

FDA Briefing Document

Dermatologic and Ophthalmic Drugs Advisory Committee Meeting

December 13, 2019

The committee will discuss biologics license application (BLA) 761143, teprotumumab solution for intravenous use, submitted by Horizon Pharma Ireland, Ltd., proposed for the treatment of active Thyroid Eye Disease (TED).

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The attached package contains background information prepared by the Food and Drug Administration (FDA) for the panel members of the advisory committee. The FDA background package often contains assessments and/or conclusions and recommendations written by individual FDA reviewers. Such conclusions and recommendations do not necessarily represent the final position of the individual reviewers, nor do they necessarily represent the final position of the Review Division or Office. We have brought the biologics license application (BLA) 761143, teprotumumab solution for intravenous use, submitted by Horizon Pharma Ireland, Ltd. and proposed for the treatment of active Thyroid Eye Disease (TED) to this Advisory Committee in order to gain the Committee's insights and opinions, and the background package may not include all issues relevant to the final regulatory recommendation and instead is intended to focus on issues identified by the Agency for discussion by the advisory committee. The FDA will not issue a final determination on the issues at hand until input from the advisory committee process has been considered and all reviews have been finalized. The final determination may be affected by issues not discussed at the advisory committee meeting.

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1. Draft Topics for Consideration

A. Efficacy

Efficacy, as demonstrated by a reduction in proptosis, has been demonstrated in two adequate and well controlled studies. Eighty-two percent (82%) of patients treated with 8 doses of teprotumumab had at least a two-millimeter reduction in ptosis compared to only 16 percent of patients treated with placebo. A two-millimeter reduction is considered clinically significant because it is expected to reduce the incidence of diplopia and improve the lid coverage over the cornea. The systemic treatment also had an effect on the non-study eye, reducing ptosis in 68% of non-study eyes compared to only 9% of patients treated with placebo.

See Erratum to FDA Briefing Document

B. Onset and Duration

The onset of ptosis reduction was evident in some patients at the first evaluation examination, 6 weeks after the first infusion (three weeks after the second infusion). The duration of the effect is still under study. An extension of Review Study #1 demonstrated that the effect continued for at least 4 weeks after the last infusion. At week 72 approximately 60% of patients who had an effective reduction in ptosis had not relapsed (lost 2 millimeter of ptosis). The extension of Review Study #2, as well as the open label treatment period for patients who had not previously responded (placebo or teprotumumab) is still ongoing. Safety and efficacy of repeated courses of treatment have not been studied.

C. Limited number of subjects studied to date

Less than 90 subjects treated with infusions of teprotumumab have been enrolled in controlled clinical trials. This is a considerably smaller database than the common safety database of greater than 300 patients treated with a course of therapy. Based on the rule of 3's, with a safety database of 90 subjects, adverse events may be expected to occur at rates of 3% without the event being observed in the prior clinical trials.

D. Indication: Active Thyroid Eye Disease (TED) vs TED

Patients enrolled in the clinical trials were euthyroid and mildly hyper- or hypothyroid. There was no requirement for progressive forward motion of the globe, although proptosis was required. The potential inclusion of the term "Active" in this setting is not well defined.

E. Hyperglycemia

The drug product is an insulin-like growth factor-1 receptor (IGF-1R) inhibitor. It has the potential to interfere with glucose regulation in the body, particularly in individuals with diabetes. Some patients receiving Teprotumumab required additional amounts of insulin to maintain glycemic control.

A total of 22 placebo subjects and 21 teprotumumab subjects had fasting glucose values available at Baseline and at other time points during the treatment period. All of the subjects in the placebo group and 17 (81.0%) of the 21 subjects in the teprotumumab group had normal fasting glucose values at Baseline with no shifts from the normal range noted during treatment. Three teprotumumab subjects with normal fasting glucose values at Baseline demonstrated elevated fasting glucose values for at least 1 visit during the treatment period. None of these 3 subjects had a history of diabetes mellitus however, 2 of the 3 subjects had somewhat elevated HbA1c values at baseline. Glucose monitoring may be warranted after initiation of teprotumumab dosing.

F. Muscle Spasms

The overall incidence of *Muscle spasms* was higher in the teprotumumab group (32%) compared to the placebo group (9.5%). In the teprotumumab group, a total of 27 TEAEs of *Muscle spasms* were reported among 13 subjects. The most common sites specified involved the lower extremities (leg: 9 events; calf: 5 events; feet/toes: 5 events); other sites included hands (2 events), back (1 event), chest (1 event) and side of body (1), and no site specified for 3 events. None of the events resulted in discontinuation of study drug. No clinically significant abnormality in electrolytes (including calcium) or aspartate aminotransferase (AST) values were noted for subjects who experienced *Muscle spasms*. A total of 6 subjects requested and received treatment for the muscle spasms including magnesium (5 subjects), calcium (1 subject), vitamins B6, B12, and folic acid (1 subject) and cyclobenzaprine hydrochloride (1 subject). Seven (7) of the 13 subjects had at least 1 adverse event of *Muscle spasms* that began either on the day of an infusion (4 subjects; 11 events) or the day following an infusion (3 subjects; 3 events).

The mechanism of action for the muscle spasms remains unknown. Physician labeling could include Warning/Precaution of the possibility of teprotumumab being associated with muscle spasms.

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G. Hearing Impairment

At least five patients reported hypoacusis/loss of hearing. In addition, patients have reported tinnitus. One subject, a 32-year-old female, experienced an adverse event of *Hypoacusis* on Day 75 that spontaneously resolved the following day. In other subjects, the event did not resolve until after completion of treatment with teprotumumab. The mechanism of action for hypoacusis remains unknown. Physician labeling could include Warning/Precaution of the possibility of teprotumumab being associated with hearing impairment.

H. Diarrhea/Irritable Bowel Syndrome

Gastrointestinal complaints were reported in clinical trials by 36% of patients. Nausea and diarrhea were each reported in 12% of patients. Abdominal pain was reported in 5% of patients. None of these events resulted in discontinuation of study drug. However, one patient discontinued teprotumumab following hospitalization for systemic *E. Coli* sepsis and dehydration and another discontinued due to an episode of inflammatory bowel disease. Any potential causal association between teprotumumab and inflammatory bowel disease is uncertain at this time. Labeling can be considered.

I. Infection Rate

The reported infection rate associated with teprotumumab was 33%, and higher than that of the placebo control in both studies. No specific site of infection was identified and the potential contribution of teprotumumab to this infection rate is not known. Labeling can be considered.

J. Benefit to Risk Ratio

Do the potential benefits of using teprotumumab as recommended outweigh the potential risks associated with the use of the drug product for the intended population?

K. Labeling Recommendations

If the product is approved are there specific recommendations for the labeling?

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2. Product Introduction

Teprotumumab (HZN-001), a fully human monoclonal antibody (mAb), is an insulin-like growth factor-1 receptor (IGF-1R) inhibitor proposed for the treatment of Active Thyroid Eye Disease.

Application Type	BLA
Application Number(s)	761143
Submitted and Received Date	July 8, 2019
Name	(b) (4) (teprotumumab- (b) (4))
Dosage Form(s)	Lyophilized powder for intravenous infusion
Applicant Proposed Dosing Regimen(s)	Intravenous infusion of 10 mg/kg for the initial dose followed by an intravenous infusion of 20 mg/kg every three weeks. The recommended course of therapy is 8 infusions.

Thyroid Eye Disease (TED), also known as thyroid-associated ophthalmopathy, Graves’ ophthalmopathy or Graves’ orbitopathy, is a rare, serious, debilitating and painful autoimmune disease associated with major comorbidities that can lead to blindness. TED is more common in women than men (16 per 100,000 versus 3 per 100,000, respectively), with no significant ethnic predisposition. Median age at diagnosis is 43 years. Risk factors for TED include female gender, middle age, and smoking. The risk of TED increases 7 to 8 times in smokers. In addition, a positive family history of TED is observed in 61% of TED patients.

Patient Experience Data Relevant to this Application

X	The patient experience data that was submitted as part of the application include:	Section where discussed, if applicable
X	Clinical outcome assessment (COA) data, such as	[Study endpoints]
	X Patient reported outcome (PRO)	
	<input type="checkbox"/> Observer reported outcome (ObsRO)	
	X Clinician reported outcome (ClinRO)	
	<input type="checkbox"/> Performance outcome (PerFO)	
	<input type="checkbox"/> Qualitative studies (e.g., individual patient/caregiver interviews, focus group interviews, expert interviews, Delphi Panel, etc.)	
	<input type="checkbox"/> Patient-focused drug development or other stakeholder meeting summary reports	
	<input type="checkbox"/> Observational survey studies designed to capture patient experience data	
	<input type="checkbox"/> Natural history studies	
	<input type="checkbox"/> Patient preference studies (e.g., submitted studies or scientific publications)	
	<input type="checkbox"/> Other: (Please specify)	
<input type="checkbox"/>	Patient experience data that were not submitted in the application, but were considered in this review:	
	<input type="checkbox"/> Input informed from participation in meetings with patient	

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	stakeholders	
	<input type="checkbox"/> Patient-focused drug development or other stakeholder meeting summary reports	
	<input type="checkbox"/> Observational survey studies designed to capture patient experience data	
	<input type="checkbox"/> Other: (Please specify)	
<input type="checkbox"/>	Patient experience data was not submitted as part of this application.	

3. Therapeutic Context

3.1. Analysis of Current Treatment Options

There are no drug or biologic products approved for the proposed indication.

4. Regulatory Background

4.1. U.S. Regulatory Actions and Marketing History

Teprotumumab was originally developed by F. Hoffman-La Roche Ltd., for the treatment of a variety of solid tumors; however, the program was terminated due to lack of efficacy. River Vision Development Corporation initiated a study of teprotumumab for the treatment of diabetic macular edema, but this program was terminated due to difficulty enrolling subjects. River Vision initiated a program in Active Thyroid Eye Disease in June 2013. Horizon Pharma USA, Inc. acquired River Vision and continued the program.

4.2. Summary of Presubmission/Submission Regulatory Activity

Teprotumumab infusion was submitted as IND 112952. Teprotumumab was granted orphan designation for the treatment of Active Thyroid Eye Disease on June 19, 2019 [12-3878/DRU-201203878]. Teprotumumab received Fast Track designation in April 2015, and Breakthrough Therapy designation in July 2016.

4.3. Foreign Regulatory Actions and Marketing History

Each of the teprotumumab studies in Active Thyroid Eye Disease was conducted in the U.S. and Europe (Germany, Italy and the United Kingdom). Of the 171 subjects randomized in the adequate and well-controlled studies of teprotumumab, 56.7% participated at sites located in the U.S. and 43.3% participated at sites in Europe. Teprotumumab is not approved for any indication anywhere in the world.

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5. Significant Issues from Other Review Disciplines Pertinent to Clinical Conclusions on Efficacy and Safety

5.1. Office of Scientific Investigations (OSI)

Routine inspections of clinical investigators and the applicant are ongoing.

5.2. Product Quality

The drug product (DP) is a sterile, preservative-free, lyophilized powder for reconstitution and dilution for infusion, which is presented as a white to off-white powder cake. Each vial delivers 500 mg of teprotumumab formulated in histidine, trehalose and polysorbate 20. The composition of the drug product is provided in Table 1. At time of use, the product is reconstituted with 10 mL of water for injection, which is supplied by the clinical pharmacy, to a final teprotumumab concentration of 50 mg/mL. The reconstituted solution for infusion is a clear to opalescent, nearly colorless to slightly brown liquid and is practically free of visible particles.

Drug Product Composition

Material	Function
Teprotumumab	Active
L-Histidine, USP/Ph. Eur./JP	Buffer
L-Histidine hydrochloride, monohydrate, Ph. Eur.	Buffer
α, α – Trehalose dihydrate, NF/Ph. Eur./JP	Bulking agent, tonicity agent
Polysorbate 20, NF/Ph. Eur./JPE	Surfactant

NF=National Formulary; Ph. Eur.=European Pharmacopeia; USP=United States Pharmacopeia; JP = Japanese Pharmacopeia; JPE = Japanese Pharmaceutical Excipients

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6. Sources of Clinical Data and Review Strategy

Trial Identity	NCT no.	Trial Design	Regimen/ schedule/ route	Treatment Duration/ Follow Up	No. of patients enrolled	Study Population
TED01RV	01868997	Randomized, double-masked, placebo-controlled, parallel-group	Teprotumumab or placebo. 10 mg/kg for first infusion; 20 mg/kg for subsequent Q3W IV infusions	24-week treatment followed by 48 week follow-up	Teprotumumab: 42 Placebo: 45	Active Thyroid Eye Disease
HZNP-TEP-301	03298867	Randomized, double-masked, placebo-controlled, parallel-group	Teprotumumab or placebo. 10 mg/kg for first infusion; 20 mg/kg for subsequent Q3W IV infusions	24-week treatment followed by 48 week follow-up and phone/email at Month 6, 12 and 15	Teprotumumab: 41 Placebo: 42	Active Thyroid Eye Disease
HZNP-TEP-302		Open-label, uncontrolled extension study	24-Week treatment period for non-responders or relapsed subjects	24-week treatment with 6 and 12 months phone/email contact	Ongoing	
DME01RV		Open-label, Phase 1, single arm	Teprotumumab 20 mg/kg Q3W	9 week treatment and 24 week follow-up	5	Diabetic macular edema
BO19373		Open-label, Phase 1, Multiple ascending dose	Teprotumumab manufactured in CHO cell line Teprotumumab manufactured in SP2/0 cell lines	6 infusions	61 SP2/0 36 CHO	Advanced solid tumors, non-Hodgkins and Hodgkins lymphoma
NO21200		Open-label, Phase 1, pediatric (2-17 years) dose finding	Teprotumumab 3 and 9 mg/kg QW or a PK-derived dose (not to exceed 16 mg/kg) 16 mg/kg Q3W or a PK-derived dose (not to exceed 25 mg/kg)		34	Advanced solid tumors
NO21157/SARC011		Open-label, Phase 2, single-arm, 2-stage design for each sub-type cohort	9 mg/kg IV QW 27 mg/kg Q3W (Expanded Ewing's Sarcoma cohort)		317	Recurrent or refractory sarcoma

