103792/5044 Approved 10/16/03

Herceptin[®] (Trastuzumab)

1 HERCEPTIN[®]

2 Trastuzumab

3 WARNINGS:

4

CARDIOMYOPATHY

5 HERCEPTIN administration can result in the development of ventricular 6 dysfunction and congestive heart failure. Left ventricular function should 7 be evaluated in all patients prior to and during treatment with 8 HERCEPTIN. Discontinuation of HERCEPTIN treatment should be 9 strongly considered in patients who develop a clinically significant 10 decrease in left ventricular function. The incidence and severity of cardiac 11 dysfunction was particularly high in patients who received HERCEPTIN 12 in combination with anthracyclines and cyclophosphamide. 13 (See WARNINGS.) HYPERSENSITIVITY REACTIONS INCLUDING ANAPHYLAXIS 14

15 **INFUSION REACTIONS**

16 **PULMONARY EVENTS**

17 HERCEPTIN administration can result in severe hypersensitivity reactions18 (including anaphylaxis), infusion reactions, and pulmonary events.

19 Rarely, these have been fatal. In most cases, symptoms occurred during or

- 20 within 24 hours of administration of HERCEPTIN. HERCEPTIN infusion
- 21 should be interrupted for patients experiencing dyspnea or clinically
- 22 significant hypotension. Patients should be monitored until signs and

23 symptoms completely resolve. Discontinuation of HERCEPTIN treatment

- 24 should be strongly considered for patients who develop anaphylaxis,
- 25 angioedema, or acute respiratory distress syndrome. (See WARNINGS.)

26 **DESCRIPTION**

- 27 HERCEPTIN (Trastuzumab) is a recombinant DNA-derived humanized
- 28 monoclonal antibody that selectively binds with high affinity in a
- 29 cell-based assay (Kd=5 nM) to the extracellular domain of the human
- 30 epidermal growth factor receptor 2 protein, HER2 (1,2). The antibody is
- 31 an IgG_1 kappa that contains human framework regions with the

32 complementarity-determining regions of a murine antibody (4D5) that33 binds to HER2.

34 The humanized antibody against HER2 is produced by a mammalian cell

35 (Chinese Hamster Ovary [CHO]) suspension culture in a nutrient medium

36 containing the antibiotic gentamicin. Gentamicin is not detectable in the

37 final product.

38 HERCEPTIN is a sterile, white to pale yellow, preservative-free

39 lyophilized powder for intravenous (IV) administration. The nominal

40 content of each HERCEPTIN vial is 440 mg Trastuzumab, 9.9 mg

41 L-histidine HCl, 6.4 mg L-histidine, 400 mg α , α -trehalose dihydrate, and

42 1.8 mg polysorbate 20, USP. Reconstitution with 20 mL of the supplied

43 **Bacteriostatic Water for Injection (BWFI), USP, containing**

44 1.1% benzyl alcohol as a preservative, yields a multi-dose solution

45 containing 21 mg/mL Trastuzumab, at a pH of approximately 6.

46 CLINICAL PHARMACOLOGY

47 General

48 The HER2 (or c-erbB2) proto-oncogene encodes a transmembrane 49 receptor protein of 185 kDa, which is structurally related to the epidermal 50 growth factor receptor (1). HER2 protein overexpression is observed in 51 25%–30% of primary breast cancers. HER2 protein overexpression can 52 be determined using immunohistochemistry (IHC) and gene amplification 53 can be determined using fluorescence in situ hybridization (FISH) of fixed 54 tumor blocks (2). In referenced studies where HERCEPTIN use was not 55 studied (3-5), approximately 96-98% of biopsy specimens that were found 56 to have protein overexpression also had gene amplification and 100% of 57 those with gene amplification also had protein overexpression (3-5). The 58 precision of the determination of protein overexpression or gene amplification, however, may vary depending on the sensitivity and 59 60 specificity of the particular assay and assay procedures used (see 61 PRECAUTIONS). When compared to the referenced studies noted above,

62 the correlation between detectable protein overexpression using

63 immunohistochemistry (IHC) and detectable gene amplification using

64 fluorescence in situ hybridization (FISH) was not as high in the studies of

65 HERCEPTIN clinical trial specimens (see CLINICAL STUDIES: HER2

66 Detection and HER2 Assay Concordance Studies and PRECAUTIONS:

67 HER2 Testing).

68 Trastuzumab has been shown, in both *in vitro* assays and in animals, to

69 inhibit the proliferation of human tumor cells that overexpress HER2 (6-

70 8).

71 Trastuzumab is a mediator of antibody-dependent cellular cytotoxicity

72 (ADCC) (9,10). In vitro, HERCEPTIN-mediated ADCC has been shown

to be preferentially exerted on HER2 overexpressing cancer cells

74 compared with cancer cells that do not overexpress HER2.

75 **Pharmacokinetics**

76 The pharmacokinetics of Trastuzumab were studied in breast cancer

77 patients with metastatic disease. Short duration intravenous infusions of

78 10 to 500 mg once weekly demonstrated dose-dependent

79 pharmacokinetics. Mean half-life increased and clearance decreased with

80 increasing dose level. The half-life averaged 1.7 and 12 days at the 10 and

81 500 mg dose levels, respectively. Trastuzumab's volume of distribution

82 was approximately that of serum volume (44 mL/kg). At the highest

83 weekly dose studied (500 mg), mean peak serum concentrations were

84 377 microgram/mL.

85 In studies using a loading dose of 4 mg/kg followed by a weekly

86 maintenance dose of 2 mg/kg, a mean half-life of 5.8 days

87 (range=1 to 32 days) was observed. Between Weeks 16 and 32,

88 Trastuzumab serum concentrations reached a steady state with mean

89 trough and peak concentrations of approximately 79 microgram/mL and

90 123 microgram/mL, respectively.

