

Deploy DANYELZA

Battle bone and bone marrow metastases

At incomplete response* to induction or relapse therapy, recruit the only FDA-approved immunotherapy to treat disease in bone and/or bone marrow.¹

*Incomplete response is defined as partial response (PR), minor response (MR), or stable disease (SD) to prior therapy.

DANYELZA[®]
 (naxitamab-gqqgk)
 40mg/10mL Injection

INDICATION

DANYELZA is indicated, in combination with granulocyte-macrophage colony-stimulating factor (GM-CSF), for the treatment of pediatric patients 1 year of age and older and adult patients with relapsed or refractory high-risk neuroblastoma in the bone or bone marrow who have demonstrated a partial response, minor response, or stable disease to prior therapy.

This indication is approved under accelerated approval based on overall response rate and duration of response. Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial(s).

IMPORTANT SAFETY INFORMATION

WARNING: SERIOUS INFUSION-RELATED REACTIONS and NEUROTOXICITY

Serious Infusion-Related Reactions

- DANYELZA can cause serious infusion reactions, including cardiac arrest, anaphylaxis, hypotension, bronchospasm, and stridor. Infusion reactions of any Grade occurred in 94-100% of patients. Severe infusion reactions occurred in 32-68% and serious infusion reactions occurred in 4-18% of patients in DANYELZA clinical studies.
- Premedicate prior to each DANYELZA infusion as recommended and monitor patients for at least 2 hours following completion of each infusion. Reduce the rate, interrupt infusion, or permanently discontinue DANYELZA based on severity.

Neurotoxicity

- DANYELZA can cause severe neurotoxicity, including severe neuropathic pain, transverse myelitis and reversible posterior leukoencephalopathy syndrome (RPLS). Pain of any Grade occurred in 94-100% of patients in DANYELZA clinical studies.
- Premedicate to treat neuropathic pain as recommended. Permanently discontinue DANYELZA based on the adverse reaction and severity.

[CONTINUE READING >](#)



Bone/BM in NB	Incomplete Response	Clinical Studies	Efficacy	MOA	Safety	Dosing & Administration	Y-mAbs Connect	References	Important Safety Information	Summary
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Half of neuroblastoma is classified as high-risk²

Bone and bone marrow are the most common sites of metastatic neuroblastoma in children presenting with metastatic disease³

70% of metastases involve bone marrow

55% of metastases involve cortical bone

2/3 of patients do not achieve a complete metastatic response during induction therapy⁴

2/5 of patients relapse despite intensive multimodal frontline therapy⁵

Most patients with metastatic disease do not have a complete response to induction therapy, or experience relapse^{4,5}

Reducing or eliminating disease in the bone and bone marrow is a goal of high-risk neuroblastoma treatment²

- Assessing metastatic disease in bone and bone marrow requires both MIBG imaging and biopsy²
- MIBG imaging quantifies disease in bone and soft tissue and is used to generate a Curie score (CS), which can have prognostic implications^{2,6}

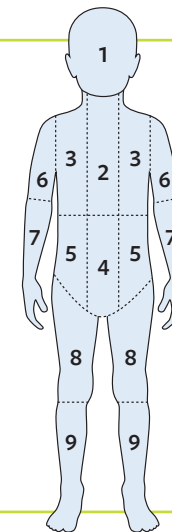
CS is determined by subdividing the body into 10 regions⁶:
9 skeletal and 1 soft tissue

Each region is designated a score of 0-3 points

Maximum collective score = 30

Skeletal regions: 27 points maximum

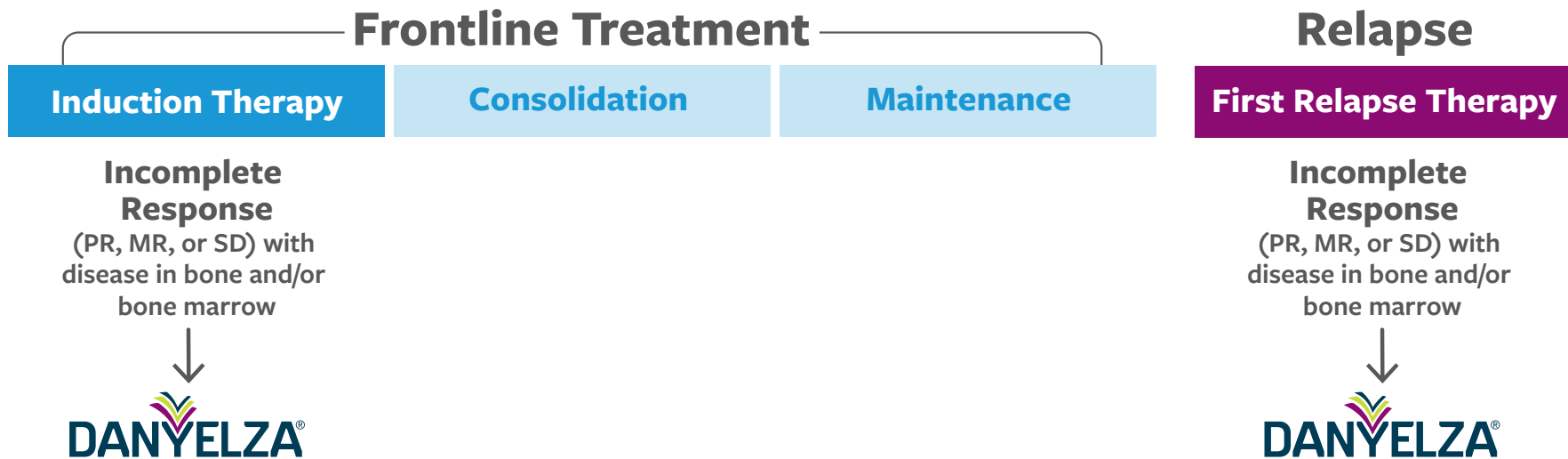
Soft tissue region: 3 points maximum



An absolute CS of 0-2 prior to transplant is shown to be more clinically prognostic than relative reduction in CS⁶⁻⁸



When response is incomplete to induction or relapse therapy, consider the only FDA-approved humanized immunotherapy for patients with high-risk neuroblastoma in the bone and/or bone marrow¹



Incomplete response is defined as partial response (PR), minor response (MR), or stable disease (SD) to prior therapy

IMPORTANT SAFETY INFORMATION

CONTRAINDICATION

DANYELZA is contraindicated in patients with a history of severe hypersensitivity reaction to naxitamab-gqgk. Reactions have included anaphylaxis.

WARNINGS AND PRECAUTIONS

Serious Infusion-Related Reactions

DANYELZA can cause serious infusion reactions requiring urgent intervention including fluid resuscitation, administration of bronchodilators and corticosteroids, intensive care unit admission, infusion rate reduction or interruption of DANYELZA infusion. Infusion-related reactions included hypotension, bronchospasm, hypoxia, and stridor. Serious infusion-related reactions occurred in 4% of patients in Study 201 and in 18% of patients in Study 12-230. Infusion-related reactions of any Grade occurred in 100% of patients in Study 201 and 94% of patients in Study 12-230. Hypotension of any grade occurred in 100% of patients in Study 201 and 89% of patients in Study 12-230. [CONTINUE READING >](#)



Bone/BM in NB	Incomplete Response	Clinical Studies	Efficacy	MOA	Safety	Dosing & Administration	Y-mAbs Connect	References	Important Safety Information	Summary
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DANYELZA with GM-CSF was granted accelerated approval based on two clinical studies¹

STUDY 12-230 (single center)¹

Phase 1/2, open-label, single-arm, single-center trial

- Efficacy analysis included only patients with evaluable disease in bone and/or bone marrow at baseline

N=72; Efficacy analysis (n=38)

STUDY 201 (multicenter)^{1,9}

Phase 2, open-label, single-arm, global trial (US, Canada, Denmark, Germany, Italy, Spain, and Hong Kong)

- Efficacy analysis included only patients with evaluable disease in bone and/or bone marrow at baseline

STUDY 201 Initial Analysis^{1*}

N=25; Efficacy analysis (n=22)

STUDY 201 Pre-specified Interim Analysis⁹

N=74; Efficacy analysis (n=52)

INCLUSION CRITERIA (both studies)^{1,9}

- High-risk neuroblastoma patients ≥12 months of age with bone and/or bone marrow involvement who had incomplete response to induction or relapse therapy
- Evaluable disease in bone and/or bone marrow
- Patients with prior anti-GD2 therapy permitted
- At least one prior systemic therapy to treat disease outside of the bone and/or bone marrow

EXCLUSION CRITERIA (both studies)¹

- Actively progressing disease
- Evaluable neuroblastoma outside of the bone/bone marrow

PRIMARY ENDPOINT^{1,9}

- Overall response rate

SECONDARY ENDPOINTS

- Duration of response
- Complete response
- Safety

*Initial analysis included trial sites in the US and Spain only.