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Trial Identity	NCT no.	Trial Design	Regimen/ schedule/ route	Treatment Duration/ Follow Up	No. of patients enrolled	Study Population
NO22068		Open label, Phase 1, 12 regimens in combination with different standard chemotherapy therapies, 13 th regimen of monotherapy of R1507 was added in amendment.	Variable regimens		104	Advanced malignancies
NO21160		Placebo controlled Phase 2, in combination with erlotinib	16 mg/kg q3W or 9 mg/kg QW in combination with erlotinib		Teprotumumab: 116 Placebo: 55	NSCLC stage IIIB/IV
NO21746		Open-label, single-arm, Phase 2 in combination with erlotinib	9 mg/kg QW in combination with erlotinib		34	NSCLC stage IIIB/IV
NO21161		Open-label, Phase ½, in combination with letrozole	16 mg/kg q3W in combination with letrozole		6	Postmenopausal with ER+ HER2-advanced letrozole nonresponsive breast cancer
NO21884		Open-label, Phase ½, multiple ascending dose in combination with mTOR inhibitor	16 mg/kg IV q3W in combination with RAD001		11	Advanced solid tumors
NO2202		Open-label, single-arm, single dose, Phase 1 study	16 mg/kg single IV dose		8	Operable breast cancer

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7. Patient Distribution at Investigational Sites:

HZNP-TEP-301 (Review Study #2)		TED01RV (Review Study #1)	
(Site No.)	# Subjects Randomized	(Site No.)	# Subjects Randomized
(US-101)	3	(001)	8
(US-102)	4	(003)	11
(US-103)	9	(004)	10
(US-104)	7	(014)	0
(US-105)	9	(013)	2
(US-106)	1	(015)	1
(US-122)	6	(016)	3
(US-129)	5	(020)	1
(DE-150)	16	(021)	1
(DE-152)	6	(022)	4
(IT-151)	7	(023)	2
(IT-154)	2	(029)	10
(IT-155)	8	Germany (050)	19
		Italy (051)	3
		Italy (054)	3
		United Kingdom (053)	10
		(002)	0
		(005)	0
		(006)	0
		(017)	0
		(018)	0
		(019)	0
		(024)	0
		(025)	0
		(026)	0
		(027)	0
		(031)	0
		Germany (052)	0

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HZNP-TEP-301 (Review Study #2)		TED01RV (Review Study #1)	
(Site No.)	# Subjects Randomized	(Site No.)	# Subjects Randomized
		Germany (055)	0
		Netherlands (056)	0

8. Review of Relevant Individual Trials Used to Support Efficacy

8.1 Review Study #1

A Multicenter, Double-masked, Placebo-controlled, Efficacy and Safety Study of Teprotumumab (HZN-001), an Insulin-like Growth Factor-1 Receptor (IGF-1R) Antagonist Antibody (fully human), administered every 3 weeks (Q3W) by Intravenous (IV) infusion in patients suffering from active Thyroid Eye Disease – TED01RV

8.1.1 Review Study #1 Design

Trial Design- Randomized, double-masked, placebo-controlled, parallel-group

Plan-

1) A Screening Phase of 4 weeks (\pm 2 weeks) with no treatment. Subjects attended the clinic once or twice, or as required, during the screening period.

2) A double-masked Treatment Phase of 24 weeks. Subjects attended clinic visits at Week 0 (baseline visit, 1st infusion), Weeks 1 and 3 (2nd infusion), 4 and 6 (3rd infusion), 9 (4th infusion), 12 (5th infusion), 15 (6th infusion), 18 (7th infusion), 21 (8th infusion), and 24 (final assessment visit). Research staff telephoned subjects focusing on safety and tolerability aspects the day after infusion for the 1st and 2nd infusions, and thereafter as required. Research staff also contacted subjects who experienced an infusion-related event the day after the infusion.

3) A Follow-up Phase of 48 weeks with no additional treatment during at least the first 12 weeks. Subjects attended clinic visits at Week 28, 36, 48, 60, and 72.

Eligible subjects who met study entry criteria were randomly assigned to the double-masked treatment phase in a 1:1 ratio to receive a starting dose of 10 mg/kg of HZN-001 or placebo once every 3 weeks (q3W) by IV. At Week 3, the dose was escalated to 20 mg/kg IV q3W. Following dose escalation, subjects continued at this dose level for all subsequent infusions.

In the case of an intolerable adverse event (AE), subjects were to be withdrawn from the study. The active treatment phase of the study was 24 weeks (8 infusions) in duration. Randomization was stratified by smoking status. During the treatment period, subjects were evaluated at clinic visits every 3 weeks and, if appropriate, by telephone contact by research staff. Measurements

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for efficacy, tolerability, safety, biomarkers, and pharmacokinetics were performed according to the assessment schedule.

Subjects were to be withdrawn from the study if they developed optic neuropathy or any condition that required surgical intervention. An independent data and safety monitoring board (DSMB) was chartered to review safety data on a regular basis; the DSMB was masked to efficacy data.

Key Inclusion Criteria:

1. Age 18 to 75 years.
2. Clinical diagnosis of Grave's disease associated with active TED with a clinical activity score of ≥ 4 for the most severely affected eye.
3. Fewer than 9 months from onset of TED.
4. Euthyroid or with mild hypo- or hyperthyroidism defined as free thyroxine and free triiodothyronine levels $< 50\%$ above or below the normal limits.

Key Exclusion Criteria:

1. Decreased best corrected visual acuity due to optic neuropathy as defined by a decrease in vision within the last 6 months of 2 lines, new visual field defect or color defect.
2. Improvement in CAS of ≥ 2 points between screening and baseline.
3. Treatment with oral or IV steroids within the previous 3 month (except doses less than 1g methylprednisolone or equivalent)
4. Previous orbital irradiation.
5. Poorly controlled diabetes.

Clinical Activity Score (CAS)

Subjects were assessed at screening and predose at Visit 1 (Week 0, baseline assessment) and at Weeks 6, 12, 18, 24, 28, and 72 using the 7-item European Group on Graves' Ophthalmopathy (EUGOGO) amended CAS [Mourits et al. 1989; Mourits et al. 1997]. The 7-point CAS scale is comprised of 2 patient-reported outcomes and 5 clinician-reported outcomes. For each item present, 1 point was given. The sum of these points was the total score.

1. Spontaneous orbital pain.
2. Gaze-evoked orbital pain.
3. Eyelid swelling that was considered to be due to active (inflammatory phase) Graves Ophthalmopathy.
4. Eyelid erythema.
5. Conjunctival redness that was considered to be due to active (inflammatory phase) Graves Ophthalmopathy (ignore "equivocal" redness).
6. Chemosis.
7. Inflammation of caruncle or plica.

Reviewer's Comments: *The Agency disagreed with the construction of the CAS score. The CAS*

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score is a composite with equal weighting of a number of factors. However, FDA’s clinical team does not consider these factors to be of equal clinical weight either to the patients or to physicians treating these patients.

Proptosis

For assessment of proptosis, the same Hertel instrument and intercanthal distance was to be used at each time point. Every effort was made for the same observer to conduct the assessment on each occasion. The Hertel values were measured for each eye at all time points.

Graves’ Ophthalmopathy Quality of Life Scale

The Graves’ Ophthalmopathy Quality of Life (GO-QOL) scale [Terwee 1998] was completed at screening, Weeks 6, 12, 24, 28, 48, 72, and early withdrawal. The GO-QOL is a 16-item self-administered questionnaire used to assess the perceived effects of TED by subjects on their daily physical and psychosocial functioning.

Clinical Measures of Severity Score (CSS)

CSS Item and Assessment Scale	Minimum change required for classifying overall response
Lid aperture (distance between the lid margins (mm) with the subject looking in the primary position sitting relaxed and with distant fixation)	Decrease ≥ 2 mm
Swelling of the eyelids (absent, mild, moderate, or severe)	Decrease ≥ 1 grade
Redness of the eyelids (absent, present)	Decrease ≥ 1 grade
Redness of the conjunctiva (absent, present)	Decrease ≥ 1 grade
Conjunctival edema (absent, present)	Decrease ≥ 1 grade
Inflammation of the caruncle or plica (absent, present)	Decrease ≥ 1 grade
Exophthalmos (measured in mm using the same Hertel exophthalmometer and same intercanthal distance for each individual subject)	Decrease ≥ 2 mm
Subjective diplopia score (0=no diplopia; 1=intermittent, i.e., diplopia in primary position of gaze, when tired or when first awakening; 2=inconstant, i.e., diplopia at extremes of gaze; 3=constant, i.e., continuous diplopia in primary or reading position)	Decrease ≥ 1 grade
Eye muscle involvement (ductions in degrees)	Increase $\geq 8^\circ$ in at least one direction of gaze
Corneal involvement (absent/punctate keratopathy/ulcer)	Decrease ≥ 1 grade
Optic nerve involvement (best corrected visual acuity, color vision, optic disc, relative afferent pupillary defect (absent, present), plus visual fields if optic nerve compression is suspected.	Change of best corrected visual acuity by ≥ 2 lines on Snellen chart, or substantial color vision change, or significant change of visual fields, or significant change in optic disc appearance, or (Dis-)appearance of relative afferent pupillary defect

Ophthalmic Examination

The ophthalmic examination included best-corrected visual acuity, pupil examination and color vision assessment, Ishihara color plates (or equivalent) or related red desaturation, intraocular pressure, and slit lamp examination. If significant abnormalities were noted compared with previous visits, including a loss of 2 lines or more of vision, development of pupil abnormalities including afferent pupillary defect, rise in intraocular pressure, development of corneal infiltrates, or other abnormalities that were of concern to the ophthalmologist, further investigations of visual function were conducted according to the ophthalmologist's decision.

Clinical Laboratory Evaluations

The following laboratory tests were performed at screening, baseline (Week 0), Weeks 3, 6, 9, 12, 18, 24, 36, 72, and early withdrawal. All sampling was performed prior to dosing (at infusion visits). Results for blood glucose and platelets from the previous visit were evaluated prior to dosing.

- Hematology: hemoglobin, platelet count, white blood cells and differential (also at Weeks 1, 4, 15, and 21)
- Renal Function: serum creatinine and blood urea nitrogen
- Hepatic function: total bilirubin, alkaline phosphatase, lactate dehydrogenase, gamma-glutamyl transferase, SGOT/AST, and alanine aminotransferase SGPT/ALT
- Electrolytes: sodium, potassium, chloride, bicarbonate, calcium, and phosphate
- Blood glucose: fasting at Weeks 1 and 4 and non-fasting at other study time points
- Hemoglobin A1c: at screening and Weeks 12, 24, 36, and 72
- Thyroid function tests: FT4, FT3 and TSH (TSH Germany only – Site 50)
- Human anti-human antibodies (HAHA): serum sample prior to dosing at Weeks 0, 3, and 9, and at Weeks 24, 36, and 72. Analysis of HAHA was performed only when all subjects had completed the masked treatment phase of the study. Any subject with treatment emergent positive HAHA titer, which was still present at Week 72, was followed. This included subjects with HAHA detected post-dose and those with positive baseline HAHA if there was an important increase in titer post-dose.
- Serum pregnancy test (for women of childbearing potential) at screening only. Urine pregnancy test at all other time points (Weeks 0, 3, 6, 9, 12, 15, 18, 21, 24, 28, 36, and early withdrawal)
- Complete urinalysis (including specific gravity, protein, blood, ketones, glucose, etc.) Hematology and blood glucose were performed for all subjects at Weeks 1, 4, 15, and 21. Subjects were fasting at Weeks 1 and 4.

Biomarker Assessments

Blood for plasma and serum was collected at baseline (Week 0) and at Weeks 12, 24, 28 and 72. Samples used for biomarker assays included thyroid stimulating immunoglobulin (TSI) and TBII; anti-thyroid peroxidase (TPO) and anti-thyroglobulin antibodies; T and B cell and fibrocyte flow cytometry for IGF-1R (Weeks 0 and 28 only) and TSHR levels; serum IL-6, IL-16, and RANTES

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(Weeks 0, 12, 24, and 72 only).

Treatment

HZN-001 or placebo was administered q3W by IV infusion over a period of 24 weeks for a total of 8 infusions. All subjects were started at a dose of 10 mg/kg. At Week 3, the dose was escalated to 20 mg/kg and remained there for the rest of the study. The first and second infusions were administered over 90 minutes. Subsequent infusions were administered over a 60-minute period, providing there were no significant infusion-related events. Subjects with intolerable AEs were to be withdrawn from the study.

Review Study #1 Primary Efficacy Endpoint

The primary efficacy endpoint was whether the subject was a responder or not (yes or no) at Week 24. A responder was defined as a subject with the following:

- A decrease in overall CAS ≥ 2 points AND
- A reduction in proptosis ≥ 2 mm, AND
- No deterioration of CAS in the Non-Study Eye (i.e., increase of CAS ≥ 2 points OR increase in proptosis ≥ 2 mm) at the 24-week evaluation.