91 Detectable concentrations of the circulating extracellular domain of the

92 HER2 receptor (shed antigen) are found in the sera of some patients with HERCEPTIN[®]—Genentech, Inc. October 2003

- HER2 overexpressing tumors. Determination of shed antigen in baseline
 serum samples revealed that 64% (286/447) of patients had detectable
- serum samples revealed that 64% (286/447) of patients had detectable
- 95 shed antigen, which ranged as high as 1880 ng/mL (median=11 ng/mL).
- 96 Patients with higher baseline shed antigen levels were more likely to have
- 97 lower serum trough concentrations. However, with weekly dosing, most
- 98 patients with elevated shed antigen levels achieved target serum
- 99 concentrations of Trastuzumab by Week 6.
- Data suggest that the disposition of Trastuzumab is not altered based on
 age or serum creatinine (up to 2.0 mg/dL). No formal interaction studies
 have been performed.
- 103 Mean serum trough concentrations of Trastuzumab, when administered in
- 104 combination with paclitaxel, were consistently elevated approximately
- 105 1.5-fold as compared with serum concentrations of Trastuzumab used in
- 106 combination with anthracycline plus cyclophosphamide. In primate
- 107 studies, administration of Trastuzumab with paclitaxel resulted in a
- 108 reduction in Trastuzumab clearance. Serum levels of Trastuzumab in
- 109 combination with cisplatin, doxorubicin or epirubicin plus
- 110 cyclophosphamide did not suggest any interactions; no formal drug
- 111 interaction studies were performed.

112 CLINICAL STUDIES

- 113 The safety and efficacy of HERCEPTIN were studied in a randomized,
- 114 controlled clinical trial in combination with chemotherapy (469 patients)
- and an open-label single agent clinical trial (222 patients). Both trials
- 116 studied patients with metastatic breast cancer whose tumors overexpress
- 117 the HER2 protein. Patients were eligible if they had 2+or 3+levels of
- 118 overexpression (based on a 0 to 3+scale) by immunohistochemical
- assessment of tumor tissue performed by a central testing lab.
- 120 A multicenter, randomized, controlled clinical trial was conducted in
- 121 469 patients with metastatic breast cancer who had not been previously
- 122 treated with chemotherapy for metastatic disease (11). Patients were

123	randomized to receive chemotherapy alone or in combination with
124	HERCEPTIN given intravenously as a 4 mg/kg loading dose followed by
125	weekly doses of HERCEPTIN at 2 mg/kg. For those who had received
126	prior anthracycline therapy in the adjuvant setting, chemotherapy
127	consisted of paclitaxel (175 mg/m ² over 3 hours every 21 days for at least
128	six cycles); for all other patients, chemotherapy consisted of anthracycline
129	plus cyclophosphamide (AC: doxorubicin 60 mg/m ² or epirubicin
130	75 mg/m ² plus 600 mg/m ² cyclophosphamide every 21 days for
131	six cycles). Compared with patients in the AC subgroups ($n=281$),
132	patients in the paclitaxel subgroup $(n=188)$ were more likely to have had
133	the following: poor prognostic factors (premenopausal status, estrogen or
134	progesterone receptor negative tumors, positive lymph nodes), prior
135	therapy (adjuvant chemotherapy, myeloablative chemotherapy,
136	radiotherapy), and a shorter disease-free interval. Sixty-five percent of
137	patients randomized to receive chemotherapy alone in this study received
138	HERCEPTIN at the time of disease progression as part of a separate
139	extension study.
140	Compared with patients randomized to chemotherapy alone, the patients
141	randomized to HERCEPTIN and chemotherapy experienced a

significantly longer median time to disease progression, a higher overall
response rate (ORR), a longer median duration of response, and a longer
median survival (see Table 1). These treatment effects were observed
both in patients who received HERCEPTIN plus paclitaxel and in those
who received HERCEPTIN plus AC, however the magnitude of the
effects was greater in the paclitaxel subgroup (see CLINICAL STUDIES:
HER2 Detection).

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Table 1	
Phase III Clinical Efficacy in First-Line Treatme	ent

	Combined Results		Paclitaxel Subgroup		AC Subgroup	
	HERCEPTIN + All Chemotherapy	All Chemotherapy	HERCEPTIN + Paclitaxel	Paclitaxel	HERCEPTIN + ACª	AC
	(n = 235)	(n = 234)	(n = 92)	(n = 96)	(n = 143)	(n = 138)
Primary Endpoint						
<u>Time to Progression</u> ^{b, c}						
Median (months)	7.2	4.5	6.7	2.5	7.6	5.7
95% confidence interval	6.9, 8.2	4.3, 4.9	5.2, 9.9	2.0, 4.3	7.2, 9.1	4.6, 7.1
p-value (log rank)	< 0.	0001	< 0.0001		0.002	
Secondary Endpoints						
Overall Response Rate ^b		:				
Rate (percent)	45	29	38	15	50	38
95% confidence interval	39, 51	23, 35	28, 48	8, 22	42, 58	30, 46
p-value (χ2-test)	< 0	.001	< 0.001		0.10	
Duration of Response ^{b,c}						
Median (months)	8.3	5.8	8.3	4.3	8.4	6.4
25%, 75% quartile	5.5, 14.8	3.9, 8.5	5.1, 11.0	3.7, 7.4	5.8, 14.8	4.5, 8.5
<u>Survival Time</u> ^c						
Median Survival (months)	25.1	20.3	22.1	18.4	26.8	21.4
95% confidence interval	22.2, 29.5	16.8, 24.2	16.9, 28.6	12.7, 24.4	23.3, 32.9	18.3, 26.6
p-value (log rank)	0.	05	0.1	7	0.1	6

^a AC = anthracycline (doxorubicin or epirubicin) and cyclophosphamide.

^b Assessed by an independent Response Evaluation Committee.

^c Kaplan-Meier Estimate.

149

150 HERCEPTIN was studied as a single agent in a multicenter, open-label,

151 single-arm clinical trial in patients with HER2 overexpressing metastatic

152 breast cancer who had relapsed following one or two prior chemotherapy

153 regimens for metastatic disease. Of 222 patients enrolled, 66% had

154 received prior adjuvant chemotherapy, 68% had received two prior

155 chemotherapy regimens for metastatic disease, and 25% had received prior

156 myeloablative treatment with hematopoietic rescue. Patients were treated

- 157 with a loading dose of 4 mg/kg IV followed by weekly doses of
- 158 HERCEPTIN at 2 mg/kg IV. The ORR (complete response+partial
- 159 response), as determined by an independent Response Evaluation
- 160 Committee, was 14%, with a 2% complete response rate and a 12% partial
- 161 response rate. Complete responses were observed only in patients with
- 162 disease limited to skin and lymph nodes (see CLINICAL STUDIES:
- 163 HER2 Detection).

