment)

Parameter	Belantamab	Belantamab Comparator Market Basket	
Progression-Free Survival, Median	2.8 Months	2.6 Months	
Overall Survival, Median	13.7 Months	7.7 Months	
Progression-Free on Therapy and Responding Utility		0.78	
Progressed Disease/Not Responding to Therapy Utility	ng to Therapy Utility 0.71		
WAC Price per Treatment Cycle [*]	\$8,277 per vial	\$20,434	
Administration and Monitoring Costs per Model Cycle	\$355	\$1,301	
Adverse Event Management Costs for First Two Model	\$2,565	\$1,259	
Cycles			
Other Management-Related Costs per Model Cycle	\$540		
Key Sources (see inputs section and supplement for all	Delforge et	Gandhi et al, 2019 ⁴¹ ;Gandhi et	
sources)	al,2020 ⁴¹ ;Lonial et al, 2020 ⁴³	al, 2019 ⁸ ;Mehra et al, 2020 ²⁶	

*Belantamab dosing based on weight distribution from trial with proportion of patients receiving 2 or 3 vials (proportion redacted); comparator market basket price assumes 15% discount for oral therapies based on Federal Supply Schedule pricing

Clinical Inputs

Base-case survival was derived from parametric fits to each intervention's available PFS and OS Kaplan-Meier curves.^{15,43,44} Tables E.2.5 and E.2.6 delineate the evidence that was used to calculate transition probabilities. The model included any grade 3/4 adverse event that occurred in at least 5% of patients for any of the treatments and comparators. Given the potentially significant impact of cytokine release syndrome on health care resource utilization and quality of life, we included all grades 1-4 for these adverse events and adjusted costs and quality of life estimates accordingly. The costs and disutility of adverse events were applied to the first two model cycles for each intervention and comparator. After cycle 2 of the model, we applied a dose adjustment factor, assuming adverse events would be resolved with lower dosing of each therapy. <u>Supplement Table E.2.8</u> lists the adverse events considered. Health state utilities were applied for each model health state to adjust for quality-of-life changes over time. Utilities were derived from publicly available sources.⁴¹ Tables 4.2 and 4.3 show health utility values by line of therapy. <u>Supplement Table E.2.9</u> describes the adverse event disutilities.

Economic Inputs

All costs used in the model were updated to 2020 US dollars using methods following the ICER reference case. The unit cost for each treatment is reported in <u>Supplement Table E.2.10</u>. The regimens used for each comparator treatment can be found in Supplement Table E.2.3 and E.2.4. We used the list price for ide-cel and, in the absence of any available market projection or other data, assumed the same price for cilta-cel. The wholesale acquisition cost for belantamab was used.⁴⁵ Comparator therapy pricing was based on WAC pricing with 15% discounts on oral

therapies based on the Federal Supply Schedule. Costs associated with additional health care utilization that occurred from administration and monitoring, and post-treatment were included in the model. <u>Supplement Table E.2.12</u> details the health care utilization unit costs used in the model and the evidence sources. AE costs were derived from reasonable treatment assumptions used in previous analyses mentioned as evidence sources in <u>Supplement Table E.2.14</u>.

4.3. Results

Base-Case Results

The total discounted costs, life years (LYs), quality-adjusted life years (QALYs), and equal value of life years gained (evLYG) over the lifetime time horizon are detailed in Tables 4.4, 4.5, and 4.6. In the triple- or quad- refractory cohort of patients treated with three or more lines of therapy, ide-cel had a total discounted cost of \$646,000 with discounted LYs, QALYs, and evLYG gained of 2.97, 2.24, and 2.40, respectively. Cilta-cel had a total discounted cost of \$617,000 with discounted LYs, QALYs, and evLYGs gained of 4.52, 3.40, and 3.74, respectively. The CAR-T therapy comparator market basket cohort had a total discounted cost of \$276,000 with discounted LYs, QALYs, and evLYGs gained of 1.50, 1.08, and 1.08, respectively. In the triple-, quad-, or penta-refractory cohort of patients treated with four or more lines of therapy, the belantamab arm had a total discounted cost of approximately \$254,000 with discounted LYs, QALYs, and evLYGs gained of 1.60, 1.15, 1.19, respectively. The belantamab comparator market basket had a total discounted cost of \$218,000 with discounted LYs, QALYs, and evLYGs gained of 1.08, 0.78, and 0.79, respectively. We note that cost differences between belantamab and its comparator were mitigated by dose reduction and/or discontinuation due to adverse events for belantamab.

Table 4.4. Results for the Base-Case for Ide-cel Compared to Triple- or Quad- Refractory MM
Comparator Market Basket (3+ prior lines of treatment)

Treatment	Intervention Cost	Other non- intervention costs*	Total Cost	Time Spent in PFS State (months)	QALYs	Life Years	evLYGs
Ide-Cel	\$466,000	\$180,000	\$646,000	16.24	2.24	2.97	2.40
CAR-T	\$153,000	\$123,000	\$276,000	5.75	1.08	1.50	1.08
Comparator							
Market							
Basket							

CAR-T: chimeric antigen receptor T-cells, evLYG: equal-value of life years gained, QALYs: quality-adjusted life years gained, PFS: progression-free survival.

*Other non-intervention costs include costs for monitoring, progressed treatment costs, physician visits, adverse event management (first two cycles only) and monthly laboratory costs for complete blood count and liver testing

Table 4.5. Preliminary Base-Case Results for Cilta-cel Compared to Triple- or Quad- RefractoryMM Comparator Market Basket (3+ prior lines of treatment)

Treatment	Intervention Cost	Other non- intervention costs*	Total Cost	Time spent in PFS State (months)	QALYs	Life Years	evLYGs
Cilta-Cel [†] (preliminar y)	\$445,000	\$172,000	\$617,000	25.82	3.40	4.52	3.74
CAR-T Comparator Market Basket	\$153,000	\$123,000	\$276,000	5.75	1.08	1.50	1.08

CAR-T: chimeric antigen receptor T-cells, evLYG: equal-value of life years gained, QALYs: quality-adjusted life years gained, PFS: progression-free survival.

*Other non-intervention costs include costs for monitoring, progressed treatment costs, physician visits, adverse event management (first two cycles only) and monthly laboratory costs for complete blood count and liver testing [†]Using placeholder price for cilta-cel

Table 4.6. Results for the Base-Case for Belantamab Compared to Triple-, Quad-, or Penta-Refractory MM Comparator Market Basket (4+ prior lines of treatment)

Treatment	Intervention Cost	Other non- intervention costs*	Total Cost	Time Spent in PFS State (months)	QALYs	Life Years	evLYGs
Belantamab	\$152,000	\$102,000	\$254,000		1.15	1.60	1.19
Belantamab	\$118,000	\$99,000	\$218,000		0.78	1.08	0.79
Comparator							
Market							
Basket							

evLYG: equal-value of life years gained, QALYs: quality-adjusted life years gained, PFS: progression-free survival. *Other non-intervention costs include costs for monitoring, progressed treatment costs, physician visits, adverse event management (first two cycles only) and monthly laboratory costs for complete blood count and liver testing

Table 4.7 presents the incremental results from the base-case analysis, which include incremental cost-effectiveness ratios for the incremental cost per LY gained, incremental cost per QALY gained, and incremental cost per evLYG gained. In the triple- or quad- refractory cohort of patients treated with three or more lines of therapy, total costs for ide-cel were approximately \$370,000 greater than total costs for the comparator arm; gains in LYs, QALYs, and evLYGs were 1.47, 1.16, and 1.32 in relation to the comparator arm. This resulted in an incremental cost-effectiveness ratio of approximately \$319,000 per QALY gained, \$250,000 per LY gained, \$280,000 per evLYG gained, and \$35,000 per additional PFS month gained for ide-cel versus the comparator arm. In the triple- or quad- refractory cohort of patients treated with three or more lines of therapy, total costs for ciltacel were approximately \$341,000 greater than total costs for the comparator arm; gains in LYs, QALYs, and evLYGs were 3.02, 2.32, and 2.66, respectively. This resulted in an incremental cost-effectiveness ratio of approximately \$147,000 per QALY gained, \$113,000 per LY gained, \$128,000

per evLYG gained, and \$17,000 per additional PFS month gained for cilta-cel versus the comparator arm. However, we note results for cilta-cel are based on very limited clinical and economic evidence and should be considered preliminary for the purposes of this analysis. Among other key inputs missing, there was no publicly available overall survival curve to inform survival extrapolations and the price of cilta-cel is a placeholder based on the price of ide-cel. In the triple-, quad-, or penta-refractory cohort treated with four or more lines of therapy, total costs for the belantamab arm were approximately \$36,000 greater than total costs for the comparator arm; gains in LYs, QALYs, and evLYGs were more than 0.52, 0.37, and 0.40 than that of the comparator arm. This resulted in an incremental cost-effectiveness ratio of approximately \$98,000 per QALY gained, \$70,000 per LY gained, \$93,000 per evLYG gained, and \$18,000 per additional PFS month gained for belantamab versus the comparator arm. As mentioned above, cost differences are relatively low for this comparison due to high rates of discontinuation in the belantamab arm.

Treatment	Comparator	Cost per QALY Gained	Cost per Life Year Gained	Cost per evLYG gained	Cost per additional PFS month gained
Ide-Cel	CAR-T Comparator Market Basket	\$319,000	\$250,000	\$280,000	\$35,000
Cilta-cel (preliminary)*	CAR-T Comparator Market Basket	\$147,000	\$113,000	\$128,000	\$17,000
Belantamab	Belantamab Comparator Market Basket	\$98,000	\$70,000	\$93,000	\$18,000

evLYG: equal-value of life years gained, QALY: quality-adjusted life year, LY: life years.

*Using placeholder price for cilta-cel

Threshold Analyses

Tables 4.8 and 4.9 present the unit price needed for each therapy to reach commonly cited costeffectiveness thresholds. The price needed to achieve these thresholds would be inclusive of both the manufacturer price and any potential hospital mark-up that may be applied. As above, we note that threshold prices differ substantially between the CAR-T therapies in part because of a paucity of available data on PFS and OS for cilta-cel.

Table 4.8. QALY-Based Threshold Analysis Results

	WAC per Unit	Net Price per Unit	Unit Price to Achieve \$50,000 per QALY	Unit Price to Achieve \$100,000 per QALY	Unit Price to Achieve \$150,000 per QALY	Unit Price to Achieve \$200,000 per QALY
Ide-Cel	\$419,500	N/A	\$140,000	\$192,000	\$245,000	\$295,000
Cilta-cel (preliminary)*	N/A	N/A	\$208,000	\$317,000	\$427,000	\$537,000
Belantamab	\$8,277		\$7,300	\$8,300	\$9,300	\$10,400

N/A: not available, evLYG: equal-value life years gained, QALY: quality-adjusted life years gained, WAC: wholesale acquisition cost.

*Using placeholder price for cilta-cel

Table 4.9. evLYG-Based Threshold Analysis Results

	WAC per Unit	Net Price per Unit	Unit Price to Achieve \$50,000 per evLYG	Unit Price to Achieve \$100,000 per evLYG	Unit Price to Achieve \$150,000 per evLYG	Unit Price to Achieve \$200,000 per evLYG
Ide-Cel	\$419,500	N/A	\$146,000	\$206,000	\$265,000	\$324,000
Cilta-cel (preliminary)*	N/A	N/A	\$224,000	\$350,000	\$475,000	\$600,000
Belantamab mafodotin	\$8,277		\$7,300	\$8,400	\$9 <i>,</i> 500	\$10,600

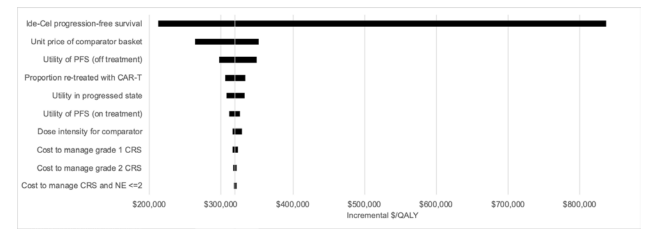
N/A: not available, evLYG: equal-value life years gained, QALY: quality-adjusted life years gained, WAC: wholesale acquisition cost.

*Using placeholder price for cilta-cel

Sensitivity Analyses

To demonstrate effects of uncertainty on both costs and health outcomes, we varied input parameters using available measures of parameter uncertainty (i.e., standard errors or plausible parameter ranges). Figure 4.2 presents an example tornado diagram resulting from the one-way sensitivity analysis for ide-cel versus the triple- or quad-refractory comparator market basket. When varying PFS, we assumed the same proportional relationship in terms of gains in OS. At lower PFS (6 months) the ICER was well above commonly cited cost-effectiveness thresholds while the upper PFS (12 months) generated an ICER of approximately \$212,500. Key drivers across all model findings included the unit price of the comparator market basket of therapies, progression-free survival for the active interventions, and utility of PFS (on or off treatment). The belantamab model in particular was extraordinarily sensitive. Small changes in any of the key drivers listed above changed model findings to a significant extent. Please see <u>Supplement Section E4 for</u> additional results from the one-way sensitivity analyses, including tornado diagrams for cilta-cel and belantamab .





With noted uncertainty outside of that modeled, a probabilistic sensitivity analysis was conducted to assess variation across all model inputs with quantified uncertainty simultaneously and to vary the results over 5,000 iterations. Tables 4.10 and 4.11 present the probability of reaching certain willingness-to-pay thresholds for ide-cel. A total of 3% of the iterations for ide-cel versus the comparator were beneath a threshold of \$150,000 per QALY gained. Similarly, 6% of the iterations for ide-cel versus the comparator were beneath a threshold of \$150,000 per evLYG gained. Sensitivity analyses for cilta-cel and belantamab are available in <u>Supplement Section E4</u>.

Table 4.10 Probabilistic Sensitivity Analysis Cost per QALY Gained Results: Ide-Cel versus Triple- or
Quad- Refractory MM Comparator Market Basket (3+ prior lines of treatment)

	Cost Effective at	Cost Effective at	Cost Effective at	Cost Effective at
	\$50,000 per	\$100,000 per	\$150,000 per	\$200,000 per
	QALY	QALY	QALY	QALY
Ide-cel	<1%	<1%	3%	15%

Table 4.11. Probabilistic Sensitivity Analysis Cost per evLYG Gained Results: Ide-Cel versus Tripleor Quad- Refractory MM Comparator Market Basket (3+ prior lines of treatment)

	Cost Effective at	Cost Effective at	Cost Effective at	Cost Effective at
	\$50,000 per	\$100,000 per	\$150,000 per	\$200,000 per
	evLYG	evLYG	evLYG	evLYG
Ide-cel	<1%	<1%	6%	24%

Scenario Analyses

We ran three main scenario analyses: 1) a scenario that assumes no retreatment costs for each CAR-T product; 2) a modified societal perspective (results presented in <u>supplement section E5</u>); and 3) a scenario analysis that adjusts the proportional relationship between PFS and OS for belantamab to be within a similar range to that suggested by a recent synthesis of the evidence (results presented in <u>supplement section E5</u>).

The base-case model assumed 22% of patients were re-treated with CAR-T as reported in KarMMa.¹² In this scenario analysis, we assume no additional retreatment charge for each CAR-T product. We include all other costs related to re-treatment for the 22% re-treated, including adverse events (i.e., hospitalizations) and monitoring. This scenario suggests re-treatment as a driver of the model results with, reductions in the incremental cost-effectiveness ratio of approximately -25% (Table 4.12).

Treatment	Comparator	Cost per QALY Gained	Cost per Life Year Gained	Cost per evLYG	Cost per Month of PFS Gained
lde-cel	CAR-T	\$247,000	\$194,000	\$217,000	\$27,000
	Comparator				
	Market Basket				
Cilta-cel	CAR-T	\$110,000	\$85,000	\$96,000	\$13,000
(preliminary)*	Comparator				
	Market Basket				

evLYG: equal-value of life years gained, QALY: quality-adjusted life year, LY: life years

*Using placeholder price for cilta-cel

Model Validation

We used several approaches to validate the model. First, we attempted multiple survival extrapolation techniques and compared estimates to findings from each intervention's most recent published paper or abstract to ensure outcomes were consistent with clinical evidence. Second, we presented preliminary results to manufacturers and clinical experts, and based on feedback from these groups, we refined data inputs or extrapolations as needed. During the model transparency process, manufacturers noted we underestimated survival for ide-cel and the CAR-T comparator market basket (which also impacted a percentage of the comparator to belantamab). These changes were reflected in the final model calculations. Third, we varied model input parameters to evaluate face validity of changes in results. Finally, we compared model results to other cost-effectiveness findings in this therapy area.

Uncertainty and Controversies

There were important uncertainties relevant to generating model outcomes. Given that evidence was abstracted from single-arm studies, there were challenges when selecting the most appropriate comparator. In order to calculate incremental costs, risks, and benefits, we compared each therapy to a contemporaneous population of RRMM patients within the MAMMOTH observational study, i.e., a triple-/quad-refractory cohort as a comparator to CAR-T therapies and a weighted average cohort of triple-, quad-, and penta-refractory treated patients as a comparator to belantamab. Advantages of using the MAMMOTH population as a comparator include generalizability to the academic medical center setting and use of currently-recommended regimens; these settings are likely to be the same ones that will provide CAR-T and belantamab therapy. This allowed us to include not only survival evidence but also the mix of therapies used to estimate a monthly cost for each comparator. However, given that the treatment landscape changes dramatically over short time periods in RRMM, and the lack of a quantitative indirect treatment comparison to inform these analyses, caution should be used when interpreting cost-effectiveness estimates. A further complication in identifying a relevant comparator to belantamab was the mix of patients exposed to

three or more lines of therapy and four or more lines of therapy, respectively. To address this limitation, we assumed a population mix from MAMMOTH to estimate weighted average outcomes and costs. Moreover, comparisons across the interventions of interest were not feasible given differences in populations from each single-arm study.

We acknowledge the challenge of interpreting incremental cost-effectiveness ratios for recently approved or investigational therapies when they are compared to existing high-cost comparators. Model outcomes were sensitive to the price of comparators as well as future health care costs for survivors. In sensitivity analyses, we varied the price of the market basket by the minimum and maximum estimated combination regimen in the market basket (which varied from approximately \$17,000 to \$37,000 per cycle of therapy). Therefore, interpretation of the cost-effectiveness of each therapy should include review of the one-way sensitivity analyses. Not surprisingly, we generally found that lower comparator prices led to less favorable cost-effectiveness estimates for the interventions.

The relationship between PFS and OS for each therapy was fairly consistent with prior meta-analysis evidence that suggests for every one month in PFS, patients gain approximately 2.5-3 months of OS.^{36,37} However, we observed a different pattern in the DREAMM-2 belantamab trial. The single arm study suggested that relationship was closer to a 5-month gain in OS for every 1-month gain in PFS. The resulting model was quite sensitive; small changes in this relationship as well as other key parameters resulted in relatively large swings in our cost-effectiveness estimates. To address this limitation and the uncertainty around these estimates, we adjusted the relationship between PFS and OS for the belantamab arm to be consistent with the most recent meta-analysis evidence in a scenario analysis.³⁶ In this scenario, belantamab was dominated (i.e., more costly, less effectiveness findings should be updated.

After our draft report was published, new data emerged in a recent publication of the results of the KarMMa trial, suggesting that >20% of ide-cel patients received a second CAR-T infusion. While these patients were treated with T cells that had already been harvested and engineered, there is uncertainty around whether there would be a second "charge" for retreatment; it is certainly the case that hospitals delivering a second infusion would incur costs of infusion, monitoring, and management of side effects. We assumed that a second charge would be levied in our base-case analyses (for both ide-cel and cilta-cel) and removed this second charge this in a scenario analysis. Our findings suggested that this would have a significant impact on the price that could be charged to achieve a \$100,000 per QALY threshold (e.g., \$191,000 vs. \$233,000 for ide-cel with and without a second charge, respectively). In addition, one of the CAR-T recipients we spoke with indicated that he was receiving high-priced maintenance medication after his infusion even though he remained in response to CAR-T, suggesting that other treatment decisions following an initial CAR-T infusion may have a significant impact on the estimated cost-effectiveness of these therapies.

Specific to cilta-cel, interpretation of the cost-effectiveness findings should be noted as a very preliminary and somewhat optimistic scenario. The evidence used in the model relies on limited clinical study evidence with a PFS estimate that has yet to reach its median and no reported estimate for OS. The only study with longer follow-up data available to us was the Phase 1 LEGEND-2 study, which focused on a younger population with fewer previous lines of treatment. Therefore, extrapolations of survival are likely overestimates of the benefit of cilta-cel. We found through sensitivity analyses that at lower PFS and OS levels, the incremental cost-effectiveness ratios became less favorable. In addition, while dosing of cilta-cel ranged in the CARTITUDE-1 study, there have been no reports of whether retreatment was necessary. In the absence of any reported data, a study publication, or input from the manufacturer, we assumed the same rate of retreatment as observed in the ide-cel KarMMa study. Finally, we found no indication of the likely price of cilta-cel and relied instead on the ide-cel price as a placeholder.

Survival curve fitting relies on assumptions that may differ substantially between parametric models (see Figures E.2.1 to E.2.4 for modeled survival extrapolations). Scenario analyses around alternative extrapolations found large variation in incremental cost-effectiveness ratios. There are further limitations to piecewise modeling approaches, such as seemingly arbitrary cut-point intervals and modeled "jumps" in the hazard that may appear clinically unjustifiable.⁴⁶ We ensured our assumptions did not lead to invalid and unrealistic survival estimates, for example the tail of the extrapolated PFS curve crossing the tail of the OS curve. We relied on reported estimates of percentage alive and in PFS and OS states in addition to fit statistics. Survival estimates were sensitive to base-case findings as shown in the one-way sensitivity analyses.

4.4. Summary and Comment

The base-case findings from our analysis suggest that CAR-T therapies provide clinical benefit in terms of gains in QALYs and LYs over current treatment options for patients exposed to three or more lines of therapy. However, the benefits of ide-cel and cilta-cel should be reviewed separately given that evidence is still emerging. Threshold pricing suggests ide-cel would meet the \$100,000 per QALY threshold at a price of around \$200,000, which represents a discount of >50% from the list price of \$419,500. Cilta-cel would meet this threshold at a price of around \$300,000, but as noted above this is likely an optimistic estimate given the limited evidence currently available. Base-case findings for belantamab suggest current list pricing is within commonly cited cost-effectiveness thresholds. However, given uncertainties with the PFS-OS relationship and other parameters in the belantamab model, updated data should be generated and incorporated into future modeling analyses. Model findings across all interventions were sensitive to the cost of comparators, PFS and OS estimates, utility of PFS (on and off treatment), and overall health care costs for multiple myeloma patients.

5. Potential Other Benefits and Contextual Considerations

Our reviews seek to provide information on potential other benefits offered by the intervention to the individual patient, caregivers, the delivery system, other patients, or the public that was not available in the evidence base nor could be adequately estimated within the cost-effectiveness model. These elements are listed in the table below, with related information gathered from patients and other stakeholders. Following the public deliberation on this report the appraisal committee will vote on the degree to which each of these factors should affect overall judgments of long-term value for money of the intervention(s) in this review.

Contextual Considerations	Relevant Information
Acuity of need for treatment of individual	The acuity of need for treatment is high. Patients
patients based on the severity of the	with heavily pre-treated MM have relatively short
condition being treated	life expectancy without treatment, and their
	treatment options are currently limited.
Magnitude of the <i>lifetime</i> impact on	MM has a moderate lifetime impact on individual
individual patients of the condition being	patients. The disease has a tremendous effect on
treated	the quality and quantity of life for affected patients.
	However, the median age at diagnosis is 69; thus,
	most patients diagnosed with MM have lived many
	decades without myeloma. Thus, unlike diseases
	such as cystic fibrosis which has a large effect on a
	patient's entire lifespan, MM has a large effect on a
	proportion of a patient's lifespan, leading to our
	assessment of moderate lifetime impact.
New mechanism of action may provide	Anti-BCMA activity of both CAR-T therapies and
benefits for patients who are	belantamab suggests that these treatments may be
unresponsive to current therapies	efficacious for patients who are unresponsive to
	other treatments.

Table 5.1. Categories of Contextual Considerations

Table 5.2. Categories of Potential Other Benefits

Patients' ability to achieve major life	For CAR-T therapy, suppressing the symptoms of MM
goals related to education, work, or	appears to support patients' ability to achieve life
family life	goals. For belantamab, substantial side effects of
	treatment balance the decrease in disease effects,
	making it less clear whether belantamab supports
	patients' ability to achieve life goals.
Caregivers' quality of life and/or ability to	For CAR-T therapy, suppressing the symptoms of MM
achieve major life goals related to	appears to support caregivers' ability to achieve life
education, work, or family life	goals. For belantamab, substantial side effects of
	treatment balance the decrease in disease effects,
	making it less clear whether belantamab supports
	caregivers' ability to achieve life goals.
Patients' ability to manage and sustain	For CAR-T therapy, patient burden may be
treatment given the complexity of	substantially less since much of the monitoring is
regimen	done immediately after infusion and many patients
	appear to need no maintenance therapy. For
	belantamab, patient burden may be less since the
	studies focused on monotherapy. However, studies
	are underway focusing on combining belantamab
	with other treatments, which may negate patient
	burden advantages with belantamab.
Society's goal of reducing health	Anti-BCMA therapies have the potential to worsen
inequities	existing disparities. Therapies with high cost or high
	side effect burden (such as the current anti-BCMA
	therapies), as well as those requiring treatment at
	specialized centers, are often utilized at lower rates
	among historically disadvantaged populations.

6. Health Benefit Price Benchmarks

The ICER health benefit price benchmark (HBPB) is a price range suggesting the highest price a manufacturer should charge for a treatment, based on the amount of improvement in overall health patients receive from that treatment, when a higher price would cause disproportionately greater losses in health among other patients due to rising overall costs of health care and health insurance. In short, it is the top price range at which a health system can reward innovation and better health for patients without doing more harm than good. Unit prices for ide-cel, cilta-cel, and belantamab that would achieve incremental cost-effectiveness ratios of \$100,000 and \$150,000 per QALY or evLYG are presented in Table 6.1. We arrive at a range of HBPB of approximately \$192,000 - \$265,000 for ide-cel, \$317,000 - \$475,000 for cilta-cel, and \$8,300 - \$9,500 for belantamab.

	WAC per unit	Unit Price at \$100,000 Threshold	Unit Price at \$150,000 Threshold	Discount from WAC to Reach Threshold Prices
Ide-cel				
QALYs Gained	\$419,500	\$192,000	\$245,000	42%-54%
evLYG	\$419,500	\$206,000	\$265,000	37%-51%
Cilta-cel (preliminary)				
QALYs Gained	\$419,500	\$317,000	\$427,000	No discount –
	(placeholder)			24%
evLYG	\$419,500	\$350,000	\$475,000	No discount –
	(placeholder)			17%
Belantamab				
QALYs Gained	\$8,277 per vial	\$8,300	\$9,300	No discount
evLYG	\$8,277 per vial	\$8,400	\$9,500	No discount

QALY: quality-adjusted life year, WAC: wholesale acquisition cost

*WAC as of March 2021

7. Potential Budget Impact

7.1. Overview of Key Assumptions

Using results from the cost-effectiveness model, we estimated the potential budgetary impact of each B-cell maturation antigen CAR-T cell and antibody drug conjugate therapy for refractory multiple myeloma. We used the price from the base-case analysis and three threshold prices (at \$50,000, \$100,000, and \$150,000 per QALY) for each intervention. Potential budget impact is defined as the total differential cost of using each new therapy rather than the relevant existing therapy for the treated population, calculated as differential health care costs (including intervention costs) minus any offsets in these costs from averted health care events. All costs were undiscounted and estimated over a five-year time horizon.

The analysis included the estimated number of individuals in the US who would be eligible for each treatment. To estimate the size of the potential candidate populations for each intervention, we used the total number of adults 18 years and older with at a minimum triple-class refractory multiple myeloma. It is estimated that 140,000 Americans are living with MM.³ Patients with MM are treated with CD38-targeting antibodies which are generally well tolerated and result in a response in approximately 30% of patients with MM. Thus, we assumed that 70% of patients with MM are refractory to CD38-targeted antibodies. Further, among MM patient's refractory to CD38targeting antibodies, 54% are triple and quad-refractory and 25% are at least penta-refractory.⁸ Therefore, we estimated approximately 98,000 MM patients are refractory to CD38-targeting antibodies, with approximately 52,900 classified as triple or quad-refractory and 24,500 classified as at least penta-refractory in the US. We assumed that 20% of these patients would initiate treatment in each of the five years, or approximately 10,580 patients eligible for CAR-T therapy and 4,900 eligible for belantamab each year. Because the CAR-T therapies will be launched (if cilta-cel is approved) within a short period of each other, the eligible population of approximately 10,580 triple or quad-refractory patients per year was split in half between the two interventions (approximately 5,290 per year per CAR-T therapy).

The aim of the potential budgetary impact analysis was to document the percentage of patients who could be treated at select prices within 5 years without crossing a potential budget impact threshold that is aligned with overall growth in the US economy. For reports begun in 2019-2020, the five-year annualized ICER potential budget impact threshold that should trigger policy actions to manage access and affordability is approximately \$819 million per year for new drugs. More detail on these methods can be found in the <u>Section F of the Report Supplement</u>.

7.2. Results

Belantamab

Figure 7.1 depicts the cumulative per-patient potential budget impact calculations for belantamab as compared to the market basket comparator, based on the wholesale acquisition cost. The average potential budgetary impact for belantamab was approximately \$5,610 per patient in year one, with cumulative net cost increasing in years two and three and beginning to plateau at year four, reaching approximately \$39,210 per patient in year five.

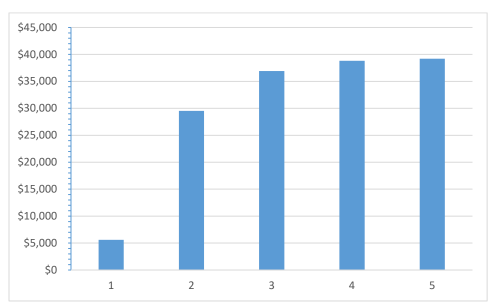


Figure 7.1. Cumulative Net Cost per Patient Treated with Belantamab at Wholesale Acquisition Cost

Assuming the wholesale acquisition cost (unit price of approximately \$8,280), all eligible patients could be treated within five years (assuming 20% uptake each year), reaching 18% of the ICER budget impact threshold of \$819 million per year over five years. Similarly using the prices to reach \$150,000, \$100,000, and \$50,000 per QALY (unit prices of approximately \$9,340, \$8,310, and \$7,290, respectively), all eligible patients could be treated within five years (assuming 20% uptake each year) without crossing the ICER budget impact threshold.

Idecabtagene vicleucel

Figure 7.2 depicts the cumulative per-patient potential budget impact calculations for ide-cel as compared to the market basket comparator, assuming the list price of \$419,500. The average potential budgetary impact was approximately \$354,340 per patient in year one, which remained relatively constant through year five where it reached approximately \$364,120 per patient.

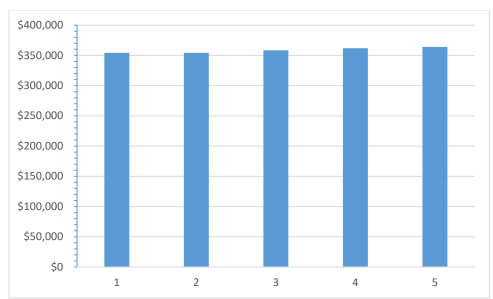


Figure 7.2. Cumulative Net Cost per Patient Treated with Ide-cel at List Price

Assuming the list price (\$419,500), only 43% of the eligible patients could be treated within five years (assuming 20% uptake each year), before crossing the ICER budget impact threshold of \$819 million per year. Similarly using the price to reach \$150,000 per QALY (price of approximately \$245,000), 95% of the eligible patients could be treated within five years (assuming 20% uptake each year) before crossing the ICER budget impact threshold. All eligible patients could be treated within five years without crossing the ICER budget impact threshold at the price to reach either \$100,000/QALY and \$50,000/QALY. Figure 7.3 depicts the potential budgetary impact of idecabtagene vicleucel.

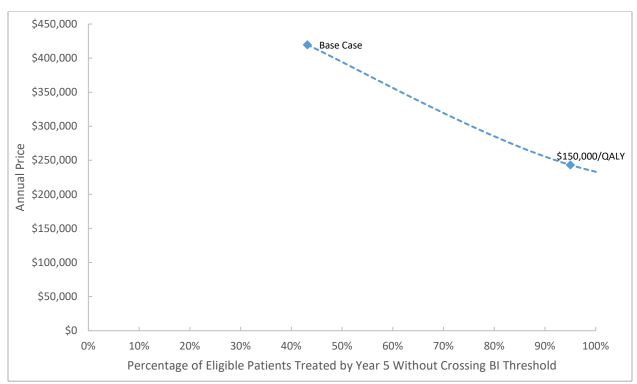


Figure 7.3. Potential Budgetary Impact of Ide-Cel in Triple- or Quad-Refractory Multiple Myeloma

Ciltacabtagene autoleucel

Figure 7.4 depicts the preliminary cumulative per-patient potential budget impact calculations for cilta-cel as compared to the market basket comparator, assuming the placeholder price of \$419,500. The average potential budgetary impact was approximately \$289,210 per patient in year one, with a small increase each year to approximately \$330,340 per patient in year five.

