TRAZIMERA- trastuzumab-qyyp TRAZIMERA- trastuzumab-qyyp injection, powder, lyophilized, for solution Pfizer Laboratories Div Pfizer Inc

HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use TRAZIMERA safely and effectively. See full prescribing information for TRAZIMERA.

TRAZIMERA™ (trastuzumab-qyyp) for injection, for intravenous use Initial U.S. Approval: 2019

TRAZIMERA (trastuzumab-qyyp) is biosimilar* to HERCEPTIN (trastuzumab).

WARNING: CARDIOMYOPATHY, INFUSION REACTIONS, EMBRYO-FETAL TOXICITY, and PULMONARY TOXICITY

See full prescribing information for complete boxed warning

Cardiomyopathy: Trastuzumab products can result in subclinical and clinical cardiac failure manifesting as CHF, and decreased LVEF, with greatest risk when administered concurrently with anthracyclines. Evaluate cardiac function prior to and during treatment. Discontinue TRAZIMERA for cardiomyopathy. (2.3, 5.1) Infusion Reactions, Pulmonary Toxicity: Discontinue TRAZIMERA for anaphylaxis, angioedema, interstitial pneumonitis, or acute respiratory distress syndrome. (5.2, 5.4)

Embryo-Fetal Toxicity: Exposure to trastuzumab products during pregnancy can result in oligohydramnios, in some cases complicated by pulmonary hypoplasia and neonatal death. Advise patients of these risks and the need for effective contraception. (5.3, 8.1, 8.3)

	RECENT	MAJOR	CHANGES
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Dosage and Administration, Preparation for Administration (2.4)

11/2020

----- INDICATIONS AND USAGE

TRAZIMERA is a HER2/neu receptor antagonist indicated for:

- The treatment of HER2-overexpressing breast cancer. (1.1, 1.2)
- The treatment of HER2-overexpressing metastatic gastric or gastroesophageal junction adenocarcinoma. (1.3)

Select patients for therapy based on an FDA-approved companion diagnostic for a trastuzumab product (1, 2.1).

----- DOSAGE AND ADMINISTRATION -----

For intravenous (IV) infusion only. Do not administer as an intravenous push or bolus. (2.2) Do not substitute TRAZIMERA (trastuzumab-qyyp) for or with ado-trastuzumab emtansine. (2.2) Perform HER2 testing using FDA-approved tests by laboratories with demonstrated proficiency. (1, 2.1) Adjuvant Treatment of HER2-Overexpressing Breast Cancer (2.2)

Administer at either:

- Initial dose of 4 mg/kg over 90 minute intravenous infusion, then 2 mg/kg over 30 minute intravenous infusion weekly for 12 weeks (with paclitaxel or docetaxel) or 18 weeks (with docetaxel and carboplatin). One week after the last weekly dose of TRAZIMERA, administer 6 mg/kg as an intravenous infusion over 30 to 90 minutes every three weeks to complete a total of 52 weeks of therapy, or
- Initial dose of 8 mg/kg over 90 minutes intravenous infusion, then 6 mg/kg over 30 to 90 minutes intravenous infusion every three weeks for 52 weeks.

Metastatic HER2-Overexpressing Breast Cancer (2.2)

• Initial dose of 4 mg/kg as a 90 minute intravenous infusion followed by subsequent weekly doses of 2 mg/kg as 30 minute intravenous infusions.

Metastatic HER2-Overexpressing Gastric Cancer (2.2) Initial dose of 8 mg/kg over 90 minutes intravenous infusion, followed by 6 mg/kg over 30 to 90 minutes intravenous infusion every 3 weeks. DOSAGE FORMS AND STRENGTHS For Injection: 150 mg lyophilized powder in a single-dose vial for reconstitution (3) For Injection: 420 mg lyophilized powder in a multiple-dose vial for reconstitution (3) CONTRAINDICATIONS None. (4) WARNINGS AND PRECAUTIONS Exacerbation of Chemotherapy-Induced Neutropenia. (5.5, 6.1)

Adjuvant Breast Cancer

• Most common adverse reactions (≥ 5%) are headache, diarrhea, nausea, and chills. (6.1)

Metastatic Breast Cancer

• Most common adverse reactions (≥ 10%) are fever, chills, headache, infection, congestive heart failure, insomnia, cough, and rash. (6.1)

------ADVERSE REACTIONS-------

Metastatic Gastric Cancer

 Most common adverse reactions (≥ 10%) are neutropenia, diarrhea, fatigue, anemia, stomatitis, weight loss, upper respiratory tract infections, fever, thrombocytopenia, mucosal inflammation, nasopharyngitis, and dysgeusia. (6.1)

See 17 for PATIENT COUNSELING INFORMATION.

* Biosimilar means that the biological product is approved based on data demonstrating that it is highly similar to an FDA-approved biological product, known as a reference product, and that there are no clinically meaningful differences between the biosimilar product and the reference product. Biosimilarity of TRAZIMERA has been demonstrated for the condition(s) of use (e.g. indication(s), dosing regimen(s)), strength(s), dosage form(s), and route(s) of administration described in its Full Prescribing Information.

Revised: 11/2020

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FULL PRESCRIBING INFORMATION

WARNING: CARDIOMYOPATHY, INFUSION REACTIONS, EMBRYO-FETAL TOXICITY, and PULMONARY TOXICITY

Cardiomyopathy

Administration of trastuzumab products can result in sub-clinical and clinical cardiac failure. The incidence and severity was highest in patients receiving trastuzumab with anthracycline-containing chemotherapy regimens.

Evaluate left ventricular function in all patients prior to and during treatment with TRAZIMERA. Discontinue TRAZIMERA treatment in patients receiving adjuvant therapy and withhold TRAZIMERA in patients with metastatic disease for clinically significant decrease in left ventricular function [see Dosage and Administration (2.3) and Warnings and Precautions (5.1)].

Infusion Reactions; Pulmonary Toxicity

Administration of trastuzumab products can result in serious and fatal infusion reactions and pulmonary toxicity. Symptoms usually occur during or within 24 hours of administration. Interrupt TRAZIMERA infusion for dyspnea or clinically significant hypotension. Monitor patients until symptoms completely resolve. Discontinue TRAZIMERA for anaphylaxis, angioedema, interstitial pneumonitis, or acute respiratory distress syndrome [see Warnings and Precautions (5.2, 5.4)].

Embryo-Fetal Toxicity

Exposure to trastuzumab products during pregnancy can result in oligohydramnios and oligohydramnios sequence manifesting as pulmonary hypoplasia, skeletal abnormalities, and neonatal death. Advise patients of these risks and the need for effective contraception [see Warnings and Precautions (5.3) and Use in Specific Populations (8.1, 8.3)].

1 INDICATIONS AND USAGE

1.1 Adjuvant Breast Cancer

TRAZIMERA is indicated for adjuvant treatment of HER2-overexpressing node positive or node negative (ER/PR negative or with one high risk feature [see Clinical Studies (14.1)]) breast cancer

- as part of a treatment regimen consisting of doxorubicin, cyclophosphamide, and either paclitaxel or docetaxel
- as part of a treatment regimen with docetaxel and carboplatin
- as a single agent following multi-modality anthracycline based therapy.

Select patients for therapy based on an FDA-approved companion diagnostic for a trastuzumab product [see Dosage and Administration (2.1)].

1.2 Metastatic Breast Cancer

TRAZIMERA is indicated:

- In combination with paclitaxel for first-line treatment of HER2-overexpressing metastatic breast cancer
- As a single agent for treatment of HER2-overexpressing breast cancer in patients who have received one or more chemotherapy regimens for metastatic disease.

Select patients for therapy based on an FDA-approved companion diagnostic for a trastuzumab product [see Dosage and Administration (2.1)].

1.3 Metastatic Gastric Cancer

TRAZIMERA is indicated, in combination with cisplatin and capecitabine or 5-fluorouracil, for the treatment of patients with HER2-overexpressing metastatic gastric or gastroesophageal junction adenocarcinoma who have not received prior treatment for metastatic disease.

Select patients for therapy based on an FDA-approved companion diagnostic for a trastuzumab product [see Dosage and Administration (2.1)].

2 DOSAGE AND ADMINISTRATION

2.1 Patient Selection

Select patients based on HER2 protein overexpression or HER2 gene amplification in tumor specimens [see Indications and Usage (1) and Clinical Studies (14)]. Assessment of HER2 protein overexpression and HER2 gene amplification should be performed using FDA-approved tests specific for breast or gastric cancers by laboratories with demonstrated proficiency. Information on the FDA-approved tests for the detection of HER2 protein overexpression and HER2 gene amplification is available at: http://www.fda.gov/CompanionDiagnostics.

Assessment of HER2 protein overexpression and HER2 gene amplification in metastatic gastric cancer should be performed using FDA-approved tests specifically for gastric cancers due to differences in gastric vs. breast histopathology, including incomplete membrane staining and more frequent heterogeneous expression of HER2 seen in gastric cancers.

Improper assay performance, including use of suboptimally fixed tissue, failure to utilize specified reagents, deviation from specific assay instructions, and failure to include appropriate controls for assay validation, can lead to unreliable results.

2.2 Recommended Doses and Schedules

- Do not administer as an intravenous push or bolus. Do not mix TRAZIMERA with other drugs.
- Do not substitute TRAZIMERA (trastuzumab-qyyp) for or with adotrastuzumab emtansine.

Adjuvant Treatment, Breast Cancer

Administer according to one of the following doses and schedules for a total of 52 weeks of TRAZIMERA therapy:

During and following paclitaxel, docetaxel, or docetaxel and carboplatin:

- Initial dose of 4 mg/kg as an intravenous infusion over 90 minutes then at 2 mg/kg as an intravenous infusion over 30 minutes weekly during chemotherapy for the first 12 weeks (paclitaxel or docetaxel) or 18 weeks (docetaxel and carboplatin).
- One week following the last weekly dose of TRAZIMERA, administer TRAZIMERA at 6 mg/kg as an intravenous infusion over 30 to 90 minutes every three weeks.

As a single agent within three weeks following completion of multi-modality, anthracycline-based chemotherapy regimens:

- Initial dose at 8 mg/kg as an intravenous infusion over 90 minutes.
- Subsequent doses at 6 mg/kg as an intravenous infusion over 30 to 90 minutes every three weeks [see Dosage and Administration (2.3)].
- Extending adjuvant treatment beyond one year is not recommended [see Adverse Reactions (6.1)].

Metastatic Treatment, Breast Cancer

 Administer TRAZIMERA, alone or in combination with paclitaxel, at an initial dose of 4 mg/kg as a 90-minute intravenous infusion followed by subsequent once weekly doses of 2 mg/kg as 30-minute intravenous infusions until disease progression.

Metastatic Gastric Cancer

 Administer TRAZIMERA at an initial dose of 8 mg/kg as a 90-minute intravenous infusion followed by subsequent doses of 6 mg/kg as an intravenous infusion over 30 to 90 minutes every three weeks until disease progression [see Dosage and Administration (2.3)].

2.3 Important Dosing Considerations

If the patient has missed a dose of TRAZIMERA by one week or less, then the usual maintenance dose (weekly schedule: 2 mg/kg; three-weekly schedule: 6 mg/kg) should be administered as soon as possible. Do not wait until the next planned cycle. Subsequent TRAZIMERA maintenance doses should be administered 7 days or 21 days later according to the weekly or three-weekly schedules, respectively.

If the patient has missed a dose of TRAZIMERA by more than one week, a re-loading dose of TRAZIMERA should be administered over approximately 90 minutes (weekly schedule: 4 mg/kg; three-weekly schedule: 8 mg/kg) as soon as possible. Subsequent TRAZIMERA maintenance doses (weekly schedule: 2 mg/kg; three-weekly schedule 6 mg/kg) should be administered 7 days or 21 days later according to the weekly or three-weekly schedules, respectively.

Infusion Reactions

[See Boxed Warning, Warnings and Precautions (5.2)]

- Decrease the rate of infusion for mild or moderate infusion reactions
- Interrupt the infusion in patients with dyspnea or clinically significant hypotension
- Discontinue TRAZIMERA for severe or life-threatening infusion reactions.

<u>Cardiomyopathy</u>

[See Boxed Warning, Warnings and Precautions (5.1)]

Assess left ventricular ejection fraction (LVEF) prior to initiation of TRAZIMERA and at

regular intervals during treatment. Withhold TRAZIMERA dosing for at least 4 weeks for either of the following:

- ≥ 16% absolute decrease in LVEF from pre-treatment values
- LVEF below institutional limits of normal and ≥ 10% absolute decrease in LVEF from pretreatment values.

TRAZIMERA may be resumed if, within 4 to 8 weeks, the LVEF returns to normal limits and the absolute decrease from baseline is $\leq 15\%$.

Permanently discontinue TRAZIMERA for a persistent (> 8 weeks) LVEF decline or for suspension of TRAZIMERA dosing on more than 3 occasions for cardiomyopathy.

2.4 Preparation for Administration

To prevent medication errors, it is important to check the vial labels to ensure that the drug being prepared and administered is TRAZIMERA (trastuzumab-qyyp) and not adotrastuzumab emtansine.

420 mg Multiple-Dose Vial

Reconstitution

Reconstitute each 420 mg vial of TRAZIMERA with 20 mL of Bacteriostatic Water for Injection (BWFI) containing 1.1% benzyl alcohol as a preservative to yield a multiple-dose solution containing 21 mg/mL trastuzumab-qyyp that delivers 20 mL (420 mg trastuzumab-qyyp). In patients with known hypersensitivity to benzyl alcohol, reconstitute with 20 mL of Sterile Water for Injection (SWFI) without preservative to yield a single use solution.