164 **HER2 Detection**

165 (See PRECAUTIONS: HER2 Testing)

166 Detection of HER2 protein overexpression is necessary for selection of

- 167 patients appropriate for HERCEPTIN therapy (see INDICATIONS).
- 168 Overexpression of HER2 by tumors was an entry criterion of the two
- 169 clinical studies described above. In those studies, a research-use-only IHC

170 assay (referred to as the Clinical Trial Assay, CTA) was used.

- 171 The commercial assays described below, HercepTest[™] (IHC assay) and
- 172 PathVysion^M (FISH assay), are appropriate assays to aid in the selection of
- 173 patients for HERCEPTIN therapy (see CLINICAL STUDIES: HER2
- 174 Detection: HER2 Protein Overexpression Detection Methods and HER2
- 175 Gene Amplification Detection Methods). The comparability of either
- assay with regard to the ability to predict clinical benefit from

177 HERCEPTIN therapy has not been prospectively studied. In addition, the

- 178 utility of either assay in patients whose tumors would score as 0 or 1+ by
- 179 the Clinical Trial Assay (CTA) has not been established because patients

180 with tumors that scored as 0 or 1+ were excluded from the clinical studies

181 described.

182 HER2 Protein Overexpression Detection Methods

- 183 HER2 protein overexpression can be established by measuring expressed
- 184 HER2 protein using IHC methodology. In the clinical trial studies
- described above, specimens were tested with the CTA and scored as 0, 1+,
- 2+, or 3+ with 3+ indicating the strongest positivity. Only patients with
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 October 2003

187 2+ or 3+ positive tumors were eligible (about 33% of those screened). 188 Data from the randomized trial suggest that the beneficial treatment effects 189 were largely limited to patients with the highest level of HER2 protein 190 overexpression (3+) (see Table 2). In an exploratory analysis, the relative 191 risk (rr) for time to progression was lower in the patients whose tumors 192 tested as CTA 3+ (rr = 0.42 with 95% CI: 0.33, 0.54) than in those tested 193 as CTA 2+ (rr = 0.76 with 95% CI: 0.50, 1.15). The relative risk 194 represents the risk of progression in the HERCEPTIN plus chemotherapy 195 arm versus the chemotherapy arm. Therefore, a lower ratio represents 196 longer time to progression in the HERCEPTIN arm. In the single-arm 197 study of HERCEPTIN as a single agent, the overall response rate in 198 patients whose tumors tested as CTA 3+ was 18% while in those that 199 tested as CTA 2+, it was 6%.

HercepTest[™], another IHC assay, was assessed for concordance with the
CTA (see HER2 Testing: Concordance Studies), but has not been used to
assess tumor specimens from the HERCEPTIN clinical studies described
above.

204 HER2 Gene Amplification Detection Methods

205 As a surrogate for protein overexpression, measurement of the number of 206 HER2 gene copies using FISH to detect gene amplification may be 207 employed. An exploratory, retrospective assessment of known CTA 2+ or 3+ tumor specimens was performed to detect HER2 gene amplification 208 209 using PathVysion[™], a FISH assay. Data from this retrospective analysis 210 involving 660 of 691 (96%) patients enrolled in the clinical studies (all 211 scoring 2+ or 3+ by the CTA) suggested that the beneficial treatment 212 effects were greater in patients whose tumors tested as FISH (+) than in 213 those that were FISH (-); however, time to progression was prolonged for 214 patients on the HERCEPTIN arm, regardless of the FISH result (see 215 Table 2). In the single arm study of HERCEPTIN as a single agent, the 216 overall response rate in patients whose tumors tested as FISH (+) was 217 20%, while in those tested as FISH (-), there were no responses.

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- These data are not sufficient to conclude whether FISH testing can 218
- distinguish a subpopulation of CTA 2+ patients who would be unlikely to 219
- benefit from HERCEPTIN therapy. In addition, there are no data 220
- correlating clinical outcome with FISH test results for patients with tumors 221
- that scored as 0 or 1+ by CTA; therefore, conclusions regarding the 222
- 223 usefulness of FISH in the general population cannot be made.
- 224
- 225 226

227

I able 2
Treatment Effect versus Level of HER2 Expression
Phase III Randomized Trial ($N = 469$):
HERCEPTIN Plus Chemotherany versus Chemotherany

m 11. A

HER2 Assay	Number of	Relative Risk** for	Relative Risk** for		
Result Patients (N)		Time to Disease	Mortality		
		Progression (95% CI)	(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	$0.62 \ (0.42, 0.94)^{\circ}$	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)		

228

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

The relative risk represents the risk of progression or death in the HERCEPTIN plus **

229 230 chemotherapy arm versus the chemotherapy arm.

HER2 Assay Concordance Studies 231

232 (See PRECAUTIONS: HER2 Testing)

Immunohistochemistry: The DAKO HercepTest[™], an IHC test for 233 detecting HER2 protein overexpression, has not been directly studied for 234 its ability to predict HERCEPTIN treatment effect, but has been compared 235 to the CTA on over 500 breast cancer histology specimens obtained from 236 237 the National Cancer Institute Cooperative Breast Cancer Tissue Resource. Based upon these results, of specimens testing 3+ (strongly positive) on 238 the HercepTest[™], 82% were 3+(i.e., the reading most associated with 239 clinical benefit), 12% were 2+, and 6% were 0 or 1+ on the CTA. The 6% 240 of HercepTest[™] 3+ specimens that were CTA 0 or 1+ would be expected 241 to represent 2% of the 0 and 1+ population. Of specimens testing 2+242 (weakly positive) on the HercepTest[™], 14% were 3+, 20% were 2+, and 243 66% were 0 or 1+ on the CTA. Of specimens testing 0 or 1+ on the 244 HercepTest^{$^{\text{M}}$}, 2% were 3+, 6% were 2+, and 92% were 0 or 1+ on the 245 246 CTA.

Fluorescence in situ Hybridization: The Vysis PathVysion[™] HER2 DNA 247 248 Probe, a FISH test for detecting HER2 gene amplification, was compared with the CTA on over 500 breast cancer histology specimens originally 249 250 submitted for potential enrollment in the HERCEPTIN trials. A HER2:CEP17 ratio of ≥ 2 was defined as FISH positive (+). Based on 251 these results, of specimens testing FISH (+) by PathVysion[™], 81% were 252 3+, 10% were 2+, and 9% were 0 or 1+ on the CTA. The 9% of FISH (+) 253 254 specimens that were CTA 0 or 1+ would be expected to represent 3% of the total CTA 0 or 1+ population. Of specimens testing FISH (-) by 255 PathVysion[™], 3% were 3+, 10% were 2+, and 87% were 0 or 1+ on the 256 257 CTA.