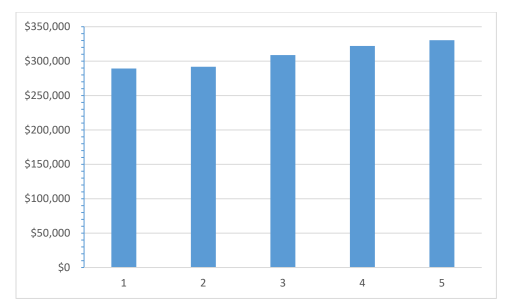
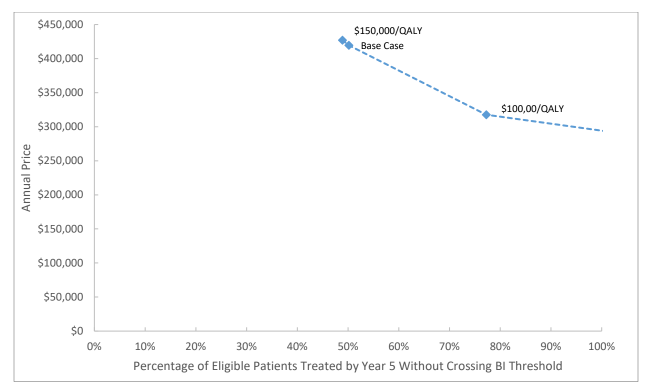


Figure 7.4. Cumulative Net Cost per Patient Treated with Cilta-cel at Placeholder Price (Preliminary)

Assuming the placeholder price of \$419,500, only 50% of the eligible population could be treated within five years (assuming 20% uptake each year) before reaching the ICER budget impact threshold of \$819 million per year. At the price to achieve a threshold of \$100,000/QALY (approximately \$317,000), only 77% of the eligible population could be treated within five years (assuming 20% uptake each year) before reaching the ICER budget impact threshold of \$819 million per year. All eligible patients could be treated at the price to reach a threshold of \$50,000 (approximately \$208,000) without reaching the ICER budget impact threshold. Figure 7.5 depicts the potential budgetary impact of ciltacabtagene autoleucel.





Additional net costs per year are presented along with cumulative net costs in <u>Section F of the</u> <u>supplement</u> for each of the three treatments.

Supplemental Materials

A. Background: Supplemental Information

A1. Definitions

Disease Definitions:

<u>Triple-Class Refractory Multiple Myeloma</u>: Multiple myeloma that has become refractory to the three common classes of myeloma medications: immunomodulators (I.e., lenalidomide), proteasome inhibitors (I.e., bortezomib) and a monoclonal antibody (I.e., daratumumab).

<u>Quad Refractory Multiple Myeloma:</u> Multiple myeloma that has become refractory to 4 commonly used myeloma medications. Most commonly, quad-refractory disease is refractory to: 1 anti-CD38 monoclonal antibody (most often daratumumab), 1 or 2 immunomodulators (most often lenalidomide +/- pomalidomide) and 1 or 2 proteasome inhibitors (most often bortezomib +/carfilzomib).

<u>Penta Refractory Multiple Myeloma</u>: Multiple myeloma that has become refractory to 5 commonly used myeloma medications. Most commonly, penta-refractory disease is refractory to 2 immunomodulators (most often lenalidomide and pomalidomide), 2 proteasome inhibitors (most often bortezomib and carfilzomib) and an anti CD38 monoclonal antibody (most often daratumumab).

<u>Extramedullary disease</u>: Multiple myeloma in which plasma cells form tumors outside of the bone marrow. Extramedullary disease is a sign of more aggressive myeloma and portends a worse prognosis.

<u>High-risk Cytogenetics</u>: A chromosomal abnormality which has been shown to increase the risk of more aggressive disease.

Intervention Definitions:

<u>CAR T-cell therapy:</u> Chimeric antigen receptors (CARs) are artificial fusion proteins constructed to recognize specific antigens. CAR T-cells are T-cell lymphocytes that have been genetically modified to express these CAR's, so that these T-cells can identify and to marshal an immune response against cancer cells that produce these antigens. The focus of this review, ide-cel and cilta-cel utilize CAR T-cells that recognize the B-cell Maturation Antigen which appears to be expressed on most malignant plasma cells. CAR T-cell therapy starts with 1) harvesting of the patient's lymphocytes with leukapheresis. 2) Lymphocytes are then modified in the laboratory to express the CAR protein. 3) These modified lymphocytes are expanded to sufficient numbers and 4) the modified, expanded lymphocytes are reinfused back into the patient.

<u>Ide-cel</u>: Idecabtagene vicleucel is a chimeric antigen receptor (CAR) T-cell therapy based on the first anti-BCMA CAR developed at the National Cancer Institute (11D5-3-CD828Z), using a mouse origin anti-BCMA moiety. CARs are artificial fusion proteins that combine a BCMA-recognition domain with a costimulatory domain⁴⁷. When reinfused into the patient, the genetically modified lymphocytes with the CAR proteins triggers a multi-pronged immune response, resulting in the destruction of cancer cells.

<u>Cilta-cel</u>: Ciltacabtagene autoleucel is a chimeric antigen receptor (CAR) T-cell therapy based on the camelid heavy chain only anti-BCMA CAR. The camelid heavy chain (LCAR-B38M) incorporates 2 BCMA recognition domains, which theoretically should increase the specificity for BCMA. Otherwise, cilta-cel has a similar mechanism of action and treatment logistics to ide-cel.⁴⁷

<u>Belantamab mafodotin</u>: a first-in-class, antibody-drug immunoconjugate consisting of an anti-BCMA monoclonal antibody and an anti-cancer drug. Belantamab mafodotin (referred to as belantamab for the remainder of the supplement) binds to BCMA-antigens and kills multiple myeloma cells via a multimodal mechanism. Belantamab induces cell apoptosis in addition to antibody-dependent cellular cytotoxicity (ADCC).^{48,49}

Outcome definitions: Studies rely on the International Myeloma Working Group (IMWG) Uniform Response Criteria definitions for outcomes.⁵⁰

<u>Complete Response (CR)</u>: Negative immunofixation on serum and urine AND disappearance of any soft tissue plasmacytomas AND <5% plasma cells in the bone marrow.

<u>Stringent Complete Response (sCR)</u>: Meets CR criteria AND normal free light chain ratio AND absence of clonal cells in bone marrow by immunohistochemistry or immunofluorescence.

Partial Response (PR):

 \geq 50% reduction of serum M-protein AND reduction in 24-hour urinary M-protein by \geq 90% or to < 200 mg/24 h

If the serum and urine M-protein are unmeasurable, $a \ge 50\%$ decrease in the difference between involved and uninvolved free light chain levels is required in place of the M-protein criteria.

If serum and urine M-protein are not measurable and serum free light chain assay is also not measurable, \geq 50% reduction in plasma cells is required in place of M-protein, provided baseline bone marrow plasma cell percentage was \geq 30%.

In addition to the above criteria, if present at baseline, $a \ge 50\%$ reduction in the size of soft tissue plasmacytomas is also required.

<u>Very Good Partial Response (vgPR)</u>: Serum and urine M-protein detectable by immunofixation but not on electrophoresis or \ge 90% reduction in serum M-protein plus urine M-protein level < 100 mg/24 h

<u>Overall Response Rate (ORR)</u>: Proportion of patients treated who had a partial response to treatment or better (PR + vgPR + CR + sCR)

Minimal Residual Disease (MRD): Small number of cancer cells that can remain after treatment. MRD status predicts relapse. In MM, MRD is assessed in patients with CR, via sensitive techniques such as next-generation flow cytometry or next-generation sequencing.⁵¹

B. Patient Perspectives: Supplemental Information

B1. Methods

ICER conducted a wide-ranging effort to engage patients and advocacy groups to develop an understanding of perspectives of patients with MM. We drew on experience from the prior ICER review of MM therapies in 2016, reaching out to patient advocacy groups and individual patients engaged in the previous review. Specifically, we had a series of conversations with the Cancer Support Community, who were able to provide us with invaluable insights into the experience of MM patients within their community. They also helped us engage with individual patients who were able and willing to speak with us. We also engaged with Patients for Affordable Drugs, who also helped us identify patients we could speak to about their experience. We purposefully tried to engage African American patients to elicit their experiences, since they are overrepresented in the MM population and may face disparities in diagnosis and access to care. We contacted the International Myeloma Foundation and the Black Women's Health Imperative for further guidance on reaching out to specific myeloma patients and advice on any relevant data resources to assess racial disparities in myeloma care. We connected with the Association of Community Cancer Centers and spoke with an expert who commented on the differences in access to new myeloma treatments between large academic medical centers and community cancer centers.

We had a series of conversations with individual patients with MM, as well as one focus group discussion. Conversations were informed by a semi-structured interview guide which focused the conversation on several themes, including:

- 1. What is your experience with different treatments that you have tried?
 - What has worked, what has not?
 - o Side-effects
 - Impact on daily life, family, work
- 2. What are the financial aspects of the treatments you have tried?
 - Any issues with insurance, paying for the treatment
- 3. Where have you received care (in what type of hospital), what doctors have you seen?
- 4. What are you hoping to get from any new treatments that become available?

- 5. What do you think are some key issues about patients' experience with multiple myeloma that are not being captured in major clinical studies or trials, including:
 - Symptoms and complications of disease
 - o Impact of disease on function and quality of life
 - Side effects of treatment
 - Effects on caregivers and family members
 - Any other issues
- 6. For patients who have experience with CAR-T therapies:
 - How well did it work for you?
 - What were the side effects?
 - What was the impact like on daily life, family, work?
 - How was it like finding these treatments, how available or accessible do you think these are for patients?
 - Compared to other treatments you have received for multiple myeloma, what was your experience like? Specifically, how did it feel to be "off" treatment after you received the CAR-T infusion?
 - Have you required any follow-up after CAR-T infusion? If so, is this routine follow-up or for complications of treatment?

We had an iterative process, where emerging themes were incorporated into subsequent conversations to determine whether these themes were universally felt by all (or most) patients or were idiosyncratic to a single (or few) patient(s). Furthermore, patients submitted individual feedback and shared their experiences via the Patient Input Questionnaire on ICER's website.

After each of these conversations, patient comments were transcribed, collated, organized, and summarized. We drew upon the themes that emerged from our conversations and our summaries for the patient perspective sections of this report.

B2. Cancer Support Community Myeloma Registry Survey

Cancer Support Community (CSC) is an international nonprofit organization that provides support, education and hope to people impacted by cancer. CSC has conducted surveys of over 14,000 cancer patients across a wide variety of cancers, including multiple myeloma. We believe that these responses provide additional insight into the patient experience with myeloma and complement the qualitative responses provided in the report. We are indebted to the CSC for allowing us access to the following data.

Table B1 highlights that side effects may be even more common than reported, as 25% of respondents reported not mentioning side effects because they didn't believe anything could be done.

Table B1. CSC Multiple Myeloma Specialty Registry Findings: Physical Symptoms and Side Effects

Physical Symptoms and Side Effects Concerns	Findings
Respondents who did not report side effects because they didn't believe anything could be done about their side effects	25%
Comfort level with speaking to their doctor about side effects and symptoms	>99% positive
	Always: 5%
Side effects impacted patients' decisions about treatments	Often: 9%
	Sometimes: 28%
How well respondents felt their health care team prepared them to manage side	Very much: 33%
effects	Quite a bit: 26%
	Somewhat: 22%

Table B2 highlights the high prevalence of fatigue in multiple myeloma. Over half of survey respondents experience fatigue often or always and for over one-third of respondents, fatigue interfered with their daily lives "quite a bit" or "very much".

Table B2. CSC Multiple Myeloma Specialty Registry Findings: Fatigue

Fatigue Concerns	Findings
Respondents experiencing fatigue in the past 7 days	70%
	Always: 20%
Respondents experienced fatigue	Often: 32%
	Sometimes: 29%
	Very much: 16%
Pain interfered with their daily lives	Quite a bit: 21%
	Somewhat: 26%
	Very much: 8%
Pain interfered with respondents' ability to participate in social activities	Quite a bit: 19%
	Somewhat: 25%

Table B3 shows that pain is also a significant concern. One-third of respondents had pain often or always and 23% reported pain interfering with their daily lives "quite a bit" or "very much".

Table B3. CSC Multiple Myeloma Specialty Registry Findings: Pain & Bone Pain

Pain & Bone Pain Concerns	Findings
Respondents experiencing bone pain in the past 7 days	48%
	Always: 19%
Respondents experienced pain	Often: 15%
	Sometimes: 25%
	Very much: 13%
Pain interfered with their daily lives	Quite a bit: 10%
	Somewhat: 18%
Pain interfered with respondents' ability to participate	Very much: 8%
in social activities	Quite a bit: 11%
	Somewhat: 12%

Table B4 highlights to importance of financial concerns to patients. Forty-two percent of respondents were always or often upset about the cost of care. Fifty-six percent of respondents felt overwhelmed by the demands of paying for care at least some of the time.

Financial Concerns	Findings
Respondents received financial assistance related to their multiple myeloma	63%
	Always: 19%
Respondents felt upset about money and the cost of care	Often: 23%
	Sometimes: 21%
	Always: 8%
Respondents felt overwhelmed by the demands of paying for medical care	Often: 19%
	Sometimes: 29%
	Always: 9%
Respondents worried they won't be able to leave assets to their families	Often: 13%
	Sometimes: 22%

 Table B4. CSC Multiple Myeloma Specialty Registry Findings: Financial Concerns

C. Clinical Guidelines

Due to the number of treatments that have recently become available, we focus on guidelines that have been published in the last 2 years.

National Comprehensive Cancer Network (NCCN) Multiple Myeloma, V5, March 2021⁵²

The NCCN convened a panel of nationally recognized expert clinicians in the care of MM to develop a consensus statement on currently accepted approaches to treatment. While there are not specific recommendations for the triple/quad/penta refractory population that is the focus of our current review, they do provide recommendations on relapsed or refractory MM.

Recommendation 1 (MYEL-7): For relapsed patients, consider a) treatments for previously treated myeloma, b) clinical trial and c) allogeneic stem cell transplant.

Recommendation 2 (MYEL-7): For patients with refractory disease and lack of treatment options, refer to palliative care.

Recommendation 3 (MYEL-F, 3 of 3): The NCCN listed a wide range of therapeutic options for patients with relapsed, previously treated MM, representing the lack of clear evidence on the preferred ordering of treatments. Specifically, they listed 9 preferred regimens, 19 other recommended regimens and 17 regimens listed as "useful in certain circumstances". Most were triplet regimens and nearly all regimens included dexamethasone. In addition, bortezomib, pomalidomide, carfilzomib, daratumumab, ixazomib, elotuzumab, selinexor and panobinostat were common components of listed regimens.

American Society of Clinical Oncology (ASCO) and Cancer Care Ontario (CCO) Joint Clinical Practice Guideline for Multiple Myeloma, Apr 2019⁵³

The American Society of Clinical Oncology and Cancer Care Ontario convened an expert panel of medical oncologists, surgeons, radiation oncologists, and patient advocates to conduct a review of the literature to develop evidence-based guidelines. While they did not address triple-class refractory patients, they did produce recommendations for myeloma patients with a first relapse.

Recommendation 7.3: Triplet therapy (3 agents including a steroid, and 2 of the following 3 classes: proteasome inhibitor, immunomodulator and monoclonal antibody) is preferred. While toxicity appears to be increased with triplet vs doublet therapy, triplet therapy leads to improved PFS, ORR and OS, even in older adults.

Recommendation 7.5: Prior therapies should be taken into consideration when selecting the treatment in patient with relapsed multiple myeloma. Patients who have been off of a particular medication for >1 year are likely to respond to a repeat course of that medication. However, patients who relapse <1 year after exposure to a medication are less likely to respond; thus, novel medications are recommended in these situations.

D. Comparative Clinical Effectiveness: Supplemental Information

D1. Detailed Methods

PICOTS

Population

The population of focus for the review is patients who have at a minimum triple-class refractory MM, defined as disease that has progressed while on an anti-CD38 antibody (e.g., daratumumab), immunomodulatory drugs (e.g., lenalidomide), and a proteasome inhibitor (e.g., bortezomib). The indication for belantamab involves a population subset with somewhat more advanced disease (at least four prior lines of treatment, triple-, quad-, or penta-refractory patients) than those in the current Ide-cel and Cilta-cel trials (at least three prior lines, mostly triple- or quad-refractory). (Ide-cel was recently approved for patients with 4+ prior lines of therapy.) We therefore did not make any explicit comparisons between belantamab and Idecabtagene vicleucel and Ciltacabtagene autoleucel, and we summarized evidence on relevant comparator therapies to match these population differences (see below). Data permitting, we included evidence across relevant subgroups, such as patients with genetic factors that put them at particularly high risk as well as subgroups defined by race.

Interventions

The full list of interventions is as follows:

- Idecabtagene Vicleucel (Ide-cel, Abecma[®], Bristol Myers Squibb and bluebird bio)
- Ciltacabtagene Autoleucel (Cilta-cel, Janssen and Legend biotech)
- Belantamab mafodotin (Blenrep[®], GlaxoSmithKline)

Comparators

We used the characteristics of patients enrolled in each of the pivotal studies of the drugs under consideration to guide the most appropriate comparator treatments (i.e., the regimens patients would have received if the drugs under consideration were not an option). Since the belantamab study focused on a more heavily pre-treated population, the comparator cohort for belantamab differs from the comparator cohort for Ide-cel and Cilta-cel.

We compared the selected interventions to commonly used regimens in triple-class refractory populations as well as to palliative care (no active anti-cancer therapy). The comparator regimens include:

- Carfilzomib + cyclophosphamide + dexamethasone (KCd)
- Pomalidomide + cyclophosphamide + dexamethasone (PCd)
- Carfilzomib + pomalidomide + dexamethasone (KPd)
- Elotuzumab + pomalidomide + dexamethasone (EPd)

Outcomes

The outcomes of interest are described in the list below.

- Patient-Important Outcomes
 - o Overall survival (OS)
 - Quality of life
 - Complete response rate
 - Progression-free survival (PFS)
 - Durability of response
 - Pain and function
 - o Treatment burden
 - o Bone fractures
 - Adverse events including:
 - cytokine response syndrome
 - fatigue/sleep disturbance
 - infection
 - peripheral neuropathy
 - ocular toxicity
 - anemia
 - gastrointestinal toxicity
 - thromboembolism
 - death
- Other Outcomes
 - o Overall response rate
 - o Partial response rate
 - Minimal residual disease
 - Blood and urine markers of disease

Timing

Evidence on intervention effectiveness and harm was derived from studies of any duration.

Settings

Evidence from all relevant settings were considered.

Table D1.1 PRISMA 2009 Checklist

		Checklist Items
TITLE		
Title	1	Identify the report as a systematic review, meta-analysis, or both.
ABSTRACT		
Structured summary	2	Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number.
INTRODUCTION		
Rationale	3	Describe the rationale for the review in the context of what is already known.
Objectives	4	Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS).
METHODS		
Protocol and registration	5	Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number.
Eligibility criteria	6	Specify study characteristics (e.g., PICOS, length of follow-up) and report characteristics (e.g., years considered, language, publication status) used as criteria for eligibility, giving rationale.
Information sources	7	Describe all information sources (e.g., databases with dates of coverage, contact with study authors to identify additional studies) in the search and date last searched.
Search	8	Present full electronic search strategy for at least one database, including any limits used, such that it could be repeated.
Study selection	9	State the process for selecting studies (i.e., screening, eligibility, included in systematic review, and, if applicable, included in the meta-analysis).
Data collection process	10	Describe method of data extraction from reports (e.g., piloted forms, independently, in duplicate) and any processes for obtaining and confirming data from investigators.
Data items	11	List and define all variables for which data were sought (e.g., PICOS, funding sources) and any assumptions and simplifications made.
Risk of bias in	12	Describe methods used for assessing risk of bias of individual studies (including specification of whether this was done at
individual studies		the study or outcome level), and how this information is to be used in any data synthesis.
Summary measures	13	State the principal summary measures (e.g., risk ratio, difference in means).
Synthesis of results	14	Describe the methods of handling data and combining results of studies, if done, including measures of consistency (e.g., I2) for each meta-analysis.
Risk of bias across studies	15	Specify any assessment of risk of bias that may affect the cumulative evidence (e.g., publication bias, selective reporting within studies).
Additional analyses	16	Describe methods of additional analyses (e.g., sensitivity or subgroup analyses, meta-regression), if done, indicating which were pre-specified.
RESULTS	1	

Study selection	17	Give numbers of studies screened, assessed for eligibility, and included in the review, with reasons for exclusions at each
		stage, ideally with a flow diagram.
Study characteristics	18	For each study, present characteristics for which data were extracted (e.g., study size, PICOS, follow-up period) and provide
		the citations.
Risk of bias within	19	Present data on risk of bias of each study and, if available, any outcome level assessment (see item 12).
studies		
Results of individual	20	For all outcomes considered (benefits or harms), present, for each study: (a) simple summary data for each intervention
studies		group (b) effect estimates and confidence intervals, ideally with a forest plot.
Synthesis of results	21	Present results of each meta-analysis done, including confidence intervals and measures of consistency.
Risk of bias across	22	Present results of any assessment of risk of bias across studies (see Item 15).
studies		
Additional analysis	23	Give results of additional analyses, if done (e.g., sensitivity or subgroup analyses, meta-regression [see Item 16]).
DISCUSSION		
Summary of evidence	24	Summarize the main findings including the strength of evidence for each main outcome; consider their relevance to key
		groups (e.g., health care providers, users, and policy makers).
Limitations	25	Discuss limitations at study and outcome level (e.g., risk of bias), and at review-level (e.g., incomplete retrieval of identified
		research, reporting bias).
Conclusions	26	Provide a general interpretation of the results in the context of other evidence, and implications for future research.
FUNDING		
Funding	27	Describe sources of funding for the systematic review and other support (e.g., supply of data); role of funders for the
		systematic review.

From: Moher D, Liberati A, Tetzlaff J, Altman DG. The PRISMA Group (2009). Preferred Reporting Items for Systematic Reviews and Meta-Analyses: The PRISMA Statement. PLoS Med 6(6): e1000097. doi:10.1371/journal.pmed1000097

Data Sources and Searches

Procedures for the systematic literature review assessing the evidence on new therapies for relapsed and refractory multiple myeloma followed established best research methods.^{54,55} We conducted the review in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines.⁵⁶ The PRISMA guidelines include a checklist of 27 items, which are described further in <u>Supplemental Table D1.1</u>.

We searched MEDLINE and EMBASE. Each search was limited to English-language studies of human subjects and excluded articles indexed as guidelines, letters, editorials, narrative reviews, case reports, or news items. We included abstracts from conference proceedings identified from the systematic literature search. All search strategies were generated utilizing the Population, Intervention, Comparator, and Study Design elements described above. The proposed search strategies included a combination of indexing terms (MeSH terms in MEDLINE and EMTREE terms in EMBASE), as well as free-text terms.

To supplement the database searches, we performed manual checks of the reference lists of included trials and systematic reviews and invited key stakeholders to share references germane to the scope of this project. We also supplemented our review of published studies with data from conference proceedings, regulatory documents, information submitted by manufacturers, and other grey literature when the evidence met ICER standards (for more information, see https://icer.org/policy-on-inclusion-of-grey-literature-in-evidence-reviews/) Where feasible and deemed necessary, we also accepted data submitted by manufacturers "in-confidence," in accordance with ICER's published guidelines on acceptance and use of such data https://icer.org/guidelines-on-icers-acceptance-and-use-of-in-confidence-data-from-manufacturers-of-pharmaceuticals-devices-and-other-health-interventions/)

Table D1.2. Search Strategies for Ovid MEDLINE(R) Epub Ahead of Print, In-Process & Other Non-Indexed Citations, Ovid MEDLINE(R) Daily, Ovid MEDLINE and Versions(R) 1946 to Present*

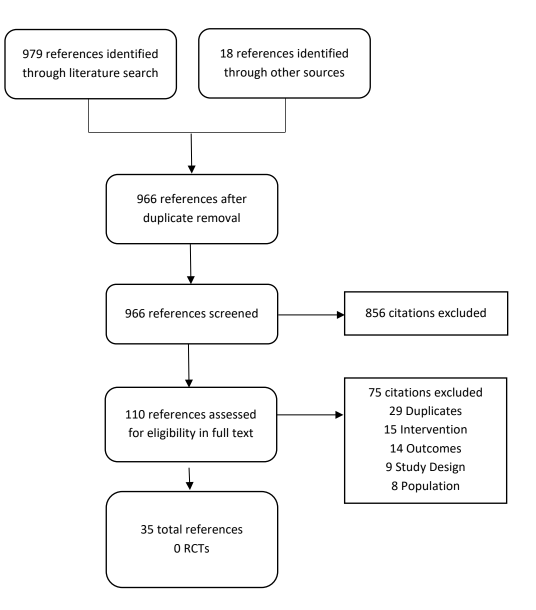
#	Search Term
1	Exp Multiple Myeloma/
2	("Multiple Myeloma" or "Multiple Myelomas" or "Myeloma, Multiple" or "Myeloma, Plasma cell" or "Plasma cell myeloma" or "Plasma cell myelomas" or "Myelomatosis" or "Kahler Disease" or "Myeloma" or "Myelomas").ti,ab OR (("relapsed" or "refractory" or "pretreated" or "high risk") and ("multiple myeloma")).ti,ab
3	1 or 2
4	("CAR T" or "CAR-T" or "chimeric antigen receptor" or "CAR-T cell" OR "Anti BCMA" or "anti-BCMA" or "CMA CAR-T" or "b-cell maturation antigen" OR "anti b-cell maturation antigen" OR "CD269").ti,ab
5	("blenrep" or "belantamab" or "belantamab mafodotin" OR "belantamab mafodotin-blmf" or "GSK2857916" or "GSK 2857916" OR "GSK-2857916" OR "gsk916" OR "gsk 916" or "gsk-916").ti,ab.
6	("bb2121" or "bb-2121" OR "bb 2121" or "idecabtagene vicleucel" or "ide-cel" OR "ide cel").ti,ab.
7	("JNJ-68284528" OR "JNJ68284528" OR "JNJ 68284528" OR "JNJ4528" OR "JNJ 4528" OR "LCAR-B38M" OR "cilta-cel" OR "cilta cel" OR "ciltacabtagene autoleucel").ti,ab.
	((("pomalidomide" or "pomalyst") and ("Cyclophosphamide" or "Cytoxan") and ("dexamethasone" or "decadron" or "Dexamethasone Intensol" or "Dexpak Taperpak" or "prednisone")) or (("Carfilzomib" or "Kyprolis") and ("pomalidomide" or "pomalyst") and ("dexamethasone" or "decadron" or "Dexamethasone Intensol" or "Dexpak Taperpak" or "prednisone")) OR (("Elotuzumab" OR "Empliciti") AND ("pomalidomide" OR "pomalyst") AND ("dexamethasone" OR "decadron" OR "Dexamethasone Intensol" OR "Dexpak Taperpak" or "prednisone")) OR (("Carfilzomib" OR "Kyprolis") AND ("Cyclophosphamide" OR "Cytoxan") AND ("dexamethasone" OR "Dexamethasone Intensol" OR "Dexpak Taperpak" or "prednisone"))).ti,ab.
9	4 or 5 or 6 or 7 or 8
10	3 and 9
11	animals not (humans and animals).sh.
12	10 not 11
13	limit 12 to English language
14	13 and ("chapter" OR "comment" OR "editorial" OR "letter" OR "note" OR "short survey" OR "review" OR "opinion").pt
15	13 not 14
* 500	rch last undated on March 08, 2021

* Search last updated on March 08, 2021.

#	Search Term
#1	'multiple myeloma'/exp
#2	('multiple myeloma' OR 'refractory multiple myeloma' OR 'relapsed multiple myeloma' OR 'Kahler disease' OR 'morbus Kahler' OR 'myeloma multiplex' OR 'Myelomatosis' OR 'Myeloma' OR 'Myelomas' OR 'plasma cell myeloma' OR (('relapsed' OR 'refractory' OR 'pretreated' OR 'high risk') and 'multiple myeloma')):ti,ab
#3	#1 or #2
#4	('car t' OR 'CAR-T' OR 'chimeric antigen receptor' OR 'anti-BCMA' OR 'anti bcma' OR 'b cell maturation antigen' OR 'anti b-cell maturation antigen' OR 'CD269'):ti,ab
#5	'belantamab mafodotin'/exp OR 'belantamab'/exp
#6	('gsk2857916' OR 'gsk 2857916' or 'gsk-2857916' or 'gsk916' or 'gsk 916' OR 'gsk-916' OR 'belantamab' or 'belantamab mafodotin' or 'belantamab mafodotin-blmf' or 'blenrep'):ti,ab
#7	#5 OR #6
#8	'idecabtagene vicleucel'/exp
#9	('bb2121' OR 'bb 2121' OR 'bb-2121' OR 'ide-cel' OR 'idecabtagene vicleucel' OR 'ide cel'):ti,ab
#10	#8 OR #9
#11	'jnj 68284528'/exp or 'ciltacabtagene autoleucel'/exp
#12	('jnj 4528' OR 'jnj68284528' OR 'lcar-b38m' or 'cilta-cel' or 'cilta cel' or 'ciltacabtagene autoleucel'):ti,ab
#13	#11 OR #12
#14	((('pomalidomide') AND ('cyclophosphamide') AND ('dexamethasone' OR 'prednisone')) OR ('carfilzomib' AND 'pomalidomide' AND ('dexamethasone' OR 'prednisone')) OR ('elotuzumab' AND ' pomalidomide' AND ('dexamethasone' or 'prednisone')) OR ('carfilzomib' AND 'cyclophosphamide' AND ('dexamethasone' OR 'prednisone'))):ti,ab
#15	#3 AND (#4 OR #7 OR #10 OR #13 OR #14)
#16	('animal'/exp OR 'nonhuman'/exp OR 'animal experiment'/exp OR 'animal model'/exp) NOT 'human'/exp
#17	#15 NOT #16
#18	#17 AND [English]/lim
#19	#18 AND ('chapter'/it or 'comment'/it OR 'editorial'/it OR 'letter'/it OR 'note'/it OR 'short survey'/it OR 'review'/it OR 'opinion'/it)
#20	#18 NOT #19

* Search last updated on March 08, 2021.

Figure D1. PRISMA flow Chart Showing Results of Literature Search for Idecabtagene vicleucel, Ciltacabtagene autoleucel, and Belantamab mafodotin



Study Selection

We performed screening at both the abstract and full-text level. Two investigators screened all abstracts identified through electronic searches according to the inclusion and exclusion criteria described earlier. We did not exclude any study at abstract-level screening due to insufficient information. For example, an abstract that did not report an outcome of interest would be accepted for further review in full text. We retrieved the citations that were accepted during abstract-level screening for full text appraisal. Two investigators reviewed full papers and provided justification for exclusion of each excluded study.

We also included FDA documents related to belantamab and ide-cel. These included the manufacturer's submission to the agency, internal FDA review documents, and the transcript of Advisory Committee deliberations and discussions. All literature that did not undergo a formal peer review process is described separately.

Data Extraction and Quality Assessment

Because we did not identify any comparative trials of the interventions, we did not assess the quality of the individual trials.

Assessment of Level of Certainty in Evidence

We used the <u>ICER Evidence Rating Matrix</u> to evaluate the level of certainty in the available evidence of a net health benefit among each of the interventions of focus (see Figure 3.1 of the main report).^{57,58}

Assessment of Bias

As part of our quality assessment, we evaluated the evidence base for the presence of potential publication bias. We performed an assessment of publication bias for Ide-cel, Cilta-cel and belantamab using the clinicaltrials.gov. We scanned the site to identify studies completed more than two years ago that would have met our inclusion criteria and for which no findings have been published and did not find any evidence of publication bias. The primary concern is the lack of peer-reviewed, published data for the KarMMa and CARTITUDE-1 trials as well as the lack of head-to-head trials of the interventions compared to usual care.

Data Synthesis and Statistical Analyses

Data on relevant outcomes were summarized in evidence tables (<u>See Supplement D3</u>) and synthesized qualitatively in the body of the review. Due to differences in study populations and limitations of study design (single-arm trials), outcomes are described for each trial separately.

D2. Additional Clinical Evidence

Evidence Base

Ide-cel

In addition to the pivotal phase II KarMMa trial described in the main report, we also identified a phase I trial of bb2121 (ide-cel), CRB-401.^{59,60} CRB-401 was a phase 1 open-label single-arm dose-escalation and dose-expansion, multi-center US-based trial. The trial enrolled 62 adults (21 in the dose-escalation and 41 in the dose-expansion phase) who had previously received three lines of therapy (including an IMiD and a PI), or were refractory to both classes. The dose-expansion phase also required exposure to daratumumab and that patients be refractory to the last line of therapy. Patients underwent leukapheresis, bridging therapy during manufacturing, and lymphodepletion with fludarabine and cyclophosphamide prior to infusion with 50, 150, 450, or 800x10⁶ CAR-T cells in the dose-escalation phase and 150 to 450x10⁶ CAR-T cells in the expansion phase. 33 patients were analyzed in the original publication and 62 patients were analyzed in the updated analysis as of January 2020. ^{59,60} The primary outcome was safety. Secondary outcomes were overall response rate (ORR) and duration of response.

More details on both KarMMa and CRB-401 are provided in Table D3.1.

