

אוגוסט 2024

Abecma / אבקמה
(Idecabtagene Vicleucel)
תרחיף לעירוי תוך ורידי / Dispersion for IV infusion

רופא/ה, רוקח/ת יקר/ה,

חברת בריסטול-מאיירס סקוויב (ישראל) מבקשת להודיע על עדכונים בעלון לרופא ובעלון לצרכן של התכשיר שבנדון.

להלן התוויית התכשיר כפי שמאושרת ע"י משרד-הבריאות:

Abecma is indicated for the treatment of adult patients with relapsed and refractory multiple myeloma who have received at least three prior therapies, including an immunomodulatory agent, a proteasome inhibitor and an anti-CD38 antibody and have demonstrated disease progression on the last therapy.

המרכיב הפעיל: Idecabtagene Vicleucel 260-500 x 10⁶ anti-BCMA CAR-positive viable T cells

השינויים בעלון לרופא ובעלון לצרכן משוקפים בעמודים הבאים.

תוספת טקסט מסומנת בקו תחתון, מחיקת טקסט בקו חוצה, החמרות מסומנות בצהוב.

למידע מלא על התרופה יש לעיין בעלון לרופא ובעלון לצרכן כפי שאושרו על ידי משרד-הבריאות.

העלון לרופא והעלון לצרכן נשלחו לפרסום במאגר התרופות שבאתר משרד-הבריאות וניתן לקבלם מודפסים על ידי פנייה לבעל הרישום בריסטול-מאיירס סקוויב (ישראל) בע"מ.

בברכה,

לנה גיטלין

מנהלת רגולציה ורוקחת ממונה
בריסטול-מאיירס סקוויב (ישראל)

1. NAME OF THE MEDICINAL PRODUCT

Abecma 260 - 500 x 10⁶ cells dispersion for infusion

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

2.1 General description

Abecma (idecabtagene vicleucel) is a genetically modified autologous cell-based product containing immunotherapy consisting of human T cells transduced ex-vivo using a replication incompetent with lentiviral vector (LVV) encoding a chimeric antigen receptor (CAR) that recognises B-cell maturation antigen (BCMA), comprising a murine-derived, anti-human BCMA single chain variable fragment (scFv) linked to a 4-1BB costimulatory domain and a CD3-zeta signalling domain.

2.2 Qualitative and quantitative composition

Each patient-specific infusion bag of Abecma contains idecabtagene vicleucel cell dispersion at a batch-dependent concentration of autologous T cells genetically modified to express an anti-BCMA chimeric antigen receptor (CAR-positive viable T cells). The finished medicinal product is packaged in one or more infusion bags overall containing a cell dispersion of 260 to 500 x 10⁶ CAR-positive viable T cells suspended in a cryopreservative solution.

Each infusion bag contains 10-30 mL, 30-70 mL or 55-100 mL of dispersion for infusion.

The cellular composition and the final cell number varies between individual patient batches. In addition to T cells, natural killer (NK) cells may be present. The quantitative information of medicinal product, including regarding the number of CAR-positive viable T cells/infusion bag(s) to be administered, are is presented in the release for infusion certificate (RfIC) documentation located inside the lid of the dry vapour cryoshipper used for transport.

Excipients with known effect

This medicinal product contains 5% dimethyl sulfoxide (DMSO), up to 752 mg sodium and up to 274 mg potassium per dose.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Dispersion for infusion.

A colourless dispersion.

4. CLINICAL PARTICULARS

The marketing of Abecma is subject to a Risk Management Plan (RMP) including a Patient Card ('Patient Safety Information Card'). The 'Patient Safety Information Card' emphasizes important safety information that the patient should be aware of before and during treatment.

Please explain to the patient the need to review the card before starting the treatment.

This product is marketed with a Healthcare Professional Guide (~~Prescriber Guide~~), providing important safety information. Please ensure you are familiar with this material as it contains important safety information.

4.1 Therapeutic indications

Abecma is indicated for the treatment of adult patients with relapsed and refractory multiple myeloma who have received at least three prior therapies, including an immunomodulatory agent, a proteasome inhibitor and an anti-CD38 antibody and have demonstrated disease progression on the last therapy.

4.2 Posology and method of administration

Abecma must be administered in a qualified treatment centre.

Abecma therapy should be initiated under the direction of and supervised by a healthcare professional experienced in the treatment of haematological malignancies and trained for the administration and management of patients treated with Abecma.

A minimum of one dose of tocilizumab for use in the event of cytokine release syndrome (CRS) and emergency equipment must be available prior to infusion of Abecma. The treatment centre must have access to an additional dose of tocilizumab within 8 hours of each previous dose. In the exceptional case where tocilizumab is not available, suitable alternative measures to treat CRS instead of tocilizumab must be available prior to infusion.

Posology

Abecma is intended for autologous use only (see section 4.4). ~~Manufacture and release of Abecma usually takes about 4-5 weeks.~~

Treatment consists of a single dose for infusion containing a dispersion of CAR-positive viable T cells in one or more infusion bags. The target dose is 420×10^6 CAR-positive viable T cells within a range of 260 to 500×10^6 CAR-positive viable T cells. See the accompanying release for infusion certificate (RfIC) for additional information pertaining to dose.

Pre-treatment (lymphodepleting chemotherapy)

Lymphodepleting chemotherapy consisting of cyclophosphamide $300 \text{ mg/m}^2/\text{day}$ intravenously (IV) and fludarabine $30 \text{ mg/m}^2/\text{day}$ IV should be administered for 3 days. See the prescribing information for cyclophosphamide and fludarabine for information on dose adjustment in renal impairment.

Abecma is to be administered 2 days after completion of lymphodepleting chemotherapy, up to a maximum of 9 days. The availability of Abecma must be confirmed prior to starting the lymphodepleting chemotherapy. If there is a delay in Abecma infusion of more than 9 days, 4-weeks between completing lymphodepleting chemotherapy and the infusion, then the patient should be re-treated with lymphodepleting chemotherapy after a minimum of 4 weeks from last lymphodepleting chemotherapy prior to receiving Abecma.

Pre-medication

It is recommended that premedication ~~To minimise the risk of infusion reactions, the patient should be pre-medicated~~ with paracetamol (500 to 1,000 mg orally) and diphenhydramine (12.5 mg IV or 25 to 50 mg orally) or another H₁-antihistamine, be administered approximately 30 to 60 minutes before the infusion of Abecma to reduce the possibility of an infusion reaction.

Prophylactic use of systemic corticosteroids should be avoided as the use may interfere with the activity of Abecma. Therapeutic doses of corticosteroids should be avoided 72 hours prior to the start of lymphodepleting chemotherapy and following Abecma infusion except for the management of CRS, neurologic toxicities and other life-threatening emergencies (see section 4.4).

Clinical assessment prior to infusion

Abecma treatment should be delayed in some patient groups at risk (see section 4.4).

Monitoring after infusion

- Patients should be monitored for the first 10 days following infusion at the qualified treatment centre for signs and symptoms of CRS, neurologic events and other toxicities.
- After the first 10 days following infusion, the patient should be monitored at the physician's discretion.
- Patients should be instructed to remain within proximity (within 2 hours of travel) of the qualified treatment centre for at least 4 weeks following infusion.

Special populations

Patients with human immunodeficiency virus (HIV), hepatitis B virus (HBV) and hepatitis C virus (HCV) infection

There is no clinical experience in patients with active HIV, HBV or HCV infection. Screening for HBV, active HIV and active HCV must be performed before collection of cells for manufacturing. Leukapheresis material from patients with active HIV or active HCV infection will not be accepted for Abecma manufacturing (see section 4.4).

Elderly

No dose adjustment is required in patients over 65 years of age (see section 5.1).

Paediatric population

The safety and efficacy of Abecma in children and adolescents below 18 years of age have not been established. No data are available.

Method of administration

Abecma is for intravenous use only.

Administration

- Do NOT use a leukodepleting filter.
- Ensure that tocilizumab or suitable alternatives, in the exceptional case where tocilizumab is not available, and emergency equipment are available prior to infusion and during the recovery period.
- Central venous access may be utilised for the infusion of Abecma and is encouraged in patients with poor peripheral access.
- Before administration, it must be confirmed that ~~Confirm~~ the patient's identity matches the unique patient information identifiers on the Abecma infusion bag and accompanying documentation. The total number of infusion bags to be administered must also be confirmed with the patient specific information on the release for infusion certificate (RfIC) (see section 4.4).

For detailed instructions on preparation, administration, measures to take in case of accidental exposure and disposal of Abecma the medicinal product see section 6.6.

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.

Contraindications of the lymphodepleting chemotherapy must be considered.

4.4 Special warnings and precautions for use

Traceability

The traceability requirements of cell-based advanced therapy medicinal products must apply. To ensure traceability the name of the product, the batch number and the name of the treated patient ~~should~~ **must** be kept for a period of 30 years after expiry date of the product.

Autologous use

~~Abecma is intended solely for autologous use and should **must not**, under ~~no any~~ circumstances, be administered to other patients. Before infusion, the patient's identity must match the patient identifiers on the Abecma infusion bag, cassette and the release for infusion certificate (RFIC). Abecma must not be administered if the information on the **product labels and the release for infusion certificate (RFIC) patient specific label** does not match the ~~intended patient's identity~~.~~

Rapidly progressing disease

Before selecting patients for Abecma treatment, physicians should consider the impact of high-risk cytogenetic abnormalities, Revised International Staging System (R-ISS) stage III, presence of extramedullary plasmacytoma or high tumour burden, particularly for patients who have rapidly progressing disease that may affect their ability to receive CAR T infusion in due time. For these patients, optimising bridging therapy may be particularly important. Some patients may not benefit from Abecma treatment due to potential increased risk of early death (see section 5.1).

Reasons to delay treatment

Due to the risks associated with Abecma treatment, infusion should be delayed up to 7 days if a patient has any of the following conditions:

- Unresolved serious adverse events (especially pulmonary events, cardiac events or hypotension) including those after preceding chemotherapies.
- Active infections or inflammatory disorders (including pneumonitis, myocarditis or hepatitis).
- Active graft-versus-host disease (GVHD).

Autologous use

~~Abecma is intended solely for autologous use and should under no circumstances be administered to other patients. Before infusion, the patient's identity must match the patient identifiers on the Abecma infusion bag, cassette and the release for infusion certificate (RFIC). Abecma must not be administered if the information on the patient specific label does not match the intended patient.~~

Concomitant disease

Patients with active central nervous system (CNS) disorder or inadequate renal, hepatic, pulmonary or cardiac function are likely to be more vulnerable to the consequences of the adverse reactions

described below and require special attention.

Central nervous system pathology

There is no experience of use of Abecma in patients with CNS involvement of myeloma or other pre-existing, clinically relevant CNS pathologies.

Prior allogeneic stem cell transplantation

It is not recommended that patients receive Abecma within 4 months after an allogeneic stem cell transplant (SCT) because of the potential risk of Abecma worsening GVHD. Leukapheresis for Abecma manufacturing should be performed at least 12 weeks after allogeneic SCT.

Prior treatment with an anti-BCMA therapy

There is limited experience with Abecma in patients exposed to prior BCMA-directed therapy.

There is limited experience of retreating patients with a second dose of Abecma. Responses after Abecma retreatment were infrequent and less durable when compared to initial treatment. Additionally, fatal outcomes were observed in retreated patients.

Cytokine release syndrome

CRS, including fatal or life-threatening reactions occurred following Abecma infusion. Nearly all patients experienced some degree of CRS. In clinical studies, the ~~The~~ median time to onset of CRS was 1 day (range: 1 to ~~12~~17) (see section 4.8).

Monitoring and management of CRS

CRS should be identified based on clinical presentation. Patients should be evaluated and treated for other causes of fever, hypoxia and hypotension. CRS has been reported to be associated with findings of haemophagocytic lymphohistiocytosis/macrophage activation syndrome (HLH/MAS) and the physiology of the syndromes may overlap. MAS is a potentially life-threatening condition, and patients should be closely monitored for evidence of MAS. Treatment of MAS should be administered per institutional guidelines.

One dose of tocilizumab per patient must be on-site and available for administration prior to Abecma infusion. The treatment centre must have access to an additional dose of tocilizumab within 8 hours of each previous dose. In the exceptional case where tocilizumab is not available, the treatment centre must have access to suitable alternative measures instead of tocilizumab to treat CRS. Patients should be monitored for the first 10 days following Abecma infusion at the qualified treatment centre for signs and symptoms of CRS. After the first 10 days following infusion, the patient should be monitored at the physician's discretion. Patients should be counselled to remain within proximity (within 2 hours of travel) of the qualified treatment centre for at least 4 weeks following infusion and to seek immediate medical attention should signs or symptoms of CRS occur at any time.

At the first sign of CRS, treatment with supportive care, tocilizumab or tocilizumab and corticosteroids - should be instituted, as indicated in Table 1. Abecma can continue to expand and persist following administration of tocilizumab and corticosteroids (see section 4.5).

Patients who experience CRS should be closely monitored for cardiac and organ functioning until resolution of symptoms. For severe or life-threatening CRS, intensive care unit level monitoring and supportive therapy should be considered.

If concurrent neurologic toxicity is suspected during CRS, ~~manage~~ the neurologic toxicity should be

managed according to the recommendations in Table 2 and use the more aggressive intervention of the two reactions specified in Tables 1 and 2.

Earlier escalation (i.e. higher corticosteroid dose, alternative anticytokine agents, anti-T cell therapies) is recommended in patients with refractory CRS within 72 hours post Abecma infusion characterised by persistent fever, end-organ toxicity (e.g. hypoxia, hypotension) and/or HLH/MAS not improving in grade within 12 hours of first line interventions.