Use appropriate aseptic technique when performing the following reconstitution steps:

- Using a sterile syringe, slowly inject the 20 mL of diluent into the vial containing the lyophilized powder of TRAZIMERA, which has a cake-like appearance. The reconstituted vial yields a solution for multiple-dose use, containing 21 mg/mL trastuzumab-qyyp.
- Swirl the vial gently to aid reconstitution. **DO NOT SHAKE.**
- Slight foaming of the product may be present upon reconstitution. Allow the vial to stand undisturbed for approximately 5 minutes.
- Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration, whenever solution and container permit. Inspect visually for particulates and discoloration. The solution should be free of visible particulates, clear to slightly opalescent and colorless to pale yellow-brown.
- <u>Store reconstituted TRAZIMERA in the refrigerator at 2°C to 8°C (36°F to 46°F)</u>; discard unused TRAZIMERA after 28 days. <u>If TRAZIMERA is reconstituted with SWFI</u> without preservative, use immediately and discard any unused portion. **Do not freeze.**

Dilution

- Determine the dose (mg) of TRAZIMERA [see Dosage and Administration (2.2)].
- Calculate the volume of the 21 mg/mL reconstituted TRAZIMERA solution needed.
- Withdraw this amount from the vial and add it to an infusion bag containing 250 mL of 0.9% Sodium Chloride Injection, USP. DO NOT USE DEXTROSE (5%) SOLUTION.
- Gently invert the bag to mix the solution.

• The solution of TRAZIMERA for infusion diluted in polyvinylchloride, polyethylene, polypropylene, or ethylene vinyl acetate bags or glass intravenous bottles containing 0.9% Sodium Chloride Injection, USP, should be stored at 2°C to 8°C (36°F to 46°F) for no more than 24 hours prior to use. **Do not freeze.**

150 mg Single-Dose Vial

Reconstitution

Reconstitute each 150 mg vial of TRAZIMERA with 7.4 mL of Sterile Water for Injection (SWFI) (not supplied) to yield a single-dose solution containing 21 mg/mL trastuzumabgyyp that delivers 7.15 mL (150 mg trastuzumab-gyyp).

Use appropriate aseptic technique when performing the following reconstitution steps:

- Using a sterile syringe, slowly inject 7.4 mL of SWFI (not supplied) into the vial containing the lyophilized powder of TRAZIMERA, which has a cake-like appearance. The reconstituted vial yields a solution for single-dose use, containing 21 mg/mL trastuzumab-qyyp.
- Swirl the vial gently to aid reconstitution. DO NOT SHAKE.
- Slight foaming of the product may be present upon reconstitution. Allow the vial to stand undisturbed for approximately 5 minutes.
- Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration, whenever solution and container permit. Inspect visually for particulates and discoloration. The solution should be free of visible particulates, clear to slightly opalescent and colorless to pale yellow-brown.
- Use the TRAZIMERA solution immediately following reconstitution with SWFI, as it contains no preservative and is intended for single-dose only. If not used immediately, store the reconstituted TRAZIMERA solution for up to 24 hours at 2°C to 8°C (36°F to 46°F); discard unused TRAZIMERA after 24 hours. **Do not freeze**.

Dilution

- Determine the dose (mg) of TRAZIMERA [see Dosage and Administration (2.2)].
- Calculate the volume of the 21 mg/mL reconstituted TRAZIMERA solution needed.
- Withdraw this amount from the vial and add it to an infusion bag containing 250 mL of 0.9% Sodium Chloride Injection, USP. DO NOT USE DEXTROSE (5%)
 SOLUTION.
- Gently invert the bag to mix the solution.
- The solution of TRAZIMERA for infusion diluted in polyvinylchloride, polyethylene, polypropylene, or ethylene vinyl acetate bags or glass intravenous bottles containing 0.9% Sodium Chloride Injection, USP, should be stored at 2°C to 8°C (36°F to 46°F) for no more than 24 hours prior to use. Discard after 24 hours. This storage time is additional to the time allowed for the reconstituted vials. **Do not freeze**.

3 DOSAGE FORMS AND STRENGTHS

- For injection: 150 mg of TRAZIMERA as a white lyophilized powder in a single-dose vial.
- For injection: 420 mg of TRAZIMERA as a white lyophilized powder in a multiple-dose vial.

4 CONTRAINDICATIONS

None.

5 WARNINGS AND PRECAUTIONS

5.1 Cardiomyopathy

Trastuzumab products can cause left ventricular cardiac dysfunction, arrhythmias, hypertension, disabling cardiac failure, cardiomyopathy, and cardiac death [see Boxed Warning]. Trastuzumab products can also cause asymptomatic decline in left ventricular ejection fraction (LVEF).

There is a 4 to 6 fold increase in the incidence of symptomatic myocardial dysfunction among patients receiving trastuzumab products as a single agent or in combination therapy compared with those not receiving trastuzumab products. The highest absolute incidence occurs when a trastuzumab product is administered with an anthracycline.

Withhold TRAZIMERA for \geq 16% absolute decrease in LVEF from pre-treatment values or an LVEF value below institutional limits of normal and \geq 10% absolute decrease in LVEF from pretreatment values [see Dosage and Administration (2.3)]. The safety of continuation or resumption of TRAZIMERA in patients with trastuzumab productinduced left ventricular cardiac dysfunction has not been studied.

Patients who receive anthracycline after stopping TRAZIMERA may also be at increased risk of cardiac dysfunction [see Drug Interactions (7) and Clinical Pharmacology (12.3)].

Cardiac Monitoring

Conduct thorough cardiac assessment, including history, physical examination, and determination of LVEF by echocardiogram or MUGA scan. The following schedule is recommended:

- Baseline LVEF measurement immediately prior to initiation of TRAZIMERA
- LVEF measurements every 3 months during and upon completion of TRAZIMERA
- Repeat LVEF measurement at 4 week intervals if TRAZIMERA is withheld for significant left ventricular cardiac dysfunction [see Dosage and Administration (2.3)]
- LVEF measurements every 6 months for at least 2 years following completion of TRAZIMERA as a component of adjuvant therapy.

In Study 1, 15% (158/1031) of patients discontinued trastuzumab due to clinical evidence of myocardial dysfunction or significant decline in LVEF after a median follow-up duration of 8.7 years in the AC-TH (anthracycline, cyclophosphamide, paclitaxel, and trastuzumab) arm. In Study 3 (one-year trastuzumab treatment), the number of patients who discontinued trastuzumab due to cardiac toxicity at 12.6 months median duration of follow-up was 2.6% (44/1678). In Study 4, a total of 2.9% (31/1056) of patients in the TCH (docetaxel, carboplatin, trastuzumab) arm (1.5% during the chemotherapy phase and 1.4% during the monotherapy phase) and 5.7% (61/1068) of patients in the AC-TH arm (1.5% during the chemotherapy phase and 4.2% during the monotherapy phase) discontinued trastuzumab due to cardiac toxicity.

Among 64 patients receiving adjuvant chemotherapy (Studies 1 and 2) who developed congestive heart failure (CHF), one patient died of cardiomyopathy, one patient died suddenly without documented etiology, and 33 patients were receiving cardiac

medication at last follow-up. Approximately 24% of the surviving patients had recovery to a normal LVEF (defined as ≥50%) and no symptoms on continuing medical management at the time of last follow-up. Incidence of congestive heart failure is presented in Table 1. The safety of continuation or resumption of TRAZIMERA in patients with trastuzumab product-induced left ventricular cardiac dysfunction has not been studied.

Table 1 Incidence of Congestive Heart Failure in Adjuvant
Breast Cancer Studies

		Incidence of CHF			
Study	Regimen	Trastuzumab	Control		
1 & 2*	AC [†] →Paclitaxel+Trastuzumab	3.2%	1.3%		
1 Q Z	AC! Facilitaxei+ II astuzui II ab	(64/2000) [‡]	(21/1655)		
3 §	Chemo → Trastuzumab	2% (30/1678)	0.3%		
23	Chemo - mastuzumab	2 /0 (30/10/0)	(5/1708)		
1	AC [†] → Docetaxel+Trastuzumab	2% (20/1068)	0.3%		
4	AC Docetaxei Trastazarilab	270 (20/1000)	(3/1050)		
1	Docataval I Carbo I Tractuzumah	0.4% (4/1056)	0.3%		
4	Docetaxel+Carbo+Trastuzumab	0.4 /0 (4/1030)	(3/1050)		

^{*} Median follow-up duration for Studies 1 and 2 combined was 8.3 years in the AC→TH arm.

In Study 3 (one-year trastuzumab treatment), at a median follow-up duration of 8 years, the incidence of severe CHF (NYHA III & IV) was 0.8%, and the rate of mild symptomatic and asymptomatic left ventricular dysfunction was 4.6%.

Table 2 Incidence of Cardiac Dysfunction* in Metastatic Breast Cancer Studies

		Incidence			
		NYHA I	I-IV	NYHA I	II-IV
Study	Event	Trastuzuma	b ControlT	rastuzuma	b Control
5 (AC) [†]	Cardiac Dysfunction	28%	7%	19%	3%
5 (paclitaxe	Cardiac l) Dysfunction	11%	1%	4%	1%
6	Cardiac Dysfunction [‡]	7%	N/A	5%	N/A

^{*} Congestive heart failure or significant asymptomatic decrease in LVEF.

In Study 4, the incidence of NCI-CTC Grade 3/4 cardiac ischemia/infarction was higher in the trastuzumab containing regimens (AC-TH: 0.3% (3/1068) and TCH: 0.2% (2/1056))

[†] Anthracycline (doxorubicin) and cyclophosphamide.

[‡] Includes 1 patient with fatal cardiomyopathy and 1 patient with sudden death without documented etiology.

[§] Includes NYHA II–IV and cardiac death at 12.6 months median duration of follow-up in the one-year trastuzumab arm.

[†] Anthracycline (doxorubicin or epirubicin) and cyclophosphamide.

[‡] Includes 1 patient with fatal cardiomyopathy.

as compared to none in AC-T.

5.2 Infusion Reactions

Infusion reactions consist of a symptom complex characterized by fever and chills, and on occasion included nausea, vomiting, pain (in some cases at tumor sites), headache, dizziness, dyspnea, hypotension, rash, and asthenia [see Adverse Reactions (6.1)].

In post-marketing reports, serious and fatal infusion reactions have been reported. Severe reactions, which include bronchospasm, anaphylaxis, angioedema, hypoxia, and severe hypotension, were usually reported during or immediately following the initial infusion. However, the onset and clinical course were variable, including progressive worsening, initial improvement followed by clinical deterioration, or delayed post-infusion events with rapid clinical deterioration. For fatal events, death occurred within hours to days following a serious infusion reaction.

Interrupt TRAZIMERA infusion in all patients experiencing dyspnea, clinically significant hypotension, and intervention of medical therapy administered (which may include epinephrine, corticosteroids, diphenhydramine, bronchodilators, and oxygen). Patients should be evaluated and carefully monitored until complete resolution of signs and symptoms. Permanent discontinuation should be strongly considered in all patients with severe infusion reactions.

There are no data regarding the most appropriate method of identification of patients who may safely be retreated with trastuzumab products after experiencing a severe infusion reaction. Prior to resumption of trastuzumab infusion, the majority of patients who experienced a severe infusion reaction were pre-medicated with antihistamines and/or corticosteroids. While some patients tolerated trastuzumab infusions, others had recurrent severe infusion reactions despite pre-medications.

5.3 Embryo-Fetal Toxicity

Trastuzumab products can cause fetal harm when administered to a pregnant woman. In post-marketing reports, use of trastuzumab during pregnancy resulted in cases of oligohydramnios and oligohydramnios sequence manifesting as pulmonary hypoplasia, skeletal abnormalities, and neonatal death.

Verify the pregnancy status of females of reproductive potential prior to the initiation of TRAZIMERA. Advise pregnant women and females of reproductive potential that exposure to TRAZIMERA during pregnancy or within 7 months prior to conception can result in fetal harm. Advise females of reproductive potential to use effective contraception during treatment and for 7 months following the last dose of TRAZIMERA [see Use in Specific Populations (8.1, 8.3) and Clinical Pharmacology (12.3)].

5.4 Pulmonary Toxicity

Trastuzumab product use can result in serious and fatal pulmonary toxicity. Pulmonary toxicity includes dyspnea, interstitial pneumonitis, pulmonary infiltrates, pleural effusions, non-cardiogenic pulmonary edema, pulmonary insufficiency and hypoxia, acute respiratory distress syndrome, and pulmonary fibrosis. Such events can occur as sequelae of infusion reactions [see Warnings and Precautions (5.2)]. Patients with symptomatic intrinsic lung disease or with extensive tumor involvement of the lungs, resulting in dyspnea at rest, appear to have more severe toxicity.

5.5 Exacerbation of Chemotherapy-Induced Neutropenia

In randomized, controlled clinical trials, the per-patient incidences of NCI-CTC Grade 3 to 4 neutropenia and of febrile neutropenia were higher in patients receiving trastuzumab in combination with myelosuppressive chemotherapy as compared to those who received chemotherapy alone. The incidence of septic death was similar among patients who received trastuzumab and those who did not [see Adverse Reactions (6.1)].

6 ADVERSE REACTIONS

The following adverse reactions are discussed in greater detail in other sections of the label:

- Cardiomyopathy [see Warnings and Precautions (5.1)]
- Infusion Reactions [see Warnings and Precautions (5.2)]
- Embryo-Fetal Toxicity [see Warnings and Precautions (5.3)]
- Pulmonary Toxicity [see Warnings and Precautions (5.4)]
- Exacerbation of Chemotherapy-Induced Neutropenia [see Warnings and Precautions (5.5)]

The most common adverse reactions in patients receiving trastuzumab products in the adjuvant and metastatic breast cancer setting are fever, nausea, vomiting, infusion reactions, diarrhea, infections, increased cough, headache, fatigue, dyspnea, rash, neutropenia, anemia, and myalgia. Adverse reactions requiring interruption or discontinuation of trastuzumab product treatment include CHF, significant decline in left ventricular cardiac function, severe infusion reactions, and pulmonary toxicity [see Dosage and Administration (2.3)].

In the metastatic gastric cancer setting, the most common adverse reactions ($\geq 10\%$) that were increased ($\geq 5\%$ difference) in patients receiving trastuzumab as compared to patients receiving chemotherapy alone were neutropenia, diarrhea, fatigue, anemia, stomatitis, weight loss, upper respiratory tract infections, fever, thrombocytopenia, mucosal inflammation, nasopharyngitis, and dysgeusia. The most common adverse reactions which resulted in discontinuation of trastuzumab treatment in the absence of disease progression were infection, diarrhea, and febrile neutropenia.