Accelerated approval is based on overall response rate and duration of response.
Continued approval may be contingent upon verification and description of clinical benefit in a confirmatory trial(s).

Treatment with DANYELZA is backed by more than a decade of clinical trial experience and was approved by the FDA in 2020¹⁰

IMPORTANT SAFETY INFORMATION

WARNINGS AND PRECAUTIONS

Serious Infusion-Related Reactions (cont)

In Study 201, 68% of patients experienced Grade 3 or 4 infusion reactions; and in Study 12-230, 32% of patients experienced Grade 3 or 4 infusion reactions. Anaphylaxis occurred in 12% of patients and two patients (8%) permanently discontinued DANYELZA due to anaphylaxis in Study 201. One patient in Study 12-230 (1.4%) experienced a Grade 4 cardiac arrest 1.5 hours following completion of DANYELZA infusion.

In Study 201, infusion reactions generally occurred within 24 hours of completing a DANYELZA infusion, most often within 30 minutes of initiation. Infusion reactions were most frequent during the first infusion of DANYELZA in each cycle. Eighty percent of patients required reduction in infusion rate and 80% of patients had an infusion interrupted for at least one infusion-related reaction. [CONTINUE READING >](#)

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Baseline patient and disease characteristics in DANYELZA with GM-CSF studies^{1,9}

	Study 201		
	STUDY 12-230 ¹ Efficacy Analysis (n=38)	Initial Analysis ¹ Efficacy Analysis (n=22)	Pre-specified Interim Analysis ⁹ Efficacy Analysis (n=52)
DISEASE TYPE			
Refractory (incomplete response to induction)	45% (n=17)	64% (n=14)	50% (n=26)
Relapsed	55% (n=21)	36% (n=8)	50% (n=26)
Median age (range)	5 years (2 to 23 years)	5 years (3 to 10 years)	6 years (2 to 18 years)
MYCN amplification	16%	14%	14%
INSS Stage 4	95%	86%	89%
DISEASE SITES			
Bone marrow only	11%	9%	4%
Bone only	50%	59%	56%
Both	39%	32%	40%
PRIOR TREATMENTS			
Surgery	100%	91%	89%
Chemotherapy	100%	95%	100%
Radiation	47%	36%	40%
ASCT	42%	18%	27%
Anti-GD2 antibody treatment	58%	18%	25%

ASCT=autologous stem cell transplant; INSS=International Neuroblastoma Staging System.

IMPORTANT SAFETY INFORMATION

WARNINGS AND PRECAUTIONS

Serious Infusion-Related Reactions (cont)

Caution is advised in patients with pre-existing cardiac disease, as this may exacerbate the risk of severe hypotension.

Premedicate with an antihistamine, acetaminophen, an H2 antagonist and corticosteroid as recommended. Monitor patients closely for signs and symptoms of infusion reactions during and for at least 2 hours following completion of each DANYELZA infusion in a setting where cardiopulmonary resuscitation medication and equipment are available.

Reduce the rate, interrupt infusion, or permanently discontinue DANYELZA based on severity and institute appropriate medical management as needed.

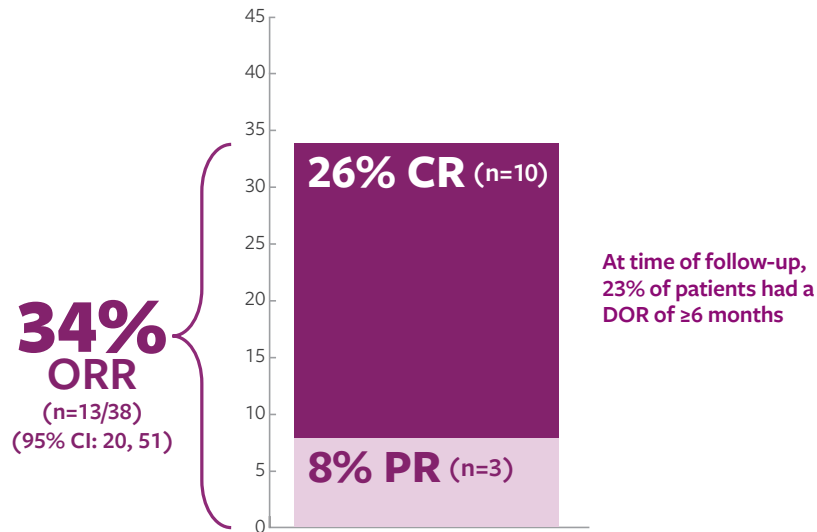
Neurotoxicity

DANYELZA can cause severe neurotoxicity, including severe neuropathic pain, transverse myelitis, and reversible posterior leukoencephalopathy syndrome. [CONTINUE READING >](#)

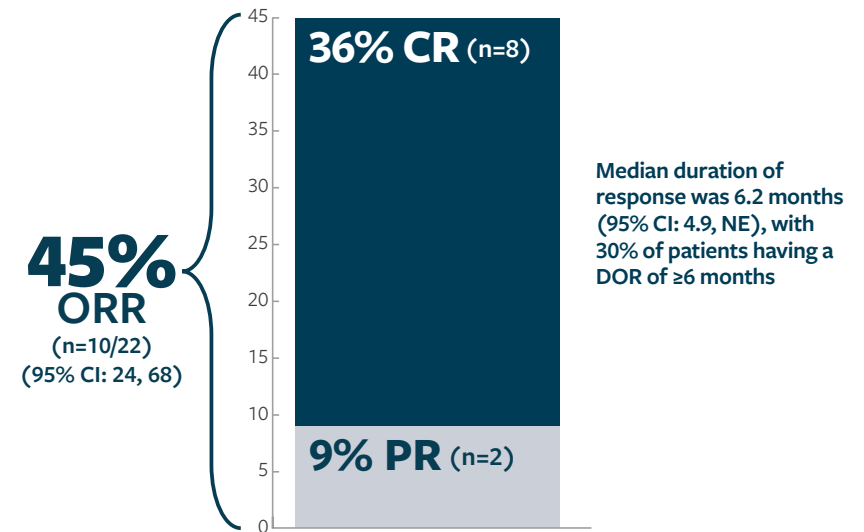
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In both registrational studies, more than 1/3 of patients responded and more than 1/4 achieved *complete* response with DANYELZA with GM-CSF¹

STUDY 12-230 (n=38)



STUDY 201 Initial Analysis (n=22)



ORR was defined as a CR or PR according to the revised INRC (2017) *and confirmed by at least 1 subsequent assessment*

Effectiveness of DANYELZA with GM-CSF was evaluated by independent pathology and imaging review. Responses were observed in the bone, bone marrow, or both bone and bone marrow.¹

CI=confidence interval; CR=complete response; DOR=duration of response; INRC=International Neuroblastoma Response Criteria; NE=not estimable; ORR=overall response rate; PR=partial response.

IMPORTANT SAFETY INFORMATION

WARNINGS AND PRECAUTIONS

Neurotoxicity (cont)

Pain

Pain, including abdominal pain, bone pain, neck pain, and extremity pain, occurred in 100% of patients in Study 201 and 94% of patients in Study 12-230. Grade 3 pain occurred in 72% of patients in Study 201. One patient in Study 201 (4%) required interruption of an infusion due to pain. Pain typically began during the infusion of DANYELZA and lasted a median of less than one day in Study 201 (range less than one day and up to 62 days).

Premedicate with drugs that treat neuropathic pain (e.g., gabapentin) and oral opioids. Administer intravenous opioids as needed for breakthrough pain.

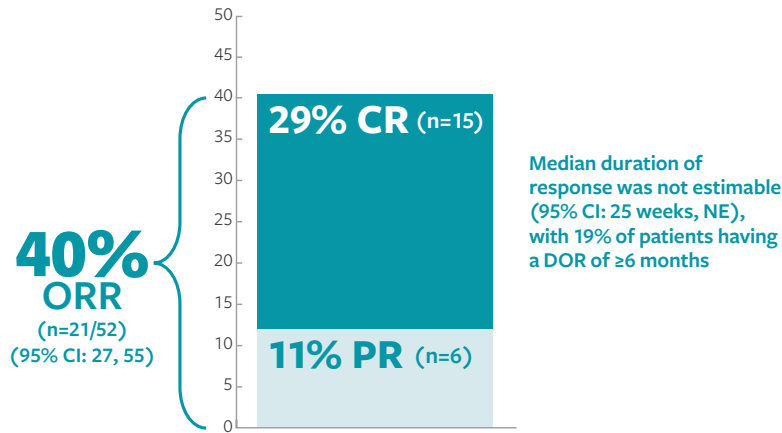
Permanently discontinue DANYELZA based on severity. [CONTINUE READING >](#)