Table 1. CRS grading and management guidance

| CRS grade^a | Tocilizumab | Corticosteroids |
|---|--|--|
| Grade 1 Symptoms require symptomatic treatment only (e.g. fever, nausea, fatigue, headache, myalgia, malaise). | If onset 72 hours or more after infusion, treat symptomatically. If onset less than 72 hours after infusion and symptoms not controlled by supportive care alone, consider tocilizumab 8 mg/kg IV over 1 hour (not to exceed 800 mg). | — |
| Grade 2 Symptoms require and respond to moderate intervention. Oxygen requirement less than 40% FiO ₂ or hypotension responsive to fluids or low dose of one vasopressor or Grade 2 organ toxicity. | Administer tocilizumab 8 mg/kg IV over 1 hour (not to exceed 800 mg). | Consider dexamethasone 10 mg IV every 12 to 24 hours. |
| Grade 3 Symptoms require and respond to aggressive intervention. Fever, oxygen requirement greater than or equal to 40% FiO ₂ or hypotension requiring high-dose or multiple vasopressors or Grade 3 organ toxicity or Grade 4 transaminitis. | Administer tocilizumab 8 mg/kg IV over 1 hour (not to exceed 800 mg). | Administer dexamethasone (e.g. 10 mg IV every 12 hours). |

| | | |
|--|---|--|
| <p>For Grade 2 and 3: <u>If no improvement within 24 hours or rapid progression, repeat tocilizumab and escalate dose and frequency of dexamethasone (20 mg IV every 6 to 12 hours).</u> <u>If no improvement within 24 hours or continued rapid progression, switch to methylprednisolone 2 mg/kg followed by 2 mg/kg divided 4 times per day.</u> <u>If steroids are initiated, continue steroids for at least 3 doses, and taper over a maximum of 7 days.</u> <u>After 2 doses of tocilizumab, consider alternative anticytokine agents. Do not exceed 3 doses tocilizumab in 24 hours or 4 doses in total.</u></p> | | |
| <p>Grade 4 <u>Life-threatening symptoms. Requirements for ventilator support, continuous veno-venous hemodialysis (CVVHD) or Grade 4 organ toxicity (excluding transaminitis).</u></p> | <p><u>Administer tocilizumab 8 mg/kg IV over 1 hour (not to exceed 800 mg).</u></p> | <p><u>Administer dexamethasone 20 mg IV every 6 hours.</u></p> |
| <p>For Grade 4: <u>After 2 doses of tocilizumab, consider alternative anticytokine agents. Do not exceed 3 doses of tocilizumab in 24 hours or 4 doses in total.</u> <u>If no improvement within 24 hours, consider methylprednisolone (1 to 2 g, repeat every 24 hours if needed; taper as clinically indicated) or anti-T cell therapies such as cyclophosphamide 1.5 g/m² or others.</u></p> | | |

^a Lee et al, 2014.

| CRS grade* | Tocilizumab | Corticosteroids |
|--|---|--|
| <p>For Grade 2 and 3: <u>If no improvement within 24 hours or rapid progression, repeat tocilizumab and escalate dose and frequency of dexamethasone (20 mg IV every 6 to 12 hours).</u> <u>If no improvement within 24 hours or continued rapid progression, switch to methylprednisolone 2 mg/kg followed by 2 mg/kg divided 4 times per day.</u> <u>If steroids are initiated, continue steroids for at least 3 doses, and taper over a maximum of 7 days.</u> <u>After 2 doses of tocilizumab, consider alternative anticytokine agents.</u> <u>Do not exceed 3 doses tocilizumab in 24 hours or 4 doses in total.</u></p> | | |
| <p>Grade 4 <u>Life-threatening symptoms. Requirements for ventilator support, continuous veno-venous hemodialysis (CVVHD) or Grade 4 organ toxicity (excluding transaminitis).</u></p> | <p><u>Administer tocilizumab 8 mg/kg IV over 1 hour (not to exceed 800 mg).</u></p> | <p><u>Administer dexamethasone 20 mg IV every 6 hours.</u></p> |

For Grade 4:

After 2 doses of tocilizumab, consider alternative anti-cytokine agents. Do not exceed 3 doses of tocilizumab in 24 hours or 4 doses in total.

If no improvement within 24 hours, consider methylprednisolone (1 to 2 g, repeat every 24 hours if needed; taper as clinically indicated) or anti-T cell therapies such as cyclophosphamide 1.5 g/m² or others.

*Lee et al, 2014.

Neurologic adverse reactions

Neurologic toxicities, such as aphasia, and encephalopathy, and immune effector cell-associated neurotoxicity syndrome (ICANS), which may be severe or life-threatening, occurred following treatment with Abecma. The median time to onset of the first event of neurotoxicity was 2-3 days (range: 1 to 40-317 days; one patient developed encephalopathy at Day 317 as a result of worsening pneumonia and *Clostridium difficile* colitis). Grade 3 parkinsonism has also been reported, with delayed onset. Neurologic toxicity may occur concurrently with CRS, after CRS resolution or in the absence of CRS (see section 4.8).

Monitoring and management of neurologic toxicities

Patients should be monitored for the first 10 days following Abecma infusion at the qualified treatment centre for signs and symptoms of neurologic toxicities. After the first 10 days following infusion, the patient should be monitored at the physician's discretion. Patients should be counselled to remain within proximity (within 2 hours of travel) of the qualified treatment centre for at least 4 weeks following infusion and to seek immediate medical attention should signs and symptoms of neurologic toxicities occur at any time.

If neurologic toxicity is suspected, manage according to the recommendations in Table 2. Other causes of neurologic symptoms should be ruled out. Intensive care supportive therapy should be provided for severe or life-threatening neurologic toxicities.

If concurrent CRS is suspected during the neurologic toxicity reaction, it should be managed according to the recommendations in Table 1 and the more aggressive intervention used for the two reactions specified in Tables 1 and 2.

Table 2. Neurologic toxicity including ICANS grading and management guidance

| Neurotoxicity grade including presenting symptoms^a | Corticosteroids and antiseizure medications |
|---|--|
| <p>Grade 1 Mild or asymptomatic.</p> <p>ICE score 7-9^b</p> <p>or</p> <p>Depressed level of consciousness^c: awakens spontaneously.</p> | <p>Start non-sedating, antiseizure medicines (e.g. levetiracetam) for seizure prophylaxis.</p> <p>If 72 hours or more after infusion, observe patient.</p> <p>If less than 72 hours after infusion, and symptoms not controlled by supportive care alone, consider dexamethasone 10 mg IV every 12 to 24 hours for 2 to 3 days.</p> |
| <p>Grade 2 Moderate.</p> <p>ICE score 3-6^b</p> <p>or</p> <p>Depressed level of consciousness: awakens to voice.</p> | <p>Start non-sedating, antiseizure medicines (e.g. levetiracetam) for seizure prophylaxis.</p> <p>Start dexamethasone 10 mg IV every 12 hours for 2 to 3 days or longer for persistent symptoms. Consider taper for a total steroid exposure of greater than 3 days. Steroids are not recommended for isolated Grade 2 headaches.</p> <p>If no improvement after 24 hours or worsening of neurologic toxicity, increase the dose and/or frequency of dexamethasone up to a maximum of 20 mg IV every 6 hours.</p> |
| <p>Grade 3 Severe or medically significant but not immediately life-threatening; hospitalization or prolongation; disabling.</p> <p>ICE score 0-2^b <i>if ICE score is 0, but the patient is arousable (e.g., awake with global aphasia) and able to perform assessment.</i></p> <p>or</p> <p>Depressed level of consciousness^c: awakens only to tactile stimulus.</p> <p>Or seizures^c, either:</p> <ul style="list-style-type: none"> • any clinical seizure, focal or generalised, that resolves rapidly, or • non-convulsive seizures on EEG that resolve with intervention. <p>Or raised ICP^c: focal/local oedema on neuroimaging.</p> | <p>Start non-sedating, antiseizure medicines (e.g. levetiracetam) for seizure prophylaxis.</p> <p>Start dexamethasone 10 to 20 mg IV every 8 to 12 hours. Steroids are not recommended for isolated Grade 3 headaches.</p> <p>If no improvement after 24 hours or worsening of neurologic toxicity, escalate to methylprednisolone (2 mg/kg loading dose, followed by 2 mg/kg divided into 4 times a day; taper within 7 days).</p> <p>If cerebral oedema is suspected, consider hyperventilation and hyperosmolar therapy. Give high-dose methylprednisolone (1 to 2 g, repeat every 24 hours if needed; taper as clinically indicated) and cyclophosphamide 1.5 g/m².</p> |

| | |
|---|---|
| <p>Grade 4</p> <p>Life- threatening.</p> <p>ICE score^b 0</p> <p>or</p> <p>Depressed level of consciousness^c either:</p> <ul style="list-style-type: none"> • patient is unarousable or requires vigorous or repetitive tactile stimuli to arouse, or • stupor or coma. <p>Or seizures^c, either:</p> <ul style="list-style-type: none"> • life-threatening prolonged seizure (>5 min), or • repetitive clinical or electrical seizures without return to baseline in between. <p>Or motor findings^c:</p> <ul style="list-style-type: none"> • deep focal motor weakness such as hemiparesis or paraparesis. <p>Or, raised ICP/cerebral oedema^c, with signs/symptoms such as:</p> <ul style="list-style-type: none"> • diffuse cerebral oedema on neuroimaging, or • decerebrate or decorticate posturing, <p>or</p> <ul style="list-style-type: none"> • cranial nerve VI palsy, or • papilledema, or • Cushing's triad. | <p>Start non-sedating, antiseizure medicines (e.g. levetiracetam) for seizure prophylaxis.</p> <p>Start dexamethasone 20 mg IV every 6 hours.</p> <p>If no improvement after 24 hours or worsening of neurologic toxicity, escalate to high-dose methylprednisolone (1 to 2 g, repeated every 24 hours if needed; taper as clinically indicated). Consider cyclophosphamide 1.5 g/m².</p> <p>If cerebral oedema is suspected, consider hyperventilation and hyperosmolar therapy. Give high-dose methylprednisolone (1 to 2 g, repeat every 24 hours if needed; taper as clinically indicated) and cyclophosphamide 1.5 g/m².</p> |
|---|---|

*NCICTCAE v.4 criteria for grading neurologic toxicities.

EEG=Electroencephalogram; ICE=Immune Effector Cell-Associated Encephalopathy; ICP=intracranial pressure

^a Management is determined by the most severe event, not attributable to any other cause.

^b If patient is arousable and able to perform ICE Assessment, assess: Orientation (oriented to year, month, city, hospital = 4 points); Naming (name 3 objects, e.g., point to clock, pen, button = 3 points); Following Commands (e.g., "show me 2 fingers" or "close your eyes and stick out your tongue" = 1 point); Writing (ability to write a standard sentence = 1 point); and Attention (count backwards from 100 by ten = 1 point). If patient is unarousable and unable to perform ICE Assessment (Grade 4 ICANS) = 0 points.

^c Attributable to no other cause.

Prolonged cytopenias

Patients may exhibit prolonged cytopenias for several weeks following lymphodepleting chemotherapy and Abecma infusion (see section 4.8). Blood counts should be monitored prior to and after Abecma infusion. Cytopenias should be managed with myeloid growth factor and blood transfusion support according to institutional guidelines.

Infections and febrile neutropenia

Abecma should not be administered to patients with active infections or inflammatory disorders. Severe infections, including life-threatening or fatal infections, have occurred in patients after receiving Abecma (see section 4.8). Patients should be monitored for signs and symptoms of infection before and after Abecma infusion and treated appropriately. Prophylactic, pre-emptive and/or therapeutic antimicrobials should be administered according to institutional guidelines.

Febrile neutropenia was observed in patients after Abecma infusion (see section 4.8) and may be concurrent with CRS. In the event of febrile neutropenia, infection should be evaluated and managed with broad-spectrum antibiotics, fluids and other supportive care as medically indicated.

Viral reactivation

Cytomegalovirus (CMV) infection resulting in pneumonia and death have occurred following Abecma administration (see section 4.8). Patients should be monitored and treated for CMV infection according to clinical guidelines.

HBV reactivation, in some cases resulting in fulminant hepatitis, hepatic failure and death, can occur in patients treated with medicinal products directed against plasma cells (see section 4.8).

Screening for CMV, HBV, active HIV and active HCV must be performed before collection of cells for manufacturing (see section 4.2).

Hypogammaglobulinaemia

Plasma cell aplasia and hypogammaglobulinaemia can occur in patients receiving treatment with Abecma (see section 4.8). Immunoglobulin levels should be monitored after treatment with Abecma and managed per institutional guidelines including infection precautions, antibiotic or antiviral prophylaxis and immunoglobulin replacement.

Secondary malignancies

Patients treated with Abecma may develop secondary malignancies. Patients should be monitored life-long for secondary malignancies. In the event that a secondary malignancy of T cell origin occurs, the company should be contacted to obtain instructions on the collection of patient samples for testing.

Hypersensitivity reactions

Allergic reactions may occur with the infusion of Abecma. Serious hypersensitivity reactions, including anaphylaxis, may be due to dimethyl sulfoxide (DMSO), an excipient in Abecma. Patients not previously exposed to DMSO should be observed closely. Vital signs (blood pressure, heart rate, and oxygen saturation) and the occurrence of any symptom should be monitored prior to the start of the infusion, approximately every ten minutes during the infusion and every hour, for 3 hours, after the infusion.

Transmission of an infectious agent

Although Abecma is tested for sterility and mycoplasma, a risk of transmission of infectious agents exists. Healthcare professionals administering Abecma must, therefore, monitor patients for signs and symptoms of infections after treatment and treat appropriately, if needed.

Interference with ~~serological-virological~~ testing

~~HIV and the lentivirus used to make Abecma have~~ Due to limited ~~and~~, short spans of identical genetic material (RNA), ~~information between the lentiviral vector used to create Abecma and HIV. Therefore,~~ some commercial HIV nucleic acid tests (NAT) may ~~yield give a false~~ positive results ~~in patients who have received Abecma.~~

Blood, organ, tissue and cell donation

Patients treated with Abecma ~~should~~ **must** not donate blood, organs, tissues and cells for transplantation.

Long-term follow-up

Patients are expected to be enrolled in a ~~registry and will be followed in the~~ registry in order to better understand the long-term safety and efficacy of Abecma.

Excipients

This medicinal product contains up to 33 mmol (752 mg) sodium per dose, equivalent to 37.6% of the WHO recommended maximum daily intake of 2 g sodium for an adult.

This medicinal product contains up to 7 mmol (274 mg) potassium per dose. To be taken into consideration by patients with reduced kidney function or patients on a controlled potassium diet.

4.5 Interaction with other medicinal products and other forms of interaction

No interaction studies have been performed.

The co-administration of agents known to inhibit T cell function has not been formally studied. The co-administration of agents known to stimulate T cell function has not been investigated and the effects are unknown.

