## 1.14.1.3 Draft Labeling Text

2 AVASTIN®

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- 3 (Bevacizumab)
- 4 For Intravenous Use

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#### | Gastrointestinal Perforations

- 7 AVASTIN administration can result in the development of gastrointestinal
- 8 perforation, in some instances resulting in fatality. Gastrointestinal
- 9 perforation, sometimes associated with intra-abdominal abscess, occurred
- 10 throughout treatment with AVASTIN (i.e., was not correlated to duration
- of exposure). The incidence of gastrointestinal perforation
- 12 (gastrointestinal perforation, fistula formation, and/or intra-abdominal
- abscess) in patients with colorectal cancer and in patients with non-small
- cell lung cancer (NSCLC) receiving AVASTIN was 2.4% and 0.9%,
- 15 respectively. The typical presentation was reported as abdominal pain
- 16 associated with symptoms such as constipation and vomiting.
- 17 Gastrointestinal perforation should be included in the differential
- diagnosis of patients presenting with abdominal pain on AVASTIN.
- 19 AVASTIN therapy should be permanently discontinued in patients with
- 20 gastrointestinal perforation. (See WARNINGS:
- 21 Gastrointestinal Perforations and DOSAGE AND
- 22 ADMINISTRATION: Dose Modifications.)

## **Wound Healing Complications**

- 24 AVASTIN administration can result in the development of wound
- dehiscence, in some instances resulting in fatality. AVASTIN therapy
- should be permanently discontinued in patients with wound dehiscence
- 27 | requiring medical intervention. The appropriate interval between
- termination of AVASTIN and subsequent elective surgery required to
- 29 avoid the risks of impaired wound healing/wound dehiscence has not been
- 30 determined. (See WARNINGS: Wound Healing Complications and
- 31 **DOSAGE AND ADMINISTRATION:** Dose Modifications.)

# 32 Hemorrhage

- 33 Fatal pulmonary hemorrhage can occur in patients with NSCLC treated
- with chemotherapy and AVASTIN. The incidence of severe or fatal
- hemoptysis was 31% in patients with squamous histology and 2.3% in
- patients with NSCLC excluding predominant squamous histology.
- 37 Patients with recent hemoptysis (≥1/2 tsp of red blood) should not receive
- 38 AVASTIN. (See WARNINGS: Hemorrhage,
- 39 ADVERSE REACTIONS: Hemorrhage, and
- 40 DOSAGE AND ADMINISTRATION: Dose Modifications.)

#### DESCRIPTION

- 42 AVASTIN® (Bevacizumab) is a recombinant humanized monoclonal
- 43 IgG1 antibody that binds to and inhibits the biologic activity of human
- 44 vascular endothelial growth factor (VEGF) in in vitro and in vivo assay
- 45 systems. Bevacizumab contains human framework regions and the
- 46 complementarity-determining regions of a murine antibody that binds to
- 47 VEGF (1). Bevacizumab is produced in a Chinese Hamster Ovary
- 48 mammalian cell expression system in a nutrient medium containing the
- 49 antibiotic gentamicin and has a molecular weight of approximately
- 50 149 kilodaltons. AVASTIN is a clear to slightly opalescent, colorless to
- 51 pale brown, sterile, pH 6.2 solution for intravenous (IV) infusion.
- 52 AVASTIN is supplied in 100 mg and 400 mg preservative-free, single-use
- vials to deliver 4 mL or 16 mL of AVASTIN (25 mg/mL). The 100 mg
- product is formulated in 240 mg  $\alpha$ ,  $\alpha$ -trehalose dihydrate, 23.2 mg sodium
- 55 phosphate (monobasic, monohydrate), 4.8 mg sodium phosphate (dibasic,
- anhydrous), 1.6 mg polysorbate 20, and Water for Injection, USP. The
- 400 mg product is formulated in 960 mg  $\alpha$ ,  $\alpha$ -trehalose dihydrate, 92.8 mg
- sodium phosphate (monobasic, monohydrate), 19.2 mg sodium phosphate
- 59 (dibasic, anhydrous), 6.4 mg polysorbate 20, and Water for Injection,
- 60 USP.

#### CLINICAL PHARMACOLOGY

62 Mechanism of Action

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- 63 Bevacizumab binds VEGF and prevents the interaction of VEGF to its
- receptors (Flt-1 and KDR) on the surface of endothelial cells. The
- 65 interaction of VEGF with its receptors leads to endothelial cell
- proliferation and new blood vessel formation in in vitro models of
- angiogenesis. Administration of Bevacizumab to xenotransplant models
- of colon cancer in nude (athymic) mice caused reduction of microvascular
- 69 growth and inhibition of metastatic disease progression.

#### 70 Pharmacokinetics

- 71 The pharmacokinetic profile of Bevacizumab was assessed using an assay
- 72 that measures total serum Bevacizumab concentrations (i.e., the assay did
- 73 not distinguish between free Bevacizumab and Bevacizumab bound to
- 74 VEGF ligand). Based on a population pharmacokinetic analysis of
- 75 491 patients who received 1 to 20 mg/kg of AVASTIN weekly, every
- 76 2 weeks, or every 3 weeks, the estimated half-life of Bevacizumab was
- approximately 20 days (range 11–50 days). The predicted time to reach
- 78 steady state was 100 days. The accumulation ratio following a dose of
- 79 10 mg/kg of Bevacizumab every 2 weeks was 2.8.
- 80 The clearance of Bevacizumab varied by body weight, by gender, and by
- 81 tumor burden. After correcting for body weight, males had a higher
- 82 Bevacizumab clearance (0.262 L/day vs. 0.207 L/day) and a larger V<sub>c</sub>
- 83 (3.25 L vs. 2.66 L) than females. Patients with higher tumor burden (at or
- 84 above median value of tumor surface area) had a higher Bevacizumab
- clearance (0.249 L/day vs. 0.199 L/day) than patients with tumor burdens
- below the median. In a randomized study of 813 patients (Study 1), there
- was no evidence of lesser efficacy (hazard ratio for overall survival) in
- males or patients with higher tumor burden treated with AVASTIN as
- 89 compared to females and patients with low tumor burden. The
- 90 relationship between Bevacizumab exposure and clinical outcomes has not
- 91 been explored.

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92	Special Populations
93	Analyses of demographic data suggest that no dose adjustments are
94	necessary for age or sex.
95	Patients with renal impairment. No studies have been conducted to
96	examine the pharmacokinetics of Bevacizumab in patients with renal
97	impairment.
98	Patients with hepatic dysfunction. No studies have been conducted to
99	examine the pharmacokinetics of Bevacizumab in patients with hepatic
100	impairment.
101	CLINICAL STUDIES
102	AVASTIN® In Metastatic Colorectal Cancer (mCRC)
103	The safety and efficacy of AVASTIN in the treatment of patients with
104	metastatic carcinoma of the colon or rectum were studied in three
105	randomized, controlled clinical trials in combination with intravenous
106	5-fluorouracil-based chemotherapy. The activity of AVASTIN in patient
107	with metastatic colorectal cancer that progressed on or after receiving both
108	irinotecan based- and oxaliplatin based-chemotherapy regimens was
109	evaluated in an open-access trial in combination with intravenous
110	5-fluorouracil-based chemotherapy.
111	AVASTIN in Combination with Bolus-IFL
112	Study 1 was a randomized, double-blind, active-controlled clinical trial
113	evaluating AVASTIN as first-line treatment of metastatic carcinoma of th
114	colon or rectum. Patients were randomized to bolus-IFL (irinotecan
115	125 mg/m <sup>2</sup> IV, 5-fluorouracil 500 mg/m <sup>2</sup> IV, and leucovorin 20 mg/m <sup>2</sup> IV
116	given once weekly for 4 weeks every 6 weeks) plus placebo (Arm 1),
117	bolus-IFL plus AVASTIN (5 mg/kg every 2 weeks) (Arm 2), or 5-FU/LV
118	plus AVASTIN (5 mg/kg every 2 weeks) (Arm 3). Enrollment in Arm 3

was discontinued, as pre-specified, when the toxicity of AVASTIN in

combination with the bolus-IFL regimen was deemed acceptable.

Of the 813 patients randomized to Arms 1 and 2, the median age was 60, 40% were female, and 79% were Caucasian. Fifty-seven percent had an ECOG performance status of 0. Twenty-one percent had a rectal primary and 28% received prior adjuvant chemotherapy. In the majority of patients, 56%, the dominant site of disease was extra-abdominal, while the liver was the dominant site in 38% of patients. Results are presented in Table 1 and Figure 1.