Reviewer's Comments: *The Agency disagreed with the inclusion of the CAS score in the primary endpoint. The CAS score is a composite with equal weighting of a number of factors. However, FDA's clinical team does not consider these factors to be of equal clinical weight either to the patients or to physicians treating these patients. The primary hallmark of patient symptoms and concerns is proptosis, and therefore the Agency considered proptosis to be the primary endpoint.*

Review Study #1 Follow-up

To evaluate subjects off-treatment following the masked 24-Week Treatment Phase with teprotumumab or placebo in subjects diagnosed with active TED. The off-treatment Follow-up Period was designed to assess safety, including evaluation of the possibility of an acute disease activity rebound effect off treatment, during the 30-day Follow-up Period (Week 24 to Week 28), and evaluate continued short-term response of teprotumumab after 4 weeks of treatment discontinuation at Week 28. Continued teprotumumab treatment effect as well as worsening of disease requiring additional treatment was evaluated in the long-term follow-up at Week 72.

The off-treatment follow-up period was for 48 weeks, with no additional treatment for TED during at least the first 12 weeks, unless medically indicated. Subjects attended clinic visits at Weeks 28, 36, 48, 60, and 72 for safety assessments; efficacy was only measured at Weeks 28 and 72.

For responder/non-responder analyses, any subject who received additional treatment for thyroid eye disease (TED) was considered a non-responder from the time of TED treatment forward. A proptosis relapse/non-relapse analysis was performed for Weeks 28 and 72 based on if a subject was a proptosis responder at Week 24 (relapse was defined as an increase in proptosis of ≥ 2 mm from Week 24).

(b) (4) (teprotumumab- (b) (4))

8.1.2 Review Study #1 Results

Patient Disposition

There were 88 subjects enrolled in the study. Of these, 87 subjects took at least 1 dose of study drug and were included in the ITT and mITT Populations.

	Placebo (N=45) n (%)	HZN-001 (N=43) n (%)
Enrolled (Informed Consent Signed)	45	43
ITT Population	45 (100%)	42 (98%)
MITT Population	45 (100%)	42 (98%)
PP Population	36 (80%)	33 (77%)
Safety Population	44 (98%)	43 (100%)
Completed the Study Treatment Reason for Early Termination	39 (87%)	37 (86%)
Adverse Event	1 (2%)	5 (12%)
Lack of Efficacy	2 (4%)	0
Pregnancy	0	0
Protocol Violation	0	0
Study Terminated by Sponsor	0	0
Death	0	0
Other – see Note below	3 (7%)	1 (2%)

Abbreviations: ITT = intent to treat, MITT=modified intent to treat, PP=per protocol.

Note: All subjects who signed informed consent were considered enrolled in the study. The percentages presented in this table are based on the ITT Population. Three subjects received the wrong treatment; these 3 subjects were excluded from the PP Population and analyzed under the first treatment actually received for the Safety Population. One subject randomized to HZN-001, terminated early, and never received any study drug.

Demographic and Baseline Characteristics (Safety Population)

Review Study #1	Placebo (N=44)	HZN-001 (N=43)
Age (years) Mean (SD)	54.2 (13.0)	51.6 (10.7)
Median	55.4	50.5
(Min, Max)	(20.4, 77.0)	(22.3, 72.6)
<65 years old	36 (80%)	39 (91%)
≥65 years old	9 (20%)	4 (9%)
Gender, n (%)		
Female	36 (82%)	28 (65%)
Ethnicity, n (%)		
Hispanic or Latino	4 (9%)	2 (5%)
Not Hispanic or Latino	40 (91%)	41 (95%)
Race, n (%)		
American Indian or Alaska Native	0	0
Asian	2 (5%)	1 (2%)
Black or African–American	4 (9%)	4 (9%)
Native Hawaiian or Other Pacific Islander	0	1 (2.3)
White	38 (86%)	37 (86%)
Weight (kg), N	44	43
Mean (SD)	78.8 (16.88)	82.5 (23.73)
Median	73.3	75.0
(Min, Max)	(53.6, 122.0)	(47.6, 168.7)
Study Eye, n(%)		
Right Eye	20 (45%)	27 (63%)
Smoking Status, n(%)		
Smoker	18 (41%)	11 (26%)
CAS Score in Study Eye at Baseline n (%)		
0 or 1	0	0
2	0	1 (2%)
3	0	0
4	6 (14%)	10 (23%)
5	23 (52%)	18 (42%)
6	13 (30%)	12 (28%)
7	2 (5%)	2 (5%)
Exophthalmos (mm), N	44	43
Mean (SD)	22.91 (2.67)	23.57 (3.36)
Median	22.5	23.0
Min, Max	(16.0, 29.0)	(17.0, 33.0)

Abbreviation: CAS=Clinical Activity Score; CSS = Clinical Measures of Severity Score; ITT=intent-to-treat; N = number; SD =standard deviation

^a Baseline was the last predose measurement.

(b) (4) (teprotumumab- (b) (4))

Efficacy Results – Review Study #1 Primary Endpoint

Responder (CAS+Proptosis)

	Placebo	Teprotumumab	Difference (95% conf)	p-value
Week 6 Study Eye	2/42 (5%)	18/39 (46%)	41% (24,58)	<0.001
Week 12 Study Eye	2/41 (5%)	23/40 (58%)	53% (36,69)	<0.001
Week 18 Study Eye	2/41 (5%)	30/39 (77%)	72% (57,87)	<0.001
Week 24 Study Eye	9/39 (23%)	29/38 (76%)	53% (34,72)	<0.001
Week 28 Study Eye*	6	31		
Week 6 Non-Study Eye	2/42 (5%)	8/39 (21%)	16% (2,30)	0.031
Week 12 Non-Study Eye	1/41 (2%)	13/40 (33%)	30% (15,45)	<0.001
Week 18 Non-Study Eye	1/41 (2%)	17/39 (44%)	41% (25,57)	<0.001
Week 24 Non-study Eye	6/39 (13%)	22/38 (58%)	42% (23,62)	<0.001
Week 28 Non-study Eye*	5	20		

p-value based on chi square

* Off treatment for 4 weeks

Reviewer's Comments: *The Agency disagreed with the primary endpoint due to the inclusion of the CAS portion of the endpoint. The CAS assigns equal weight to a number of components which have different clinical value to both patients and clinicians. The Agency requested an endpoint which included only Proptosis.*

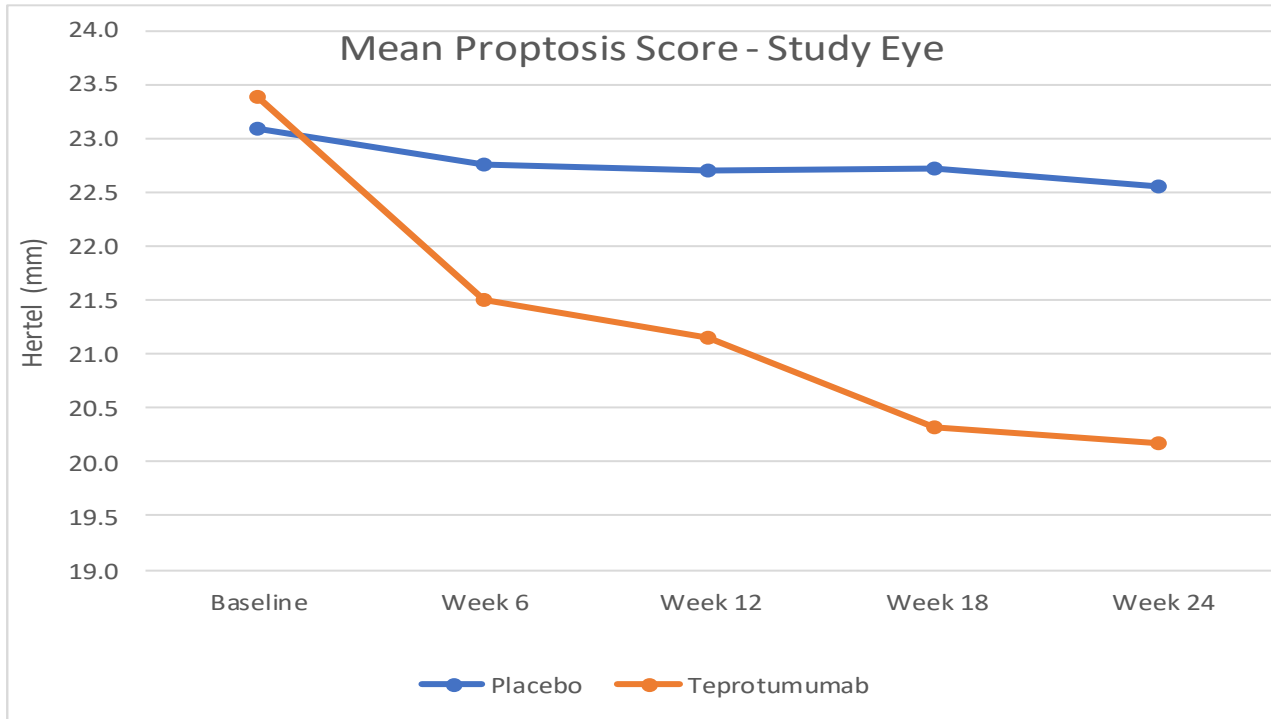
Agency's Requested – Primary Endpoint: % patients with 2 mm or more decrease in Proptosis

Proptosis	Placebo	Teprotumumab	Difference (95% conf)	p-value
Week 6 Study Eye	4/42 (10%)	22/40 (55%)	45%	<0.001
Week 12 Study Eye	2/41 (5%)	24/40 (60%)	55%	<0.001
Week 18 Study Eye	4/41 (10%)	32/39 (82%)	72%	<0.001
Week 24 Study Eye	9/39 (23%)	30/38 (79%)	56%	<0.001
Week 28 Study Eye*	6	31		
Week 6 Non-study Eye	3/42 (7%)	9/40 (23%)	16%	<0.001
Week 12 Non-study Eye	3/41 (7%)	15/40 (38%)	31%	<0.001
Week 18 Non-study Eye	4/41 (10%)	21/39 (54%)	44%	<0.001
Week 24 Non-study Eye	6/39 (15%)	26/38 (68%)	53%	<0.001

* Off treatment for 4 weeks

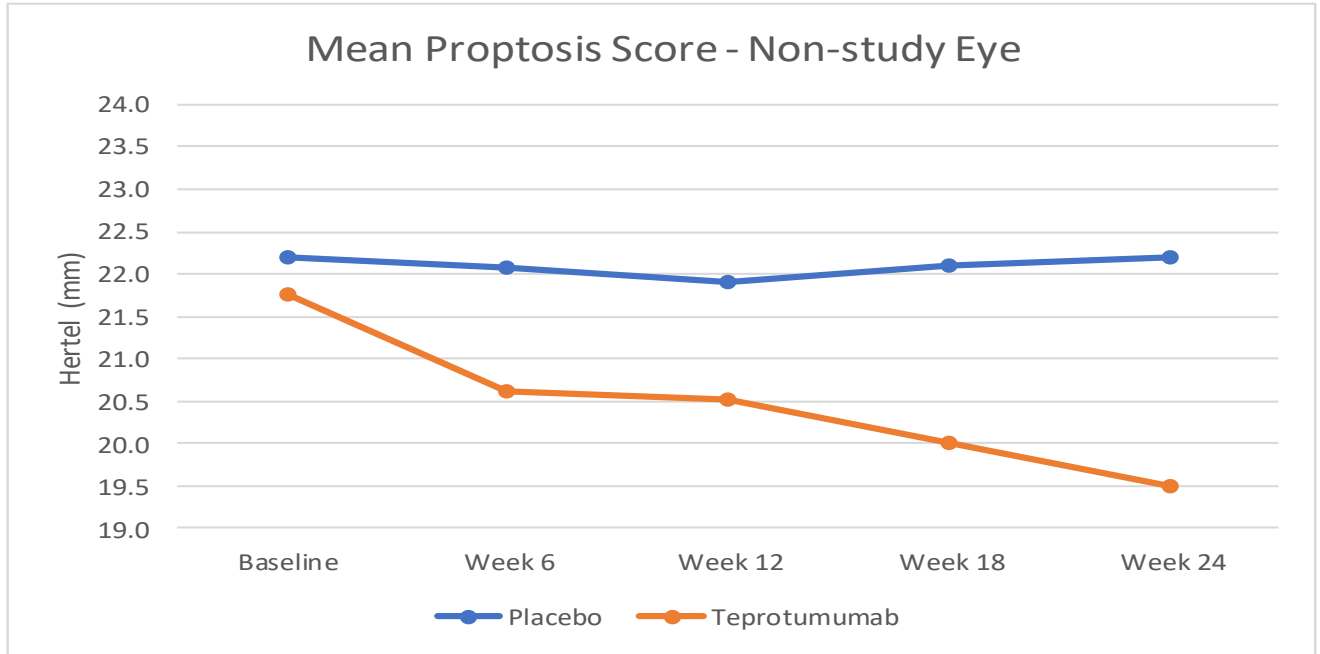
Reviewer's Comments: *By the first evaluation period at Week 6, there is a clinically significant reduction in proptosis (i.e., greater than 2 mm) in both eyes which continues through the treatment period.*

Review Study #1



Reviewer's Comments: *By the first evaluation period at Week 6, there is a reduction in proptosis which continues through the treatment period.*