Tocilizumab ~~or siltuximab~~ and corticosteroid use

Some patients required tocilizumab ~~or siltuximab~~ and/or corticosteroid for the management of CRS (see section 4.8). The use of tocilizumab ~~or siltuximab~~ and/or ~~corticosteroids~~ for CRS management was more common in patients with higher cellular expansion.

~~Patients~~ In the KarMMA-3 study, patients with CRS treated with tocilizumab or siltuximab had higher Abecma cellular expansion levels, as measured by 3.1-fold and 2.9-fold higher median C_{max} (N = 156) and $AUC_{0-28days}$ (N = 155), respectively, compared to patients who did not receive tocilizumab or siltuximab (N = 64 for C_{max} and N = 63 for $AUC_{0-28days}$). Patients with CRS treated with corticosteroids had higher Abecma cellular expansion levels, as measured by 2.3-fold and 2.4-fold higher median C_{max} (N = 60) and $AUC_{0-28days}$ (N = 60), respectively, compared to patients who did not receive corticosteroids (N = 160 for C_{max} and N = 158 for $AUC_{0-28days}$).

Similarly, in the KarMMA study, patients with CRS treated with tocilizumab had higher Abecma cellular expansion levels, as measured by 1.4-fold and 1.6-fold higher median C_{max} (N = 66) and $AUC_{0-28days}$ (N = 65), respectively, compared to patients who did not receive tocilizumab (N = 61 for C_{max} and N = 60 for $AUC_{0-28days}$). ~~Similarly,~~ patients with CRS treated with corticosteroids had higher Abecma cellular expansion levels, as measured by 1.7-fold and 2.2-fold higher median C_{max} (N = 18) and $AUC_{0-28days}$ (N = 18), respectively, compared to patients who did not receive corticosteroids (N = 109 for C_{max} and N = 107 for $AUC_{0-28days}$).

Live vaccines

The safety of immunisation with live viral vaccines during or following ~~Abecma~~ treatment with Abecma has not been studied. As a precautionary measure, vaccination ~~Vaccination~~ with live ~~virus~~ vaccines is not recommended for at least 6 weeks prior to the start of lymphodepleting chemotherapy, during Abecma treatment and until immune recovery following treatment ~~with Abecma~~.

4.6 Fertility, pregnancy and lactation

Women of childbearing potential/Contraception in males and females

Pregnancy status for women of childbearing potential should be verified using a pregnancy test prior to starting treatment with Abecma.

See the prescribing information for fludarabine and cyclophosphamide for information on the need for effective contraception in patients who receive the lymphodepleting chemotherapy.

There are insufficient exposure data to provide a recommendation concerning duration of contraception following treatment with Abecma.

Pregnancy

There are no data from the use of idecabtagene vicleucel in pregnant women. No animal reproductive and developmental toxicity studies have been conducted with idecabtagene vicleucel to assess whether it can cause foetal harm when administered to a pregnant woman (see section 5.3).

It is not known if idecabtagene vicleucel has the potential to be transferred to the foetus. Based on the mechanism of action, if the transduced cells cross the placenta, they may cause foetal toxicity, including plasma cell aplasia or hypogammaglobulinaemia. Therefore, Abecma is not recommended for women who are pregnant or for women of childbearing potential not using contraception. Pregnant women should be advised on the potential risks to the foetus. Pregnancy after Abecma therapy should be discussed with the treating physician.

Assessment of immunoglobulin levels in newborn infants of mothers treated with Abecma should be considered.

Breast-feeding

It is unknown whether idecabtagene vicleucel cells are excreted in human milk or transferred to the breast-feeding child. A risk to the breast-fed infant cannot be excluded. Women who are breast-feeding should be advised of the potential risk to the breast-fed child.

Fertility

There are no data on the effect of idecabtagene vicleucel on fertility. Effects of idecabtagene vicleucel on male and female fertility have not been evaluated in animal studies.

4.7 Effects on ability to drive and use machines

Abecma may have major influence on the ability to drive and use machines.

Due to the potential for neurologic adverse reactions, including altered mental status or seizures with Abecma, patients receiving Abecma should refrain from driving or operating heavy or potentially dangerous machines for at least 8 weeks after Abecma infusion or until resolution of neurologic adverse reactions.

4.8 Undesirable effects

Summary of the safety profile

The safety data described in this section reflect the exposure to Abecma in the KarMMa₁ and CRB-401 and KarMMa-3 studies in which 184 409 patients with relapsed and refractory multiple myeloma received Abecma. ~~The~~ In KarMMa (N = 128) and CRB-401 (N = 56), the median duration of follow-up (from Abecma infusion to data cutoff date) was 15.520.8 months. In KarMMa-3 (N = 225), the median duration of follow-up was 29.3 months.

The most common adverse reactions ($\geq 20\%$) included CRS (84.6%), neutropenia (91.380.0%), ~~CRS (81.0%),~~ anaemia (70.763.6%), thrombocytopenia (66.855.0%), ~~infections - pathogen unspecified (53.8%), leucopenia (48.4%), fatigue (39.1%), diarrhoea (36.4%), hypokalaemia (34.243.8%), hypophosphataemia (32.633.3%), diarrhoea (33.0%), leukopenia (32.8%), hypokalaemia (32.0%), fatigue (29.8%), nausea (32.628.1%), lymphopenia (31.526.9%), pyrexia (28.824.7%), cough (27.2%), hypocalcaemia (26.6%), infections - viral (26.123.2%), headache (23.922.5%), hypocalcaemia (22.0%), hypomagnesaemia (22.1.3%), ~~upper respiratory tract infection (21.7%), arthralgia (20.7%), oedema peripheral (20.1%), decreased appetite (19.6%), hypogammaglobulinaemia (19.6%) and febrile neutropenia (16.3%)~~ and arthralgia (20.0%); other common adverse events occurring at lower frequency and considered clinically important included hypotension (18.6%), upper respiratory tract infection (15.6%), hypogammaglobulinemia (13.7%), febrile neutropenia (11.2%), pneumonia (10.11.03%), tremor (8.25.6%), somnolence (5.46%), aphasia (4.3%), encephalopathy (4.33.4%), ~~and~~ syncope (4.33.2%) and aphasia (2.9%).~~

Serious adverse reactions occurred in 70.157.2% of patients. The most common serious adverse reactions ($\geq 5\%$) included CRS (17.410.3 %) and pneumonia (7.1%;%), ~~febrile neutropenia (6.0%) and pyrexia (6.0%);~~ other serious adverse events occurring at lower frequency and considered clinically important include febrile neutropenia (4.32%), pyrexia (3.7%), neutropenia (2.7%), sepsis (3.82.7%), thrombocytopenia (3.8%), confusional state (2.22.4%), haemophagocytic lymphohistiocytosis (1.7%), thrombocytopenia (1.5%), encephalopathy (1.5%), dyspnoea (2.21.5%), seizure (1.0%), hypoxia (1.6%), mental status changes (1.60%), hypoxia (0.7%) and disseminated intravascular coagulation (0.5%) encephalopathy (1.6%).

The most common Grade 3 or 4 adverse reactions ($\geq 5\%$) were neutropenia (88.677.3%), anaemia (58.250.9%), thrombocytopenia (53.42.5%), leukopenia (45.131.5%), lymphopenia (30.425.9%), hypophosphataemia (19.8%), infections - pathogen unspecified (17.915.2%), hypophosphataemia (17.4%), febrile neutropenia (14.10.75%), hypocalcaemia (7.1%), infections - viral (7.16%), pneumonia (6.08%), CRS (5.4%), hypertension (5.6.46%), hypocalcaemia (5.6%) and infections - bacterial (5.4%) and hyponatraemia (5.4%).

Grade 3 or 4 adverse reactions were more often observed within the initial 8 weeks post-infusion (97.893.2%) compared to after 8 weeks post-infusion (60.858.1%). The most frequently reported Grade 3 or 4 adverse reactions reported within the first 8 weeks after infusion were neutropenia (87.075.8%), anaemia (56.47.4.0%), thrombocytopenia (48.38.6.4%), leukopenia (44.030.3%) lymphopenia (27.723.5%) and hypophosphataemia (16.18.3%).

Tabulated list of adverse reactions

Table 3 summarises the adverse reactions observed in ~~the clinical studies of 409, 128 and 56~~ patients treated with Abecma ~~across~~within the ~~target-allowed~~ dose ~~levels-range~~ of 150 to ~~450-540~~ x 10⁶ CAR-positive T cells (see Table 4-6 in section 5.1 for the corresponding dose range of CAR-positive viable T cells) ~~in the KarMMA and CRB 401 studies, respectively) and from post-marketing reports.~~

Adverse reactions are presented by MedDRA system organ class and by frequency.

Frequencies are defined as: very common (≥ 1/10), common (≥ 1/100 to < 1/10), uncommon (≥ 1/1,000 to < 1/100), rare (≥ 1/10,000 to < 1/1,000), very rare (< 1/10,000) and not known (cannot be estimated from available data). Within each frequency grouping, adverse reactions are presented in order of decreasing seriousness.

Table 3. Adverse reactions observed in patients treated with Abecma

| System organ class | Adverse reaction | All grades frequency |
|--|---|---|
| Infections and infestations ^a | Infections – bacterial Infections – viral Infections – pathogen unspecified Infections – fungal | Very common Very common Very common Common |
| Blood and lymphatic system disorders | Neutropenia Leucopenia <u>Leukopenia</u> Thrombocytopenia Febrile neutropenia Lymphopenia Anaemia Disseminated intravascular coagulation | Very common Very common Very common Very common Very common Very common Common |
| Immune system disorders | Cytokine release syndrome Hypogammaglobulinaemia Haemophagocytic lymphohistiocytosis* | Very common Very common Common |
| Metabolism and nutrition disorders | Hypophosphataemia Hypokalaemia Hyponatraemia Hypocalcaemia Hypoalbuminaemia Decreased appetite Hypomagnesaemia | Very common Very common Very common Very common Very common Very common Very common |
| Psychiatric disorders | <u>Insomnia</u> Delirium ^b Insomnia | <u>Very c</u> Common Common |

| System organ class | Adverse reaction | All grades frequency |
|--|---|--|
| Nervous system disorders | Encephalopathy ^c Headache* Dizziness ^d <u>Aphasia^c</u> <u>Ataxia^f</u> <u>Motor dysfunction^g</u> <u>Tremor</u> Seizure Hemiparesis <u>Immune effector cell-associated neurotoxicity syndrome**</u> <u>Aphasia^e</u> <u>Ataxia^f</u> <u>Motor-dysfunction^g</u> <u>Tremor</u> | Very common Very common Very common Common Common Common Common Common Common Uncommon <u>Uncommon</u> |
| Cardiac disorders | Tachycardia* Atrial fibrillation* | Very common Common |
| Vascular disorders | Hypertension Hypotension* ^h | Very common Very common |
| Respiratory, thoracic, and mediastinal disorders | Dyspnoea Cough Pulmonary oedema Hypoxia* | Very common Very common Common Common |
| Gastrointestinal disorders | Vomiting Diarrhoea Nausea Constipation Gastrointestinal haemorrhage ⁱ | Very common Very common Very common Very common Common |
| Musculoskeletal and connective tissue disorders | Arthralgia Myalgia | Very common Common |
| General disorders and administration site conditions | Pyrexia* Fatigue* ^j <u>Oedema^k</u> <u>Chills*</u> Asthenia <u>Oedema^k</u> <u>Chills*</u> | Very common Very common Very common Very common Very Common |
| Investigations | Alanine aminotransferase increased Aspartate aminotransferase increased Blood alkaline phosphatase increased C-reactive protein increased* | Very common Very common Very common Common Common |

* Event that has been reported as a manifestation of CRS.

** Event was not systematically collected in clinical trials.

^a Infections and infestations system organ class adverse events are grouped by pathogen type and selected clinical syndromes.

^b Delirium includes delirium, disorientation, agitation, hallucination, restlessness.

^c Encephalopathy includes amnesia, bradyphrenia, cognitive disorder, confusional state, depressed level of consciousness, disturbance in attention, dyscalculia, dysgraphia, encephalopathy, incoherent, lethargy, memory impairment, mental impairment, mental status changes, metabolic encephalopathy, neurotoxicity, somnolence, toxic encephalopathy stupor.

^d Dizziness includes dizziness, presyncope, syncope, vertigo.

- ^e Aphasia includes aphasia, dysarthria, slow speech, and speech disorder.
- ^f Ataxia includes ataxia, dysmetria, gait disturbance.
- ^g Motor dysfunction includes motor dysfunction, muscular spasms, muscular weakness, parkinsonism.
- ^h Hypotension includes hypotension, orthostatic hypotension.
- ⁱ Gastrointestinal haemorrhage includes gastrointestinal haemorrhage, gingival bleeding, haematochezia, haemorrhoidal haemorrhage, melaena, mouthhaemorrhage.
- ^j Fatigue includes fatigue, malaise.
- ^k Oedema includes oedema, oedema peripheral, face oedema, generalised oedema, peripheral oedema, peripheral swelling.

Description of selected adverse reactions

Cytokine release syndrome

In the pooled studies (KarMMa, ~~and~~ CRB-401 and KarMMa-3), CRS occurred in 81.084.6% of patients receiving Abecma. Grade 3 or higher CRS (Lee et al, 2014) occurred in 5.41% of patients, with fatal (Grade 5) CRS reported in 0.57% of patients. The median time-to-onset, any grade, was 1 day (range: 1 to 17) and the median duration of CRS was 5.4 days (range: 1 to 63).

The most common manifestations of CRS ($\geq 10\%$) included pyrexia (78.382.6%), hypotension (32.429.1%), tachycardia (25.524.7%), chills (23.418.8%), hypoxia (16.315.9%), ~~C-reactive protein increased (16.3%),~~ headache (14.711.2%) and ~~fatigue (10.9%), increased C-reactive protein (10.5%).~~ Grade 3 or higher events that may be observed in association with CRS included atrial fibrillation, capillary leak syndrome, hypotension, hypoxia and HLH/MAS.