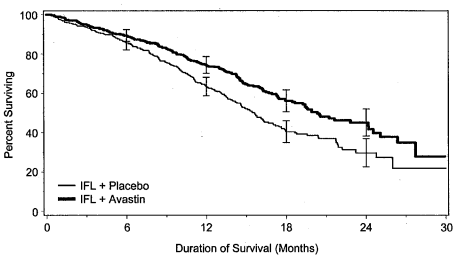
**Table 1**Study 1 Efficacy Results

	IFL+Placebo	IFL+AVASTIN 5 mg/kg q 2 wks
Number of Patients	411	402
Overall Survival <sup>a</sup>		
Median (months)	15.6	20.3
Hazard ratio		0.66
Progression-free Survivala		
Median (months)	6.2	10.6
Hazard ratio		0.54
Overall Response Rate <sup>b</sup>		
Rate (percent)	35%	45%
Duration of Response		
Median (months)	7.1	10.4

<sup>&</sup>lt;sup>a</sup>p<0.001 by stratified logrank test.

 $<sup>^{</sup>b}p<0.01$  by  $\chi^{2}$  test.

**Figure 1**Duration of Survival in Study 1



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Error bars represent 95% confidence intervals.

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The clinical benefit of AVASTIN, as measured by survival in the two principal arms, was seen in the subgroups defined by age (<65 yrs,  $\ge65$  yrs) and gender.

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Among the 110 patients enrolled in Arm 3, median overall survival was 18.3 months, median progression-free survival was 8.8 months, overall response rate was 39%, and median duration of response was 8.5 months.

# 140 AVASTIN in Combination with 5-FU/LV Chemotherapy

Study 2 was a randomized, active-controlled clinical trial testing AVASTIN in combination with 5-FU/LV as first-line treatment of metastatic colorectal cancer. Patients were randomized to receive 5-FU/LV (5-fluorouracil 500 mg/m², leucovorin 500 mg/m² weekly for 6 weeks every 8 weeks) or 5-FU/LV plus AVASTIN (5 mg/kg every 2 weeks) or 5-FU/LV plus AVASTIN (10 mg/kg every 2 weeks). The primary endpoints of the trial were objective response rate and progression-free survival. Results are presented in Table 2.

Table 2 Study 2 Efficacy Results

	5-FU/LV	5-FU/LV+AVASTIN 5 mg/kg	5-FU/LV+AVASTIN 10 mg/kg
Number of Patients	36	35	33
Overall Survival  Median (months)	13.6	17.7	15.2
Progression-free Survival Median (months)	5.2	9.0	7.2
Overall Response Rate Rate (percent)	17	40	24

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Progression-free survival was significantly longer in patients receiving 150

5-FU/LV plus AVASTIN at 5 mg/kg when compared to those not 151

receiving AVASTIN. However, overall survival and overall response rate

were not significantly different. Outcomes for patients receiving 5-FU/LV 153

154 plus AVASTIN at 10 mg/kg were not significantly different than for

patients who did not receive AVASTIN. 155

# AVASTIN in Combination with 5-FU/LV and Oxaliplatin Chemotherapy

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Study 3 was an open-label, randomized, 3-arm, active-controlled,

159 multicenter clinical trial evaluating AVASTIN alone, AVASTIN in

160 combination with 5-FU/LV and oxaliplatin (FOLFOX4), and FOLFOX4

alone in the second-line treatment of metastatic carcinoma of the colon or

162 rectum. Patients were previously treated with irinotecan and 5-FU for

initial therapy for metastatic disease or as adjuvant therapy. Patients were 163

randomized to FOLFOX4 (Day 1: oxaliplatin 85 mg/m<sup>2</sup> and leucovorin 164

200 mg/m<sup>2</sup> concurrently IV, then 5-FU 400 mg/m<sup>2</sup> IV bolus followed by 165

600 mg/m<sup>2</sup> continuously IV: Day 2: leucovorin 200 mg/m<sup>2</sup> IV, then 5-FU 166

400 mg/m<sup>2</sup> IV bolus followed by 600 mg/m<sup>2</sup> continuously IV; repeated

every 2 weeks), FOLFOX4 plus AVASTIN, or AVASTIN monotherapy. 168

AVASTIN was administered at a dose of 10 mg/kg every 2 weeks and for 169

1/0	patients in the FOLFOX4 plus AVASTIN arm, prior to the FOLFOX4
171	chemotherapy on Day 1.
172	Of the 829 patients randomized to the three arms, the median age was
173	61 years, 40% were female, 87% were Caucasian, and 49% had an ECOG
174	performance status of 0. Twenty-six percent had received prior radiation
175	therapy, and 80% received prior adjuvant chemotherapy. Ninety-nine
176	percent received prior irinotecan, with or without 5-FU for metastatic
177	colorectal cancer, and 1% received prior irinotecan and 5-FU as adjuvant
178	therapy.
179	The AVASTIN monotherapy arm of Study 3 was closed to accrual after
180	enrollment of 244 of the planned 290 patients following a planned interim
181	analysis by the data monitoring committee (DMC), based on evidence of
182	decreased survival in the AVASTIN alone arm as compared to the
183	FOLFOX4 alone arm. In the two remaining study arms, overall survival
184	(OS) was significantly longer in patients receiving AVASTIN in
185	combination with FOLFOX4 as compared to those receiving FOLFOX4
186	alone (median OS 13.0 mos vs. 10.8 mos; hazard ratio 0.75 [95% CI 0.63
187	0.89], p=0.001 stratified log rank test). In addition, patients treated with
188	AVASTIN in combination with FOLFOX4 were reported to have
189	significantly longer progression-free survival and a higher overall
190	response rate based on investigator assessment. The clinical benefit of
191	AVASTIN, as measured by survival, was seen in the subgroups defined by
192	age (<65 yrs, ≥65 yrs) and gender.
193	AVASTIN in Third-Line Metastatic Colorectal Cancer
194	Study 4 was an open access, multicenter, single arm study that evaluated
195	the activity of AVASTIN in combination with bolus or infusional
196	5-FU/LV in 339 patients with metastatic colorectal cancer with disease
197	progression following both irinotecan- and oxaliplatin-containing
198	chemotherapy regimens. The majority (73%) of patients received
199	concurrent 5-FU/LV according to a bolus regimen.

200	There was one objective partial response in the first 100 evaluable patients
201	for an overall response rate of 1% (95% CI 0-5.5%).
202 203	AVASTIN <sup>®</sup> In Unresectable Non-Squamous, Non-Small Cell Lung Cancer (NSCLC)
204	The safety and efficacy of AVASTIN as first-line treatment of patients
205	with locally advanced, metastatic, or recurrent non-squamous, NSCLC
206	was studied in a single, large, randomized, active-controlled, open-label,
207	multicenter study (Study 5, n=878), supported by a randomized, dose
208	ranging, active controlled Phase 2 study (Study 6, n=98).
209	In Study 5, chemotherapy-naïve patients with locally advanced, metastatic
210	or recurrent non-squamous NSCLC were randomized (1:1) to receive six
211	cycles of paclitaxel 200 mg/m <sup>2</sup> and carboplatin AUC=6.0, both by IV
212	infusion on day 1 (PC) or PC in combination with AVASTIN at a dose of
213	15 mg/kg by IV infusion on day 1 (PC plus AVASTIN). After completion
214	or upon discontinuation of chemotherapy, patients in the PC plus
215	AVASTIN arm continued to receive AVASTIN alone until disease
216	progression or until unacceptable toxicity. Cycles were repeated every
217	21 days. Patients with predominant squamous histology (mixed cell type
218	tumors only), central nervous system (CNS) metastasis, gross hemoptysis
219	(≥1/2 tsp of red blood), or unstable angina and those receiving therapeutic
220	anticoagulation were excluded. The main outcome measure of the study
221	was duration of survival.
222	Among the 878 patients randomized to the two treatment arms, the median
223	age was 63, 46% were female, 43% were $\geq$ age 65, and 28% had $\geq$ 5%
224	weight loss at study entry. Eleven percent had recurrent disease and of the
225	remaining 89% with newly diagnosed NSCLC, 12% had Stage IIIB with
226	malignant pleural effusion and 76% had Stage IV disease. The survival
227	curves are presented in Figure 2. Overall survival was statistically
228.	significantly higher among patients receiving PC plus AVASTIN
229	compared with those receiving PC alone; median OS was 12.3 mos vs.
230	10.3 mos (hazard ratio 0.80 [repeated 95% CI 0.68, 0.94], final p-value

0.013, stratified log-rank test). Based on investigator assessment which was not independently verified, patients were reported to have longer progression-free survival with AVASTIN in combination with PC compared to PC alone.

