A Multicenter Phase 2 Study of AMG 337 in Patients With MET-

2 Amplified Gastric/Gastroesophageal Junction/Esophageal

3 Adenocarcinoma and Other Solid Tumors

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18	Running title:	AMG 337 Phase 2 Study in MET-Amplified Tumors
19 20 21 22 23 24 25 26 27 28	Keywords:	Keywords from dropdown menu: Gastrointestinal cancers: stomach; Gastrointestinal cancers: other; CB02 CELL GROWTH/SIGNALING PATHWAYS:_Protein tyrosine kinases; CL01 Phase I-III CLINICAL TRIALS:_ Gastrointestinal cancers: stomach; CL01 Phase I-III CLINICAL TRIALS:_ Gastrointestinal cancers: other; ET05 SMALL MOLECULES AND OTHER THERAPEUTIC AGENTS_Kinase and phosphatase inhibitors User-defined keywords: AMG 337, MET inhibitor, advanced solid tumors
29	Financial support:	This study was funded by Amgen Inc.
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36 37 38 39 40 41 42 43 44	Conflicts of interest:	Eric Van Cutsem reports consulting or advisory roles for Bayer, Lilly, Roche, and SERVIER and has received research funding from Amgen, Bayer, Boehringer Ingelheim, Lilly, Novartis, Roche, Sanofi, Celgene, Ipsen, Merck, Merck KGaA, and SERVIER. Veena Shankaran has received research funding during the study (paid to institution) from Amgen Inc., and research funding outside of the submitted work (paid to institution) from Bayer, EMD-Serono, Merck, and Bristol-Myers Squibb. Salvatore Siena has participated in advisory boards

1 2 3 4 5 6 7 8 9 10 11 12 13		for Amgen, Roche, Bayer, Merck, Sanofi, Novartis, and Merrimack. He is supported by research grants from Fondazione Oncologia Niguarda Onlus, Associazione Italiana Ricerca Cancro, and MoTriColor European Union Horizon 2020 research and innovation programme under grant agreement N.635342. Ning F. Go and Hui Yang are employees and stockholders of Amgen Inc. Marco Schupp is an employee of Amgen (Europe) GmbH and is a stockholder of Amgen Inc. David Cunningham has received grants from 4SC, AstraZeneca, Amgen, Bayer, Celgene, Clovis, Eli Lilly, Janssen, Sanofi, Merrimack, Medimmune, and Merck Serono. Boguslawa Karaszewska, Yoon-Koo Kang, and Hyun Cheol Chung declare no conflicts of interest.
14	Word count:	3738 (limit, 5000)
15	Tables/figures:	4/2 (limit, 6 total)
16	Article Type:	Research Article, Cancer Therapy: Clinical
17	Clinical trial registration:	ClinicalTrials.gov; NCT02016534
18		

1 TRANSLATIONAL RELEVANCE

2 A number of mesenchymal-epithelial transition (MET) pathway inhibitors have been 3 assessed in clinical trials, but those trials have mainly focused on patients with high levels of 4 MET protein expression. In this study, we assessed AMG 337, a highly selective small-5 molecule MET inhibitor, in patients with MET gene amplification, a relatively rare event. 6 AMG 337 monotherapy in heavily pretreated patients with advanced stage MET-amplified 7 tumors showed an objective response rate of 18% in the cohort of 45 patients with 8 gastric/gastroesophageal junction/esophageal tumors and measurable disease. No 9 responses were observed in patients with other solid tumors. The study was terminated 10 after a protocol-permitted review showed lower-than-expected activity in a separate first-in-11 human study of AMG 337. Future studies are necessary to determine which biomarker(s) 12 would be predictive of response to MET-targeted therapy, which signaling pathways 13 contribute to resistance, and whether combination therapy would show greater efficacy than 14 was observed in this study.