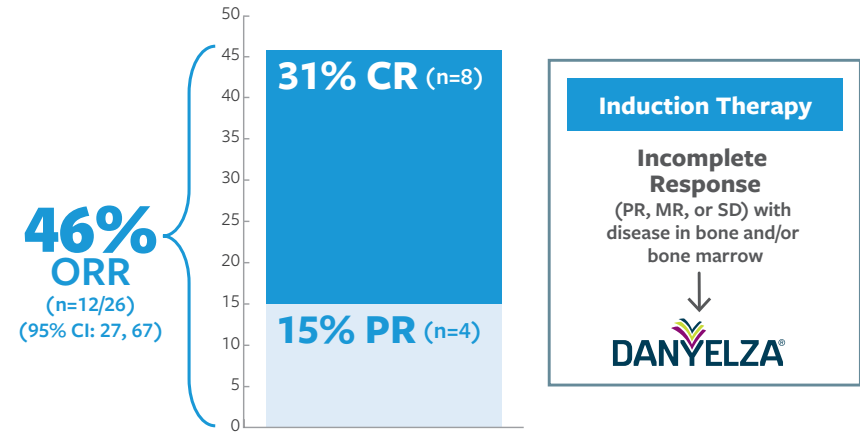
Bone/BM in NB	Incomplete Response	Clinical Studies	Efficacy	MOA	Safety	Dosing & Administration	Y-mAbs Connect	References	Important Safety Information	Summary
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Study 201 pre-specified interim analysis⁹

All Efficacy-Evaluable Patients* (n=52)



Patients with Incomplete Response to Induction[†] (n=26)



ORR was defined as a CR or PR according to the revised INRC (2017) *and confirmed by at least 1 subsequent assessment*

Effectiveness of DANYELZA with GM-CSF was evaluated by independent pathology and imaging review. Responses were observed in the bone, bone marrow, or both bone and bone marrow.⁹

*Median follow-up: 5.9 months (range: 0.6–17.8).

For the primary endpoint, a sample size of at least 37 patients in the efficacy population is sufficient to ensure at least 90% power to exclude an ORR of 20% or less at the two-sided 5% level.⁹

Limitations: Interim analysis may not be representative of the final analysis.

[†]Study design: These data underwent pre-specified analyses, including subgroup analyses of the primary endpoint.⁹

Limitations: These subgroup results are based on small sample sizes and could represent chance findings, and they were not adjusted for multiplicity; interpret with caution.⁹

IMPORTANT SAFETY INFORMATION

WARNINGS AND PRECAUTIONS

Neurotoxicity (cont)

Transverse Myelitis

Transverse myelitis has occurred with DANYELZA. Permanently discontinue DANYELZA in patients who develop transverse myelitis.

Reversible Posterior Leukoencephalopathy Syndrome (RPLS)

Reversible posterior leukoencephalopathy syndrome (RPLS) (also known as posterior reversible encephalopathy syndrome or PRES) occurred in 2 (2.8%) patients in Study 12-230. Events occurred 2 and 7 days following completion of the first cycle of DANYELZA. Monitor blood pressure during and following DANYELZA infusion and assess for neurologic symptoms. Permanently discontinue DANYELZA in case of symptomatic RPLS. [CONTINUE READING >](#)




Bone/BM in NB	Incomplete Response	Clinical Studies	Efficacy	MOA	Safety	Dosing & Administration	Y-mAbs Connect	References	Important Safety Information	Summary
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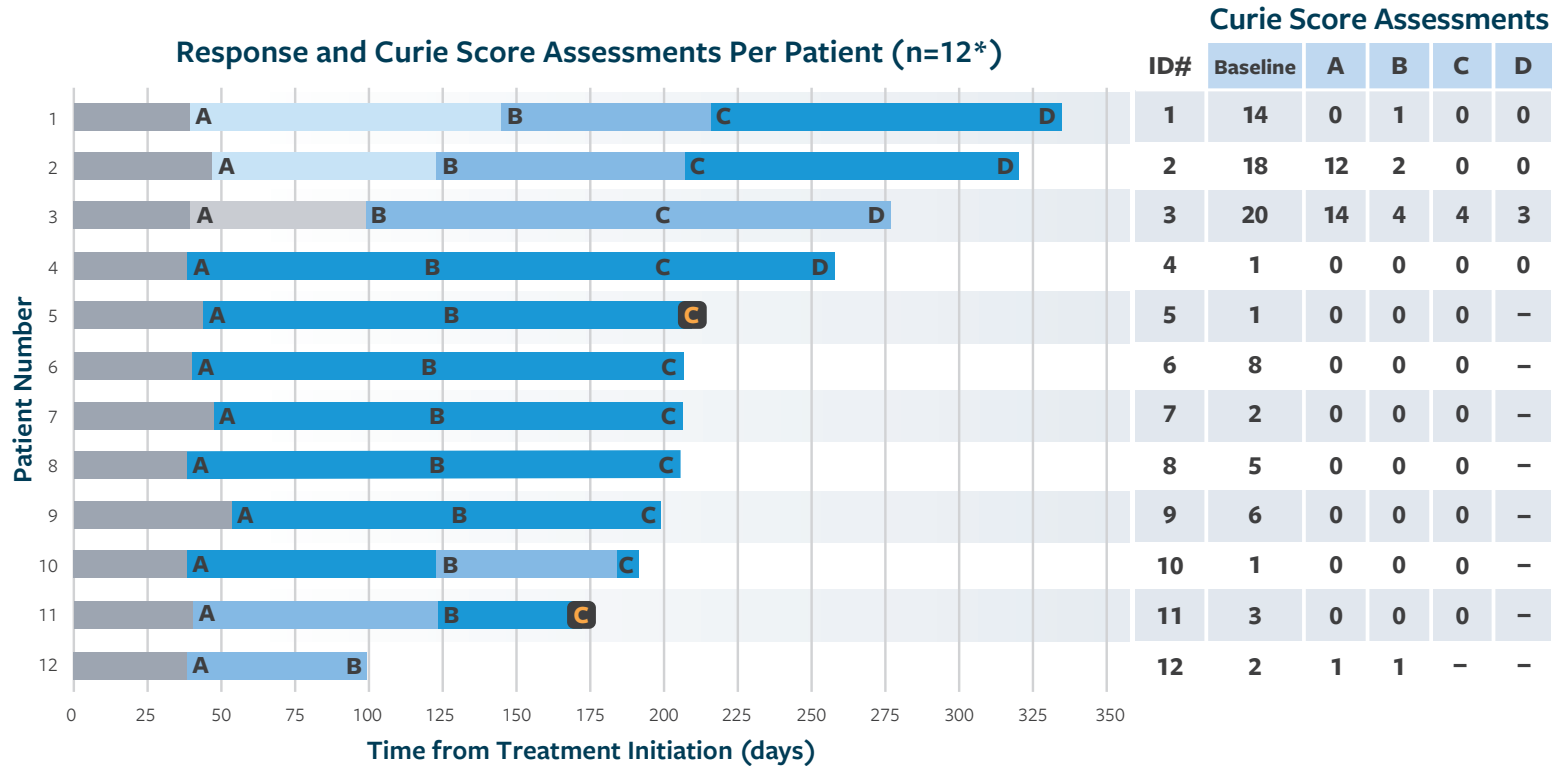
Study 201 pre-specified interim analysis: swimmer plot of patients with incomplete response to induction⁹

Induction Therapy

Incomplete Response
(PR, MR, or SD) with disease in bone and/or bone marrow



- No response assessment
- Stable disease
- Minor response
- Partial response
- Complete response
- Progressive disease



*Patients with a best response of minor response (MR), stable disease (SD), or progressive disease (PD) to DANYELZA with GM-CSF are excluded from the swimmer plot.

Limitations: Patient-level data are for descriptive purposes and should not be considered indicative of typical product efficacy or duration; interpret with caution.

IMPORTANT SAFETY INFORMATION

WARNINGS AND PRECAUTIONS

Neurotoxicity (cont)

Peripheral Neuropathy

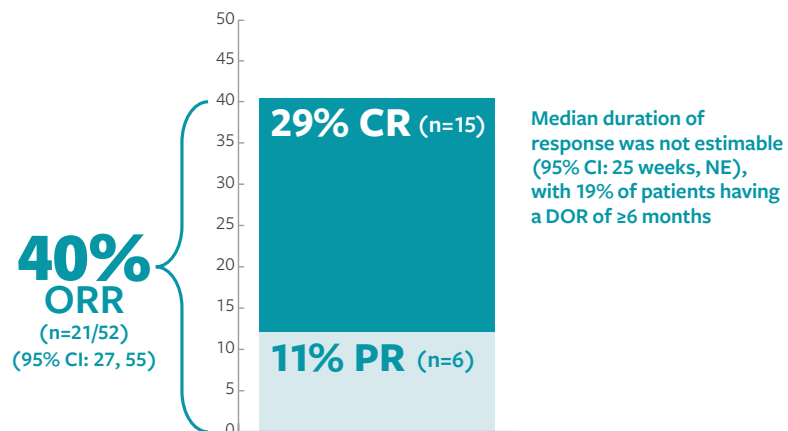
Peripheral neuropathy, including peripheral sensory neuropathy, peripheral motor neuropathy, paresthesia, and neuralgia, occurred in 32% of patients in Study 201 and in 25% of patients in Study 12-230. Most signs and symptoms of neuropathy began on the day of the infusion and neuropathy lasted a median of 5.5 days (range 0 to 22 days) in Study 201 and 0 days (range 0 to 22 days) in Study 12-230.