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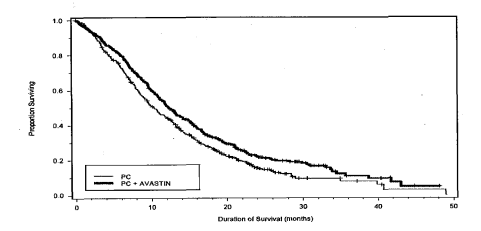
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Figure 2
Duration of Survival in Study 5



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In an exploratory analyses across patient subgroups, the impact of

240 AVASTIN on overall survival was less robust in the following: women

241 [HR=0.99 (95% CI: 0.79, 1.25)], age  $\geq$  65 years [HR=0.91 (95% CI:

242 0.72, 1.14)] and patients with  $\geq$ 5% weight loss at study entry [HR=0.96

243 (95% CF: 0.73, 1.26)].

# INDICATIONS AND USAGE

AVASTIN®, in combination with intravenous 5-fluorouracil-based

246 chemotherapy, is indicated for first- or second-line treatment of patients

with metastatic carcinoma of the colon or rectum.

AVASTIN®, in combination with carboplatin and paclitaxel, is indicated

249 for first-line treatment of patients with unresectable, locally advanced,

recurrent or metastatic non-squamous, non-small cell lung cancer.

**CONTRAINDICATIONS** 

252	None.
253	WARNINGS
254 255	Gastrointestinal Perforations (See DOSAGE AND ADMINISTRATION: Dose Modifications)
256	Gastrointestinal perforation complicated by intra-abdominal abscesses or
257	fistula formation and in some instances with fatal outcome, occurs at an
258	increased incidence in patients receiving AVASTIN as compared to
259	controls. In Studies 1, 2, and 3, the incidence of gastrointestinal
260	perforation (gastrointestinal perforation, fistula formation, and/or
261	intra-abdominal abscess) in patients receiving AVASTIN was 2.4%.
262	These episodes occurred with or without intra-abdominal abscesses and at
263	various time points during treatment. The typical presentation was
264	reported as abdominal pain associated with symptoms such as constipation
265	and emesis.
266	In post-marketing clinical studies and reports, gastrointestinal perforation,
267	fistula formation in the gastrointestinal tract (eg. gastrointestinal,
268	enterocutaneous, esophageal, duodenal, rectal), and/or intra-abdominal
269	abscess occurred in patients receiving AVASTIN for colorectal and for
270	other types of cancer. The overall incidence in clinical studies was 1%,
271	but may be higher in some cancer settings. Of the reported events,
272	approximately 30% were fatal. Patients with gastrointestinal perforation,
273	regardless of underlying cancer, typically present with abdominal pain,
274	nausea and fever. Events were reported at various time points during
275	treatment ranging from one week to greater than 1 year from initiation of
276	AVASTIN, with most events occurring within the first 50 days.
277	Permanently discontinue AVASTIN in patients with gastrointestinal
278	perforation (gastrointestinal perforation, fistula formation, and/or
279	intra-abdominal abscess).

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281	ADMINISTRATION: Dose Modifications)
282	Non-gastrointestinal fistula formation has been reported in patients treated
283	with AVASTIN in controlled clinical studies (with an incidence of
284	< 0.3%) and in post-marketing experience, in some cases with fatal
285	outcome. Fistula formation involving the following areas of the body
286	other than the gastrointestinal tract have been reported:
287	tracheo-esophageal, bronchopleural, biliary, vagina and bladder. Events
288	were reported throughout treatment with Avastin, with most events
289	occurring within the first 6 months.
290	Permanently discontinue AVASTIN in patients with fistula formation
291	involving an internal organ.
292	Wound Healing Complications (See DOSAGE AND
292	ADMINISTRATION: Dose Modifications)
294	AVASTIN impairs wound healing in animal models. In clinical studies of
295	AVASTIN, patients were not allowed to receive AVASTIN until at least
296	28 days had elapsed following surgery. In clinical studies of AVASTIN in
297	combination with chemotherapy, there were 6 instances of dehiscence
298	among 788 patients (0.8%).
299	The appropriate interval between discontinuation of AVASTIN and
300	subsequent elective surgery required to avoid the risks of impaired wound
301	healing has not been determined. In Study 1, 39 patients who received
302	bolus-IFL plus AVASTIN underwent surgery following AVASTIN
303	therapy; of these patients, six (15%) had wound healing/bleeding
304	complications. In the same study, 25 patients in the bolus-IFL arm
305	underwent surgery; of these patients, one of 25 (4%) had wound
306	healing/bleeding complications. The longest interval between last dose of
307	study drug and dehiscence was 56 days; this occurred in a patient on the
308	bolus-IFL plus AVASTIN arm.

309	The interval between termination of AVASTIN and subsequent elective
310	surgery should take into consideration the calculated half-life of
311	AVASTIN (approximately 20 days).
312	Discontinue AVASTIN in patients with wound healing complications
313	requiring medical intervention.
314	Hemorrhage (See DOSAGE AND ADMINISTRATION:
315	Dose Modifications)
316	Two distinct patterns of bleeding have occurred in patients receiving
317	AVASTIN. The first is minor hemorrhage, most commonly NCI-CTC
318	Grade 1 epistaxis. The second is serious, and in some cases fatal,
319	hemorrhagic events.
320	In Study 6, four of 13 (31%) AVASTIN-treated patients with squamous
321	cell histology and two of 53 (4%) AVASTIN-treated patients with
322	histology other than squamous cell, experienced serious or fatal
323	pulmonary hemorrhage as compared to none of the 32 (0%) patients
324	receiving chemotherapy alone. Of the patients experiencing pulmonary
325	hemorrhage requiring medical intervention, many had cavitation and/or
326	necrosis of the tumor, either pre-existing or developing during AVASTI
327	therapy. In Study 5, the rate of pulmonary hemorrhage requiring medica
328	intervention for the PC plus AVASTIN arm was 2.3% (10 of 427)
329	compared to 0.5% (2 of 441) for the PC alone arm. There were seven
330	deaths due to pulmonary hemorrhage reported by investigators in the PC
331	plus AVASTIN arm as compared to one in the PC alone arm. Generally,
332	these serious hemorrhagic events presented as major or massive
333	hemoptysis without an antecedent history of minor hemoptysis during
334	Avastin therapy. Do not administer AVASTIN to patients with recent
335	history of hemoptysis of $\geq 1/2$ tsp of red blood. Other serious bleeding
336	events occurring in patients receiving AVASTIN across all indications
337	include gastrointestinal hemorrhage, subarachnoid hemorrhage, and
338	hemorrhagic stroke. Some of these events were fatal. (See ADVERSE
339	REACTIONS: Hemorrhage.)

340	The risk of central nervous system (CNS) bleeding in patients with CNS
341	metastases receiving AVASTIN has not been evaluated because these
342	patients were excluded from late stage clinical studies following
343	development of CNS hemorrhage in a patient with a CNS metastasis in a
344	Phase 1 study.
345	Discontinue AVASTIN in patients with serious hemorrhage (i.e., requiring
346	medical intervention) and initiate aggressive medical management.
347	(See ADVERSE REACTIONS: Hemorrhage.)
348 349 350	Arterial Thromboembolic Events (see DOSAGE AND ADMINISTRATION: Dose Modifications and PRECAUTIONS: Geriatric Use)
351	Arterial thromboembolic events (ATE) occurred at a higher incidence in
352	patients receiving AVASTIN in combination with chemotherapy as
353	compared to those receiving chemotherapy alone. ATE included cerebral
354	infarction, transient ischemic attacks (TIAs), myocardial infarction (MI),
355	angina, and a variety of other ATE. These events were fatal in some
356	instances.
357	In a pooled analysis of randomized, controlled clinical trials involving
358	1745 patients, the incidence of ATE was 4.4% among patients treated with
359	AVASTIN in combination with chemotherapy and 1.9% among patients
360	receiving chemotherapy alone. Fatal outcomes for these events occurred
361	in 7 of 963 patients (0.7%) who were treated with AVASTIN in
362	combination with chemotherapy, compared to 3 of 782 patients (0.4%)
363	who were treated with chemotherapy alone. The incidences of both
364	cerebrovascular arterial events (1.9% vs. 0.5%) and cardiovascular arterial
365	events (2.1% vs. 1.0%) were increased in patients receiving AVASTIN
366	compared to chemotherapy alone. The relative risk of ATE was greater in
367	patients 65 and over (8.5% vs. 2.9%) as compared to those less than 65
368	(2.1% vs. 1.4%). (See PRECAUTIONS: Geriatric Use.)
369	The safety of resumption of AVASTIN therapy after resolution of an ATE
370	has not been studied. Permanently discontinue AVASTIN in patients who