1 ABSTRACT

2 **Purpose:** *MET* gene amplification is associated with poor prognosis in

3 gastric/gastroesophageal junction/esophageal (G/GEJ/E) cancers. We determined

- 4 antitumor activity, safety, and pharmacokinetics of the small-molecule MET inhibitor
- 5 AMG 337 in *MET*-amplified G/GEJ/E adenocarcinoma or other solid tumors.
- 6 **Experimental Design:** In this phase 2, single-arm study, adults with *MET*-amplified
- 7 G/GEJ/E adenocarcinoma (Cohort 1) or other *MET*-amplified solid tumors (Cohort 2)
- 8 received AMG 337 300 mg/d orally in 28-day cycles. The primary endpoint was objective

9 response rate (ORR; Cohort 1). Secondary endpoints included ORR (Cohort 2),

- 10 progression-free survival (PFS), overall survival (OS), and safety.
- 11 **Results:** Of 2101 patients screened for *MET* amplification, 132 were *MET*-amplified and 60
- 12 were enrolled: 45 in Cohort 1, and 15 in Cohort 2. Fifty-six patients (97%) had metastatic
- 13 disease; 57 had prior lines of therapy (1 prior line, 29%; ≥ 2 prior lines, 69%). A protocol-
- 14 permitted review showed efficacy that was lower-than-expected based on preliminary data
- 15 from a first-in-human study, and enrollment was stopped. Fifty-eight patients received ≥1
- 16 AMG 337 dose. ORR in Cohort 1 was 18% (8 partial responses). No responses were
- 17 observed in Cohort 2. Of 54 evaluable patients, median (95% CI) PFS and OS were 3.4
- 18 (2.2–5.0) and 7.9 (4.8–10.9) months, respectively. The most frequent adverse events (AEs)
- 19 were headache (60%), nausea (38%), vomiting (38%), and abdominal pain, decreased
- 20 appetite, and peripheral edema (33% each); 71% had grade ≥3 AEs and 59% had serious
- 21 AEs.
- 22 Conclusions: AMG 337 showed antitumor activity in MET-amplified G/GEJ/E
- adenocarcinoma but not in *MET*-amplified non–small-cell lung cancer.
- 24

1 INTRODUCTION

Gastric and esophageal cancers are the fifth and eighth most common types of cancer
worldwide, respectively (1). They are typically diagnosed at the locally advanced or
advanced stage, when surgery is not an option (2). Systemic chemotherapy remains the
primary mode of treatment for advanced disease; however, median overall survival (OS) for
first-line treatment is approximately 9 to 11 months (3,4).

7 The mesenchymal-epithelial transition (MET) receptor tyrosine kinase regulates cell survival, 8 proliferation, and migration (5-9). MET overexpression and gene amplification have been 9 observed in multiple solid tumors (10-14). MET overexpression has been reported in 46% to 10 74% of patients with gastric and esophageal cancers (15-18); MET amplification has been 11 reported in 2% to 10% of this patient population (16-18). MET overexpression and 12 amplification have been associated with poor prognosis, and MET overexpression has been 13 correlated with depth of tumor invasion and lymph node metastasis, advanced stage, and 14 shortened survival (18,19); thus, MET inhibition represents a rational therapeutic strategy. 15 Furthermore, MET pathway inhibitors (eq, monoclonal antibodies and tyrosine kinase 16 inhibitors) have shown activity in MET-overexpressing and MET-amplified gastric cancer 17 (16, 20).

18 AMG 337 is a highly selective and potent small-molecule inhibitor of MET receptor signaling 19 (21). In preclinical studies, AMG 337 inhibited phosphorylation of MET and downstream 20 effectors in multiple MET-amplified cell lines, inhibited MET-dependent cell growth and induced apoptosis in those cell lines, and reduced tumor growth in MET-dependent 21 22 xenograft models (21). In the phase 1 AMG 337 first-in-human study in solid tumors, the 23 maximum tolerated and recommended phase 2 dose was determined to be 300 mg orally 24 once daily (QD), and the most common treatment-related adverse events (AEs) were 25 headache, fatigue, nausea, and vomiting (22). In that study, AMG 337 showed an objective 26 response rate (ORR) of 9.9% (11/111) in all patients, regardless of MET-amplification status,

- 1 with a higher ORR (29.6%; 8/27) among *MET*-amplified patients. Based on the heightened
- 2 antitumor activity in *MET*-amplified patients and acceptable toxicity profile observed in the
- 3 first-in-human study, a decision was made to evaluate AMG 337 in additional trials, including
- 4 the phase 2 study in patients with *MET*-amplified solid tumors reported here.
- 5 The objective of this phase 2, multicenter, single-arm, two-cohort study was to determine the
- 6 antitumor activity, safety, and pharmacokinetics of AMG 337 in MET-amplified
- 7 gastric/gastroesophageal junction/esophageal (G/GEJ/E) adenocarcinoma or other MET-
- 8 amplified solid tumors (ClinicalTrials.gov Identifier; NCT02016534).
- 9