258 INDICATIONS AND USAGE

HERCEPTIN as a single agent is indicated for the treatment of patients
with metastatic breast cancer whose tumors overexpress the HER2 protein
and who have received one or more chemotherapy regimens for their
metastatic disease. HERCEPTIN in combination with paclitaxel is
indicated for treatment of patients with metastatic breast cancer whose
tumors overexpress the HER2 protein and who have not received

- 265 chemotherapy for their metastatic disease. HERCEPTIN should be used
- 266 in patients whose tumors have been evaluated with an assay validated to
- 267 predict HER2 protein overexpression (see PRECAUTIONS: HER2
- 268 Testing and CLINICAL STUDIES: HER2 Detection).

269 CONTRAINDICATIONS

270 None known.

271 (WARNINGS)

272 **Cardiotoxicity**:

- 273 Signs and symptoms of cardiac dysfunction, such as dyspnea, increased
- 274 cough, paroxysmal nocturnal dyspnea, peripheral edema, S₃ gallop, or
- 275 reduced ejection fraction, have been observed in patients treated with
- 276 HERCEPTIN. Congestive heart failure associated with HERCEPTIN
- therapy may be severe and has been associated with disabling cardiac
- failure, death, and mural thrombosis leading to stroke (see BOXED
- 279 WARNINGS: CARDIOMYOPATHY). The clinical status of patients in
- 280 the trials who developed congestive heart failure was classified for
- 281 severity using the New York Heart Association classification system
- 282 (I-IV, where IV is the most severe level of cardiac failure). (See Table 3.)

Table 3

Incidence and Severity of Cardiac Dysfunction

	HERCEPTIN ^a alone n=213	HERCEPTIN +Paclitaxel ^b n=91	Paclitaxel ^b n=95	HERCEPTIN + Anthracycline + cyclophosphamide ^b n=143	Anthracycline + cyclophosphamide ^b n=135
Any Cardiac Dysfunction	7%	11%	1%	28%	7%
Class III-IV	5%	4%	1%	19%	3%

^a Open-label, single-agent Phase II study (94% received prior anthracyclines).

^b Randomized Phase III study comparing chemotherapy plus HERCEPTIN to chemotherapy alone, where chemotherapy is either anthracycline/cyclophosphamide or paclitaxel.

283

- 284 Candidates for treatment with HERCEPTIN should undergo thorough
- 285 baseline cardiac assessment including history and physical exam and one

- 286 or more of the following: EKG, echocardiogram, and MUGA scan. There
- are no data regarding the most appropriate method of evaluation for the
- 288 identification of patients at risk for developing cardiotoxicity. Monitoring
- 289 may not identify all patients who will develop cardiac dysfunction.
- <u>Extreme caution</u> should be exercised in treating patients with pre-existing
 cardiac dysfunction.
- 292 Patients receiving HERCEPTIN should undergo frequent monitoring for293 deteriorating cardiac function.
- 294 The probability of cardiac dysfunction was highest in patients who
- 295 received HERCEPTIN concurrently with anthracyclines. The data suggest
- that advanced age may increase the probability of cardiac dysfunction.
- 297 Pre-existing cardiac disease or prior cardiotoxic therapy
- 298 (e.g., anthracycline or radiation therapy to the chest) may decrease the
- ability to tolerate HERCEPTIN therapy; however, the data are not
- 300 adequate to evaluate the correlation between HERCEPTIN-induced
- 301 cardiotoxicity and these factors.
- 302 Discontinuation of HERCEPTIN therapy should be strongly considered in
- 303 patients who develop clinically significant congestive heart failure. In the
- 304 clinical trials, most patients with cardiac dysfunction responded to
- 305 appropriate medical therapy often including discontinuation of
- 306 HERCEPTIN. The safety of continuation or resumption of HERCEPTIN
- 307 in patients who have previously experienced cardiac toxicity has not been
- 308 studied. There are insufficient data regarding discontinuation of
- 309 HERCEPTIN therapy in patients with asymptomatic decreases in ejection
- 310 fraction; such patients should be closely monitored for evidence of clinical
- 311 deterioration.
- 312 Hypersensitivity Reactions Including Anaphylaxis:
- 313 Severe hypersensitivity reactions have been infrequently reported in
- 314 patients treated with HERCEPTIN (see BOXED WARNINGS:
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- 315 HYPERSENSITIVITY REACTIONS INCLUDING ANAPHYLAXIS).
- 316 Signs and symptoms include anaphylaxis, urticaria, bronchospasm,
- 317 angioedema, and/or hypotension. In some cases, the reactions have been
- 318 fatal. The onset of symptoms generally occurred during an infusion, but
- there have also been reports of symptom onset after the completion of an
- 320 infusion. Reactions were most commonly reported in association with the
- 321 initial infusion.
- 322 **HERCEPTIN** infusion should be interrupted in all patients with
- 323 severe hypersensitivity reactions. In the event of a hypersensitivity
- 324 reaction, appropriate medical therapy should be administered, which may
- 325 include epinephrine, corticosteroids, diphenhydramine, bronchodilators,
- 326 and oxygen. Patients should be evaluated and carefully monitored until
- 327 complete resolution of signs and symptoms.
- 328 There are no data regarding the most appropriate method of identification
- 329 of patients who may safely be retreated with HERCEPTIN after
- 330 experiencing a severe hypersensitivity reaction. HERCEPTIN has been
- 331 readministered to some patients who fully recovered from a previous
- 332 severe reaction. Prior to readministration of HERCEPTIN, the majority of
- 333 these patients were prophylactically treated with pre-medications
- including antihistamines and/or corticosteroids. While some of these
- 335 patients tolerated retreatment, others had severe reactions again despite the
- 336 use of prophylactic pre-medications.
- 337 Infusion Reactions:
- 338 In the postmarketing setting, rare occurrences of severe infusion reactions
- 339 leading to a fatal outcome have been associated with the use of
- 340 HERCEPTIN. (See BOXED WARNINGS: INFUSION REACTIONS.)
- 341 In clinical trials, infusion reactions consisted of a symptom complex
- 342 characterized by fever and chills, and on occasion included nausea,
- 343 vomiting, pain (in some cases at tumor sites), headache, dizziness,