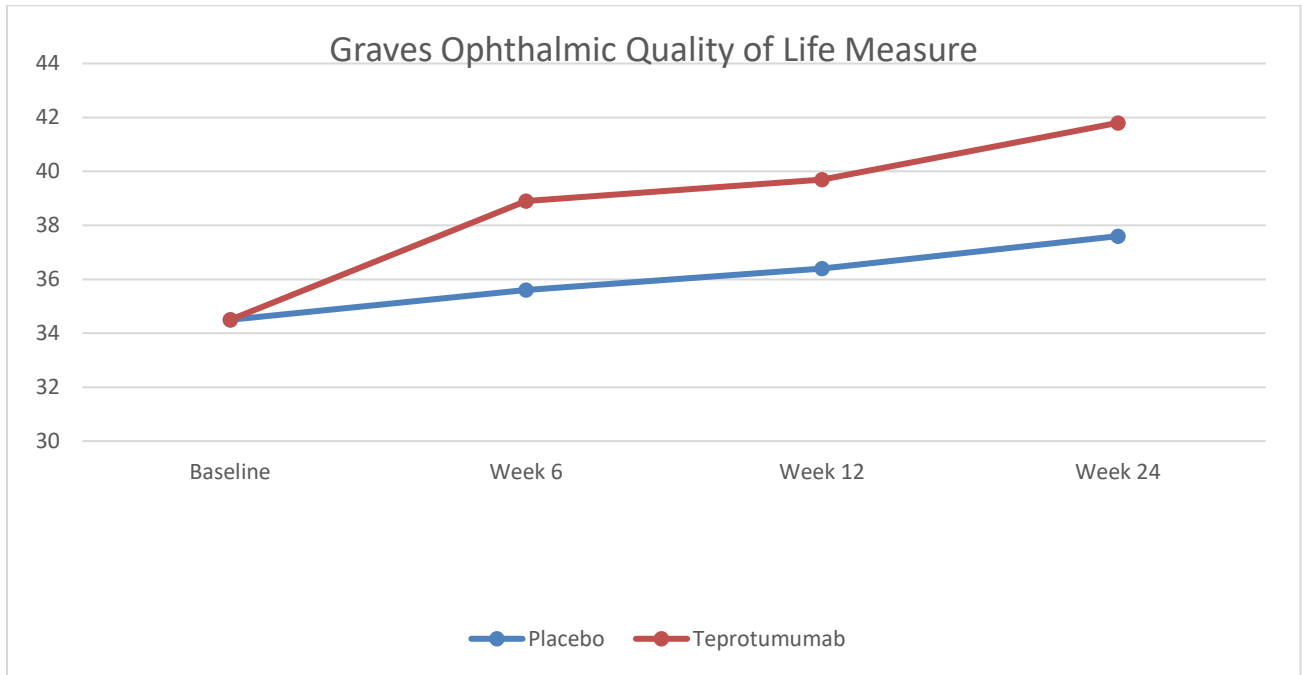
Review Study #1



Reviewer's Comments: *By the first evaluation period at Week 6, there is a reduction in proptosis in both eyes which continues through the treatment period.*

(b) (4) (teprotumumab- (b) (4))

Review Study #1: Secondary Endpoints

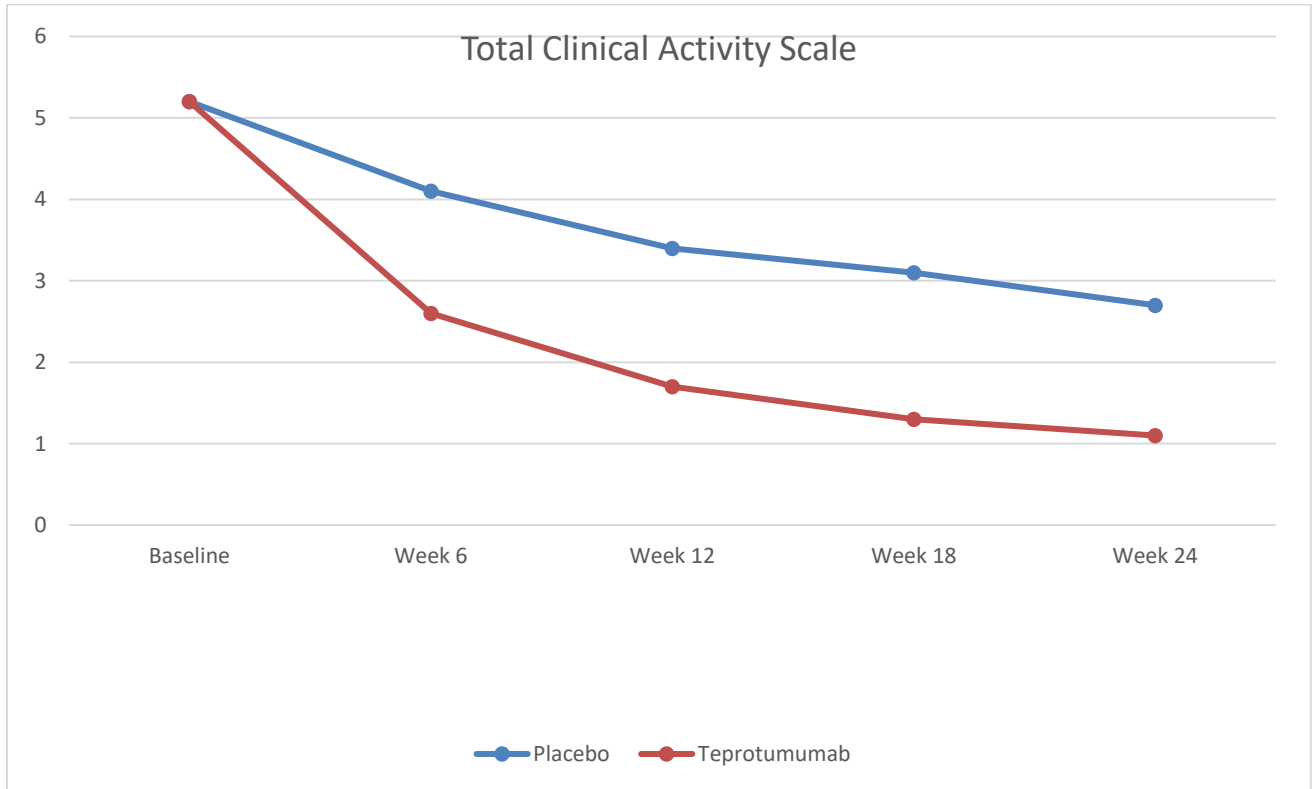


Grave’s Ophthal Quality of Life Score	Placebo	Teprotumumab	Difference
Baseline	34.5 (6.8)	34.5 (7.4)	0
Week 6	35.6 (6.3)	38.9 (7.2)	3.3
Week 12	36.4 (6.9)	39.7 (6.2)	3.3
Week 24	37.6 (6.9)	41.8 (6.4)	4.2

Reviewer's Comments: *Validation information for the Grave’s Ophthalmology Quality of Life Score has not been submitted and therefore interpretation of the scores is not possible.*

(b) (4) (teprotumumab- (b) (4))

Review Study #1

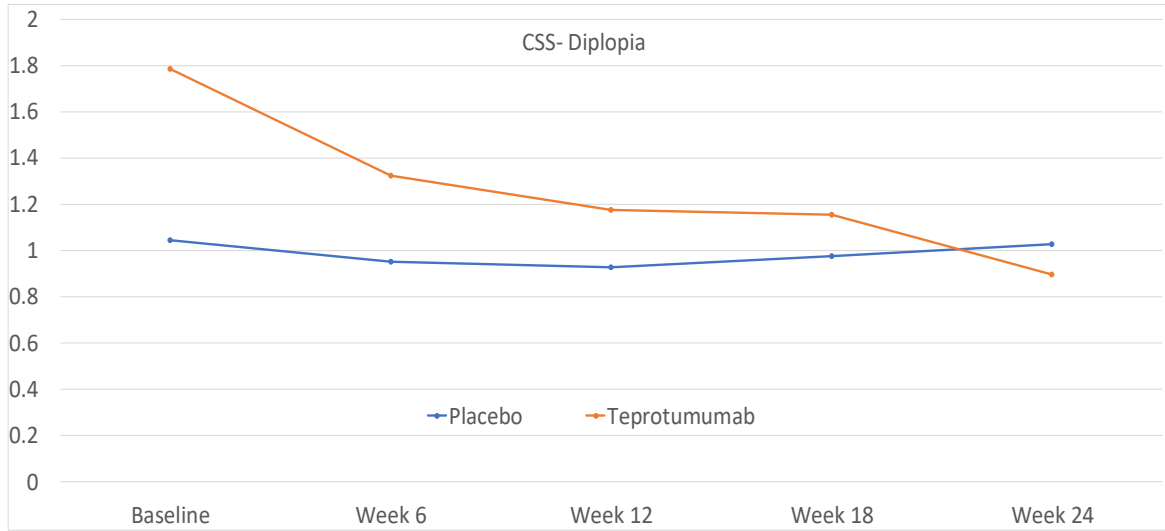


CAS Change from Baseline	Placebo	Teprotumumab	Difference
Baseline	5.2	5.1	0.1
Week 6	-1.1	-2.5	-1.5
Week 12	-1.8	-3.4	-1.6
Week 18	-2.1	-3.8	-1.7
Week 24	-2.5	-4.0	-1.6

Reviewer's Comments: *Clinical Activity Scale is not accepted because there is not necessarily equal weight for each component.*

(b) (4) (teprotumumab- (b) (4))

Review Study #1 Subjective Diplopia measured on CSS Score (ITT Population)



	Placebo (N=45)	Teprotumumab (N=42)
Baseline		
0, no diplopia	14 (31.1)	4 (9.5)
1, intermittent	19 (42.2)	16 (38.1)
2, inconstant	8 (17.8)	7 (16.7)
3, constant	4 (8.9)	15 (35.7)
Week 6, n (%)		
0, no diplopia	18 (40.0)	13 (31.0)
1, intermittent	12 (26.7)	11 (26.2)
2, inconstant	8 (17.8)	6 (14.3)
3, constant	4 (8.9)	10 (23.8)
CSS responder, n (%)	9 (21.4)	17 (42.5)
Week 12, n (%)		
0, no diplopia	20 (44.4)	16 (38.1)
1, intermittent	9 (20.0)	8 (19.0)
2, inconstant	7 (15.6)	9 (21.4)
3, constant	5 (11.1)	7 (16.7)
CSS responder, n (%)	10 (24.4)	24 (60.0)
Week 18, n (%)		
0, no diplopia	19 (42.2)	16 (38.1)
1, intermittent	9 (20.0)	8 (19.0)
2, inconstant	8 (17.8)	8 (19.0)
3, constant	5 (11.1)	7 (16.7)
CSS responder, n (%)	11 (26.8)	23 (59.0)
Week 24, n (%)		
0, no diplopia	18 (40.0)	21 (50.0)
1, intermittent	8 (17.8)	4 (9.5)
2, inconstant	7 (15.6)	9 (21.4)
3, constant	6 (13.3)	4 (9.5)
CSS responder, n (%)	10 (25.6)	26 (68.4)

Reviewer's Comments: *As a single question about diplopia, the response demonstrates an improvement in diplopia.*

(b) (4) (teprotumumab- (b) (4))

8.1.3 Subject Disposition from Week 24 to Week 72 (All Subjects)

	Placebo, N=45 n (%)	Teprotumumab, N=42 n (%)
ITT Population	45	42
Completed Study Treatment (Week 24)	39	37
Completed Extended Follow-up (Week 72)	38	36
Reason for Extended Follow-up Termination		
Adverse Event	2	5
Lack of Efficacy	2	0
Other ^a	3	1

Source: Table 14.1.1 Abbreviations: ITT = intent-to-treat; TED = thyroid eye disease

- a. Placebo group: left eye optic disc edema; incorrect treatment given; and subject decision to withdraw.
Teprotumumab group: elective TED surgery

Subjects who received additional TED treatment during the off-treatment follow-up period

Site	Group	Study Week	Corticosteroids	Rituximab	Orbital Decompression
001	Placebo	26	Yes		Yes
050	Placebo	25	Yes		
001	Placebo	28	Yes		Yes
001	Placebo	29	Yes	Yes	Yes
013	Placebo	60			Yes
022	Placebo	57			Yes
001 ^a	Teprotumumab	47	Yes		
001 ^a	Teprotumumab	50	Yes		
050 ^a	Teprotumumab	62	Yes		Yes
004 ^a	Teprotumumab	70			Yes ^b

Source: Listing 16.2.3.7.3

- a. Responder at Week 24
b. Elective TED surgery at Week 70; no proptosis data collected at Week 72

Proportions of Proptosis Responders Who Relapsed (≥ 2 mm) from Week 24 through Week 72

	Placebo	Teprotumumab
Week 28, n		
Relapse ^a	1 (11%)	0
No Relapse	8 (89%)	29 (97%)
Missing ^b	0	1 (3%)
Week 72, n		
Relapse ^a	3 (33%)	11 (37%)
No Relapse	6 (67%)	18 (60%)
Missing ^c	0	1 (3%)

Source: [Table 14.2.2.2.6](#)

Includes Week 24 proptosis responders. Subjects who received TED treatment in the off-treatment Follow-up Period were treated as relapsed from the time of TED treatment forward.

- a. Relapse was defined as an increase in proptosis of ≥ 2 mm from Week 24 in the Study Eye only.
- b. One Subject, Week 28 proptosis value is missing.
- c. One Subject had elective TED surgery at Week 70; therefore, the Week 72 proptosis value is missing.

Reviewer's Comments: *Relapses occur in the follow-up period after treatment. However, approximately 60% of patients did not relapse in the year following treatment.*

(b) (4) (teprotumumab- (b) (4))

8.2 Review Study #2

A Phase 3, Randomized, Double-Masked, Placebo-Controlled, Parallel-Group, Multicenter Study Evaluating Teprotumumab (HZN-001) Treatment in Subjects with Active Thyroid Eye Disease

Short title: Treatment of Graves' Orbitopathy to Reduce Proptosis with Teprotumumab Infusions in a Randomized, Placebo-Controlled, Clinical Study (OPTIC)

8.2.1 Review Study #2 Study Design

Trial Design- Same as Review Study #1

Plan- Same as Review Study #1

Key Inclusion Criteria: Same as Review Study #1 except maximum age of 80.