Permanently discontinue DANYELZA based on severity. [CONTINUE READING >](#)

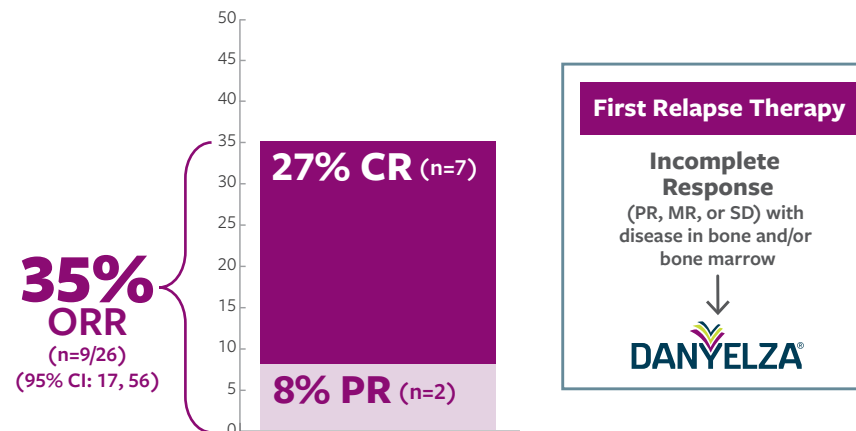


Study 201 pre-specified interim analysis⁹

All Efficacy-Evaluable Patients* (n=52)



Patients with Incomplete Response to Relapse Therapy[†] (n=26)



ORR was defined as a CR or PR according to the revised INRC (2017) *and confirmed by at least 1 subsequent assessment*

Effectiveness of DANYELZA with GM-CSF was evaluated by independent pathology and imaging review. Responses were observed in the bone, bone marrow, or both bone and bone marrow.⁹

*Median follow-up: 5.9 months (range: 0.6–17.8).

For the primary endpoint, a sample size of at least 37 patients in the efficacy population is sufficient to ensure at least 90% power to exclude an ORR of 20% or less at the two-sided 5% level.⁹

Limitations: Interim analysis may not be representative of the final analysis.

[†]Study design: These data underwent pre-specified analyses, including subgroup analyses of the primary endpoint.⁹

Limitations: These subgroup results are based on small sample sizes and could represent chance findings, and they were not adjusted for multiplicity; interpret with caution.⁹

IMPORTANT SAFETY INFORMATION

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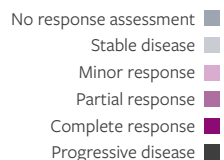
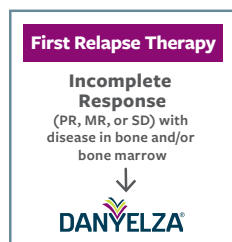
Neurotoxicity (cont)

Neurological Disorders of the Eye

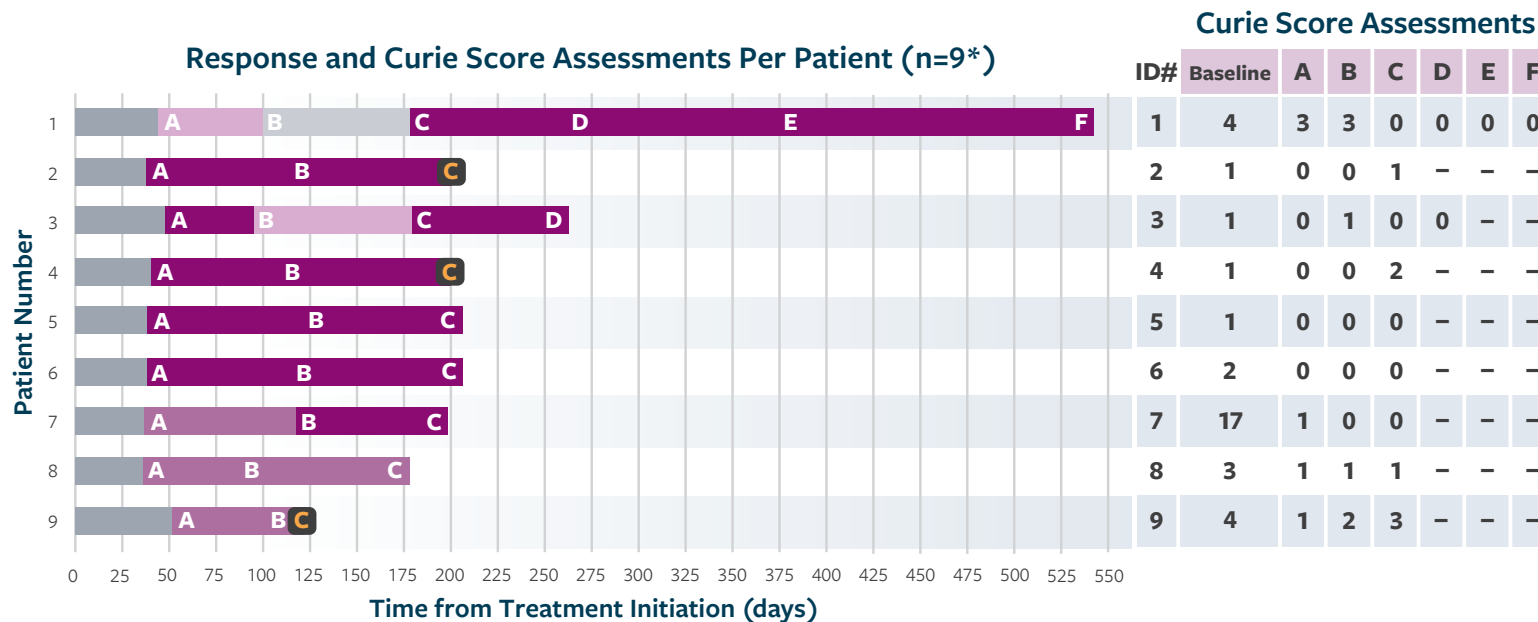
Neurological disorders of the eye including unequal pupils, blurred vision, accommodation disorder, mydriasis, visual impairment, and photophobia occurred in 24% of patients in Study 201 and 19% of patients in Study 12-230. Neurological disorders of the eye lasted a median of 17 days (range 0 to 84 days) in Study 201 with two patients (8%) experiencing an event that had not resolved at the time of data cutoff, and a median of 1 day (range less than one day to 21 days) in Study 12-230. Permanently discontinue DANYELZA based on severity. [CONTINUE READING >](#)

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Study 201 pre-specified interim analysis: swimmer plot of patients with incomplete response to relapse therapy⁹



Response and Curie Score Assessments Per Patient (n=9*)



*Patients with a best response of minor response (MR), stable disease (SD), or progressive disease (PD) to DANYELZA with GM-CSF are excluded from the swimmer plot.

Limitations: Patient-level data are for descriptive purposes and should not be considered indicative of typical product efficacy or duration; interpret with caution.

IMPORTANT SAFETY INFORMATION

WARNINGS AND PRECAUTIONS

Neurotoxicity (cont)

Prolonged Urinary Retention

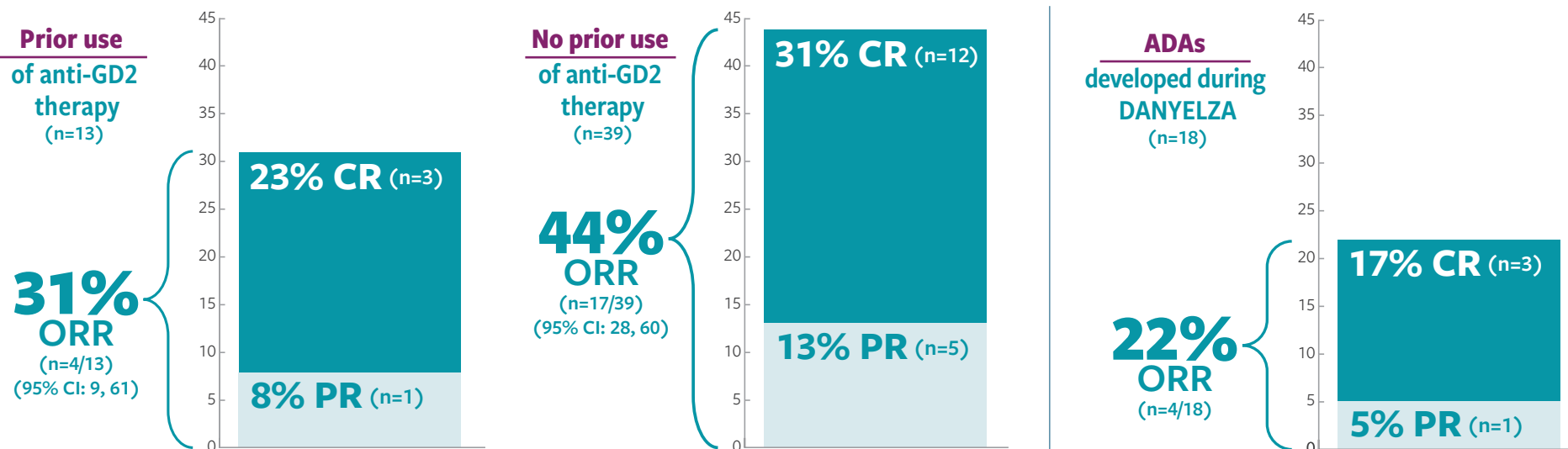
Urinary retention occurred in 1 (4%) patient in Study 201 and in 3 patients (4%) in Study 12-230. All events in both studies occurred on the day of an infusion of DANYELZA and lasted between 0 and 24 days. Permanently discontinue DANYELZA in patients with urinary retention that does not resolve following discontinuation of opioids.