371	experience a severe ATE during treatment. (See DOSAGE AND
372	ADMINISTRATION: Dose Modifications and PRECAUTIONS:
373	Geriatric Use.)
374	Hypertension (See DOSAGE AND ADMINISTRATION:
375	Dose Modifications)
376	The incidence of severe hypertension was increased in patients receiving
377	AVASTIN as compared to controls. Across clinical studies the incidence
378	of NCI-CTC Grade 3 or 4 hypertension ranged from 8-18%.
379	Medication classes used for management of patients with NCI-CTC
380	Grade 3 hypertension receiving AVASTIN included
381	angiotensin-converting enzyme inhibitors, beta blockers, diuretics, and
382	calcium channel blockers. Development or worsening of hypertension can
383	require hospitalization or require discontinuation of AVASTIN in up to
384	1.7% of patients. Hypertension can persist after discontinuation of
385	AVASTIN. Complications can include hypertensive encephalopathy
386	(in some cases fatal) and CNS hemorrhage.
387	In the post-marketing experience, acute increases in blood pressure
388	associated with initial or subsequent infusions of AVASTIN have been
389	reported (see PRECAUTIONS: Infusion Reactions). Some cases were
390	serious and associated with clinical sequelae.
391	Permanently discontinue AVASTIN in patients with hypertensive crisis or
392	hypertensive encephalopathy. Temporarily suspend AVASTIN in patients
393	with severe hypertension that is not controlled with medical management.
394	(See DOSAGE AND ADMINISTRATION: Dose Modifications.)
395	Reversible Posterior Leukoencephalopathy Syndrome (RPLS)
396	(See DOSAGE AND ADMINISTRATION: Dose Modifications)
397	RPLS has been reported in clinical studies (with an incidence of <0.1%)
398	and in post-marketing experience. RPLS is a neurological disorder which
399	can present with headache, seizure, lethargy, confusion, blindness and
400	other visual and neurologic disturbances. Mild to severe hypertension

401	may be present, but is not necessary for diagnosis of RPLS. Magnetic
402	Resonance Imaging (MRI) is necessary to confirm the diagnosis of RPLS.
403	The onset of symptoms has been reported to occur from 16 hours to 1 year
404	after initiation of AVASTIN.
405	In patients developing RPLS, discontinue AVASTIN and initiate
406	treatment of hypertension, if present. Symptoms usually resolve or
407	improve within days, although some patients have experienced ongoing
408	neurologic sequelae. The safety of reinitiating AVASTIN therapy in
409	patients previously experiencing RPLS is not known.
410 411	Neutropenia and Infection (See PRECAUTIONS: Geriatric Use and ADVERSE REACTIONS: Neutropenia and Infection)
412	Increased rates of severe neutropenia, febrile neutropenia, and infection
413	with severe neutropenia (including some fatalities) have been observed in
414	patients treated with myelosuppressive chemotherapy plus AVASTIN.
415	(See PRECAUTIONS: Geriatric Use and ADVERSE REACTIONS:
416	Neutropenia and Infection.)
417 418	Proteinuria (See DOSAGE AND ADMINISTRATION: Dose Modifications)
419	The incidence and severity of proteinuria is increased in patients receiving
420	AVASTIN as compared to control. In Studies 1, 3 and 5 the incidence of
421	NCI-CTC Grade 3 and 4 proteinuria, characterized as >3.5 gm/24 hours,
422	ranged up to 3.0% in AVASTIN-treated patients.
423	Nephrotic syndrome occurred in seven of 1459 (0.5%) patients receiving
424	AVASTIN in clinical studies. One patient died and one required dialysis.
425	In three patients, proteinuria decreased in severity several months after
426	discontinuation of AVASTIN. No patient had normalization of urinary
427	protein levels (by 24-hour urine) following discontinuation of AVASTIN.
428	The highest incidence of proteinuria was observed in a dose-ranging,
429	placebo-controlled, randomized study of AVASTIN in patients with
430	metastatic renal cell carcinoma, an indication for which AVASTIN is not

BL125085/131 Amendment: Bevacizumab—Genentech, Inc. 16 of 38/Regional (Fistula): 1-14-1-3.doc

431	approved, 24-hour urine collections were obtained in approximately half		
432	the patients enrolled. Among patients in whom 24-hour urine collections		
433	were obtained, four of 19 (21%) patients receiving AVASTIN at 10 mg/kg		
434	every two weeks, two of 14 (14%) patients receiving AVASTIN at		
435	3 mg/kg every two weeks, and none of the 15 placebo patients		
436	experienced NCI-CTC Grade 3 proteinuria (>3.5 gm protein/24 hours).		
437	Discontinue AVASTIN in patients with nephrotic syndrome. The safety		
438	of continued AVASTIN treatment in patients with moderate to severe		
439	proteinuria has not been evaluated. In most clinical studies, AVASTIN		
440	was interrupted for ≥2 grams of proteinuria/24 hours and resumed when		
441	proteinuria was <2 gm/24 hours. Patients with moderate to severe		
442	proteinuria based on 24-hour collections should be monitored regularly		
443	until improvement and/or resolution is observed. (See DOSAGE AND		
444	ADMINISTRATION: Dose Modifications.)		
445	Congestive Heart Failure		
446	Congestive heart failure (CHF), defined as NCI-CTC Grade 2-4 left		
447	ventricular dysfunction, was reported in 25 of 1459 (1.7%) patients		
448	receiving AVASTIN in clinical studies. The risk of CHF appears to be		
449			
	higher in patients receiving AVASTIN who have received prior or		
450	higher in patients receiving AVASTIN who have received prior or concurrent anthracyclines. In a controlled study in patients with breast		
450 451			
	concurrent anthracyclines. In a controlled study in patients with breast		
451	concurrent anthracyclines. In a controlled study in patients with breast cancer (an unlabelled indication), the incidence of CHF was higher in the		
451 452	concurrent anthracyclines. In a controlled study in patients with breast cancer (an unlabelled indication), the incidence of CHF was higher in the AVASTIN plus chemotherapy arm as compared to the chemotherapy		
451 452 453	concurrent anthracyclines. In a controlled study in patients with breast cancer (an unlabelled indication), the incidence of CHF was higher in the AVASTIN plus chemotherapy arm as compared to the chemotherapy alone arm. Congestive heart failure occurred in 13 of 299 (4%) patients		
451 452 453 454	concurrent anthracyclines. In a controlled study in patients with breast cancer (an unlabelled indication), the incidence of CHF was higher in the AVASTIN plus chemotherapy arm as compared to the chemotherapy alone arm. Congestive heart failure occurred in 13 of 299 (4%) patients who received prior anthracyclines and/or left chest wall irradiation.		
451 452 453 454 455	concurrent anthracyclines. In a controlled study in patients with breast cancer (an unlabelled indication), the incidence of CHF was higher in the AVASTIN plus chemotherapy arm as compared to the chemotherapy alone arm. Congestive heart failure occurred in 13 of 299 (4%) patients who received prior anthracyclines and/or left chest wall irradiation. Congestive heart failure occurred in six of 44 (14%) patients with relapsed		
451 452 453 454 455 456	concurrent anthracyclines. In a controlled study in patients with breast cancer (an unlabelled indication), the incidence of CHF was higher in the AVASTIN plus chemotherapy arm as compared to the chemotherapy alone arm. Congestive heart failure occurred in 13 of 299 (4%) patients who received prior anthracyclines and/or left chest wall irradiation. Congestive heart failure occurred in six of 44 (14%) patients with relapsed acute leukemia (an unlabelled indication) receiving AVASTIN and		

460	PRECAUTIONS
461	General
462	Use AVASTIN with caution in patients with known hypersensitivity to
463	AVASTIN or any component of this drug product.
464	Infusion Reactions
465	In clinical studies, infusion reactions with the first dose of AVASTIN
466	were uncommon (<3%) and severe reactions occurred in 0.2% of patients.
467	Infusion reactions reported in the clinical trials and post-marketing
468	experience include hypertension, hypertensive crises associated with
469	neurologic signs and symptoms, wheezing, oxygen desaturation,
470	NCI-CTC Grade 3 hypersensitivity, chest pain, headaches, rigors, and
471	diaphoresis. Adequate information on rechallenge is not available.
472	AVASTIN infusion should be interrupted in all patients with severe
473	infusion reactions and appropriate medical therapy administered.
474	There are no data regarding the most appropriate method of identification
475	of patients who may safely be retreated with AVASTIN after experiencing
476	a severe infusion reaction.
477	Surgery
478	AVASTIN therapy should not be initiated for at least 28 days following
479	major surgery. The surgical incision should be fully healed prior to
480	initiation of AVASTIN. Because of the potential for impaired wound
481	healing, AVASTIN should be suspended prior to elective surgery.
482	The appropriate interval between the last dose of AVASTIN and elective
483	surgery is unknown; however, the half-life of AVASTIN is estimated to be
484	20 days (see CLINICAL PHARMACOLOGY: Pharmacokinetics) and
485	the interval chosen should take into consideration the half-life of the drug.
486	(See WARNINGS: Gastrointestinal Perforations and
487	Wound Healing Complications.)