Key Exclusion Criteria: Same as Review Study #1

Clinical Activity Score (CAS)- Same as Review Study #1

Proptosis- Same as Review Study #1

Graves' Ophthalmopathy Quality of Life Scale- Same as Review Study #1

Clinical Measures of Severity Score (CSS)- Same as Review Study #1

Treatment- Same as Review Study #1

Ophthalmic Examination

The ophthalmic examination included pupil examination, color vision assessment, intraocular pressure, and slit lamp examination. If significant abnormalities were noted compared with previous visits, including a loss of 2 lines or more of vision.

Clinical Laboratory Evaluations- see Schedule of Events Table

Criteria for Responders Who Relapse

Subjects who met the response criteria at Week 24, but subsequently experience a disease relapse

during the 48-week Follow-up Period will have the option to enter the open-label extension study

(HZNP-TEP-302) and receive 8 infusions of teprotumumab. Determination of relapse is based on the following criteria:

- Increase in proptosis of ≥ 2 mm in the study eye since Week 24, or
- An increase in CAS of ≥ 2 points since Week 24 with an absolute CAS of ≥ 4 in the study eye following the Week 24 Visit.
- In addition to one of the bullet points above, the Investigator should consider the subject's symptomology to ensure a relapse has occurred (e.g., new onset of double vision).

Review Study #2 Schedule of Events: Screening, Treatment and Follow-up Periods

Study Visit	Screening ¹	Treatment Period ²										Follow-Up Period ³						Follow-Up Contact ⁴	
	S1/S2/ S3	1	2	3	4	5	6	7	8	9	10	11/ PW1 ⁵	12	13	14	15	16/ PW2 ⁶	17	18
Week (W)/Month(M)	-42 to -14 days	Day 1 ⁷	W1	W3	W4/M 1	W6	W9	W12/ M3	W15	W18	W21	W24/ M6	W28/ M7	W36/ M9	W48/ M12	W60/ M15	W72/ M18	W96/ M24	W120/ M30
Visit Window (± days)		(±3)	(±1)	(±3)	(±1)	(±3)	(±3)	(±3)	(±3)	(±3)	(±3)	(±7)	(±7)	(±7)	(±7)	(±7)	(±7)	(±14)	(±14)
Informed consent	X																		
Review inc/exc criteria	X	X																	
Demographics	X																		
Medical history 8	X ⁹	X																	
Weight ¹⁰	X							X				X		X	X	X	X		
Randomization ¹¹		X ⁷																	
Study drug infusion		X		X		X	X	X	X	X	X								
Phone (email) contact for		X		X															
Safety 24 hours postdose ¹²																			
Efficacy assessments																			
Clinical Activity Score ¹³	X	X ¹⁴				X		X		X		X	X	X	X	X	X		
Clinical Measures of Severity - includes proptosis, diplopia and motility restriction	X	X ¹⁵				X		X		X		X	X	X	X	X	X		
Pregnancy test ¹⁶	X	X		X		X	X	X	X	X	X	X	X	X	X		X ¹⁷		
Physical examination ¹⁸	X ¹⁹	X ¹⁸	X			X		X		X		X ¹⁸			X		X ¹⁸		
Ophthalmic examination ²⁰	X ²¹	X	X			X		X		X		X			X		X		
Vital signs ²²	X	X ²²	X	X ²²	X	X	X	X	X	X	X	X	X	X	X	X	X	X	
12-lead ECG	X	X		X		X		X				X						X	
Clinical laboratory tests ²³																			
Chemistry (excl. glucose)	X ²⁴	X		X		X	X	X		X		X		X			X		
Thyroid (F3, FT4,TSH) ²⁵	X	X		X		X	X	X		X		X		X			X		
Hematology	X	X	X		X	X	X	X	X	X	X	X		X			X		
Glucose ²³	X	X	X	X	X	X	X	X	X	X	X	X		X			X		
HbA1c ²⁶	X							X				X		X			X		
Urinalysis	X	X		X		X	X	X		X		X		X			X		
ADA/NAb samples ²⁷		X		X			X					X ²⁸		X			X		
AE/ SAE assessment ²⁹	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X	
Concomitant medications	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X	
GO-QoL Questionnaire		X				X		X				X	X		X		X		
Biomarker samples ³⁰		X						X				X							

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(b) (4) (teprotumumab- (b) (4))

Study Visit	Screening ¹ S1/S2/ S3	Treatment Period ²										Follow-Up Period ³						Follow-Up Contact ⁴	
		1	2	3	4	5	6	7	8	9	10	11/ PW1 ⁵	12	13	14	15	16/ PW2 ⁶	17	18
Week (W)/Month(M)	-42 to -14 days	Day 1 ⁷	W1	W3	W4/M 1	W6	W9	W12/ M3	W15	W18	W21	W24/ M6	W28/ M7	W36/ M9	W48/ M12	W60/ M15	W72/ M18	W96/ M24	W120/ M30
Visit Window (± days)		(±3)	(±1)	(±3)	(±1)	(±3)	(±3)	(±3)	(±3)	(±3)	(±3)	(±7)	(±7)	(±7)	(±7)	(±7)	(±7)	(±14)	(±14)
Pharmacokinetic samples ³¹		X	X	X	X		X					X ²⁸							
Contact (phone/ email) to assess additional TED Treatment ³²																		X	X

ADA = anti-drug antibody; AE = adverse event; CAS = Clinical Activity Score; ECG = electrocardiogram; exc = exclusion; FT3 = free triiodothyronine; FT4 = free thyroxine; GO-QoL = Graves’ Ophthalmopathy Quality of Life; HbA1c = glycated hemoglobin; IL = interleukin; inc = inclusion; INFγ = interferon gamma; micRNA = microribonucleic acid; NAb = neutralizing antibody; PW = premature withdrawal; q3W = every 3 weeks; S = Screening; SAE = serious adverse event; sIL-1RA = secretory interleukin-1 receptor antagonist; TED = thyroid eye disease; TGFβ = transforming growth factor beta; TNFα = tumor necrosis factor alpha; TSH = thyroid-stimulating hormone; TSH-R-Ab = thyroid-stimulating-hormone-receptor stimulating, blocking and binding antibody; ULN = upper limit of normal.

- Screening procedures could have taken place over more than 1 day/clinic visit provided consent was obtained first and all assessments were completed within the designated window.
- Double-Masked Treatment Period. Subjects who were proptosis_non-responders at Week 24 were eligible to enroll in an open-label extension study in which all subjects receive teprotumumab 20 mg/kg (10 mg/kg for the first infusion and 20 mg/kg for the remaining 7 infusions).
- Proptosis responders and non-responders who chose not to enroll in the open-label extension study participated in a Follow-Up Period.
- Subjects who completed the Week 72 Visit were contacted via phone or email by research staff to enquire if any treatment for TED had been received since last study contact.
- If a subject prematurely discontinued study drug during the Double-Masked Treatment Period, they returned for a clinic visit and underwent the Week 24 assessments, with the exception of the collection of blood samples for pharmacokinetic and ADA evaluations. Subjects were encouraged to continue study participation in the Follow-Up Period.
- If a subject prematurely discontinued from the study during the Follow-Up Period, they returned for a clinic visit and underwent the Week 72 assessments prior to discharge.
- On Day 1 (Baseline), subjects were randomized and received the first dose of study drug; however, Baseline assessments were performed prior to dosing.
- Medical history included tobacco use history and Graves’ disease and treatment history.
- TED must have been moderate to severe in intensity (non-sight threatening but appreciable impact on daily life) with an onset of symptoms (as determined by subject records) within 9 months prior to the Baseline Visit for study enrollment.
- Dosing was adjusted if there was a change in weight during the Double-Masked Treatment Period. The weight obtained at Week 12 could have been used in dose calculations beginning at Week 12 or Week 15.
- Subjects were randomized in a 1:1 ratio (stratified by tobacco use status) to receive either: a) teprotumumab (10 mg/kg on Day 1 followed by 20 mg/kg q3W for the remaining 7 infusions) or b) placebo (q3W for all 8 infusions).
- Phone (or email) contact by research staff focusing on safety and tolerability aspects was made the day after infusion for the first and second infusions, and thereafter as deemed appropriate. In addition, subjects who experienced an infusion-associated event after any subsequent infusion were also contacted by phone (or email) by research staff the day after the infusion, and thereafter as deemed appropriate.
- CAS must have been ≥4 for enrollment and randomization.
- Subjects whose CAS in the study eye decreased 2 or more points from Screening were not eligible for randomization.
- Subjects who had a ≥2 mm decrease in proptosis in the study eye from Screening were not eligible for randomization.
- Serum pregnancy test at Screening and urine pregnancy tests prior to dosing at all other visits, as applicable. Performed for female subjects of childbearing potential (including those with an onset of menopause <2 years prior to Screening, non-therapy-induced amenorrhea for <12 months prior to Screening, or not surgically sterile [absence of ovaries and/or uterus]).
- Pregnancy test only performed for female subjects of childbearing potential who entered the Follow-Up Period but discontinued study participation prior to Week 48.

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(b) (4) (teprotumumab)- (b) (4)

18. Physical examination included assessment of presence or absence of pretibial myxedema on Day 1 and Week 24 (or PW) of the Double-Masked Treatment Period and Week 72 (or PW) of the Follow-Up Period. If present, measurements of instep and calf were taken.
19. Height was measured at Screening only.
20. Ophthalmic examination: best corrected visual acuity, pupil examination, color vision assessment, Ishihara color plates (or equivalent) or related red desaturation, intraocular pressure and slit lamp examination. If significant abnormalities, including a loss of 2 lines or more of vision, development of pupil abnormalities including afferent pupillary defect, rise in intraocular pressure, development of corneal infiltrates or other abnormalities not specified here but of concern to the ophthalmologist, were noted compared to previous visits, further investigations of visual function were conducted according to the ophthalmologist's decision.
21. Subjects who had decreased best-corrected visual acuity due to optic neuropathy (defined by a decrease in vision of 2 lines on the Snellen chart, new visual field defect, or color defect secondary to optic nerve involvement within the last 6 months) were not eligible for randomization.
22. Vital signs (heart rate, blood pressure, respiratory rate, temperature) were measured at all clinic visits. Vital signs were measured pre- and post-infusion on Day 1 and Week 3, and predose on all other infusion days. Additional vital signs were monitored if infusion-associated AEs occurred.
23. Non-diabetic subjects were fasting at Weeks 1 and 4 only. Diabetic subjects were fasting at each visit blood glucose was evaluated.
24. Alanine aminotransferase/aspartate aminotransferase was to be $\leq 3 \times$ ULN and serum creatinine was to be $< 1.5 \times$ ULN according to age to be eligible for randomization.
25. Hypothyroidism (defined as FT4 and FT3 levels $< 50\%$ above or below the normal limits). Every effort was made to correct the mild hypo- or hyperthyroidism promptly and to maintain the euthyroid state for the full duration of the clinical trial.
26. HbA1c was $< 9.0\%$ for randomization. If the HbA1c was elevated and considered clinically significant at any time point after Screening, it was repeated approximately every 45 days until it returned to normal or Baseline value.
27. If a sample was positive in the ADA test, after confirmatory and reactive titer testing, the sample was then tested for NAb. If the subject tested positive for NAb, he/she was followed until levels either reverted to Baseline or the subject's value decreased or remained stable. Any subject with a positive NAb test at Week 72 (or PW) during the Follow-Up Period continued to be followed until the subject's value decreased or remained stable.
28. Not collected for subjects who prematurely discontinued from the Double-Masked Treatment Period.
29. AEs that occurred within 2 weeks prior to Day 1 and prior to dosing on Day 1 were considered Baseline signs/symptoms. AEs occurring or worsening after the first dose on Day 1 through the end of the Double-Masked Treatment Period were considered treatment-emergent. AEs occurring or worsening during the Follow-Up Period were considered postdose AEs. All SAEs that occurred from the signing of informed consent through 30 days after study discontinuation were recorded.
30. Serum was obtained on Day 1 and Weeks 12 and 24 of the Double-Masked Treatment Period for possible analysis of IL-4, IL-6, IL-10, IL-12, IL-13, IL-17, IL-23, IL-1 β , sIL-1RA, INF γ , TGF β , TNF α , micRNA and TSH-R-Ab. Based on the results of the assays, other similar serum biomarkers may have been assayed to further explore drug and disease mechanisms.
31. Pharmacokinetic samples were collected prior to and at the end of the infusion on Day 1 and Weeks 3 and 9 of the Double-Masked Treatment Period; additional single samples were collected at Weeks 1, 4 and 24.
32. If TED treatment had been received since last contact, the subject was questioned regarding type of treatment and outcome/response.

(b) (4) (teprotumumab- (b) (4))

Review Study #2 Primary Efficacy Endpoint- The primary efficacy endpoint was the proptosis responder rate (percentage of subjects with a ≥ 2 mm reduction from Baseline in proptosis in the study eye, without deterioration [≥ 2 mm increase] of proptosis in the fellow eye) at Week 24.

Reviewer's Comments: *The Agency was in agreement with the primary endpoint.*

Secondary Efficacy Endpoints (hierarchical testing)

1. Overall responder rate (percentage of subjects with ≥ 2 mm reduction in proptosis AND ≥ 2 point reduction in CAS from Baseline in the study eye, provided there was no corresponding deterioration [≥ 2 mm/point increase] in proptosis or CAS in the fellow eye) at Week 24.
2. Percentage of subjects with a CAS value of 0 or 1 (no or minimal inflammatory symptoms) in the study eye at Week 24.
3. Mean change from Baseline to Week 24 in proptosis measurement in the study eye.
4. Diplopia responder rate (percentage of subjects with Baseline diplopia grade >0 in the study eye who had a reduction of ≥ 1 grade with no corresponding deterioration [≥ 1 grade worsening] in the fellow eye) at Week 24.
5. Mean change from Baseline to Week 24 in the GO-QoL questionnaire overall score.