Myocarditis

Myocarditis has occurred in adolescent patients receiving DANYELZA in clinical trials and expanded access programs. Myocarditis occurred within days of receiving DANYELZA requiring drug interruption. Monitor for signs and symptoms of myocarditis during treatment with DANYELZA. Withhold, reduce the dose, or permanently discontinue DANYELZA based on severity. [CONTINUE READING >](#)



Study 201 pre-specified interim analysis: patients by prior use of anti-GD2 therapy and those who developed anti-drug antibodies (ADAs)⁹



ORR was defined as a CR or PR according to the revised INRC (2017) *and confirmed by at least 1 subsequent assessment*

Effectiveness of DANYELZA with GM-CSF was evaluated by independent pathology and imaging review. Responses were observed in the bone, bone marrow, or both bone and bone marrow.⁹

DANYELZA is the only FDA-approved anti-GD2 immunotherapy approved for this patient population (ie, relapsed or refractory high-risk neuroblastoma in the bone or bone marrow).¹

Study design: These data underwent pre-specified analyses, including subgroup analyses of the primary endpoint.⁹

Limitations: These subgroup results are based on small sample sizes and could represent chance findings, and they were not adjusted for multiplicity; interpret with caution.⁹

IMPORTANT SAFETY INFORMATION

WARNINGS AND PRECAUTIONS

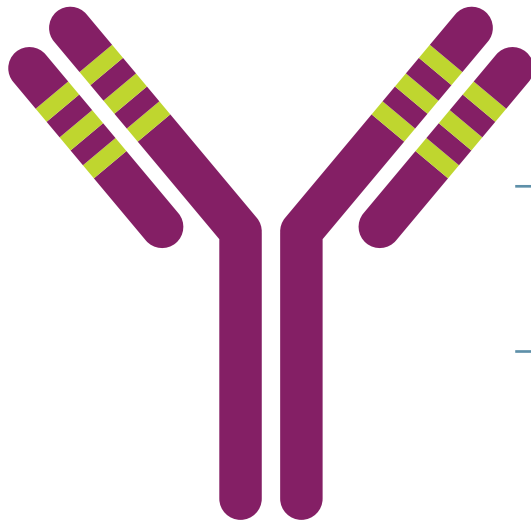
Hypertension

Hypertension occurred in 44% of patients in Study 201 and 28% of patients in Study 12-230 who received DANYELZA. Grade 3 or 4 hypertension occurred in 4% of patients in Study 201 and 7% of patients in Study 12-230. Four patients (6%) in Study 12-230 permanently discontinued DANYELZA due to hypertension. In both studies, most events occurred on the day of DANYELZA infusion and occurred up to 9 days following an infusion of DANYELZA.

Do not initiate DANYELZA in patients with uncontrolled hypertension. Monitor blood pressure during infusion, and at least daily on Days 1 to 8 of each cycle of DANYELZA and evaluate for complications of hypertension including RPLS. Interrupt DANYELZA infusion and resume at a reduced rate, or permanently discontinue DANYELZA based on the severity. [CONTINUE READING >](#)

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DANYELZA, the only humanized GD2-binding monoclonal antibody approved by the FDA,* is structurally distinct^{1,11†}



Neuroblastoma is characterized by an overexpression of GD2, a disialoganglioside also found in the central nervous system and on peripheral nerves¹

Antibody structure¹²

- 92% human framework
- 8% murine framework

~10-fold higher binding affinity to the GD2 receptor due to a slower off-rate than approved chimeric anti-GD2 antibodies shown in *in vitro* studies¹¹

Clinical significance and product comparisons of efficacy or safety should not be inferred

*In refractory or relapsed high-risk neuroblastoma.

†Based on *in vitro* data.

IMPORTANT SAFETY INFORMATION

WARNINGS AND PRECAUTIONS

Orthostatic Hypotension

Orthostatic hypotension has occurred in patients receiving DANYELZA in clinical trials and expanded access programs. Severe orthostatic hypotension, including cases requiring hospitalization, have occurred. Cases occurred within hours to 6 days of DANYELZA infusions in any cycle.

In patients with symptoms of orthostatic hypotension, monitor postural blood pressure prior to initiating treatment with DANYELZA and as clinically indicated with subsequent dosing. Withhold, reduce dose, or permanently discontinue DANYELZA based on severity. [CONTINUE READING >](#)

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Safety analysis of patients who received DANYELZA with GM-CSF

DANYELZA can cause serious infusion reactions, including hypotension, bronchospasm, hypoxia, and stridor, as well as severe neurotoxicity, including pain¹:

- Any-grade infusion-related reactions occurred in **94%–100%** of patients
 - Any-grade hypotension occurred in **89%–100%** of patients
- Any-grade pain occurred in **94%–100%** of patients

The most common ARs in Studies 12-230 and 201 (both analyses) (≥25% in either study)^{1,9}

- | | |
|-----------------------------|---------------------------|
| ■ Infusion-related reaction | ■ Erythema multiforme |
| ■ Pain | ■ Peripheral neuropathy |
| ■ Tachycardia | ■ Urticaria |
| ■ Vomiting | ■ Pyrexia |
| ■ Cough | ■ Headache |
| ■ Pruritus | ■ Injection site reaction |
| ■ Nausea | ■ Edema |
| ■ Diarrhea | ■ Anxiety |
| ■ Decreased appetite | ■ Localized edema |
| ■ Hypertension | ■ Irritability |
| ■ Fatigue | ■ Anemia |

Total DANYELZA exposure across Studies 12-230 and 201^{1,9}

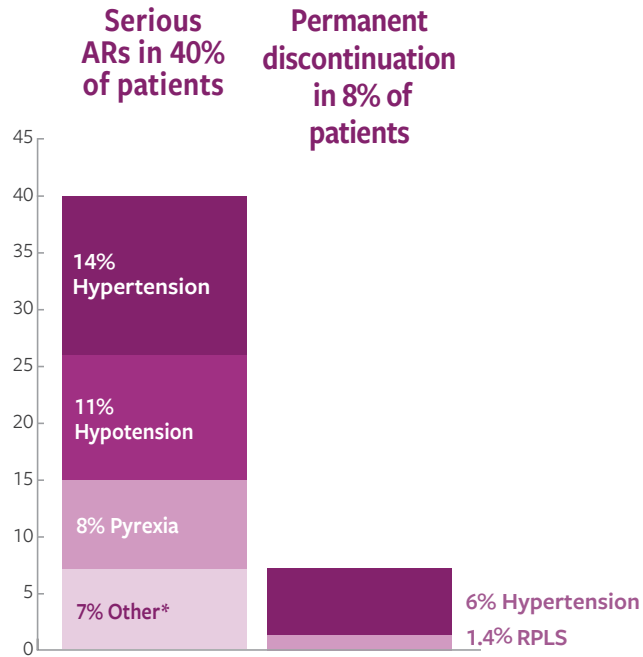
- Of the 72 patients in Study 12-230, 32% were exposed to DANYELZA with GM-CSF for ≥6 months and 8% for >1 year¹
- Of the 25 patients in Study 201 (initial analysis), an ongoing multicenter trial, 12% were exposed to DANYELZA with GM-CSF for ≥6 months and 0% for >1 year¹
- Of the 74 patients in the Study 201 pre-specified interim analysis, 18% were exposed to DANYELZA with GM-CSF for ≥6 months and 3% for ≥1 year⁹

AR=adverse reaction.