Of the 184.409 patients, 45.459.7% of patients received tocilizumab; 3237.62% received a single dose while 4222.5% received more than 1 dose of tocilizumab for treatment of CRS. Overall, ~~across the target dose levels,~~ 15.822.7% of patients received at least 1 dose of corticosteroids for treatment of CRS. Of the 92 patients, in KarMMa and CRB-401 who received at the target dose of 450×10^6 CAR-positive T cells, 54.3% of patients received tocilizumab and 22.8% received at least 1 dose of corticosteroids for treatment of CRS. Of the 225 patients in KarMMa-3 who received Abecma infusion, 71.6% of patients received tocilizumab and 28.4% received at least 1 dose of corticosteroids for the treatment of CRS. See section 4.4 for monitoring and management guidance.

Neurologic adverse reactions including ICANS

In the pooled studies, of the 184.409 patients, independent of investigator attribution of neurotoxicity, the most frequent neurologic or psychiatric adverse reactions ($\geq 5\%$) included headache (28.822.5%), dizziness (15.12.5%), confusional state (13.11.0%), insomnia (9.810.3%), anxiety (8.25.9%), tremor (8.25.6%), and somnolence (6.55.6%). Other neurological adverse reactions occurring at a lower frequency and considered clinically important included ~~aphasia (4.3%) and~~ encephalopathy (4.33.4%) ~~and aphasia (2.9%)~~.

Neurotoxicity identified by the investigators, which was the primary method of assessing CAR T cell-associated neurotoxicity in the KarMMa and KarMMa-3 studies ~~study only~~, occurred in 57 (48.16.01%) of the 128.353 patients receiving Abecma, including Grade 3 or 4 in 3.1% of patients (with no Grade ~~4 or~~ 5 events). The median time to onset of the first event was 2.3 days (range: 1 to 317; one patient developed encephalopathy at Day 317 as a result of worsening pneumonia and Clostridium difficile colitis), 40). The median duration was 3 days (range: 1 to 252; one patient developed neurotoxicity [highest Grade 3] 43 days after ide-cel infusion which resolved after 252 days) 26). Overall, 7.81% of patients received at least 1 dose of corticosteroid for treatment of CAR T cell-associated neurotoxicity ~~:-~~.

In KarMMa, across the target dose levels, 7.8% of patients received at least 1 dose of corticosteroid for treatment of CAR T cell-associated neurotoxicity, while at the target dose of 450×10^6 CAR-positive T cells, 14.8% of patient-s received at least 1 dose of corticosteroids.

In KarMMa-3, across all patients who received Abecma infusion at the target dose range, 6.7% of patients received at least 1 dose of corticosteroid for treatment of CAR T cell-associated neurotoxicity.

Of the 353 patients in the KarMMa and KarMMa-3 studies, the most common manifestations of investigator identified neurotoxicity ($\geq 2\%$) included confusional state (9.48.5%), encephalopathy (5.53.4%), somnolence (2.8%), aphasia (42.75%), hallucination (3.1%), and mental status changes (3.1%), tremor (2.3%), disturbance in attention (2.0%) and dysgraphia (2.0%). See section 4.4 for monitoring and management guidance.

Febrile neutropenia and infections

In the pooled studies, infections occurred in 71.262.8% of patients. Grade 3 or 4 infections occurred in 23.423.2% of patients. Grade 3 or 4 infections with an unspecified pathogen occurred in 17.915.2%, viral infections in 7.16%, bacterial infections in 3.84.6% and fungal infections in 0.51.2% of patients. Fatal infections of unspecified pathogen were reported in 1.62.0% of patients and 0.57% of patients had fatal fungal or viral infection and 0.2% of patients had fatal bacterial infection. See section 4.4 for monitoring and management guidance.

Febrile neutropenia (Grade 3 or 4) was observed in 14.710.8% of patients after Abecma infusion. Febrile neutropenia may be concurrent with CRS. See section 4.4 for monitoring and management guidance.

Prolonged cytopenia

Patients may exhibit prolonged cytopenias following lymphodepleting chemotherapy and Abecma infusion. In the pooled studies, 34.838.2% of the 178.395 patients who had Grade 3 or 4 neutropenia and 72.771.3% of the 110.230 patients who had Grade 3 or 4 thrombocytopenia during the first month following Abecma infusion had not resolved by last assessment during the first month. Among the 62.151 patients with neutropenia not resolved by month 1, 82.388.7% recovered from Grade 3 or 4 neutropenia with a median time to recovery from Abecma infusion of 1.9 months. Of the 80.164 patients with thrombocytopenia not resolved by month 1, 71.379.9% recovered from Grade 3 or 4 thrombocytopenia with the median time to recovery of 2.2-0 months. See section 4.4 for monitoring and management guidance.

Hypogammaglobulinaemia

Hypogammaglobulinaemia was reported in 19.613.7% of patients treated with Abecma in the pooled studies with a median time to onset of 100.90 days (range 15-1 to 326). See section 4.4 for monitoring and management guidance.

Immunogenicity

Abecma has the potential to induce anti-CAR antibodies. In clinical studies, humoral immunogenicity of Abecma was measured by determination of anti-CAR antibody in serum pre- and post-administration. In the pooled studies of KarMMa, CRB-401 and KarMMa-4.3, 3.2% of patients tested positive for pre-infusion anti-CAR antibodies and post-infusion anti-CAR antibodies were detected in 50.556.2% of the patients. There is no evidence that the presence of pre-existing or post-infusion anti-CAR antibodies impact the cellular expansion, safety or effectiveness of Abecma.

Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product.

Any suspected adverse events should be reported to the Ministry of Health according to the National Regulation by using an online form <https://sideeffects.health.gov.il/>

4.9 Overdose

There are limited data regarding overdose with Abecma.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: ~~Other antineoplastic agents, not yet assigned,~~

ATC code: ~~L01XL07 not yet assigned~~

Mechanism of action

Abecma is a chimeric antigen receptor (CAR)-positive T cell therapy targeting B-cell maturation antigen (BCMA), which is expressed on the surface of normal and malignant plasma cells. The CAR construct includes an anti-BCMA scFv-targeting domain for antigen specificity, a transmembrane domain, a CD3-zeta T cell activation domain, and a 4-1BB costimulatory domain. Antigen-specific activation of Abecma results in CAR-positive T cell proliferation, cytokine secretion and subsequent cytolytic killing of BCMA-expressing cells.

Clinical efficacy and safety

KarMMA-3

KarMMA-3 was an open-label, multicentre, randomised, controlled study that evaluated the efficacy and safety of Abecma, compared to standard regimens, in adult patients with relapsed and refractory multiple myeloma who had received two to four prior antimyeloma regimens including an immunomodulatory agent, a proteasome inhibitor, and daratumumab, and were refractory to the most recent prior antimyeloma regimen. A standard regimen was assigned to each patient prior to randomisation, contingent upon the patient's most recent antimyeloma treatment. The standard regimens consisted of daratumumab, pomalidomide, dexamethasone (DPd), daratumumab, bortezomib, dexamethasone (DvD), ixazomib, lenalidomide, dexamethasone (IRd), carfilzomib, dexamethasone (Kd), or elotuzumab, pomalidomide, dexamethasone (EPd). In patients randomised to the Abecma arm, the assigned standard regimen was to be used as bridging therapy, if clinically indicated.

The study included patients who achieved a response (minimal response or better) to at least 1 prior treatment regimen and had ECOG performance status of 0 or 1. The study excluded patients with CNS involvement of myeloma, history of CNS disorders (such as seizures), prior allogeneic SCT or prior treatment with any gene therapy-based therapeutic for cancer, investigational cellular therapy for cancer or BCMA targeted therapy, ongoing treatment with immunosuppressants, serum creatinine clearance < 45 mL/min, serum aspartate aminotransferase (AST) or alanine aminotransferase (ALT) > 2.5 times upper limit of normal, and left ventricular ejection fraction (LVEF) < 45%. Patients were also excluded if absolute neutrophil count < 1000/ μ L and platelet count < 75,000/ μ L in patients in whom < 50% of bone marrow nucleated cells are plasma cells and platelet count < 50,000/ μ L in patients in whom > 50% of bone marrow nucleated cells are plasma cells.

Patients were randomised 2:1 to receive either Abecma (N = 254) or standard regimens (N = 132) for relapsed and refractory multiple myeloma. Randomisation was stratified by age, number of prior antimyeloma regimens and high-risk cytogenetics abnormalities. Patients receiving standard regimens were allowed to receive Abecma upon confirmed disease progression.

Patients randomised to Abecma were to receive lymphodepleting chemotherapy consisting of cyclophosphamide (300 mg/m² IV infusion daily for 3 days) and fludarabine (30 mg/m² IV infusion daily for 3 days) starting 5 days prior to the target infusion date of Abecma. Up to 1 cycle of DPd, DvD, IRd, Kd, or EPd anticancer therapy for disease control (bridging therapy) was permitted between apheresis and until 14 days before the start of lymphodepleting chemotherapy.

Of the 254 patients randomised to Abecma, 249 (98%) patients underwent leukapheresis, and 225 (88.6%) patients received Abecma. Of the 225 patients, 192 (85.3%) patients received bridging therapy. Twenty-nine patients did not receive Abecma due to death (n = 4), adverse event (n = 5), patient withdrawal (n = 2), physician decision (n = 7), failure to meet lymphodepleting chemotherapy treatment

criteria (n = 8) or manufacturing failure (n = 3).

The allowed dose range was 150 to 540 x 10⁶ CAR-positive T cells. The median actual received dose was 445.3 x 10⁶ CAR-positive T cells (range: 174.9 to 529.0 x 10⁶ CAR-positive T cells). The median time from leukapheresis to product availability was 35 days (range: 24 to 102 days) and the median time from leukapheresis to infusion was 49 days (range: 34 to 117 days).

Of the 132 patients randomised to standard regimens, 126 (95.5%) patients received treatment. Six patients discontinued without receiving treatment due to disease progression (n = 1), patient withdrawal (n = 3), or physician decision (n = 2). Patients receiving standard regimens were allowed to receive Abecma at investigator's request, upon confirmed disease progression by the independent review committee (IRC) based on the International Myeloma Working Group (IMWG) criteria and confirmed eligibility. Of the eligible patients, 69 (54.8%) underwent leukapheresis and 60 (47.6%) received Abecma.

Table 4 summarises the baseline patient and disease characteristics in KarMMa-3 study.

Table 4. Baseline demographic/disease characteristics for patients in KarMMa-3 study

| Characteristic | Abecma (N = 254) | Standard regimens (N = 132) |
|---|-----------------------------|--|
| Age (years) | | |
| Median (min, max) | 63 (30, 81) | 63 (42, 83) |
| ≥ 65 years, n (%) | 104 (40.9) | 54 (40.9) |
| ≥ 75 years, n (%) | 12 (4.7) | 9 (6.8) |
| Gender, male, n (%) | 156 (61.4) | 79 (59.8) |
| Race, n (%) | | |
| Asian | 7 (2.8) | 5 (3.8) |
| Black | 18 (7.1) | 18 (13.6) |
| White | 172 (67.7) | 78 (59.1) |
| ECOG performance status, n (%)^a | | |
| 0 | 120 (47.2) | 66 (50.0) |
| 1 | 133 (52.4) | 62 (47.0) |
| 2 | 0 | 3 (2.3) |
| 3 | 1 (0.4) | 1 (0.8) |
| Patients with extramedullary plasmacytoma, n (%) | 61 (24.0) | 32 (24.2) |
| Time since initial diagnosis (year) | | |
| n | 251 | 131 |
| median (min, max) | 4.1 (0.6, 21.8) | 4.0 (0.7, 17.7) |
| Prior stem cell transplant, n (%) | 214 (84.3) | 114 (86.4) |
| Baseline cytogenetic abnormality, n (%)^b | | |
| High risk ^c | 107 (42.1) | 61 (46.2) |
| Non-high risk | 114 (44.9) | 55 (41.7) |
| Not evaluable/Missing | 33 (13.0) | 16 (12.1) |
| Revised ISS stage at baseline (derived)^d, n (%) | | |
| Stage I | 50 (19.7) | 26 (19.7) |

| | | |
|--|--------------------------|-------------------------|
| <u>Stage II</u> | <u>150 (59.1)</u> | <u>82 (62.1)</u> |
| <u>Stage III</u> | <u>31 (12.2)</u> | <u>14 (10.6)</u> |
| <u>Unknown</u> | <u>23 (9.1)</u> | <u>10 (7.6)</u> |
| <u>Distribution of prior anti-myeloma regimens, n (%)</u> | | |
| <u>2</u> | <u>78 (30.7)</u> | <u>39 (29.5)</u> |
| <u>3</u> | <u>95 (37.4)</u> | <u>49 (37.1)</u> |
| <u>4</u> | <u>81 (31.9)</u> | <u>44 (33.3)</u> |
| <u>Refractory status to prior classes of therapy, n (%)</u> | | |
| <u>IMiD</u> | <u>224 (88.2)</u> | <u>124 (93.9)</u> |
| <u>Proteasome inhibitor (PI)</u> | <u>189 (74.4)</u> | <u>95 (72.0)</u> |
| <u>Anti-CD38 antibodies</u> | <u>242 (95.3)</u> | <u>124 (93.9)</u> |
| <u>Triple refractory^e, n (%)</u> | <u>164 (64.6)</u> | <u>89 (67.4)</u> |

ECOG = Eastern Cooperative Oncology Group; IMiD = immunomodulatory agents; ISS = International Staging System; max = maximum; min = minimum

^a All subjects had ECOG score 0 or 1 at screening, but the ECOG score may be >1 at baseline.

^b Baseline cytogenetic abnormality was based on baseline cytogenetics from central laboratory if available. If central laboratory was not available or was unknown, cytogenetics prior to screening was used.

^c High-risk defined as deletion in chromosome 17p (del[17p]), translocation involving chromosomes 4 and 14 (t[4;14]) or translocation involving chromosomes 14 and 16 (t[14;16]).

^d Revised ISS was derived using baseline ISS stage, cytogenetic abnormality and serum lactate dehydrogenase.

^e Triple refractory is defined as refractory to an immunomodulatory agent, a proteasome inhibitor and an anti-CD38 antibody.