	Herceptin [®] (Trastuzumab)
344	dyspnea, hypotension, rash, and asthenia. These reactions were usually
345	mild to moderate in severity. (See ADVERSE REACTIONS.)
346	However, in postmarketing reports, more severe adverse reactions to
347	HERCEPTIN infusion were observed and included bronchospasm,
348	hypoxia, and severe hypotension. These severe reactions were usually
349	associated with the initial infusion of HERCEPTIN and generally occurred
350	during or immediately following the infusion. However, the onset and
351	clinical course were variable. For some patients, symptoms progressively
352	worsened and led to further pulmonary complications. (See WARNINGS:
353	Pulmonary Events.) In other patients with acute onset of signs and
354	symptoms, initial improvement was followed by clinical deterioration.
355	Delayed post-infusion events with rapid clinical deterioration have also
356	been reported. Rarely, severe infusion reactions culminated in death
357 [.]	within hours or up to one week following an infusion.
358	Some severe reactions have been treated successfully with interruption of
359	the HERCEPTIN infusion and supportive therapy including oxygen,
360	intravenous fluids, beta-agonists, and corticosteroids.
361	There are no data regarding the most appropriate method of identification
362	of patients who may safely be retreated with HERCEPTIN after
363	experiencing a severe infusion reaction. HERCEPTIN has been
364	readministered to some patients who fully recovered from the previous
365	severe reaction. Prior to readministration of HERCEPTIN, the majority of
366	these patients were prophylactically treated with pre-medications
367	including antihistamines and/or corticosteroids. While some of these
368	patients tolerated retreatment, others had severe reactions again despite the

use of prophylactic pre-medications. 369

Exacerbation of Chemotherapy Induced Neutropenia 370

- 371
- 372 In randomized, controlled clinical trials designed to assess the impact of
- 373 the addition of HERCEPTIN on chemotherapy, the per-patient incidences

374	of moderate to severe neutropenia and of febrile neutropenia were higher
375	in patients receiving HERCEPTIN in combination with myelosuppressive
376	chemotherapy as compared to those who received chemotherapy alone. In
377	the postmarketing setting, deaths due to sepsis in patients with severe
378	neutropenia have been reported in patients receiving HERCEPTIN and
379	myelosuppressive chemotherapy, although in controlled clinical trials
380	(pre- and post-marketing), the incidence of septic deaths was not
381	significantly increased. The pathophysiologic basis for exacerbation of
382	neutropenia has not been determined; the effect of HERCEPTIN on the
383	pharmacokinetics of chemotherapeutic agents has not been fully evaluated
384	(See ADVERSE REACTIONS: Anemia and Leukopenia; ADVERSE
385	REACTIONS: Infection).

386 **Pulmonary Events:**

- 387 Severe pulmonary events leading to death have been reported rarely with
- 388 the use of HERCEPTIN in the postmarketing setting. Signs, symptoms
- and clinical findings include dyspnea, pulmonary infiltrates, pleural
- 390 effusions, non-cardiogenic pulmonary edema, pulmonary insufficiency
- and hypoxia, and acute respiratory distress syndrome. These events may
- 392 or may not occur as sequelae of infusion reactions. (See WARNINGS:
- 393 Infusion Reactions.) Patients with symptomatic intrinsic lung disease or
- 394 with extensive tumor involvement of the lungs, resulting in dyspnea at
- 395 rest, may be at greater risk of severe reactions.
- 396 Other severe events reported rarely in the postmarketing setting include
- 397 pneumonitis and pulmonary fibrosis.

398 **PRECAUTIONS**

399 General:

- 400 HERCEPTIN therapy should be used with caution in patients with known
- 401 hypersensitivity to Trastuzumab, Chinese Hamster Ovary cell proteins, or
- 402 any component of this product.

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403 **HER2 Testing:**

- 404 Assessment for HER2 overexpression should be performed by laboratories
- 405 with demonstrated proficiency in the specific technology being utilized.
- 406 Improper assay performance, including use of suboptimally fixed tissue,
- 407 failure to utilize specified reagents, deviation from specific assay
- 408 instructions, and failure to include appropriate controls for assay
- 409 validation, can lead to unreliable results. Refer to the HercepTest[™] and
- 410 PathVysion[™] package inserts for full instructions on assay performance
- 411 (see CLINICAL STUDIES: HER2 Detection).

412 **Drug Interactions:**

- 413 There have been no formal drug interaction studies performed with
- 414 HERCEPTIN in humans. Administration of paclitaxel in combination
- 415 with HERCEPTIN resulted in a two-fold decrease in HERCEPTIN
- 416 clearance in a non-human primate study and in a 1.5-fold increase in
- 417 HERCEPTIN serum levels in clinical studies.
- 418 (See PHARMACOKINETICS.)

419 Benzyl Alcohol:

- 420 For patients with a known hypersensitivity to benzyl alcohol (the
- 421 preservative in Bacteriostatic Water for Injection) reconstitute
- 422 HERCEPTIN with Sterile Water for Injection (SWFI), USP. DISCARD
- 423 THE SWFI-RECONSTITUTED HERCEPTIN VIAL FOLLOWING A
- 424 SINGLE USE.

425 Carcinogenesis, Mutagenesis, Impairment of Fertility:

426 **Carcinogenesis:**

427 HERCEPTIN has not been tested for its carcinogenic potential.

428 Mutagenesis:

- 429 No evidence of mutagenic activity was observed in Ames tests using
- 430 six different test strains of bacteria, with and without metabolic activation,
- 431 at concentrations of up to 5000 µg/mL Trastuzumab. Human peripheral
- 432 blood lymphocytes treated *in vitro* at concentrations of up to 5000 µg/plate
- Trastuzumab, with and without metabolic activation, revealed no evidence
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 October 2003

- 434 of mutagenic potential. In an *in vivo* mutagenic assay (the micronucleus
- 435 assay), no evidence of chromosomal damage to mouse bone marrow cells
- 436 was observed following bolus intravenous doses of up to 118 mg/kg
- 437 Trastuzumab.

438 **Impairment of Fertility:**

- 439 A fertility study has been conducted in female cynomolgus monkeys at
- 440 doses up to 25 times the weekly human maintenance dose of 2 mg/kg
- 441 HERCEPTIN and has revealed no evidence of impaired fertility.