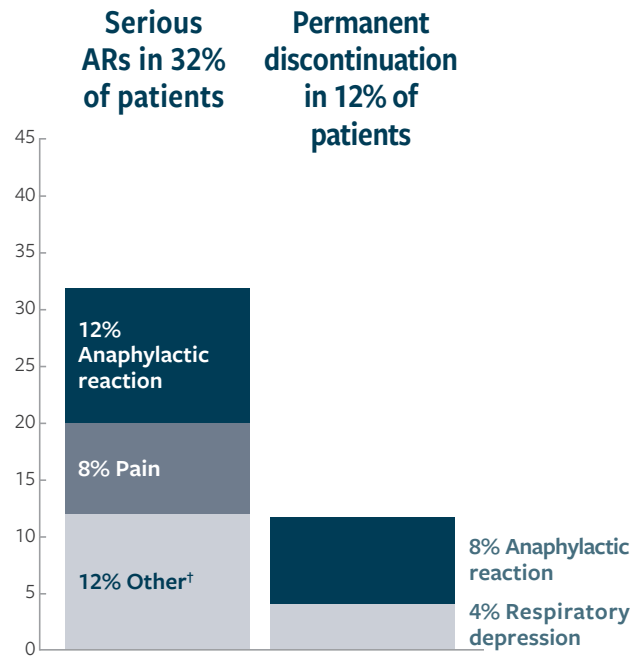


Some patients experienced serious adverse reactions that led to permanent discontinuation^{1,9}

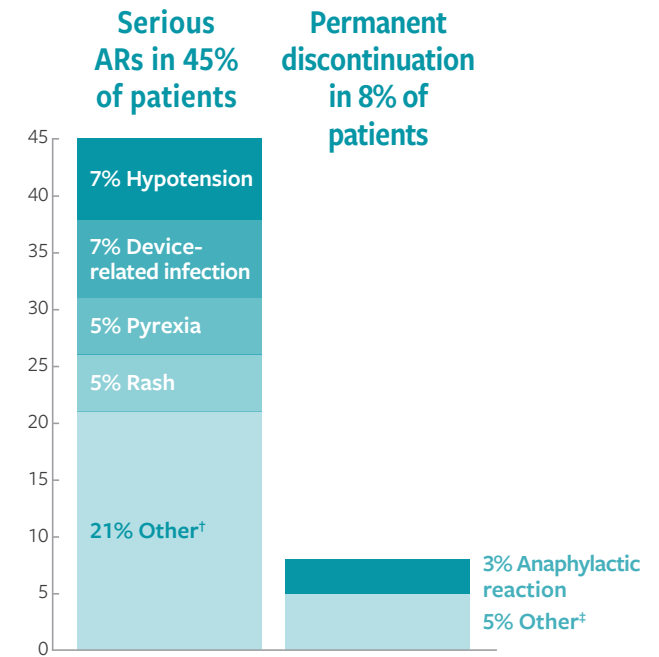
STUDY 12-230 (N=72)¹



STUDY 201 Initial Analysis (N=25)¹



STUDY 201 Pre-specified Interim Analysis (N=74)⁹



- In the Study 201 initial analysis, dose interruptions due to an AR occurred in 84% of patients. ARs requiring dosage interruption in >10% of patients included hypotension and bronchospasm¹
- In the Study 201 pre-specified interim analysis, dose interruptions due to an AR occurred in 69% of patients. ARs requiring dosage interruption in >10% of patients included hypotension, pain, and bronchospasm⁹

*Serious ARs occurring in <5% of patients.

† Serious ARs occurring in only 1 patient.

‡ 1% each: respiratory depression, myocarditis, hypotension, RPLS, and urticaria.

RPLS=reversible posterior leukoencephalopathy syndrome.

When to permanently discontinue DANYELZA¹

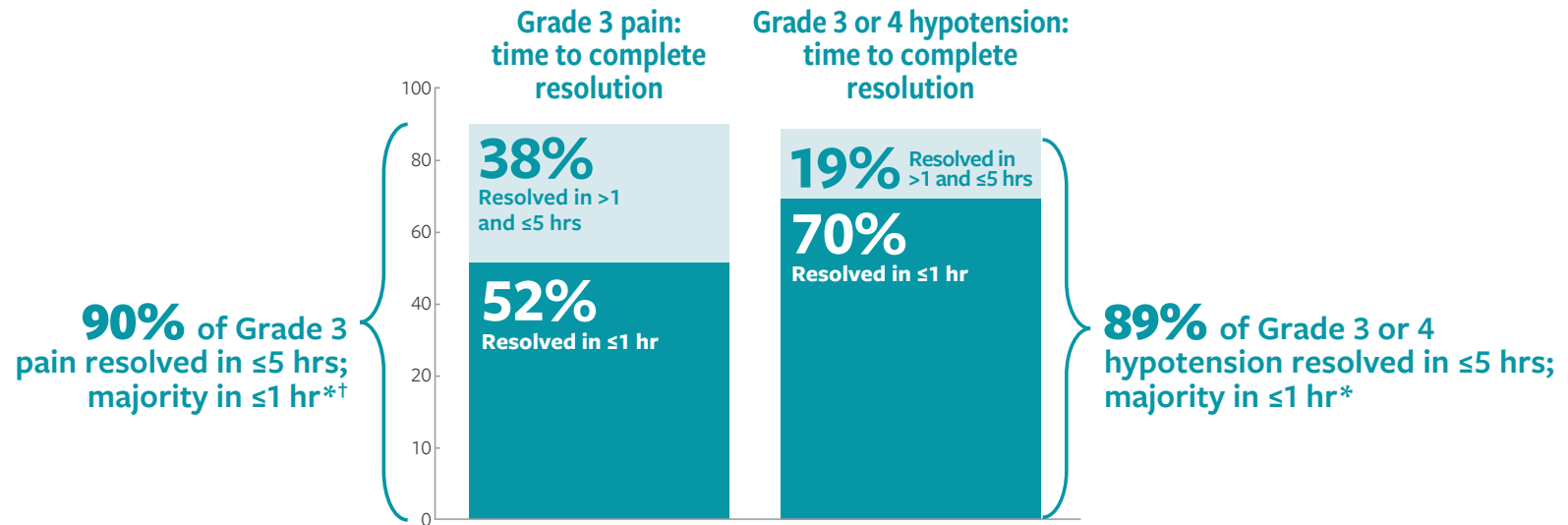
DANYELZA should be discontinued in the case of*:

Infusion-related reactions	<ul style="list-style-type: none"> Grade 4, Grade 3 and not responding to medical intervention, or Grade 3-4 anaphylaxis 	Prolonged urinary retention	<ul style="list-style-type: none"> Persisting following discontinuation of opioids
Pain	<ul style="list-style-type: none"> Grade 3 and unresponsive to maximum supportive measures 	Myocarditis	<ul style="list-style-type: none"> Grade 4, Grade 2 or 3 based on severity and duration
Reversible posterior leukoencephalopathy syndrome (RPLS)	<ul style="list-style-type: none"> All grades 	Hypertension	<ul style="list-style-type: none"> Grade 4, or Grade 3 and not responding to medical intervention
Transverse myelitis	<ul style="list-style-type: none"> All grades 	Orthostatic hypotension	<ul style="list-style-type: none"> Any grade not resolved within 1 week
Peripheral neuropathy	<ul style="list-style-type: none"> Grade ≥2 motor neuropathy or Grade 3-4 sensory neuropathy 	Other ARs	<ul style="list-style-type: none"> Grade 4, or Grade 3 not resolving to Grade ≤2 within 2 weeks
Neurological disorders of the eye	<ul style="list-style-type: none"> Grade 2-4 not resolving within 2 weeks or upon recurrence; any grade with subtotal or total vision loss 		

*Based on Common Terminology Criteria for Adverse Events (CTCAE) v5.0.



Study 201 pre-specified interim analysis: resolution of select Grade 3 or 4 adverse reactions⁹



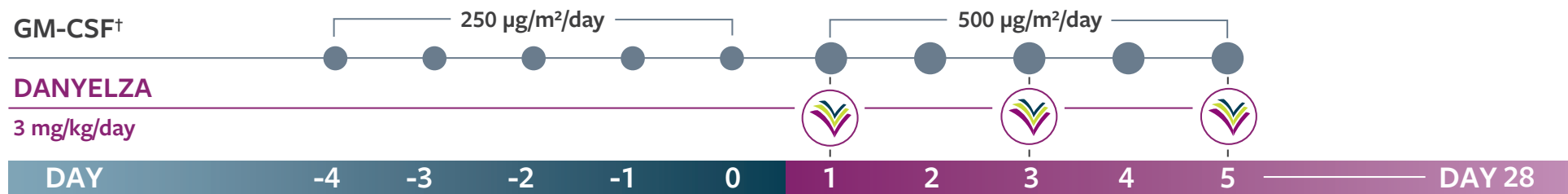
*Incidence of events related to DANYELZA or DANYELZA with GM-CSF occurring on day of infusion, after start of infusion.

[†]Excludes procedural pain and vessel puncture site pain.