6.1 Clinical Trials Experience

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to rates in the clinical trials of another drug and may not reflect the rates observed in practice.

<u>Adjuvant Breast Cancer Studies</u>

The data below reflect exposure to one-year trastuzumab therapy across three randomized, open-label studies, Studies 1, 2, and 3, with (n = 3678) or without (n = 3363) trastuzumab in the adjuvant treatment of breast cancer.

The data summarized in Table 3 below, from Study 3, reflect exposure to trastuzumab in 1678 patients; the median treatment duration was 51 weeks and median number of infusions was 18. Among the 3386 patients enrolled in the observation and one-year trastuzumab arms of Study 3 at a median duration of follow-up of 12.6 months in the trastuzumab arm, the median age was 49 years (range: 21 to 80 years), 83% of

Table 3 Adverse Reactions for Study 3*, All Grades†

	One Year	
	Trastuzumab	Observation
Adverse Reaction	(n = 1678)	(n = 1708)
Cardiac		
Hypertension	64 (4%)	35 (2%)
Dizziness	60 (4%)	29 (2%)
Ejection Fraction Decreased	58 (3.5%)	11 (0.6%)
Palpitations	48 (3%)	12 (0.7%)
Cardiac Arrhythmias [‡]	40 (3%)	17 (1%)
Cardiac Failure Congestive	30 (2%)	5 (0.3%)
Cardiac Failure	9 (0.5%)	4 (0.2%)
Cardiac Disorder	5 (0.3%)	0 (0%)
Ventricular Dysfunction	4 (0.2%)	0 (0%)
Respiratory Thoracic Medias		3 (0/0)
Cough	81 (5%)	34 (2%)
Influenza	70 (4%)	9 (0.5%)
Dyspnea	57 (3%)	26 (2%)
URI	46 (3%)	20 (1%)
Rhinitis	36 (2%)	6 (0.4%)
Pharyngolaryngeal Pain	32 (2%)	8 (0.5%)
Sinusitis	26 (2%)	5 (0.3%)
Epistaxis	25 (2%)	1 (0.06%)
<u> </u>		
Pulmonary Hypertension Interstitial Pneumonitis	4 (0.2%)	0 (0%)
	4 (0.2%)	0 (0%)
Gastrointestinal Disorders	122 /70/\	16 (10/)
Diarrhea	123 (7%)	16 (1%)
Nausea	108 (6%)	19 (1%)
Vomiting	58 (3.5%)	10 (0.6%)
Constipation	33 (2%)	17 (1%)
Dyspepsia	30 (2%)	9 (0.5%)
Upper Abdominal Pain	29 (2%)	15 (1%)
Musculoskeletal & Connectiv		
Arthralgia	137 (8%)	98 (6%)
Back Pain	91 (5%)	58 (3%)
Myalgia	63 (4%)	17 (1%)
Bone Pain	49 (3%)	26 (2%)
Muscle Spasm	46 (3%)	3 (0.2%)
Nervous System Disorders		
Headache	162 (10%)	49 (3%)
Paraesthesia	29 (2%)	11 (0.6%)
Skin & Subcutaneous Tissue	e Disorders	

ь .	70 (40/)	10 (0 60()
Rash	70 (4%)	10 (0.6%)
Nail Disorders	43 (2%)	0 (0%)
Pruritus	40 (2%)	10 (0.6%)
General Disorders		
Pyrexia	100 (6%)	6 (0.4%)
Edema Peripheral	79 (5%)	37 (2%)
Chills	85 (5%)	0 (0%)
Asthenia	75 (4.5%)	30 (2%)
Influenza-like Illness	40 (2%)	3 (0.2%)
Sudden Death	1 (0.06%)	0 (0%)
Infections		
Nasopharyngitis	135 (8%)	43 (3%)
UTI	39 (3%)	13 (0.8%)
Immune System Disorders		
Hypersensitivity	10 (0.6%)	1 (0.06%)
Autoimmune Thyroiditis	4 (0.3%)	0 (0%)

^{*} Median follow-up duration of 12.6 months in the one-year trastuzumab treatment arm.

In Study 3, a comparison of 3-weekly trastuzumab treatment for two years versus one year was also performed. The rate of asymptomatic cardiac dysfunction was increased in the 2-year trastuzumab treatment arm (8.1% versus 4.6% in the one-year trastuzumab treatment arm). More patients experienced at least one adverse reaction of Grade 3 or higher in the 2-year trastuzumab treatment arm (20.4%) compared with the one-year trastuzumab treatment arm (16.3%).

The safety data from Studies 1 and 2 were obtained from 3655 patients, of whom 2000 received trastuzumab; the median treatment duration was 51 weeks. The median age was 49 years (range: 24 to 80); 84% of patients were White, 7% Black, 4% Hispanic, and 3% Asian.

In Study 1, only Grade 3 to 5 adverse events, treatment-related Grade 2 events, and Grade 2 to 5 dyspnea were collected during and for up to 3 months following protocol-specified treatment. The following non-cardiac adverse reactions of Grade 2 to 5 occurred at an incidence of at least 2% greater among patients receiving trastuzumab plus chemotherapy as compared to chemotherapy alone: fatigue (29.5% vs. 22.4%), infection (24.0% vs. 12.8%), hot flashes (17.1% vs. 15.0%), anemia (12.3% vs. 6.7%), dyspnea (11.8% vs. 4.6%), rash/desquamation (10.9% vs. 7.6%), leukopenia (10.5% vs. 8.4%), neutropenia (6.4% vs. 4.3%), headache (6.2% vs. 3.8%), pain (5.5% vs. 3.0%), edema (4.7% vs. 2.7%), and insomnia (4.3% vs. 1.5%). The majority of these events were Grade 2 in severity.

In Study 2, data collection was limited to the following investigator-attributed treatment-related adverse reactions: NCI-CTC Grade 4 and 5 hematologic toxicities, Grade 3 to 5 non-hematologic toxicities, selected Grade 2 to 5 toxicities associated with taxanes (myalgia, arthralgias, nail changes, motor neuropathy, and sensory neuropathy) and Grade 1 to 5 cardiac toxicities occurring during chemotherapy and/or trastuzumab

[†] The incidence of Grade 3 or higher adverse reactions was <1% in both arms for each listed term.

[‡] Higher level grouping term.

treatment. The following non-cardiac adverse reactions of Grade 2 to 5 occurred at an incidence of at least 2% greater among patients receiving trastuzumab plus chemotherapy as compared to chemotherapy alone: arthralgia (12.2% vs. 9.1%), nail changes (11.5% vs. 6.8%), dyspnea (2.4% vs. 0.2%), and diarrhea (2.2% vs. 0%). The majority of these events were Grade 2 in severity.

Safety data from Study 4 reflect exposure to trastuzumab as part of an adjuvant treatment regimen from 2124 patients receiving at least one dose of study treatment [AC-TH: n=1068; TCH: n=1056]. The overall median treatment duration was 54 weeks in both the AC-TH and TCH arms. The median number of infusions was 26 in the AC-TH arm and 30 in the TCH arm, including weekly infusions during the chemotherapy phase and every three week dosing in the monotherapy period. Among these patients, the median age was 49 years (range 22 to 74 years). In Study 4, the toxicity profile was similar to that reported in Studies 1, 2, and 3 with the exception of a low incidence of CHF in the TCH arm.

Metastatic Breast Cancer Studies

The data below reflect exposure to trastuzumab in one randomized, open-label study, Study 5, of chemotherapy with (n = 235) or without (n = 234) trastuzumab in patients with metastatic breast cancer, and one single-arm study (Study 6; n = 222) in patients with metastatic breast cancer. Data in Table 4 are based on Studies 5 and 6.

Among the 464 patients treated in Study 5, the median age was 52 years (range: 25 to 77 years). Eighty-nine percent were White, 5% Black, 1% Asian, and 5% other racial/ethnic groups. All patients received 4 mg/kg initial dose of trastuzumab followed by 2 mg/kg weekly. The percentages of patients who received trastuzumab treatment for \geq 6 months and \geq 12 months were 58% and 9%, respectively.

Among the 352 patients treated in single agent studies (213 patients from Study 6), the median age was 50 years (range 28 to 86 years), 86% were White, 3% were Black, 3% were Asian, and 8% in other racial/ethnic groups. Most of the patients received 4 mg/kg initial dose of trastuzumab followed by 2 mg/kg weekly. The percentages of patients who received trastuzumab treatment for \geq 6 months and \geq 12 months were 31% and 16%, respectively.

Table 4 Per-Patient Incidence of Adverse Reactions Occurring in ≥5% of Patients in Uncontrolled Studies or at Increased Incidence in the Trastuzumab Arm (Studies 5 and 6)

	Single Agent* n = 352	Trastuzumab + Paclitaxel n = 91		Trastuzumab + AC [†] n = 143	AC [†] Alone n = 135
Body as a Whole					
Pain	47%	61%	62%	57%	42%
Asthenia	42%	62%	57%	54%	55%
Fever	36%	49%	23%	56%	34%
Chills	32%	41%	4%	35%	11%
Headache	26%	36%	28%	44%	31%

Abdominal pain	22%	34%	22%	23%	18%
Back pain	22%	34%	30%	27%	15%
Infection	20%	47%	27%	47%	31%
Flu syndrome	10%	12%	5%	12%	6%
Accidental injury	6%	13%	3%	9%	4%
Allergic reaction	3%	8%	2%	4%	2%
Cardiovascular Tachycardia	5%	12%	4%	10%	5%
Congestive	J /0	12 /0	4 /0	10 /0	J /0
heart failure	7%	11%	1%	28%	7%
Digestive					
Nausea	33%	51%	9%	76%	77%
Diarrhea	25%	45%	29%	45%	26%
Vomiting	23%	37%	28%	53%	49%
Nausea and vomiting	8%	14%	11%	18%	9%
Anorexia	14%	24%	16%	31%	26%
Heme & Lymphatic					
Anemia	4%	14%	9%	36%	26%
Leukopenia	3%	24%	17%	52%	34%
Metabolic					
Peripheral edema	10%	22%	20%	20%	17%
Edema	8%	10%	8%	11%	5%
Musculoskeletal					
Bone pain	7%	24%	18%	7%	7%
Arthralgia	6%	37%	21%	8%	9%
Nervous					
Insomnia	14%	25%	13%	29%	15%
Dizziness	13%	22%	24%	24%	18%
Paresthesia	9%	48%	39%	17%	11%
Depression	6%	12%	13%	20%	12%
Peripheral neuritis	2%	23%	16%	2%	2%
Neuropathy	1%	13%	5%	4%	4%
Respiratory	_,0	10,0	3,0	. , 0	.,0
Cough	26%	41%	22%	43%	29%
increased	220/	270/	260/	420/	250/
Dyspnea	22%	27%	26%	42%	25%
Rhinitis	14%	22%	5%	22%	16%
Pharyngitis	12%	22%	14%	30%	18%
Sinusitis	9%	21%	7%	13%	6%

Skin					
Rash	18%	38%	18%	27%	17%
Herpes simplex	2%	12%	3%	7%	9%
Acne	2%	11%	3%	3%	< 1%
Urogenital					
Urinary tract infection	5%	18%	14%	13%	7%

^{*} Data for trastuzumab single agent were from 4 studies, including 213 patients from Study 6.

Metastatic Gastric Cancer

The data below are based on the exposure of 294 patients to trastuzumab in combination with a fluoropyrimidine (capecitabine or 5-FU) and cisplatin (Study 7). In the trastuzumab plus chemotherapy arm, the initial dose of trastuzumab 8 mg/kg was administered on Day 1 (prior to chemotherapy) followed by 6 mg/kg every 21 days until disease progression. Cisplatin was administered at 80 mg/m² on Day 1 and the fluoropyrimidine was administered as either capecitabine 1000 mg/m² orally twice a day on Days 1 to 14 or 5-fluorouracil 800 mg/m²/day as a continuous intravenous infusion Days 1 through 5. Chemotherapy was administered for six 21-day cycles. Median duration of trastuzumab treatment was 21 weeks; median number of trastuzumab infusions administered was eight.

Table 5 Study 7: Per-Patient Incidence of Adverse
Reactions of All Grades (Incidence ≥ 5% between Arms)
or Grade 3/4 (Incidence > 1% between Arms) and Higher
Incidence in Trastuzumab Arm

	Trastuzumab FC (N = 294) N (%)		FC (N = 290) N (%)	
Body System/Adverse	All	Grades	All	Grades
Event	Grades	3/4	Grades	3/4
Investigations				
Neutropenia	230 (78)	101 (34)	212 (73)	83 (29)
Hypokalemia	83 (28)	28 (10)	69 (24)	16 (6)
Anemia	81 (28)	36 (12)	61 (21)	30 (10)
Thrombocytopenia	47 (16)	14 (5)	33 (11)	8 (3)
Blood and Lymphatic Syste	em Disord	ers		
Febrile Neutropenia	_	15 (5)	_	8 (3)
Gastrointestinal Disorders				
Diarrhea	109 (37)	27 (9)	80 (28)	11 (4)
Stomatitis	72 (24)	2 (1)	43 (15)	6 (2)
Dysphagia	19 (6)	7 (2)	10 (3)	1 (≤ 1)
Body as a Whole				
Fatigue	102 (35)	12 (4)	82 (28)	7 (2)

[†] Anthracycline (doxorubicin or epirubicin) and cyclophosphamide.

i					
54 (18)	3 (1)	36 (12)	0 (0)		
37 (13)	6 (2)	18 (6)	2 (1)		
23 (8)	1 (≤ 1)	0 (0)	0 (0)		
isorders					
69 (23)	6 (2)	40 (14)	7 (2)		
l					
56 (19)	0 (0)	29 (10)	0 (0)		
37 (13)	0 (0)	17 (6)	0 (0)		
^S					
53 (18)	8 (3)	42 (15)	5 (2)		
Nervous System Disorders					
28 (10)	0 (0)	14 (5)	0 (0)		
	37 (13) 23 (8) Disorders 69 (23) 56 (19) 37 (13) 5 53 (18)	37 (13) 6 (2) 23 (8) 1 (≤ 1) Disorders 69 (23) 6 (2) 56 (19) 0 (0) 37 (13) 0 (0) 5 53 (18) 8 (3)	$37 (13)$ $6 (2)$ $18 (6)$ $23 (8)$ $1 (\le 1)$ $0 (0)$ Disorders $69 (23)$ $6 (2)$ $40 (14)$ $56 (19)$ $0 (0)$ $29 (10)$ $37 (13)$ $0 (0)$ $17 (6)$ $53 (18)$ $8 (3)$ $42 (15)$		

The following subsections provide additional detail regarding adverse reactions observed in clinical trials of adjuvant breast cancer, metastatic breast cancer, metastatic gastric cancer, or post-marketing experience.