488	Cardiovascular Disease			
489	Patients were excluded from participation in AVASTIN clinical trials if, in			
490	the previous year, they had experienced clinically significant			
<b>49</b> 1	cardiovascular disease. In an exploratory analysis pooling the data from			
492	five randomized, placebo-controlled, clinical trials conducted in patients			
493	without a recent history of clinically significant cardiovascular disease, the			
494	overall incidence of arterial thromboembolic events, the incidence of fatal			
495	arterial thromboembolic events, and the incidence of cardiovascular			
496	thromboembolic events were increased in patients receiving AVASTIN			
497	plus chemotherapy as compared to chemotherapy alone.			
498	Laboratory Tests			
499	Blood pressure monitoring should be conducted every two to three weeks			
500	during treatment with AVASTIN. Patients who develop hypertension on			
501	AVASTIN may require blood pressure monitoring at more frequent			
502	intervals. Patients with AVASTIN-induced or -exacerbated hypertension			
503	who discontinue AVASTIN should continue to have their blood pressure			
504	monitored at regular intervals.			
505	Patients receiving AVASTIN should be monitored for the development or			
506	worsening of proteinuria with serial urinalyses. Patients with a 2+ or			
507	greater urine dipstick reading should undergo further assessment, e.g., a			
508	24-hour urine collection. (See WARNINGS: Proteinuria and DOSAGE			
509	AND ADMINISTRATION: Dose Modifications.)			
510	Drug Interactions			
511	No formal drug interaction studies with anti-neoplastic agents have been			
512	conducted. In Study 1, patients with colorectal cancer were given			
513	irinotecan/5-FU/leucovorin (bolus-IFL) with or without AVASTIN.			
514	Irinotecan concentrations were similar in patients receiving bolus-IFL			
515	alone and in combination with AVASTIN. The concentrations of SN38,			
516	the active metabolite of irinotecan, were on average 33% higher in patients			
517	receiving bolus-IFL in combination with AVASTIN when compared with			
518	bolus-IFL alone. In Study 1, patients receiving bolus-IFL plus AVASTIN			

519	had a higher incidence of NCI-CTC Grade 3–4 diarrhea and neutropenia.		
520	Due to high inter-patient variability and limited sampling, the extent of the		
521	increase in SN38 levels in patients receiving concurrent irinotecan and		
522	AVASTIN is uncertain.		
523	In Study 6, based on limited data, there did not appear to be a difference in		
524	the mean exposure of either carboplatin or paclitaxel when each was		
525	administered alone or in combination with AVASTIN. However, 3 of the		
526	8 patients receiving AVASTIN plus paclitaxel/carboplatin had		
527	substantially lower paclitaxel exposure after four cycles of treatment (at		
528	Day 63) than those at Day 0, while patients receiving		
529	paclitaxel/carboplatin without AVASTIN had a greater paclitaxel		
530	exposure at Day 63 than at Day 0.		
531	Carcinogenesis, Mutagenesis, Impairment of Fertility		
532	No carcinogenicity data are available for AVASTIN in animals or		
533	humans.		
534	AVASTIN may impair fertility. Dose-related decreases in ovarian and		
535	uterine weights, endometrial proliferation, number of menstrual cycles,		
536	and arrested follicular development or absent corpora lutea were observed		
537	in female cynomolgus monkeys treated with 10 or 50 mg/kg of AVASTIN		
538	for 13 or 26 weeks. Following a 4- or 12-week recovery period, which		
539	examined only the high-dose group, trends suggestive of reversibility		
540	were noted in the two females for each regimen that were assigned to		
541	recover. After the 12-week recovery period, follicular maturation arrest		
542	was no longer observed, but ovarian weights were still moderately		
543	decreased. Reduced endometrial proliferation was no longer observed at		
544	the 12-week recovery time point, but uterine weight decreases were still		
545	notable, corpora lutea were absent in 1 out of 2 animals, and the number of		
546	menstrual cycles remained reduced (67%).		

547	Pregnancy Category C			
548	AVASTIN has been shown to be teratogenic in rabbits when administered			
549	in doses that approximate the human dose on a mg/kg basis. Observed			
550	effects included decreases in maternal and fetal body weights, an			
551	increased number of fetal resorptions, and an increased incidence of			
552	specific gross and skeletal fetal alterations. Adverse fetal outcomes were			
553	observed at all doses tested.			
554	Angiogenesis is critical to fetal development and the inhibition of			
555	angiogenesis following administration of AVASTIN is likely to result in			
556	adverse effects on pregnancy. There are no adequate and well-controlled			
557	studies in pregnant women. AVASTIN should be used during pregnancy			
558	or in any woman not employing adequate contraception only if the			
559	potential benefit justifies the potential risk to the fetus. All patients should			
560	be counseled regarding the potential risk of AVASTIN to the developing			
561	fetus prior to initiation of therapy. If the patient becomes pregnant while			
562	receiving AVASTIN, she should be apprised of the potential hazard to the			
563	fetus and/or the potential risk of loss of pregnancy. Patients who			
564	discontinue AVASTIN should also be counseled concerning the prolonged			
565	exposure following discontinuation of therapy (half-life of approximately			
566	20 days) and the possible effects of AVASTIN on fetal development.			
567	Nursing Mothers			
568	It is not known whether AVASTIN is secreted in human milk. Because			
569	human IgG1 is secreted into human milk, the potential for absorption and			
570	harm to the infant after ingestion is unknown. Women should be advised			
571	to discontinue nursing during treatment with AVASTIN and for a			
572	prolonged period following the use of AVASTIN, taking into account the			
573	half-life of the product, approximately 20 days [range 11–50 days].			
574	(See CLINICAL PHARMACOLOGY: Pharmacokinetics.)			
575	Pediatric Use			
576	The safety and effectiveness of AVASTIN in pediatric patients has not			
577	been studied. However, physeal dysplasia was observed in juvenile			

578	cynomolgus monkeys with open growth plates treated for four weeks with		
579	doses that were less than the recommended human dose based on mg/kg		
580	and exposure. The incidence and severity of physeal dysplasia were		
581	dose-related and were at least partially reversible upon cessation of		
582	treatment.		
583	Geriatric Use		
584	In Study 1, NCI-CTC Grade 3-4 adverse events were collected in all		
585	patients receiving study drug (396 bolus-IFL plus placebo; 392 bolus-IFL		
586	plus AVASTIN; 109 5-FU/LV plus AVASTIN), while NCI-CTC Grade 1		
587	and 2 adverse events were collected in a subset of 309 patients. There		
588	were insufficient numbers of patients 65 years and older in the subset in		
589	which NCI-CTC Grade 1-4 adverse events were collected to determine		
590	whether the overall adverse event profile was different in the elderly as		
591	compared to younger patients. Among the 392 patients receiving		
592	bolus-IFL plus AVASTIN, 126 were at least 65 years of age. Severe		
593	adverse events that occurred at a higher incidence (≥2%) in the elderly		
594	when compared to those less than 65 years were asthenia, sepsis, deep		
595	thrombophlebitis, hypertension, hypotension, myocardial infarction,		
596	congestive heart failure, diarrhea, constipation, anorexia, leukopenia,		
597	anemia, dehydration, hypokalemia, and hyponatremia. The effect of		
598	AVASTIN on overall survival was similar in elderly patients as compared		
599	to younger patients.		
600	In Study 3, patients age 65 and older receiving AVASTIN plus FOLFOX4		
601	had a greater relative risk as compared to younger patients for the		
602	following adverse events: nausea, emesis, ileus, and fatigue.		
603	In Study 5 patients age 65 and older receiving carboplatin, paclitaxel, and		
604	AVASTIN had a greater relative risk for proteinuria as compared to		
605	younger patients.		
606	Of the 742 patients enrolled in Genentech-sponsored clinical studies in		
607	which all adverse events were captured, 212 (29%) were age 65 or older		