Reviewer's Comments: *The Agency disagreed with the inclusion of the CAS score as an endpoint. The CAS score is a composite with equal weighting of a number of factors. However, FDA's clinical team does not consider these factors to be of equal weight either to the patients or to physician's treating these patients.*

8.2.2 Review Study #2 Study Results

See Erratum to FDA Briefing Document

Patient Disposition

There were 88 subjects enrolled in the study. Of these, 87 subjects took at least 1 dose of study drug and were included in the ITT and mITT Populations.

	Placebo (N=45) n (%)	HZN-001 (N=43) n (%)
Enrolled (Informed Consent Signed)	45	43
ITT Population	45 (100%)	42 (98%)
MITT Population	45 (100%)	42 (98%)
PP Population	36 (80%)	33 (77%)
Safety Population	44 (98%)	43 (100%)
Completed the Study Treatment Reason for Early Termination	39 (87%)	37 (86%)
Adverse Event	1 (2%)	5 (12%)
Lack of Efficacy	2 (4%)	0
Pregnancy	0	0
Protocol Violation	0	0
Study Terminated by Sponsor	0	0
Death	0	0
Other – see Note below	3 (7%)	1 (2%)

Abbreviations: ITT = intent to treat, MITT=modified intent to treat, PP=per protocol.

Note: All subjects who signed informed consent were considered enrolled in the study. The percentages presented in this table are based on the ITT Population. Three subjects received the wrong treatment; these 3 subjects were excluded from the PP Population and analyzed under the first treatment actually received for the Safety Population. One subject, randomized to HZN-001, terminated early, and never received any study drug.

(b) (4) (teprotumumab- (b) (4))

Review Study #2 Demographic and Baseline Characteristics (Safety Population)

Review Study #2	Placebo (N=42)	Teprotumumab (N=41)
Age (years) Mean (SD)	48.9 (13.0)	51.6 (12.6)
Median	51.5	53
(Min, Max)	(20, 73)	(31,79)
<65 years old	38 (90%)	32 (78%)
≥65 years old	4 (10%)	9 (22%)
Gender, n (%)		
Female	31 (74%)	29 (71%)
Ethnicity, n (%)		
Hispanic or Latino	1 (2%)	2 (5%)
Not Hispanic or Latino	41 (98%)	39 (95%)
Race, n (%)		
American Indian or Alaska Native	0	0
Asian	1 (2%)	2 (5%)
Black or African–American	2 (5%)	4 (10%)
Native Hawaiian or Other Pacific Islander	0	0
White	37 (88%)	35 (85%)
Mixed	2 (5%)	0
Weight (kg), N	44	43
Mean (SD)	75.8 (18.5)	75.0 (16.5)
Median	74.5	73.9
(Min, Max) Kilograms	45.0, 122.9	49.4, 110.0
Study Eye, n(%)		
Right Eye	20 (48%)	22 (54%)
Smoking Status, n(%)		
Smoker (current or former)	17 (40%)	18 (44%)
Time since diagnosis of Active TED (months)		
Mean (SD)	6.4 (2.4)	6.2 (2.3)
Median	6.8	6.3
Min, Max	1.1, 10.3	0.9, 9.7

Abbreviation: CAS=Clinical Activity Score; CSS = Clinical Measures of Severity Score; ITT=intent-to-treat; N = number; SD =standard deviation

^a Baseline was the last predose measurement.

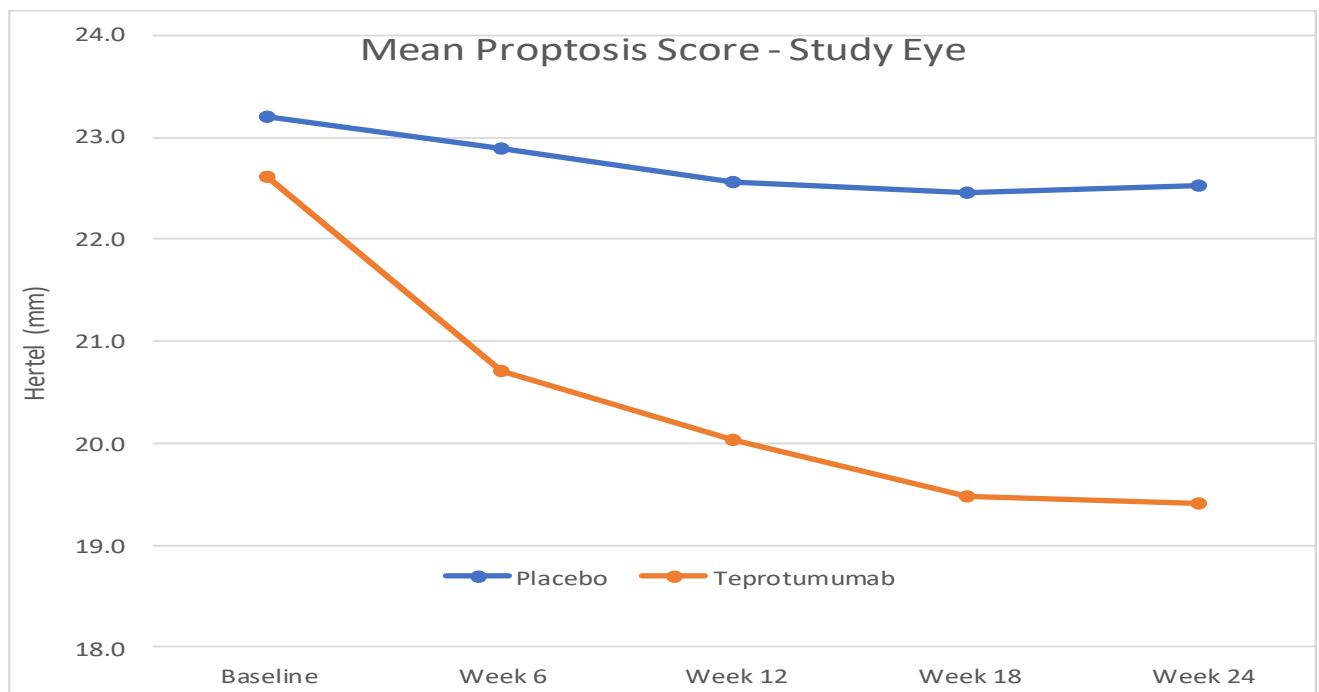
See Erratum to FDA Briefing Document

(b) (4) (teprotumumab- (b) (4))

Review Study #2 Efficacy Results – Primary Endpoint: % patients with 2 mm or more decrease in Proptosis

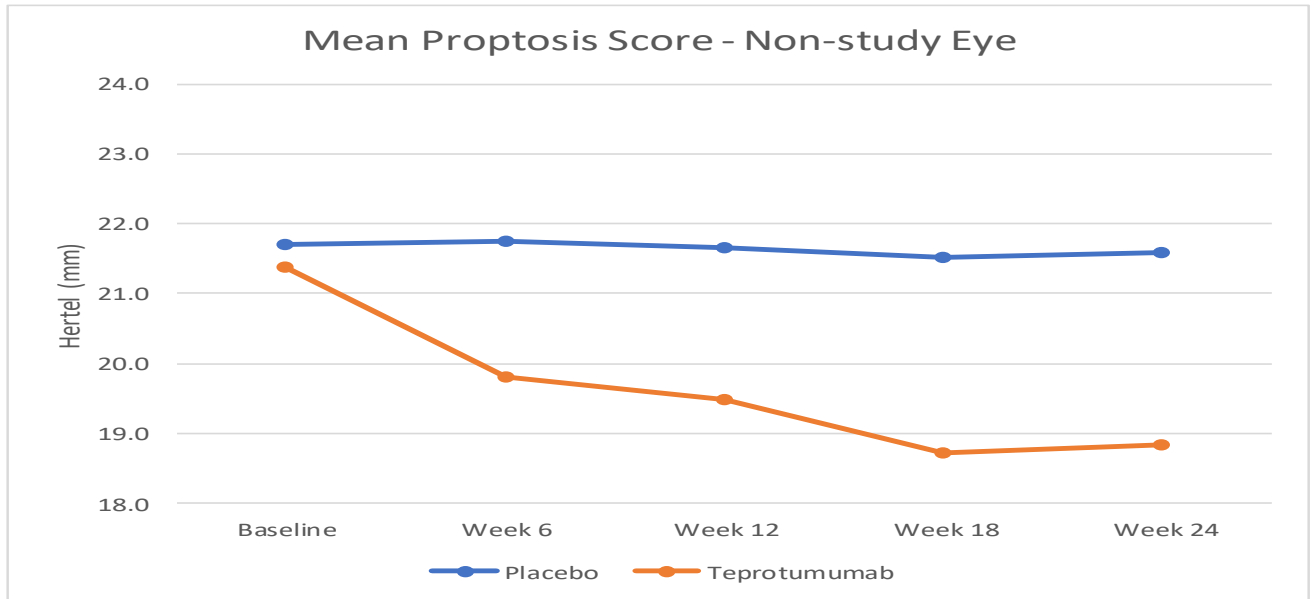
Proptosis	Placebo	Teprotumumab	Difference	p-value
Week 6 Study Eye	3/42 (7%)	23/40 (58%)	51%	<0.001
Week 12 Study Eye	6/41 (15%)	31/39 (80%)	65%	<0.001
Week 18 Study Eye	6/40 (15%)	34/39 (87%)	72%	<0.001
Week 24 Study Eye	4/40 (10%)	34/40 (85%)	75%	<0.001
Week 6 Non-study Eye	0/42	22/39 (55%)	55%	<0.001
Week 12 Non-study Eye	2/41 (5%)	24/39 (62%)	57%	<0.001
Week 18 Non-study Eye	2/40 (5%)	29/39 (74%)	69%	<0.001
Week 24 Non-study Eye	1/40 (3%)	27/40 (68%)	65%	<0.001

Reviewer's Comments: *By the first evaluation period at Week 6, there is a clinically significant reduction in proptosis (i.e., greater than 2 mm) in both eyes which continues through the treatment period.*



Reviewer's Comments: *The clinical effect continues to improve over the course of treatment.*

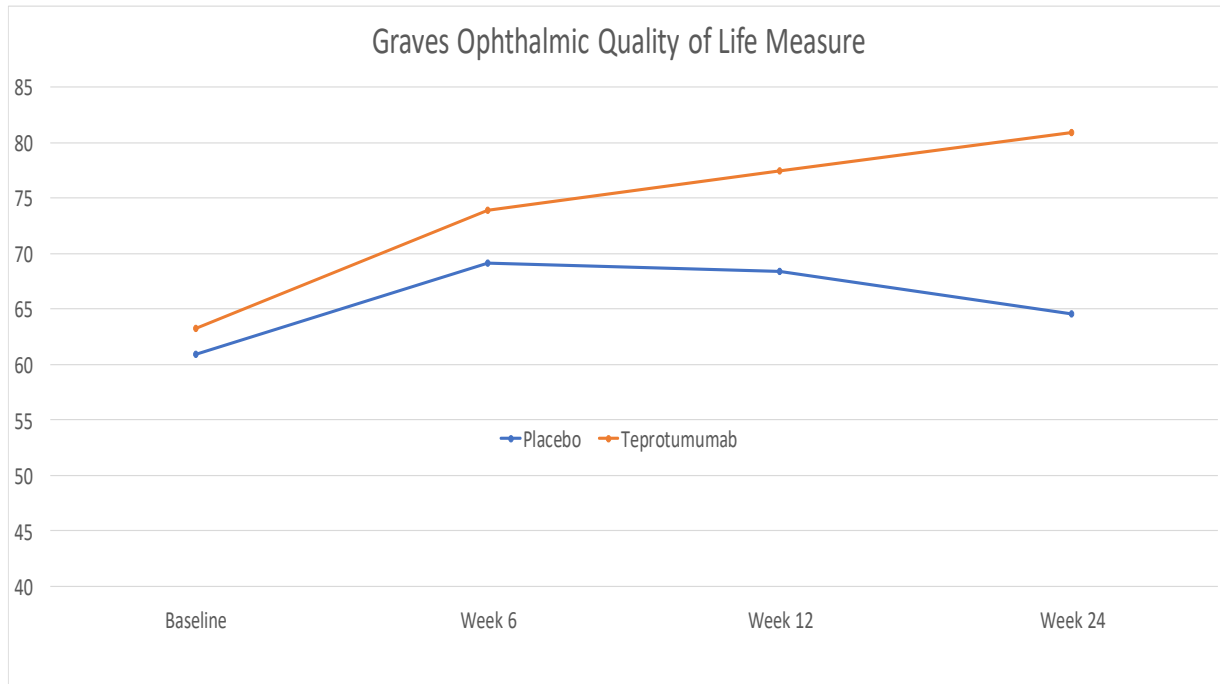
Review Study #2 – Non-study Eye Proptosis



Reviewer's Comments: *By the first evaluation period at Week 6, there is a reduction in proptosis in both eyes which continues through the treatment period.*

(b) (4) (teprotumumab- (b) (4))