Cardiomyopathy

Serial measurement of cardiac function (LVEF) was obtained in clinical trials in the adjuvant treatment of breast cancer. In Study 3, the median duration of follow-up was 12.6 months (12.4 months in the observation arm; 12.6 months in the 1-year trastuzumab arm); and in Studies 1 and 2, 7.9 years in the AC-T arm, 8.3 years in the AC-TH arm. In Studies 1 and 2, 6% of all randomized patients with post-AC LVEF evaluation were not permitted to initiate trastuzumab following completion of AC chemotherapy due to cardiac dysfunction (LVEF < LLN or ≥ 16 point decline in LVEF from baseline to end of AC). Following initiation of trastuzumab therapy, the incidence of new-onset dose-limiting myocardial dysfunction was higher among patients receiving trastuzumab and paclitaxel as compared to those receiving paclitaxel alone in Studies 1 and 2, and in patients receiving one-year trastuzumab monotherapy compared to observation in Study 3 (see Table 6, Figures 1 and 2). The per-patient incidence of newonset cardiac dysfunction, as measured by LVEF, remained similar when compared to the analysis performed at a median follow-up of 2.0 years in the AC-TH arm. This analysis also showed evidence of reversibility of left ventricular dysfunction, with 64.5% of patients who experienced symptomatic CHF in the AC-TH group being asymptomatic at latest follow-up, and 90.3% having full or partial LVEF recovery.

Table 6* Per-patient Incidence of New Onset Myocardial Dysfunction (by LVEF) Studies 1, 2, 3 and 4

	LVEF <50% and Absolute Decrease from Baseline			Absolute LVEF Decrease	
		LVEF < ≥ 10% ≥ 16% 50% decrease decrease			≥ 20%
Studies 1 & 2 ^{†,‡}					

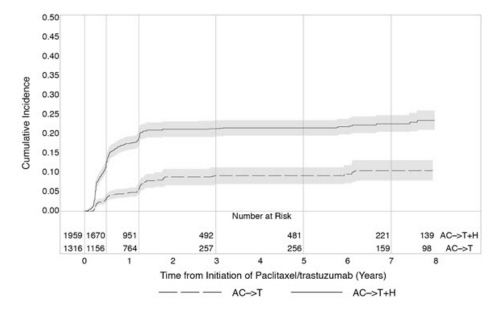
AC→TH	23.1%	18.5%	11.2%	37.9%	8.9%		
(n = 1856)	(428)	(344)	(208)	(703)	(166)		
AC→T	11.7%	7.0%	3.0%	22.1%	3.4%		
(n = 1170)	(137)	(82)	(35)	(259)	(40)		
Study 3§							
Trastuzumab	8.6%	7.0%	3.8%	22.4%	3.5%		
(n = 1678)	(144)	(118)	(64)	(376)	(59)		
Observation	2.7%	2.0%	1.2%	11.9%	1.2%		
(n = 1708)	(46)	(35)	(20)	(204)	(21)		
Study 4 [¶]							
TCH	8.5%	5.9%	3.3%	34.5%	6.3%		
(n = 1056)	(90)	(62)	(35)	(364)	(67)		
AC→TH	17%	13.3%	9.8%	44.3%	13.2%		
(n = 1068)	(182)	(142)	(105)	(473)	(141)		
AC→T	9.5%	6.6%	3.3%	34%	5.5%		
(n = 1050)	(100)	(69)	(35)	(357)	(58)		
* For Ctudioc 1	Ear Studies 1, 2 and 2, events are solunted from the haginning of						

^{*} For Studies 1, 2 and 3, events are counted from the beginning of trastuzumab treatment. For Study 4, events are counted from the date of randomization.

- † Studies 1 and 2 regimens: doxorubicin and cyclophosphamide followed by paclitaxel (AC→T) or paclitaxel plus trastuzumab (AC→TH).
- ‡ Median duration of follow-up for Studies 1 and 2 combined was 8.3 years in the AC→TH arm.
- § Median follow-up duration of 12.6 months in the one-year trastuzumab treatment arm.
- ¶ Study 4 regimens: doxorubicin and cyclophosphamide followed by docetaxel (AC→T) or docetaxel plus trastuzumab (AC→TH); docetaxel and carboplatin plus trastuzumab (TCH).

Figure 1

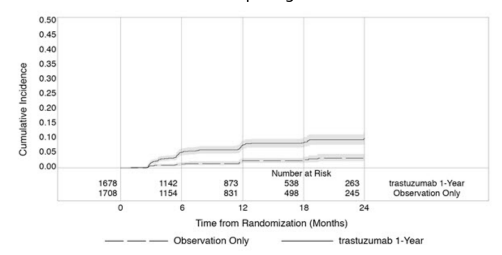
Studies 1 and 2: Cumulative Incidence of Time to First LVEF Decline of ≥ 10 Percentage Points from Baseline and to Below 50% with Death as a Competing Risk Event



Time 0 is initiation of paclitaxel or trastuzumab + paclitaxel therapy.

Figure 2

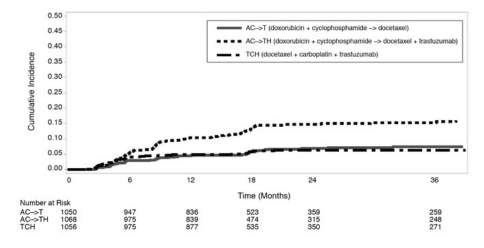
Study 3: Cumulative Incidence of Time to First LVEF Decline of ≥ 10 Percentage Points from Baseline and to Below 50% with Death as a Competing Risk Event



Time 0 is the date of randomization.

Figure 3

Study 4: Cumulative Incidence of Time to First LVEF Decline of ≥10 Percentage Points from Baseline and to Below 50% with Death as a Competing Risk Event



Time 0 is the date of randomization.

The incidence of treatment emergent congestive heart failure among patients in the metastatic breast cancer trials was classified for severity using the New York Heart Association classification system (I–IV, where IV is the most severe level of cardiac failure) (see Table 2). In the metastatic breast cancer trials, the probability of cardiac dysfunction was highest in patients who received trastuzumab concurrently with anthracyclines.

In Study 7, 5.0% of patients in the trastuzumab plus chemotherapy arm compared to 1.1% of patients in the chemotherapy alone arm had LVEF value below 50% with a \geq 10% absolute decrease in LVEF from pretreatment values.

<u>Infusion Reactions</u>

During the first infusion with trastuzumab, the symptoms most commonly reported were chills and fever, occurring in approximately 40% of patients in clinical trials. Symptoms were treated with acetaminophen, diphenhydramine, and meperidine (with or without reduction in the rate of trastuzumab infusion); permanent discontinuation of trastuzumab for infusion reactions was required in < 1% of patients. Other signs and/or symptoms may include nausea, vomiting, pain (in some cases at tumor sites), rigors, headache, dizziness, dyspnea, hypotension, elevated blood pressure, rash, and asthenia. Infusion reactions occurred in 21% and 35% of patients, and were severe in 1.4% and 9% of patients, on second or subsequent trastuzumab infusions administered as monotherapy or in combination with chemotherapy, respectively. In the postmarketing setting, severe infusion reactions, including hypersensitivity, anaphylaxis, and angioedema have been reported.

Anemia

In randomized controlled clinical trials, the overall incidence of anemia (30% vs. 21% [Study 5]), of selected NCI-CTC Grade 2 to 5 anemia (12.3% vs. 6.7% [Study 1]), and of anemia requiring transfusions (0.1% vs. 0 patients [Study 2]) were increased in patients receiving trastuzumab and chemotherapy compared with those receiving chemotherapy alone. Following the administration of trastuzumab as a single agent (Study 6), the incidence of NCI-CTC Grade 3 anemia was < 1%. In Study 7 (metastatic gastric cancer), on the trastuzumab containing arm as compared to the chemotherapy alone arm, the overall incidence of anemia was 28% compared to 21% and of NCI-CTC Grade 3/4 anemia was 12.2% compared to 10.3%.

Neutropenia

In randomized controlled clinical trials in the adjuvant setting, the incidence of selected NCI-CTC Grade 4 to 5 neutropenia (1.7% vs. 0.8% [Study 2]) and of selected Grade 2 to 5 neutropenia (6.4% vs. 4.3% [Study 1]) were increased in patients receiving trastuzumab and chemotherapy compared with those receiving chemotherapy alone. In a randomized, controlled trial in patients with metastatic breast cancer, the incidences of NCI-CTC Grade 3/4 neutropenia (32% vs. 22%) and of febrile neutropenia (23% vs. 17%) were also increased in patients randomized to trastuzumab in combination with myelosuppressive chemotherapy as compared to chemotherapy alone. In Study 7 (metastatic gastric cancer) on the trastuzumab containing arm as compared to the chemotherapy alone arm, the incidence of NCI-CTC Grade 3/4 neutropenia was 36.8% compared to 28.9%; febrile neutropenia 5.1% compared to 2.8%.

<u>Infection</u>

The overall incidences of infection (46% vs. 30% [Study 5]), of selected NCI-CTC Grade 2 to 5 infection/febrile neutropenia (24.3% vs. 13.4% [Study 1]) and of selected Grade 3 to 5 infection/febrile neutropenia (2.9% vs. 1.4% [Study 2]) were higher in patients receiving trastuzumab and chemotherapy compared with those receiving chemotherapy alone. The most common site of infections in the adjuvant setting involved the upper respiratory tract, skin, and urinary tract.

In Study 4, the overall incidence of infection was higher with the addition of trastuzumab to AC-T but not to TCH [44% (AC-TH), 37% (TCH), 38% (AC-T)]. The incidences of NCI-CTC Grade 3 to 4 infection were similar [25% (AC-TH), 21% (TCH), 23% (AC-T)] across the three arms.

In a randomized, controlled trial in treatment of metastatic breast cancer, the reported incidence of febrile neutropenia was higher (23% vs. 17%) in patients receiving trastuzumab in combination with myelosuppressive chemotherapy as compared to chemotherapy alone.

Pulmonary Toxicity

Adjuvant Breast Cancer

Among women receiving adjuvant therapy for breast cancer, the incidence of selected NCI-CTC Grade 2 to 5 pulmonary toxicity (14.3% vs. 5.4% [Study 1]) and of selected NCI-CTC Grade 3 to 5 pulmonary toxicity and spontaneous reported Grade 2 dyspnea (3.4% vs. 0.9% [Study 2]) was higher in patients receiving trastuzumab and chemotherapy compared with chemotherapy alone. The most common pulmonary toxicity was dyspnea (NCI-CTC Grade 2 to 5: 11.8% vs. 4.6% [Study 1]; NCI-CTC Grade 2 to 5: 2.4% vs. 0.2% [Study 2]).

Pneumonitis/pulmonary infiltrates occurred in 0.7% of patients receiving trastuzumab compared with 0.3% of those receiving chemotherapy alone. Fatal respiratory failure occurred in 3 patients receiving trastuzumab, one as a component of multi-organ system failure, as compared to 1 patient receiving chemotherapy alone.

In Study 3, there were 4 cases of interstitial pneumonitis in the one-year trastuzumab treatment arm compared to none in the observation arm at a median follow-up duration of 12.6 months.

Metastatic Breast Cancer

Among women receiving trastuzumab for treatment of metastatic breast cancer, the incidence of pulmonary toxicity was also increased. Pulmonary adverse events have been reported in the post-marketing experience as part of the symptom complex of infusion reactions. Pulmonary events include bronchospasm, hypoxia, dyspnea, pulmonary infiltrates, pleural effusions, non-cardiogenic pulmonary edema, and acute respiratory distress syndrome. For a detailed description, see *Warnings and Precautions* (5.4).

Thrombosis/Embolism

In 4 randomized, controlled clinical trials, the incidence of thrombotic adverse events was higher in patients receiving trastuzumab and chemotherapy compared to chemotherapy alone in three studies (2.6% vs. 1.5% [Study 1], 2.5% and 3.7% vs. 2.2% [Study 4] and 2.1% vs. 0% [Study 5]).

Diarrhea

Among women receiving adjuvant therapy for breast cancer, the incidence of NCI-CTC Grade 2 to 5 diarrhea (6.7% vs. 5.4% [Study 1]) and of NCI-CTC Grade 3 to 5 diarrhea (2.2% vs. 0% [Study 2]), and of Grade 1 to 4 diarrhea (7% vs. 1% [Study 3; one-year trastuzumab treatment at 12.6 months median duration of follow-up]) were higher in patients receiving trastuzumab as compared to controls. In Study 4, the incidence of Grade 3 to 4 diarrhea was higher [5.7% AC-TH, 5.5% TCH vs. 3.0% AC-T] and of Grade

1 to 4 was higher [51% AC-TH, 63% TCH vs. 43% AC-T] among women receiving trastuzumab. Of patients receiving trastuzumab as a single agent for the treatment of metastatic breast cancer, 25% experienced diarrhea. An increased incidence of diarrhea was observed in patients receiving trastuzumab in combination with chemotherapy for treatment of metastatic breast cancer.