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608	and 43 (6%) were age 75 or older. Adverse events of any severity that		
609	occurred at a higher incidence in the elderly as compared to younger		
610	patients, in addition to those described above, were dyspepsia,		
611	gastrointestinal hemorrhage, edema, epistaxis, increased cough, and voice		
612	alteration.		
613	In an exploratory, pooled analysis of 1745 patients treated in		
614	five randomized, controlled studies, there were 618 (35%) patients age		
615	65 or older and 1127 patients less than 65 years of age. The overall		
616	incidence of arterial thromboembolic events was increased in all patients		
617	receiving AVASTIN with chemotherapy as compared to those receiving		
618	chemotherapy alone, regardless of age. However, the increase in arterial		
619	thromboembolic events incidence was greater in patients 65 and over		
620	(8.5% vs. 2.9%) as compared to those less than 65 (2.1% vs. 1.4%).		
621	(See WARNINGS: Arterial Thromboembolic Events.)		
622	ADVERSE REACTIONS		
623	The most serious adverse reactions in patients receiving AVASTIN were		
624			
	• Gastrointestinal Perforations (see WARNINGS)		
625	<ul> <li>Gastrointestinal Perforations (see WARNINGS)</li> <li>Non-Gastrointestinal Fistula Formation (see WARNINGS)</li> </ul>		
625 626			
	Non-Gastrointestinal Fistula Formation (see WARNINGS)		
626	<ul> <li>Non-Gastrointestinal Fistula Formation (see WARNINGS)</li> <li>Wound Healing Complications (see WARNINGS)</li> </ul>		
626 627	<ul> <li>Non-Gastrointestinal Fistula Formation (see WARNINGS)</li> <li>Wound Healing Complications (see WARNINGS)</li> <li>Hemorrhage (see WARNINGS)</li> </ul>		
626 627 628	<ul> <li>Non-Gastrointestinal Fistula Formation (see WARNINGS)</li> <li>Wound Healing Complications (see WARNINGS)</li> <li>Hemorrhage (see WARNINGS)</li> <li>Arterial Thromboembolic Events (see WARNINGS)</li> </ul>		
626 627 628 629 630	<ul> <li>Non-Gastrointestinal Fistula Formation (see WARNINGS)</li> <li>Wound Healing Complications (see WARNINGS)</li> <li>Hemorrhage (see WARNINGS)</li> <li>Arterial Thromboembolic Events (see WARNINGS)</li> <li>Hypertensive Crises (see WARNINGS: Hypertension)</li> <li>Reversible Posterior Leukoencephalopathy Syndrome</li> </ul>		
626 627 628 629 630 631	<ul> <li>Non-Gastrointestinal Fistula Formation (see WARNINGS)</li> <li>Wound Healing Complications (see WARNINGS)</li> <li>Hemorrhage (see WARNINGS)</li> <li>Arterial Thromboembolic Events (see WARNINGS)</li> <li>Hypertensive Crises (see WARNINGS: Hypertension)</li> <li>Reversible Posterior Leukoencephalopathy Syndrome (see WARNINGS)</li> </ul>		
626 627 628 629 630 631 632	<ul> <li>Non-Gastrointestinal Fistula Formation (see WARNINGS)</li> <li>Wound Healing Complications (see WARNINGS)</li> <li>Hemorrhage (see WARNINGS)</li> <li>Arterial Thromboembolic Events (see WARNINGS)</li> <li>Hypertensive Crises (see WARNINGS: Hypertension)</li> <li>Reversible Posterior Leukoencephalopathy Syndrome (see WARNINGS)</li> <li>Neutropenia and Infection (see WARNINGS)</li> </ul>		

The most common adverse events in patients receiving AVASTIN were

asthenia, pain, abdominal pain, headache, hypertension, diarrhea, nausea,

637	vomiting, anorexia, stomatitis, constipation, upper respiratory infection,
638	epistaxis, dyspnea, exfoliative dermatitis, and proteinuria.
639	Adverse Reactions in Clinical Trials
640	Because clinical trials are conducted under widely varying conditions,
641	adverse reaction rates observed in the clinical trials of a drug cannot be
642	directly compared to rates in the clinical trials of another drug and may not
643	reflect the rates observed in practice. The adverse reaction information
644	from clinical trials does, however, provide a basis for identifying the
645	adverse events that appear to be related to drug use and for approximating
646	rates.
647	The data described below reflect exposure to AVASTIN in 1529 patients,
648	including 665 receiving AVASTIN for at least 6 months and 199 receiving
649	AVASTIN for at least one year. AVASTIN was studied primarily in
650	placebo- and active-controlled trials (n=501, and n=1028, respectively).
651	Gastrointestinal Perforation
652	The incidence of gastrointestinal perforation across all studies ranged from
653	0-3.7%. The incidence of gastrointestinal perforation, in some cases fatal
654	in patients with mCRC receiving AVASTIN alone or in combination with
655	chemotherapy was 2.4% compared to 0.3% in patients receiving only
656	chemotherapy. The incidence of gastrointestinal perforation in NSCLC
657	patients receiving AVASTIN was 0.9% compared to 0% in patients
658	receiving only chemotherapy. (See WARNINGS:
659	Gastrointestinal Perforations and DOSAGE AND
660	ADMINISTRATION: Dose Modifications.)
661	Non-Gastrointestinal Fistula Formation
662	(See WARNINGS: Non-Gastrointestinal Fistula Formation,
663	<b>DOSAGE AND ADMINISTRATION: Dose Modifications.)</b>

664	Wound Healing Complications			
665	The incidence of post-operative wound healing and/or bleeding			
666	complications was increased in patients with mCRC receiving AVASTIN			
667	as compared to patients receiving only chemotherapy. Among patients			
668	requiring surgery on or within 60 days of receiving study treatment,			
669	wound healing and/or bleeding complications occurred in 15% (6/39) of			
670	patients receiving bolus-IFL plus AVASTIN as compared to 4% (1/25) of			
671	patients who received bolus-IFL alone. In the same study, the incidence			
672	of wound dehiscence was also higher in the AVASTIN-treated patients			
673	(1% vs. 0.5%).			
674	Hemorrhage			
675	Severe or fatal hemorrhages, including hemoptysis, gastrointestinal			
676	bleeding, hematemesis, CNS hemorrhage, epistaxis, and vaginal bleeding			
677	occurred up to five-fold more frequently in AVASTIN treated patients			
678	compared to patients treated with chemotherapy alone. NCI-CTC			
679	Grade 3-5 hemorrhagic events occurred in 4.7% of NSCLC patients and			
680	5.2% of mCRC patients receiving AVASTIN compared to 1.1% and 0.7%			
681	for the control groups respectively. (See WARNINGS: Hemorrhage.)			
682	The incidence of epistaxis was higher (35% vs. 10%) in patients with			
683	mCRC receiving bolus-IFL plus AVASTIN compared with patients			
684	receiving bolus-IFL plus placebo. These events were generally mild in			
685	severity (NCI-CTC Grade 1) and resolved without medical intervention.			
686	Additional mild to moderate hemorrhagic events reported more frequently			
687	in patients receiving bolus-IFL plus AVASTIN when compared to those			
688	receiving bolus-IFL plus placebo included gastrointestinal hemorrhage			
689	(24% vs. 6%), minor gum bleeding (2% vs. 0), and vaginal hemorrhage			
690	(4% vs. 2%). (See WARNINGS: Hemorrhage and DOSAGE AND			
691	ADMINISTRATION: Dose Modifications.)			
692	Arterial Thromboembolic Events			
693	The incidence of arterial thromboembolic events was increased in NSCLC			
694	patients receiving PC plus AVASTIN (3.0%) compared with patients			

695	receiving PC alone (1.4%). Five events were fatal in the PC plus
696	AVASTIN arm, compared with 1 event in the PC alone arm. This
697	increased risk is consistent with that observed in patients with mCRC.
698	(See WARNINGS: Arterial Thromboembolic Events, DOSAGE AND
699	ADMINISTRATION: Dose Modifications, and PRECAUTIONS:
700	Geriatric Use.)
701	Venous Thromboembolic Events
702	The incidence of NCI-CTC Grade 3–4 venous thromboembolic events
703	was higher in patients with mCRC or NSCLC receiving AVASTIN with
704	chemotherapy as compared to those receiving chemotherapy alone. In
705	addition, in patients with mCRC the risk of developing a second
706	subsequent thromboembolic event in patients receiving AVASTIN and
707	chemotherapy is increased compared to patients receiving chemotherapy
708	alone. In Study 1, 53 patients (14%) on the bolus-IFL plus AVASTIN arm
709	and 30 patients (8%) on the bolus-IFL plus placebo arm received full dose
710	warfarin following a venous thromboembolic event. Among these
711	patients, an additional thromboembolic event occurred in 21% (11/53) of
712	patients receiving bolus-IFL plus AVASTIN and 3% (1/30) of patients
713	receiving bolus-IFL alone.
714	The evenual incidence of NGI CTC Conde 2. A venue of throughous holic
714	The overall incidence of NCI-CTC Grade 3–4 venous thromboembolic
715	events in Study 1 was 15.1% in patients receiving bolus-IFL plus
716	AVASTIN and 13.6% in patients receiving bolus-IFL plus placebo.
717	In Study 1, the incidence of the following NCI-CTC Grade 3 and 4 venous
718	thromboembolic events was higher in patients receiving bolus-IFL plus
719	AVASTIN as compared to patients receiving bolus-IFL plus placebo:
720	deep venous thrombosis (34 vs. 19 patients) and intra-abdominal venous
721	thrombosis (10 vs. 5 patients).
722	Hypertension
723	Fatal CNS hemorrhage complicating AVASTIN induced hypertension can

724

occur.

725 In Study 1, the incidences of hypertension and of severe hypertension

were increased in patients with mCRC receiving AVASTIN compared to

727 those receiving chemotherapy alone (see Table 3).