The primary efficacy endpoint was progression free survival (PFS) according to the IMWG Uniform Response Criteria for Multiple Myeloma as determined by an Independent Review Committee (IRC). Other efficacy measures included overall response rate (ORR), overall survival (OS) and patient-reported outcomes. At a pre-specified interim analysis at 80% information fraction with a median follow up time of 18.6 months, Abecma demonstrated a statistically significant improvement in PFS compared to the standard regimens arm; HR = 0.493 (95% CI: 0.38, 0.65, two-sided p-value < 0.0001). The results of the subsequent primary analysis (shown in Table 5 and Figure 1), with a median follow-up time of 30.9 months, were consistent with the interim analysis.

Table 5. Summary of efficacy results from KarMMA-3 (intent-to-treat population)

| | <u>Abecma arm</u> <u>(N = 254)</u> | <u>Standard regimens arm</u> <u>(N = 132)</u> |
|--|---|--|
| <u>Progression free survival</u> | | |
| <u>Number of events, n (%)</u> | <u>184 (72.4)</u> | <u>105 (79.5)</u> |
| <u>Median, months [95% CI]^a</u> | <u>13.8 [11.8, 16.1]</u> | <u>4.4 [3.4, 5.8]</u> |
| <u>Hazard ratio [95% CI]^b</u> | <u>0.49 [0.38, 0.63]</u> | |
| <u>Overall response rate</u> | | |
| <u>n (%)</u> | <u>181 (71.3)</u> | <u>56 (42.4)</u> |
| <u>95% CI (%)^c</u> | <u>(65.7, 76.8)</u> | <u>(34.0, 50.9)</u> |
| <u>CR or better (sCR+CR)</u> | <u>111 (43.7)</u> | <u>7 (5.3)</u> |
| <u>sCR</u> | <u>103 (40.6)</u> | <u>6 (4.5)</u> |
| <u>CR</u> | <u>8 (3.1)</u> | <u>1 (0.8)</u> |
| <u>VGPR</u> | <u>45 (17.7)</u> | <u>15 (11.4)</u> |

| | <u>Abecma arm</u> <u>(N = 254)</u> | <u>Standard regimens arm</u> <u>(N = 132)</u> |
|---|---------------------------------------|--|
| <u>PR</u> | <u>25 (9.8)</u> | <u>34 (25.8)</u> |
| <u>DOR if best response is CR</u> | | |
| <u>N</u> | <u>111</u> | <u>7</u> |
| <u>Median, months [95% CI]</u> | <u>15.7 [12.1, 22.1]</u> | <u>24.1 [4.6, NA]</u> |
| <u>DOR if best response is PR</u> | | |
| <u>N</u> | <u>181</u> | <u>56</u> |
| <u>Median, months [95% CI]</u> | <u>16.5 [12.0, 19.4]</u> | <u>9.7 [5.4, 15.5]</u> |
| <u>MRD-negative status by NGS and ≥ CR</u> | | |
| <u>MRD negativity rate, n (%)^d</u> | <u>57 (22.4)</u> | <u>1 (0.8)</u> |
| <u>95% CI (%)^c</u> | <u>(17.3, 27.6)</u> | <u>(0.0, 2.2)</u> |

CI=confidence interval; CR=complete response; DOR=duration of response; MRD=minimal residual disease; PR=partial response; sCR=stringent complete response; VGPR=very good partial response.

^a Kaplan-Meier estimate.

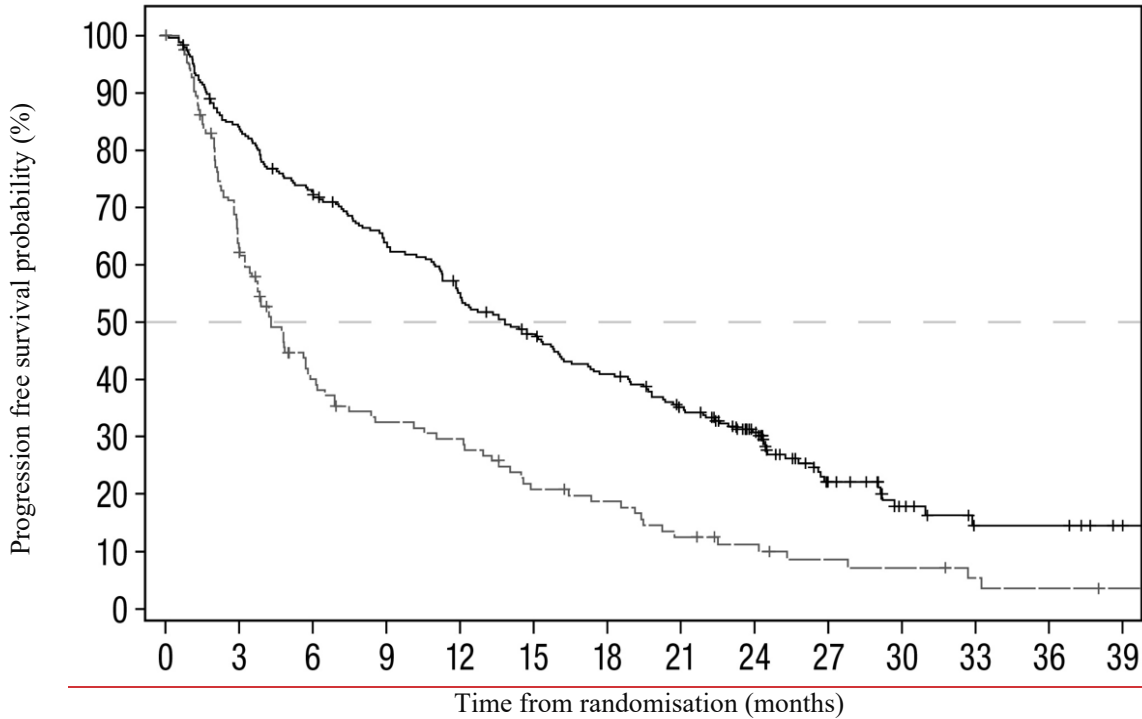
^b Based on stratified univariate Cox proportional hazards model.

^c Two-sided Wald confidence interval.

^d MRD negativity was defined as the proportion of all patients in the ITT population who achieved CR or stringent CR and are MRD negative at any timepoint within 3 months prior to achieving CR or stringent CR until the time of progression or death. Based on a threshold of 10⁻⁵ using ClonoSEQ, a next-generation sequencing (NGS) assay.

Figure 1. Kaplan-Meier plot of progression-free survival based on IRC assessment in KarMMa-

3 study (intent-to-treat population)



Number of subjects at risk

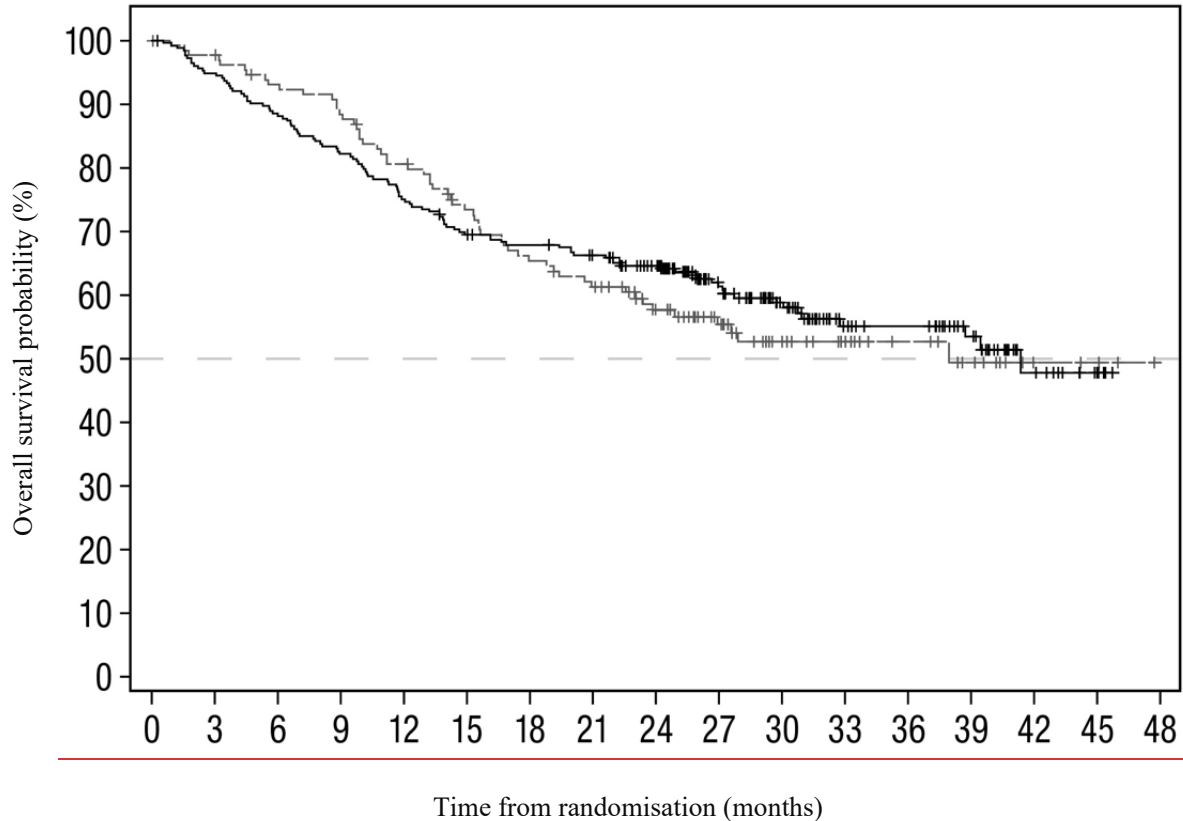
| | 0 | 3 | 6 | 9 | 12 | 15 | 18 | 21 | 24 | 27 | 30 | 33 | 36 | 39 |
|-------------------------|-----|-----|-----|-----|-----|-----|----|----|----|----|----|----|----|----|
| — Abecma | 254 | 206 | 177 | 153 | 131 | 111 | 94 | 77 | 54 | 25 | 14 | 7 | 7 | 2 |
| - - - Standard regimens | 132 | 76 | 43 | 34 | 31 | 21 | 18 | 12 | 9 | 6 | 5 | 3 | 2 | 1 |

At the time of the final analysis for PFS, 74% of planned OS events were reached. Patients receiving standard regimens were allowed to receive Abecma upon confirmed disease progression; the OS data are therefore confounded by 74 (56.1%) patients from the standard regimen arm who received Abecma as a subsequent therapy. The median OS for Abecma was 41.4 months (95% CI: 30.9, NR) versus standard regimens 37.9 months (95% CI: 23.4, NR); HR = 1.01 (95% CI: 0.73, 1.40). Figure 2 shows the Kaplan-Meier curve for OS in the intent-to-treat population (not corrected for cross-over).

Compared to the standard regimens arm (9/132; 6.8%), a higher proportion of patients experienced death within 6 months after randomisation in the Abecma arm (30/254; 11.8%). Of the 30 patients with an early death event in the Abecma arm, 17 patients never received Abecma treatment, and 13 of these 17 died of disease progression. High-risk factors such as high-risk cytogenetic abnormalities, R-ISS stage III, presence of extramedullary plasmacytoma or high tumour burden (see section 4.4 on rapidly progressing disease) are associated with higher risk of early death.

Figure 2. Kaplan-Meier plot of overall survival based on IRC assessment in KarMMA-3 study

(intent-to-treat population)



Number of subjects at risk

| | | | | | | | | | | | | | | | | | |
|------------------------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|----|----|----|----|----|---|---|
| <u>—</u> Abecma | 254 | 240 | 223 | 208 | 190 | 175 | 169 | 161 | 143 | 103 | 75 | 48 | 44 | 30 | 13 | 4 | 0 |
| <u>---</u> Standard regimens | 132 | 128 | 120 | 114 | 103 | 91 | 81 | 75 | 59 | 45 | 32 | 24 | 18 | 11 | 4 | 3 | 0 |

KarMMA

KarMMA was an open-label, single-arm, multicentre study that evaluated the efficacy and safety of Abecma in adult patients with relapsed and refractory multiple myeloma who had received at least 3 prior antimyeloma therapies including an immunomodulatory agent, a proteasome inhibitor and an anti-CD38 antibody and who were refractory to the last treatment regimen. Patients with CNS involvement of myeloma, a history of other BCMA targeting therapies, allogeneic SCT or prior genetherapy based or other genetically modified T cell therapy were excluded. Patients with a history of CNS disorders (such as seizures), inadequate hepatic, renal, bone marrow function, cardiac, pulmonary function or ongoing treatment with immunosuppressants were excluded.

The study consisted of pre-treatment (screening, leukapheresis and bridging therapy [if needed]); treatment (lymphodepleting chemotherapy and Abecma infusion); and posttreatment (ongoing) for a minimum of 24 months following Abecma infusion or until documented disease progression, whichever was longer. The lymphodepleting chemotherapy period was one 3-day cycle of cyclophosphamide (300 mg/m² IV infusion daily for 3 days) and fludarabine (30 mg/m² IV infusion daily for 3 days) starting 5 days prior to the target infusion date of Abecma. Patients were hospitalised for 14 days after infusion of Abecma to monitor and manage potential CRS and neurotoxicity.

Of the 140 patients who were enrolled (i.e. underwent leukapheresis), 128 patients received the Abecma

infusion. Out of the 140 patients, only one did not receive the product due to manufacturing failure. Eleven other patients were not treated with Abecma, due to physician decision (n = 3), patient withdrawal (n = 4), adverse events (n = 1), progressive disease (n = 1) or death (n = 2) prior to receiving Abecma.

Anticancer therapy for disease control (bridging) was permitted between apheresis and lymphodepletion with the last dose being administered at least 14 days prior to initiation of lymphodepleting chemotherapy. Of the 128 patients treated with Abecma, most patients (87.5%) received anticancer therapy for disease control at the discretion of the investigator.

The doses targeted in the clinical study were 150, 300 or 450 x 10⁶ CAR-positive T cells per infusion. The allowed dose range was 150 to 540 x 10⁶ CAR-positive T cells. Table 4-6 below shows the target dose levels used in the clinical study based on total CAR-positive T cells and the corresponding range of actual dose administered defined as CAR-positive viable T cells.

Table 46. Total CAR-positive T cells dose with the corresponding dose range of CAR-positive viable T cells (x10⁶) - KarMMa study

| Target dose based on total CAR-positive T cells, including both viable and non-viable cells (x10 ⁶) | CAR-positive viable T cells (x10 ⁶) (min, max) |
|---|--|
| 150 | 133 to 181 |
| 300 | 254 to 299 |
| 450 | 307 to 485 |

Table 5-7 summarises the baseline patient and disease characteristics for the enrolled and treated population in study.