Renal Toxicity

In Study 7 (metastatic gastric cancer) on the trastuzumab-containing arm as compared to the chemotherapy alone arm the incidence of renal impairment was 18% compared to 14.5%. Severe (Grade 3/4) renal failure was 2.7% on the trastuzumab-containing arm compared to 1.7% on the chemotherapy only arm. Treatment discontinuation for renal insufficiency/failure was 2% on the trastuzumab-containing arm and 0.3% on the chemotherapy only arm.

In the post-marketing setting, rare cases of nephrotic syndrome with pathologic evidence of glomerulopathy have been reported. The time to onset ranged from 4 months to approximately 18 months from initiation of trastuzumab therapy. Pathologic findings included membranous glomerulonephritis, focal glomerulosclerosis, and fibrillary glomerulonephritis. Complications included volume overload and congestive heart failure.

6.2 Immunogenicity

As with all therapeutic proteins, there is potential for immunogenicity. The detection of antibody formation is highly dependent on the sensitivity and specificity of the assay. Additionally, the observed incidence of antibody (including neutralizing antibody) positivity in an assay may be influenced by several factors, including assay methodology, sample handling, timing of sample collection, concomitant medications, and underlying disease. For these reasons, comparison of the incidence of antibodies in the studies described below with the incidence of antibodies in other studies or to other trastuzumab products may be misleading.

Among 903 women with metastatic breast cancer, human anti-human antibody (HAHA) to trastuzumab was detected in one patient using an enzyme-linked immunosorbent assay (ELISA). This patient did not experience an allergic reaction. Samples for assessment of HAHA were not collected in studies of adjuvant breast cancer.

6.3 Post-Marketing Experience

The following adverse reactions have been identified during post-approval use of trastuzumab. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

- Infusion reaction [see Warnings and Precautions (5.2)]
- Oligohydramnios or oligohydramnios sequence, including pulmonary hypoplasia, skeletal abnormalities, and neonatal death [see Warnings and Precautions (5.3)]
- Glomerulopathy [see Adverse Reactions (6.1)]
- Immune thrombocytopenia
- Tumor lysis syndrome (TLS): Cases of possible TLS have been reported in patients treated with trastuzumab products. Patients with significant tumor burden (e.g. bulky metastases) may be at a higher risk. Patients could present with hyperuricemia, hyperphosphatemia, and acute renal failure which may represent possible TLS. Providers should consider additional monitoring and/or treatment as clinically

indicated.

7 DRUG INTERACTIONS

Patients who receive anthracycline after stopping trastuzumab products may be at increased risk of cardiac dysfunction because of trastuzumab's long washout period based on population PK analysis [see Clinical Pharmacology (12.3)]. If possible, physicians should avoid anthracycline-based therapy for up to 7 months after stopping trastuzumab products. If anthracyclines are used, the patient's cardiac function should be monitored carefully.

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Risk Summary

Trastuzumab products can cause fetal harm when administered to a pregnant woman. In post-marketing reports, use of trastuzumab during pregnancy resulted in cases of oligohydramnios and of oligohydramnios sequence, manifesting as pulmonary hypoplasia, skeletal abnormalities, and neonatal death (see Data). Apprise the patient of the potential risks to a fetus. There are clinical considerations if a trastuzumab product is used in a pregnant woman or if a patient becomes pregnant within 7 months following the last dose of a trastuzumab product (see Clinical Considerations).

The estimated background risk of major birth defects and miscarriage for the indicated population is unknown. In the U.S. general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnancies is 2% to 4% and 15% to 20%, respectively.

Clinical Considerations

Fetal/Neonatal Adverse Reactions

Monitor women who received TRAZIMERA during pregnancy or within 7 months prior to conception for oligohydramnios. If oligohydramnios occurs, perform fetal testing that is appropriate for gestational age and consistent with community standards of care.

Data

Human Data

In post-marketing reports, use of trastuzumab during pregnancy resulted in cases of oligohydramnios and of oligohydramnios sequence, manifesting in the fetus as pulmonary hypoplasia, skeletal abnormalities and neonatal death. These case reports described oligohydramnios in pregnant women who received trastuzumab either alone or in combination with chemotherapy. In some case reports, amniotic fluid index increased after trastuzumab was stopped. In one case, trastuzumab therapy resumed after amniotic index improved, and oligohydramnios recurred.

Animal Data

In studies where trastuzumab was administered to pregnant cynomolgus monkeys during the period of organogenesis at doses up to 25 mg/kg given twice weekly (up to

25 times the recommended weekly human dose of 2 mg/kg), trastuzumab crossed the placental barrier during the early (Gestation Days 20 to 50) and late (Gestation Days 120 to 150) phases of gestation. The resulting concentrations of trastuzumab in fetal serum and amniotic fluid were approximately 33% and 25%, respectively, of those present in the maternal serum but were not associated with adverse developmental effects.

8.2 Lactation

Risk Summary

There is no information regarding the presence of trastuzumab products in human milk, the effects on the breastfed infant, or the effects on milk production. Published data suggest human IgG is present in human milk but does not enter the neonatal and infant circulation in substantial amounts. Trastuzumab was present in the milk of lactating cynomolgus monkeys but not associated with neonatal toxicity (see Data). Consider the developmental and health benefits of breastfeeding along with the mother's clinical need for TRAZIMERA treatment and any potential adverse effects on the breastfed child from TRAZIMERA or from the underlying maternal condition. This consideration should also take into account the trastuzumab product wash out period of 7 months [see Clinical Pharmacology (12.3)].

Data

In lactating cynomolgus monkeys, trastuzumab was present in breast milk at about 0.3% of maternal serum concentrations after pre-(beginning Gestation Day 120) and post-partum (through Post-partum Day 28) doses of 25 mg/kg administered twice weekly (25 times the recommended weekly human dose of 2 mg/kg of trastuzumab products). Infant monkeys with detectable serum levels of trastuzumab did not exhibit any adverse effects on growth or development from birth to 1 month of age.

8.3 Females and Males of Reproductive Potential

Pregnancy Testing

Verify the pregnancy status of females of reproductive potential prior to the initiation of TRAZIMERA.

Contraception

Females

Trastuzumab products can cause embryo-fetal harm when administered during pregnancy. Advise females of reproductive potential to use effective contraception during treatment with TRAZIMERA and for 7 months following the last dose of TRAZIMERA [see Use in Specific Populations (8.1) and Clinical Pharmacology (12.3)].

8.4 Pediatric Use

The safety and effectiveness of trastuzumab products in pediatric patients have not been established.

8.5 Geriatric Use

Trastuzumab has been administered to 386 patients who were 65 years of age or over (253 in the adjuvant treatment and 133 in metastatic breast cancer treatment settings).

The risk of cardiac dysfunction was increased in geriatric patients as compared to younger patients in both those receiving treatment for metastatic disease in Studies 5 and 6, or adjuvant therapy in Studies 1 and 2. Limitations in data collection and differences in study design of the 4 studies of trastuzumab in adjuvant treatment of breast cancer preclude a determination of whether the toxicity profile of trastuzumab in older patients is different from younger patients. The reported clinical experience is not adequate to determine whether the efficacy improvements (ORR, TTP, OS, DFS) of trastuzumab treatment in older patients is different from that observed in patients <65 years of age for metastatic disease and adjuvant treatment.

In Study 7 (metastatic gastric cancer), of the 294 patients treated with trastuzumab, 108 (37%) were 65 years of age or older, while 13 (4.4%) were 75 and over. No overall differences in safety or effectiveness were observed.

10 OVERDOSAGE

There is no experience with overdosage in human clinical trials. Single doses higher than 8 mg/kg have not been tested.

11 DESCRIPTION

Trastuzumab-qyyp is a humanized IgG1 kappa monoclonal antibody that selectively binds with high affinity to the extracellular domain of the human epidermal growth factor receptor 2 protein, HER2. Trastuzumab-qyyp is produced by recombinant DNA technology in a mammalian cell (Chinese Hamster Ovary) culture.

TRAZIMERA (trastuzumab-qyyp) for injection is a sterile, white, preservative-free lyophilized powder with a cake-like appearance, for intravenous administration.

Each multiple-dose vial of TRAZIMERA delivers 420 mg trastuzumab-qyyp, 7.9 mg L-histidine, 9.5 mg L-histidine HCl monohydrate, 1.7 mg polysorbate 20, and 386 mg sucrose. Reconstitution with 20 mL of the appropriate diluent (BWFI or SWFI) yields a solution containing 21 mg/mL trastuzumab-qyyp that delivers 20 mL (420 mg trastuzumab-qyyp), at a pH of approximately 6. If TRAZIMERA is reconstituted with SWFI without preservative, the reconstituted solution is considered single-dose.

Each single-dose vial of TRAZIMERA delivers 150 mg trastuzumab-qyyp, 2.8 mg L-histidine, 3.4 mg L-histidine HCl monohydrate, 0.6 mg polysorbate 20, and 138 mg sucrose. Reconstitution with 7.4 mL of SWFI yields a solution containing 21 mg/mL trastuzumab-qyyp that delivers 7.15 mL (150 mg trastuzumab-qyyp), at a pH of approximately 6.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

The HER2 (or c-erbB2) proto-oncogene encodes a transmembrane receptor protein of 185 kDa, which is structurally related to the epidermal growth factor receptor. Trastuzumab products have been shown, in both *in vitro* assays and in animals, to inhibit the proliferation of human tumor cells that overexpress HER2.

Trastuzumab products are mediators of antibody-dependent cellular cytotoxicity (ADCC). *In vitro*, trastuzumab product-mediated ADCC has been shown to be preferentially exerted on HER2 overexpressing cancer cells compared with cancer cells that do not overexpress HER2.

12.2 Pharmacodynamics

Cardiac Electrophysiology

The effects of trastuzumab on electrocardiographic (ECG) endpoints, including QTc interval duration, were evaluated in patients with HER2 positive solid tumors. Trastuzumab had no clinically relevant effect on the QTc interval duration and there was no apparent relationship between serum trastuzumab concentrations and change in QTcF interval duration in patients with HER2 positive solid tumors.

12.3 Pharmacokinetics

The pharmacokinetics of trastuzumab were evaluated in a pooled population pharmacokinetic (PK) model analysis of 1,582 subjects with primarily breast cancer and metastatic gastric cancer (MGC) receiving intravenous trastuzumab. Total trastuzumab clearance increases with decreasing concentrations due to parallel linear and non-linear elimination pathways.

Although the average trastuzumab exposure was higher following the first cycle in breast cancer patients receiving the three-weekly schedule compared to the weekly schedule of trastuzumab, the average steady-state exposure was essentially the same at both dosages. The average trastuzumab exposure following the first cycle and at steady state as well as the time to steady state was higher in breast cancer patients compared to MGC patients at the same dosage; however, the reason for this exposure difference is unknown. Additional predicted trastuzumab exposure and PK parameters following the first trastuzumab cycle and at steady state exposure are described in Tables 7 and 8, respectively.

Population PK based simulations indicate that following discontinuation of trastuzumab, concentrations in at least 95% of breast cancer patients and MGC patients will decrease to approximately 3% of the population predicted steady-state trough serum concentration (approximately 97% washout) by 7 months [see Warnings and Precautions (5.1) and Use in Specific Populations (8.1, 8.3)].

Table 7 Population Predicted Cycle 1 PK Exposures (Median with 5th to 95th Percentiles) in Breast Cancer and MGC Patients

Schedule	Primary tumor type	N	C _{min} (μg/mL)	C _{max} (μg/mL)	AUC ₀₋₂₁ days (μg.day/mL)
8 mg/kg + 6 mg/kg q3w	Breast cancer	1195	29.4 (5.8 to 59.5)	178 (117 to 291)	1373 (736 to 2245)
	MGC	274	23.1 (6.1 to 50.3)	132 (84.2 to 225)	1109 (588 to 1938)

1 ma/ka 1	Drooct	1195	37.7	88.3	1066
4 mg/kg +			(12.3 to	(58 to	(586 to
2 mg/kg qw	Caricei		70.9)	144)	1754)

Table 8 Population Predicted Steady State PK Exposures (Median with 5th to 95th Percentiles) in Breast Cancer and MGC Patients

Schedule	Primary tumor type		C _{min,ss} * (μg/mL)	C _{max,ss} † (μg/mL)	ΔΠ(` Δ 31	Time to steady- state (week)	range					
	Breast		47.4	179	1794		0.173 to					
8 mg/kg	cancer	1195	1195	1195	1195	1195	1195	(5 to	(107 to	(673 to	12	0.283
+ 6	Carreer		115)	309)	3618)		0.203					
mg/kg			32.9	131	1338		0.189 to					
q3w	MGC	274	(6.1 to	(72.5 to	(557 to	9	0.169 (0					
			88.9)	251)	2875)		0.557					
4 mg/kg	Droost	Droost		66.1	109	1765		0.201 to				
+ 2	Breast	1195	(14.9 to	(51.0 to	(647 to	12	0.244					
mg/kg qw	cancer		142)	209)	3578)		0.244					

^{*} Steady-state trough serum concentration of trastuzumab

Specific Populations

Based on a population pharmacokinetic analysis, no clinically significant differences were observed in the pharmacokinetics of trastuzumab based on age (<65 (n = 1294); ≥65 (n = 288)), race (Asian (n = 264); non-Asian (n = 1324)) and renal impairment (mild (creatinine clearance [CLcr] 60 to 90 mL/min) (n = 636) or moderate (CLcr 30 to 60 mL/min) (n = 133)). The pharmacokinetics of trastuzumab products in patients with severe renal impairment, end-stage renal disease with or without hemodialysis, or hepatic impairment are unknown.

<u>Drug Interaction Studies</u>

There have been no formal drug interaction studies performed with trastuzumab products in humans. Clinically significant interactions between trastuzumab and concomitant medications used in clinical trials have not been observed.

Paclitaxel and Doxorubicin

Concentrations of paclitaxel and doxorubicin and their major metabolites (i.e., $6-\alpha$ hydroxyl-paclitaxel [POH], and doxorubicinol [DOL], respectively) were not altered in the presence of trastuzumab when used as combination therapy in clinical trials. Trastuzumab concentrations were not altered as part of this combination therapy.