Table 3
Incidence of Hypertension and Severe Hypertension in Study 1

	Arm 1 IFL+Placebo (n=394)	Arm 2 IFL+AVASTIN (n=392)	Arm 3 5-FU/LV+AVASTIN (n=109)
Hypertension <sup>a</sup> (>150/100 mmHg)	43%	60%	67%
Severe Hypertension <sup>a</sup> (>200/110 mmHg)	2%	7%	10%

<sup>&</sup>lt;sup>a</sup> This includes patients with either a systolic or diastolic reading greater than the cutoff value on one or more occasions.

728

Among patients with severe hypertension in the AVASTIN arms, slightly

over half the patients (51%) had a diastolic reading greater than

731 110 mmHg associated with a systolic reading less than 200 mmHg.

732 Similar results were seen in patients receiving AVASTIN alone or in

733 combination with FOLFOX4 or carboplatin and paclitaxel.

734 (See WARNINGS: Hypertension and DOSAGE AND

735 ADMINISTRATION: Dose Modifications.)

736 Neutropenia and Infection

An increased incidence of neutropenia has been reported in patients

738 receiving AVASTIN and chemotherapy compared to chemotherapy alone.

739 In Study 1, the incidence of NCI-CTC Grade 3 or 4 neutropenia was

740 increased in patients with mCRC receiving IFL+AVASTIN (21%)

compared to patients receiving IFL alone (14%). In Study 5, the incidence

of NCI-CTC Grade 4 neutropenia was increased in patients with NSCLC

receiving PC plus AVASTIN (26.2%) compared with patients receiving

PC alone (17.2%). Febrile neutropenia was also increased (5.4% for PC

745 plus AVASTIN vs. 1.8% for PC alone). There were 19 (4.5%) infections

with NCI-CTC Grade 3 or 4 neutropenia in the PC plus AVASTIN arm of

747	which 3 were fatal compared to 9 (2%) neutropenic infections in patients
748	receiving PC alone, of which none were fatal. During the first 6 cycles of
749	treatment the incidence of serious infections including pneumonia, febrile
750	neutropenia, catheter infections and wound infections was increased in the
751	PC plus AVASTIN arm [58 patients (13.6%)] compared to the PC alone
752	arm [29 patients (6.6%)].
753	Proteinuria
754	(See WARNINGS: Proteinuria, DOSAGE AND
755	ADMINISTRATION: Dose Modifications, and PRECAUTIONS:
756	Geriatric Use.)
757	Immunogenicity
758	As with all therapeutic proteins, there is a potential for immunogenicity.
759	The incidence of antibody development in patients receiving AVASTIN
760	has not been adequately determined because the assay sensitivity was
761	inadequate to reliably detect lower titers. Enzyme-linked immunosorbent
762	assays (ELISAs) were performed on sera from approximately 500 patients
763	treated with AVASTIN, primarily in combination with chemotherapy.
764	High titer human anti-AVASTIN antibodies were not detected.
765	Immunogenicity data are highly dependent on the sensitivity and
766	specificity of the assay. Additionally, the observed incidence of antibody
767	positivity in an assay may be influenced by several factors, including
768	sample handling, timing of sample collection, concomitant medications,
769	and underlying disease. For these reasons, comparison of the incidence of
770	antibodies to AVASTIN with the incidence of antibodies to other products
771	may be misleading.
772	Metastatic Carcinoma of the Colon and Rectum
773	The data in Tables 4 and 5 were obtained in Study 1. All NCI-CTC
774	Grade 3 and 4 adverse events and selected NCI-CTC Grade 1 and 2
775	adverse events (hypertension, proteinuria, thromboembolic events) were

reported for the overall study population. The median age was 60, 60%

787

788

777 were male, 79% were Caucasian, 78% had a colon primary lesion, 56% 778 had extra-abdominal disease, 29% had prior adjuvant or neoadjuvant 779 chemotherapy, and 57% had ECOG performance status of 0. The median 780 duration of exposure to AVASTIN was 8 months in Arm 2 and 7 months 781 in Arm 3. Severe and life-threatening (NCI-CTC Grade 3 and 4) adverse 782 events, which occurred at a higher incidence ( $\geq 2\%$ ) in patients receiving 783 bolus-IFL plus AVASTIN as compared to bolus-IFL plus placebo, are 784 presented in Table 4.

Table 4

NCI-CTC Grade 3 and 4 Adverse Events in Study 1

(Occurring at Higher Incidence (≥2%) AVASTIN vs. Control)

	IFL+	rm 1 Placebo =396)	IFL+A	rm 2 NASTIN =392)
NCI-CTC Grade 3-4 Events	295	(74%)	340	(87%)
Body as a Whole				
Asthenia	28	(7%)	38	(10%)
Abdominal Pain	20	(5%)	32	(8%)
Pain	21	(5%)	30	(8%)
Cardiovascular				
Hypertension	10	(2%)	46	(12%)
Deep Vein Thrombosis	19	(5%)	34	(9%)
Intra-Abdominal Thrombosis	5	(1%)	13	(3%)
Syncope	4	(1%)	11	(3%)
Digestive				
Diarrhea	99	(25%)	133	(34%)
Constipation	9	(2%)	14	(4%)
Hemic/Lymphatic				
Leukopenia	122	(31%)	145	(37%)
Neutropenia	41	(14%)	58	(21%)

<sup>&</sup>lt;sup>a</sup> Central laboratories were collected on Days 1 and 21 of each cycle. Neutrophil counts are available in 303 patients in Arm 1 and 276 in Arm 2.

NCI-CTC Grade 1–4 adverse events which occurred at a higher incidence (≥5%) in patients receiving bolus-IFL plus AVASTIN as compared to the bolus-IFL plus placebo arm, are presented in Table 5.

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Table 5

NCI-CTC Grade 1-4 Adverse Events in Study 1
(Occurring at Higher Incidence (≥5%) in IFL+AVASTIN vs. IFL)

	IFL	arm 1 -Placebo 1=98)	IFL+A	arm 2 AVASTIN = 102)	5-FU/LV	Arm 3 7+AVASTIN =109)
Body as a Whole						
Pain	54	(55%)	62	(61%)	67	(62%)
Abdominal Pain	54	(55%)	62	(61%)	55	(50%)
Headache	19	(19%)	27	(26%)	30	(26%)
Cardiovascular						
Hypertension	14	(14%)	23	(23%)	37	(34%)
Hypotension	7	(7%)	15	(15%)	8	(7%)
Deep Vein Thrombosis	3	(3%)	9	(9%)	6	(6%)
Digestive						
Vomiting	46	(47%)	53	(52%)	51	(47%)
Anorexia	29	(30%)	44	(43%)	38	(35%)
Constipation	28	(29%)	41	(40%)	32	(29%)
Stomatitis	18	(18%)	33	(32%)	33	(30%)
Dyspepsia	15	(15%)	25	(24%)	19	(17%)
GI Hemorrhage	6	(6%)	25	(24%)	21	(19%)
Weight Loss	. 10	(10%)	15	(15%)	18	(16%)
Dry Mouth	2	(2%)	7	(7%)	4	(4%)
Colitis	1	(1%)	6	(6%)	1	(1%)
Hemic/Lymphatic						
Thrombocytopenia		0	5	(5%)	5	(5%)
<u>Nervous</u>						
Dizziness	20	(20%)	27	(26%)	21	(19%)

Table 5 (cont'd)
NCI-CTC Grade 1–4 Adverse Events in Study 1

(Occurring at Higher Incidence (≥5%) in IFL+AVASTIN vs. IFL)

	IFL+	rm 1 -Placebo 1=98)	IFL+A	arm 2 AVASTIN =102)	5-FU/LV	arm 3 +AVASTIN =109)
Respiratory						
Upper Respiratory Infection	38	(39%)	48	(47%)	44	(40%)
Epistaxis	10	(10%)	36	(35%)	35	(32%)
Dyspnea	15	(15%)	26	(26%)	27	(25%)
Voice Alteration	2	(2%)	. 9	(9%)	6	(6%)
Skin/Appendages						
Alopecia	25	(26%)	33	(32%)	. 6	(6%)
Skin Ulcer	1	(1%)	6	(6%)	7	(6%)
Special Senses						
Taste Disorder	9	(9%)	14	(14%)	23	(21%)
Urogenital	•					
Proteinuria	24	(24%)	. 37	(36%)	39	(36%)