Cilta-cel

In addition to the pivotal phase Ib/II CARTITUDE-1 trial described in the main report, we also identified a phase I trial of cilta-cel (LEGEND-2). LEGEND-2 is a Phase I single-arm trial conducted at four sites in China.^{14,61,62} The trial enrolled 74 adults with TCRMM who had progressive disease after at least 3 prior therapies (including a PI, an IMiD, and anti-CD38 antibody). The trial explored the differences between a single and three CAR-T cell infusion approach with varying doses (0.2-2.0×10⁶ CAR-T cells/kg) as well as cyclophosphamide alone versus in combination with fludarabine during conditioning. For this report, we will include data from the largest study site (Xi'an, N=57) because data was not aggregated across all four sites. The primary outcome was AEs; the secondary outcome was CR.

More details on both trials are provided in Table D3.1.

Belantamab

In addition to the pivotal phase II, open-label, two-arm, multicenter trial of belantamab (DREAMM-2), we identified a pooled post-hoc analysis of DREAMM-1 and DREAMM-2 and one expanded access study. Neither of the two additional studies had been published at the time of this report, and data was only available in form of conference abstracts/posters.

<u>DREAMM-2</u>: DREAMM-2, the pivotal trial of belantamab in adults with triple-class refractory multiple myeloma, is an ongoing global, open-label, phase II randomized multicenter trial comparing the efficacy and safety of two doses of belantamab (2.5 mg/kg and 3.4 mg/kg).¹⁷ Patients were treated with intravenous belantamab every three weeks until disease progression, or unacceptable toxicity occurred. Dosing delays or reductions were permitted for the management of adverse events. Efficacy and safety outcomes were assessed every three weeks after the first course of treatment had been administered.¹⁷ Due to the risk for ocular toxicity of belantamab, ophthalmic testing was required prior to each round of treatment.¹⁷ To further mitigate corneal events, patients were administered prophylactic corticosteroid eye drops as well as artificial tears.

Pooled post-hoc analysis: Trudel 2020 presented pooled tolerability and safety data from the DREAMM-1 and DREAMM-2 trials.⁶³ DREAMM-1 was an open-label phase I trial and included adult patients with RRMM, who had previously failed 3 or more lines of treatment, and were refractory to an alkylator, PI, and IMiD. DREAMM-2 has been described previously in this report. A total of 264 patients were randomized to either 2.5 mg/kg (N=103) or 3.4 mg/kg (N=161) every three weeks.⁶³ For the purpose of this review, we will only discuss the arm that received the FDA approved dose of 2.5 mg/kg.

The median age of patients was 65 years and 50% of the randomized patients were male. The study population also included a subset of RRMM patients at higher risk of more aggressive disease, including 40% with ISS stage III, 20% who exhibited extramedullary disease (EMD), and 27% with high-risk cytogenetic features. The median number of prior lines of therapy was 7 (range 3 to 21).

<u>Expanded Access Program</u>: This multicenter, observational study included 32 patients with RRMM who were treated under the expanded access compassionate care program at 6 Israeli multiple myeloma centers.⁶⁴ The primary outcome assessed was progression-free survival. Secondary outcomes included overall response rate, overall survival, as well as safety and tolerability.

Between July 2019 and February 2020, 32 patients were treated with at least one dose of belantamab (median 3; range 1-11) and identified for inclusion in this study. Median follow-up duration was 5.7 months (range 0.5 – 13.8 months) and data were obtained from medical charts. A total of 13 patients received the 2.5 mg/kg dose, and 17 patients received the 3.4 mg/kg dose, respectively. Of note, this study did not present stratified results of the two separate doses, and therefore, the results presented contain both doses. Thus, these results should be synthesized cautiously with other studies which focused solely on patients receiving the FDA approved 2.5 mg/kg dose.

Patients included had a median age of 70 years, over half were male (59%), and roughly 20% were considered to have high-risk cytogenetics. The heavily pre-treated patients had received a median of 6 prior lines of treatments (range 3-11), with a majority having been previously exposed to bortezomib (94%), carfilzomib (74%), lenalidomide (91%), pomalidomide (87%), or daratumumab

(97%). The overwhelming majority of included participants (97%) had also received an autologous stem-cell transplant.

Usual Care

The main report discusses the primary source of outcomes data to inform our comparison of the interventions to usual care, the MAMMOTH study.⁸ Two additional retrospective studies were identified with sufficient numbers of triple-class refractory patients (Mehra 2020 and Goldsmith 2020). In Mehra 2020, patient data were abstracted from a US-based electronic health record system from January 2011 to October 2019.²⁶ A total of 251 patients with at least triple-class refractory multiple myeloma were included in the analysis. Of those, 73 (29%) were "penta-refractory". Primary outcomes were overall survival (OS), progression free survival (PFS), and time to next treatment (TTNT). In Goldsmith 2019, data from 58 patients were abstracted from health records at a single US-based academic center from January 2013 and August 2018.²⁵ Patients were either quad or penta-refractory and treated with at least one cycle of bendustamine/prednisone (BP) or dexamethasone, cyclophosphamide, etoposide, cisplatin (DCEP). Primary outcomes were PFS and OS. Additional information on the study design of the usual care studies are available in Table D3.18 and baseline characteristics are available in Table D3.19.

Effectiveness

Ide-cel

Outcomes from the pivotal phase II trial of ide-cel (KarMMa) are described in the main report. In the phase 1 dose-escalation/expansion trial, CRB-401 (n=62), as of January 2020, median follow-up was 14.7 months, as-treated median PFS was 8.8 months (95% CI: 5.9-11.9 months), and as-treated median OS was 34.2 months (95% CI: 19.2-not estimable). As-treated ORR was 75.8% and as-treated stringent or complete response (sCR or CR) was 38.7%. As with KarMMa, ORR appeared to be dose-related in CRB-401. As-treated ORR in the higher dose (450x10⁶) was 90% compared to 50% in the lower dose (150x10⁶). As-treated median duration of response was 10.3 months overall (95% CI: 7.7-13.7 months).⁶⁰ More detailed outcomes from both the KarMMa and CRB-401 trial are provided in Table D3.3. Details on HRQoL outcomes from KarMMa are provided in Table D3.10.

Cilta-cel

The main report provides outcomes from the pivotal phase Ib/II trial (CARTITUDE-1) In the Phase I trial (LEGEND-2, Xi'an study site), with a median follow-up time of 25 months, median PFS was 19.9 months, median OS was 36 months, and ORR was 87.7%.^{65,66} Because all patients in LEGEND-2 who were enrolled were leukapheresed and underwent infusion, an ITT analysis was not applicable, however comparisons based on data from LEGEND-2 should be interpreted with caution due to differences in the patient population. Detailed outcomes from both trials are provided in Table D3.4. Details on HRQoL outcomes from CARTITUDE-1 are provided in Table D3.10.

Belantamab

<u>DREAMM-2</u>: The DREAMM-2 study reported ITT results; thus, the following results are all calculated on an ITT basis. At 13-month follow-up (data cut-off date: January 2020), 13 and 11 patients treated with 2.5 mg/kg belantamab achieved PR and vgPR, respectively. The median duration of response was 11 months (95% CI: 4.2 to not reached). Time to initial response and time to CR was not reported.²² Out of 97 participants, 59 had progressive disease at 6.3 months of follow-up.¹⁷ More detailed outcomes from DREAMM-2 are provided in Table D3.5 and for additional data on HRQoL please refer to Table D3.10.

<u>Expanded Access Program</u>: Overall, the median duration of PFS achieved by DREAMM-2 trial participants was 2.6 months. Twelve out of 29 patients who were evaluable, achieved an ORR (with 3 participants achieving PR, 8 achieving VGPR, and one participant achieving CR). OS at 6 months was 68%. Statistical analyses were not reported. A subgroup analysis showed that ORR and PFS were comparable between patient who had been previously treated with \leq 5 prior lines of treatments, and those who had previously received more than 5.⁶⁴ More detailed outcomes from the Expanded Access Program can be found in Table D3.17.

Usual Care

Outcomes from the retrospective MAMMOTH study are discussed in the main report.⁸ We also identified two additional retrospective studies (Goldsmith 2020 and Mehra 2020) to inform our comparison of usual care to the interventions. In these trials, median PFS varied from 1.4 months to 4.8 months, median OS varied from 6.2 months to 11.0 months, and ORR was around 31%^{25,26}. The patient populations of these studies varied, particularly in the percentage of patients with penta-refractory disease (29% in Mehra and 78% in Goldsmith). Additional outcomes data for the MAMMOTH study is available in Table D3.20 and for the two additional retrospective studies is available in Table D3.21.

Harms

Ide-cel

AEs from the pivotal phase II trial of ide-cel (KarMMa) are described in the main report. In the phase 1 dose-escalation/expansion trial, CRB-401 (n=62), the most frequent grade three or higher AEs were neutropenia (92%), leukopenia (61%), anemia (57%), and thrombocytopenia (57%). CRS was also common, occurring in 76% of patients and requiring tocilizumab in 21%. Sixty-nine percent had low to moderate (Grade 1 or 2) CRS, 7% had severe (Grade 3) CRS, and none had a grade 4 CRS. As in the KarMMa trial, risk of CRS in CRB-401 was dose-related. 92% of patients in the higher dose group (450×10^6) reported any grade CRS compared to 11% in the lower dose group (150×10^6). 82% of patients ≥ 65 experienced any grade CRS. As of the January 2020 cutoff date, 49 (79%) had discontinued due to progressive disease (58%), withdrawal by patient (10%), or death (10%).⁶⁷ More detailed safety data from both the KarMMa and CRB-401 trials are provided in Table D3.11.

Cilta-cel

The main report provides adverse events from the pivotal phase Ib/II trial of cilta-cel (CARTITUDE-1). In the phase I trial (LEGEND-2, Xi'an site) of cilta-cel, CRS was common (89.5%) and with 25 months of follow-up, 17 deaths (29.8%) were reported, 14 (24.6%) due to progressive disease and 2 (3.5%) due to AEs. Other important AEs included thrombocytopenia (49.1%) and neurotoxicity (1.8%).^{65,66} Detailed safety data from both trials are provided in Table D3.12.

Belantamab

<u>DREAMM-2</u>: Most dose delays, as well as dose reductions were due to keratopathy (45/51 and 24/33 patients, respectively).²² Adverse events grade 3 or above were reported by 84% of patients treated with belantamab, and the most commonly reported events were keratopathy (46%), thrombocytopenia (22%), and anemia (21%).

Median time to onset of the first corneal event was 37 days and lasted for a median duration of 86.5 days.¹⁸ The occurrence of ocular toxicities increased with increasing number of doses; 25% of patients reported ocular toxicities after the first dose, 69% reported their first corneal event after by the fourth dose. Only two patients developed a corneal event subsequent to having received 4 doses. Overall, 24 patients reported experiencing blurred vision and 14 patients experienced dry eyes. CRS or neurotoxicity was not reported by any patient in the DREAMM-2 trial. Please refer to Table D3.13 for a more detailed description of safety data.

<u>Pooled Analysis</u>: Patients received a median of 3 courses of treatment with 2.5 mg/kg belantamab (range 1 - 15). Most patients (98%) experienced at least one AE, of which 90% were considered treatment related. Serious adverse events (SAEs) were reported by 42 participants, with 13

reporting SAEs related to the study treatment. One participant died due a treatment-related SAE (sepsis). Keratopathy, which was only assessed in the DREAMM-2 trial, was the most commonly reported AE of any grade (66%), followed by anemia (26%) and thrombocytopenia (23%). Blurred vision and dry eye were reported by 20 and 12 patients, respectively. Grade 3/4 adverse events most frequently experienced by patients were keratopathy (27%), anemia (18%) and thrombocytopenia (17%). Four patients experienced grade 3/4 blurred vision, and no patients reported experiencing grade 3/4 dry eye. Generally, adverse events were managed by means of dose reductions (32%) and/or delays in treatment administration (51%). Keratopathy was the most frequently cited reason for delays or reduction in dosing by 45% and 24% of patients, respectively, as well as for treatment discontinuation (2%).

Expanded Access Program: Seventeen study participants (53%) were still receiving treatment at the time of data cut-off, while the remaining 15 had discontinued treatment. The most commonly cited reason for treatment discontinuation was progression or death (13 patients). Twenty out of 31 evaluable patients experienced ocular toxicity (keratopathy) of any grade, and eight reported grade \geq 3 keratopathy. 62.5% of patients who experienced grade \geq 3 ocular toxicity reported an improvement to grade \leq 2 ocular toxicity, and one person discontinued treatment. Other adverse events commonly reported by \geq 20% of study participants were thrombocytopenia (30%), neutropenia (22%), and infections (22%). Thrombocytopenia, neutropenia, and infections grade 3 or higher were reported by three, four, and three patients, respectively. Adverse events were managed by means of dosing delays (13 patients) and dosing reductions (11 patients). Ocular toxicity was most commonly cited as requiring dosing delays or reductions (9 patients).

Please refer to Table D3.17 for a more detailed description of safety data for the Pooled Analysis and Expanded Access Program.

Subgroup Analyses

<u>Ide-cel</u>

Subgroup data from the pivotal phase II trial of ide-cel (KarMMa) are described in the main report. In the phase 1 dose-escalation/expansion trial, CRB-401, subgroup efficacy data was only available from the original publication (n=33).⁵⁹ At a median of 11.3 months of follow-up, as-treated ORR was 85% (28 patients had a response out of 33 infused). Response appeared to be dose-related, with the highest dose (>150x10⁶ CAR-T cells) achieving the highest as-treated ORR (96%). Those with high cytogenetic risk (n=15) had a lower as-treated ORR (73%).⁵⁹

More detailed subgroup data from both the KarMMa and CRB-401 trial are provided in Tables D3.6 and D3.7.

<u>Cilta-cel</u>

At the time of the report, subgroup data from the pivotal phase Ib/II trial (CARTITUDE-1) of cilta-cel was not available. In the Phase I trial (LEGEND-2, Xi'an site), as-treated median PFS for cilta-cel for patients with EMD was significantly lower (8.1 months) than for patients without EMD (25 months, p<0.001).⁶⁸ Additional subgroup data is presented in Table D3.8.

<u>Belantamab</u>

In DREAMM-2 the probability of PFS reaching a duration of 6 months or more was 35% for patients who had received 3 to 6 prior lines of treatment, and 30% for both those who had received 7 prior lines of therapy or more, and for patients with high-risk cytogenetics. Median OS, which was only reported for the subgroup considered to have high-risk cytogenetics, was 9.4 months. The probability of OS at 12 months was 45%.

At 12.4 months of median follow up time, 34% of the patients who had received three to six prior treatments achieved an overall response compared to 30% of patients who had received seven prior lines of treatments or more. Very good partial response was achieved by 17% of those who had received three to six prior lines of therapies and by 20% of those who had received seven or more.

ORR at nine months was achieved by 27% who had a high-risk cytogenetic risk profile, with 22% achieving a vgPR. Of the patients who had mild to moderate renal impairment, ORR was achieved by 31.3% and 33.3%, respectively. The median duration of response achieved by patients who had been treated with three to six prior lines of therapy was 11 months versus 13.1 months for those who had received seven or more. The probability of a response lasting six months or longer was 63% and 73%, respectively. At nine months of follow-up, the median duration of response for the high-risk cytogenetics and renal impairment subgroups had not yet been reached. The probability of a response lasting for six months or more was 52% for the high-risk cytogenetics subgroup. For the renal impairment subgroups (mild to moderate), the probability of a duration of response of six months or more was not reported.

More patients with moderate renal impairment experienced serious adverse events (50%) compared to those with mild renal impairment (33.3%). Of those with high-risk cytogenetics, 46.3% reported experiencing SAEs. For more information regarding subgroup safety outcomes refer to Table D3.13.

<u>Usual Care</u>

As discussed in the main report, the retrospective studies we selected to represent the effectiveness of usual care did not provide sufficient information on the harms of the treatment regimens. See Table D3.22 for what safety data was available in two of the retrospective studies.^{8,25}

Therefore we selected representative prospective trials of commonly used treatments that make up components of the market basket of therapies in the economic model (Elo-Pom-Dex: elotuzumab-pomalidomide-dexamethasone, Car-Cy-Dex: carfilzomib-cyclophosphamide-dexamethasone, Ixa-Len-Dex: ixazomib-lenalidomide-dexamethasone).28-30 In these prospective trials, treatment-related AEs were reported by 7-8% of patients. Discontinuation rates varied from 14 to 18%. Grade 3 or 4 AEs were common (57-74%). The most common grade 3 or 4 AEs were neutropenia (13-22%), anemia (9-11%), and thrombocytopenia (4-19%). Mortality ranged from 4.2% to 21.7%, however follow-up time varied (median 9 to 23 months). Deaths were most likely due to progressive disease (3.6-13.3%) followed by AEs (7.1-8.3%). Differences in harms between these regimes and that of the interventions should be interpreted with caution, however, as the trials were generally conducted in less heavily pre-treated populations (median of 2-3 prior therapies). See Table D3.23 for more details on harms of the usual care treatment regimens.

D3. Evidence Tables

Interventions

Table D3.1. Study Design: Interventions

Trial (NCT)	Study Design & Location	Treatment	Inclusion Criteria	Exclusion Criteria
			lde-cel	
KarMMa ^{11,13} (NCT03361748)	Phase II, open label, two-part, single-arm, multicenter trial N (enrolled) = 149 N (leukapheresed) = 140 N (treated) = 128 <u>Location</u> : Global	- 150 x 10 ⁶ CAR+ T cells/kg - 300 x 10 ⁶ CAR+ T cells/kg - 450 x 10 ⁶ CAR+ T cells/kg Single infusion	 2 18 years of age Documented diagnosis of multiple myeloma Received ≥ 3 prior treatment regimens, including a PI, IMiD, and an anti-CD38 antibody Must have undergone ≥2 consecutive treatment cycles for each regimen Must be refractory to last treatment regimen ECOG status of 0-1 Subjects must have measurable disease 	 History of clinically relevant central nervous system pathology Active or history of plasma cell leukemia Solitary plasmacytomas or non-secretory myeloma without other evidence of measurable disease Inadequate organ function Ongoing treatment with chronic immunosuppressants Previous allogeneic hematopoietic SCT; or treatment with any gene therapy-based therapy for cancer or investigational cellular therapy for cancer or BCMA targeted therapy HIV, hepatitis B, hepatitis C History of class III or IV, CHF or severe non-ischemic cardiomyopathy, history of stroke, unstable angina, myocardial infarction, or ventricular arrhythmia within the previous 6 months
CRB-401 ^{59,67,69} (NCT02658929)	Phase I, open- label, single- group, multicenter trial N = 62 <u>Location</u> : United States	Dose-escalation phase (N=21): – 150 x 10 ⁶ CAR+ T cells/kg – 450 x 10 ⁶ CAR+ T cells/kg *50 x 10 ⁶ and 800 x 10 ⁶ CAR+ T cells/kg arms were not abstracted	 18 years of age and older ≥3 lines of therapy (including a Pl and IMiD) ECOG status of 0-1 Measurable disease or more than 30% bone marrow plasma cells 	 Secondary malignancies in addition to myeloma Known CNS disease Inadequate hepatic, renal, bone marrow function Presence of active infection within 72 hours Malignancies in addition to myeloma if the second malignancy has required therapy in the last 3 years or is not in complete remission History of class III or IV CHF or non-ischemic cardiomyopathy, unstable angina, myocardial infarction, or ventricular arrhythmia requiring

Trial (NCT)	Study Design & Location	Treatment	Inclusion Criteria	Exclusion Criteria
		Expansion phase (N=41):		medication or mechanical control within the
		-150×10^6 CAR+ T cells/kg		previous 6 months
		-450×10^6 CAR+ T cells/kg		– HIV
				 Plasma cell leukemia or clinically significant
				amyloidosis
			Cilta-cel	
CARTITUDE-	Phase Ib/II,	<u>Target dose</u> :	$- \ge 18$ years of age	 Prior CAR-T treatment directed at any target
1 ^{16,19}	single-group,	-0.75×10^{6} CAR+ T	 Documented MM diagnosis 	 Prior therapy that is targeted to B-cell maturation
(NCT03548207)	open-label,	cells/kg (Range: 0.5–1.0	per IMWG criteria	antigen (BCMA)
	multi-center	× 10 ⁶)	– ECOG status ≤1	 NYHA stage III or IV CHF; myocardial infarction or
	study		 Measurable disease 	CABG within 6 months; history of clinically
			− Previously received \geq 3	significant ventricular arrhythmia or unexplained
	N (enrolled) =		therapies (including PI, IMiD,	syncope; history of severe non-ischemic
	126	Median Dose:	anti-CD38 antibody therapy)	cardiomyopathy; impaired cardiac function
	N (apheresed) =	-0.71 x 10 ⁶ CAR+ T	or are double refractory to	 Received a cumulative dose of corticosteroids
	113	cells/kg (Range: 0.51-	an IMiD and PI	equivalent to >= 70 mg of prednisone within 7 days
		0.95 x 10 ⁶)	 Documented evidence of 	prior to apheresis
	N (treated) = 97		progressive disease per	 Received an allogenic SCT within 6 months or an
		Single infusion	IMWG criteria or within 12	autologous SCT within 12 weeks
	Location: Japan,		months of most recent	 History of CNS involvement or clinical signs of
	United States		therapy	meningeal involvement of multiple myeloma
LEGEND-2 ^{62,70}	Phase I, open-	(Xi'an site) Median dose:	 – 18-75 years of age 	 NYHA Stage III-IV CHF, myocardial infarction, or
(NCT03090659)	label, single-	-0.5×10^{6} CAR+ T cells/kg	 Documented initial diagnosis 	CABG ≤6 months prior
	group,	(Range: 0.07 - 2.1 × 10 ⁶)	of multiple myeloma	 History of ventricular arrythmia or unexplained
	multicenter trial		according to IMWG	syncope
		<u>(Changzheng, Jiangsu,</u>	diagnostic criteria	 Impaired cardiac function (LVEF <45%)
	N = 74	Ruijin sites) Mean dose:	 Measurable disease at 	 Systemic corticosteroid therapy of greater than 5
		-0.7 x 10 ⁶ CAR+ T cells/kg	screening	mg/day of prednisone (or equivalent dose of
	Location: China		- Received at least 3 prior lines	another corticosteroid) within 2 weeks
		3 separate infusions within	of treatment for multiple	 Received autologous SCT within 12 weeks
		7 days (One clinical site	myeloma (incl. PI and/or	 Received allogeneic SCT
		administered the	IMiD)	
		treatment in one single-	 Documented disease 	
		dose)	progression during/within 12	

Trial (NCT)	Study Design & Location	Treatment	Inclusion Criteria	Exclusion Criteria
			months of most recent anti-	
			myeloma therapy	
			 ECOG status of 0 -2 	
			Belantamab	
DREAMM-2 ^{17,71} (NCT03525678)	Phase II, open- label, multicenter study N (total) = 196 N (2.5mg/kg dose) = 97 <u>Location</u> : Global	2.5 mg/kg every three weeks The studied 3.4 mg/kg dose was not approved by the FDA and will not be presented here	 ≥ 18 years of age Confirmed relapsed or refractory multiple myeloma according to IMWG ECOG status of 0-2 Received ≥3 previous lines of anti-myeloma treatments Refractory to an PI and IMiD, refractory and or intolerant to an anti-CD38 monoclonal antibody Autologous SCT ineligible or transplantation >100 days Adequate organ system function 	 Previous BCMA therapies, systemic high-dose corticosteroids, or investigational drug Received allogeneic SCT Current corneal epithelial disease (except mild punctate keratopathy) Active renal condition; active mucosal or internal bleeding POEMS syndrome

BCMA: B-cell maturation antigen, CABG: coronary artery bypass graft, CAR+: chimeric antigen receptor positive, CHF: congestive heart failure, CNS: central nervous system, ECOG: Eastern Cooperative Oncology Group, IMWG: International Myeloma Working Group, IMiD: immunomodulatory drug, kg: kilogram, LVEF: left ventricular ejection fraction, mg: milligram, N: total number, NYHA: New York Heart Association, PI: proteasome inhibitor, POEMS: polyneuropathy, organomegaly, endocrinopathy, myeloma protein, and skin changes, SCT: stem cell transplant

Int	tervention			Ide-cel				Cilta-cel		Belantamab	
	Trial		KarN	IMa ^{11,12}		CRB-401 ^{59,67}	CARTITUDE- 1 ¹⁹	LEGEND-2⁶² Xi'an	LEGEND-2⁶² Changzheng, Ruijin, Jiangsu	DREAMM- 2 ¹⁷	
	Arms		300 x10 ⁶ CAR+ T cells/kg	450 x10 ⁶ CAR+ T cells/kg	Overall	Overall	0.71x10 ⁶ CAR+ T cells/kg	0.5 × 10 ⁶ CAR+ T cells/kg	0.71x10 ⁶ CAR+ T cells/kg	Belantamab (2.5 mg/kg)	
	Ν	4	70	54	128	62**	97	57	17	97	
Age, Med	ian Years (Range)	54.0 (49.0- 69.0)	61.0 (33.0- 76.0)	62.0 (43.0- 78.0)	61.0 (33.0- 78.0)	61.0 (NR)	61.0 (43.0- 78.0)	54.0 (27.0- 72.0)	55.1 (35.0- 73.0)	65.0 (IQR: 60-70)	
Μ	ale, n (%)	4 (100)	38 (54.3)	34 (63.0)	76 (59.4)	21/33 (63.6)	57 (58.8)	34 (59.6)	11 (64.7)	51 (52.6)	
Race, n (%)	White	NR	NR	NR	NR	NR	NR	NR	NR	72 (74.2)	
Race, II (76)	Black	INIT								16 (16.5)	
	Time since Diagnosis, Median Years (Range)		7.0 (2.0- 18.0)	6.0 (1.0- 17.0)	6.0 (1.0-18.0)	5.0 (1.0– 36.0); N=33	5.9 (1.6- 18.2)	4.0 (1.0-9.0)	NR	5.5 (IQR: 4.0-7.0)	
Tumor BCM	A Expression [*] , n (%)	4 (100)	60 (85.7)	45 (83.3)	109 (85.2)	23/33 (69.7)	57 (91.9)	NR	16 (94.1)	NR	
	Extramedullary Disease	0 (0)	34 (48.6)	16 (29.6)	50 (39.1)	9/33 (27.2)	13 (13.4)	17 (29.8)			22 (22.7)
	Received Bridging Therapy Prior to Lymphodepletion	4 (100)	61 (87.1)	47 (87.0)	112 (87.5)	14/33 (42.4)¤	73 (75.2)	NR		NR	
High Risk Population,	High Cytogenetic Risk	1 (25.0)	20 (28.6)	24 (44.4)	45 (35.2)	15/33 (45.5) [§]	23 (23.7)	NR	NR	41 (42.3)	
n (%)	High Tumor Burden	3 (75.0)	34 (48.6)	28 (51.9)	65 (50.8)	16/33 (48.5)	NR	NR		NR	
	ISS Disease Stage III	1 (25.0)	12 (17.1)	8 (14.8)	21 (16.4)	8/33 (24.2)	NR	21 (36.8)		42 (43.3)	
	>1 Treatment Regimen per Year	2 (50.0)	36 (51.4)	22 (40.7)	60 (46.9)	NR	NR	NR		NR	
	of Prior Regimens, lian (Range)	9 (4-12)	6 (3-16)	5 (3-13)	6 (3-16)	7 (3–23); N=33	6 (3-18)	3 (1-9)	4 (3-11)	7 (3-21)	
Triple-	exposed, n (%)	128 (100)#	128 (100)#	128 (100)#	128 (100)#	62/62 (100)	97 (100)	NR	17 (100)	97 (100)	
Triple-re	efractory, n (%)	4 (100)	60 (85.7)	44 (81.5)	108 (84.4)	NR	85 (87.6)	NR	NR	97 (100)	

Table D3.2. Baseline Characteristics: Ide-cel, Cilta-cel, and Belantamab

	ervention			Ide-cel			Cilta-cel			Belantamab
	Trial		KarN	IMa ^{11,12}		CRB-401 ^{59,67}	CARTITUDE- 1 ¹⁹	LEGEND-2⁶² Xi'an	LEGEND-2 ⁶² Changzheng, Ruijin, Jiangsu	DREAMM- 2 ¹⁷
	Arms	150x10 ⁶ CAR+ T cells/kg	300 x10 ⁶ CAR+ T cells/kg	450 x10 ⁶ CAR+ T cells/kg	Overall	Overall	0.71x10 ⁶ CAR+ T cells/kg	0.5 × 10 ⁶ CAR+ T cells/kg	0.71x10 ⁶ CAR+ T cells/kg	Belantamab (2.5 mg/kg)
Ν		4	70	54	128	62**	97	57	17	97
Penta-e	exposed, n (%)	NR	NR	NR	77 (60.2)	26/33 (78.8)	81 (83.5)	NR	7 (41.2)	
Penta-re	efractory, n (%)	1 (25.0)	24 (34.3)	8 (14.8)	33 (25.8)	6/33 (18.2)	41 (42.3)	NR	NR	
Anti-CD38 A	b-refractory, n (%)	4 (100)	66 (94.3)	50 (92.6)	120 (93.8)	NR	96 (99.0)	NR	NR	97 (100)
	Bortezomib					33/33 (100)		39 (68.4)	14 (82.4)	95 (97.9)
Prior Therapies Received, n (%)	Carfilzomib				NR	30/33 (90.9)	_	1 (1.8)	2 (11.8)	74 (76.3)
	Lenalidomide	- NR	NR	NR		33/33 (100)	NR	25 (43.9)	10 (58.8)	97 (100)
	Pomalidomide					31/33 (93.9)	INK	2 (3.6)	1 (5.9)	89 (91.8)
	Daratumumab					56/62 (90.0)		NR	NR	97 (100)
	Isatuximab					NR		NR	1 (5.9)	3 (3.1)
	Bortezomib	NR	NR	NR	NR	20/33 (60.6)	NR			74 (76.3)
Refractory	Carfilzomib	NR	NR	NR	NR	19/33 (57.6)	63 (64.9)			63 (64.9)
to Prior	Lenalidomide	NR	NR	NR	NR	24/33 (72.7)	NR		ND	87 (89.7)
Therapies,	Pomalidomide	NR	NR	NR	NR	26/33 (78.8)	81 (83.5)	NR	NR	84 (86.6)
n (%)	Daratumumab	3 (75.0)	61 (87.1)	45 (83.3)	109 (85.2)	48/62 (77.0)	NR			97 (100)
	Isatuximab	NR	NR	NR	NR	NR	NR			3 (3.1)
	e of Therapy due to oriness, n (%)	4 (100)#	70 (100)#	54 (100)#	128 (100)#	21/33 (63.6) [‡]	96 (99.0)	NR	NR	NR
Received Autologous	1	4 (100)	67 (95.7)	49 (90.7)	120 (93.8)	32/33 (97.0)	87 (89.7)	10 (17.5)	8 (47.1)	
SCT, n (%)	≥1	3 (75.0)	23 (32.9)	18 (33.3)	44 (34.4)				0 (47.1)	
EORTC QLQ-C30,	Fatigue Pain	NR	NR	NR	39.3 (24.4 ⁺) 39.9 (28.1 ⁺)	NR	NR	NR	NR	NR

Int	ervention			Ide-cel				Cilta-cel		Belantamab
	Trial Arms		KarMMa ^{11,12}				CARTITUDE- 1 ¹⁹	LEGEND-2⁶² Xi'an	LEGEND-2 ⁶² Changzheng, Ruijin, Jiangsu	DREAMM- 2 ¹⁷
			300 x10 ⁶ CAR+ T cells/kg	450 x10 ⁶ CAR+ T cells/kg	Overall	Overall	0.71x10 ⁶ CAR+ T cells/kg	0.5 × 10 ⁶ CAR+ T cells/kg	0.71x10 ⁶ CAR+ T cells/kg	Belantamab (2.5 mg/kg)
	N	4	70	54	128	62**	97	57	17	97
Mean Score (SD)	Physical Functioning				69.4 (25.1 ⁺)					
	Global Health/QoL				60.7 (20.6 ⁺)					
EORTC	Disease Symptoms				32.4 (24.1) ⁺	ND	ND		ND	
QLQ-MY20	Side Effects				82.0 (15.3) ⁺	NR	NR	NR	NR	NR
	0	3 (75.0)	31 (44.3)	23 (42.6)	57 (44.5)	10/33 (30.3)	0 (0)#	21 (36.8)		
ECOG PS, n (%)	1	1 (25.0)	38 (54.3)	29 (53.7)	68 (53.1)	21/33 (63.6)	07 (100)#	27 (47.4)	17 (100)#	97 (100)#
11 (%)	2	0 (0)	1 (1.4)	2 (3.7)	3 (2.3)	2/33 (6.1)	97 (100)#	9 (15.8)	1	

Data not reported for the following baseline characteristics: Height, weight, age at diagnosis, D/C last line of therapy due to side effects

BCMA: B-cell maturation antigen, CAR+: chimeric antigen receptor positive, ECOG PS: Eastern Cooperative Oncology Group performance status, EORTC: European

Organization for Research and Treatment of Cancer, D/C: discontinued, kg: kilogram, mg: milligram, n: number, N: total number, NR: not reported, SCT: stem cell transplant, SD: standard deviation, QLQ: Quality of Life questionnaire

* Defined as ≥50% BCMA+

⁺ Data are digitized and should be interpreted with caution

‡ Progressive disease during most recent line of therapy

§ Defined by the presence of the following abnormalities: del(17p), t(4;14), or t(14;16)

Assumption made based on study protocol

× Administered after leukapheresis and before lymphodepletion

** N of 62 represents the entire treated CRB-401 population, N of 33 represents the first 33 patients to received Ide-cel treatment within the population