**Review Study #2 Secondary Endpoints
Graves Ophthalmic Quality of Life Measure**



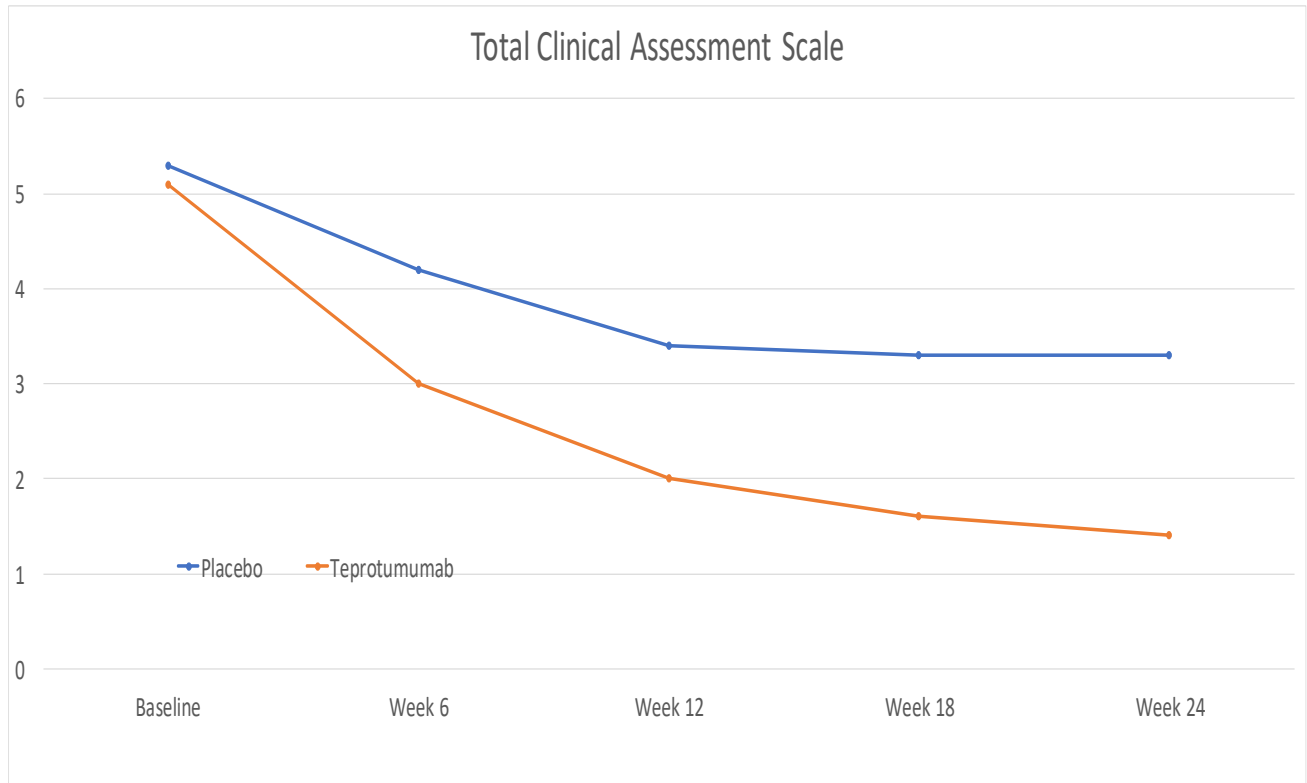
Transformed score = [(sum of each score – number of completed items) / (2 * number of completed items)] * 100.

Grave’s Ophthal Quality of Life Score	Placebo	Teprotumumab	Difference
Baseline	60.9 (19.4)	63.3 (22.1)	2.4
Week 6	69.1 (16.3)	73.9 (21.0)	4.8
Week 12	68.4 (16.5)	77.5 (21.9)	9.1
Week 24	64.6 (18.7)	80.9 (17.6)	16.3

Reviewer’s Comments: *Validation information for the Grave’s Ophthalmology Quality of Life Score has not been submitted and therefore interpretation of the scores is not possible.*

(b) (4) (teprotumumab- (b) (4))

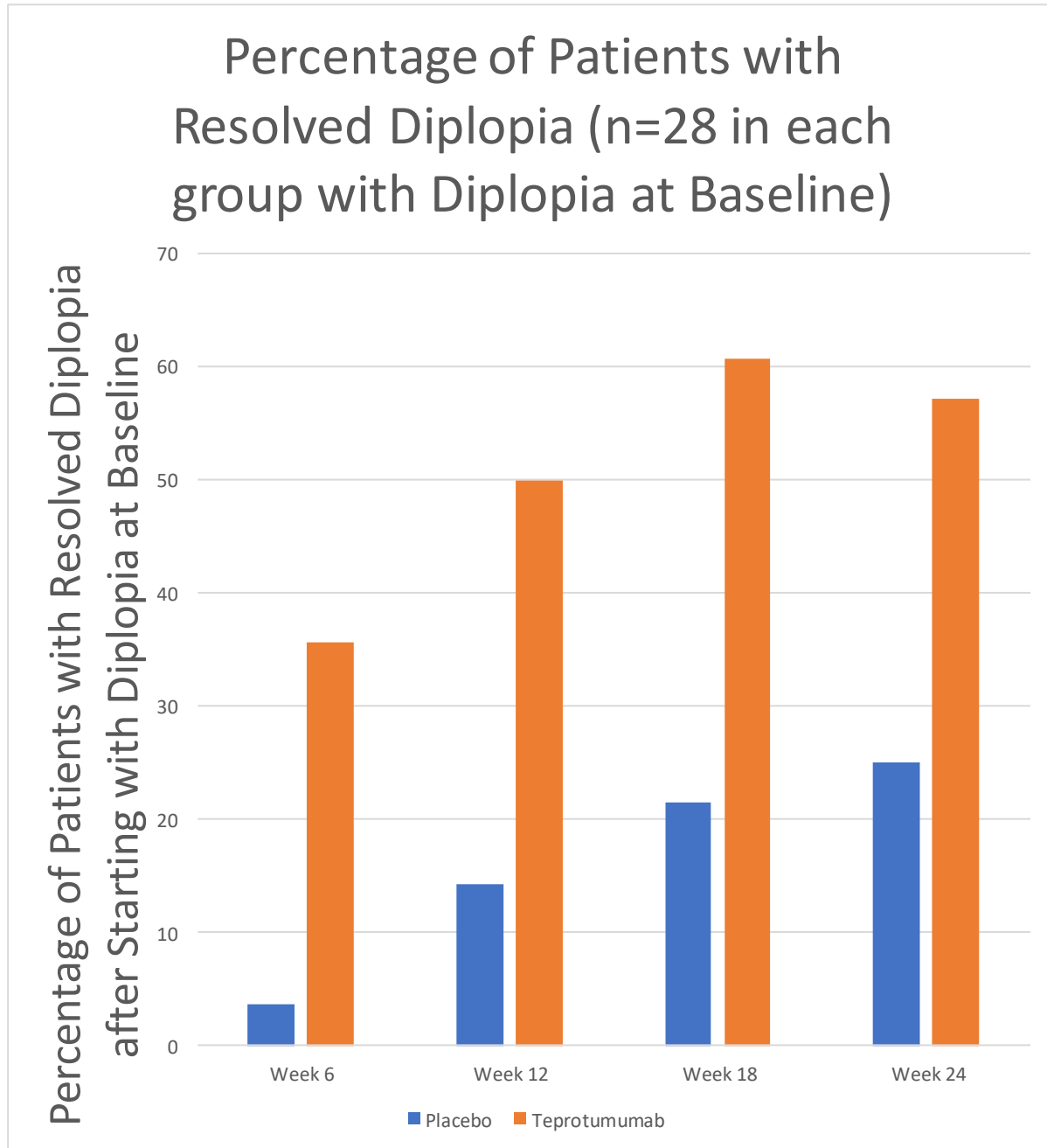
Review Study #2 – Clinical Assessment Scale



CAS Change from Baseline	Placebo	Teprotumumab	Difference
Baseline	5.3 (1.0)	5.1 (0.9)	0.2
Week 6	4.2 (1.5)	3.0 (1.7)	1.2
Week 12	3.4 (1.6)	2.0 (1.5)	1.4
Week 18	3.3 (1.9)	1.6 (1.7)	1.7
Week 24	3.3 (1.9)	1.4 (1.4)	1.9

Reviewer's Comments: *Clinical Activity Scale is not accepted because there is not necessarily equal weight for each component.*

Review Study #2- Resolution of Diplopia



Reviewer's Comments: *Diplopia was resolved in approximately 57% of patients with diplopia at the start of the clinical trial.*

8.2.3 Extension Period – From Safety Update

The analysis presented in the initial submission included data collected from 2 randomized, double-masked, placebo-controlled, parallel-group studies (Phase 2 Study TED01RV (Review Study #1) and Phase 3 Study HZNP-TEP-301; OPTIC (Review Study #2)); both study designs included a 24-Week Treatment Period and a Follow-up Period with no additional study treatment. The Treatment and Follow-up Periods of Study TED01RV had been completed; however, the Follow-up Period of Study HZNP-TEP-301 was ongoing and safety data through 19 February 2019, were included in the submission. In addition, safety data from an ongoing open-label extension study (Study HZNP-TEP-302; OPTIC-X) through 27 February 2019, were also included in the initial submission.

Among the 121 subjects in the All Teprotumumab Population, 97 (80.2%) had received at least 8 doses of teprotumumab at the time of the data cutoff. Among the 9 subjects who received teprotumumab in Study HZNP-TEP-301 and in Study HZNP-TEP-302, the total number of teprotumumab doses received across both studies was 9 for 2 subjects, 10 for 1 subject, 11 for 1 subject, 12 for 1 subject, 14 for 1 subject, 15 for 1 subject, and 16 for 2 subjects. Among the 121 subjects in the All Teprotumumab Population, 63 (52.1%) have been followed for at least 24 weeks off treatment and 29 (24.0%) have been followed for at least 48 weeks off treatment.

Since the initial submission, 1 additional subject experienced a serious adverse event during the Treatment Period (life-threatening *Cerebral haemorrhage*) and 2 additional subjects experienced serious adverse events severe *Intercostal neuralgia* and severe *Optic neuropathy* during the Follow-up Period.

(b) (4) (teprotumumab- (b) (4))

9. Review of Safety

9.1 Deaths – none

9.2 Serious Adverse Events

Listing of Serious Treatment-Emergent Adverse Events

See Erratum to FDA Briefing Document

Review Study #1

Site	Preferred Term	Start Date/ End Date	Outcome	Study Medication Action Taken	Other Action Taken
Teprotumumab					
(b) (6)	Hashimoto's encephalopathy	(b) (6)	Unknown	Drug interrupted	Hospitalization, (b) (6)
	Urinary retention		Resolved	Dose not changed	Medication and hospitalization
	Diarrhoea ^b		Resolved	Drug withdrawn	Hospitalization
	Escherichia sepsis		Unknown	Drug	Hospitalization
	Inflammatory bowel disease		Resolved With Sequelae	Drug withdrawn	Medication and hospitalization
Placebo					
(b) (6)	Optic neuropathy	(b) (6)	Resolved	Not applicable	Medication

(b) (6)

Subject had medical history of colitis with abdominal cramping and bloody diarrhea in the 7 months preceding randomization. After 3 months on study, subject had a colonoscopy and was diagnosed with ulcerative colitis. Source: [Appendix 16.2, Listing 16.3.1.2.](#)

Review Study #2

Site	Preferred Term	Day of Onset	Outcome	Study Medication Action Taken	Other Action Taken
Teprotumumab					
(b) (6)	Infusion related reaction	1	Resolved	Drug withdrawn	Concomitant medication discontinued study
(b) (6)	Pneumothorax	113	Recovering	Dose not changed	Hospitalization
Placebo					
(b) (6)	Visual field defect	64	Recovering	Drug withdrawn	Orbital decompression surgery

Source: [Listing 16.2.4.1 and 16.2.7.2.](#)

9.3 Dropouts and/or Discontinuations

No treatment – Review Study #1

Site (b) (6) (b) (6) Subject voluntarily withdrew from the study due to difficulties with placing an IV and withdrawing blood at the baseline visit. No infusions were administered to the patient.

Teprotumumab

Review Study #1 Day

Site (b) (6)	130	AE Hospitalized for altered mental status. Suspect Hashimoto’s Encephalopathy
Site	43	AE Facial flushing, heart palpitations, elevated BP & Heart rate
Site	253	AE Diarrhea
Site	79	The subject was dispensed the incorrect treatment assignment at Week 3 in error and an administrative decision on the part of the sponsor was made to discontinue the subject
Site	45	AE Hospitalization for systemic E. Coli Sepsis and dehydration
Site	169	AE Inflammatory bowel disease
Site	127	Back pain, back surgery

Review Study #2

Site (b) (6)	1	AE Infusion Reaction
Site (b) (6)	43	AE Skin itchy and red after 3 rd dose, concerned about the risk of allergy

Placebo

Review Study #1

Site (b) (6)	47	Left eye, Optic disc edema
Site	170	Lack of Efficacy
Site	64	Lack of Efficacy
Site	1	AE Vasovagal Attack
Site	302	Patient decision to withdraw

Review Study #2

Site (b) (6)	43	Worsening Visual Field
Site (b) (6)	85	Subject Decision

(b) (4) (teprotumumab- (b) (4))