Docetaxel and Carboplatin

When trastuzumab was administered in combination with docetaxel or carboplatin, neither the plasma concentrations of docetaxel or carboplatin nor the plasma

[†] Maximum steady-state serum concentration of trastuzumab

concentrations of trastuzumab were altered.

Cisplatin and Capecitabine

In a drug interaction substudy conducted in patients in Study 7, the pharmacokinetics of cisplatin, capecitabine and their metabolites were not altered when administered in combination with trastuzumab.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Trastuzumab products have not been tested for carcinogenic potential.

No evidence of mutagenic activity was observed when trastuzumab was tested in the standard Ames bacterial and human peripheral blood lymphocyte mutagenicity assays at concentrations of up to 5000 mcg/mL. In an *in vivo* micronucleus assay, no evidence of chromosomal damage to mouse bone marrow cells was observed following bolus intravenous doses of up to 118 mg/kg of trastuzumab.

A fertility study was conducted in female cynomolgus monkeys at doses up to 25 times the weekly recommended human dose of 2 mg/kg of trastuzumab and has revealed no evidence of impaired fertility, as measured by menstrual cycle duration and female sex hormone levels.

14 CLINICAL STUDIES

14.1 Adjuvant Breast Cancer

The safety and efficacy of trastuzumab in women receiving adjuvant chemotherapy for HER2 overexpressing breast cancer were evaluated in an integrated analysis of two randomized, open-label, clinical trials (Studies 1 and 2) with a total of 4063 women at the protocol-specified final overall survival analysis, a third randomized, open-label, clinical trial (Study 3) with a total of 3386 women at definitive Disease-Free Survival analysis for one-year trastuzumab treatment versus observation, and a fourth randomized, open-label clinical trial with a total of 3222 patients (Study 4).

Studies 1 and 2

In Studies 1 and 2, breast tumor specimens were required to show HER2 overexpression (3+ by IHC) or gene amplification (by FISH). HER2 testing was verified by a central laboratory prior to randomization (Study 2) or was required to be performed at a reference laboratory (Study 1). Patients with a history of active cardiac disease based on symptoms, abnormal electrocardiographic, radiologic, or left ventricular ejection fraction findings or uncontrolled hypertension (diastolic > 100 mm Hg or systolic > 200 mm Hg) were not eligible.

Patients were randomized (1:1) to receive doxorubicin and cyclophosphamide followed by paclitaxel (AC→paclitaxel) alone or paclitaxel plus trastuzumab (AC→paclitaxel + trastuzumab). In both trials, patients received four 21-day cycles of doxorubicin 60 mg/m² and cyclophosphamide 600 mg/m². Paclitaxel was administered either weekly (80 mg/m²) or every 3 weeks (175 mg/m²) for a total of 12 weeks in Study 1; paclitaxel was administered only by the weekly schedule in Study 2. Trastuzumab was administered at

4 mg/kg on the day of initiation of paclitaxel and then at a dose of 2 mg/kg weekly for a total of 52 weeks. Trastuzumab treatment was permanently discontinued in patients who developed congestive heart failure, or persistent/recurrent LVEF decline [see Dosage and Administration (2.3)]. Radiation therapy, if administered, was initiated after the completion of chemotherapy. Patients with ER+ and/or PR+ tumors received hormonal therapy. The primary endpoint of the combined efficacy analysis was Disease-Free Survival (DFS), defined as the time from randomization to recurrence, occurrence of contralateral breast cancer, other second primary cancer, or death. The secondary endpoint was overall survival (OS).

A total of 3752 patients were included in the joint efficacy analysis of the primary endpoint of DFS following a median follow-up of 2.0 years in the AC→paclitaxel + trastuzumab arm. The pre-planned final OS analysis from the joint analysis included 4063 patients and was performed when 707 deaths had occurred after a median follow-up of 8.3 years in the AC→paclitaxel + trastuzumab arm. The data from both arms in Study 1 and two of the three study arms in Study 2 were pooled for efficacy analyses. The patients included in the primary DFS analysis had a median age of 49 years (range, 22 to 80 years; 6% > 65 years), 84% were white, 7% black, 4% Hispanic, and 4% Asian/Pacific Islander. Disease characteristics included 90% infiltrating ductal histology, 38% T1, 91% nodal involvement, 27% intermediate and 66% high grade pathology, and 53% ER+ and/or PR+ tumors. Similar demographic and baseline characteristics were reported for the efficacy evaluable population, after 8.3 years of median follow-up in the AC→paclitaxel + trastuzumab arm.

Study 3

In Study 3, breast tumor specimens were required to show HER2 overexpression (3+ by IHC) or gene amplification (by FISH) as determined at a central laboratory. Patients with node-negative disease were required to have ≥ T1c primary tumor. Patients with a history of congestive heart failure or LVEF < 55%, uncontrolled arrhythmias, angina requiring medication, clinically significant valvular heart disease, evidence of transmural infarction on ECG, poorly controlled hypertension (systolic > 180 mm Hg or diastolic > 100 mm Hg) were not eligible.

Study 3 was designed to compare one and two years of three-weekly trastuzumab treatment versus observation in patients with HER2 positive EBC following surgery, established chemotherapy and radiotherapy (if applicable). Patients were randomized (1:1:1) upon completion of definitive surgery, and at least four cycles of chemotherapy to receive no additional treatment, or one year of trastuzumab treatment or two years of trastuzumab treatment. Patients undergoing a lumpectomy had also completed standard radiotherapy. Patients with ER+ and/or PgR+ disease received systemic adjuvant hormonal therapy at investigator discretion. Trastuzumab was administered with an initial dose of 8 mg/kg followed by subsequent doses of 6 mg/kg once every three weeks. The main outcome measure was Disease-Free Survival (DFS), defined as in Studies 1 and 2.

A protocol-specified interim efficacy analysis comparing one-year trastuzumab treatment to observation was performed at a median follow-up duration of 12.6 months in the trastuzumab arm and formed the basis for the definitive DFS results from this study. Among the 3386 patients randomized to the observation (n = 1693) and trastuzumab one-year (n = 1693) treatment arms, the median age was 49 years (range 21 to 80), 83% were Caucasian, and 13% were Asian. Disease characteristics: 94%

infiltrating ductal carcinoma, 50% ER+ and/or PgR+, 57% node positive, 32% node negative, and in 11% of patients, nodal status was not assessable due to prior neo-adjuvant chemotherapy. Ninety-six percent (1055/1098) of patients with node-negative disease had high-risk features: among the 1098 patients with node-negative disease, 49% (543) were ER- and PgR-, and 47% (512) were ER and/or PgR+ and had at least one of the following high-risk features: pathological tumor size greater than 2 cm, Grade 2 to 3, or age < 35 years. Prior to randomization, 94% of patients had received anthracycline-based chemotherapy regimens.

After the definitive DFS results comparing observation to one-year trastuzumab treatment were disclosed, a prospectively planned analysis that included comparison of one year versus two years of trastuzumab treatment at a median follow-up duration of 8 years was performed. Based on this analysis, extending trastuzumab treatment for a duration of two years did not show additional benefit over treatment for one year [Hazard Ratios of two-years trastuzumab versus one-year trastuzumab treatment in the intent to treat (ITT) population for Disease-Free Survival (DFS) = 0.99 (95% CI: 0.87, 1.13), p-value = 0.90 and Overall Survival (OS) = 0.98 (0.83, 1.15); p-value = 0.78].

Study 4

In Study 4, breast tumor specimens were required to show HER2 gene amplification (FISH+ only) as determined at a central laboratory. Patients were required to have either node-positive disease, or node-negative disease with at least one of the following high-risk features: ER/PR-negative, tumor size > 2 cm, age < 35 years, or histologic and/or nuclear Grade 2 or 3. Patients with a history of CHF, myocardial infarction, Grade 3 or 4 cardiac arrhythmia, angina requiring medication, clinically significant valvular heart disease, poorly controlled hypertension (diastolic > 100 mm Hg), any T4 or N2 or known N3 or M1 breast cancer were not eligible.

Patients were randomized (1:1:1) to receive doxorubicin and cyclophosphamide followed by docetaxel (AC-T), doxorubicin and cyclophosphamide followed by docetaxel plus trastuzumab (AC-TH), or docetaxel and carboplatin plus trastuzumab (TCH). In both the AC-T and AC-TH arms, doxorubicin 60 mg/m² and cyclophosphamide 600 mg/m² were administered every 3 weeks for four cycles; docetaxel 100 mg/m² was administered every 3 weeks for four cycles. In the TCH arm, docetaxel 75 mg/m² and carboplatin (at a target AUC of 6 mg/mL/min as a 30- to 60-minute infusion) were administered every 3 weeks for six cycles. Trastuzumab was administered weekly (initial dose of 4 mg/kg followed by weekly dose of 2 mg/kg) concurrently with either T or TC, and then every 3 weeks (6 mg/kg) as monotherapy for a total of 52 weeks. Radiation therapy, if administered, was initiated after completion of chemotherapy. Patients with ER+ and/or PR+ tumors received hormonal therapy. Disease-Free Survival (DFS) was the main outcome measure.

Among the 3222 patients randomized, the median age was 49 (range 22 to 74 years; $6\% \ge 65$ years). Disease characteristics included 54% ER+ and/or PR+ and 71% node positive. Prior to randomization, all patients underwent primary surgery for breast cancer.

The results for DFS for the integrated analysis of Studies 1 and 2, Study 3, and Study 4 and OS results for the integrated analysis of Studies 1 and 2, and Study 3 are presented in Table 9. For Studies 1 and 2, the duration of DFS following a median follow-up of 2.0 years in the AC→TH arm is presented in Figure 4, and the duration of OS after a median follow-up of 8.3 years in the AC→TH arm is presented in Figure 5. The duration of DFS

for Study 4 is presented in Figure 6. Across all four studies, at the time of definitive DFS analysis, there were insufficient numbers of patients within each of the following subgroups to determine if the treatment effect was different from that of the overall patient population: patients with low tumor grade, patients within specific ethnic/racial subgroups (Black, Hispanic, Asian/Pacific Islander patients), and patients > 65 years of age. For Studies 1 and 2, the OS hazard ratio was 0.64 (95% CI: 0.55, 0.74). At 8.3 vears of median follow-up [AC→TH], the survival rate was estimated to be 86.9% in the $AC \rightarrow TH$ arm and 79.4% in the AC $\rightarrow T$ arm. The final OS analysis results from Studies 1 and 2 indicate that OS benefit by age, hormone receptor status, number of positive lymph nodes, tumor size and grade, and surgery/radiation therapy was consistent with the treatment effect in the overall population. In patients ≤ 50 years of age (n = 2197), the OS hazard ratio was 0.65 (95% CI: 0.52, 0.81) and in patients > 50 years of age (n = 1866), the OS hazard ratio was 0.63 (95% CI: 0.51, 0.78). In the subgroup of patients with hormone receptor-positive disease (ER-positive and/or PR-positive) (n = 2223), the hazard ratio for OS was 0.63 (95% CI: 0.51, 0.78). In the subgroup of patients with hormone receptor-negative disease (ER-negative and PR-negative) (n = 1830), the hazard ratio for OS was 0.64 (95% CI: 0.52, 0.80). In the subgroup of patients with tumor size \leq 2 cm (n = 1604), the hazard ratio for OS was 0.52 (95% CI: 0.39, 0.71). In the subgroup of patients with tumor size > 2 cm (n = 2448), the hazard ratio for OS was 0.67 (95% CI: 0.56, 0.80).

Table 9 Efficacy Results from Adjuvant Treatment of Breast Cancer (Studies 1 + 2, Study 3, and Study 4)

	DFS events	DFS Hazard ratio (95% CI) p-value	Deaths (OS events)	ratio
Studies 1 + 2*				
AC→TH (n = 1872) [†] (n = 2031) [‡]	133 [†]	0.48 ^{†,§} (0.39, 0.59) p< 0.0001 [¶]	289 [‡]	0.64 ^{‡,§} (0.55, 0.74) p< 0.0001 [¶]
AC→T (n = 1880) [†] (n = 2032) [‡]	261 [†]		418 [‡]	
Study 3 [#]				
Chemo→Trastuzumab (n = 1693)	127	0.54 (0.44, 0.67) p< 0.0001 ^b	31	0.75 p = NS ^ß
Chemo→Observation (n = 1693)	219		40	
Study 4 ^à				
TCH (n = 1075)	134	0.67 (0.54 - 0.84) p=0.0006 [¶] ,è		
AC→TH (n = 1074)	121	0.60 (0.48 - 0.76) p< 0.0001 ^{¶,à}	49	

AC→T	180	80	
(n = 1073)	100	00	

CI = confidence interval.

- * Studies 1 and 2 regimens: doxorubicin and cyclophosphamide followed by paclitaxel (AC→T) or paclitaxel plus trastuzumab (AC→TH).
- † Efficacy evaluable population, for the primary DFS analysis, following a median follow-up of 2.0 years in the AC→TH arm.
- ‡ Efficacy evaluable population, for the final OS analysis, following 707 deaths (8.3 years of median follow-up in the AC→TH arm).
- § Hazard ratio estimated by Cox regression stratified by clinical trial, intended paclitaxel schedule, number of positive nodes, and hormone receptor status.
- ¶ stratified log-rank test.
- # At definitive DFS analysis with median duration of follow-up of 12.6 months in the one-year trastuzumab treatment arm.
- Þ log-rank test.
- β NS = non-significant.
- à Study 4 regimens: doxorubicin and cyclophosphamide followed by docetaxel (AC→T) or docetaxel plus trastuzumab (AC→TH); docetaxel and carboplatin plus trastuzumab (TCH).
- è A two-sided alpha level of 0.025 for each comparison.