442 **Pregnancy Category B:**

443 Reproduction studies have been conducted in cynomolgus monkeys at 444 doses up to 25 times the weekly human maintenance dose of 2 mg/kg 445 HERCEPTIN and have revealed no evidence of impaired fertility or harm 446 to the fetus. However, HER2 protein expression is high in many 447 embryonic tissues including cardiac and neural tissues; in mutant mice 448 lacking HER2, embryos died in early gestation (12). Placental transfer of 449 HERCEPTIN during the early (Days 20–50 of gestation) and late 450 (Days 120-150 of gestation) fetal development period was observed in 451 monkeys. There are, however, no adequate and well-controlled studies in 452 pregnant women. Because animal reproduction studies are not always 453 predictive of human response, this drug should be used during pregnancy 454 only if clearly needed.

455 Nursing Mothers:

456 A study conducted in lactating cynomolgus monkeys at doses 25 times the 457 weekly human maintenance dose of 2 mg/kg HERCEPTIN demonstrated 458 that Trastuzumab is secreted in the milk. The presence of Trastuzumab in 459 the serum of infant monkeys was not associated with any adverse effects 460 on their growth or development from birth to 3 months of age. It is not 461 known whether HERCEPTIN is excreted in human milk. Because human 462 IgG is excreted in human milk, and the potential for absorption and harm 463 to the infant is unknown, women should be advised to discontinue nursing

464 during HERCEPTIN therapy and for 6 months after the last dose of

465 HERCEPTIN.

466 **Pediatric Use:**

467 The safety and effectiveness of HERCEPTIN in pediatric patients have not468 been established.

469 **Geriatric Use:**

470 HERCEPTIN has been administered to 133 patients who were 65 years of

471 age or over. The risk of cardiac dysfunction may be increased in geriatric

472 patients. The reported clinical experience is not adequate to determine

473 whether older patients respond differently from younger patients.

474 (ADVERSE REACTIONS)

475 The most serious adverse reactions caused by HERCEPTIN include

476 cardiomyopathy, hypersensitivity reactions including anaphylaxis,

477 infusion reactions, pulmonary events, and exacerbation of chemotherapy-

478 induced neutropenia. Please refer to the BOXED WARNINGS and/or

479 WARNINGS sections for detailed descriptions of these reactions. The

480 most common adverse reactions associated with HERCEPTIN use are

481 fever, diarrhea, infections, chills, increased cough, headache, rash and

482 insomnia.

483 Because clinical trials are conducted under widely varying conditions,

484 adverse reaction rates observed in the clinical trials of a drug cannot be

485 directly compared to rates in the clinical trials of another drug and may not

486 reflect the rates observed in practice. The adverse reaction information

487 from clinical trials does, however, provide a basis for identifying the

adverse events that appear to be related to drug use and for approximating

489 rates.

490 Additional adverse reactions have been identified during post-marketing

491 use of HERCEPTIN. Because these reactions are reported voluntarily

492 from a population of uncertain size, it is not always possible to reliably

493 estimate their frequency or establish a causal relationship to HERCEPTIN
 HERCEPTIN[®]—Genentech, Inc.
 18
 October 2003

494 exposure. Decisions to include these reactions in labeling are typically

495 based on one or more of the following factors: (1) seriousness of the

496 reaction, (2) frequency of reporting, or (3) strength of causal connection to

497 HERCEPTIN.

498 Where specific percentages are noted, these data are based on clinical

499 studies of HERCEPTIN alone or in combination with chemotherapy in

500 clinical trials. Data in Table 4 are based on the experience with the

501 recommended dosing regimen for HERCEPTIN in a randomized

502 controlled clinical trial of 234 patients who received HERCEPTIN in

503 combination with chemotherapy and four open-label studies of

504 HERCEPTIN as a single agent in 352 patients at doses of 10-500 mg

administered weekly. Data regarding serious adverse events are based on

506 experience in 958 patients enrolled in all clinical trials of HERCEPTIN

507 conducted prior to marketing approval.

508 **Cardiac Failure/Dysfunction:**

509 For a description of cardiac toxicities, see BOXED WARNINGS:

510 CARDIOMYOPATHY and WARNINGS: Cardiotoxicity.

511 Anemia and Leukopenia:

512 In a randomized, controlled trial (see CLINICAL STUDIES), the per-

513 patient incidences of anemia (30% vs. 21%) and leukopenia (53% vs.

514 37%) were higher in patients receiving HERCEPTIN in combination with

515 chemotherapy as compared to those receiving chemotherapy alone. The

516 majority of these cytopenic events were mild to moderate in intensity,

517 reversible, and none resulted in discontinuation of therapy with

518 HERCEPTIN.

519 In a randomized, controlled trial conducted in the post-marketing setting,

- 520 there were also increased incidences of NCI-CTC Grade 3/4 neutropenia
- 521 (32% [29/92] vs. 22% [21/94]) and of febrile neutropenia (23% [21/91] vs.
- 522 17% [16/94]) in patients randomized to HERCEPTIN in combination with

523 mylosuppressive chemotherapy as compared to chemotherapy alone (See524 ADVERSE REACTIONS: Infection).

- 525 Hematologic toxicity is infrequent following the administration of
- 526 HERCEPTIN as a single agent, with an incidence of Grade III toxicities
- 527 for WBC, platelets, hemoglobin all <1%. No Grade IV toxicities were
- 528 observed.

529 **Diarrhea**:

530 Of patients treated with HERCEPTIN as a single agent, 25% experienced

531 diarrhea. An increased incidence of diarrhea, primarily mild to moderate

- in severity, was observed in patients receiving HERCEPTIN in
- 533 combination with chemotherapy.

534 **Infection:**

- 535 In a randomized, controlled trial (see CLINICAL STUDIES), the
- 536 incidence of infections, primarily mild upper respiratory infections of
- 537 minor clinical significance or catheter infections, was higher (46% vs.
- 538 30%) in patients receiving HERCEPTIN in combination with
- 539 chemotherapy as compared to those receiving chemotherapy alone.
- 540 In a randomized, controlled trial conducted in the post-marketing setting,
- 541 the reported incidence of febrile neutropenia was higher (23% [21/92] vs.
- 542 17% [16/94] in patients receiving HERCEPTIN in combination with
- 543 mylosuppresive chemotherapy as compared to chemotherapy alone.
- 544 In the postmarketing setting there have also been reports of febrile
- 545 neutropenia and infection with neutropenia culminating in death
- 546 associated with the use of HERCEPTIN and myelosuppressive
- 547 chemotherapy (See WARNINGS: Exacerbation of Chemotherapy-Induced
- 548 Neutropenia).