DANYELZA offers the flexibility of outpatient or inpatient administration, at the treating physician's discretion⁹

>90% of infusions were given in an outpatient setting in the Study 201 pre-specified interim analysis^{9*}



Pretreatment¹

- **Five days before first infusion in each cycle:** initiate GM-CSF and a 12-day course (Day -4 through Day 7) of gabapentin or other prophylactic medication for neuropathic pain

Infusion days¹

- DANYELZA is given on **Days 1, 3, and 5 of each 28-day cycle** until disease progression or unacceptable toxicity
- In preparation for each DANYELZA dose:
 - **2 hours to 30 min before DANYELZA:** premedicate (see following pages)
 - **≥1 hour before infusion** on Days 1, 3, 5 of each cycle: administer GM-CSF
- Administer DANYELZA 3 mg/kg/infusion (up to 150 mg/day) on Days 1, 3, 5 (9 mg/kg/cycle), given as IV infusion after dilution and in combination with GM-CSF subcutaneously. Do not administer DANYELZA as IV push or bolus
 - **60-min first infusion** (Cycle 1, Day 1) and subsequently 30-60 min as tolerated
 - **Observation required for at least 2 hours after** the DANYELZA infusion in a setting where cardiopulmonary resuscitation medication and equipment are available

*Out of 1,237 infusions, 92.5% (1,144) were outpatient and 7.5% (93) were inpatient.⁹

¹For more details, refer to the GM-CSF Prescribing Information.

IV=intravenous.

IMPORTANT SAFETY INFORMATION

WARNINGS AND PRECAUTIONS

Embryo-Fetal Toxicity

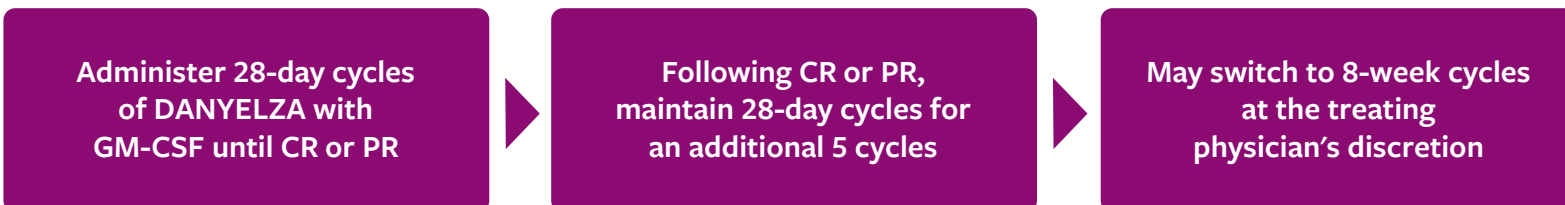
Based on its mechanism of action, DANYELZA may cause fetal harm when administered to a pregnant woman. Advise females of reproductive potential, including pregnant women, of the potential risk to a fetus. Advise females of reproductive potential to use effective contraceptive during treatment with DANYELZA and for two months after the last dose. [CONTINUE READING >](#)



Bone/BM in NB	Incomplete Response	Clinical Studies	Efficacy	MOA	Safety	Dosing & Administration	Y-mAbs Connect	References	Important Safety Information	Summary
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Administer DANYELZA until CR or PR and follow treatment course shown below¹

Treatment Course¹



Discontinue for disease progression or unacceptable toxicity

If a DANYELZA dose is missed¹

- Administer the missed dose the following week by Day 10
- Administer GM-CSF 500 µg/m²/day on the first day of the DANYELZA infusion, and on the day before and the days of the second and third infusions (ie, a total of 5 days with 500 µg/m²/day)

IMPORTANT SAFETY INFORMATION

ADVERSE REACTIONS

The most common adverse reactions in Studies 201 and 12-230 (≥25% in either study) were infusion-related reaction, pain, tachycardia, vomiting, cough, nausea, diarrhea, decreased appetite, hypertension, fatigue, erythema multiforme, peripheral neuropathy, urticaria, pyrexia, headache, injection site reaction, edema, anxiety, localized edema and irritability. The most common Grade 3 or 4 laboratory abnormalities (≥5% in either study) were decreased lymphocytes, decreased neutrophils, decreased hemoglobin, decreased platelet count, decreased potassium, increased alanine aminotransferase, decreased glucose, decreased calcium, decreased albumin, decreased sodium and decreased phosphate. [CONTINUE READING >](#)

DANYELZA[®]
(naxitamab-ggqk)
40mg/10mL Injection



Y-mAbs Connect[®] is a patient support program that provides information about access, insurance, financial support, and other resource programs for qualifying patients



Your link to patient support

ymabsconnect.com or 1-833-33YMABS, option 2



Y-mAbs Connect

Healthcare professionals get help with:

- Summary of Benefits for health insurance coverage of DANYELZA, including assistance in determining when a prior authorization or appeal may be needed
- Information on ordering DANYELZA

Patients get help with:

- Determining eligibility for Y-mAbs Connect Patient Support Programs
- Information on third-party organizations* that may help with logistical and other support

*Third-party organizations are not associated with Y-mAbs Therapeutics, Inc.; specific details and eligibility requirements may vary by organization.

DANYELZA J-code: J9348

Information about Y-mAbs Connect can be found at ymabsconnect.com or by calling Y-mAbs Connect at 1-833-33YMABS, option 2 between 8:00 am – 8:00 pm ET, Monday – Friday. Closed on weekends and major holidays.

DANYELZA[®]
(naxitamab-ggqk)
40mg/10mL Injection

References

- 1.** DANYELZA[®] [package insert]. New York, NY: Y-mAbs Therapeutics, Inc.; 2024. Available online at <https://labeling.ymabs.com/danyelza>.
- 2.** Park JR, Bagatell R, Cohn SL, et al. *J Clin Oncol*. 2017;35(22):2580-2587.
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- 4.** Garaventa A, Poetschger U, Valteau-Couanet D, et al. *J Clin Oncol*. 2021;39(23):2552-2563.
- 5.** Pinto N, Naranjo A, Hibbitts E, et al. *Eur J Cancer*. 2019;112:66-79.
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- 7.** Yanik GA, Parisi MT, Shulkin BL, et al. *J Nucl Med*. 2013;54(4):541-548.
- 8.** Streby KA, Parisi MT, Shulkin BL, et al. *Pediatr Blood Cancer*. 2023;e30418. <https://doi.org/10.1002/pbc.30418>.
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- 10.** NIH US National Library of Medicine. <https://clinicaltrials.gov/ct2/show/NCT01419834?term=NCT01419834&draw=2&rank=1>. Accessed April 22, 2024.
- 11.** Lisby S, Liebenberg N, Bukrinski J, et al. Presented at the SIOP virtual congress. Abstract #945. October 16, 2020.
- 12.** Cheung N-KV, Guo H, Hu J, et al. *Oncoimmunology*. 2012;1(4):477-486.





Indication and Important Safety Information

INDICATION

DANYELZA is indicated, in combination with granulocyte-macrophage colony-stimulating factor (GM-CSF), for the treatment of pediatric patients 1 year of age and older and adult patients with relapsed or refractory high-risk neuroblastoma in the bone or bone marrow who have demonstrated a partial response, minor response, or stable disease to prior therapy.

This indication is approved under accelerated approval based on overall response rate and duration of response. Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial(s).

IMPORTANT SAFETY INFORMATION

WARNING: SERIOUS INFUSION-RELATED REACTIONS and NEUROTOXICITY

Serious Infusion-Related Reactions

- DANYELZA can cause serious infusion reactions, including cardiac arrest, anaphylaxis, hypotension, bronchospasm, and stridor. Infusion reactions of any Grade occurred in 94-100% of patients. Severe infusion reactions occurred in 32-68% and serious infusion reactions occurred in 4-18% of patients in DANYELZA clinical studies.
- Premedicate prior to each DANYELZA infusion as recommended and monitor patients for at least 2 hours following completion of each infusion. Reduce the rate, interrupt infusion, or permanently discontinue DANYELZA based on severity.

Neurotoxicity

- DANYELZA can cause severe neurotoxicity, including severe neuropathic pain, transverse myelitis and reversible posterior leukoencephalopathy syndrome (RPLS). Pain of any Grade occurred in 94-100% of patients in DANYELZA clinical studies.
- Premedicate to treat neuropathic pain as recommended. Permanently discontinue DANYELZA based on the adverse reaction and severity.

CONTRAINDICATION

DANYELZA is contraindicated in patients with a history of severe hypersensitivity reaction to naxitamab-gqgk. Reactions have included anaphylaxis.

WARNINGS AND PRECAUTIONS

Serious Infusion-Related Reactions

DANYELZA can cause serious infusion reactions requiring urgent intervention including fluid resuscitation, administration of bronchodilators and corticosteroids, intensive care unit admission, infusion rate reduction or interruption of DANYELZA infusion. Infusion-related reactions included hypotension, bronchospasm, hypoxia, and stridor.