9.4 Treatment Emergent Adverse Events and Adverse Reactions in at least 5%

System Organ Class	Study 1 Placebo (N=44)	Study 2 Placebo (N=42)	Study 1 Teprotumumab (N=43)	Study 2 Teprotumumab (N=41)
Preferred Term	n (%)	n (%)	n (%)	n (%)
Any TEAE ^a	32 (73%)	29 (69%)	32 (74%)	35 (85%)
Gastrointestinal Disorders	6 (14%)	9 (21%)	16 (37%)	18 (44%)
Nausea	4 (9%)	4 (10%)	8 (19%)	6 (15%)
Diarrhea	2 (5%)	5 (12%)	6 (14%)	4 (10%)
Abdominal pain upper		3 (7%)		2 (5%)
Stomatitis		1 (2%)		3 (7%)
Infections and Infestations	9 (21%)	10 (24%)	13 (30%)	16 (40%)
Upper respiratory tract infection	4 (9%)		0	
Influenza		3 (7%)		1 (2%)
Respiratory, thoracic and mediastinal disorders		4 (10%)		6 (15%)
Cough		3 (7%)		2 (5%)
Skin and Subcutaneous Tissue Disorders	9 (20%)	11 (26%)	11 (26%)	15 (37%)
Alopecia	2 (5%)	5 (12%)	3 (7%)	8 (20%)
Dry skin	0	0	3 (7%)	4 (10%)
Rash	4 (9%)		3 (7%)	
Musculoskeletal and Connective Tissue Disorders	7 (16%)	5 (12%)	12 (28%)	16 (39%)
Muscle spasms	2 (5%)	4 (10%)	8 (19%)	13 (32%)
Nervous System Disorders	9 (20%)	8 (19%)	10 (23%)	14 (34%)
Dizziness	4 (9%)	0	0	3 (7%)
Dysgeusia	0	0	3 (7%)	4 (10%)
Headache	2 (5%)	4 (10%)	3 (7%)	4 (10%)
Paresthesia	0		3 (7%)	
Somnolence	3 (7%)		0	
Investigations	7 (16%)		9 (21%)	
Weight decreased	0		3 (7%)	
Metabolism and Nutrition Disorders	2 (5%)		10 (23%)	
Hyperglycemia	2 (5%)	0	5 (12%)	2 (5%)
Reproductive system and breast disorders		0		4 (10%)
Amenorrhea		0		3 (7%)
General Disorders and Site Conditions	10 (23%)	4 (10%)	6 (14%)	8 (20%)
Fatigue	6 (14%)	1 (2%)	3 (7%)	5 (12%)

Abbreviations: MedDRA = Medical Dictionary for Regulatory Activities; TEAE = treatment-emergent adverse event. Note: The denominator for the percentages is the number of subjects in each treatment group. At each level of summarization, subjects who experienced more than 1 TEAE were counted only once. All TEAEs were coded using MedDRA, Version 14.0. ^a A TEAE was defined as an AE with onset at the time of or following the start of treatment with study drug or an AE starting before the start of treatment but increasing in severity following the start of treatment. [Table 14.3.1.1](#)

Reviewer's Comments: *Interpretability is difficult because of the low number of subjects enrolled in the clinical trials. There appear to be increased trends in the teprotumumab groups for gastrointestinal disorders, infections, muscle spasms, hyperglycemia and reproductive system and breast disorders.*

(b) (4) (teprotumumab- (b) (4))

9.5 Laboratory Findings

With the exception of elevated glucose and hemoglobin A1c levels in some patients, no significant shifts in laboratory findings were noted.

9.6 Vital Signs

No clinically significant changes were reported.

9.7 Electrocardiograms (ECGs)

No clinically significant changes were reported.

9.8 QT

Monoclonal antibodies would not be expected to have a significant risk of inducing QT changes. A traditional thorough QT study (based on International Conference on Harmonization of Technical Requirements for the Registration of Pharmaceuticals for Human Use Guidance: E14) was considered unnecessary and was not conducted. In Review Studies #1 and #2, all of the teprotumumab-treated subjects had electrocardiogram (ECG) results at Baseline and throughout the Treatment Period. None demonstrated clinically significant findings.

9.9 Immunogenicity

No clinically significant changes were reported.

10. Pediatrics

Teprotumumab was granted orphan drug designation for the treatment of Active Thyroid Eye Disease (Orphan Drug Designation 12-3878). Submission of a pediatric assessment is not required for an application to market a product for an orphan-designated indication. In addition, Thyroid Eye Disease occurs very rarely, if at all in pediatric patients.

(b) (4) (teprotumumab- (b) (4))

11. Safety Update Summary of Reported Adverse Events

Teportumumab N=121

System Organ Class or Verbatim Term	Subjects	(%)	Events
Any TEAEs	101	(83%)	553
Musculoskeletal and connective tissue disorders	49	(40%)	94
Gastrointestinal disorders	43	(36%)	88
Skin and subcutaneous tissue disorders	43	(36%)	72
Infections and infestations	40	(33%)	49
Nervous system disorders	32	(26%)	55
Ear and labyrinth disorders	21	(17%)	25
General disorders and administration site conditions	20	(17%)	31
Metabolism and nutrition disorders	17	(14%)	20
Respiratory, thoracic and mediastinal disorders	15	(12%)	23
Investigations	13	(11%)	16
Injury, poisoning and procedural complications	11	(9%)	11
Reproductive system and breast disorders	10	(8%)	14
Eye disorders	10	(8%)	15
Psychiatric disorders	7	(6%)	8
Renal and urinary disorders	7	(6%)	8
Vascular disorders	6	(5%)	7
Cardiac disorders	4	(3%)	5
Blood and lymphatic system disorders	4	(3%)	4
Neoplasms benign, malignant and unspecified (incl cysts and polyps)	3	(2.5%)	4
Endocrine disorders	2	(2%)	2
Muscle spasms	39	(32%)	75
Alopecia	18	(15%)	21
Nausea	15	(12%)	24
Diarrhea	15	(12%)	20
Fatigue	12	(10%)	15
Dry skin	12	(10%)	12
Dysgeusia	10	(8%)	15
Headache	8	(7%)	10
Rash	8	(7%)	10
Ear discomfort	8	(7%)	9
Urinary tract infection	7	(6%)	7
Amenorrhoea	6	(5%)	7
Hyperglycaemia	6	(5%)	7
Abdominal pain upper	6	(5%)	6
Influenza	5	(4%)	6
Pain in extremity	5	(4%)	6
Bronchitis	5	(4%)	5
Hypoacusis	5	(4%)	5
Onychoclasis	5	(4%)	5

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(b) (4) (teprotumumab- (b) (4))

Epistaxis	4	(3%)	7
Sinusitis	4	(3%)	5
Abdominal pain	4	(3%)	4
Cystitis	4	(3%)	4
Dizziness	4	(3%)	4
Rhinorrhoea	4	(3%)	4
Tremor	4	(3%)	4
Weight decreased	4	(3%)	4
Madarosis	3	(2.5%)	4
Tinnitus	3	(2.5%)	4
Blood glucose increased	3	(2.5%)	3
Blood pressure increased	3	(2.5%)	3
Cough	3	(2.5%)	3
Decreased appetite	3	(2.5%)	3
Depression	3	(2.5%)	3
Dry mouth	3	(2.5%)	3
Feeling hot	3	(2.5%)	3
Gastroesophageal reflux disease	3	(2.5%)	3
Hypertension	3	(2.5%)	3
Nasopharyngitis	3	(2.5%)	3
Noninfective gingivitis	3	(2.5%)	3
Palpitations	3	(2.5%)	3
Paraesthesia	3	(2.5%)	3
Stomatitis	3	(2.5%)	3
Vomiting	3	(2.5%)	3
Asthenia	2	(2%)	4
Abdominal distension	2	(2%)	2
Arthralgia	2	(2%)	2
Chest pain	2	(2%)	2
Deafness	2	(2%)	2
Dry eye	2	(2%)	2
Dysuria	2	(2%)	2
Gingival pain	2	(2%)	2
Gingival recession	2	(2%)	2
Glossodynia	2	(2%)	2
Hair growth abnormal	2	(2%)	2
Ingrowing nail	2	(2%)	2
Localised infection	2	(2%)	2
Myalgia	2	(2%)	2
Nail disorder	2	(2%)	2
Nasal congestion	2	(2%)	2
Nasal dryness	2	(2%)	2
Nightmare	2	(2%)	2
Osteopenia	2	(2%)	2
Rash pruritic	2	(2%)	2
Tachycardia	2	(2%)	2
Thrombocytopenia	2	(2%)	2
Vertigo	2	(2%)	2
Muscle contractions involuntary	1	(1%)	4
Butterfly rash	1	(1%)	2

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(b) (4) (teprotumumab- (b) (4))

Disturbance in attention	1	(1%)	2
Inflammatory bowel disease	1	(1%)	2
Metrorrhagia	1	(1%)	2
Nail discolouration	1	(1%)	2
Rectal haemorrhage	1	(1%)	2
Trichiasis	1	(1%)	2
Abnormal faeces	1	(1%)	1
Abnormal sensation in eye	1	(1%)	1
Acarodermatitis	1	(1%)	1
Acne	1	(1%)	1
Adverse drug reaction	1	(1%)	1
Aphasia	1	(1%)	1
Arthropod sting	1	(1%)	1
Bacterial vaginosis	1	(1%)	1
Balance disorder	1	(1%)	1
Biotin deficiency	1	(1%)	1
Blepharospasm	1	(1%)	1
Blood bilirubin increased	1	(1%)	1
Blood creatinine increased	1	(1%)	1
Blood urine present	1	(1%)	1
Cerebral haemorrhage	1	(1%)	1
Chills	1	(1%)	1
Chronic kidney disease	1	(1%)	1
Confusional state	1	(1%)	1
Contusion	1	(1%)	1
Corneal abrasion	1	(1%)	1
Corneal erosion	1	(1%)	1
Dehydration	1	(1%)	1
Dermal cyst	1	(1%)	1
Dermatitis	1	(1%)	1
Diabetes mellitus	1	(1%)	1
Diplopia	1	(1%)	1
Dysarthria	1	(1%)	1
Dysmenorrhoea	1	(1%)	1
Dyspepsia	1	(1%)	1
Ear infection	1	(1%)	1
Erectile dysfunction	1	(1%)	1
Erythema of eyelid	1	(1%)	1
Escherichia sepsis	1	(1%)	1
Eustachian tube dysfunction	1	(1%)	1
Eustachian tube patulous	1	(1%)	1
Eye irritation	1	(1%)	1
Eye pain	1	(1%)	1
Eyelid retraction	1	(1%)	1
Feeling cold	1	(1%)	1
Flushing	1	(1%)	1
Gastroenteritis	1	(1%)	1
Gingival bleeding	1	(1%)	1
Gout	1	(1%)	1
Haemangioma of skin	1	(1%)	1

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(b) (4) (teprotumumab- (b) (4))

Haematochezia	1	(1%)	1
Haematoma	1	(1%)	1
Haematuria	1	(1%)	1
Haemoglobin decreased	1	(1%)	1
Hashimoto's encephalopathy	1	(1%)	1
Heart rate increased	1	(1%)	1
Helicobacter infection	1	(1%)	1
Hyperacusis	1	(1%)	1
Hypersensitivity	1	(1%)	1
Hyperthyroidism	1	(1%)	1
Hypogeusia	1	(1%)	1
Hypokalaemia	1	(1%)	1
Hyposmia	1	(1%)	1
Hypotension	1	(1%)	1
Hypothyroidism	1	(1%)	1
Immune system disorders	1	(1%)	1
Increased appetite	1	(1%)	1
Influenza like illness	1	(1%)	1
Infusion related reaction	1	(1%)	1
Inguinal hernia	1	(1%)	1
Iron deficiency	1	(1%)	1
Joint stiffness	1	(1%)	1
Jugular vein distension	1	(1%)	1
Lenticular opacities	1	(1%)	1
Ligament sprain	1	(1%)	1
Limb injury	1	(1%)	1
Loss of consciousness	1	(1%)	1
Lymphadenopathy	1	(1%)	1
Medial tibial stress syndrome	1	(1%)	1
Memory impairment	1	(1%)	1
Migraine	1	(1%)	1
Migraine with aura	1	(1%)	1
Migraine without aura	1	(1%)	1
Muscle injury	1	(1%)	1
Muscle rupture	1	(1%)	1
Muscle strain	1	(1%)	1
Muscular weakness	1	(1%)	1
Musculoskeletal pain	1	(1%)	1
Musculoskeletal stiffness	1	(1%)	1
Nail infection	1	(1%)	1
Nasal discomfort	1	(1%)	1
Night sweats	1	(1%)	1
Oedema peripheral	1	(1%)	1
Oral herpes	1	(1%)	1
Oropharyngeal pain	1	(1%)	1
Osteoporosis	1	(1%)	1
Otitis media	1	(1%)	1
Paranasal sinus discomfort	1	(1%)	1
Pelvic discomfort	1	(1%)	1
Periodontitis	1	(1%)	1

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(b) (4) (teprotumumab- (b) (4))

Peripheral swelling	1	(1%)	1
Petechiae	1	(1%)	1
Pneumonia chlamydial	1	(1%)	1
Pneumothorax	1	(1%)	1
Pollakiuria	1	(1%)	1
Polydipsia	1	(1%)	1
Polyuria	1	(1%)	1
Procedural nausea	1	(1%)	1
Pruritus	1	(1%)	1
Pruritus generalized	1	(1%)	1
Pyogenic granuloma	1	(1%)	1
Rhinitis	1	(1%)	1
Seborrhoeic keratosis	1	(1%)	1
Sinusitis bacterial	1	(1%)	1
Sleep disorder	1	(1%)	1
Sneezing	1	(1%)	1
Spontaneous haematoma	1	(1%)	1
Squamous cell carcinoma	1	(1%)	1
Strabismus	1	(1%)	1
Subcutaneous abscess	1	(1%)	1
Tendonitis	1	(1%)	1
Tension	1	(1%)	1
Thirst	1	(1%)	1
Tonsillitis	1	(1%)	1
Tooth abscess	1	(1%)	1
Tooth development disorder	1	(1%)	1
Tooth infection	1	(1%)	1
Toothache	1	(1%)	1
Trichorrhexis	1	(1%)	1
Type 2 diabetes mellitus	1	(1%)	1
Urinary retention	1	(1%)	1
Urine odour abnormal	1	(1%)	1
Vaginal discharge	1	(1%)	1
Vaginal haemorrhage	1	(1%)	1
Vascular injury	1	(1%)	1
Vision blurred	1	(1%)	1
Visual field defect	1	(1%)	1
Vitamin D deficiency	1	(1%)	1
Weight increased	1	(1%)	1

Reviewer's Comments: *Using the verbatim term for the reported adverse event has served to separate terms which may otherwise refer to a very similar adverse event. For example, Tooth abscess, Tooth infection, and Toothache should be included as the same adverse event.*