Table D3.3. Efficacy Outcomes: Ide-cel

	Trial		KarM	Ma ^{11,12}			CRB-401 ⁶⁷	
	Arms	150x10 ⁶ CAR+ T cells/kg	300 x10 ⁶ CAR+ T cells/kg	450 x10 ⁶ CAR+ T cells/kg	Overall	150 x10 ⁶ CAR+ T cells/kg	450 x10 ⁶ CAR+ T cells/kg	Overall
Γ	l (as-treated)	4	70	54	128	18	38	62#
Me	dian Follow-Up		13.3	Months		14.7 Months		
Median Time to Initial Response, Months (Range)		NR	NR	NR	1 (0.5-8.8)	NR	NR	1 (0.5-3.0); N=33
	Median Time to CR or better, Months (Range)	NR	NR	NR	2.8 (1.0-11.8)	NR	NR	NR
Response	Median Duration, Months (95%Cl)	NR (2.8, NE) [*]	9.9 (5.4 <i>,</i> 1.1.0)	11.3 (10.3, 11.4)	10.7 (9.0 <i>,</i> 11.3)	13.7 (2.9, 39.6)	10.0 (6.3, 14.8)	10.3 (7.7 <i>,</i> 13.7)
	ORR, n (%); [95%Cl]	2 (50.0)	48 (68.6)	44 (81.5)	94 (73.4); [65.8, 81.1]	9 (50.0); NR	34 (89.5)	47 (75.8); NR
	sCR/CR, n (%)	1 (25.0)	20 (28.6)	21 (38.9)	42 (32.8)	7 (38.9)	14 (36.8)	24 (38.7)
	vgPR, n (%)	1 (25.0)	10 (14.3)	14 (25.9)	25 (19.5)	7 (38.9)	30 (78.9)	40 (64.5)
	PR, n (%)	0 (0)	18 (25.7)	9 (16.7)	27 (21.1)	NR	NR	NR
	Median Duration, Months (95%Cl)				19.4 (18.2, NE)	NE (10.8, NE)	34.2 (23.2, NE)	34.2 (19.2 <i>,</i> NE)
Overall	OS at 3 Months, n (%)				NR (95.2) ⁺	NR	NR	NR
Survival	OS at 6 Months, n (%)	NR	NR	NR	NR (90.0) ⁺	NR	NR	NR
	OS at 9 Months, n (%)				NR (84.1) ⁺	NR	NR	NR
	OS at 12 Months, n (%)				NR (78.0)	NR	NR	NR
	Median Duration, Months (95%Cl)	2.8 (1.0, NE)	5.8 (4.2, 8.9)	12.1 (8.8, 12.3)	8.8 (5.6, 11.6)	4.5 (2.0, 12.0)	9 (7.2, 12.2)	8.8 (5.9, 11.9)
Progression	PFS at 3 Months, n (%)	NR (50.6) [†]	NR $(69.1)^{+}$	NR (80.9) ⁺	NR (72.0) ⁺	NR	NR	NR
-Free Survival	PFS at 6 Months, n (%)	NR (24.0) ⁺	NR (48.7) ⁺	NR (69.3) ⁺	NR (56.7) [†]	NR	NR	NR
Suivival	PFS at 9 Months, n (%)	NR (24.0) [†]	NR (36.4) ⁺	NR (60.3) [†]	NR (45.2) [†]	NR	NR	NR
	PFS at 12 Months, n (%)	NR (24.0) ⁺	NR (27.4) ⁺	NR (49.3) ⁺	NR (37.9) ⁺	NR	NR	NR
MRD-neg	MRD-negativity, n (%); [95%CI]		17 [§] (24.0); [14.8, 36.0]	15 [§] (28.0); [16.5, 41.6]	33 [§] (26.0); [18.5, 34.3]	NR	NR	30/37 [‡] (81.0)

	Trial		KarN	1Ma ^{11,12}		CRB-401 ⁶⁷			
Arms		150x10 ⁶ CAR+ T cells/kg	300 x10 ⁶ CAR+ T cells/kg	450 x10 ⁶ CAR+ T cells/kg	Overall	150 x10 ⁶ CAR+ T cells/kg	450 x10 ⁶ CAR+ T cells/kg	Overall	
Median peak CAR-T cell Expansion, Days (SE)		14 (NR)	11 (NR)	11 (NR)	11 (NR)	NR	NR	NR	
CAR-T cells	At 6 Months, n (%)	ND		NR	29/49 (59.2)	- NR	NR	13/23 (57.0)	
detectable	At 12 Months, n (%)	NR	NR		4/11 (36.4)			2/10 (20.0)	
Disease Progression, n (%)		NR	NR	NR	8 (6.3)	NR	NR	36/62 (58.0)	
Retreatment [¤] , n (%)		NR	NR	NR	28 (21.9)	NR	NR	NR	

95% CI: 95% confidence interval, CAR: chimeric antigen receptor, CAR+: chimeric antigen receptor positive, CR: complete response, kg: kilogram, MRD: minimal residual disease, n: number, N: total number, NE: not estimable, NR: not reported, ORR: overall response rate, PFS: progression-free survival, PR: partial response, sCR: stringent complete response, SE: standard error, vgPR: very good partial response

* Due to small N

⁺ Data are digitized and should be interpreted with caution

‡ Evaluable patients

§ MRD-negativity assessed in patients with complete response or better

N of 62 represents the entire treated CRB-401 population, N of 33 represents the first 33 patients to received Ide-cel treatment within the population

× Patients were retreated after disease progression

Table D3.4. Efficacy Outcomes: Cilta-cel

	Trial	CARTITUDE-1 ^{15,19}	LEGEND-2 ^{65,66} Xi'an	LEGEND-2 ^{65,72} Changzheng, Ruijin, Jiangsu	
	Arms	Overall	0.5x10 ⁶ CAR+ T cells/kg	0.7x10 ⁶ CAR+ T cells/kg	
Ν	N (as-treated)	97	57	17	
Me	dian Follow Up	12.4 Months	25 Months	26 Months	
	Median Time to Initial Response, Months (Range)	1 (0.9-8.5)	1.0 (0.4-3.5)	1.0 (NR)	
	Median Duration, Months (95%Cl)	Not reached [‡]	27 (NR)	NR	
Response	ORR, n (%); [95%Cl]	94 (96.9)	50 (87.7); [76.0, 95.0]	15 (88.2); [64.0, 99.0]	
	sCR, n (%)	65 (67.0)	NR	NR	
	CR, n (%)	0 (0)	42 (73.7)	14 (82.4)	
	vgPR, n (%)	25 (25.8)	2 (3.5)	1 (5.9)	
	PR, n (%)	4 (4.1)	6 (10.5)	NR	
	Median Duration, Months (95%Cl)	Not reached [‡]	36.1 (26.4, NE)	Not reached [¤]	
	At 3 months, n (%)	NR	NR (97.8) ⁺	NR (89.1) [†]	
Overall Survival	At 6 months, n (%)	NR (93.8) [‡]	NR (92.4) ⁺	NR (89.1) ⁺	
-	At 9 months, n (%)	NR	NR (81.5) ⁺	NR (89.1) ⁺	
-	At 12 months, n (%)	NR (88.5)	NR (78.3) ⁺	NR (82.3)	
	Median Duration, Months (95%Cl)	Not reached	19.9 (9.6, 31.0)	18.0 (NR)	
Progression-	At 3 months, n (%)	NR (98.0) ⁺	NR (86.4) ⁺	NR (95.0) ⁺	
Free Survival	At 6 months, n (%)	NR (87.5) ⁺	NR (81.2) ⁺	NR (83.4) ⁺	
	At 9 months, n (%)	NR (80.3) ⁺	NR (65.3) ⁺	NR (75.6) ⁺	
	At 12 months, n (%)	n (76.6)	NR (60.1) ⁺	5/9 (57.0)	
MRD	-negativity, n (%)	53# (54.6)	39§ (68.4) [¤]	NR	
Median Peak (CAR-T Cell Expansion, Days (Range)	13 (9-55)	NR	NR	
Disease	e Progression, n (%)	NR	18/50 ⁺⁺ (36.0) [¤]	11 (64.7)	

Trial	CARTITUDE-1 ^{15,19}	LEGEND-2 ^{65,66}	LEGEND-2 ^{65,72}	
Iriai	CARTIODE-1	Xi'an	Changzheng, Ruijin, Jiangsu	
Arms	Overall	0.5x10 ⁶ CAR+ T cells/kg	0.7x10 ⁶ CAR+ T cells/kg	

Data not reported for the following efficacy outcomes: Median time to complete response, CAR-T cells detectable at 6 and 12 months

95% CI: 95% confidence interval, CAR: chimeric antigen receptor, CAR+: chimeric antigen receptor positive, CR: complete response, kg: kilogram, MRD: minimal residual disease, n: number, N: total number, N/A: not available, NE: not estimable, NR: not reported, ORR: overall response rate, PFS: progression-free survival, PR: partial response, sCR: stringent complete response, SE: standard error, vgPR: very good partial response

* 95.7% confidence interval

⁺ Data are digitized and should be interpreted with caution

‡ 8.8-month follow up time

§ MRD-negativity assessed in patients with complete response

MRD-negativity assessed among evaluable patients

¤ 12-month follow up time

++ Patients with partial response or better

Table D3.5. Efficacy Outcomes: Belantamab

	Trial	DREAMM-2 ²²
	Arms	2.5 mg/kg
	N (ITT)	97
Medi	an Follow Up	13 Months
	Median Duration, Months (95%CI)	11.0 (4.2, not reached)
	ORR, n (%); [95%Cl]	31 (32.0); [21.7, 43.6]*
Response	sCR, n (%)	2 (2.1)
Response	CR, n (%)	5 (5.2)
	vgPR, n (%)	11 (11.3)
	PR, n (%)	13 (13.4)
	Median Duration, Months (95%CI)	13.7 (9.9, not reached)
	At 3 Months, n (%)	63/77 (82.2) [†]
Overall Survival	At 6 Months, n (%)	48/66 (72.6) [†]
	At 9 Months, n (%)	37/66 (63.0) [†]
	At 12 Months, n (%)	28/49 (56.9) [†]
	Median Duration, Months (95%CI)	2.8 (1.6, 3.6)
	At 3 Months, n (%)	
Progression-Free Survival	At 6 Months, n (%)	
	At 9 Months, n (%)	
	At 12 Months, n (%)	
Disease P	rogression, n (%)	59 (60.8) [‡]

Data not reported for the following efficacy outcomes: Median time to complete response, CAR-T cells detectable at 6 and 12 months, median time to initial response, MRD-negativity

95% CI: 95% confidence interval, ITT: Intention to treat, CR: complete response, kg: kilogram, mg: milligram, MRD: minimal residual disease, n: number, N: total number, N/A: not available, NE: not estimable, NR: not reported, ORR: overall response rate, PFS: progression-free survival, PR: partial response, sCR: stringent complete response, SE: standard error, vgPR: very good partial response

* 95.7% confidence interval

⁺ Data are digitized and should be interpreted with caution

‡ 6.3-month follow-up time

Table D3.6. Subgroup Efficacy Data: Ide-cel

Trial		KarMMa ^{12,31,32}									
				Age Groups							
Subgroups	Extramedullary Disease	Received Bridging Therapy	High Cytogenetic Risk	High Tumor Burden	ISS Disease Stage III	>1 Treatment Regimen per Year	< 65 years	≥ 65 years	≥ 70 years		
Median Follow-		11.3 Months									
Up											
N (as-treated)	50	112	45	65	21	60	83	45	20		
ORR, n (%),	35 (70.0);	80 (71.0);	31 (69.0);	46 (71.0);	10 (48.0);	39 (65.0);	56 (67.5);	38 (84.4);	18 (90.0);		
[95%CI]	[55.4 <i>,</i> 82.1] [*]	[62.1, 79.6]	[55.4, 82.4]	[58.2 <i>,</i> 81.4] [*]	[25.7, 70.2]	[51.6, 76.9]	[56.2 <i>,</i> 77.1] [†]	[70.5, 93.5]	[76.9, 100]		
CR, n (%);	12 (24.0);	38 (34.0);	14 (31.0);	19 (29.0);	2 (10.0);	18 (30.0);	28 (33.4);	14 (31.1);	7 (35.0);		
[95%CI]	[13.1, 38.2]	[25.3 <i>,</i> 43.5]	[17.6 <i>,</i> 44.6]	[18.6, 41.8]	[1.2, 30.4]	[18.8, 43.2]	[23.7 <i>,</i> 44.3] [†]	[18.2 <i>,</i> 46.6]	[14.1, 55.9]		
Median DOR,	9.2	10.9	10.7	10.4	6.9	10.5	ND	10.9	11.0		
Months [95%CI]	[5.4, 11.3]	[9.0, 11.4]	[6.5 <i>,</i> NE]	[6.1, 11.3]	[1.9, 10.3]	[9.0, 11.3]	NR	[4.5, 11.4]	[3.9, 11.4]		
Median PFS,	7.9	8.8	8.2	7.5	4.9	8.9	ND	8.6	10.2		
Months [95%CI]	[5.1, 10.9]	[5.5, 11.6]	[4.8, 11.9]	[4.9, 11.3]	[1.8 <i>,</i> 8.2]	[3.1, 11.1]	NR	[4.9, 12.2]	[3.1, 12.3]		
Subgroup data not	reported for the fo	ollowing: probat	oility of DOR ≥6	months, probabi	lity of PFS at 6 r	nonths, median (OS, OS at 12 mo	nths; vgPR, PR			

95% CI: 95% confidence interval, CR: complete response, DOR: duration of response, n: number, N: total number, ORR: overall response rate, OS: overall response, PFS: progression-free survival, PR: partial response, vgPR: very good partial response.

* Did not significantly differ between patients with versus patients without risk factors

⁺ Data are digitized and should be interpreted with caution

Table D3.7. Subgroup Efficacy Data: Ide-cel II

Trial		CRB-40	1 ⁵⁹	
	Extramedullary Disease	Received Bridging Therapy	High Cytogenetic Risk	High Tumor Burden
Median Follow-Up		11.3 Mor	nths	
N (as-treated)	9	14	15	16
ORR, n (%); [95% Cl]	8 (88.9); [51.8, 99.7]	14 (100); [76.8, 100]	11 (73.3); [44.9, 92.2)	12 (75.0); [47.6, 92.7]
CR, n (%)	1 (11.1)	2 (14.3)	1 (6.7)	0 (0)
vgPR, n (%)	2 (22.2)	4 (28.6)	3 (20.0)	4 (25.0)
PR, n (%)	2 (22.2)	1 (7.1)	1 (6.7)	1 (6.3)

95% CI: 95% confidence interval, BCMA: B-cell maturation antigen, CR: complete response, DOR: duration of response, EMD: extramedullary disease, n: number, N: total number, NE: not estimable, NR: not reported, ORR: overall response rate, OS: overall response, PFS: progression-free survival, PR: partial response, vgPR: very good partial response

Table D3.8. Subgroup Efficacy Data: Cilta-cel

Trial			: ND-2^{65,68} Xi'an	
	BCMA Expression	High Risk	Complete R	esponse
Subgroups	≥40%	EMD	Achieved	Not Achieved
Median Follow-Up	8 months		25 months	
N (as-treated)	27/53	17	42	15
ORR, n (%); [95% Cl]	22 (81.5); [NR]	14 (82.4); [NR]	NR	NR
CR, n (%)	17 (63.0)	10 (60.0)*	NR	NR
vgPR, n (%)	1 (3.7)	2 (11.6)*	NR	NR
PR, n (%)	4 (14.8)	2 (11.6)*	NR	NR
Median DOR, Months [95%CI]	NR	NR	29.1 [NR]	NR
Median PFS, Months [95%CI]	11 [6-NE]	8.1 [NR]	28.2 [19.9, NE]	3.2 [1.7, 6.4]
Median OS, Months [95%CI]	Not reached	13.9 [NR]	Not reached [35.0, NE]	7.5 [3.8, 13.1]
OS at 12 Months, n (%)	NR	NR	NR (92.9)	NR

(Changzheng, Ruijin, Jiangsu sites)

95% CI: 95% confidence interval, BCMA: B-cell maturation antigen, CR: complete response, DOR: duration of response, EMD: extramedullary disease, n: number, N: total number, NE: not estimable, NR: not reported, ORR: overall response rate, OS: overall response, PFS: progression-free survival, PR: partial response, VGPR: very good partial response

* Data are digitized and should be interpreted with caution

Table D3.9. Subgroup Efficacy Data: Belantamab

Trial				DRE	AMM-2 ^{17,33-35}				
	Prior Th	erapies	High Risk	Renal Imp	airment†	A	ge	Race / Ethnicity	
Subgroups	3-6 Therapies	≥ 7 Therapies	High Risk Cytogenetics	Mild	Moderate	65 to < 75 years	≥75 years	White	Black
Median Follow-Up	12.4 N	lonths	9 Months	9 Mc	onths	6.3 Months		6.3 M	onths
N (ITT)	47	50	41	48	24	39	13	76	16
ORR, n (%); [95%Cl]	16 (34.0); [19.3, 51.4]*	15 (30.0); [16.5, 46.6]*	11 (27.0); [14.2, 42.9]*	15 (31.3); [18.7, 46.3]	8 (33.3); [15.6, 55.3]	17 (43.6); [27.8, 60.4]	1 (7.7); [0.2, 36.0]	24 (31.6); [21.4, 43.3]‡	6 (37.5); [15.2 <i>,</i> 64.6]§
vgPR, n (%)	8 (17.0)	10 (20.0)	9 (22.0)	NR	NR				
Median DOR, Months [95%CI]	11.0 [4.2, Not reached]	13.1 [4.0, Not reached]	Not reached [1.4, Not reached]	Not reached	Not reached				
Probability DOR ≥6 Months, % [95%CI]	63 [31, 83]	73 [44, 89]	52 [20, 77]	NR	NR				
Median PFS, Months [95%CI]	2.9 [1.5, 5.7]	2.2 [1.2, 3.6]	2.1 [0.8, 3.7]	2.2 [2.1, 3.6]	3.7 [1.0, Not reached]	N	R	Ν	R
Probability of 6-Month PFS, % [95%CI]	35 [20, 50]	30 [17, 43]	30 [16, 45]	NR	NR				
Median OS, Months [95%CI]	NR	NR	9.4 [4.3, 13.1]	NR	NR				
Probability of 12- Month OS, % [95% CI]	NR	NR	45 [27, 61]	NR	NR				
Subgroup data not report	ed for the follow	ving outcomes: C	CR, PR						

95% CI: 95% confidence interval, ITT: Intention to treat, CR: complete response, DOR: duration of response, GFR: glomerular filtration rate, m²: meters squared, min: minute, mL: milliliter, n: number, N: total number, NR: not reported, ORR: overall response rate, OS: overall response, PFS: progression-free survival, PR: partial response, VGPR: very good partial response

* 97.5% confidence interval

+ Mild renal impairment defined as GFR ≥60-<90 mL/min/1.73 m², moderate renal impairment defined as GFR ≥30-<60 mL/min/1.73 m²

‡ ORR at 13 months of follow-up was received as academic in confidence (

§ ORR at 13 months of follow-up was received as academic in confidence (

	Intervent	ion	Ide-cel (a	as-treated)	Cilta-cel (a	as-treated)		Belan	tamab (ITT)			
	Trial		KarN	MMa ²⁰	CARTIT	UDE-1 ²¹		DRE	AMM-2 ^{23,24}			
	Arms (N	J)	Overal	(N=128)	Overal	l (N=68)	2.5 mg/kg (N=97)					
	Follow-U	Ъ	Day 1	Month 9	Day 100 Day 184 Week 7 Week 13 Week 19 N					Week 25		
		Ν	111	59								
EQ-5D- 5L	Impr	ovement	33 (29.7)	32 (54.2)		IR	ND					
Index	No	Change	57 (51.4)	22 (37.3)				NR				
	Dete	erioration	49 (44.1)	5 (8.5)								
		Ν	111	59								
EQ VAS	Impr	ovement	46 (41.4)	48 (81.3)		ID			NR			
EQ VAS	No	Change	16 (14.4)	1 (1.8)	NR		I INK					
	Dete	rioration	49 (44.1)	10 (16.9)								
		Ν	110	59	47	30	46	29	19	19		
	Physical Functioning	Change from BL, Mean Score (95%Cl)	-1.2 (-4.3, 2.0)†	13.2 (7.9, 17.9)†	NR	NR	5.1 (0.5, 9.9)†	3.0 (-2.8, 8.7)†	0.3 (-6.3, 6.9)†	-0.1 (-5.3, 5.3)†		
		Improvement‡	40 (36.4)	35 (59.3)	34 (72.1)	NR	13 (28.3)	8 (27.6)	3 (15.8)	4 (21.1)		
		No Change	31 (28.2)	19 (32.2)	NR	NR	NR	NR	NR	NR		
		Deterioration	39 (35.5)	5 (8.5)	NR	NR	NR	NR	NR	NR		
EORTC QLQ-	Cognitive	Change from BL, Mean Score (95%Cl)	0.8 (-3.1, 3.4)†	6.6 (2.7, 11.2)†								
C30	Functioning	Improvement‡	22 (20.0)	42 (71.2)	N	IR			NR			
	_	No Change	38 (34.5)	10 (16.9)								
	Role	Deterioration	50 (45.5)	7 (11.9)								
		Change from BL, Mean Score (95%CI)	NR	NR			1.9 (-8.1, 11.9)†	7.3 (-5.7, 20.4)†	-4.4 (-19.3, 10.7)†	8.0 (-4.9, 20.7)†		
	Functioning	Improvement‡	23 (20.9)	34 (57.6)	N	IR	NR	NR	NR	NR		
		No Change	49 (44.5)	19 (32.2)			NR	NR	NR	NR		
		Deterioration	38 (34.5)	6 (10.2)			NR	NR	NR	NR		

Table D3.10. Patient Reported Outcomes: Ide-cel, Cilta-cel, Belantamab

Interve	tion	Ide-cel (as-treated)	Cilta-cel (as-treated)		Belar	ntamab (ITT)	
Tria	l	Karl	MMa ²⁰	CARTIT	UDE-1 ²¹		DRE	AMM-2 ^{23,24}	
Arms	N)	Overal	l (N=128)	Overal	ll (N=68)		2.5 m	ng/kg (N=97)	
Follow	-Up	Day 1	Month 9	Day 100	Day 184	Week 7	Week 13	Week 19	Week 25
Fur etternel	Improvement‡	32 (29.1)	30 (50.8)						
Emotional Functioning	No Change	34 (30.9)	17 (28.8)	NR				NR	
T unctioning	Deterioration	44 (40.0)	12 (20.3)						
Casial	Improvement‡	25 (22.7)	36 (61.0)						
Social Functioning	No Change	42 (38.2)	16 (27.1)	1	NR			NR	
Tunctioning	Deterioration	43 (39.1)	7 (11.9)						
	Change from BL#, mean score (95%Cl)	7.7 (3.7, 11.4)†	-22.8 (-29.1, -17.1)†	-1.5 (-9.1, 5.1)†	-9.2 (-16.4, -2.0)†	-3.9 (-11.1, 2.5)†	-7.3 (-16.6, 1.8)†	-0.8 (-11.2, 10.6)†	3.6 (-7.6, 14.6)†
Fatigue	Improvement‡	28 (25.5)	39 (66.1)	NR (62.2)	NR	21 (45.7)	12 (41.4)	6 (31.6)	6 (31.6)
	No Change	32 (28.8)	17 (28.8)	NR	NR	NR	NR	NR	NR
	Deterioration	51 (46.4)	3 (5.1)	NR	NR	NR	NR	NR	NR
	Change from BL#, mean score (95%Cl)	-8.7 (-13.0, 5.6)†	-23.8 (-30.2, -18.3)†	-8.9 (-16.6, NR)†	-8.9, (-17.6, -0.3)†	-4.7 (-13.0, 3.5)†	-4.4 (-14.4, 6.0)†	4.9 (-4.9, 14.2)†	2.6 (-6.5, 11.4)†
Pain	Improvement‡	38 (34.5)	36 (61.0)	33 (71.1)	NR	14 (30.4)	9 (31.0)	4 (21.1)	3 (15.8)
	No Change	48 (43.6)	17 (28.8)	NR	NR	NR	NR	NR	NR
	Deterioration	24 (21.8)	6 (10.2)	NR	NR	NR	NR	NR	NR
	Improvement‡	7 (6.4)	15 (25.4)						
Nausea/ Vomiting	No Change	41 (37.3)	38 (64.4)	1	NR			NR	
	Deterioration	62 (56.4)	6 (10.2)						
	Improvement‡	9 (8.2)	11 (18.6)						
Constipation	No Change	60 (54.5)	45 (76.3)	1	NR			NR	
	Deterioration	41 (37.3)	3 (5.1)						
	Improvement‡	17 (15.5)	13 (22.0)						
Diarrhea	No Change	76 (69.1)	44 (74.6)	1	NR			NR	
	Deterioration	17 (15.5)	2 (3.4)						
Insomnia	Improvement‡	20 (18.2)	24 (40.7)	1	NR			NR	

	Intervent	ion	Ide-cel (a	as-treated)	Cilta-cel (as-treated)		Belan	tamab (ITT)	
	Trial		Karl	MMa ²⁰	CARTIT	UDE-1 ²¹		DRE	AMM-2 ^{23,24}	
	Arms (N	1)	Overal	l (N=128)	Overal	l (N=68)		2.5 m	g/kg (N=97)	
	Follow-U	Jp	Day 1	Month 9	Day 100	Day 184	Week 7	Week 13	Week 19	Week 25
		No Change	73 (66.4)	29 (49.2)						
		Deterioration	17 (15.5)	6 (10.2)						
		Improvement‡	24 (21.8)	17 (28.8)						
	Dyspnea	No Change	77 (70.0)	37 (62.7)	1	NR			NR	
		Deterioration	9 (8.2)	5 (8.5)						
		Improvement‡	7 (6.4)	18 (30.5)						
	Appetite Loss	No Change	47 (42.7)	35 (59.3)	1	NR			NR	
	LUSS	Deterioration	56 (50.9)	6 (10.2)						
		Improvement‡	11 (10.0)	10 (16.9)						
	Financial	No Change	79 (71.8)	40 (67.8)	٦	NR			NR	
	Difficulties	Deterioration	20 (18.2)	9 (15.3)						
	Global	Change from BL, Mean Score (95%Cl)	-5.1 (-2.5, -7.7)†	15.4 (9.8, 20.9)†	NR	NR	0.4 (-5.7, 6.4)†	-3.2 (-10.0, 3.8)†	-2.3 (-8.9, 4.5)†	-4.7 (-12.1, 2.8)†
	Health / QoL	Improvement‡	23 (20.9)	24 (40.7)	24 (51.1)	NR	NR	NR	NR	NR
		No Change	64 (58.2)	29 (49.2)	NR	NR	NR	NR	NR	NR
		Deterioration	23 (20.9)	6 (10.2)	NR	NR	NR	NR	NR	NR
		N	109	57	47	NR	45	28	18	18
	Disease	Change from BL, Mean Score (95%Cl)	-1.14 (-3.4, 1.7)†	-20.00 (-14.3, -9.7)†	NR	NR	-2.5 (-8.1, 3.8)†	-0.8 (-7.2, 5.5)†	1.1. (NR, 8.5)†	0.0 (-7.5, 7.5)†
EORTC	Symptoms	Improvement‡	15 (13.8)	25 (43.9)	NR	NR	17 (37.8)	8 (28.6)	5 (27.8)	6 (33.3)
QLQ-		No Change	72 (66.1)	28 (49.1)	NR	NR	NR	NR	NR	NR
MY20		Deterioration	22 (20.2)	4 (7.0)	NR	NR	NR	NR	NR	NR
	Future	Improvement‡	32 (29.4)	34 (59.6*)			0 (0)	0 (0)	0 (0)	0 (0)
	Future Perspectives	No Change	41 (37.6)	14 (24.6)	١	NR	NR	NR	NR	NR
	i erspectives	Deterioration	36 (33.0)	9 (15.8)			NR	NR	NR	NR
	Body Image	Improvement‡	13 (11.9)	17 (29.8)	1	NR			NR	

	Intervent	ion	Ide-cel (as-treated)	Cilta-cel (a	as-treated)		Belar	itamab (ITT)	
	Trial		Karl	MMa ²⁰	CARTIT	UDE-1 ²¹		DRE	AMM-2 ^{23,24}	
	Arms (N	1)	Overal	l (N=128)	Overall (N=68)		2.5 mg/kg (N=97)			
	Follow-U	Jp	Day 1	Month 9	Day 100	Day 184	Week 7 Week 13 Week 19 V		Week 25	
		No Change	79 (72.5)	36 (63.2)						
		Deterioration	17 (15.6)	4 (7.0)						
		Change from BL, Mean Score (95%CI)	-0.57 (-5.14, - 2.86)†	6.29 (3.43, 9.14)†						
	Side Effects	Improvement‡	35 (32.1)	3 (5.3)	N	IR			NR	
		No Change	62 (56.9)	37 (64.9)						
		Deterioration	12 (11.0)	17 (29.8)						
		N		NR	N	IR	92		NR	
OSDI	Vision- related	Change from BL, Mean Score (95%Cl) Deterioration					NR 46 (49.5)§			
	Functioning Domain	from Baseline Improvement from Worst Severity post-BL		NR	N	IR	33 (72.0)		NR	
		No Change					NR			

Patient reported outcomes not reported for the following trails: CRB-401, LEGEND-2

95% CI: 95% confidence interval, ITT: Intention to treat, EORTC: European Organization for Research and Treatment of Cancer, EQ-5D-5L: EuroQol 5 dimensions 5 levels, kg: kilogram, mg: milligram, MY20: Myeloma Module questionnaire, n: number, N: total number, NR: not reported, OSDI: Ocular Surface Disease Index, QoL: quality of life, QLQ-C30: Quality of Life C30 questionnaire, VAS: visual analog scale

* Statistically significant improvement, p<0.05

⁺ Data are digitized and should be interpreted with caution

‡ Change of ≥10 points

§ Change of ≥12.5-points

Negative changes indicate a reduction in pain or fatigue

Table D3.11. Safety Outcomes: Ide-cel

	Trial		KarM	Ma ^{11,12,27}			CRB-401 ^{59,67}	7
	Arms	150x10 ⁶ CAR+ T cells/kg	300 x10 ⁶ CAR+ T cells/kg	450 x10 ⁶ CAR+ T cells/kg	Overall	150 x10 ⁶ CAR+ T cells/kg	450 x10 ⁶ CAR+ T cells/kg	Overall
Γ	l (as-treated)	4	70	54	128	18	38	62†
	Overall				128 (100)			33/33 (100)
Any AEs n (%)	Grade 3/4		NR		127 (99.2)	NF	ł	32/33 (96.7)
Treatme	nt-related SAEs, n (%)				NR			NR
	Overall	2 (50.0)	27 (38.6)	15 (27.8)	44 (34.4)			6 (10.0)
Mortality, n (%) [#]	Disease Progression	2 (50.0)	18 (25.7)	7 (13.0)	27 (21.1)			NR
wortality, n (%)"	AEs	0 (0)	5 (7.1)	4 (7.4)	9 (7.0)	NR		NR
	Other	0 (0)	3 (4.3)	4 (7.4)	7 (5.5)			NR
	ICU [§]				19/107 (17.8)			
Hospitalizations, n (%)	Mean Length of Stay, Days (Range)		NR		NR (6-30)	NF	NR	
	Overall				12 (9.4)			3/36 (8.3)
Study	Death				2 (1.6)			0/36 (0)
Discontinuation	Disease Progression		NR		1 (0.8)	NF)	3/36 (8.3)
prior to Treatment,	Adverse Event		INIT		1 (0.8)	INF	N	0/36 (0)
n (%)	Patient Withdrawal				4 (3.1)			0/36 (0)
	Manufacturing Failure				1 (0.8)			0/36 (0)
	Overall				66 (51.6) [‡]			49 (79.0)
Discontinuation,	Death		NR		41 (32.0)	NF		6 (10.0)
n (%)	Disease Progression		NK		23 (18.0)	INF	i	36 (58.0)
	Patient Withdrawal				2 (1.6)			6 (10.0)
	Median Onset, Days (Range)	7 (2-12)	2 (1-12)	1 (1-10)	1 (1-12)	NR	NR	2 (1-25); N=33
	Median Duration, Days (Range)	5 (3-7)	4 (2-28)	7 (1-63)	5 (1-63)	NR	NR	5 (1-32); N=33
CRS, n (%)	Overall	2 (50.0)	53 (75.7)	52 (96.3)	107 (83.6)	7 (11.3)	35 (92.1)	47 (76.0)
	Grade 1	1 (25.0)	33 (47.1)	27 (50.0)	61 (47.7)	NR	NR	43 (69.3)

	Trial		KarM	Ma ^{11,12,27}			CRB-401 ^{59,67}	7
	Arms	150x10 ⁶ CAR+ T cells/kg	300 x10 ⁶ CAR+ T cells/kg	450 x10 ⁶ CAR+ T cells/kg	Overall	150 x10 ⁶ CAR+ T cells/kg	450 x10 ⁶ CAR+ T cells/kg	Overall
	Grade 2	1 (25.0)	16 (22.9)	22 (40.7)	39 (30.5)	NR	NR	
	Grade 3	0 (0)	2 (2.9)	3 (4.3)	5 (3.9)	0 (0)	3 (7.9)	4 (6.5)
	Grade 4	0 (0)	1 (1.4)	0 (0)	1 (0.8)	0 (0)	0 (0)	0 (0)
	Grade 5	0 (0)	1 (1.4)	0 (0)	1 (0.8)	0 (0)	0 (0)	0 (0)
	Progression from Grade 1 to ≥2	NR	NR	NR	30 (65.2)*	NR	NR	NR
Required	Overall	1 (25.0)	30 (42.8)	36 (66.7)	67 (52.3)			7/33 (21.2)
Tocilizumab for	1 Dose	1 (25.0)	21 (30.0)	22 (40.7)	44 (34.3)	NI	२	NR
CRS <i>,</i> n (%)	≥1 Dose	0 (0)	9 (12.9)	14 (25.9)	23 (18.0)			NR
	Median Onset, Days (Range)	N/A	3 (1-10)	2 (1-5)	2 (1–10)	NR	NR	NR
	Median Duration, Days (Range)	N/A	3 (2–26)	5 (1-22)	3 (1–26)	NR NR		NR
-	Overall	0 (0)	12 (17.1)	11 (20.4)	23 (18.0)	5 (27.8)	20 (52.6)	27 (43.5)
Neurotoxicity, n	Grade 1	0 (0)	7 (10.0)	5 (9.3)	12 (9.4)	NR	NR	25 (20.4)
(%)	Grade 2	0 (0)	4 (5.7)	3 (5.6)	7 (5.5)	NR	NR	25 (39.1)
	Grade 3	0 (0)	1 (1.4)	3 (5.6)	4 (3.1)	0 (0)	2 (5.3)	1 (1.6)
	Grade 4	0 (0)	0 (0)	0 (0)	0 (0)	NR	NR	1 (1.6)
	Grade 5	0 (0)	0 (0)	0(0)	0 (0)	0 (0)	0 (0)	0 (0)
$C_{\rm c}$ to make m (0/)	Overall	4 (100)	67 (95.7)	53 (98.1)	124 (97.0)			NR
Cytopenia, n (%)	≥Grade 3	NR	NR	NR	NR			NR
	Overall	4 (100)	66 (94.3)	51 (94.4)	117 (91.4)			57 (92.0)
Neutropenia, n (%)	≥Grade 3	NR	NR	NR	114 (89.1)			55 (89.0)
Anomio n (9/)	Overall	4 (100)	51 (72.9)	34 (63.0)	89 (69.5)		`	47 (76.0)
Anemia, n (%)	≥Grade 3	NR	NR	NR	77 (60.2)	NR		35 (57.0)
Thrombocytopenia,	Overall	4 (100)	45 (64.3)	35 (64.8)	81 (63.3)			46 (74.0)
n (%)	≥Grade 3	NR	NR	NR	67 (52.3)		35 (57.0)	
Infactions = (0/)	Overall	3 (75.0)	47 (67.1)	38 (70.4)	88 (68.8)			14/33 (42.4
Infections, n (%)	≥Grade 3	NR	NR	NR	28 (21.9)			2/33 (6.1)

Trial		KarM	Ma ^{11,12,27}			CRB-401 ^{59,67}	
Arms	150x10 ⁶ CAR+ T	300 x10 ⁶ CAR+ T	450 x10 ⁶ CAR+ T	Overall	150 x10 ⁶ CAR+ T	450 x10 ⁶ CAR+ T	Overall
	cells/kg	cells/kg	cells/kg		cells/kg	cells/kg	

Data not reported for the following safety outcomes: Any SAEs, treatment-related AEs, discontinuation due to AE and lack of efficacy, bone disease, hypercalcemia, hyper viscosity, renal failure, thrombosis, corneal events

AE: adverse event, CAR+: chimeric antigen receptor positive, CRS: cytokine release syndrome, ICU: intensive care unit, kg: kilogram, mg: milligram, n: number, N: total number, N/A: not applicable, NR: not reported, SAE: serious adverse event.