Serious infusion-related reactions occurred in 4% of patients in Study 201 and in 18% of patients in Study 12-230. Infusion-related reactions of any Grade occurred in 100% of patients in Study 201 and 94% of patients in Study 12-230. Hypotension of any grade occurred in 100% of patients in Study 201 and 89% of patients in Study 12-230.

In Study 201, 68% of patients experienced Grade 3 or 4 infusion reactions; and in Study 12-230, 32% of patients experienced Grade 3 or 4 infusion reactions. Anaphylaxis occurred in 12% of patients and two patients (8%) permanently discontinued DANYELZA due to anaphylaxis in Study 201. One patient in Study 12-230 (1.4%) experienced a Grade 4 cardiac arrest 1.5 hours following completion of DANYELZA infusion.

In Study 201, infusion reactions generally occurred within 24 hours of completing a DANYELZA infusion, most often within 30 minutes of initiation. Infusion reactions were most frequent during the first infusion of DANYELZA in each cycle. Eighty percent of patients required reduction in infusion rate and 80% of patients had an infusion interrupted for at least one infusion-related reaction.

Caution is advised in patients with pre-existing cardiac disease, as this may exacerbate the risk of severe hypotension.



Bone/BM in NB	Incomplete Response	Clinical Studies	Efficacy	MOA	Safety	Dosing & Administration	Y-mAbs Connect	References	Important Safety Information	Summary
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Important Safety Information (cont)

WARNINGS AND PRECAUTIONS

Serious Infusion-Related Reactions (cont)

Premedicate with an antihistamine, acetaminophen, an H2 antagonist and corticosteroid as recommended. Monitor patients closely for signs and symptoms of infusion reactions during and for at least 2 hours following completion of each DANYELZA infusion in a setting where cardiopulmonary resuscitation medication and equipment are available.

Reduce the rate, interrupt infusion, or permanently discontinue DANYELZA based on severity and institute appropriate medical management as needed.

Neurotoxicity

DANYELZA can cause severe neurotoxicity, including severe neuropathic pain, transverse myelitis, and reversible posterior leukoencephalopathy syndrome.

Pain

Pain, including abdominal pain, bone pain, neck pain, and extremity pain, occurred in 100% of patients in Study 201 and 94% of patients in Study 12-230. Grade 3 pain occurred in 72% of patients in Study 201. One patient in Study 201 (4%) required interruption of an infusion due to pain. Pain typically began during the infusion of DANYELZA and lasted a median of less than one day in Study 201 (range less than one day and up to 62 days).

Premedicate with drugs that treat neuropathic pain (e.g., gabapentin) and oral opioids. Administer intravenous opioids as needed for breakthrough pain. Permanently discontinue DANYELZA based on severity.

Transverse Myelitis

Transverse myelitis has occurred with DANYELZA. Permanently discontinue DANYELZA in patients who develop transverse myelitis.

Reversible Posterior Leukoencephalopathy Syndrome (RPLS)

Reversible posterior leukoencephalopathy syndrome (RPLS) (also known as posterior reversible encephalopathy syndrome or PRES) occurred in 2 (2.8%) patients in Study 12-230. Events occurred 2 and 7 days following completion of the first cycle of DANYELZA. Monitor blood pressure during and following DANYELZA infusion and assess for neurologic symptoms. Permanently discontinue DANYELZA in case of symptomatic RPLS.

Peripheral Neuropathy

Peripheral neuropathy, including peripheral sensory neuropathy, peripheral motor neuropathy, paresthesia, and neuralgia, occurred in 32% of patients in Study 201 and in 25% of patients in Study 12-230. Most signs and symptoms of neuropathy began on the day of the infusion and neuropathy lasted a median of 5.5 days (range 0 to 22 days) in Study 201 and 0 days (range 0 to 22 days) in Study 12-230.

Permanently discontinue DANYELZA based on severity.

Neurological Disorders of the Eye

Neurological disorders of the eye including unequal pupils, blurred vision, accommodation disorder, mydriasis, visual impairment, and photophobia occurred in 24% of patients in Study 201 and 19% of patients in Study 12-230. Neurological disorders of the eye lasted a median of 17 days (range 0 to 84 days) in Study 201 with two patients (8%) experiencing an event that had not resolved at the time of data cutoff, and a median of 1 day (range less than one day to 21 days) in Study 12-230. Permanently discontinue DANYELZA based on severity.

Prolonged Urinary Retention

Urinary retention occurred in 1 (4%) patient in Study 201 and in 3 patients (4%) in Study 12-230. All events in both studies occurred on the day of an infusion of DANYELZA and lasted between 0 and 24 days. Permanently discontinue DANYELZA in patients with urinary retention that does not resolve following discontinuation of opioids.





Important Safety Information (cont)

WARNINGS AND PRECAUTIONS

Myocarditis

Myocarditis has occurred in adolescent patients receiving DANYELZA in clinical trials and expanded access programs. Myocarditis occurred within days of receiving DANYELZA requiring drug interruption. Monitor for signs and symptoms of myocarditis during treatment with DANYELZA. Withhold, reduce the dose, or permanently discontinue DANYELZA based on severity.

Hypertension

Hypertension occurred in 44% of patients in Study 201 and 28% of patients in Study 12-230 who received DANYELZA. Grade 3 or 4 hypertension occurred in 4% of patients in Study 201 and 7% of patients in Study 12-230. Four patients (6%) in Study 12-230 permanently discontinued DANYELZA due to hypertension. In both studies, most events occurred on the day of DANYELZA infusion and occurred up to 9 days following an infusion of DANYELZA.

Do not initiate DANYELZA in patients with uncontrolled hypertension. Monitor blood pressure during infusion, and at least daily on Days 1 to 8 of each cycle of DANYELZA and evaluate for complications of hypertension including RPLS. Interrupt DANYELZA infusion and resume at a reduced rate, or permanently discontinue DANYELZA based on the severity.

Orthostatic Hypotension

Orthostatic hypotension has occurred in patients receiving DANYELZA in clinical trials and expanded access programs. Severe orthostatic hypotension, including cases requiring hospitalization, have occurred. Cases occurred within hours to 6 days of DANYELZA infusions in any cycle.

In patients with symptoms of orthostatic hypotension, monitor postural blood pressure prior to initiating treatment with DANYELZA and as clinically indicated with subsequent dosing. Withhold, reduce dose, or permanently discontinue DANYELZA based on severity.

Embryo-Fetal Toxicity

Based on its mechanism of action, DANYELZA may cause fetal harm when administered to a pregnant woman. Advise females of reproductive potential, including pregnant women, of the potential risk to a fetus. Advise females of reproductive potential to use effective contraceptive during treatment with DANYELZA and for two months after the last dose.

ADVERSE REACTIONS

The most common adverse reactions in Studies 201 and 12-230 ($\geq 25\%$ in either study) were infusion-related reaction, pain, tachycardia, vomiting, cough, nausea, diarrhea, decreased appetite, hypertension, fatigue, erythema multiforme, peripheral neuropathy, urticaria, pyrexia, headache, injection site reaction, edema, anxiety, localized edema and irritability. The most common Grade 3 or 4 laboratory abnormalities ($\geq 5\%$ in either study) were decreased lymphocytes, decreased neutrophils, decreased hemoglobin, decreased platelet count, decreased potassium, increased alanine aminotransferase, decreased glucose, decreased calcium, decreased albumin, decreased sodium and decreased phosphate.

Please [click](#) for full Prescribing Information and Patient Information for DANYELZA including Boxed Warning on serious infusion-related reactions and neurotoxicity.

To review important state-specific disclosure information for licensed healthcare practitioners, please visit <https://www.ymabs.com/information-for-prescribers>



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In the battle against relapsed/refractory high-risk neuroblastoma

- Reducing or eliminating disease in the **bone and bone marrow** is a goal of high-risk neuroblastoma treatment²
- DANYELZA is the **only FDA-approved therapy** indicated to treat high-risk neuroblastoma in the bone and/or bone marrow when response to induction or relapse therapy is incomplete¹
- DANYELZA is a **structurally distinct, humanized** anti-GD2 monoclonal antibody that provides another immunotherapeutic option^{1,11}
- DANYELZA offers the **flexibility** to be administered in either an **outpatient or inpatient** hospital setting, at the treating physician's discretion⁹

When response to induction or relapse therapy is incomplete, **DEPLOY DANYELZA**

**Backed by >10 years of clinical trial experience
and approved by the FDA in 2020¹⁰**

For coverage and access information, visit ymabsconnect.com

Administered at an expanding
nationwide network⁹

60+ US Healthcare Institutions⁹
AND GROWING

IMPORTANT SAFETY INFORMATION

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Learn more at danyelzahcp.com

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DANYELZA[®]
(naxitamab-ggqk)
40mg/10mL Injection

25

Bone/BM
in NB

Incomplete
Response

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Efficacy

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