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791 The data in Table 6 were obtained in Study 3. Only NCI-CTC Grade 3–5 non-hematologic and Grade 4-5 hematologic adverse events related to 792 793 treatment were reported. The median age was a 61 years, 40% were 794 female, 87% were Caucasian, 99% received prior chemotherapy for 795 metastatic colorectal cancer, 26% had received prior radiation therapy, and 796 the 49% had an ECOG performance status of 0. Selected NCI-CTC Grade 3-5 non-hematologic and Grade 4-5 hematologic adverse events 797 798 which occurred at a higher incidence in patients receiving FOLFOX4 plus AVASTIN as compared to those who received FOLFOX4 alone, are 799 presented in Table 6. These data are likely to under-estimate the true 800 adverse event rates due to the reporting mechanisms used in Study 3. 801

Table 6

NCI-CTC Grade 3–5 Non-Hematologic and
Grade 4–5 Hematologic Adverse Events in Study 3

(Occurring at Higher Incidence (≥2%)

with AVASTIN+FOLFOX4 vs. FOLFOX4)

		FOLFOX4+	
•	FOLFOX4	AVASTIN	AVASTIN
	(n=285)	(n=287)	(n=234)
Patients with at least one event	171 (60%)	219 (76%)	87 (37%)
Gastrointestinal			
Diarrhea	36 (13%)	51 (18%)	5 (2%)
Nausea	13 (5%)	35 (12%)	14 (6%)
Vomiting	11 (4%)	32 (11%)	15 (6%)
Dehydration	14 (5%)	29 (10%)	15 (6%)
Ileus	4 (1%)	10 (4%)	11 (5%)
Neurology			
Neuropathy-sensory	26 (9%)	48 (17%)	2 (1%)
Neurologic-other	8 (3%)	15 (5%)	3 (1%)
Constitutional symptoms			
Fatigue	37 (13%)	56 (19%)	12 (5%)
Pain			
Abdominal pain	13 (5%)	24 (8%)	19 (8%)
Headache	0 (0%)	8 (3%)	4 (2%)
Cardiovascular (general)			
Hypertension	5 (2%)	26 (9%)	19 (8%)
Hemorrhage			
Hemorrhage	2 (1%)	15 (5%)	9 (4%)

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## Non-Squamous, Non-Small Cell Lung Cancer

The data in Table 7 were obtained in Study 5. Only NCI-CTC Grade 3–5 non-hematologic and Grade 4–5 hematologic adverse events were reported. The median age was 63, 46% were female, no patients had received prior chemotherapy, 76% had Stage IV disease, 12% had Stage IIIB disease with malignant pleural effusion, 11% had recurrent disease, and 40% had an ECOG performance status of 0. The median duration of exposure to AVASTIN was 4.9 months.

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- NCI-CTC Grade 3, 4, and 5 adverse events that occurred at a  $\geq$ 2% higher
- 812 incidence in patients receiving PC plus AVASTIN as compared with PC
- alone are presented in Table 7.

Table 7

NCI-CTC Grade 3-5 Non-Hematologic and
Grade 4 and 5 Hematologic Adverse Events in Study 5

(Occurring at a ≥2% Higher Incidence in

AVASTIN-Treated Patients Compared with Control)

No. (%) of NSCLC Patients		
NCI-CTC Category Term <sup>a</sup>	PC (n=441)	PC + AVASTIN (n=427)
Any event	286 (65%)	334 (78%)
Blood/bone marrow		
Neutropenia	76 (17%)	113 (27%)
Constitutional symptoms		
Fatigue	57 (13%)	67 (16%)
Cardiovascular (general)		
Hypertension	3 (0.7%)	33 (8%)
Vascular		
Venous thrombus/embolism	14 (3%)	23 (5%)
Infection/febrile neutropenia		
Infection without neutropenia	12 (3%)	30 (7%)
Infection with NCI-CTC Grade 3 or 4 neutropenia	9 (2%)	19 (4%)
Febrile neutropenia	8 (2%)	23 (5%)
Pulmonary/upper respiratory		
Pneumonitis/pulmonary infiltrates	11 (3%)	21 (5%)
Metabolic/laboratory		
Hyponatremia	5 (1%)	16 (4%)
Pain		
Headache	2 (0.5%)	13 (3%)
Renal/genitourinary		
Proteinuria	0 (0%)	13 (3%)

<sup>&</sup>lt;sup>a</sup> Events were reported and graded according to NCI-CTC, Version 2.0. Per protocol, investigators were required to report NCI-CTC Grade 3–5 non-hematologic and Grade 4 and 5 hematologic events.

815	Other Serious Adverse Events
816	The following additional serious adverse events occurred in at least one
817	subject treated with AVASTIN in clinical studies or post-marketing
818	experience:
819	Body as a Whole: polyserositis
820 821	Digestive: intestinal necrosis, mesenteric venous occlusion, anastomotic ulceration
822	Hemic and lymphatic: pancytopenia
823	Respiratory: nasal septum perforation
824	OVERDOSAGE
825	The highest dose tested in humans (20 mg/kg IV) was associated with
826	headache in nine of 16 patients and with severe headache in three of
827	16 patients.
828	DOSAGE AND ADMINISTRATION
829	Do not initiate AVASTIN until at least 28 days following major surgery.
830	The surgical incision should be fully healed prior to initiation of
831	AVASTIN.
832	Metastatic Carcinoma of the Colon or Rectum
833	AVASTIN, used in combination with intravenous 5-FU-based
834	chemotherapy, is administered as an intravenous infusion (5 mg/kg or
835	10 mg/kg) every 14 days.
836	The recommended dose of AVASTIN, when used in combination with
837	bolus-IFL, is 5 mg/kg.
838	The recommended dose of AVASTIN, when used in combination with
839	FOLFOX4, is 10 mg/kg.
840	Non-Squamous, Non-Small Cell Lung Cancer
841	The recommended dose of AVASTIN is 15 mg/kg, as an IV infusion
842	every 3 weeks.

843	Dose Wodifications
844	There are no recommended dose reductions for the use of AVASTIN.
845	If needed, AVASTIN should be either discontinued or temporarily
846	suspended as described below.
847	AVASTIN should be permanently discontinued in patients who develop
848	gastrointestinal perforation (gastrointestinal perforation, fistula formation
849	in the gastrointestinal tract, intra-abdominal abscess), fistula formation
850	involving an internal organ, wound dehiscence requiring medical
851	intervention, serious bleeding, a severe arterial thromboembolic event,
852	nephrotic syndrome, hypertensive crisis or hypertensive encephalopathy.
853	In patients developing RPLS, discontinue AVASTIN and initiate
854	treatment of hypertension, if present. (See WARNINGS:
855	Reversible Posterior Leukoencephalopathy Syndrome.)
856	Temporary suspension of AVASTIN is recommended in patients with
857	evidence of moderate to severe proteinuria pending further evaluation and
858	in patients with severe hypertension that is not controlled with medical
859	management. The risk of continuation or temporary suspension of
860	AVASTIN in patients with moderate to severe proteinuria is unknown.
861	AVASTIN should be suspended at least several weeks prior to elective
862	surgery. (See WARNINGS: Gastrointestinal Perforation and
863	Wound Healing Complications and PRECAUTIONS: Surgery.)
864	AVASTIN should not be resumed until the surgical incision is fully
865	healed.
866	Preparation for Administration
867	AVASTIN should be diluted for infusion by a healthcare professional
868	using aseptic technique. Withdraw the necessary amount of AVASTIN to
869	obtain the required dose and dilute in a total volume of 100 mL of 0.9%
870	Sodium Chloride Injection, USP. Discard any unused portion left in a
871	vial, as the product contains no preservatives. Parenteral drug products

872	should be inspected visually for particulate matter and discoloration prior
873	to administration.
874	Diluted AVASTIN solutions for infusion may be stored at 2°C-8°C
875	(36°F-46°F) for up to 8 hours. No incompatibilities between AVASTIN
876	and polyvinylchloride or polyolefin bags have been observed.
877	AVASTIN infusions should not be administered or mixed with
878	dextrose solutions.
879	Administration
880	DO NOT ADMINISTER AS AN IV PUSH OR BOLUS. The initial
881	AVASTIN dose should be delivered over 90 minutes as an IV infusion
882	following chemotherapy. If the first infusion is well tolerated, the second
883	infusion may be administered over 60 minutes. If the 60-minute infusion
884	is well tolerated, all subsequent infusions may be administered over
885	30 minutes.
886	Stability and Storage
887	AVASTIN vials must be refrigerated at 2–8°C (36–46°F). AVASTIN
888	vials should be protected from light. Store in the original carton until tim
889	of use. DO NOT FREEZE. DO NOT SHAKE.
890	HOW SUPPLIED
891	AVASTIN is supplied as 4 mL and 16 mL of a sterile solution in
892	single-use glass vials to deliver 100 and 400 mg of Bevacizumab per vial,
893	respectively.
894	Single unit 100 mg carton: Contains one 4 mL vial of AVASTIN
895	(25 mg/mL). NDC 50242-060-01
896	Single unit 400 mg carton: Contains one 16 mL vial of AVASTIN
897	(25 mg/mL). NDC 50242-061-01