549 **Infusion Reactions:**

- 550 During the first infusion with HERCEPTIN, a symptom complex most
- 551 commonly consisting of chills and/or fever was observed in about 40% of

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552	patients in clinical trials. The symptoms were usually mild to moderate in
553	severity and were treated with acetaminophen, diphenhydramine, and
554	meperidine (with or without reduction in the rate of HERCEPTIN
555	infusion). HERCEPTIN discontinuation was infrequent. Other signs
556	and/or symptoms may include nausea, vomiting, pain (in some cases at
557	tumor sites), rigors, headache, dizziness, dyspnea, hypotension, elevated
558	blood pressure, rash and asthenia. The symptoms occurred infrequently
559	with subsequent HERCEPTIN infusions. (See BOXED WARNINGS:
560	INFUSION REACTIONS and WARNINGS: Infusion Reactions.)
561	Additional adverse reactions have been identified during postmarketing
561 562	Additional adverse reactions have been identified during postmarketing use of HERCEPTIN. Because these reactions are reported voluntarily
561 562 563	Additional adverse reactions have been identified during postmarketing use of HERCEPTIN. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably
561 562 563 564	Additional adverse reactions have been identified during postmarketing use of HERCEPTIN. 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 HERCEPTIN
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561 562 563 564 565 566 567	Additional adverse reactions have been identified during postmarketing use of HERCEPTIN. 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 HERCEPTIN exposure. Decisions to include these reactions in labeling are typically based on one or more of the following factors: (1) seriousness of the reaction, (2) frequency of reporting, or (3) strength of causal connection to
561 562 563 564 565 566 567 568	Additional adverse reactions have been identified during postmarketing use of HERCEPTIN. 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 HERCEPTIN exposure. Decisions to include these reactions in labeling are typically based on one or more of the following factors: (1) seriousness of the reaction, (2) frequency of reporting, or (3) strength of causal connection to HERCEPTIN.

569 Hypersensitivity Reactions Including Anaphylaxis

570 **Pulmonary Events:**

571 In the postmarketing setting, severe hypersensitivity reactions (including

anaphylaxis), infusion reactions, and pulmonary adverse events have been

- 573 reported (see BOXED WARNINGS: HYPERSENSITIVITY
- 574 REACTIONS INCLUDING ANAPHYLAXIS and WARNINGS:
- 575 Hypersensitivity Reactions Including Anaphylaxis). These events include
- 576 anaphylaxis, angioedema, bronchospasm, hypotension, hypoxia, dyspnea,
- 577 pulmonary infiltrates, pleural effusions, non-cardiogenic pulmonary
- 578 edema and acute respiratory distress syndrome. For a detailed description,
- 579 see WARNINGS.

580 **Glomerulopathy:**

- 581 In the postmarketing setting, rare cases of nephrotic syndrome with
- 582 pathologic evidence of glomerulopathy have been reported. The time to
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- 583 onset ranged from 4 months to approximately 18 months from initiation of
- 584 HERCEPTIN therapy. Pathologic findings included membranous
- 585 glomerulonephritis, focal glomerulosclerosis and fibrillary
- 586 glomerulonephritis. Complications included volume overload and
- 587 congestive heart failure.

Table 4

Adverse Events Occurring in \geq 5% of Patients or at Increased Incidence in the HERCEPTIN Arm of the Randomized Study (Percent of Patients)

		i ereem er i u	101105)		
	Single Agent n=352	HERCEPTIN+ Paclitaxel n=91	Paclitaxel Alone n=95	HERCEPTIN +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 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

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Table 4 (cont'd)

Adverse Events Occurring in \geq 5% of Patients or at Increased Incidence in the HERCEPTIN Arm of the Randomized Study (Percent of Patients)

· · · · · · · · · · · · · · · · · · ·	<u>`</u>				
	Single	HERCEPTIN	Paclitaxel	HERCEPTIN	
	Agent	+Paclitaxel	Alone	+AC	AC Alone
	n=352	n=91	n=95	n=143	n=135
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					
Cough increased	26	41	22	43	29
Dyspnea	22	27	26	42	25
Rhinitis	14	22	5	22	16
Pharyngitis	12	22	14	30	18
Sinusitis	9	21	7	13	6
<u>Skin</u>					
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

588

589 Other Serious Adverse Events

590 The following other serious adverse events occurred in at least one of the

591 958 patients treated with HERCEPTIN in clinical studies:

592 Body as a Whole: cellulitis, anaphylactoid reaction, ascites,

593 hydrocephalus, radiation injury, deafness, amblyopia

594 <u>Cardiovascular</u>: vascular thrombosis, pericardial effusion, heart arrest,

595 hypotension, syncope, hemorrhage, shock, arrhythmia

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596 <u>Digestive</u>: hepatic failure, gastroenteritis, hematemesis, ileus, intestinal

597 obstruction, colitis, esophageal ulcer, stomatitis, pancreatitis, hepatitis

598 Endocrine: hypothyroidism

599 <u>Hematological</u>: pancytopenia, acute leukemia, coagulation disorder,
600 lymphangitis

601 Metabolic: hypercalcemia, hypomagnesemia, hyponatremia,

602 hypoglycemia, growth retardation, weight loss

603 <u>Musculoskeletal</u>: pathological fractures, bone necrosis, myopathy

604 <u>Nervous</u>: convulsion, ataxia, confusion, manic reaction

605 <u>Respiratory</u>: apnea, pneumothorax, asthma, hypoxia, laryngitis

- 606 <u>Skin</u>: herpes zoster, skin ulceration
- 607 Urogenital: hydronephrosis, kidney failure, cervical cancer, hematuria,
- 608 hemorrhagic cystitis, pyelonephritis

609 **Immunogenicity**:

610 Of 903 patients who have been evaluated, human anti-human antibody

611 (HAHA) to Trastuzumab was detected in one patient, who had no allergic

- 612 manifestations.
- 613 The data reflect the percentage of patients whose test results were

614 considered positive for antibodies to HERCEPTIN in the HAHA assay for

- 615 Trastuzumab, and are highly dependent on the sensitivity and specificity
- 616 of the assay. Additionally, the observed incidence of antibody positivity
- 617 in an assay may be influenced by several factors including sample
- 618 handling, timing of sample collection, concomitant medications, and
- 619 underlying disease. For these reasons, comparison of the incidence of
- 620 antibodies to HERCEPTIN with the incidence of antibodies to other
- 621 products may be misleading.