* Patients categorized as having Grade ≥2 CRS

⁺ N of 62 represents the entire treated CRB-401 population, N of 33 represents the first 33 patients to received Ide-cel treatment within the population

‡ includes 18 patients who were retreated

Up to 24 months of follow-up for KarMMa

§ due to CRS with or without neurologic event

Table D3.12. Safety Outcomes: Cilta-cel

	Trial	CARTITUDE-1 ¹⁹	LEGEND-2^{61,65,66} Xi'an	LEGEND-2 ^{14,65,72} Changzheng, Ruijin, Jiangsu
	Arms	0.71x10 ⁶ CAR+ T cells/kg	0.5x10 ⁶ CAR+ T cells/kg	0.7x10 ⁶ CAR+ T cells/kg
N (a	s-treated)	97	57	17
Any	AE, n (%)	97 (100)	57 (100)	
Treatment	related AE, n (%)	NR	NR	NR
Any	SAEs, n (%)	NR	37 (64.9)*	NR
Treatment-r	elated SAEs, n (%)	NR	NR	
	Overall	14 (14.4)	17 (29.8)	
Mantality a (0/)	Disease Progression	5 (5.2)	14 (24.6)	
Mortality, n (%)	AEs	9 (9.3)	2 (3.5)	NR
	Other	NR	1 (1.8)	
	Overall	16/113 (14.2)		
Study	Death	9/113 (8.0)		
Discontinuation Prior to Treatment,	Disease Progression	2/113 (1.8)	NR	NR
n (%)	Patient Withdrawal	5/113 (4.4)		
(/	Manufacturing Failure	0 (0)		
	Overall	14 (14.4)	NR	
	Death	14 (14.4)	17 (29.8)	
Discontinuation, n (%)	Disease Progression	NR	NR	NR
(70)	AEs	NR	NR	
	Lack of Efficacy	NR	NR	
	Median Onset, Days (Range)	7 (1-12)	9 (1-19)	NR
	Median Duration, Days (Range)	4 (1-97)	9 (3-57)	NR
CRS, n (%)	Overall	92 (94.8)	51 (89.5)	17 (100)
	Grade 1	49 (50.5)	27 (47.4)	10 (58.8)
	Grade 2	38 (39.2)	20 (35.1)	10 (58.8)
	Grade 3	3 (3.1)	4 (7.0)	6 (35.3)

	Trial	CARTITUDE-1 ¹⁹	LEGEND-2^{61,65,66} Xi'an	LEGEND-2 ^{14,65,72} Changzheng, Ruijin, Jiangsu
	Arms	0.71x10 ⁶ CAR+ T cells/kg	0.5x10 ⁶ CAR+ T cells/kg	0.7x10 ⁶ CAR+ T cells/kg
	Grade 4	1 (1.0)	0 (0)	0 (0)
	Grade 5	1 (1.0)	0 (0)	1 (5.9)
	Progression from Grade 1 to ≥ 2	NR	NR	NR
	Required tocilizumab, n (%)	67 (69.1)	26 (45.6)†	9 (52.9)
	Overall	20 (20.6)	1 (1.8)	
	Grade 1	10 (10 2)	1 (1 0)	
Normatoriaitor a (0/)	Grade 2	10 (10.3)	1 (1.8)	0 (0)
Neurotoxicity, n (%)	Grade 3	0 (0 2)		0 (0)
	Grade 4	9 (9.3)	0 (0)	
	Grade 5	1 (1.0)		
Outononia n (%)	Overall	NR	NR	14 (82.4)
Cytopenia, n (%)	≥Grade 3	NR	NR	10 (58.8)
Noutrononia n (9/)	Overall	93 (95.9)	NR	NR
Neutropenia, n (%)	≥Grade 3	92 (94.8)	NR	NR
Anomia $n(9/)$	Overall	79 (81.4)	17 (29.8)	ND
Anemia, n (%)	≥Grade 3	66 (68.0)	10 (17.5)	– NR
Thrombocytopenia,	Overall	77 (79.4)	28 (49.1)	– NR
n (%)	≥ Grade 3	58 (59.8)	13 (22.8)	
Infactions n (P/)	Overall	56 (57.7)	NR	NR
Infections, n (%)	≥ Grade 3	19 (19.6)	NR	INK

AE: adverse event, CAR+: chimeric antigen receptor positive, CRS: cytokine release syndrome, kg: kilogram, n: number, N: total number, N/A: not available, NR: not reported, SAE: serious adverse events, SE: standard error

* Grade ≥3 AEs

⁺ Median 8-month follow up time

Table D3.13. Safety Outcomes: Belantamab

Trial		DREAMM-2 ^{17,18,22}
Arms		Belantamab 2.5 mg/kg
N (Safety Population)		95
Any AE, n (%)		93 (97.9)
Treatment-related AEs, n (%)		84 (88.4)
Any SAEs, n (%)		40 (42.1)
Treatment-related SAEs, n (%)		11 (11.6)
Mortality, n (%)	Overall	31 (32.6)†
	Disease Progression	25 (26.3)†
	AEs	3 (3.2)
	Other	3 (3.2)†
Discontinuation, n (%)	Overall	73 (76.8)†
	Death	31 (32.6)†
	Disease Progression	59 (60.8)†
	AEs	9 (9.5)
	Corneal Events	3 (3.2)
	Lack of Efficacy	1 (1.1)†
CRS, n (%)		0 (0)
Neurotoxicity, n (%)		0 (0)
Infections, n (%)		1 (1.1)*†
Renal Failure, n (%)		1 (1.1)†
Hyper viscosity, n (%)		0 (0)†
Hypercalcemia, n (%)	Overall	13 (13.7)†
	≥Grade 3	7 (7.4)
Neutropenia, n (%)	Overall	13 (13.7)†
	≥Grade 3	9 (9.3)
Anemia, n (%)	Overall	23 (24.2)†
	≥Grade 3	20 (21.1)
Fhrombocytopenia, n (%)	Overall	36 (37.9)
	≥ Grade 3	21 (22.1)

Trial Arms		DREAMM-2 ^{17,18,22} Belantamab 2.5 mg/kg
Grade 1	8 (8.4)	
Grade 2	16 (16.8)	
Grade 3	43 (45.3)	
Grade 4	1 (1.1)	
Keratopathy (MECs) Grade ≥2, n (%)	Median Time to Onset, Days (Range)	37.0 (19.0-143.0)
	Median Duration, Days (Range)	86.5 (8.0-358.0)
	Recovered [‡] from First Occurrence, n (%)	46/60 (76.7)§
Blurred Vision, n (%)		24 (25.3)
Dry Eye, n (%)		14 (14.7)
Permanent Vision Loss, n (%)		0 (0)
Changes in BCVA	Overall, n (%)	51 (53.7)
	Grade 1	7 (7.4)
	Grade 2	15 (15.8)
	Grade 3	28 (29.5)
	Grade 4	1 (1.1)
	Median Time to Onset, Days (Range)	64.0 (20–213)
	Median Duration, Days (Range)	33.0 (8–127)
	Recovered from First Occurrence, n (%)§	34/44 (77)
Clinically Meaningful Changes in BCVA (BCVA of 20/50 or worse in the better-seeing eye)	Overall, n (%)	17 (17.9)
	Median Time to Onset, Days (Range)	66.0 (20-442)
	Median Time to Resolution [#] , Days (Range)	21.5 (7–64); N=14
	Not recovered as of last follow-up, n (%)	3/17 (17.6)
	Overall, n (%)	1 (1.1)
Clinically Meaningful Changes in BCVA (BCVA of 20/200 or worse in the better-seeing eye)	Median Time to Onset, Days (Range)	21.0 (21–21)
	Median Time to Resolution [#] , Days (Range)	22.0 (22–22); N=1
	Not Recovered as of Last Follow-up, n (%)	0 (0)

AE: adverse event, BCVA: best corrected visual acuity, CAR+: chimeric antigen receptor positive, CRS: cytokine release syndrome, kg: kilogram, MECs: Microcyst-like epithelial changes, mg: milligram, n: number, N: total number, N/A: not available, NR: not reported, SAE: serious adverse events, SE: standard error.

* Viral infection

+ 6.3-month follow up time

‡ Any Grade 1 eye exam findings/no exam findings

§ Data for Grade \geq 2 events per the keratopathy and visual acuity (KVA) scale

 ${\tt x}$ in patients who recovered as of last follow-up

Table D3.14. Subgroup Safety Data: Ide-cel

Tria	l	KarMI	/la ³¹		11.3 Months 14 16	
<u>Cuban</u>		Elderly P	atients	Elderly Patients	High	ı Risk
Subgro	oups	≥ 65 years	≥ 70 years	≥ 65 years	Bridging Therapy	High Tumor Burden
Median Fo	llow-Up	11.3 M	onths		11.3 Months	
N		45	20	11	14 16	
	Overall	40 (88.9)	20 (100)	9 (82.0)	11 (79.0)	12 (75.0)
CRS, n (%)	Grade ≥3	2 (4.4)	2 (10.0)	NR	NR	NR
Neurotoxicity, n	Overall	11 (24.4)	6 (30.0)	NR	NR	NR
(%)	Grade ≥3	4 (8.9)	1 (5.0)	NR	NR	NR

Subgroup data not reported for the following outcomes: Any adverse events, any serious adverse events, keratopathy, dry eye, blurred vision, thrombocytopenia, anemia; Subgroup safety data not available for the following: KarMMa high risk subgroups, CARTITUDE-1, LEGEND-2

CRS: cytokine release syndrome, n: number, N: total number, NR: not reported, SAE: serious adverse event

Table D3.15. Subgroup Safety Data: Belantamab

Trial				DREAMM-2 ³³⁻³⁵		
		Prior Th	erapies	High Risk	Renal Im	pairment*
Subgrou	ups	3-6 Therapies	≥ 7 Therapies	High Risk Cytogenetics	Mild	Moderate
N		47	50	41	48	24
Median Fol	low-Up	12.4 N	lonths	9 Months	9 M	onths
Any SAEs,	n (%)	N	R	19 (46.3)	16 (33.3) 12 (50.0	
Kanatanathar	Overall	NR	NR	24 (58.5)	33 (68.8)	15 (62.5)
Keratopathy	Grade ≥3	16 (33.0)	14 (27.0)	NR	12 (25.0)	8 (33.3)
Dry eye	Overall	NR	NR	5 (12.2)	9 (18.8)	1 (4.2)
Blurred Vision	Overall	NR	NR	8 (19.5)	10 (20.8)	4 (1.7)
Thursday and an an in	Overall	NR	NR	17 (41.5)	11 (22.9)	6 (25.0)
Thrombocytopenia	Grade ≥3	8 (17.0)	10 (20.0)	NR	9 (18.8)	6 (25.0)
Amorria	Overall	NR	NR	10 (24.4)	13 (27.1)	7 (29.2)
Anemia	Grade ≥3	5 (11.0)	16 (31.0)	NR	9 (18.8)	6 (25.0)

GFR: glomerular filtration rate, m²: meters squared, min: minute, mL: milliliter, n: number, N: total number, NR: not reported, NT: neurotoxicity, SAEs: serious adverse events

* Mild renal impairment defined as GFR ≥60-<90 mL/min/1.73 m², moderate renal impairment defined as GFR ≥30-<60 mL/min/1.73 m²

Table D3.16. Secondary Analyses: Ide-cel

	Source		Rodriguez-Ote	ro EHA 2020 ⁷³	Jagannath	ASCO 2020 ⁷⁴	Shah AS	H 2020 ⁴⁴
	Trials		KarMMa vs.	DREAMM-2		KarMMa Real orld*	KarMMa vs.	маммотн
	Arms		KarMMa (N=128)	DREAMM-2 2.5 mg/kg (N=97)	KarMMa (N=128)	RW EC (N=190)	Ide-cel (ESS=67) Ide-cel (ESS=67) 450x10 ⁶ CAI cells/kg (ESS	
Med	ian Follow Up	o, Months	13.3	Study level	11.3	10.2	13	3.3
	Age, M	ledian Years	61 (33-78)	65	61	64		
	Ma	lle, n (%)	76 (59.4)	51 (52.6)	76 (59.4)	NR		
	Extramedu	ıllary Disease, n (%)	50 (39.1)	22 (22.7)	50 (39.1)	NR		D
	ISS sta	age 3, n (%)	21 (16.4)	42 (43.3)	21 (16.4)	8 (4)		R
		Prior Regimens /ed (Range)	6 (3-16)	7 (3-21)	6 (3-16)	5 (NR)		
Baseline	High-risk Cy	togenetics, n (%)	45 (35.2)	41 (42.3)	45 (35.2)	57 (30.0)		
Dusenne		Bortezomib	NR	95 (97.9)				
	Prior	Carfilzomib	NR	74 (76.3)	NR		NR	
	Therapies	Lenalidomide	NR	97 (100)				
	Received,	Pomalidomide	NR	89 (91.8)	ľ	NK		ĸ
	n (%)	Daratumumab	NR	97 (100)				
		Isatuximab	NR	3 (3.1)				
	Triple-re	fractory, n (%)	108 (84.0)	97 (100)	108 (84.0)	82 (43.0)	N	R
	ORR,	OR (95%CI)	5.12 (2.3	5, 11.13)	RR: 2.4 (1.7,	3.3), p<0.0001	5.11 (2.92, 8.94), p<0.001	6.96 (2.94, 16.49)
Efficacy	vgPR,	RR (95%CI)	N	R	4.2 (2.4, 7.	2), p<0.0001	NR	NR
Outcomes	PFS, I	HR (95%CI)	0.45 (0.2	27,0.76)	0.48 (0.33, 0	.69), p<0.0001	0.55 (0.42, 0.73), p<0.001	0.37 (0.23, 0.57)
	OS, H	IR (95%CI)	0.36 (0.1	15, 0.86)	1	NR	0.36 (0.24, 0.54), p<0.001	0.35 (0.18, 0.67), p=0.002
Data not rep	OS, F ported for safe		0.36 (0.1	15, U.86)	ľ	١ĸ	p<0.001	p=0.00

95% CI: 95% confidence interval, AE: adverse event, CAR+: chimeric antigen receptor positive, ESS: effective sample size, HR: hazard ratio, kg: kilogram, mg: milligram, n: number, N: total number, NR: not reported, OR: odds ratio, ORR: overall response rate, PFS: progression-free survival, RR: risk ratio, SAE: serious adverse events, vgPR: very good partial response

* Real world patients meeting KarMMa eligibility criteria (noninterventional, retrospective study (KarMMa-RW), patient-level data from clinical sites)

	Source		Trudel EHA 2020 ⁶³	Shragai EHA 2020 ⁶⁴	
	Trials		Pooled DREAMM-1 and DREAMM-2	Real World	
	Arms		Belantamab 2.5mg/kg (N=103)	Belantamab 2.5 and 3.4 mg/kg (N=32)	
Medi	an Follow Up, Mor	iths	NR	5.7 (0.5-13.8)	
	Age, m	edian years	65.0 (39.0-85.0)	69.6 (49-88)	
	Ma	le, n (%)	52 (50.5)	19 (59.4)	
	Extramedulla	ary Disease, n (%)	21 (20.4)	NR	
	ISS Sta	ge 3, n (%)	41 (39.8)	NR	
	Median Prior Regimens (Range)		7 (3-21)	6 (3-11)	
Baseline	High-risk Cytogenetics, n (%)		28 (27.2)	7 (21.9)	
		Bortezomib	NR	30 (93.8)	
	Prior Therapies	Carfilzomib		25 (78.1)	
		Lenalidomide		29 (90.6)	
	Received, n (%)	Pomalidomide		28 (87.5)	
		Daratumumab		31 (96.9)	
		Isatuximab		6 (18.8)	
	Triple ref	ractory, n (%)	NR	NR	
	ORR, (DR (95%CI)		n/N (%): 12/29 (41.4)	
ficacy Outcomes	vgPR,	RR (95%CI)	NR	n/N (%): 8/29 (27.6)	
incacy Outcomes	Median PFS,	months (95%Cl)		2.6 (NR)	
	OS at 6 m	nonths, n (%)		22 (68.0)	
Safety	A	ny AEs	101 (98.1)		
Outcomes,	Treatmei	nt related AEs	91 (88.3)	NR	
n (%)	Ar	y SAEs	42 (40.8)		

Table D3.17. Secondary Analyses: Belantamab

	Source		Trudel EHA 2020 ⁶³	Shragai EHA 2020 ⁶⁴
	Trials		Pooled DREAMM-1 and DREAMM-2	Real World
	Arms		Belantamab 2.5mg/kg (N=103)	Belantamab 2.5 and 3.4 mg/kg (N=32)
	Treatmen	t-related SAEs	13 (12.6)	
	Fatal Treatm	ent-related SAEs	1 (0.9)	
	Anemia	Overall	27 (26.2)	
		Grade ≥3	19 (18.4)	NR
	Thrombo-	Overall	24 (23.3)	7/23 (30.4)
	cytopenia	≥ Grade 3	18 (17.5)	3/23 (13.0)
	Kanatanatha	Overall	68 (66.0)	20/31 (64.5)
	Keratopathy	≥ Grade 3	28 (27.2)	8/31 (25.8)
	Diama d Maian	Overall	20 (19.4)	
	Blurred Vision	≥ Grade 3	4 (3.9)	NR
	Dura	Overall	12 (11.7)	
	Dry Eye	≥ Grade 3	0 (0)	NR

AE: adverse event, CAR+: chimeric antigen receptor positive, ESS: effective sample size, HR: hazard ratio, kg: kilogram, mg: milligram, n: number, N: total number, NR: not reported, OR: odds ratio, ORR: overall response rate, PFS: progression-free survival, RR: risk ratio, SAE: serious adverse events, VGPR: very good partial response

Usual Care Regimens

Table D3.18. Study Design: Usual Care Regimens

Trial/Study	Design	Patient Characteristics
MAMMOTH ⁸	Retrospective cohort study N = 275	 Diagnosis of active MM Refractory to daratumumab or isatuximab (administered alone or in combination) Treatment for at least 4 weeks with a CD38 MoAB-containing treatment regimen and with evidence of progressive disease
Mehra 2020 ²⁶	Real-world treatment patterns; Flatiron Health electronic health records N =251	 MM diagnosis Received at least 3 lines of therapy (including at least one PI, one IMiD, and an anti-CD38 MoAB)
Goldsmith 2020 ²⁵	Single-center, retrospective cohort study N = 58	 Quad/Penta-refractory MM Received at least one cycle of bendamustine/prednisone (BP) and/or dexamethasone, cyclophosphamide, etoposide, and cisplatin (DCEP)

IMiD: immunomodulatory drug, MM: multiple myeloma, MoAB: monoclonal antibody, N: total number, PI: protease inhibitor

Trial/	Study		MAMMOTH ⁸		Mehra 2020 ²⁶	Goldsmi	th 2020 ²⁵
Ar	ms	Overall	Triple/Quad- refractory	Penta-refractory	Triple-refractory	ВР	DCEP
I	N	275	148	70	251	27	31
	w-Up, Months nge)		10.6 (1.9-42.3)	-	6.0 (IQR: 2.7-10.8)	Ν	IR
Age, Median	Years (Range)	65.0 (27.0–90.0)	60.0 (23.0–85.0)	58.5 (35.0–76.0)	67.9 (IQR: 61.3-75.7)	61 (38-85)	60 (38-73)
Male,	n (%)	152 (55.3)	85 (57.4)	39 (55.7)	136 (54.2)	12 (44.4)	15 (48.4)
	White	202 (73.5)	116 (78.4)	47 (67.1)	173 (68.9)	23 (85.2)	27 (87.1)
Race/Ethnicity,	Black	45 (16.4)	19 (12.8)	13 (18.6)	38 (15.1)	4 (14.8)	3 (9.7)
n (%)	Hispanic	6 (2.2)	1 (0.7)	2 (2.9)	20 (12 0)	0.(0)	1 /2 2)
	Other	22 (8.0)	12 (8.1)	8 (11.4)	30 (12.0)	0 (0)	1 (3.2)
	ince Diagnosis, Range)	4.5 (0.4–19.4)	4.4 (0.4–19.4)	5.7 (0.6–14.4)	44.8 (IQR: 28.0, 62.4)	4.7 (1.1-23.1)	4.5 (1.1-2.32)
High Risk Cyto	genetics, n (%)	80 (29.1)	42 (28.4)	25 (35.7)	62 (24.7)	NR	NR
Renal Func	tion†, n (%)	22 (8.0)	11 (7.4)	5 (7.1)	NR	NR	NR
	Stage 1	69 (25.1)	40 (27.0)	17 (24.3)	49 (19.5)	4 (14.8)	9 (29.0)
ISS Disease Stage, n (%)	Stage 2	84 (30.5)	40 (27.0)	24 (34.3)	59 (23.5)	9 (33.3)	12 (38.7)
Stage, 11 (76)	Stage 3	80 (29.1)	47 (31.8)	13 (18.6)	62 (24.7)	7 (25.9)	5 (16.1)
ECOG	Grade 0				124 (52.4)		
Performance	Grade 1		NR		134 (53.4)	NR	
Status	Grade 2	-			N/A		
Received autolo	ogous SCT, n (%)	198 (72.0)	104 (70.3)	47 (67.1)	141 (56.2)*	22 (81.5)	28 (90.3)
	or Regimens, ange)	4 (1–16)	4 (1–11)	5 (2–16)	4 (IQR: 3, 6)	6 (4-15)	8 (4-15)
Penta-exp	osed, n (%)	157 (57.1)	70 (47.3)	70 (100)	117 (46.6)	NR	NR
Quad-refra	ctory, n (%)	NR	NR	0 (0)	NR	5 (18.5)	8 (25.8)
Penta-refra	ctory, n (%)	70 (25.4)	0 (0)	70 (100)	73 (29.1)	22 (81.5)	23 (74.2)
Exposed,	Daratumumab	at least 256 (93.1)	at least 138 (93.2)	at least 68 (97.1)	251 (100)	NR	NR
n (%)	Lenalidomide	270 (98.2)	146 (98.6)	70 (100)	244 (97.2)	27 (100)	31 (100)

Table D3.19. Baseline Characteristics: Usual Care Regimens

Tria	l/Study		MAMMOTH ⁸			Goldsmith 2020 ²⁵	
ŀ	Arms	Overall	Triple/Quad- refractory	Penta-refractory	Triple-refractory	BP	DCEP
	Bortezomib	271 (98.6)	146 (98.6)	69 (98.6)	226 (90.0)	27 (100)	31 (100)
	Pomalidomide	189 (68.7)	91 (61.5)	69 (98.6)	173 (68.9)	27 (100)	31 (100)
	Carfilzomib	178 (64.8)	85 (57.4)	68 (97.1)	145 (57.8)	27 (100)	31 (100)
	Elotuzumab	NR	NR	NR	24 (9.6)	NR	NR
	Ixazomib	38 (13.9)	24 (16.2)	12 (17.1)	NR	NR	NR
	Thalidomide	55 (20.0)	26 (17.6)	23 (32.9)	NR	NR	NR
	Daratumumab	256 (93.1)	138 (93.2)	68 (97.1)	-	22 (81.5)	23 (74.2)
	Lenalidomide	211 (76.7)	117 (79.1)	69 (98.6)		27 (100)	31 (100)
	Bortezomib	188 (68.4)	107 (72.3)	68 (97.1)		27 (100)	31 (100)
Refractory,	Pomalidomide	179 (65.1)	87 (58.8)	69 (98.6)	ND	27 (100)	31 (100)
n (%)	Carfilzomib	130 (47.3)	57 (38.5)	67 (95.7)	NR	27 (100)	31 (100)
	Elotuzumab	NR	NR	NR		NR	NR
	Ixazomib	34 (12.4)	23 (15.5)	10 (14.3)		NR	NR
	Thalidomide	23 (8.4)	6 (4.1)	14 (20.0)		NR	NR

Data not reported on the following baseline characteristics: Height, weight BP: bendamustine/prednisone, DCEP: dexamethasone, cyclophosphamide, etoposide, and cisplatin, dL: deciliter, ECOG: Eastern Cooperative Oncology Group, IQR: interquartile range, mg: milligram, n: number, N: total number, N/A: not available, NR: not reported, SCT: stem cell transplant.

* Type of SCT not specified.

+ Creatinine > 2mg/dL

Table D3.20. Efficacy Outcomes: Usual Care Regimens I

	Trial		MAMMOTH ⁸				
	Arms		Overall	Triple/Quad-refractory	Penta-refractor		
	N		275	148	70		
Median Fo	ollow-Up, Ma	nths (Range)	10.	6 (1.9-42.3)	•		
		Overall	NR				
	Received ≥ 1 Subsequent LOT ⁺		3.4 (2.8, 4.0)				
Median PFS, Months		Carfilzomib-based	4.2 (HR 0.60, p=0.004); [N=68]	NR	NR		
(95%Cl or HR)	LOT1 [†]	Carfilzomib + alkylator	5.7 (1.6-9.7); [N=19]	INK	INK		
	LUII	Daratumumab + IMiD	4.5 (2.8-6.3); [N=41]				
		Elotuzumab + IMiD	2.6 (1.1-4.1); [N=19]				
		Overall	8.6 (7.2, 9.9)	9.2 (7.1, 11.2)	5.6 (3.5, 7.8)		
	High-Risk Cytogenetics		5.6,(NR) p=0.025‡; [N=80]				
	Impaired Renal Function*		3.7 (NR) p=0.031; [N=22]				
Median OS, Months	Received ≥ 1 subsequent LOT		9.31 (8.1, 10.6)				
(95%CI)	Carfilzomib-based		10.9 (9.5-12.4); [N=68]	NR	NR		
	LOT1 [†]	Carfilzomib + alkylator	12.7 (5.9-19.5); [N=19]				
	1011	Daratumumab + IMiD	12.6 (8.5-16.6); [N=41]				
		Elotuzumab + IMiD	8.3 (1.9-14.6); [N=19]				
		Overall	116/249 (46.6)	NR	NR		
	Hig	h-Risk Cytogenetics	OR=0.14 (95%CI: 0.03, 0.65)	NR	NR		
		Overall	78/249 (31.0)	57/197 (29)	19/63 (30)		
		Carfilzomib-based	22/68 (32.4)				
ORR, n (%)	LOT1 ⁺	Carfilzomib + alkylator	9/19 (47.0)				
		Daratumumab + IMiD	15/41 (36.6)	NR	NR		
		Elotuzumab + IMiD	4/19 (21.1)	INIT			
		LOT2 ⁺	34/158 (21.5)				
		LOT3 ⁺	22/87 (25.3)				
ORR after LOT1 ⁺		sCR/CR, n (%)	5/249 (2.0)	NR	NR		
		vgPR, n (%)	22/249 (8.8)	INK	INK		

	Trial		MAMMOTH ⁸	
	Arms	Overall	Triple/Quad-refractory Penta-re	
	PR, n (%)	51/249 (20.5)		
Prog	ressive Disease, n (%)	77 (28.0)	38 (25.7)	27 (38.6)
	N	249	134	63
Patients Receiving	Median LOT received, n (range)	2 (1-10)	NR	NR
Treatment After Becoming anti-CD38	Carfilzomib-based Treatment, n (%)	68 (27.3)	43 (32.1)	8 (12.7)
MoAB Refractory	Elotuzumab-based Treatments, n (%)	19 (7.6)	NR	NR
,	Daratumumab-based Treatments, n (%)	57 (22.9)	49 (36.6)	9 (50.9)

Data not reported for the following efficacy outcomes: LOT received (Pomalidomide-based, Bortezomib-based, Lenalidomide-based, Ixazomib-based, Thalidomide-based, Monotherapy, Doublet regimen, Triplet regimens)

95% CI: 95% confidence interval, CR: complete response, HR: hazard ratio, IMiD: immunomodulatory drug, LOT: line of therapy (after becoming CD38 MoAB refractory), MoAB: monoclonal antibody, n: number, N: total number, NR: not reported, OR: odds ratio, ORR: overall response rate, OS: overall survival, PFS: progression-free survival, PR: partial response, sCR: stringent complete response, vgPR: very good partial response,

* Creatinine > 2mg/dl.

+ First (LOT1), second (LOT2), or third (LOT3) line of therapy after becoming CD38 MoAB refractory.

‡ Versus standard risk.