Table 57. Baseline demographic/disease characteristics for study population- KarMMa study

| Characteristic | Total enrolled (N = 140) | Total treated (N = 128) |
|--|--------------------------|-------------------------|
| Age (years) | | |
| Median (min, max) | 60.5 (33, 78) | 60.5 (33, 78) |
| ≥ 65 years, n (%) | 48 (34.3) | 45 (35.2) |
| ≥ 75 years, n (%) | 5 (3.6) | 4 (3.1) |
| Gender, male, n (%) | 82 (58.6) | 76 (59.4) |
| Race, n (%) | | |
| Asian | 3 (2.1) | 3 (2.3) |
| Black | 8 (5.7) | 6 (4.7) |
| White | 113 (80.7) | 103 (80.5) |
| ECOG performance status, n (%) | | |
| 0 | 60 (42.9) | 57 (44.5) |
| 1 | 77 (55.0) | 68 (53.1) |
| 2 ^a | 3 (2.1) | 3 (2.3) |
| Patients with extramedullary plasmacytoma, n (%) | 52 (37.1) | 50 (39.1) |
| Time since initial diagnosis (years), median (min, max) | 6 (1.0, 17.9) | 6 (1.0, 17.9) |
| Prior stem cell transplant, n (%) | 131 (93.6) | 120 (93.8) |
| Baseline cytogenetic high risk^{b,c} | 46 (32.9) | 45 (35.2) |

| | | |
|--|------------|------------|
| Revised ISS stage at baseline (derived)^d, n (%) | | |
| Stage I | 14 (10.0) | 14 (10.9) |
| Stage II | 97 (69.3) | 90 (70.3) |
| Stage III | 26 (18.6) | 21 (16.4) |
| Unknown | 3 (2.1) | 3 (2.3) |
| Number of prior anti-myeloma therapies^e, median (min, max) | 6 (3, 17) | 6 (3, 16) |
| Triple refractory^f, n (%) | 117 (83.6) | 108 (84.4) |
| Creatinine clearance (mL/min), n (%) | | |
| < 30 | 3 (2.1) | 1 (0.8) |
| 30 to < 45 | 9 (6.4) | 8 (6.3) |
| 45 to < 60 | 13 (9.3) | 10 (7.8) |
| 60 to < 80 | 38 (27.1) | 36 (28.1) |
| ≥ 80 | 77 (55.0) | 73 (57.0) |

max = maximum; min = minimum

^a These patients had ECOG scores of < 2 at screening for eligibility but subsequently deteriorated to ECOG scores of ≥ 2 at baseline prior to start of LD chemotherapy.

^b Baseline cytogenetic abnormality was based on baseline cytogenetics from central laboratory if available. If central laboratory was not available or was unknown, cytogenetics prior to screening was used.

^c High-risk defined as deletion in chromosome 17p (del[17p]), translocation involving chromosomes 4 and 14 (t[4;14]) or translocation involving chromosomes 14 and 16 (t[14;16]).

^d Revised ISS was derived using baseline ISS stage, cytogenetic abnormality and serum lactate dehydrogenase.

^e Induction with or without haematopoietic stem cell transplant and with or without maintenance therapy was considered a single therapy.

^f Triple refractory is defined as refractory to an immunomodulatory agent, a proteasome inhibitor and an anti-CD38 antibody.

The median time from leukapheresis to product availability was 32 days (range: 24 to 55 days) and the median time from leukapheresis to infusion was 40 days (range: 33 to 79 days). The median actual dose received across all doses targeted in the clinical study was 315.3 x 10⁶ CAR-positive T cells (range 150.5 to 518.4).

Efficacy was assessed on the basis of overall response rate (ORR), complete response (CR) rate and duration of response (DOR), as determined by an independent review committee. Other efficacy endpoints included minimal residual disease (MRD) using next-generation sequencing (NGS).

Efficacy results across doses targeted in the clinical study (150 to 450 x 10⁶ CAR-positive T cells) are shown in the Table 68. Median follow-up was 19.9 months for all Abecma treated patients.

Table 68. Summary of efficacy based on the KarMMA study

| | Enrolled ^a (N = 140) | Treated population Target dose of Abecma (CAR-positive T cells) | | | |
|--|------------------------------------|--|-----------------------------------|-----------------------------------|--|
| | | 150 x 10 ^{6b} (N = 4) | 300 x 10 ⁶ (N = 70) | 450 x 10 ⁶ (N = 54) | Total 150 to 450 x 10 ⁶ (N = 128) |
| Overall response rate (sCR+CR+VGPR+PR), n (%) | 94 (67.1) | 2 (50.0) | 48 (68.6) | 44 (81.5) | 94 (73.4) |
| 95% CI ^c | 59.4, 74.9 | 6.8, 93.2 | 56.4, 79.1 | 68.6, 90.7 | 65.8, 81.1 |
| CR or better, n (%) | 42 (30.0) | 1 (25.0) | 20 (28.6) | 21 (38.9) | 42 (32.8) |

| | Enrolled ^a (N = 140) | Treated population Target dose of Abecma (CAR-positive T cells) | | | |
|---|------------------------------------|--|-----------------------------------|-----------------------------------|--|
| | | 150 x 10 ^{6b} (N = 4) | 300 x 10 ⁶ (N = 70) | 450 x 10 ⁶ (N = 54) | Total 150 to 450 x 10 ⁶ (N = 128) |
| 95% CI ^c | 22.4, 37.6 | 0.6, 80.6 | 18.4, 40.6 | 25.9, 53.1 | 24.7, 40.9 |
| VGPR or better, n (%) | 68 (48.6) | 2 (50.0) | 31 (44.3) | 35 (64.8) | 68 (53.1) |
| 95% CI ^c | 40.3, 56.9 | 6.8, 93.2 | 32.4, 56.7 | 50.6, 77.3 | 44.5, 61.8 |
| MRD-negative status^d and ≥ CR | | | | | |
| Based on treated patients | – | 4 | 70 | 54 | 128 |
| n (%) | – | 1 (25.0) | 17 (24.3) | 14 (25.9) | 32 (25.0) |
| 95% CI | – | 0.6, 80.6 | 14.8, 36.0 | 15.0, 39.7 | 17.8, 33.4 |
| Time to response, n | 94 | 2 | 48 | 44 | 94 |
| Median (months) | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 |
| Min, max | 0.5, 8.8 | 1.0, 1.0 | 0.5, 8.8 | 0.9, 2.0 | 0.5, 8.8 |
| Duration of response (PR or better)^e, n | 94 | 2 | 48 | 44 | 94 |
| Median (months) | 10.6 | 15.8 | 8.5 | 11.3 | 10.6 |
| 95% CI | 8.0, 11.4 | 2.8, 28.8 | 5.4, 11.0 | 10.3, 17.0 | 8.0, 11.4 |

CAR = chimeric antigen receptor; CI = confidence interval; CR = complete response; MRD = minimal residual disease; NE = not estimable; PR = partial response; sCR = stringent complete response; VGPR = very good partial response.

^a All patients who underwent leukapheresis.

^b The 150 x 10⁶ CAR-positive T cell dose is not part of the approved dose range.

^c For “Total (Treated population” and “Enrolled population”): Wald CI; for individual target dose levels: Clopper-Pearson exact CI.

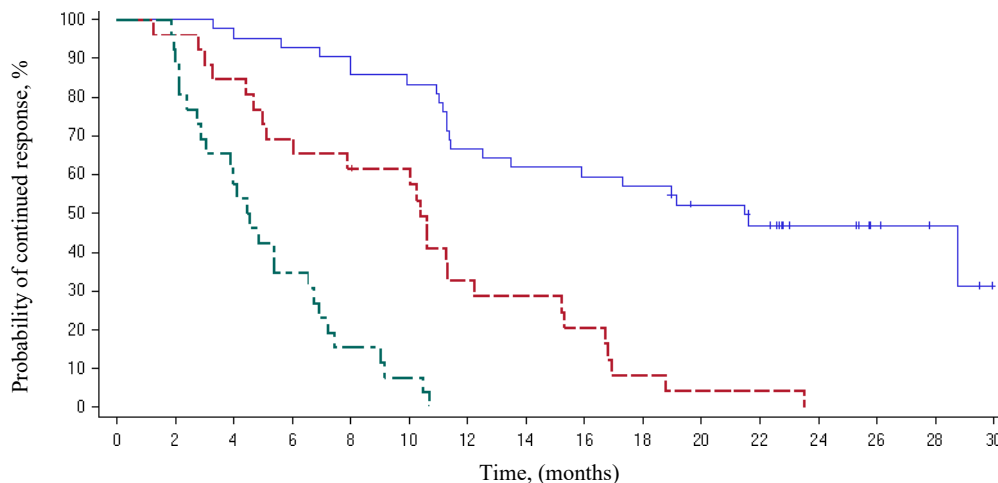
^d Based on a threshold of 10⁻⁵ using a next-generation sequencing assay. 95% CI for percentage of MRD negativity use Clopper-Pearson exact CI for individual target dose levels as well as for Treated population.

^e Median and 95% CI are based on the Kaplan-Meier approach.

Note: The target dose is 450 x 10⁶ CAR-positive T cells within a range of 150 to 540 × 10⁶ CAR-positive T cells. The 150 x 10⁶ CAR-positive T cell dose is not part of the approved dose range.

The Kaplan-Meier curve of duration of response by best overall response is shown in Figure 13.

Figure 13. Kaplan-Meier curve of duration of response based on independent response committee review according to IMWG criteria – by best overall response (Abecma-treated population- KarMMA study)



| | | | | | | | | | | | | | | | | |
|--------------|----|----|----|----|----|----|----|----|----|----|----|----|----|---|---|---|
| CR or better | 42 | 42 | 40 | 39 | 36 | 35 | 28 | 26 | 25 | 24 | 20 | 17 | 10 | 6 | 3 | 0 |
| VGPR | 26 | 25 | 22 | 18 | 16 | 15 | 8 | 7 | 5 | 2 | 1 | 1 | 0 | 0 | 0 | 0 |
| PR | 26 | 23 | 15 | 9 | 4 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |

— CR or better: Subjects: 42; Events: 23; Median: 21.45 (95% CI: 12.52, NE)
- - - VGPR: Subjects: 26; Events: 25; Median: 10.38 (95% CI: 5.09, 12.22)
- - - PR: Subjects: 26; Events: 26; Median: 4.50 (95% CI: 2.86, 6.54)

CI= confidence interval; IMWG = International Myeloma Working Group; NE = not estimable. Two patients with 150×10^6 CAR-positive T cell dose, which is not part of the approved dose range, are included in Figure 13.

Special populations

Elderly

In the clinical trials of Abecma, 48-163 (34.3-39.9%) patients in the KarMMA study were 65 years of age or older and 5-17 (3.6-4.2%) were 75 years of age or older (see Table 5). No clinically important differences in the safety or effectiveness of Abecma were observed between these patients and patients younger than 65 years of age.

5.2 Pharmacokinetic properties

Following Abecma infusion, the CAR-positive T cells proliferate and undergo rapid multi-log expansion followed by a bi-exponential decline. The median time of maximal expansion in peripheral blood (T_{max}) occurred 11 days after infusion.

Abecma can persist in peripheral blood for up to 1 year post-infusion.

Abecma transgene levels were positively associated with objective tumour response (partial response or better). In patients who received Abecma in the KarMMA-3 study, the median C_{max} levels in responders (N = 180) were approximately 5.4-fold higher compared to the corresponding levels in non-responders (N = 40). Median $AUC_{0-28days}$ in responders (N = 180) was approximately 5.5-fold higher than non-responders (N = 38). In patients who received Abecma in the KarMMA study, the median C_{max} levels in responders (N = 93) were approximately 4.5-fold higher.

compared to the corresponding levels in non-responders (N = 34). Median AUC_{0-28days} in responding patients (N = 93) was approximately 5.5-fold higher than non-responders (N = 32).

Special populations

Renal and hepatic impairment

Hepatic and renal impairment studies of Abecma were not conducted.

Effects of age, weight, gender or race

Age (range: ~~33-30~~ to ~~78-81~~ years) had no impact on Abecma expansion parameters. The pharmacokinetics of Abecma in patients less than 18 years of age have not been evaluated.

Patients with lower body weight had higher cellular expansion. Due to high variability in pharmacokinetic cellular expansion, the overall effect of weight on the expansion parameters of Abecma is considered not to be clinically relevant.

Gender had no impact on Abecma expansion parameters.

Race and ethnicity had no significant impact on Abecma expansion parameters.

5.3 Preclinical safety data

Abecma comprises engineered human T cells, therefore there are no representative *in vitro* assays, *ex vivo* models, or *in vivo* models that can accurately address the toxicological characteristics of the human product. Hence, traditional toxicology studies used for drug development were not performed. Genotoxicity assays and carcinogenicity studies were not conducted.

In vitro expansion studies from healthy donors and patients showed no evidence for transformation and/or immortalisation and no preferential integration near genes of concern in Abecma T cells.

Given the nature of the product, non-clinical studies on fertility, reproduction and development were not conducted.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

CryoStor CS10 freeze media (~~containing~~ contains 10% DMSO)

Sodium chloride

Sodium gluconate

Sodium acetate trihydrate

Potassium chloride

Magnesium chloride

Water for injections

6.2 Incompatibilities

In the absence of compatibility studies, this medicinal product must not be mixed with other medicinal products.

6.3 Shelf life

The expiry date of the product is indicated on the packaging materials.

Abecma is stable for 12 months when stored in the vapour phase of liquid nitrogen ($\leq -130^{\circ}\text{C}$).

Each bag must be infused within 1 hour from start of thaw. After thawing, the volume of the product intended for infusion should be kept at room temperature (20°C – 25°C).

6.4 Special precautions for storage

~~Store and transport frozen~~ Abecma **must be stored** in the vapour phase of liquid nitrogen ($\leq -130^{\circ}\text{C}$) and must remain frozen until the patient is ready for treatment to ensure viable ~~live autologous~~ cells are available for patient administration. ~~Product must~~ **Thawed medicinal product should not NOT** be refrozen ~~following thaw~~.