Figure 4
Duration of Disease-Free Survival in Patients with Adjuvant
Treatment of Breast Cancer (Studies 1 and 2)

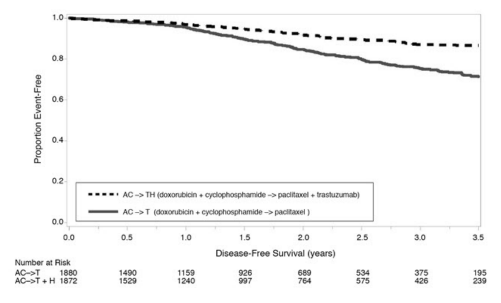


Figure 5

Duration of Overall Survival in Patients with Adjuvant
Treatment of Breast Cancer (Studies 1 and 2)

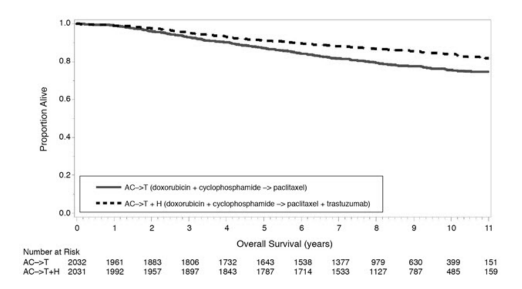
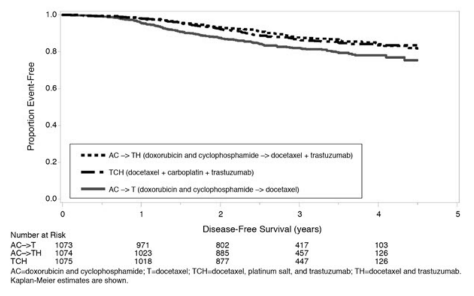


Figure 6
Duration of Disease-Free Survival in Patients with Adjuvant
Treatment of Breast Cancer (Study 4)



Exploratory analyses of DFS as a function of HER2 overexpression or gene amplification were conducted for patients in Studies 2 and 3, where central laboratory testing data were available. The results are shown in Table 10. The number of events in Study 2 was small with the exception of the IHC 3+/FISH+ subgroup, which constituted 81% of those with data. Definitive conclusions cannot be drawn regarding efficacy within other subgroups due to the small number of events. The number of events in Study 3 was adequate to demonstrate significant effects on DFS in the IHC 3+/FISH unknown and the FISH +/IHC unknown subgroups.

Table 10 Treatment Outcomes in Studies 2 and 3 as a Function of HER2 Overexpression or Amplification

	Stu	ıdy 2	Study 3*		
HER2 Assay	Number	Hazard	Number	Hazard	

Result [†]	Patients	(95% CI)	Patients	(95% CI)
IHC 3+	1			
FISH (+)	1170	0.42 (0.27, 0.64)	91	0.56 (0.13, 2.50)
FISH (-)	51	0.71 (0.04, 11.79)	8	_
FISH Unknown	51	0.69 (0.09, 5.14)	2258	0.53 (0.41, 0.69)
IHC < 3+ / FISH (+)	174	1.01 (0.18, 5.65)	299 [‡]	0.53 (0.20, 1.42)
IHC unknown / FISH (+)	_	_	724	0.59 (0.38, 0.93)

^{*} Median follow-up duration of 12.6 months in the one-year trastuzumab treatment arm.

14.2 Metastatic Breast Cancer

The safety and efficacy of trastuzumab in treatment of women with metastatic breast cancer were studied in a randomized, controlled clinical trial in combination with chemotherapy (Study 5, n = 469 patients) and an open-label single agent clinical trial (Study 6, n = 222 patients). Both trials studied patients with metastatic breast cancer whose tumors overexpress the HER2 protein. Patients were eligible if they had 2 or 3 levels of overexpression (based on a 0 to 3 scale) by immunohistochemical assessment of tumor tissue performed by a central testing lab.

<u>Previously Untreated Metastatic Breast Cancer (Study 5)</u>

Study 5 was a multicenter, randomized, open-label clinical trial conducted in 469 women with metastatic breast cancer who had not been previously treated with chemotherapy for metastatic disease. Tumor specimens were tested by IHC (Clinical Trial Assay, CTA) and scored as 0, 1+, 2+, or 3+, with 3+ indicating the strongest positivity. Only patients with 2+ or 3+ positive tumors were eligible (about 33% of those screened). Patients were randomized to receive chemotherapy alone or in combination with trastuzumab given intravenously as a 4 mg/kg loading dose followed by weekly doses of trastuzumab at 2 mg/kg. For those who had received prior anthracycline therapy in the adjuvant setting, chemotherapy consisted of paclitaxel (175 mg/m² over 3 hours every 21 days for at least six cycles); for all other patients, chemotherapy consisted of anthracycline plus cyclophosphamide (AC: doxorubicin 60 mg/m² or epirubicin 75 mg/m² plus 600 mg/m² cyclophosphamide every 21 days for six cycles). Sixty-five percent of patients randomized to receive chemotherapy alone in this study received trastuzumab at the time of disease progression as part of a separate extension study.

Based upon the determination by an independent response evaluation committee, the patients randomized to trastuzumab and chemotherapy experienced a significantly longer median time to disease progression, a higher overall response rate (ORR), and a longer median duration of response as compared with patients randomized to chemotherapy alone. Patients randomized to trastuzumab and chemotherapy also had a

[†] IHC by HercepTest, FISH by PathVysion (HER2/CEP17 ratio ≥ 2.0) as performed at a central laboratory.

[‡] All cases in this category in Study 3 were IHC 2+.

longer median survival (see Table 11). These treatment effects were observed both in patients who received trastuzumab plus paclitaxel and in those who received trastuzumab plus AC; however the magnitude of the effects was greater in the paclitaxel subgroup.

Table 11 Study 5: Efficacy Results in First-Line Treatment for Metastatic Breast Cancer

	Combine	d Results	Paclitaxel S	ubgroup	AC Subgroup		
	Trastuzumab + All Chemotherapy (n = 235)	All Chemotherapy (n = 234)	Trastuzumab + Paclitaxel (n = 92)	Paclitaxe (n = 96)		AC (n = 138)	
Primary E	ndpoint						
Median TTP (mos) ^{†,‡}	7.2	4.5	6.7	2.5	7.6	5.7	
95% CI	7, 8	4, 5	5, 10	2, 4	7, 9	5, 7	
p-value§	< 0.0	0001	< 0.00	01	0.002		
Secondar	y Endpoints				T		
Overall Response Rate [†]	45	29	38	15	50	38	
95% CI	39, 51	23, 35	28, 48	8, 22	42, 58	30, 46	
p-value [¶]	< 0.	001	< 0.00	01	0.10		
Median Resp Duration (mos) ^{†,‡}	8.3	5.8	8.3	4.3	8.4	6.4	
25%, 75% Quartile	6, 15	4, 8	5, 11	4, 7	6, 15	4, 8	
Med Survival (mos) [‡]	25.1	20.3	22.1	18.4	26.8	21.4	
95% CI	22, 30	17, 24	17, 29	13, 24	23, 33	18, 27	
p-value§	0.0	05	0.17		0.16		

^{*} AC = Anthracycline (doxorubicin or epirubicin) and cyclophosphamide.

Data from Study 5 suggest that the beneficial treatment effects were largely limited to patients with the highest level of HER2 protein overexpression (3+) (see Table 12).

[†] Assessed by an independent Response Evaluation Committee.

[‡] Kaplan-Meier Estimate.

[§] log-rank test.

[¶] χ 2-test.

HER2 Overexpression or Amplification

HER2 Assay Result	Number of Patients (N)	Relative Risk* for Time to Disease Progression (95% CI)	Relative Risk* for Mortality (95% CI)
CTA 2+ or 3+	469	0.49 (0.40, 0.61)	0.80 (0.64, 1.00)
FISH (+) [†]	325	0.44 (0.34, 0.57)	0.70 (0.53, 0.91)
FISH (–) [†]	126	0.62 (0.42, 0.94)	1.06 (0.70, 1.63)
CTA 2+	120	0.76 (0.50, 1.15)	1.26 (0.82, 1.94)
FISH (+)	32	0.54 (0.21, 1.35)	1.31 (0.53, 3.27)
FISH (-)	83	0.77 (0.48, 1.25)	1.11 (0.68, 1.82)
CTA 3+	349	0.42 (0.33, 0.54)	0.70 (0.51, 0.90)
FISH (+)	293	0.42 (0.32, 0.55)	0.67 (0.51, 0.89)
FISH (-)	43	0.43 (0.20, 0.94)	0.88 (0.39, 1.98)

^{*} The relative risk represents the risk of progression or death in the trastuzumab plus chemotherapy arm versus the chemotherapy arm.

Previously Treated Metastatic Breast Cancer (Study 6)

Trastuzumab was studied as a single agent in a multicenter, open-label, single-arm clinical trial (Study 6) in patients with HER2 overexpressing metastatic breast cancer who had relapsed following one or two prior chemotherapy regimens for metastatic disease. Of 222 patients enrolled, 66% had received prior adjuvant chemotherapy, 68% had received two prior chemotherapy regimens for metastatic disease, and 25% had received prior myeloablative treatment with hematopoietic rescue. Patients were treated with a loading dose of 4 mg/kg IV followed by weekly doses of trastuzumab at 2 mg/kg IV.

The ORR (complete response + partial response), as determined by an independent Response Evaluation Committee, was 14%, with a 2% complete response rate and a 12% partial response rate. Complete responses were observed only in patients with disease limited to skin and lymph nodes. The overall response rate in patients whose tumors tested as CTA 3+ was 18% while in those that tested as CTA 2+, it was 6%.

14.3 Metastatic Gastric Cancer

The safety and efficacy of trastuzumab in combination with cisplatin and a fluoropyrimidine (capecitabine or 5-fluorouracil) were studied in patients previously untreated for metastatic gastric or gastroesophageal junction adenocarcinoma (Study

[†] FISH testing results were available for 451 of the 469 patients enrolled on study.

7). In this open-label, multi-center trial, 594 patients were randomized 1:1 to trastuzumab in combination with cisplatin and a fluoropyrimidine (FC+H) or chemotherapy alone (FC). Randomization was stratified by extent of disease (metastatic vs. locally advanced), primary site (gastric vs. gastroesophageal junction), tumor measurability (yes vs. no), ECOG performance status (0,1 vs. 2), and fluoropyrimidine (capecitabine vs. 5-fluorouracil). All patients were either HER2 gene amplified (FISH+) or HER2 overexpressing (IHC 3+). Patients were also required to have adequate cardiac function (e.g., LVEF > 50%).

On the trastuzumab-containing arm, trastuzumab was administered as an IV infusion at an initial dose of 8 mg/kg followed by 6 mg/kg every 3 weeks until disease progression. On both study arms cisplatin was administered at a dose of 80 mg/m² Day 1 every 3 weeks for 6 cycles as a 2 hour IV infusion. On both study arms, capecitabine was administered at 1000 mg/m² dose orally twice daily (total daily dose 2000 mg/m²) for 14 days of each 21 day cycle for 6 cycles. Alternatively, continuous intravenous infusion (CIV) 5-fluorouracil was administered at a dose of 800 mg/m²/day from Day 1 through Day 5 every three weeks for 6 cycles.

The median age of the study population was 60 years (range: 21 to 83); 76% were male; 53% were Asian, 38% Caucasian, 5% Hispanic, 5% other racial/ethnic groups; 91% had ECOG PS of 0 or 1; 82% had primary gastric cancer and 18% had primary gastroesophageal adenocarcinoma. Of these patients, 23% had undergone prior gastrectomy, 7% had received prior neoadjuvant and/or adjuvant therapy, and 2% had received prior radiotherapy.

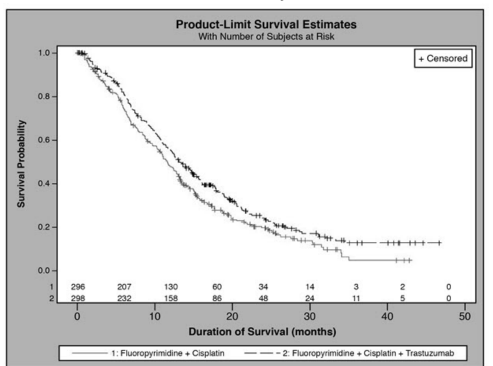
The main outcome measure of Study 7 was overall survival (OS), analyzed by the unstratified log-rank test. The final OS analysis based on 351 deaths was statistically significant (nominal significance level of 0.0193). An updated OS analysis was conducted at one year after the final analysis. The efficacy results of both the final and the updated analyses are summarized in Table 13 and Figure 7.

Table 13 Study 7: Overall Survival in ITT Population

	FC Arm N = 296	FC + H Arm N = 298		
Definitive (Second Interim) Overall Survival				
No. Deaths (%)	184 (62.2%)	167 (56.0%)		
Median	11.0	13.5		
95% CI (mos.)	(9.4, 12.5)	(11.7, 15.7)		
Hazard Ratio	0.73			
95% CI	(0.60, 0.91)			
p-value*, two-sided	0.0038			
Updated Overall Survi	val			
No. Deaths (%)	227 (76.7%)	221 (74.2%)		
Median	11.7	13.1		
95% CI (mos.)	(10.3, 13.0) (11.9, 15.1)			
Hazard Ratio	0.80			
95% CI (0.67, 0.97)		0.97)		

^{*} Comparing with the nominal significance level of 0.0193.

Figure 7
Updated Overall Survival in Patients with Metastatic Gastric Cancer (Study 7)



An exploratory analysis of OS in patients based on HER2 gene amplification (FISH) and protein overexpression (IHC) testing is summarized in Table 14.