Table D3.21. Efficacy Outcomes: Usual Care Regimens II

	Study		Mehr	a 2020 ²⁶		Goldsmit	h 2020 ²⁵
	Arms	Overall	LOT1*	LOT2*	LOT3*	BP	DCEP
	Ν	251	251	138	54	27	31
Median Follow	/-Up, Months (Range)		6.0 (IQR	: 2.7-10.8)		N	R
Median PF	S, Months (95%Cl)	4.8 (3.7, 6.1)	NR	NR	NR	1.4 (1.1-1.6)	2.7 (1.5-3.8)
Median OS	6, Months (95%CI)	11.0 (8.7, 13.6)	NR	NR	NR	8.7 (2.3-15.0)	6.2 (4.4-7.8)
0	RR, n (%)	NR	NR	NR	NR	7 (25.9)	11 (35.5)
ORR after First	sCR/CR, n (%)					CR: 0 (0)	CR: 1 (3.2)
Subsequent	vgPR, n (%)		I	NR		4 (14.8)	1 (3.2)
Line	PR, n (%)					3 (11.1)	9 (29.0)
Progressiv	ve Disease, n (%)	NR	NR	NR	NR	NR	NR
	to Next Treatment, ths (95%Cl)	NR	NR	NR	NR	_	
-	tment After Becoming -refractory, n	251	251	138	54		
	After Becoming Triple- , Median (Range)	2 (1-8)	NR	NR	NR		
	Carfilzomib-based	111 (44.2)	81 (32.3)	25 (18.1)	12 (22.2)		
	Pomalidomide-based	98 (39.0)	76 (30.3)	37 (26.8)	9 (16.7)		
	Bortezomib-based	83 (33.1)	47 (18.7)	32 (23.2)	12 (22.2)		
	Elotuzumab-based	64 (25.5)	41 (16.3)	19 (13.8)	7 (13.0)	N	D
Lines of	Lenalidomide-based	57 (22.7)	39 (15.5)	19 (13.8)	7 (13.0)		n
Therapy Received, n	Ixazomib-based	NR	22 (8.8)	25 (18.1)	5 (9.3)		
(%)	Thalidomide-based	NR	12 (4.8)	2 (1.4)	4 (74)		
. ,	Daratumumab-based	NR	5 (2.0)	19 (13.8)	8 (14.8)		
	Monotherapy Doublet Regimen		123 (49.0)	58 (42.0)	26 (48.1)		
			85 (33.9)	54 (39.1)	13 (24.1)		
	Triplet Regimen		7 (2.8)	4 (2.9)	4 (7.4)		
Retreatment	Carfilzomib	39/145 (26.9)	ND	ND	ND		
with	Pomalidomide	49/173 (28.3)	NR	NR	NR		

	Study		Mehra	2020 ²⁶		Goldsmith 2020 ²⁵	
	Arms		LOT1*	LOT2*	LOT3*	BP	DCEP
Previously	Bortezomib	72/226 (31.9)					
	Elotuzumab	1/24 (4.2)					
Regimen, n/N (%)	Lenalidomide	55/244 (22.5)					
(70)	Ixazomib	NR					
	Thalidomide	NR					
	Daratumumab	34/251 (13.5)					

95% CI: 95% confidence interval, BP: bendamustine/prednisone, CR: complete response, DCEP: dexamethasone, cyclophosphamide, etoposide, and cisplatin, LOT: line of therapy, n: number, N: total number, NR: not reported, ORR: overall response rate, OS: overall survival, PFS: progression-free survival, PR: partial response, sCR: stringent complete response, vgPR: very good partial response

*Subgroup of patients receiving a first (LOT1), second (LOT2), or third (LOT3) line of therapy after becoming triple-class refractory.

Table D3.22. Safety Outcomes: Usual Care Regimens

Trial/Study		MAMMOTH ⁸	Golds	smith 2020 ²⁵
	Arms	Overall	BP DCEP 27 31	
	N	275		
Median Follow-Up, Months (Range)		10.6 (1.9-42.3)	NR	
Risk of PD or Death,	Carfilzomib-based Regimens	0.60 (0.42, 0.85)	NR	
HR (95%CI)	Daratumumab + IMiD Regimens	0.64 (0.43, 0.94)		
Neutronomia	Grade 4		5 (18.5)	NR
Neutropenia	Neutropenic fevers		NR	5 (16.1)
Thrombocytopenia	Grade 4	NR	9 (33.3)	NR
Tumo	or Lysis Syndrome		1 (3.7)	1 (3.2)
Sepsis			1 (3.7)	1 (3.2)
Safety data not report	ted for MAMMOTH triple/quad-refra	ctory or penta-refractory arms, or any Meh	ra 2020 arm.	

95% CI: 95% confidence interval, BP: bendamustine/prednisone, DCEP: dexamethasone, cyclophosphamide, etoposide, and cisplatin, HR: hazard ratio, IMiD: immunomodulatory drug, n: number, N: total number, NR: not reported, PD: progressive disease

Trial/Study		ELOQUENT-3 ³⁰	TOURMALINE-MM1 ²⁸	Bringhen 2014 ²⁹
Treatm	nent	Elo + Pom + Dex	lxa + Len + Dex	Car + Cy + Dex
Median Fo	llow-up	Minimum 9.1 Months	23.3 Months	18 Months
Safety Popu	lation, N	60	361	56
Age, Mediar	n (Range)	69 (43-81)	66 (38-91)	71 (IQR: 68-75)
Prior Therapies, N	Aedian (Range)	3 (2-8)	NR	NR
Double-refrac	tory, n (%)	41 (68)	NR	NR
AEs, n	(%)	58 (96.7)	355 (98.3)	44 (78.6)*
Grade ≥3 A	Es, n (%)	34 (56.7)	267 (74.0)	15 (26.8)*
SAEs, n	(%)	32 (53)	168 (46.5)	NR
	Overall	13 (21.7)	15 (4.2)	7 (12.5)
Mortality, n (%)	Disease Progression	8 (13.3)	NR	2 (3.6)
	AE	5 (8.3)	NR	4 (7.1)
Discontinuation de	ue to AEs, n (%)	11 (18)	60 (16.6)	8 (14)
	Overall	14 (23.3)	118 (32.7)	20 (35.7)
Neutropenia, n (%)	Grade 3/4	8 (13.3)	81 (22.4)	11 (19.6)
Amountin (9/)	Overall	15 (25.0)	103 (28.5)	39 (69.6)
Anemia, n (%)	Grade 3/4	6 (10.0)	34 (9.4)	6 (10.7)
Thrombocytopenia, n	Overall	9 (15.0)	112 (31.0)	21 (37.5)
(%)	Grade 3/4	5 (8.3)	69 (19.1)	2 (3.6)
1	Overall	6 (10.0)	NR	NR
Lymphopenia, n (%)	Grade 3/4	5 (8.3)	NR	NR
	Overall	12 (20.0)	NR	7 (12.5)
Hyperglycemia, n (%)	Grade 3/4	5 (8.3)	NR	1 (1.8)
Infections = (0/)	Overall	39 (65.0)	83 (23.0)	10 (17.9)
Infections, n (%)	Grade 3/4	8 (13.3)	2 (0.6)	3 (5)
	Overall	9 (15.0)	106 (29.4)	11 (20)
Fatigue, n (%)	Grade 3/4	0 (0.0)	13 (3.6)	1 (1.8)
Neuropathy, n (%)	Overall	NR	97 (26.9)	5 (8.9)

Table D3.23. Safety Outcomes: Additional Usual Care Regimens⁺

Trial/Study		ELOQUENT-3 ³⁰	TOURMALINE-MM1 ²⁸	Bringhen 2014 ²⁹
Treatm	ent	Elo + Pom + Dex	lxa + Len + Dex	Car + Cy + Dex
	Grade 3/4 NR		9 (2.5)	0 (0)

AEs: adverse events, Car: carfilzomib, Cy: cyclophosphamide, Dex: dexamethasone, Elo: elotuzumab, IQR: interquartile range, Ixa: ixazomib, Len: lenalidomide, n: number, N: total number, NR: not reported, Pom: pomalidomide, SEA: serious adverse events

* Hematologic adverse events.

[†] Representative prospective trials of commonly used treatments selected externally to the systematic review meant to supplement the insufficient safety data in the three retrospective studies selected to represent the effectiveness of usual care regimens.

D4. Ongoing Studies

Table D4.1. Ongoing Studies: Ide-cel

Title / Trial Sponsor	Study Design	Treatment Arms	Patient Population	Outcomes	Estimated Completion
Study of bb2121 in	Two-part, Non-	Dose Escalation	Inclusions:	Primary:	November
Multiple Myeloma	randomized, Phase	Phase (Part A):	 ECOG status of 0 or 1 	 Incidence of 	30, 2023
	I Clinical Trial	– Ide-cel (dose	 Measurable disease 	adverse events	
Celgene		range: 150 – 450 x	 Diagnosis of relapsed or refractory MM with 	(including dose	
	Actual enrollment:	106 CAR+ T cells)	at least 3 different prior lines of therapy	limiting toxicities)	
NCT02658929	N = 67		including PI & IMiD, or be "double-		
		Expansion Phase	refractory" to a PI and IMiD or previous	Secondary:	
		(Part B):	(Part A)	– ORR, CR, vgPR, PR	
		-Ide-cel	 Diagnosis of relapsed/refractory MM with 		
		(recommended	previous PI, IMiD, and dara exposure	[60 Months]	
		dose)			
			Exclusions:		
			 Known CNS disease 		
			 Inadequate renal, hepatic, bone marrow 		
			function		
			 Significant co-morbid conditions, second 		
			malignancies, history of class III or IV		
			congestive heart failure etc.		
Efficacy and Safety	Single Arm, Phase	Intervention:	Inclusions:	Primary:	November
Study of bb2121 in	II Clinical Trial	– Ide-cel (dose	– ≥3 prior MM treatments with at least 2	– ORR	1, 2024
Subjects with Relapsed		range: 150 – 450 x	consecutive cycles of treatment for each		
and Refractory	Actual enrollment:	10 ⁶ CAR+ T cells)	regimen	Secondary:	
Multiple Myeloma	N = 149		 Received PI, IMiD, and an anti-CD38 	– CR, Time to	
(KarMMa)			antibody, refractory to last treatment	response, DOR,	
			 ECOG status 0 or 1 	PFS, OS, MRD	
Celgene			 Measurable disease 	– Adverse events	

Title / Trial Sponsor	Study Design	Treatment Arms	Patient Population	Outcomes	Estimated Completion
				– EORTC-QLQ-C30,	
NCT03361748			Exclusions:	EuroQol Group	
			 Known CNS involvement with myeloma or 	EQ-5D-5L, EORTC-	
			presence of relevant CNS pathology	QLQ-MY20	
			 Active or history of plasma cel leukemia 		
			 Inadequate organ function 	[≥24 months]	
			 History of allogeneic hematopoietic SCT 		
An Efficacy and Safety	Single Arm, Multi-	Intervention:	Inclusions:	Primary:	May 13,
Study of bb2121 in	cohort Phase II	– Ide-cel (dose	 Measurable disease 	– Cohort 1: ORR	2026
Subjects with Relapsed	Clinical Trial	range: 150 – 450 x	 Cohort-specific requirements: (Cohort 1) 	 Cohort 2: CR 	
and Refractory		10 ⁶ CAR+ T cells)	relapsed/refractory MM subjects with ≥3		
Multiple Myeloma and	<u>Estimated</u>		prior treatment regimens; (Cohort 2) subject	Secondary:	
in Subjects with High-	<u>enrollment</u> :		with 1 prior treatment regimen	– Cohort 1: CR	
Risk Multiple Myeloma	N = 181		– ECOG status ≤1	– Cohort 2: ORR	
(KarMMa-2)			 Grade 1 or baseline non-hematological 	 DOR, time to 	
			toxicities due to prior treatments	response, PFS,	
Celgene				OS, time to	
			Exclusions:	progression, MRD	
<u>NCT03601078</u>			 Receiving investigational dugs, 	 Adverse events 	
			plasmapheresis, major surgery, radiation	[≥5 years]	
			therapy, systemic anti-myeloma therapy 14	– EORTC-QLQ-C30,	
			days prior to leukapheresis	EuroQoL Group	
			 History of relevant CNS pathology, CNS 	EQ-5D-5L, EORTC-	
			involvement with myeloma	QLQ-MY20 [≥5	
			- Active plasma cell leukemia, Waldenstrom's	years]	
			macroglobulinemia, POEMS syndrome,		
			clinically significant amyloidosis	[≥2 years]	
			 Previous allogeneic SCT 		

Title / Trial Sponsor	Study Design	Treatment Arms	Patient Population	Outcomes	Estimated Completion
Efficacy and Safety	Randomized,	Intervention:	Inclusions:	Primary:	November
Study of bb2121	Phase III Clinical	– Ide-cel (dose	 Measurable disease 	– PFS	6, 2025
Versus Standard	Trial	range: 150-450 x	 ECOG status of 0 or 1 		
Regimens in Subjects		10 ⁶ CAR+ T cells)	– Received ≥2 but not >4 prior MM regimens	Secondary:	
with Relapsed and	Estimated		– Prior treatment with dara, a PI, and an IMiD-	 OS, event-free 	
Refractory Multiple	<u>enrollment</u> :	Standard Regimens:	containing regiment for ≥2 consecutive	survival, ORR,	
Myeloma (RRMM)	N = 381	– Dara/Pom/Dex	cycles	MRD, CR, DOR,	
(KarMMa-3) Efficacy		– Dara/Bor/Dex	 Refractory to last treatment regimen 	time to response	
and Safety Study of		– Ixa/Len/Dex	 Achieved minimal response or better to ≥1 	 Adverse events 	
bb2121 Versus		– Car/Dex	prior treatment	– EORTC-QLQ-C30,	
Standard Regimens in		– Elo/Pom/Dex		EuroQoL Group	
Subjects with Relapsed		-, - , -	Exclusions:	EQ-5D-5L, EORTC-	
and Refractory			– Non-secretory MM	QLQ-MY20	
Multiple Myeloma			 History of malignancies, history of or active 	$[\geq 5 years]$	
(RRMM) (KarMMa-3)			plasma cell leukemia, Waldenstrom's		
			macroglobulinemia, POEMS syndrome,		
Celgene			amyloidosis, etc.		
			 Subject treated with one of the comparator 		
NCT03651128					
			regimens as most-recent regimen cannot		
			receive it again but may receive one of the		
			others		
			 Autologous SCT 12 weeks prior to 		
			randomization		
A Study to Evaluate	Single Group,	Intervention:	Inclusions:	Primary:	January 15,
the Safety of bb2121 in	Phase 1 Clinical	– Ide-cel (dose	 New diagnosis of symptomatic MM 	 Dose-limiting 	2025
Subjects with High	Trial	range: 150-800 x	 Subject has measurable disease 	toxicity [Up to 2	
Risk, Newly Diagnosed		10 ⁶ CAR+ T cells)	 Subject has high-risk MM 	years]	
Multiple Myeloma	<u>Estimated</u>		- ECOG status of ≤ 1	 Adverse events 	
(NDMM) (KarMMa-4)	<u>enrollment</u> : N = 60		 Has received ≤ 3 cycles of induction anti- 	[Up to 5 years]	
			myeloma therapy		

Title / Trial Sponsor	Study Design	Treatment Arms	Patient Population	Outcomes	Estimated Completion
Celgene				Secondary: [2.5	
			Exclusions:	years]	
NCT04196491			 Non-secretory MM 	- CR, ORR, DOR,	
			 Subject received any treatments of MM 	PFS, OS	
			other than up to 3 cycles of induction	 Time to 	
			therapy	maintenance	
			 Clinically significant CNS pathology 	therapy	
			 High risk developing deep vein 		
			thrombosis/pulmonary embolus & cannot		
			undergo anti-thrombotic therapy		
			 Moderate or severe pulmonary 		
			hypertension		
			 Subject has cardiac or pulmonary 		
			conditions; needs chronic		
			immunosuppressants		
			 History of primary immunodeficiency 		

AE: adverse event, Bor: bortezomib, Car: carfilzomib, CNS: central nervous system, CR: complete response, Dara: daratumumab, Dex: dexamethasone, DOR: duration of response, ECOG: Eastern Cooperative Oncology Group, Elo: elotuzumab, IMiD: immunomodulatory drug, Ixa: Ixazomib, Len: lenalidomide, MM: multiple myeloma, MRD: minimal residual disease, N: total number, ORR: overall response rate, OS: overall survival, PFS: progression-free survival, PI: protease inhibitor, POEMS: polyneuropathy, organomegaly, endocrinopathy, myeloma protein, and skin changes, Pom: pomalidomide, SCT: stem cell transplant, vgPR: very good partial response. Source: www.ClinicalTrials.gov (NOTE: studies listed on site include both clinical trials and observational studies)

Table D4.2. Ongoing Studies: Cilta-cel

Title / Trial Sponsor	Study Design	Treatment Arms	Patient Population	Outcomes	Estimated Completion
LCAR-B38M Cells in	Single Arm Phase	Intervention:	Inclusions:	Primary: [1 month]	December
Treating	I/II Clinical Trial	 Cilta-cel (dose 	 IMWG confirmed diagnosis of active MM 	 Treatment related 	31, 2021
Relapsed/Refractory		range: 0.5-5 x 10 ⁶	 Refractory MM (≥3 prior regimens, 	adverse events	
(R/R) Multiple	Estimated	CAR+ T cells/kg)	including Bortezomib)		
Myeloma (LEGEND-2)	<u>enrollment</u> :		 Relapse criteria in NCCN clinical practice 	Secondary: [36 months]	
	N = 100		guidelines	 Changes in aberrant 	
Nanjing Legend				immunoglobin in	
Biotech Co.			Exclusions:	serum	
			 Systemic corticosteroid therapy greater 	 Multiple myeloma cells 	
<u>NCT03090659</u>			than 5 mg/day prednisone or equivalent of another corticosteroid within 2 weeks of leukapheresis or chemotherapy regimen	in bone marrow	
			 History of allogeneic SCT (active acute or 		
			chronic GVHD or require		
			immunosuppressant medication for GVHD		
			within 6 months of enrollment		
			 Active autoimmune diseases or CNS 		
			metastases or symptomatic CNS		
			involvement		
A Study of JNJ-	Single Arm, Phase	Intervention:	Inclusions:	Primary:	April 30,
68284528, a Chimeric	1b-2 Clinical Trial	 Cilta-cel (Dose: NR) 	 Measurable disease 	 Phase 1b: Adverse 	2022
Antigen Receptor T			 ECOG status of 0 or 1 	events	
Cell (CAR-T) Therapy	Estimated		 Received ≥3 prior therapies (including PI, 	– Phase 2: ORR	
Directed Against B-Cell	<u>enrollment</u> :		IMiD, and an anti-CD38 antibody), or		
Maturation Antigen	N = 126		double refractory to IMiD and PI	Secondary:	
(BCMA) in Participants			 Evidence of progressive disease 	 Phase 2: Adverse 	
with Relapsed or				events	
Refractory Multiple			Exclusions:	 vgPR or better, DOR, 	
Myeloma (CARTITUDE-			 Prior CAR-T therapy at any target, therapy 	PFS, OS, MRD	
1)			targeted to BCMA	 EORTC QLQ-C30 and 	
			 Allogenic SCT within 6 months before 	QLQ-MY20, EQ-5D-5L	
Janssen Research &			apheresis; autologous SCT within 12 weeks		
Development, LLC			before apheresis	[≥2 years]	

Title / Trial Sponsor	Study Design	Treatment Arms	Patient Population	Outcomes	Estimated Completion
			 Known active/prior history of CNS or 		
NCT03548207			meningeal involvement of MM		
A Study of LCAR-B38M	Single Arm, Phase	Intervention:	Inclusions:	Primary:	November
CAR-T Cells, a Chimeric	II Clinical Trial	- Cilta-cel (Dose: NR)	 Measurable disease 	- ORR	30, 2022
Antigen Receptor T-			 ECOG status of 0 or 1 		
cell (CAR-T) Therapy	Estimated		 Received ≥3 prior lines of MM treatment 	Secondary:	
Directed Against B-cell	<u>enrollment</u> : N =		(≥1 complete cycle of treatment for each	 vgPR or better, DOR, 	
Maturation Antigen	60		line; received a PI and IMiD)	PFS, OS, MRD	
(BCMA) in Chinese			 Evidence of progressive disease 	 Adverse events 	
Participants with					
Relapsed or Refractory			Exclusions:	[≥2 years]	
Multiple Myeloma			 Prior CAR-T therapy at any target, any 		
(CARTIFAN-1)			therapy targeted to BCMA		
			 Allogeneic SCT for MM; Autologous SCT 12 		
Nanjing Legend			weeks prior to apheresis		
Biotech Co.			 Diagnosed or treated for non-MM invasive 		
			malignancies		
NCT03758417			 Prior antitumor therapy, insufficient 		
			washout period		
A Study of JNJ-	Single Group,	Intervention:	Inclusions:	Primary:	July 25,
68284528, a Chimeric	single arm, Phase	– Cohort A: cilta-cel	 Cohort A: Received 1-3 lines of prior 	$-$ MRD [≥ 1 year]	2024
Antigen Receptor T	2 Clinical Trial	– Cohort B: cilta-cel	therapy		
Cell (CAR-T) Therapy		– Cohort C: cilta-cel	 Cohort B: One line of therapy, early relapse 	Secondary:	
Directed Against B-cell	Estimated	- Cohort D: cilta-cel +	 Cohort C: Treated with PI, IMiD, anti-CD38 	 ORR, vgPR or better, 	
Maturation Antigen	enrollment:	lenalidomide (only	monoclonal antibody and BCMA-directed	CBR, DOR	
(BCMA) in Participants	N = 100	some participants)	therapy	 Adverse events 	
with Multiple		– Cohort E: cilta-cel +	 Cohort D: Newly diagnosed MM with 	- Auverse events	
Myeloma (CARTITUDE-		Dara/Bor/Len/Dex	history of 4-8cycles initial therapy	[2] upgral	
2)		Dara Dor Len Dex	 Cohort E: Newly diagnosed, no prior 	[≥2 years]	
			therapy		
Janssen Research &			(incrapy		
Development, LLC			Evolucione		
• • • •			Exclusions:		
NCT04133636			 Prior treatment with CAR T therapy for any terrest 		
			target	1	

			 Ongoing toxicity Grade 1 or less (except 		
			alopecia or peripheral neuropathy) – Prednisone (≥70mg) within 7 days – History of CNS or meningeal involvement of multiple myeloma		
Pomalidomide, Bortezomib and	Interventional: Randomized, Parallel Assignment, Phase 3 Clinical Trial <u>Estimated</u> <u>enrollment</u> : N = 400	Intervention: – Cilta-cel (target dose of 0.75 * 10^6 CAR+ T cells/kg) Standard Regimens: – Bor/Pom/Dex – Dara/Pom/Dex	 Inclusions: Measurable disease Received 1-3 prior lines of therapy Evidence of progressive disease Refractory to lenalidomide Have clinical laboratory values meeting screening phase criteria Exclusions: Prior CAR T-cell therapy directed at any target Previous therapy targeting BCMA Ongoing toxicity from previous anticancer therapy Monoclonal antibody treatment within 21 days; Cytotoxic therapy or Proteasome inhibitory therapy within 14 days; Immunomodulatory drug therapy within 7 days 	Primary: – PFS Secondary: – ORR, CR/sCR, OS, MRD – Adverse events [Up to 6 years]	April 10, 2026

BCMA: b-cell maturation antigen, Bor: bortezomib, CAR: chimeric antigen receptor, CBR: clinical benefit rate, CNS: central nervous system, CR: complete response, Dara: daratumumab, Dex: dexamethasone, DOR: duration of response, ECOG: Eastern Cooperative Oncology Group, GVHD: graft versus host disease, IMiD: immunomodulatory drug, IMWG: International Myeloma Working Group, Len: lenalidomide, MM: multiple myeloma, mg: milligram, MRD: minimal residual disease, N: total number, NCCN: National Comprehensive Cancer Network, ORR: overall response rate, OS: overall survival, PFS: progression-free survival, PI: protease inhibitor, Pom: pomalidomide, sCR: stringent complete response, SCT: stem cell transplant, VGPR: very good partial response Source: www.ClinicalTrials.gov (NOTE: studies listed on site include both clinical trials and observational studies)

Table D4.3. Ongoing Studies: Belantamab

Title / Trial Sponsor	Study Design	Treatment Arms	Patient Population	Outcomes	Estimated Completion
A Study to Investigate	Randomized, Two-	Interventions:	Inclusions:	Primary:	November
the Efficacy and Safety	Arm Phase II Clinical	– Arm 1: 2.5 mg/kg	 ECOG status of 0-2 	– ORR	30, 2020
of Two Doses of	Trial	frozen	 Histologically/cytologically confirmed MM; 		
GSK2857916 in		belantamab	has undergone autologous SCT or	Secondary:	
Participants with	Actual enrollment:	every three	transplant-ineligible; has failed ≥2 prior	– CBR, DOR, PFS, OS	
Multiple Myeloma	N = 221	weeks	lines of therapy including IMiD and PI		
Who Have Failed Prior		-Arm 2: 3.4 mg/kg	 Measurable disease 	[Up to 48 weeks]	
Treatment with an		frozen	 History of autologous SCT only if 100 days 		
Anti-CD38 Antibody		belantamab	prior to study enrollment & no active		
(DREAMM 2)		– Arm 3: 3.4 mg/kg	infections		
		lyophilized	 Adequate organ system function 		
GlaxoSmithKline		belantamab			
			Exclusions:		
<u>NCT03525678</u>			 Systemic anti-myeloma therapy, 		
			treatment with high dose steroids,		
			investigational drug within 14 days		
			 Symptomatic amyloidosis, active POEMS, 		
			active plasma cell leukemia at time of		
			screening		
			 Prior allogeneic SCT 		
			 Current corneal epithelial disease, active 		
			renal condition, active mucosal/internal		
			bleeding, unstable liver/biliary disease,		
			other malignancies etc.		
Characterization of	Parallel, Non-	Intervention:	Inclusions:	Primary:	November
Corneal Epithelial	Randomized, Phase	– Arm 1:	 Age 18 years or older 	 Abnormal corneal 	30, 2020
Changes in Participants	3 Clinical Trial	Participants	 Patients with relapsed/refractory MM who 	epithelium composition	
Treated with		undergoing	receives/has received treatment with	 Abnormal pathologic 	
Belantamab Mafodotin	<u>Estimated</u>	Impression	Belantamab and microcyst-like epithelial	characteristics	
	<u>enrollment</u> : N = 25	cytology +	changes diagnosis		
GlaxoSmithKline		belantamab	 If undergoing superficial keratectomy, 	Secondary:	
		– Arm 2:	must not pose excessive risk to patient	– Adverse events, serious	
NCT04549363		Participants		adverse events	

Title / Trial Sponsor	Study Design	Treatment Arms	Patient Population	Outcomes	Estimated Completion
		undergoing Superficial keratectomy + belantamab	 Exclusions: Serious/unstable medical or psychiatric disorder Excess risk of delayed wound healing Eye infections Active uveitis Permanent legal blindness in the nonstudy eye 	 Abnormal BCVA scores, corneal symptoms, and corneal epithelial lesions [Up to 5 weeks] 	
A Study of Belantamab	Non-randomized,	Intervention:	Inclusions:	Primary:	July 31,
Mafodotin to Investigate Safety, Tolerability, Pharmacokinetics, Immunogenicity and Clinical Activity in Participants with Relapsed/Refractory Multiple Myeloma (RRMM) GlaxoSmithKline <u>NCT04177823</u>	Single Group, Phase 1 Clinical Trial Estimated enrollment: N = 12 Actual enrollment: N = 5	 2.5 mg/kg belantamab every 3 weeks 3.4 mg/kg belantamab every 3 weeks 	 ECOG status of 0-2 Histological/cytologically confirmed MM diagnosis Has undergone stem cell transplant/stem cell transplant not deemed feasible Failed at least 2 prior lines of antimyeloma treatment Refractory to an immunomodulatory drug and proteasome inhibitor Measurable disease Adequate organ system functions All prior treatment-related toxicities must be ≤ Grade 1, Grade 2 peripheral neuropathy 	 Adverse events, serious adverse events Dose-limiting toxicities Secondary: Systolic and diastolic blood pressure Hematology and chemistry parameters [Up to 15 months] 	2021
			Exclusions:		
			 Prior allogenic stem cell transplant Systemic anti-myeloma therapy or investigational drugs within 14 days; plasmapheresis within 7 days Symptomatic amyloidosis, POEMS syndrome Active renal condition, unstable liver, or biliary disease 		

Title / Trial Sponsor Study Design		Treatment Arms	Patient Population	Outcomes	Estimated Completion
			 Malignancies other than disease under study 		
An Open-label, Dose	Single Group, Dose-	Part 1: Belantamab	Inclusions:	Primary:	February
Escalation Study in	escalation, Phase I	monotherapy	 Age 20 years or older 	 Dose-limiting toxicities 	28, 2023
Japanese Subjects with	Clinical Trial		 ECOG status of 0-2 	[Day 21]	
Relapsed/Refractory		Part 2:	 Measurable disease 	 Adverse events 	
Multiple Myeloma	<u>Estimated</u>	– Belantamab +	 Autologous stem cell transplant > 100 days 	 Abnormal hematology, 	
Who Have Failed Prior	<u>enrollment</u> : N = 14	Bor/Dex	prior	clinical chemistry, urine	
Anti Myeloma		– Belantamab +	– Prior toxicities ≤ Grade 1, except alopecia	parameters	
Treatments		Pom/Dex	& peripheral neuropathy Grade 2	 Abnormal vital signs, 	
				ECG, physical and ocular	
GlaxoSmithKline			Exclusions:	examination	
			 Systemic anti-tumor therapy within 14 		
NCT03828292			days, plasmapheresis within 7 days of first	[Up to 2.2 years]	
			dose		
			 Symptomatic amyloidosis, active POEMS 		
			syndrome, active plasma cell leukemia at		
			time of screening		
			 Allogeneic SCT 		
			 Active renal condition, corneal epithelial 		
			disease, active mucosal or internal		
			bleeding, severe or uncontrolled systemic		
			disease, etc.		
A Study of Belantamab	Non-Randomized,	Intervention (Part	Inclusions:	Primary:	May 6,
Mafodotin	Parallel, Phase I	1):	– ECOG status of 0-2	 Max observed plasma 	2024
(GSK2857916) in	Clinical Trial	– Participants with	 Histologically/cytologically confirmed MM; 	concentration	
Multiple Myeloma		normal hepatic	has undergone autologous SCT or	 Predose plasma 	
Participants with	Estimated	function: 2.5	transplant-ineligible; has failed ≥2 prior	concentration	
Normal and Impaired	<u>enrollment</u> : N = 40	mg/kg	lines of therapy including IMiD and PI	 AUC for plasma 	
Hepatic Function		belantamab	 Measurable disease 	concentration-time	
(DREAMM 13)		every 3 weeks	 History of autologous SCT only if 100 days 	 AUC over the dosing 	
		 Participants with 	prior to study enrollment & no active	interval of belantamab	
GlaxoSmithKline		moderate	infections		
		hepatic	 Adequate organ system function 	[throughout 21-day cycle]	

Title / Trial Sponsor	Study Design	Treatment Arms	Patient Population	Outcomes	Estimated Completion
<u>NCT04398680</u>		impairment: 2.5 mg/kg belantamab every 3 weeks Intervention (Part 2): - Patients with severe hepatic function: 2.5 mg/kg or 1.9 mg/kg belantamab	 Exclusions: Active plasma cell leukemia, symptomatic amyloidosis, active POEMS syndrome, Waldenstroem Macroglobulinemia Prior allogeneic SCT Investigational drug or strong organic anion transporting polypeptide inhibitor received 2 weeks prior ≥2 Grade toxicity from previous treatment except alopecia or peripheral neuropathy up to Grade 2 Previous or concurrent malignancies 	Secondary: – Adverse events, serious adverse events [Up to 4 years]	
Study of Single Agent	Randomized,	every 3 weeks Intervention:	unless medically stable for at least 2 years Inclusions:	Drimany	November
Study of Single Agent Belantamab Mafodotin Versus Pomalidomide Plus Low-dose Dexamethasone (Pom/Dex) in Participants with Relapsed/Refractory Multiple Myeloma (RRMM) GlaxoSmithKline <u>NCT04162210</u>	Parallel, Phase 3 Clinical Trial <u>Estimated</u> <u>enrollment</u> : N = 380	 - 2.5 mg/kg belantamab every 3 weeks Comparator: - Pom/Dex (low dose) 	 ECOG status of 0-2 Histological/cytologically confirmed MM Has undergone stem cell transplant/stem cell transplant not deemed feasible Failed at least 2 prior lines of antimyeloma treatment Refractory to an immunomodulatory drug and proteasome inhibitor Measurable disease Adequate organ system functions All prior treatment-related toxicities must be ≤ Grade 1, Grade 2 peripheral neuropathy 	Primary: – PFS [Up to 20 months] Secondary: – OS, ORR, CBR, DOR – Adverse events [Up to 55 months]	November 21, 2024
			 Exclusions: Prior allogenic stem cell transplant Systemic anti-myeloma therapy or investigational drugs within 14 days; plasmapheresis within 7 days 		

Title / Trial Sponsor	Study Design	Treatment Arms	Patient Population	Outcomes	Estimated Completion
A Study of Belantamab	Non-Randomized,	Intervention (Part	 Symptomatic amyloidosis, POEMS syndrome, plasma cell leukemia Active renal condition, unstable liver, or biliary disease Malignancies other than disease under study Inclusions: 	Primary:	March 7,
Mafodotin (GSK2857916) in Multiple Myeloma Participants with Normal and Varying Degree of Impaired Renal Function (DREAMM 12) GlaxoSmithKline <u>NCT04398745</u>	Phase I Clinical Trial <u>Estimated</u> <u>enrollment</u> : N = 36	 1): Patients with normal/mild impaired renal function: 2.5 mg/kg belantamab every 3 weeks Patients with severe renal impairment: 2.5 mg/kg belantamab every 3 weeks Intervention (Part 2): Patients with ESRD (not on dialysis): 2.5 mg/kg or 1.9 mg/kg belantamab every 3 weeks 	 ECOG status of 0-2 Histologically/cytologically confirmed MM; has undergone autologous SCT or transplant-ineligible; has failed ≥2 prior lines of therapy including IMiD and PI Measurable disease History of autologous SCT only if 100 days prior to study enrollment & no active infections Adequate organ system function Exclusions: Active plasma cell leukemia, symptomatic amyloidosis, active POEMS syndrome, Waldenstroem Macroglobulinemia Prior allogeneic SCT Investigational drug, belantamab, strong organic anion transporting polypeptide inhibitor received 2 weeks prior ≥2 Grade toxicity from previous treatment except alopecia or peripheral neuropathy up to Grade 2 Previous or concurrent malignancies unless medically stable for at least 2 years 	 Max. observed plasma concentration Concentration belantamab at end of infusion Pre-dose plasma concentration AUC over the dosing interval of belantamab [Up to 3 21-day cycles] Secondary: Adverse events, serious adverse events [Up to 4 years] 	2025

Title / Trial Sponsor	Study Design	Treatment Arms	Patient Population	Outcomes	Estimated Completion
		mg/kg belantamab every 3 weeks			
Platform Study of	Randomized,	Intervention:	Inclusions:	Primary:	February
Belantamab Mafodotin as Monotherapy and in Combination with Anti-cancer Treatments in Participants with Relapsed/Refractory Multiple Myeloma	Sequential Assignment, Phase I/II Clinical Trial <u>Estimated</u> <u>enrollment</u> : N = 464	 Substudy 1: Belantamab + GSK3174998 Substudy 2: Belantamab + GSK3359609 Substudy 3: Belantamab + 	 3 prior lines of anti-myeloma treatments (including IMiD, PI, anti-CD38 monoclonal antibody) History of autologous stem cell transplant >100 days prior to enrollment ECOG status 0-1 Measurable disease 	 Dose-limiting toxicities Adverse events Abnormality in vital signs Changes in hematology, clinical chemistry, and urinalysis lab parameters 	24, 2028
(RRMM) (DREAMM 5)		Nirogacestat	Exclusions:	– ORR	
GlaxoSmithKline		– Substudy 4: Belantamab + Dostarlimab	 Corneal epithelial disease Prior radiotherapy within 2 weeks, prior allogenic transplant, prior CAR T therapy 	Secondary: – CBR, PR, vgPR, CR, sCR,	
		Active Comparator: – Belantamab monotherapy	 within 3 months, prior investigational agent treatment within 2 weeks Delayed hypersensitivity reaction or idiosyncrasy to drugs chemically similar to Belantamab 	PFS, DOR, OS – Adverse events, serious adverse events [Up to 36 months]	

AUC: area under the curve, BCVA: Best Corrected Visual Acuity, CAR: chimeric antigen receptor, CBR: clinical benefit rate, CNS: central nervous system, CR: complete response, Dex: dexamethasone, DLT: dose limiting toxicities, DOR: duration of response, ECG: echocardiogram, ECOG: Eastern Cooperative Oncology Group, ESRD: end-stage renal disease, IMiD: immunomodulatory drug, kg: kilogram, max: maximal, MM: multiple myeloma, mg: milligram, MRD: minimal residual disease, N: total number, ORR: overall response rate, OS: overall survival, PFS: progression-free survival, PI: protease inhibitor, POEMS: polyneuropathy, organomegaly, endocrinopathy, myeloma protein, and skin changes, Pom: pomalidomide, PR: partial response, sCR: stringent complete rate, SCT: stem cell transplant

Source: www.ClinicalTrials.gov (NOTE: studies listed on site include both clinical trials and observational studies)

D5. Previous Systematic Reviews and Technology Assessments

We identified two ongoing health technology assessments, one of belantamab and one of ide-cel by the National Institute for Health and Care Excellence (NICE) summarized below. We also identified one systematic review and meta-analysis of BCMA CAR-T therapies and one systematic review that aimed to put the ide-cel and cilta-cel clinical trials into an appropriate comparative context.