For storage conditions after thawing of the medicinal product, see section 6.3.

6.5 Nature and contents of container

Ethylene vinyl acetate cryopreservation bag(s) with sealed addition tube containing 10-30 mL (50 mL bag), 30-70 mL (250 mL bag) or 55-100 mL (500 mL bag) of cell dispersion.

Each cryopreservation bag is individually packed in a metal cassette.

One individual treatment dose is comprised of one or more infusion bags of the same size and fill volume.

6.6 Special precautions for disposal and other handling

Precautions to be taken before handling or administering the medicinal product

Abecma ~~should~~ **must** be transported within the ~~treatment centre~~ facility in closed, break-proof, leak-proof containers.

This medicinal product contains human blood cells. Healthcare professionals handling Abecma ~~should~~ **must** take appropriate precautions (wearing gloves and glasses) to avoid potential transmission of infectious diseases.

Preparation prior to administration

Prior to Abecma infusion, it must be confirmed that the patient's identity matches the patient identifiers on the Abecma cassette(s), the infusion bag(s) and the release for infusion certificate (RfIC). The Abecma infusion bag must not be removed from the cassette if the information on the patient-specific label does not match the intended patient. The company must be contacted immediately if there are any discrepancies between the labels and the patient identifiers.

If more than one infusion bag has been received for treatment, thaw each infusion bag one at a time. The timing of thaw of Abecma and infusion should be coordinated. The infusion start time should be confirmed in advance and adjusted for thaw so that Abecma is available for infusion when the patient is ready.

Thawing

- Remove the Abecma infusion bag from the cassette and inspect the infusion bag for any breaches of container integrity such as breaks or cracks before thawing. If the infusion bag appears to have been damaged or to be leaking, it should not be infused and should be disposed of according to local guidelines on handling of waste of human-derived material.
- Place the infusion bag inside a second sterile bag.
- Thaw Abecma at approximately 37°C using an approved thaw device or water bath until there is no visible ice in the infusion bag. Gently mix the contents of the bag to disperse clumps of

cellular material. If visible cell clumps remain, continue to gently mix the contents of the bag. Small clumps of cellular material should disperse with gentle manual mixing. Do not wash, spin down and/or resuspend Abecma in new media prior to infusion.

Abecma administrationAdministration

- Prime the tubing of the infusion set with sodium chloride 9 mg/mL (0.9%) solution for injection prior to infusion.
- Infuse Abecma within 1 hour from start of thaw as quickly as tolerated by gravity flow.
- After the entire content of the infusion bag is infused, rinse the tubing with sodium chloride 9 mg/mL (0.9%) solution for injection at the same infusion rate to ensure all product is delivered.
- Follow the same procedure for all subsequent infusion bags for the identified patient.

Measures to take in case of Accidental exposure

In case of accidental exposure local guidelines on handling of human-derived material should must be followed. Work surfaces and materials which have potentially been in contact with Abecma must be decontaminated with appropriate disinfectant.

Precautions to be taken for disposal of the medicinal product

Unused medicinal product and all material that has been in contact with Abecma (solid and liquid waste) ~~should~~ **must** be handled and disposed of as potentially infectious waste in accordance with local guidelines on handling of human-derived material.

~~Accidental exposure~~

~~In case of accidental exposure local guidelines on handling of human-derived material should be followed. Work surfaces and materials which have potentially been in contact with Abecma must be decontaminated with appropriate disinfectant.~~

7. REGISTRATION HOLDER

Bristol Myers Squibb (Israel) Ltd.
18 Aharon Bart
P.O Box 3361,
Kiryat Arie
Petach Tikva 4951448

8. MANUFACTURER

Celgene Corporation
Building S12 556 Morris Ave
Summit, NJ, 07901
USA

9. REGISTRATION NUMBER

171-53-37039-00

ApprovedRevised on in 020708/20232024

Abecma_NPI_Aug 2024

עלון לצרכן לפי תקנות הרוקחים (תכשירים) התשמ"ו - 1986

התרופה משווקת על פי מרשם רופא בלבד

אבקמה

תרחיף לעירוני תוך ורידי המכיל $10^6 \times 500 - 260$ תאים

חומר פעיל

אידקבטג'ן ויקלאוסל (idecabtagene vicleucel-) התרחיף מכיל $10^6 \times 500 - 260$ תאי T עצמיים שעברו שינוי גנטי לביטוי קולטן לאנטיגן כימרי נגד BCMA (תאי T חיוניים חיוביים ל-CAR)

חומרים בלתי פעילים ואלרגניים: ראה פרק 2 סעיף "מידע חשוב על חלק מהמרכיבים של התרופה" ופרק 6 "מידע נוסף".

קרא בעיון את העלון עד סופו בטרם תשתמש בתרופה. עלון זה מכיל מידע תמציתי על התרופה. אם יש לך שאלות נוספות, פנה אל הרופא או אל הרוקח. תרופה זו נרשמה עבורך. אל תעביר אותה לאחרים. היא עלולה להזיק להם אפילו אם נראה לך כי מצבם הרפואי דומה.

בנוסף לעלון זה, לתכשיר אבקמה קיים כרטיס **מידע בטיחותי**-למטופל. כרטיס זה מכיל מידע בטיחותי חשוב, שעליך לדעת לפני התחלת הטיפול ובמהלך הטיפול באבקמה ולפעול על פיו. יש לעיין בכרטיס **מידע בטיחותי**-למטופל ובעלון לצרכן בטרם תחילת השימוש בתכשיר. יש לשמור את הכרטיס לעיון נוסף במידת הצורך.

1. למה מיועדת התרופה?

אבקמה מיועדת לטיפול במבוגרים עם מיאלומה נפוצה חוזרת ועמידה, אשר קיבלו לפחות שלושה טיפולים קודמים, כולל טיפול בתכשיר אימונומודולטורי, במעכב פרוטאזום ובנוגדן אנטי CD38 והראו התקדמות של המחלה בטיפול האחרון.

קבוצה תרפויטית: **תרופות אנטי סרטניות אחרות.**

אבקמה היא סוג של תרופה המכונה "טיפול בתאים שעברו שינוי גנטי". החומר הפעיל בתרופה, אידקבטג'ן ויקלאוסל, מיוצר מתאי הדם הלבנים העצמיים (תאים שלך), המכונים תאי T.

כיצד פועלת אבקמה

תאי הדם הלבנים הנלקחים מהדם שלך עוברים שינוי גנטי על מנת שיוכלו להתמקד בתאי המיאלומה בגוף שלך. כאשר אבקמה תינתן בעירוני לדם, תאי הדם הלבנים שעברו שינוי ייהרגו את תאי המיאלומה.

2. לפני השימוש בתרופה

אין להשתמש בתרופה אם:

- אתה רגיש (אלרגי) לחומר הפעיל או לכל אחד מהמרכיבים הנוספים אשר מכילה התרופה (ראה פרק 6). אם אתה חושב שאתה עלול להיות אלרגי, פנה לרופא לייעוץ.
- אם אתה אלרגי לאחד המרכיבים בתרופות אותן תקבל במסגרת כימותרפיה להפחתת לימפוציטים, אשר משמשת להכנת הגוף לטיפול באבקמה.

אזהרות מיוחדות הנוגעות לשימוש בתרופה

לפני הטיפול באבקמה, ספר לרופא אם:

- אתה סובל מבעיות כלשהן בריאות או בלב.
- אתה סובל מלחץ דם נמוך.

- עברת השתלת תאי גזע במהלך 4 החודשים האחרונים.
- אתה סובל מסימנים או מתסמינים של מחלת השתל נגד המאכסן. הדבר מתרחש כאשר התאים המושתלים תוקפים את הגוף וגורמים לתסמינים כגון פריחה, בחילות, הקאות, שלשול וצואה דמית.
- **אתה סובל מזיהום או תהליך דלקתי פעיל (כולל דלקת ברקמת הריאה, דלקת שריר הלב או דלקת בכבד).** הזיהום יטופל בטרם תקבל אבקמה.
- **אתה סובל מבעיות במערכת העצבים המרכזית, או מתפקוד לקוי של הכליות או הכבד.**
- אתה מבחין בהחמרת תסמיני הסרטן. תסמינים אלה במיאלומה עשויים לכלול חום, תחושת חולשה, כאב בעצמות, ירידה במשקל שאינה מוסברת.
- סבלת מזיהום בציטומגלו-וירוס (CMV), בדלקת כבד נגיפית (הפטיטיס) B או C או בנגיף הכשל החיסוני האנושי (HIV).
- קיבלת חיסון במהלך 6 השבועות הקודמים או שאתה מתכנן לקבל חיסון במהלך החודשים הקרובים.

אם אתה חושב שאחד מהמקרים מעלה רלוונטי לגביך (או שאינך בטוח), דבר עם הרופא לפני **מתן שאתה מקבל אבקמה**

ילדים ומתבגרים

אין לתת אבקמה לילדים ולמתבגרים מתחת לגיל 18 שנים. לא קיים מידע לגבי בטיחות ויעילות השימוש בתרופה בילדים ומתבגרים.

בדיקות ומעקב

בטרם תקבל אבקמה, הרופא:

- יבדוק לך את הריאות, הלב ולחץ הדם.
- יחפש סימני זיהום; כל זיהום יטופל בטרם תקבל אבקמה.
- יבדוק האם הסרטן מחמיר.
- יבדוק אם יש לך זיהום ב-CMV, בדלקת כבד נגיפית (הפטיטיס) B, בדלקת כבד נגיפית (הפטיטיס) C או זיהום ב-HIV.

לאחר קבלת אבקמה

- קיימות תופעות לוואי רציניות עליהן אתה חייב לדווח מייד לרופא או לאחות ואשר עלולות לאלץ אותך לקבל עזרה רפואית מיידית. ראה פרק 4 תחת "תופעות לוואי רציניות".
- הרופא יבדוק את ספירות הדם שלך באופן סדיר מאחר שמספר תאי הדם עלול לרדת.
- עליך להישאר קרוב למרכז הרפואי בו קיבלת אבקמה במשך לפחות 4 שבועות. ראה פרקים 3 ו-4.
- אין לתרום דם, איברים, רקמות או תאים להשתלה.
- **מטופלים אשר טופלו באבקמה עלולים לפתח סוגים חדשים של סרטן. תצטרך להיות במעקב אצל רופא לכל החיים.**

אינטראקציות/תגובות בין תרופתיות

אם אתה לוקח או אם לקחת לאחרונה תרופות אחרות כולל תרופות ללא מרשם ותוספי תזונה, ספר על כך לרופא או לרוקח.

תרופות המשפיעות על מערכת החיסון

טרם קבלת אבקמה, דווח לרופא או לאחות אם אתה נוטל תרופות כלשהן המחלישות את מערכת החיסון, כגון קורטיקוסטרואידים. הסיבה לכך היא שתרופות אלו עלולות לשבש את ההשפעה של אבקמה.

ראה פרק 3 למידע על התרופות אותן תקבל טרם קבלת אבקמה.

חיסונים

- אין לקבל חיסונים מסוימים המכונים חיסונים חיים:
 - במהלך 6 השבועות הקודמים לקבלת מחזור טיפול קצר בכימותרפיה (מכונה כימותרפיה להפחתת לימפוציטים) על מנת להכין את הגוף לאבקמה.

- במהלך הטיפול באבקמה.
- לאחר הטיפול, במהלך התאוששות מערכת החיסון.

שוחח עם הרופא אם אתה צריך לקבל חיסונים כלשהם.

היריון והנקה

אם את בהיריון או מניקה, חושבת שאת עשויה להיות בהיריון או מתכננת להרות, פני לרופא ליעוץ טרם קבלת תרופה זו. הסיבה לכך היא שההשפעות של אבקמה על נשים הרות או מניקות אינן ידועות והיא עלולה להזיק לעובר או לתינוק יונק.

- אם את בהיריון או חושבת שאת עשויה להיות בהיריון לאחר הטיפול באבקמה, שוחחי מייד עם הרופא.
- עלייך לעבור בדיקת היריון טרם תחילת הטיפול. **תקבלי** טיפול באבקמה **ינתן** רק אם התוצאות מראות שאינך בהיריון.

שוחחי על היריון עם הרופא אם קיבלת אבקמה.

נהיגה ושימוש במכונות

אין לנהוג, להשתמש במכונות או להשתתף בפעילויות המחייבות אותך להיות ערני במשך לפחות 8 שבועות לאחר הטיפול או עד אשר הרופא יודיע לך שהתאוששת לחלוטין. אבקמה עלולה לגרום לך לתחושת ישנוניות, לבלבול או לפרכוסים.

מידע חשוב על חלק מהמרכיבים של התרופה

אבקמה מכילה נתרן, אשלגן ודימתיל סולפוקסיד (DMSO)

תרופה זו מכילה עד 752 מ"ג נתרן (המרכיב העיקרי של מלח ביסול/שולחן) למנה. כמות זו היא שוות ערך ל- 37.6% מצריכת הנתרן היומית המקסימלית המומלצת עבור מבוגר.

תרופה זו מכילה עד 274 מ"ג אשלגן למנה. על מטופלים עם תפקוד כליות ירוד או מטופלים הצורכים דיאטה בעלת תכולת אשלגן מבוקרת להתחשב בכך.

אם לא נחשפת לדימתיל סולפוקסיד בעבר, עליך להיות במעקב צמוד במהלך הדקות הראשונות של מתן העירוי.

3. כיצד תשתמש בתרופה?

יש להשתמש בתכשיר תמיד בהתאם להוראות הרופא. עליך לבדוק עם הרופא או הרוקח אם אינך בטוח בנוגע למינון ואופן הטיפול בתכשיר. המינון ואופן הטיפול יקבעו על ידי הרופא בלבד.

איסוף דם לייצור אבקמה מתאי הדם הלבנים שלך

- הרופא ייקח ממך דם באמצעות צינורית (צנתר) המוחדרת לווריד. חלק מתאי הדם הלבנים יופרדו מן הדם ויתרת הדם תוחזר לגופך. התהליך מכונה "לויקופריזיס" והוא עשוי להימשך 3 עד 6 שעות. ייתכן כי יהיה צורך לחזור על תהליך זה.
- לאחר מכן תאי הדם הלבנים שלך יוקפאו וישלחו לייצור אבקמה.