Table 14 Exploratory Analyses by HER2 Status Using Updated Overall Survival Results

	FC (N = 296)*	FC + H (N = 298) [†]
FISH+ / IHC 0, 1+ subgrou	up (N=133)	
No. Deaths / n (%)	57/71 (80%)	56/62 (90%)
Median OS Duration (mos.)	8.8	8.3
95% CI (mos.)	(6.4, 11.7)	(6.2, 10.7)
Hazard ratio (95% CI)	1.33 (0.9	92, 1.92)
FISH+ / IHC2+ subgroup (N=160)	
No. Deaths / n (%)	65/80 (81%)	64/80 (80%)
Median OS Duration (mos.)	10.8	12.3
95% CI (mos.)	(6.8, 12.8)	(9.5, 15.7)
Hazard ratio (95% CI)	0.78 (0.5	55, 1.10)
FISH+ or FISH- / IHC3+‡ s	ubgroup (N=294)	
No. Deaths / n (%)	104/143 (73%)	96/151 (64%)
Median OS Duration (mos.)	13.2	18.0
95% CI (mos.)	(11.5, 15.2)	(15.5, 21.2)

Hazard ratio (95% CI)

0.66 (0.50, 0.87)

- * Two patients on the FC arm who were FISH+ but IHC status unknown were excluded from the exploratory subgroup analyses.
- † Five patients on the trastuzumab-containing arm who were FISH+, but IHC status unknown were excluded from the exploratory subgroup analyses.
- ‡ Includes 6 patients on chemotherapy arm, 10 patients on trastuzumab arm with FISH-, IHC3+ and 8 patients on chemotherapy arm, 8 patients on trastuzumab arm with FISH status unknown, IHC 3+.

16 HOW SUPPLIED/STORAGE AND HANDLING

16.1 How Supplied

420 mg Multiple-dose vial

TRAZIMERA (trastuzumab-qyyp) for injection 420 mg/vial is supplied in a multiple-dose vial as a sterile, white lyophilized powder. Each carton contains one multiple-dose vial of TRAZIMERA and one vial (20 mL) of Bacteriostatic Water for Injection (BWFI) containing 1.1% benzyl alcohol as a preservative.

NDC 0069-0305-01.

150 mg Single-dose vial

TRAZIMERA (trastuzumab-qyyp) for injection 150 mg/vial is supplied in a single-dose vial as a sterile, white lyophilized powder. Each carton contains one single-dose vial of TRAZIMERA.

NDC 0069-0308-01.

16.2 Storage

Store TRAZIMERA vials in the refrigerator at 2°C to 8°C (36°F to 46°F) in the original carton to protect from light.

If needed, unopened TRAZIMERA vials may be removed from the refrigerator and stored at room temperature up to 30°C (86°F) for a single period of up to 3 months in the original carton to protect from light. Once removed from the refrigerator, do not return to the refrigerator and discard after 3 months or by the expiration date stamped on the vial, whichever occurs first. Write the revised expiration date in the space provided on the carton labeling.

17 PATIENT COUNSELING INFORMATION

Cardiomyopathy

 Advise patients to contact a health care professional immediately for any of the following: new onset or worsening shortness of breath, cough, swelling of the ankles/legs, swelling of the face, palpitations, weight gain of more than 5 pounds in 24 hours, dizziness or loss of consciousness [see Boxed Warning].

Embryo-Fetal Toxicity

 Advise pregnant women and females of reproductive potential that TRAZIMERA exposure during pregnancy or within 7 months prior to conception can result in fetal harm. Advise female patients to contact their healthcare provider with a known or suspected pregnancy [see Use in Specific Populations (8.1)].

 Advise females of reproductive potential to use effective contraception during treatment and for 7 months following the last dose of TRAZIMERA [see Use in Specific Populations (8.3)].

This product's labeling may have been updated. For the most recent prescribing information, please visit www.TRAZIMERA.com.

Manufactured by: Pfizer Ireland Pharmaceuticals Cork, Ireland US License No. 2060



Distributed by Pfizer Labs Division of Pfizer Inc New York, NY 10017

LAB-0995-4.0

PRINCIPAL DISPLAY PANEL - 420 mg Vial Label

NDC 0069-0306-01 Rx only

Trazimera[™] (trastuzumab-qyyp) For Injection

Mfg. by Pfizer Ireland Pharmaceuticals Cork, Ireland

US License No. 2060

420 mg/vial

For intravenous infusion after reconstitution

Multiple-dose vial

Reconstitution, Dosage, and Administration: For IV administration only.

See prescribing information for preparation instructions, dosage, and administration.

Storage: Refrigerate at 2°C to 8°C (36°F to 46°F). Unopened vial can be stored up to 30°C

(86°F) for a single period of up to 3 months but not exceeding the expiry date.

Reconstitute with 20 mL Bacteriostatic Water for Injection (1.1% benzyl alcohol) to yield a

21 mg/mL multiple-dose solution that delivers 20 mL (420 mg trastuzumab-qyyp). Store

reconstituted solution in refrigerator at 2°C to 8°C (36°F to 46°F). Do Not Freeze. Do Not Shake After Reconstitution. Discard unused reconstituted solution after 28 days.

Discard after ___/___/

	NDC 0069-0306-01 Rx only Trazimera ™ (trastuzumab-qyyp) For Injection	Mfg. by Pfizer Ireland Pharmaceuticals Cork, Ireland US License No. 2060
		on: For IV administration only. n instructions, dosage, and administration. 46°F). Unopened vial can be stored up to 30°C
LOT/EXP:	21 mg/mL multiple-dose solution that deliv reconstituted solution in refrigerator at 2°0	ater for Injection (1.1% benzyl alcohol) to yield a ers 20 mL (420 mg trastuzumab-qyyp). Store C to 8°C (36°F to 46°F). Do Not Freeze. Id unused reconstituted solution after 28 days.

PRINCIPAL DISPLAY PANEL - 20 mL Vial Label

BACTERIOSTATIC WATER FOR INJECTION

NOT FOR USE IN NEWBORNS NDC 0069-0307-01

For Drug Diluent Use Only Use 20 mL as DILUENT for reconstitution of Trazimera

Each mL contains: 1.1% benzyl alcohol; Water for Injection q.s. pH 4.5 to 8.0 See enclosed full prescribing information. Use only if solution is clear and seal intact. Sterile. Nonpyrogenic.

Store at controlled room temperature 2°C to 30°C (36°F to 86°F).

20 mL VIAL

PAA140335



PRINCIPAL DISPLAY PANEL - Kit Carton

NDC 0069-0305-01

Trazimera™ (trastuzumab-qyyp) For Injection

420 mg/vial

For intravenous infusion after reconstitution

Multiple-dose vial

Rx only

Pfizer Oncology

Reconstitution, Dosage, and Administration: For IV administration only. See prescribing information for preparation instructions, dosage, and administration.

Storage: Refrigerate at 2°C to 8°C (36°F to 46°F). Unopened vial can be stored at up to 30°C (86°F) for a single period of up to 3 months but not exceeding the expiry date. Discard after

Reconstitute with 20 mL Bacteriostatic Water for Injection (1.1% benzyl alcohol) to yield a 21 mg/mL multiple-dose solution that delivers 20 mL (420 mg trastuzumab-qyyp). Store reconstituted solution in refrigerator at 2°C to 8°C (36°F to 46°F). Do Not Freeze. Do Not Shake After Reconstitution. Discard unused reconstituted solution after 28 days.



K1010 SC029 Black Cool Gray 11 Cool Gray 2 PU7452 NDC 0069-0305-01

Trazimera™

(trastuzumab-qyyp) For Injection

420 mg/vial

For intravenous infusion after reconstitution Multiple-dose vial

Rx only

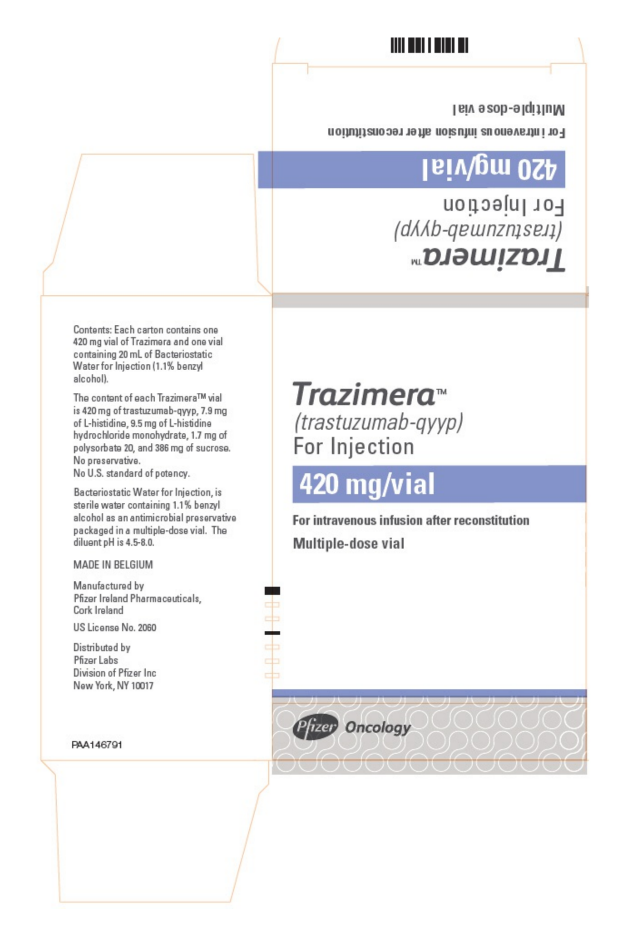
Pfizer Oncology

GTIN: 00300690305011 LOT/EXP/SN:

> Data matrix area







PRINCIPAL DISPLAY PANEL - 150 mg Vial Label

Trazimera[™] (trastuzumab-qyyp) For Injection

Mfg. by Pfizer Ireland Pharmaceuticals Cork, Ireland

US License No. 2060

150 mg/vial

For intravenous infusion after reconstitution Single-dose vial

For Reconstitution, Dilution and Dosage: See Prescribing Information.

Storage: Refrigerate at 2°C to 8°C (36°F to 46°F) in the original carton to protect from light. Unopened vial can be stored up to 30°C (86°F) for a single period of up to 3 months but not exceeding the expiry date. Discard after ___/____. Do Not Freeze. Do Not Shake After Reconstitution. Discard unused reconstituted solution after 24 hours.



PRINCIPAL DISPLAY PANEL - 150 mg Vial Carton

NDC 0069-0308-01

Trazimera™ (trastuzumab-qyyp)

For Injection

150 mg/vial

For intravenous infusion after

reconstitution
Single-dose vial
Rx only
Pfizer Oncology



TRAZIMERA

trastuzumab-qyyp kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0069-0305

Packaging

ı					
I	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
ı	1 N	IDC:0069-0305-01	1 in 1 KIT	02/24/2020	

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	1 VIAL, MULTI-DOSE	20 mL
Part 2	1 VIAL	20 mL

Part 1 of 2

TRAZIMERA

trastuzumab-qyyp injection, powder, lyophilized, for solution

Product Information

Item Code (Source)	NDC:0069-0306
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
TRASTUZUMAB (UNII: P188ANX8CK) (TRASTUZUMAB - UNII:P188ANX8CK)	TRASTUZ UMAB	420 mg in 20 mL

Inactive Ingredients Ingredient Name Strength SUCROSE (UNII: C151H8M554) HISTIDINE MONOHYDROCHLORIDE MONOHYDRATE (UNII: X573657P6P) HISTIDINE (UNII: 4QD397987E) POLYSORBATE 20 (UNII: 7T1F30V5YH) Strength 386 mg in 20 mL 7.9 mg in 20 mL

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:0069- 0306-01	20 mL in 1 VIAL, MULTI-DOSE; Type 1: Convenience Kit of Co-Package		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA761081	02/24/2020	

Part 2 of 2

BACTERIOSTATIC WATER

water injection, solution

Product Information

Item Code (Source)NDC:0069-0307Route of AdministrationINTRAVENOUS

Inactive Ingredients Ingredient Name Strength WATER (UNII: 059QF0KO0R) BENZYL ALCOHOL (UNII: LKG8494WBH)

Packaging				
# I	Item Code	Package Description	Marketing Start Date	Marketing End Date
1 NE 01		20 mL in 1 VIAL; Type 1: Convenience Kit of Co-Package		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA761081	02/24/2020	

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
BLA	BLA761081	02/24/2020			

TRAZIMERA

trastuzumab-qyyp injection, powder, lyophilized, for solution

Product Information

Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0069-0308
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety Ingredient Name Basis of Strength TRASTUZUMAB (UNII: P188ANX8CK) (TRASTUZUMAB - UNII: P188ANX8CK) TRASTUZUMAB 150 mg in 7.15 mL

Inactive Ingredients					
Ingredient Name	Strength				
HISTIDINE (UNII: 4QD397987E)	2.8 mg in 7.15 mL				
HISTIDINE MONOHYDROCHLORIDE MONOHYDRATE (UNII: X573657P6P)	3.4 mg in 7.15 mL				
POLYSORBATE 20 (UNII: 7T1F30V5YH)	0.6 mg in 7.15 mL				
SUCROSE (UNII: C151H8M554)	138 mg in 7.15 mL				

P	Packaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date			
1	NDC:0069-0308- 01	1 in 1 CARTON	03/10/2021				
1		7.15 mL in 1 VIAL; Type 0: Not a Combination Product					

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
BLA	BLA761081	03/10/2021			

Labeler - Pfizer Laboratories Div Pfizer Inc (134489525)

Establishment					
Name	Address	ID/FEI	Business Operations		
Boehringer Ingelheim Pharma GmbH and Co. KG		340700520	API MANUFACTURE(0069-0305) , ANALYSIS(0069-0305)		

Establishment			
Name	Address	ID/FEI	Business Operations
Pfizer Ireland Pharmaceuticals		985586408	ANALYSIS(0069-0305, 0069-0308), API MANUFACTURE(0069-0305, 0069-0308)

Establishment					
Name	Address	ID/FEI	Business Operations		
Pfizer			ANALYSIS(0069-0305, 0069-0308) . MANUFACTURE(0069-0305, 0069-0308) .		

	3/012020/	PACK(0069-0305,	0069-0308),	LABEL(0069-0305,	0069-0308)
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Establishment						
Name	Address	ID/FEI	Business Operations			
Wyeth BioPharma Division of Wyeth Pharmaceuticals LLC		174350868	ANALYSIS(0069-0305, 0069-0308), API MANUFACTURE(0069-0305, 0069-0308)			

Manuracturing Belgium NV

Revised: 2/2022 Pfizer Laboratories Div Pfizer Inc