622 **OVERDOSAGE**

- 623 There is no experience with overdosage in human clinical trials. Single
- 624 doses higher than 500 mg have not been tested.

625 **DOSAGE AND ADMINISTRATION**

626 Usual Dose

- 627 The recommended initial loading dose is 4 mg/kg Trastuzumab
- 628 administered as a 90-minute infusion. The recommended weekly
- 629 maintenance dose is 2 mg/kg Trastuzumab and can be administered as a
- 630 30-minute infusion if the initial loading dose was well tolerated.
- 631 HERCEPTIN may be administered in an outpatient setting. HERCEPTIN
- 632 is to be diluted in saline for IV infusion. DO NOT ADMINISTER AS
- 633 AN IV PUSH OR BOLUS. (See DOSAGE AND ADMINISTRATION:
- 634 Administration.)

635 **Preparation for Administration**

- 636 The diluent provided has been formulated to maintain the stability and
- 637 sterility of HERCEPTIN for up to 28 days. Other diluents have not been
- 638 shown to contain effective preservatives for HERCEPTIN. Each vial of
- 639 HERCEPTIN should be reconstituted with 20 mL of BWFI, USP,
- 640 1.1% benzyl alcohol preserved, as supplied, to yield a multi-dose solution
- 641 containing 21 mg/mL Trastuzumab. Immediately upon reconstitution with
- 642 BWFI, the vial of HERCEPTIN must be labeled in the area marked "Do
- not use after:" with the future date that is 28 days from the date of
- 644 reconstitution.
- 645 If the patient has known hypersensitivity to benzyl alcohol, HERCEPTIN
- 646 must be reconstituted with Sterile Water for Injection.
- 647 (See PRECAUTIONS.) HERCEPTIN WHICH HAS BEEN
- 648 RECONSTITUTED WITH SWFI MUST BE USED IMMEDIATELY
- 649 AND ANY UNUSED PORTION DISCARDED. USE OF OTHER
- 650 RECONSTITUTION DILUENTS SHOULD BE AVOIDED.

651	Shaking the reconstituted HERCEPTIN or causing excessive foaming					
652	during the addition of diluent may result in problems with dissolution and					
653	the amount of HERCEPTIN that can be withdrawn from the vial.					
654	Use appropriate aseptic technique when performing the following					
655	reconstitution steps:					
656 657 658	a. Using a sterile syringe, slowly inject the 20 mL of diluent into the vial containing the lyophilized cake of Trastuzumab. The stream of diluent should be directed into the lyophilized cake.					
659 660 661	b. Swirl the vial gently to aid reconstitution. Trastuzumab may be sensitive to shear-induced stress, e.g., agitation or rapid expulsion from a syringe. DO NOT SHAKE.					
662 663 664 665	 c. Slight foaming of the product upon reconstitution is not unusual. Allow the vial to stand undisturbed for approximately 5 minutes. The solution should be essentially free of visible particulates, clear to slightly opalescent and colorless to pale yellow. 					
666	Determine the number of mg of Trastuzumab needed, based on a loading					
667	dose of 4 mg Trastuzumab/kg body weight or a maintenance dose of					
668	2 mg Trastuzumab/kg body weight. Calculate the volume of 21 mg/mL					
669	Trastuzumab solution and withdraw this amount from the vial and add it to					
670	an infusion bag containing 250 mL of 0.9% Sodium Chloride Injection,					
671	USP. DEXTROSE (5%) SOLUTION SHOULD NOT BE USED.					
672	Gently invert the bag to mix the solution. The reconstituted preparation					
673	results in a colorless to pale yellow transparent solution. Parenteral drug					
674	products should be inspected visually for particulates and discoloration					
675	prior to administration.					
676	No incompatibilities between HERCEPTIN and polyvinylchloride or					
677	polyethylene bags have been observed.					
678	Administration					
679	Treatment may be administered in an outpatient setting by administration					
680	of a 4 mg/kg Trastuzumab loading dose by intravenous (IV) infusion over					
681	90 minutes. DO NOT ADMINISTER AS AN IV PUSH OR BOLUS.					
682	Patients should be observed for fever and chills or other					
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- 683 infusion-associated symptoms. (See BOXED WARNINGS,
- 684 WARNINGS, and ADVERSE REACTIONS.) If prior infusions are well
- tolerated, subsequent weekly doses of 2 mg/kg Trastuzumab may be
- 686 administered over 30 minutes.
- 687 HERCEPTIN should not be mixed or diluted with other drugs.

688 HERCEPTIN infusions should not be administered or mixed with

689 **Dextrose solutions.**

690 **Stability and Storage**

- 691 Vials of HERCEPTIN are stable at 2–8°C (36–46°F) prior to
- 692 reconstitution. Do not use beyond the expiration date stamped on the vial.
- 693 A vial of HERCEPTIN reconstituted with BWFI, as supplied, is stable for
- 694 28 days after reconstitution when stored refrigerated at $2-8^{\circ}C$ (36–46°F),
- and the solution is preserved for multiple use. Discard any remaining
- 696 multi-dose reconstituted solution after 28 days. If unpreserved SWFI (not
- 697 supplied) is used, the reconstituted HERCEPTIN solution should be used
- 698 immediately and any unused portion must be discarded. DO NOT
- 699 FREEZE HERCEPTIN THAT HAS BEEN RECONSTITUTED.
- 700 The solution of HERCEPTIN for infusion diluted in polyvinylchloride or
- 701 polyethylene bags containing 0.9% Sodium Chloride Injection, USP, may
- be stored at 2-8°C (36-46°F) for up to 24 hours prior to use. Diluted
- HERCEPTIN has been shown to be stable for up to 24 hours at room
- 704 temperature (2-25°C). However, since diluted HERCEPTIN contains no
- r05 effective preservative, the reconstituted and diluted solution should be
- 706 stored refrigerated $(2-8^{\circ}C)$.

707 HOW SUPPLIED

- 708 HERCEPTIN is supplied as a lyophilized, sterile powder nominally
- 709 containing 440 mg Trastuzumab per vial under vacuum.
- 710 Each carton contains one vial of 440 mg HERCEPTIN[®] (Trastuzumab)
- and one vial containing 20 mL of Bacteriostatic Water for Injection, USP,
- 712 1.1% benzyl alcohol. NDC 50242-134-68.
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