תרופות אחרות אותן תקבל לפני אבקמה

- מספר ימים טרם קבלת אבקמה, תקבל מחזור טיפול קצר בכימותרפיה. הדבר מיועד לסלק את תאי הדם הלבנים הקיימים שלך.
- זמן קצר טרם קבלת אבקמה, תקבל פרצטמול ותרופה אנטי-היסטמינית. הדבר מיועד להפחית את הסיכון לתגובות לעירוי ולחום.

כיצד ניתנת אבקמה

- הרופא יוודא כי אבקמה הוכנה מהדם שלך באמצעות וידוא כי פרטי זיהוי המטופל על תוויות התרופה תואמים לפרטים שלך.
- אבקמה ניתנת כעירוי בטפטוף דרך צינורית המוחדרת לווריד.

לאחר מתן אבקמה

- עליך להישאר קרוב למרכז הרפואי בו קיבלת אבקמה במשך לפחות 4 שבועות.

- ייתכן כי תעבור מעקב יומיומי במרכז הרפואי במשך לפחות 10 ימים על מנת לבדוק האם הטיפול עלילעובד וכדי לעזור לך אם תסבול מתופעות לוואי כלשהן. ראה פרקים 2 ו-4.
- אין לתרום דם, איברים, רקמות או תאים להשתלה.

אם החמצת ביקור

התקשר לרופא או למרכז הרפואי בהקדם האפשרי על מנת לקבוע ביקור אחר.

יש להתמיד בטיפול כפי שהומלץ על ידי הרופא.

אין ליטול תרופות בחושך! בדוק התווית והמנה בכל פעם שאתה לוקח תרופה. הרכב משקפיים אם אתה זקוק להם.
אם יש לך שאלות נוספות בנוגע לשימוש בתרופה, היוועץ ברופא או ברוקח.

4. תופעות לוואי

כמו בכל תרופה, השימוש באבקמה עלול לגרום לתופעות לוואי בחלק מהמשתמשים. אל תיבהל למקרא רשימת תופעות הלוואי. ייתכן שלא תסבול מאף אחת מהן.

תופעות לוואי רציניות

דווח מייד לרופא אם תפתח אחת מתופעות הלוואי הבאות לאחר קבלת אבקמה. הן בדרך כלל מופיעות במהלך 8 השבועות הראשונים לאחר העירוי, אך עלולות להתפתח גם במועד מאוחר יותר:
תופעות לוואי שכיחות מאוד - (very common) תופעות שמופיעות ביותר ממשמש אחד מתוך עשרה:

- חום, צמרמורת, קושי בנשימה, סחרחורת או תחושת סחרור, בחילות, כאב ראש, קצב לב מהיר, לחץ דם נמוך או תשישות – אלה עלולים להיות תסמינים של תסמונת שחרור ציטוקינים או CRS, מצב רציני שעלול להיות קטלני.
- **בלבול, קושי בזיכרון, קושי בדיבור או דיבור מואט, קושי בהבנת דיבור, איבוד שיווי משקל או קואורדינציה, חוסר התמצאות, ירידה בערנות (ירידה ברמת ההכרה) או ישנוניות מופרזת, איבוד הכרה, מצב בלבולי חריף (דליריום), פרכוסים, רעד או חולשה עם אובדן תנועה בצד אחד של הגוף.**
- סימני זיהום כלשהם, העלולים לכלול חום, צמרמורת או רעד, שיעול, קוצר נשימה, נשימה מהירה ודופק מהיר.
- תחושת עייפות קיצונית או חולשה או קוצר נשימה – אשר עלולים להיות סימנים לרמות נמוכות של תאי דם אדומים (אנמיה).
- דימום או הופעת חבורות בקלות רבה יותר ללא סיבה, לרבות דימומים מהאף או דימום מהפה או מהמעיים, אשר עלולים להיות סימן של רמות נמוכות של טסיות בדם שלך.

תופעות לוואי שכיחות (common) תופעות שמופיעות ב 10-1 משתמשים מתוך 100:

- **רעידות, חולשה עם אובדן תנועה של צד אחד של הגוף, רעד, תנועות איטיות או נוקשות - אלה עלולים להיות תסמינים של פרקינסוניזם**

תופעות לוואי לא שכיחות (uncommon) תופעות שמופיעות ב 10-1 משתמשים מתוך 1000:

- **בלבול, קושי בזיכרון, קושי בדיבור או האטה בדיבור, קושי בהבנת דיבור, איבוד שיווי משקל או קואורדינציה, חוסר התמצאות, ירידה בערנות (ירידה ברמת ההכרה) או ישנוניות מופרזת, איבוד הכרה, מצב בלבולי חריף (דליריום), פרכוסים – אלה עלולים להיות תסמינים של מצב רפואי הנקרא רעילות נירולוגית מסוג (immune effector cell-associated neurotoxicity syndrome (ICANS**

דווח מייד לרופא אם אתה סובל מאחת מתופעות הלוואי שלעיל, מאחר שיתכן כי אתה זקוק לטיפול רפואי דחוף.

תופעות לוואי נוספות

תופעות לוואי שכיחות מאוד - (very common) תופעות שמופיעות ביותר ממשתמש אחד מתוך עשרה:

- חוסר אנרגיה
- לחץ דם גבוה
- תיאבון ירוד
- עצירות
- נפיחות בקרסוליים, בזרועות, ברגליים ובפנים
- כאבי מפרקים
- קשיי שינה
- ספירה נמוכה של תאי דם לבנים (נויטרופילים, לויקוציטים ולימפוציטים), אשר עלולה להגביר את הסיכון לזיהום
- זיהומים, לרבות דלקת ריאות או זיהומים בדרכי הנשימה, בפה, בעור, בדרכי השתן או בדם, אשר עשויים להיות חיידקיים, וירליים או פטרייתיים
- תוצאות בדיקות מעבדה המראות רמות נמוכות של נוגדנים, המכונים אימונוגלובולינים (היפוגמגלובולינמיה), שחשובים במלחמה בזיהומים
- תוצאות בדיקות מעבדה המראות רמות מופחתות של סידן, נתרן, מגנזיום, אשלגן, פוספט או אלבומין, אשר עלולות לגרום לתשישות, לחולשת שרירים או לעוויתות שרירים, או לקצב לב בלתי סדיר
- תוצאות בדיקות מעבדה המראות רמות מוגברות של אנזימי כבד (ממצאים חריגים בבדיקת תפקודי כבד) או רמה גבוהה יותר של חלבון (C-reactive protein) בדם העשויה להעיד על דלקת.

תופעות לוואי שכיחות (common) תופעות שמופיעות ב 10-1 משתמשים מתוך 100:

- דלקת חמורה בשל שפעול מערכת החיסון, אשר עלולה להוביל לנזק רציני בגוף
- קשיי שינה
- כאבי שרירים
- תנועות גוף חריגות או חוסר קואורדינציה
- קצב לב משתנה או בלתי סדיר
- נזל בריאות
- רמת חמצן נמוכה בדם, אשר עלולה לגרום לקוצר נשימה, לבלבול או לנמנום.

אם הופיעה תופעת לוואי, אם אחת מתופעות הלוואי מחמירה או כאשר אתה סובל מתופעת לוואי שלא צוינה בעלון, עליך להתייעץ עם הרופא.

ניתן לדווח על תופעות לוואי למשרד הבריאות באמצעות לחיצה על הקישור "דיווח על תופעות לוואי עקב טיפול תרופתי" שנמצא בדף הבית של אתר משרד הבריאות (www.health.gov.il) המפנה לטופס המקוון לדיווח על תופעות לוואי, או ע"י כניסה לקישור: <https://sideeffects.health.gov.il/>

5. כיצד לאחסן את התרופה?

מנע הרעלה! תרופה זו וכל תרופה אחרת יש לשמור במקום סגור מחוץ להישג ידם וטווח ראייתם של ילדים ו/או תינוקות ועל ידי כך תמנע הרעלה. אין לגרום להקאה ללא הוראה מפורשת מרופא.

המידע שלהלן מיועד לרופאים בלבד.

אין להשתמש בתרופה זו לאחר תאריך התפוגה המצוין על תווית הקסטה ושקית העירוי אחרי "EXP".

תנאי אחסון: יש לאחסן ~~הלשנע~~ במצב קפוא בפאזת האדים של חנקן נוזלי ($\geq -130^{\circ}\text{C}$). אין להפשיר את התרופה עד אשר תהיה מוכן להשתמש בה. אין להקפיא מחדש.

אין להשתמש בתרופה זו אם שקית העירוי פגומה או דולפת.

~~תרופה זו מכילה תאי דם אנושיים שעברו שינוי גנטי. יש לפעול על פי ההנחיות לטיפול בפסולת המכילה חומר המופק מבני אדם עבור תרופה שנתרה ללא שימוש או חומר פסולת.~~

6. מידע נוסף

כל שקית עירוי של אבקמה מכילה תרחיף תאים של idecabtagene vicleucel בריכוז תלוי-אצווה; התרחיף מכיל תאי T עצמיים שעברו שינוי גנטי לביטוי קולטן לאנטיגן כימרי נגד BCMA (תאי T חיוניים חיוביים ל-CAR). שקית עירוי אחת או מספר שקיות עירוי מכילות בסה"כ 260×10^6 עד 500×10^6 תאי T חיים חיוביים ל-CAR.

• נוסף על המרכיב הפעיל, התרופה מכילה גם:

Cryostor CS10 freeze media (containing 40% DMSO), sodium chloride, sodium gluconate, sodium acetate trihydrate, potassium chloride, magnesium chloride, water for injection

תרופה זו מכילה תאי דם אנושיים שעברו שינוי גנטי.

• כיצד נראית התרופה ומה תוכן האריזה

אבקמה היא תרחיף חסר צבע של תאים לעירוי, המסופק בשקית עירוי אחת או במספר שקיות עירוי הארוזות בנפרד בקסטות מתכתיות. כל שקית מכילה 10 מ"ל עד 100 מ"ל תרחיף תאים.

• **בעל הרישום וכתובתו:** בריסטול-מאירס סקוויב (ישראל) בע"מ, רח' אהרון ברט 18, ת.ד. 3361, קריית אריה, פתח תקווה 4951448.

• שם היצרן וכתובתו:

Celgene Corporation
Building S12, 556 Morris Avenue Summit, New Jersey 07901, USA

אושר-נערך באוגוסט 2024-דצמבר 2022

מספר רישום התרופה בפנקס התרופות הממלכתי במשרד הבריאות: ~~XX-171-XXX-53-37039-XX~~

לשם הפשטות ולהקלת הקריאה עלון זה נוסח בלשון זכר. על אף זאת, התרופה מיועדת לבני שני המינים.

The following information is intended for healthcare professionals only:

It is important that you read the entire content of this procedure prior to administering Abecma.

Precautions to be taken before handling or administering the medicinal product

- Abecma ~~should~~-must be transported within the ~~treatment centre~~-facility in closed, break-proof, leak-proof containers.
- This medicinal product contains human blood cells. Healthcare professionals handling Abecma ~~should~~-must take appropriate precautions (wearing gloves and glasses) to avoid potential transmission of infectious diseases.

Preparation prior to administration

- Prior to Abecma infusion, it must be confirmed that the patient's identity matches the patient identifiers on the Abecma cassette(s), the infusion bag(s) and the release for infusion certificate (RfIC).
- The Abecma infusion bag must not be removed from the cassette if the information on the patient-specific label does not match the intended patient. The company must be contacted immediately if there are any discrepancies between the labels and the patient identifiers.
- If more than one infusion bag has been received for treatment, thaw each infusion bag one at a time. The timing of thaw of Abecma and infusion should be coordinated. The infusion start time should be confirmed in advance and adjusted for thaw so that Abecma is available for infusion when the patient is ready.

Thawing

- Remove the Abecma infusion bag from the cassette and inspect the infusion bag for any breaches of container integrity such as breaks or cracks before thawing. If the infusion bag appears to have been damaged or to be leaking, it should not be infused and should be disposed of according to local guidelines on handling of waste of human-derived material.
- Place the infusion bag inside a second sterile bag.
- Thaw Abecma at approximately 37°C using an approved thaw device or water bath until there is no visible ice in the infusion bag. Gently mix the contents of the bag to disperse clumps of cellular material. If visible cell clumps remain, continue to gently mix the contents of the bag. Small clumps of cellular material should disperse with gentle manual mixing. Do not wash, spin down and/or resuspend Abecma in new media prior to infusion.

Abecma administration Administration

- Do NOT use a leukodepleting filter.
- Intravenous infusion of Abecma should only be administered by a healthcare professional experienced with immunosuppressed patients and prepared to manage anaphylaxis.
- Ensure that tocilizumab and emergency equipment are available prior to infusion and during the recovery period. In the exceptional case where tocilizumab is not available, ensure that suitable alternative measures to treat CRS instead of tocilizumab are available on-site.
- Central venous access may be utilised for the infusion of Abecma and is encouraged in patients with poor peripheral access.
- Before administration, it must be confirmed that ~~Confirm~~ the patient's identity matches the unique patient identifiers_—on information on the Abecma infusion bag and accompanying documentation.-. The total number of infusion bags to be administered must also be confirmed with the patient specific information on the release for infusion certificate (RfIC).
- Prime the tubing of the infusion set with sodium chloride 9 mg/mL (0.9%) solution for injection prior to infusion.
- Infuse Abecma within 1 hour from start of thaw as quickly as tolerated by gravity flow.
- After the entire content of the infusion bag is infused, rinse the tubing with sodium chloride 9 mg/mL (0.9%) solution for injection at the same infusion rate to ensure all product is delivered.

- Follow the same procedure for all subsequent infusion bags for the identified patient.

Disposal of Abecma

Measures to take in case of accidental exposure

- In case of accidental exposure, local guidelines on handling of human-derived material must be followed. Work surfaces and materials which have potentially been in contact with Abecma must be decontaminated with appropriate disinfectant.

Precautions to be taken for the disposal of the medicinal product

Unused medicinal product and all material that has been in contact with Abecma (solid and liquid waste) ~~should~~must be handled and disposed of as potentially infectious waste in accordance with local guidelines on handling of human-derived material.

Accidental exposure

~~In case of accidental exposure local guidelines on handling of human-derived material should be followed. Work surfaces and materials which have potentially been in contact with Abecma must be decontaminated with appropriate disinfectant.~~