NICE Technology Assessments

Belantamab mafodotin for treating relapsed or refractory multiple myeloma after 3 therapies [ID2701]

NICE is currently conducting an appraisal of the clinical and cost effectiveness of belantamab in relapsed and refractory multiple myeloma patients who have received three prior therapies. The expected publication date is to be confirmed.

Idecabtagene vicleucel for treating relapsed and refractory multiple myeloma in people who have received at least 3 prior therapies [ID1442]

NICE is currently conducting an appraisal of the clinical and cost effectiveness of ide-cel in relapsed and refractory multiple myeloma patients who have received at least three prior therapies. Comparators in the final scope include pomalidomide-dexamethasone and panobinostatbortezomib-dexamethasone combinations, reflecting a difference in preferred treatment in England as compared to the U.S. The expected publication date is to be confirmed.

Previous Systematic Reviews

Roex G, Timmers M, Wouters K, et al. Safety and clinical efficacy of BCMA CAR-T-cell therapy in multiple myeloma. *J Hematol Oncol*. 2020;13(1):164.

This systematic review and meta-analysis assess the safety and clinical efficacy of BCMA-targeted CAR-T-cell therapies in patients with multiple myeloma. Including a total of 27 clinical studies pertaining to 23 different BCMA CAR-T-cell therapies, it is the most comprehensive review to date. One study for each of the two most advanced therapies (ide-cel and cilta-cel) were identified.

For all BCMA CAR-T patients evaluable for clinical response, high response rates were achieved. Response rates for high dose ide-cel and cilta-cel were comparable (ORR 82% and 88%, respectively). The median PFS among evaluable patients treated with high-dose ide-cel was 12.1 months (95% CI 8.8, 12.3) and 19.9 months for patients treated with cilta-cel (95% CI 9.6, 31.0). Although high response rates were achieved across BCMA CAR-T studies, toxicity was also high. 80.3% patients evaluable for safety experienced CRS with 14.1% experiencing CRS of grade ≥3. High dose ide-cel and cilta-cel had higher than average rates of CRS overall (96.3% and 89.5%, respectively) but lower rates of grade \geq 3 CRS (7.0% and 5.6%, respectively). Rates of neurotoxicity differed considerably between studies. The population included in the ide-cel study was generally older and more heavily pretreated, possibly contributing to its higher neurotoxicity rate.

Overall, BCMA CAR-T-cell therapies prove to have high response rates, but equally as high toxicity. Despite toxicities, this meta-analysis provides robust evidence that BCMA CAR-T therapies are considered highly efficacious even in heavily pretreated MM patients.

Shah N, Sussman M, Crivera C, et al. Comparative effectiveness research for CAR-T therapies in multiple myeloma: appropriate comparisons require careful considerations of data sources and patient populations. *Clin Drug Investig.* Published online February 18, 2021. https://doi.org/10.1007/s40261-021-01012-x

This systematic literature review sought to evaluate the most appropriate data sources and populations for comparison to the novel CAR-T therapies ide-cel and cilta-cel, specifically their single-arm KarMMa and CARTITUDE-1 trials. Investigators reviewed clinical trials of regimens preferred by the National Comprehensive Cancer Network (NCCN) for previously treated multiple myeloma and assessed them for comparability to the CAR-T trials' patient populations. None of the clinical trials identified were conducted in patient with triple class exposed (TCE) or triple class refractory (TCR) multiple myeloma exclusively and thus were not suitable for comparison. Additionally, investigators systematically reviewed real-world studies of patients with TCE and/or TCR multiple myeloma. Five real-world studies were included, with two exclusively focusing on the TCR patient population. Based on currently available data, real world studies with matching TCE and/or TCR patient populations make the most appropriate comparators for the single-arm trials of ide-cel and cilta-cel. Of note, the two real world studies identified by this systematic review, Gandhi 2019⁸ and Mehra 2020²⁶, were included to define the comparator population in this report.

E. Long-Term Cost-Effectiveness: Supplemental Information

E1. Detailed Methods

Table E.1. Impact Inventory

Sector	Type of Impact (Add additional domains, as	Included Analysis fr Perspec	om []	Notes on Sources (if quantified), Likely Magnitude	
	relevant)	Health Care Sector	Societal	& Impact (if not)	
Formal Health C	Care Sector		-	-	
Health	Longevity effects	Х	Х		
Outcomes	Health-related quality of life	Х	Х		
	effects				
	Adverse events	Х	Х		
Medical Costs	Paid by third-party payers	Х	Х		
	Paid by patients out-of-pocket				
	Future related medical costs	Х	Х		
	Future unrelated medical costs				
Informal Health	Care Sector				
Health-	Patient time costs	NA			
Related Costs	Unpaid caregiver-time costs	NA			
	Transportation costs	NA	Х		
Non-Health Car	e Sector	•			
Productivity	Labor market earnings lost	NA	Х		
	Cost of unpaid lost productivity	NA	Х		
	due to illness				
	Cost of uncompensated household	NA			
	production				
Consumption	Future consumption unrelated to health	NA			
Social services	Cost of social services as part of intervention	NA			
Legal/Criminal	Number of crimes related to	NA			
Justice	intervention				
	Cost of crimes related to	NA			
	intervention				
Education	Impact of intervention on educational achievement of population	NA			

Housing	Cost of home improvements,	NA	
	remediation		
Environment	Production of toxic waste pollution	NA	
	by intervention		
Other	Other impacts (if relevant)	NA	

NA: not applicable

Adapted from Sanders et al⁷⁵

Target Population

The model focused on an intention-to-treat analysis, with a hypothetical cohort of patients in two different populations: 1) triple- or quad-refractory with 3+ prior lines of treatment being treated with ide-cel or cilta-cel, and 2) triple-, quad-, penta-refractory with 4+ prior lines of treatment being treated with belantamab. Cohort characteristics for each treatment group are described in E.1.2 and E.1.3.

Table E.1.2. Baseline Population Characteristics: Triple- or Quad-Refractory MM (3+ prior lines of treatment)

Triple-Class Refractory MM	Idecabtagene vicleucel	Ciltacabtagene autoleucel	Triple and Quad- Refractory Comparator
Median Age	61	61	60
Percent Male	59%	59%	57%
Refractory Status, %	Anti-CD38 Ab-refractory: 94% Triple-refractory: 84%	Triple-refractory: 88%	Anti-CD38 Ab-refractory: 100%
Median Prior Lines of Treatment	6	6	4
Source	Munshi et al, 2020 ¹¹	Madduri et al, 2020 ¹⁵	Gandhi et al, 2019 ⁸

Table E.1.3. Baseline Population Characteristics: Triple-, Quad-, or Penta- Refractory MM (4+ prior
lines of treatment)

Penta-Refractory MM	Belantamab	Triple/Quad/Penta-Refractory Therapies
Median Age	65	59
Percent Male	53%	56%
Refractory Status, %	Anti-CD38 Ab-refractory: 100% in 2.5 mg/kg dose	Anti-CD38 Ab-refractory: 100%
Median Prior Lines of Treatment	7	5
Source	Lonial et al, 2020 ⁴³	Gandhi et al, 2019 ⁸

Comparators

The primary comparator for each type of therapy is listed below. Because of differences in population indications, interventions were not compared to each other. Given the numerous available therapies used by clinicians at various lines of therapy, a market basket approach was used to compare to each intervention based on refractory status using the MAMMOTH study.⁸ The market basket composition was approximated by both broad-therapy and specific-therapy estimations. For belantamab, we used a weighted average of MAMMOTH subcohorts so that the proportion of penta vs triple/quad refractory patients from the MAMMOTH comparator cohort matched that in the DREAMM-2 study. MAMMOTH results for these cohorts were used to inform comparator PFS and OS in the model. Specific regimens that are commonly employed in the relevant populations were also identified for the purpose of estimating market basket costs. For the triple- or quad-refractory population, the comparator market basket included (comparator for ide-cel and cilta-cel, sub-population within belantamab also compared using this market basket):

- Carfilzomib + cyclophosphamide + dexamethasone (KCd)
- Pomalidomide + cyclophosphamide + dexamethasone (PCd)
- Carfilzomib + pomalidomide + dexamethasone (KPd)
- Elotuzumab + pomalidomide + dexamethasone (EPd)
- Ixazomib, lenalidomide, and dexamethasone (IRd)

Comparator market basket for the penta-refractory population (used in weighted average comparator basket along with triple/quad for belantamab):

- Carfilzomib + cyclophosphamide + dexamethasone (KCd)
- Pomalidomide + cyclophosphamide + dexamethasone (PCd)
- Ixazomib, pomalidomide, and dexamethasone (IPd)
- Elotuzumab + pomalidomide + dexamethasone (EPd)
- Ixazomib, lenalidomide, and dexamethasone (IRd)
- Bendamustine, prednisone, dexamethasone, cyclophosphamide, etoposide, and cisplatin (DCEP)

Within each progressed state, we also applied a proportion of patients on palliative chemotherapy consistent with current evidence. Tables E.2.2 and E.2.3 provide dosing, administration schedules, and costs for each market basket of comparators.

E2. Model Inputs

Model inputs were estimated from the clinical review, published literature, and information from expert stakeholders. The inputs that informed the model are described below. The base case

analysis took a health care system perspective and focused on direct medical care costs only. Outcomes were estimated over a lifetime time horizon using a monthly cycle to capture the potential impacts of short-term and ongoing morbidity and mortality. Costs and outcomes were discounted at 3% per year.

Treatment Regimen Inputs

Treatment regimens for the interventions are described in Tables E2.1. The market basket of therapy regimens for three or more lines of therapy and four or more lines of therapy are described in tables E2.2 and E2.3.

Category/Therapy	Idecabtagene vicleucel	Ciltacabtagene autoleucel	Belantamab
Brand Name	Abecma	TBD	Blenrep
Manufacturer	Bristol Myers Squibb/bluebird bio	Janssen/Legend Biotech	GlaxoSmithKline
Route of Administration	Single infusion	Single infusion	Infusion once every 3 weeks
Dosing	150-450 x 10º CAR+ T cells	0.75 x 10 (range: 0.5 - 1.0 x10) CAR+ T cells/kg	2.5 mg/kg of body weight
Use of Lymphocyte Depleting Chemotherapy	Fludarabine (30 mg per square meter of body surface per day) and cyclophosphamide (300 mg per square meter per day) on days -5, -4, -3	Cyclophosphamide (300 mg per square meter per day) on days -5, -4, -3	N/A

Table E2.1.Treatment Regimen Recommended Dosage

Table E2.2. Triple- or Quad-Refractory MM Comparator Market Basket (3+ prior lines of treatment) Recommended Dosage and Total Cost

Therapy Combination	Days/Cycle	Cycle 1 Dose	Admin. Days	Proportion of Patients on Each Combination	Per Cycle Cost
Carfilzomib, Cyclophosphamide, Dexamethasone				18%	\$22,676
Carfilzomib	28	20 mg/m2	1, 2, 8, 9, 15, 16 (cycle 1);		
			1, 2, 15, 16 (remaining cycles)		
Cyclophosphamide	28	300mg/m2	1,8,15		
Dexamethasone	28	20 mg	1, 2, 8, 9, 15 and 16		
Pomalidomide, Cyclophosphamide, Dexamethasone				34%	\$17,083
Pomalidomide	28	4mg/day	1-21		
Cyclophosphamide	28	300mg/day	1, 8, 15, 22		
Dexamethasone	28	40mg/day	1-4 and 15-18		
Carfilzomib, Pomalidomide, Dexamethasone				18%	\$30,471
Carfilzomib	28	20mg/m2; 27 mg/m2	1, 2; 8, 9, 15, 16 (cycle 1) 1, 2, 8, 9, 15, 16 (cycles 2-12)		
Pomalidomide	28	4mg/day	1-21		
Dexamethasone	28	20 mg	1,8,15,22		
Elotuzumab, Pomalidomide, Dexamethasone		•	·	19%	\$36,676
Elotuzumab	28	10 mg/kg	1, 8, 15, and 22		
Pomalidomide	28	4mg/day	1-21		
Dexamethasone	28	28mg oral+8mg iv	1,8,15,22		
Ixazomib, Lenalidomide, Dexamethasone				10%	\$22,391
Ixazomib	28	4mg/day	1,8,15		
Lenalidomide	28	25mg/day	1-21		
Dexamethasone	28	40mg/day	1,8,15,22		
Weighted average administration costs					\$3,130
Weighted average adverse event cost management (applied for 2 cycles)					\$1,531
Weighted average dose intensity (applied after 2 cycles)					96%
Weighted average total cost					\$29,490

Table E.2.3. Triple-, Quad-, or Penta- Refractory MM Comparator Market Basket (4+ prior lines of treatment) Recommended Dosage and Total Cost

Therapy Combination	Days/Cycle	Cycle 1 Dose	Admin. Days	Proportion of Patients on Each Combination	Per Cycle Cost
Carfilzomib, Cyclophosphamide, Dexamethasone				22%	\$22,456
Carfilzomib	28	20 mg/m2	1, 2, 8, 9, 15, 16 (cycle 1);		
1, 2, 15, 16 (remaining cycles)					
Cyclophosphamide	28	300mg/m2	1,8,15		
Dexamethasone	28	20 mg	1, 2, 8, 9, 15 and 16		
Pomalidomide, Cyclophosphamide, Dexamethasone				17%	\$17,083
Pomalidomide	28	4mg/day	1-21		
Cyclophosphamide	28	300mg/day	1, 8, 15, 22		
Dexamethasone	28	40mg/day	1-4 and 15-18		
Ixazomib, Pomalidomide, Dexamethasone				9%	\$27,323
Ixazomib	28	4mg/day	1,8,15		
Pomalidomide	28	4mg/day	1-21		
Dexamethasone	28	40mg/day	1,8,15,22		
Elotuzumab, Pomalidomide, Dexamethasone				13%	\$36,676
Elotuzumab	28	10 mg/kg	1, 8, 15, and 22		
Pomalidomide	28	4mg/day	1-21		
Dexamethasone	28	28mg oral+8mg iv	1,8,15,22		
Ixazomib, Lenalidomide, Dexamethasone				10%	\$22,393
Ixazomib	28	4mg/day	1,8,15		
Lenalidomide	28	25mg/day	1-21		
Dexamethasone	28	40mg/day	1,8,15,22		
Bendamustine, prednisone, dexamethasone, cyclophosphamide, etoposide, and cisplatin				29%	\$9,907
Bendamustine	28	90mg/m2			
Prednisone			1,2,3,4		
Dexamethasone	28	40mg/day	1,2,3,4		
Cyclophosphamide	28	400mg/m2	1,2,3,4		
Etoposide	28	40mg/m2	1,2,3,4		

Cisplatin	28	10mg/m2	1,2,3,4	
Weighted average administration costs				\$1,301
Weighted average adverse event cost				\$1,259
management (applied for 2 cycles)				
Weighted average dose intensity (applied after 2				96%
cycles)				
Weighted average total cost				\$22,994

Detailed Description of Curve Digitization and Survival Extrapolation

Kaplan-Meier curves from the evidence were digitized using the algorithm by Guyot and colleagues to impute patient-level time-to-event data.⁷⁶ Base-case survival was derived from parametric fits to each intervention's available PFS and OS Kaplan-Meier curves.^{12,15,43} Table E.2.4 delineates the evidence that was used to calculate transition probabilities. The comparator OS evidence was derived from the MAMMOTH study using triple- and quad-refractory curves (for comparison to CAR-Ts) and a weighted average of triple-/quad-refractory and penta-refractory curves (for comparator populations (i.e., triple-, quad-, or penta-refractory in Gandhi et al.), we assumed similar distribution parameters as other curves in triple-, quad-, or penta-refractory MM and adjusted to fit a PFS to OS relationship as observed in meta-analyses.^{36,37}

The model curves considered included distributional forms Weibull, exponential, log-normal, loglogistic, and Gompertz. The base-case distributional form was selected separately for each curve based on the best model fit using the Akaike information criterion (AIC) values, visual comparison, and clinical plausibility. Transition probabilities were then calculated for each time period in the model (monthly cycles). In some cases, we used piecewise modeling techniques to fit survival distributional forms at different time points after examining hazard functions.⁴⁶ Table E.2.4 presents sources for each curve.

Survival Estimate	ldecabtagene vicleucel	Ciltacabtagene autoleucel	Triple- and Quad- Refractory (Comparator to CAR-Ts)*	Belantamab	Penta-Refractory (Comparator to Belantamab)*
Progression- free survival	Phase II KarMMa results	CARTITUDE-1	PFS curve fit to proportional relationship reported in previous meta- analyses	Updated DREAMM-2 PFS curve	PFS curve fit to proportional relationship reported in previous meta-analyses
Overall survival Sources	Phase II KarMMa results Munshi et al, 2021 ¹²	CARTITUDE-1 Madduri et al, 2020 ¹⁵	MAMMOTH triple/quad- refractory Gandhi et al, 2019 ⁸	Updated DREAMM-2 OS curve Academic in confidence	MAMMOTH penta- refractory Gandhi et al, 2019 ⁸

* Comparator to belantamab weights outcomes from both populations exposed to three or more lines of therapy and populations exposed to four or more lines of therapy

Table E.2.5 presents the final distributions chosen for the model based on visual inspection and Akaike information criterion (AIC). The shape and scale parameters were used to generate time-dependent transition probabilities for each curve. We provide alternative extrapolations to approximated KM curves in Figures E.4.1 to E.4.4. Figures showing belantamab extrapolations are not shown due to academic in confidence considerations. Extrapolations based on assumptions from other curve fittings are also not shown (i.e., Cilta-cel OS, MAMMOTH PFS).

	Outcome (Distribution Chose)	AIC	Shape/ Dist 1	Scale/ Dist 2	Source	Notes	Modeled % at 12 months
lde-Cel	Progression Free Survival (Log-normal)	185.9	2.31	1.00	Munshi et al, 2021 ¹²	N/A	40%
	Overall Survival (Log- normal)	204.7	3.24	0.93	Munshi et al, 2021 ¹²	N/A	78%
Cilta-Cel	Progression Free Survival (Weibull)	218.6	1.61	29.24	Madduri et al, 2020 ¹⁵	Median PFS not reached, therefore calibrated based on % alive and PFS at 12 months	77%
	Overall Survival (Weibull)	218.6	1.61	76.49	Madduri et al, 2020 ¹⁵	Assumed same shape as PFS curve with adjustment to scale parameter	94%
Triple- or Quad- Refractory Comparator	Progression Free Survival (Log-normal)	N/A	1.22	1.00	Dimopoulos et al, 2017 ³⁶ Gandhi et al, 2019 ⁸ Felix et al, 2013 ³⁷	PFS curve fit to proportional relationship reported in previous meta-analyses	10%
	Overall Survival (Log- normal)	426.5	2.13	1.30	Gandhi et al, 2019 ⁸	Figure 1a triple/quad refractory	39%

Table E.2.5. Survival Curve Fit, Shape, and Scale Parameters for Final Model

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	Outcome (Distribution Chose)	AIC	Shape	Scale	Source	Notes	Modeled % at 12 Months
Belantamab*	Progression Free Survival (Log-normal)	Academic in confidence Academic in confidence					
	Overall Survival (Log- normal)						
Penta-Refractory Comparator	Progression Free Survival (Log-normal)	N/A	1.0	0.71	Dimopoulos et al, 2017 ³⁶ Gandhi et al, 2019 ⁸	PFS curve fit to proportional relationship reported in previous meta-analyses	1.5%
	Overall Survival (Weibull)	329.0	1.53	7.61	Gandhi et al, 2019 ⁸	Figure 1a penta- refractory	11%

*Comparator to belantamab weights outcomes from both populations exposed to three or more lines of therapy and populations exposed to four or more lines of therapy

Progression Free Survival

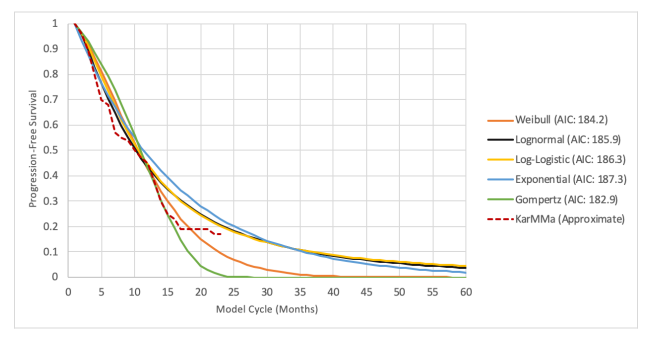
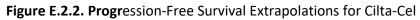
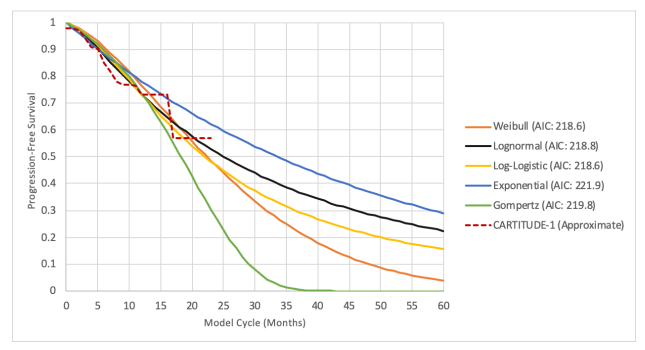


Figure E.2.1. Progression-Free Survival Extrapolations for Ide-Cel





Overall Survival

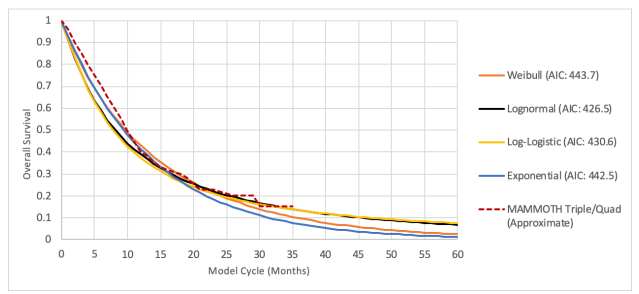


Figure E.2.3. Overall Survival Extrapolations for MAMMOTH Triple- or Quad-Refractory Cohort*

*Gompertz not shown due to poor fit of observed data from MAMMOTH

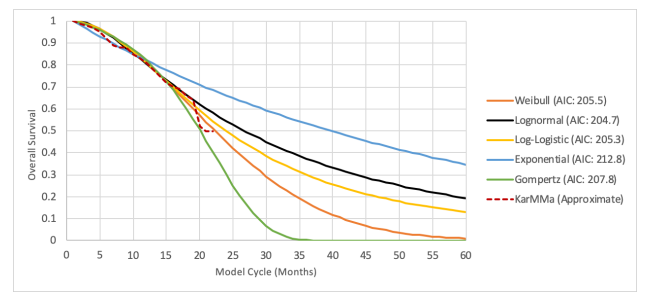


Figure E.2.4. Overall Survival Extrapolations for Ide-Cel

Health State Utilities

The most current and best available evidence on health utilities comes from the KarMMa study, with elicitation of utilities using the European Quality of Life-5 dimensions 5 levels (EQ-5D-5L) health state classification instrument.⁴¹ The analysis elicited utilities from the US, UK, and Canadian populations across different time points including baseline, pre-progression, and post-progression.

We applied the baseline utility value to the progressed state, the highest utility value elicited for the progression-free off therapy state, and the first month pre-progression overall utility for the progression-free on therapy state. Given feedback from RRMM patients, we applied a separate utility to the progression-free off therapy state for both ide-cel and cilta-cel to reflect the benefits of being off therapies for a disease that commonly continues patients on therapies until death. See table E.2.6 for health state utilities applied in the model.

Adverse event disutilities were applied for two cycles in the model (i.e., two months) as evidence suggested most adverse events were resolved within 1-3 cycles with additional dose adjustments. Consistent with previous health technology assessments,^{77,78} a utility score of 0 was applied for grade 3 or higher cytokine release syndrome for a duration. Table E.2.7 details the disutility estimates applied for adverse event disutilities.

Parameter	Three or More Lines of Therapy
Progression-free on Therapy and Responding	0.78
Progression-free off Therapy and Responding	0.82
Progressed Disease/not Responding	0.71
to Treatment	
Source	Delforge et al, 2020 ⁴⁰

Table E.2.6. Utility Values for Health States

<u>Adverse Events</u>

The model included any grade 3/4 adverse events that occur in 5% of patients in any of the treatments and comparators. Given the potentially significant impact of cytokine release syndrome on health care resource utilization and quality of life, we included all grades 1-4 for these adverse events and adjusted costs and quality of life estimates accordingly. The costs and disutility of adverse events were applied to the first two cycles for each intervention and comparator. After cycle 2 of the model, we applied a dose adjustment factor, assuming adverse events would be resolved with lower dosing of each therapy. Table E.2.7 includes the proportions of patients with adverse events applied in the model.

Parameter	Idecabtagene vicleucel ¹¹	Ciltacabtagene autoleucel ¹⁵	Belantamab 43
Proportion with	41%	49%	N/A
grade 1 CRS			
Proportion with	23%	39%	N/A
grade 2 CRS			
Proportion with	2%	3%	N/A
grade 3 CRS			
Proportion with	1%	1%	N/A
grade 4 CRS			
Proportion with	12%	NR	N/A
CRS and NE <=2			
Proportion with	3%	NR	N/A
CRS>= 3 and NE <=2			
Proportion with	3%	NR	N/A
CRS<=2 and NE=>3			
Anemia	See above categories	68%	20%
Neutropenia		95%	9.5%
Thrombocytopenia		60%	20%
Lymphopenia		49%	16.8%
Leukopenia		61%	
Keratopathy	1	N/A	27%
Hypercalcemia	1	NR	7.4%
Hypophosphatemia		NR	5.3%

NR: Not reported

Adverse event disutilities or utilities are described in Table E.2.8. Adverse event disutilities were applied for two cycles in the model (i.e., two months). Consistent with previous health technology assessments,^{77,78} a utility score of 0 were applied for grade 3 or higher cytokine release syndrome for 8 days in the first cycle.

Table E.2.8. Adverse Event Disutilities

Adverse Event Parameter	Disutility	Source
Anemia	-0.31	Brown et al. 2013 79
Neutropenia	-0.15	Brown et al. 2013 79
Thrombocytopenia	-0.31	Brown et al. 2013 79
Lymphopenia	-0.07	NICE TA 510 ⁸⁰
Cytokine release syndrome	0.00*	Hettle et al. 2017 77
Keratopathy	-0.05	Sullivan 2006 (ICD-9 369) 81
Hypercalcemia	-0.04	Sullivan 2006 (ICD-9 289) 81
Hypophosphatemia	-0.04	Sullivan 2006 (ICD-9 289) 81

*This value corresponds to a utility, not a disutility.

Economic Inputs

Cost Inputs

All costs used in the model were updated to 2020 dollars.

Drug Acquisition Costs

For CAR-T therapies, the base-case findings use the list price for ide-cel. There is not a price yet for cilta-cel; therefore, we assume the same price as a placeholder. For belantamab we used WAC pricing for the base-case findings. Comparator therapy prices were a function of one or more therapies on the market, inclusive of discounts, rebates (15% discount for comparator oral therapies based on FSS pricing schedule), patient assistance programs, and concessions to wholesalers and distributors. Patients that discontinued the CAR-T treatment before receiving the CAR-T infusion were not charged the CAR-T costs. Costs for subsequent therapies, including a proportion on palliative care, were assigned to the progressed state for 4 cycles using the appropriate comparator therapies for each population. The progressed state costs were consistent across treatment comparisons. Infusion therapies were subject to ASP + 6% pricing.

Table E.2.9. Drug Costs

Intervention (Dosage)	WAC/List Price per Unit or per Time Period*	Net Price per Unit or per Time Period	Source
Idecabtagene vicleucel	\$419,500	N/A	Market analyst estimates ⁸²
Ciltacabtagene autoleucel	Assumed same as ide- cel	N/A	Assumption
Belantamab	\$8,277 per 100mg package		Micromedex Solutions ⁴⁵
Comparator therapies	See Table E.2.3 and Table E.2.4	See Table E.2.3 and Table E.2.4	Multiple

*WAC as of March 25, 2021

Administration and Monitoring Costs

Tables E.2.10 through E2.12 detail administration and monitoring utilization and costs applied in the model. Table E.2.10 includes pre-infusion regimens and unit prices for CAR-T therapies. Table E.2.11 includes administration and monitoring utilization applied at different stages of the model. We used recent evidence in heavily pre-treated patients with multiple myeloma to inform average utilization inputs per cycle.⁸³ We then applied unit prices from Table E.2.12 to each utilization parameter estimate. For hospital admissions we applied a fee-for-service approach.

Table E.2.10. Pre-Infusion Regimens for CAR-T Therapies

Treatment	Regimen	Unit Price	Source
Cyclophosphamide	300 mg/mg(2) on days -5, -4, -3	\$33	
Fludarabine	30 mg/m(2) on days -5, -4, -3	\$50	Munshi et al,
Cytarabine	500 mg/m(2) for 2 days a week, 2 weeks total	\$1	202011
Methotrexate	1000 mg/m(2) for 1 day a week, 2 weeks total	\$2	

Model Stage	Idecabtagene vicleucel	Ciltacabtagene autoleucel	Belantamab	Comparator Market Basket of Therapies
Prior to and during therapy administration	 Leukapheresis CRS-related treatment Inpatient days (2 in ICU) Neurotoxicity adverse events 	 Leukapheresis CRS-related treatment Inpatient days (2 in ICU) Neurotoxicity adverse events 	 Ophthalmic examinations at baseline and prior to each dose Other AE- related costs 	 IV administration costs Other AE-related costs
Post-therapy monitoring; progression-free ⁸³	 Complete blood count testing Liver function testing 	 Complete blood count testing Liver function testing 	 Treatment- specific outpatient visits per cycle Complete blood count testing each outpatient visit Liver function testing 	 Treatment-specific outpatient visits per cycle Complete blood count testing each outpatient visit Liver function testing
Progressed disease ⁸³	 4 cycles of subsequent treatment administration with market basket Complete blood count testing Liver function testing 	 4 cycles of subsequent treatment administration with market basket Complete blood count testing Liver function testing 	 4 cycles of subsequent treatment administration with market basket Complete blood count testing Liver function testing 	 4 cycles of subsequent treatment administration with market basket Complete blood count testing Liver function testing

Table E.2.11. Administration and Monitoring Utilization

	Value	Source
Cost per hospital day*	\$3,190	HCUP Statistical Brief #125 ⁸⁴
Cost per day in ICU	\$5,563	Dasta, 2005 ⁸⁵
Office visit	\$74	Physicians' Fee and Coding Guide (HCPCS code 99213) ⁸⁶
Leukapheresis (CAR-T only)	\$1,323	Physicians' Fee and Coding Guide HCPCS code 36511 ⁸⁶
Intravenous treatment administration (first hour)	\$140	Physicians' Fee and Coding Guide (HCPCS code 96413) ⁸⁶
Intravenous treatment administration (each additional hour)	\$29	Physicians' Fee and Coding Guide (HCPCS code 96415) ⁸⁶
Visual acuity test	\$31	Physicians' Fee and Coding Guide (HCPCS code 99173) ⁸⁶
Complete blood count test	\$44	Physicians' Fee and Coding Guide (HCPCS code 85027) ⁸⁶
Slit lamp exam	\$110	Physicians' Fee and Coding Guide (HCPCS code 92285) ⁸⁶
Liver function test	\$62	Physicians' Fee and Coding Guide (HCPCS code 80076) ⁸⁶

Table E.2.12. Other Administration and Monitoring Unit Prices

*Inflated to 2020 USD

Adverse Event Costs

The unit cost of adverse events applied to patients experiencing these events are shown in Table E.2.13. Adverse event costs were applied for the first two cycles of the model. Specific to CAR-T therapies, we relied on recent evidence that combined CRS and neurotoxicity events in different categories.²⁷

Table E.2.13. Adverse Event Unit Costs

Adverse Event Parameter	Mean Cost	Source
Anemia	\$2,007	Roy et al. 2015 ⁸⁷
Neutropenia	\$1,791	Roy et al. 2015 ⁸⁷
Thrombocytopenia	\$1,764	Roy et al. 2015 ⁸⁷
Leukopenia	\$3,045	Roy et al. 2015 ⁸⁷
Lymphopenia	\$3,102	Roy et al. 2015 ⁸⁷
Cytokine release syndrome	\$18,500 (grade 1) - \$121,500 (grade 4)	Hari et al. 2020 ²⁷
Hypercalcemia	\$193	Roy et al. 2015 ⁸⁷
Hypophosphatemia	\$193	Roy et al. 2015 ⁸⁷

*Inflated to 2020 USD

Indirect Costs

A modified societal perspective was explored in a scenario analysis, and the below inputs informed that analysis to assess the impact on model outcomes.

Table E.2.14. Indirect Cost Inputs for Modified Societal Perspective
--

Category	Mean	Source
Average hourly wage	\$25.72	US BLS ⁸⁸
Transportation cost per administration*	\$18.52	Ailawadhi et al. Clin Ther 2019 ⁸⁹
Patient workdays missed per administration	3 hours lost for each medical visit; 103 hours lost across 6 different regimens (average 17.17 hours)	Ailawadhi et al. Clin Ther 2019 ⁸⁹
Employment rate	60.8%	US BLS ⁸⁸

Description of evLYG Calculations

The cost per evLYG considers any extension of life at the same "weight" no matter what treatment is being evaluated. Below are the stepwise calculations used to derive the evLYG.

- 1. First, we attribute a utility of 0.851, the age- and gender-adjusted utility of the general population in the US that are considered healthy. ⁹⁰
- For each cycle (Cycle I) in the model where using the intervention results in additional years of life gained, we multiply this general population utility with the additional life years gained (ΔLYG).
- 3. We sum the product of the life years and average utility (cumulative LYs/cumulative QALYs) for Cycle I in the comparator arm with the value derived in Step 2 to derive the equal value of life years (evLY) for that cycle.

- 4. If no life years were gained using the intervention versus the comparator, we use the conventional utility estimate for that Cycle I.
- 5. The total evLY is then calculated as the cumulative sum of QALYs gained using the above calculations for each arm.
- 6. We use the same calculations in the comparator arm to derive its evLY.

Finally, the evLYG is the incremental difference in evLY between the intervention and the comparator arms.

E3. Undiscounted Results

Tables E.3.1 – E.3.3 present undiscounted results for all interventions and comparators.

Table E.3.1. Base-Case Undiscounted Results for Ide-Cel Compared to Population to Triple- orQuad-Refractory MM Comparator Market Basket (3+ prior lines of treatment)

Treatment	Intervention	Other non-	Total	OALVe	Life	avilVCa	Incremental Results		ılts
Treatment	Cost	intervention costs	Cost	QALYs	Years	evLYGs	Cost/QALY gained	Cost/LY gained	Cost per evLYG gained
lde-Cel	\$466,000	\$188,000	\$654,000	2.47	3.30	2.66	\$224,000	\$286,000	\$251,000
CAR-T Comparator Market Basket	\$157,000	\$127,000	\$284,000	1.18	1.64	1.19		-	

evLYG: equal-value of life years gained, QALYs: quality-adjusted life years gained

Table E.3.2. Preliminary Results for the Base-Case for Cilta-Cel Compared to Triple- or Quad-Refractory MM Comparator Market Basket (3+ prior lines of treatment)

	Intervention	Other non-	Total	O ALV-	Life		Incremental Results		ults
Treatment	Cost	intervention costs	Cost	QALYs	Years	evLYGs	Cost/QALY gained	Cost/LY gained	Cost per evLYG gained
Cilta-Cel*	\$445,000	\$184,000	\$629,000	3.80	5.06	4.19	\$132,000	\$101,000	\$115,000
CAR-T Comparator Market Basket	\$157,000	\$127,000	\$284,000	1.18	1.64	1.19		-	

*using placeholder price for cilta-cel

evLYG: equal-value of life years gained, QALYs: quality-adjusted life years gained

Table E.3.3. Results for the Base-Case for Belantamab Compared to Triple-, Quad-, or Penta-
Refractory MM Comparator Market Basket (4+ prior lines of treatment)

	Intervention	Other non-	Total		Life		Increr	nental Res	ults
Treatment	Cost	intervention costs	Cost	QALYs	Years	evLYGs	Cost/QALY gained	Cost/LY gained	Cost per evLYG gained
Belantamab	\$155,000	\$105,000	\$260,000	1.21	1.68	1.24	\$95,000	\$68,000	\$90,000
Belantamab Comparator Market Basket	\$120,000	\$101,000	\$221,000	0.83	1.15	0.85		-	

evLYG: equal-value of life years gained, QALYs: quality-adjusted life years gained

E4. Sensitivity Analyses

To demonstrate effects of uncertainty on both costs and health outcomes, we varied input parameters using available measures of parameter uncertainty (i.e., standard errors) or reasonable ranges to evaluate changes in cost per QALY.

We conducted one-way sensitivity analyses to identify the impact of parameter uncertainty and key drivers of model outcomes. Probabilistic sensitivity analyses were also be performed by jointly varying model parameters over 5000 simulations, then calculating 95% credible range estimates for each model outcome based on the results. We also performed threshold analyses for drug costs across a range of incremental cost-effectiveness ratios (\$50,000, \$100,000, \$150,000, and \$200,000 per QALY and evLYG).

Important input parameter drivers of model findings include progression-free survival for the active interventions, the unit price of the comparator market basket of therapies, and health state utility

values. When varying PFS, we assumed the same proportional relationship in terms of gains in OS. In the case of Ide-cel (Figure E.4.1 and Table E.4.1), higher PFS (and thus higher OS) leads to more favorable cost-effectiveness estimates but still do not meet commonly cited cost-effectiveness thresholds whereas lower PFS leads to less favorable cost-effectiveness estimates that do not meet commonly cited cost-effectiveness thresholds. When changing the price of the comparator market basket to the lower range, we find increases in the incremental cost-effectiveness ratio, and separately, decreases in the incremental cost-effectiveness ratio when the price of the comparator market basket is at the higher range. Utility of PFS (off treatment) was also a driver that led to increases in the incremental cost-effectiveness ratio at upper levels of the PFS utility (off treatment) and decreases in the incremental cost-effectiveness ratio at lower levels of utility. After varying multiple inputs simultaneously while running multiple iterations of the model, we found Ide-cel ranged from a lower bound incremental-cost effectiveness ratio that meets commonly cited costeffectiveness thresholds to significantly higher incremental cost-effectiveness ratios above \$1,000,000 per QALY (Table E.4.2 and Figure E.4.2).

Finally, given the sensitivity of progression-free and overall survival on model outcomes, we provide cost-effectiveness estimates using alternative survival extrapolations (Table E.4.3). Varying the distribution used to extrapolate survival resulted in incremental cost-effectiveness ratios of -36% from the base-case to above +490%. This sensitivity analysis should be interpreted with caution as some distributions may provide implausible estimates of long-run survival.

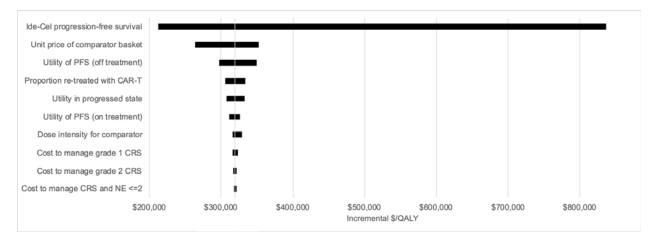


Figure E.4.1. Tornado Diagram for Ide-Cel

Table E.4.1. Tornado Diagram Inputs and Results for Ide-Cel versus Triple- or Quad-RefractoryMM Comparator Market Basket (3+ prior lines of treatment)

	Lower Cost- Effectiveness Estimate	Upper Cost- Effectiveness Estimate	Lower Input*	Upper Input*
Ide-Cel	\$212,478	\$836,482	6	12
progression-free survival				
Unit price of comparator basket	\$264,180	\$352,644	\$17,083	\$37,508
Proportion re- treated with CAR-T	\$305,420	\$333,805	0.16	0.24
Utility in progressed state	\$307,436	\$332,635	0.64	0.78
Utility of PFS (on treatment)	\$311,466	\$326,151	0.70	0.85
Dose intensity for comparator	\$315,781	\$329,000	0.83	1.00
Cost to manage grade 1 CRS	\$315,981	\$323,069	\$10,573	\$28,601
Cost to manage grade 2 CRS	\$316,896	\$321,902	\$13,309	\$36,005
Cost to manage CRS and NE <=2	\$317,460	\$321,182	\$18,967	\$51,310

*Note lower input may reflect either upper or lower ICER value depending on the direction that the input has on the ICER output.

Table E.4.2. Results of Probabilistic Sensitivity Analysis for Ide-Cel versus Triple- or Quad-Refractory MM Comparator Market Basket (3+ prior lines of treatment)

	lde-Cel		Compa	ator Basket	Incr	emental	
	Mean	Credible Range	Mean		Mean	Credible Range	
Total							
Total Costs	\$642,000	\$500,000 - \$710,000	\$277,000	\$226,000 - \$340,000	\$365,000	\$208,000 - \$454,000	
Total QALYs	2.30	1.30 - 3.67	1.08	1.00-1.15	1.22	0.20-2.55	
ICER					\$93,000	\$138,000 - \$1,200,000	

Figure E.4.2. Incremental Cost-Effectiveness Cloud for Ide-Cel versus Triple- or Quad-Refractory MM Comparator Market Basket (3+ prior lines of treatment)

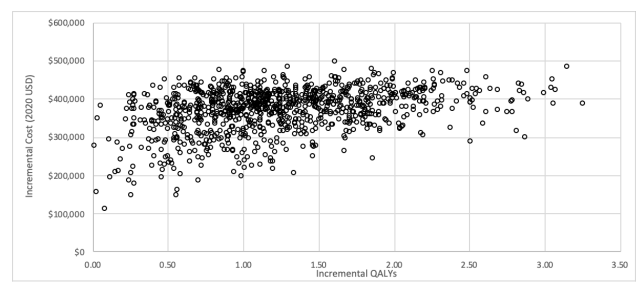


Table E.4.3. Incremental Cost-Effectiveness Ratios under Alternative Survival Extrapolations for
lde-Cel

Curve	Distribution	Incremental Cost- Effectiveness Ratio versus Comparator (per QALY)	Relative percentage difference from base-case
PFS			
	Log-normal (base-case)	\$319,000	Reference
	Exponential	\$320,000	+0.3%
	Log-logistic	\$320,000	+3.4%
	Weibull	\$330,000	+3.4%
	Gompertz	\$332,000	+4.0%
Overall Survival			
	Log-normal (base-case)	\$319,000	Reference
	Exponential	\$203,000	-36%
	Log-logistic	\$410,000	+28%
	Weibull	\$887,000	+178%
	Gompertz	\$1,900,000	+495%

In the case of cilta-cel (Figure E.4.3 and Table E.4.4), higher PFS (and thus higher OS) leads to more favorable cost-effectiveness estimates that meet commonly cited cost-effectiveness thresholds whereas lower PFS leads to less favorable cost-effectiveness estimates that do not meet commonly cited cost-effectiveness thresholds. Changing the price of the comparator market basket to the upper range, we find decreases in the incremental cost-effectiveness ratio that meet commonly cited cost-effectiveness ratios and increases in the incremental cost-effectiveness ratio above commonly cited cost-effectiveness thresholds when the price of the comparator market basket is at the lower range. After varying multiple inputs simultaneously while running multiple iterations of the model, we found cilta-cel ranged from \$118,000 per QALY to \$171,000 per QALY (Table E.4.5).

Based on limited clinical evidence, 60% of iterations for cilta-cel were below a threshold of \$150,000 per QALY gained and 98% were below a threshold of \$150,000 per evLYG gained (Tables E.4.6, E.4.7, and Figure E.4.4). However, this probabilistic analysis is limited in its incorporation of overall survival, therefore underestimates the range of uncertainty around lifetime outcomes and costs.

Finally, given the sensitivity of progression-free survival on model outcomes, we provide incremental cost-effectiveness ratios using alternative survival extrapolations (Table E.4.8). Varying the distribution used to extrapolate survival resulted in varying incremental cost-effectiveness ratios of -7% from the base-case of \$147,000 to above 3%. This sensitivity analysis should be interpreted with caution as some distributions may provide clinically implausible estimates of progression-free survival. Furthermore, we did not capture all uncertainty in extrapolations given a publicly available overall survival curve was not available. The potential credible range around QALYs gained for cilta-cel would likely be larger given inclusion of uncertainty around overall survival. This limitation is in part addressed through the one-way sensitivity analysis that increased median OS at upper levels of PFS (Figure E.4.3). Please note for all cilta-cel sensitivity analyses that a placeholder price is used based on the list price of ide-cel.

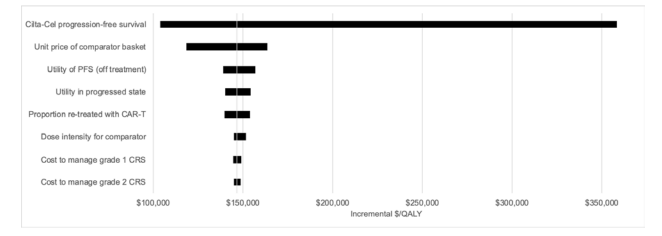


Figure E.4.3. Tornado Diagram for Cilta-Cel

	Lower Cost- Effectiveness Estimate	Upper Cost- Effectiveness Estimate	Lower Input*	Upper Input*
Cilta-Cel	\$103,721	\$358,262	13	33
progression-free				
survival				
Unit price of	\$118,543	\$163,616	\$17,083	\$37,508
comparator basket				
Utility of PFS (off	\$138,819	\$156,895	0.73	0.89
treatment)				
Utility in	\$139,988	\$154,252	0.64	0.78
progressed state				
Proportion re-	\$139,714	\$153,847	0.16	0.24
treated with CAR-T				
Dose intensity for comparator	\$144,834	\$151,570	0.83	1.00
Cost to manage grade 1 CRS	\$144,698	\$148,851	\$10,573	\$28,601
Cost to manage grade 2 CRS	\$144,742	\$148,796	\$13,309	\$36,005
Utility of PFS (on treatment)	\$144,689	\$148,182	0.70	0.85
Cost to manage grade 4 CRS	\$146,034	\$147,148	\$69,468	\$187,925

Table E.4.4. Tornado Diagram Inputs and Preliminary Results for Cilta-Cel versus Triple- or Quad-Refractory MM Comparator Market Basket (3+ prior lines of treatment)

*Note lower input may reflect either upper or lower ICER value depending on the direction that the input has on the ICER output.

	Cil	Cilta-Cel		Comparator Basket		mental
	Mean	Credible Range	Mean	Credible Range	Mean	Credible Range
	•		Total		•	
Total Costs	\$617,000	\$600,000 - \$636,000	\$277,000	\$226,000 - \$340,000	\$340,000	\$277,000 - \$386,000
Total QALYs	3.41	3.18 - 3.64	1.08	1.00-1.15	2.33	2.13-2.51
ICER					\$146,000	\$118,000 - \$171,000

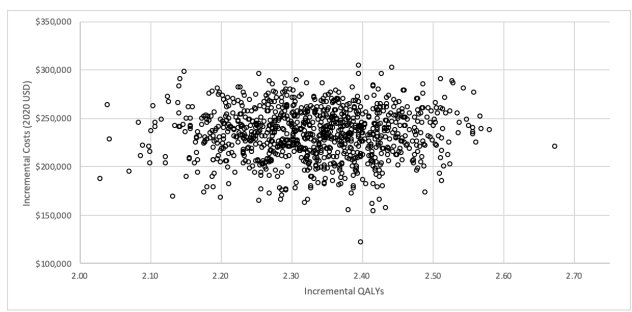
Table E.4.5. Preliminary Results of Probabilistic Sensitivity Analysis for Cilta-Cel versus Triple- orQuad-Refractory MM Comparator Market Basket (3+ prior lines of treatment)

Table E.4.6. Preliminary Probabilistic Sensitivity Analysis Cost per QALY Gained Results: Cilta-Celversus Triple-Class Refractory MM Comparator Market Basket (3+ prior lines of treatment)

	Cost Effective at \$50,000 per QALY	Cost Effective at \$100,000 per QALY	Cost Effective at \$150,000 per QALY	Cost Effective at \$200,000 per QALY
Cilta-cel	0%	0%	64%	100%

Table E.4.7. Preliminary Probabilistic Sensitivity Analysis Cost per evLYG Gained Results: Cilta-Celversus Triple-Class Refractory MM Comparator Market Basket (3+ prior lines of treatment)

	Cost Effective at	Cost Effective at	Cost Effective at	Cost Effective at
	\$50,000 per	\$100,000 per	\$150,000 per	\$200,000 per
	evLYG	evLYG	evLYG	evLYG
Cilta-cel	0%	1%	98%	100%



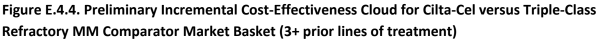


Table E.4.8. Preliminary Incremental Cost-Effectiveness Ratios under Alternative Survival
Extrapolations for Cilta-Cel

Curve	Distribution	Incremental Cost- Effectiveness Ratio versus Comparator (per QALY)	Relative percentage difference from base-case
PFS*			
	Weibull (base-case)	\$147,000	Reference
	Exponential	\$136,300	-7.2%
	Log-logistic	\$141,000	-3.8%
	Log-normal	\$138,000	-5.8%
	Gompertz	\$151,000	+3.1%

*Variation in overall survival not included; OS was modeled based on PFS relationship given there was no available overall survival curve

In the case of belantamab (Figure E.4.5 and Table E.4.11), the model results were most sensitive to progression-free survival and the price of the comparator market basket, specifically the market basket of triple- or quad-refractory comparators. Higher PFS (and thus higher survival overall) led to decreases in the incremental cost-effectiveness ratio whereas lower PFS resulted in incremental cost-effectiveness ratios that exceeded commonly cited cost-effectiveness thresholds. Higher prices for the comparator market basket drove the incremental cost-effectiveness ratio toward a scenario where belantamab was less costly and more effective whereas lower prices drove the incremental cost-effectiveness thresholds. Note that specific parameters for the triple-, quad- and penta-refractory market basket subsets are presented separately because the proportion of belantamab patients who were penta-refractory was submitted as academic-in-confidence).

After varying multiple inputs simultaneously while running multiple iterations of the model, we found belantamab ranged from less costly and more effective to incremental cost-effectiveness ratios that exceeded commonly cited cost-effectiveness thresholds (Table E.4.10, Table E.4.11, Table E.4.12, and Figure E.4.6). Overall model sensitivity is also illustrated here; for example, despite a base case estimate of \$98,000 per QALY, only 50% of iterations in the probabilistic analysis yielded a similar result (i.e., \$100,000 per QALY).

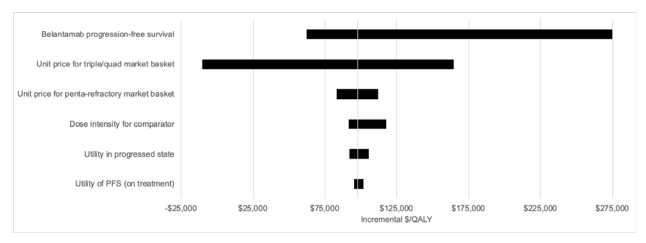


Figure E.4.5. Tornado Diagram for Belantamab

Table E.4.9. Tornado Diagram Inputs and Results for Belantamab versus Triple-, Quad-, or Penta-Refractory MM Comparator Market Basket (4+ prior lines of treatment)

	Lower Cost- Effectiveness Estimate	Upper Cost- Effectiveness Estimate	Lower Input*	Upper Input*
Belantamab	\$62,300	\$276,000	2	4
Progression-Free				
Survival (months)				
Unit Price of	Dominant (less costly,	\$164,000	\$17,083	\$37,508
triple/quad	more effective)			
comparator basket				
Unit Price of penta-	\$83,400	\$112,100	\$16,626	\$24,629
refractory				
comparator basket				
Dose intensity for	\$92,800	\$117,000	0.83	1.00
triple/quad				
comparator				
Utility in progressed	\$92,200	\$105,400	0.64	0.78
state				
Utility of PFS (on	\$95,000	\$102,000	0.70	0.85
treatment)				

^{*}Note lower input may reflect either upper or lower ICER value depending on the direction that the input has on the ICER output.

Table E.4.10. Results of Probabilistic Sensitivity Analysis for Belantamab versus Triple-, Quad-, orPenta- Refractory MM Comparator Market Basket (4+ prior lines of treatment)

	Belantamab		Compara	Comparator Basket		mental
	Mean	Credible Range	Mean	Credible Range	Mean	Credible Range
Total						
Total Costs	\$254,000	\$236,000 - \$270,000	\$217,000	\$151,000 - \$291,000	\$36,000	-\$5,000 - \$68,000
Total QALYs	1.16	0.91 - 1.41	0.78	0.72-0.84	0.38	0.14-0.60
ICER					\$97,000	Less costly, more effective - \$275,000

Table E.4.11. Probabilistic Sensitivity Analysis Cost Per QALY Gained Results: Belantamab versus Triple-, Quad-, or Penta- Refractory MM Comparator Market Basket (4+ prior lines of treatment)

	Cost Effective at \$50,000 per QALY	Cost Effective at \$100,000 per QALY	Cost Effective at \$150,000 per QALY	Cost Effective at \$200,000 per QALY
Belantamab	17%	50%	78%	91%

QALY: quality-adjusted life years gained

Table E.4.12. Probabilistic Sensitivity Analysis Cost Per evLYG Gained Results: Belantamab versus Triple-, Quad-, or Penta- Refractory MM Comparator Market Basket (4+ prior lines of treatment)

	Cost Effective at	Cost Effective at	Cost Effective at	Cost Effective at
	\$50,000 per	\$100,000 per	\$150,000 per	\$200,000 per
	evLYG	evLYG	evLYG	evLYG
Belantamab	18%	55%	81%	92%

evLYG: equal-value life years gained

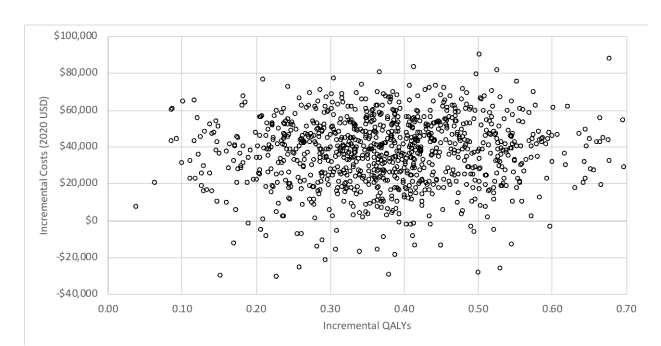


Figure E.4.6. Incremental Cost-Effectiveness Cloud for Belantamab versus Triple-, Quad-, or Penta-Refractory MM Comparator Market Basket (4+ prior lines of treatment)

E5. Scenario Analyses

We conducted scenario analyses that include:

- Assuming no additional charge for CAR-T retreatment, while continuing to assume administration, monitoring, and side-effect management costs for patients receiving a second infusion
- 2. Modified societal perspective that includes components such as productivity losses
- 3. Adjusting the proportional relationship between PFS and OS for belantamab to be within a similar range as a recent meta-analyses.³⁶

Scenario Analysis 1

Scenario analysis 1 is presented in the main report.

Scenario Analysis 2

We ran a modified societal perspective that included productivity losses and transportation time to and from health care appointments (see Table E.2.15 for unit costs).

Treatment	Comparator	Cost per QALY Gained	Cost per Life Year Gained	Cost per evLYG	Cost per Month of PFS Gained
lde-cel	CAR-T	\$319,000	\$250,000	\$280,000	\$35,000
	Comparator				
	Market Basket				
Cilta-cel	CAR-T	\$146,000	\$113,000	\$128,000	\$17,000
	Comparator				
	Market Basket				
Belantamab	Belantamab	\$99,000	\$70,000	\$93,000	\$18,000
	Comparator				
	Market Basket				

Table E.5.1. Incremental Cost-Effectiveness Ratios for Scenario Analysis 1

evLYG: equal-value of life years gained, QALY: quality-adjusted life year, LY: life years

Scenario Analysis 3

Recent evidence synthesis in multiple myeloma suggests a proportional relationship between PFS and OS consistent with a 2.5-3.0 month gain in OS for every 1 month gain in PFS.^{36,37} While both CAR-T therapies were within the range of these proportional relationships, evidence from the DREAMM-2 trial suggests a nearly five month gain for every one month gain in PFS. We ran a scenario analysis that adjusted the relationship between PFS and OS for belantamab to be consistent with prior published data. In this scenario, belantamab adjusted OS estimates were set at an approximate median of 9 months (vs. 13.7 months from the trial). This scenario suggests belantamab is more costly and less effective than the market basket comparator. (Table E.5.2).

Treatment	Comparator	Cost per QALY Gained	Cost per Life Year Gained	Cost per evLYG	Cost per Month of PFS Gained
Belantamab	Belantamab Comparator	Dominated	Dominated	Dominated	\$17,000
	Market Basket	(more costly,	(more costly,	(more costly,	
		less effective)	less effective)	less effective)	

evLYG: equal-value of life years gained, QALY: quality-adjusted life year, LY: life years, PFS: progression-free survival

E6. Heterogeneity and Subgroups

We considered estimating costs and health outcomes among relevant subgroups, such as patients with genetic factors that put them at particularly high risk as well as subgroups defined by race. Due to small patient numbers and the lack of data on survival, health-related quality of life, and health care costs stratified by these subgroups, we were not able to conduct subgroup analyses.

E7. Model Validation

Model validation followed standard practices in the field. We tested all mathematical functions in the model to ensure they were consistent with the report (and supplemental materials). We also conducted sensitivity analyses with null input values to ensure the model was producing findings consistent with expectations. Further, independent modelers tested the mathematical functions in the model as well as the specific inputs and corresponding outputs.

Model validation was also conducted in terms of comparisons to other model findings. We searched the literature to identify models that were similar to our analysis, with comparable populations, settings, perspective, and treatments.

We used several approaches to validate the model. First, we provided preliminary model structure, methods and assumptions to manufacturers, patient groups, and clinical experts. Based on feedback from these groups, we refined data inputs used in the model, as needed. We also performed calibration techniques that fit survival estimates to observed findings from trials and other evidence in MM. Second, we varied model input parameters to evaluate face validity of changes in results. We performed model verification for model calculations using internal reviewers. As part of ICER's efforts in acknowledging modeling transparency, we also are sharing the model with the relevant manufacturers for external verification around the time of publishing the draft report for this review. During the model transparency process manufacturers noted we underestimated survival for ide-cel and the CAR-T comparator market basket (which also impacted a percentage of the comparator to belantamab). These changes were reflected in the final model calculations. Finally, we compared results to other cost-effectiveness models in this therapy area. The outputs from the model were validated against the trial/study data of the interventions and also any relevant observational datasets.

E8. Prior Economic Models

We found prior economic models in multiple myeloma through two recent systematic literature reviews.^{38,91} While the most recent review identified 17 publications,³⁸ the majority were assessing therapies in the first or second line setting. One publication that assessed the third line of therapy was based on the prior ICER review in multiple myeloma.³⁹ Carlson et al. estimated the cost-effectiveness of second- and third-line therapies in RRMM. The Carlson et al. analysis had the advantage of a network meta-analysis to combine evidence on indirect comparisons across regimens of interest which was not available in this analysis. Therefore, this previous study allowed for direct application of hazard ratios to PFS and OS curves. Our analysis instead separately estimated and extrapolated survival outcomes by treatment arm in the model given the lack of an available indirect treatment comparison. For sensitivity analyses, instead of applying a hazard ratio we varied the shape and scale parameters on the active interventions, while fixing the comparator

arm. While the approach to estimate our base-case and sensitivity analyses was different than Carlson et al., both studies found considerable uncertainty on whether therapies were cost-effective at various commonly cited thresholds. It should be noted the Carlson et al. analysis was based on second- and third-lines of therapy versus this analysis that is focused on TCRMM patients that failed at least three lines of therapy. Therefore, direct comparisons on QALYs, LYs, and other outcomes should not be made between studies.

Studies by Ailawadhi et al. and Roy et al. informed our market basket cost calculations.^{87,89} Specifically, the approach used to identify regimen dosing each cycle along with identifying and costing adverse events informed our market basket calculation. In some cases, the combination of therapies was different so caution should be used when comparing the total cost estimates between studies. The most recent evidence from Ailawadhi et al. found total costs ranging from approximately \$93,000 to \$315,000 for common second-line regimens. While these estimates are for earlier line therapies, some of the reported regimens are approved for later lines of therapy and were used in our market basket calculations. For example, our total cost estimation for the CAR-T market basket of comparators was approximately \$150,000 (discounted) over a lifetime horizon which is within the range reported by Ailawadhi et al. We further varied the market basket cost and found the cost of comparators to be a key driver of model results.

F. Potential Budget Impact: Supplemental Information

ICER's methods for estimating potential budget impact are described in detail elsewhere and have recently been updated.^{92,93} The intent of our revised approach to potential budgetary impact is to document the percentage of patients that could be treated at selected prices without crossing a budget impact threshold that is aligned with overall growth in the US economy.

Briefly, we evaluate whether a new drug would take market share from one or more existing treatments and calculate the blended potential budget impact associated with displacing use of existing therapies with the new intervention. In this analysis, we assumed that patients eligible for the interventions under review in this analysis would otherwise have been treated with the comparator treatment(s).

Using this approach to estimate potential budget impact, we then compared our estimates to an updated budget impact threshold that represents a potential trigger for policy mechanisms to improve affordability, such as changes to pricing, payment, or patient eligibility. As described in ICER's methods presentation (<u>https://icer-review.org/methodology/icers-methods/icer-value-assessment-framework-2/</u>), this threshold is based on an underlying assumption that health care costs should not grow much faster than growth in the overall national economy. From this foundational assumption, our potential budget impact threshold is derived using an estimate of growth in US gross domestic product (GDP) +1%, the average number of new drug approvals by the FDA over the most recent two-year period, and the contribution of spending on retail and facility-based drugs to total health care spending. For reports begun in 2019-2020, the five-year annualized potential budget impact threshold that should trigger policy actions to manage access and affordability is calculated to total approximately \$819 million per year for new drugs.

All costs used in the potential budget impact model were undiscounted and estimated over a fiveyear time horizon. This potential budget impact analysis included the estimated number of individuals in the US who would be eligible for treatment described earlier in the report. We assumed 20% of these patients would initiate treatment in each of the five years

In the final evidence report, ICER will include an "affordability and access alert" if discussion among clinical experts at the public meeting of ICER's independent appraisal committees suggests that full, "clinically optimal" utilization at estimated net pricing (or at the \$150,000 per QALY threshold price if estimated net price is not available) would exceed the ICER annual potential budget impact threshold, without active intervention by insurers and others to manage access to the treatment.

Results

Table F.1. Cumulative Net Cost per Patient Treated with Belantamab at Wholesale AcquisitionCost Over a Five-Year Time Horizon

	Belantamab	
Year	Additional Costs per Year (non-cumulative)	Cumulative Cost
Year 1	\$5,613	\$5,613
Year 2	\$23,914	\$29,527
Year 3	\$7,384	\$36,911
Year 4	\$1,908	\$38,819
Year 5	\$390	\$39,209

 Table F.2. Cumulative Net Cost per Patient Treated with Ide-cel at List Price

	lde-cel	
Year	Additional Costs per Year (non-cumulative)	Cumulative Cost
Year 1	\$354,339	\$354,339
Year 2	-\$152	\$354,187
Year 3	\$4,361	\$358,548
Year 4	\$3,297	\$361,845
Year 5	\$2,270	\$364,115

 Table F.3. Cumulative Net Cost per Patient Treated with Cilta-cel at Placeholder Price

	Cilta-cel	
Year	Additional Costs per Year (non-cumulative)	Cumulative Cost
Year 1	\$289,205	\$289,205
Year 2	\$2,604	\$291,809
Year 3	\$17,039	\$308,848
Year 4	\$13,242	\$322,090
Year 5	\$8,245	\$330,335

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