1 MATERIALS AND METHODS

2 Patients

3 Adults with pathologically confirmed advanced G/GEJ/E adenocarcinoma or other solid 4 tumors who had received prior therapy, for whom no standard therapy was available, or who 5 had refused standard therapy, were included. Patients had tumor MET amplification as 6 determined by central testing. MET gene amplification status was determined at a central 7 laboratory; MET amplification was defined as a MET/CEN-7 ratio ≥ 2.0 . Patients also had 8 measurable disease per Response Evaluation Criteria in Solid Tumors (RECIST) version 9 1.1, Eastern Cooperative Oncology Group (ECOG) performance status 0 or 1, and adequate 10 organ function. Patients were excluded if they had known central nervous system 11 metastases, arterial thrombosis, vascular ischemic events, venous thromboembolic events, 12 peripheral edema grade >1, acute hepatitis B or detectable hepatitis C virus, or history of 13 other malignancy within the previous 3 years. Patients with human epidermal growth factor 14 receptor 2 (HER2)-positive tumors were not excluded. All patients provided written informed 15 consent. This study was conducted in accordance with the principles of the applicable 16 country, US Food and Drug Administration, and International Conference on Harmonization 17 (ICH) Good Clinical Practice (GCP) regulations/guidelines. Compliance with ICH GCP 18 guidelines provides public assurance that the rights, safety and well-being of trial subjects 19 are protected, consistent with the principles that have their origin in the Declaration of 20 Helsinki. The protocol was approved by an institutional review board or independent ethics 21 committee at each study site.

22 Study Design

This was a phase 2, multicenter, single-arm cohort study. During screening, formalin-fixed,
paraffin-embedded tumor samples were submitted for *MET*-amplification testing by a central
laboratory. Tumor tissue submitted for testing was recent (preferred) or archival. Eligible
patients with *MET*-amplified tumors were subsequently enrolled into two cohorts: Cohort 1

1 included patients with MET-amplified G/GEJ/E adenocarcinoma with measurable disease 2 per RECIST version 1.1 (planned enrollment, n=100). Cohort 2 included patients with other MET-amplified mixed solid tumors with measurable disease per RECIST version 1.1 3 4 (planned enrollment, n=40); this cohort could include ≤ 10 patients with G/GEJ/E 5 adenocarcinoma with nonmeasurable disease per RECIST version 1.1 (Cohort 2A), ≤10 6 patients with G/GEJ/E adenocarcinoma with measurable disease who had received prior 7 MET antibody therapy (Cohort 2B), and patients with other types of MET-amplified solid 8 tumors (Cohort 2C).

9 Each treatment cycle consisted of a 28-day (±3 days) period. All patients self-administered 10 AMG 337 300 mg orally QD on an empty stomach; at first, no food or drink (except water) 11 was permitted 2 hours before/after administration. The protocol was later amended to allow 12 caffeine (eg, coffee) intake because caffeine use before dosing or during headache onset in 13 the AMG 337 first-in-human study reduced the incidence of grade ≥3 headaches. Treatment 14 continued for 12 months or until disease progression (per RECIST version 1.1), intolerance, 15 consent withdrawal, initiation of a new systemic anticancer therapy, or study termination.

Treatment was withheld for patients who experienced grade ≥3 toxicity for which AMG 337 could not be excluded as the cause or grade ≥3 peripheral edema or headache until toxicity resolved. If resolution occurred within 4 weeks, patients resumed treatment at 200 mg QD. If toxicity recurred at the 200-mg QD dose, treatment was again withheld and patients could resume treatment at 150 mg QD. If resolution did not occur within 4 weeks or if toxicity occurred after the second dose reduction, treatment was discontinued.

22 Endpoints

23 The primary endpoint was ORR (proportion of patients with a complete response [CR] or

24 partial response [PR]) per RECIST version 1.1 in Cohort 1. Secondary endpoints included

- 25 ORR in Cohort 2, duration of response (DOR; time from first response to disease
- 26 progression or death) and time to response (TTR; time from first dose to first response) in

Cohort 1 and patients from Cohort 2 with measurable disease at baseline, progression-free
 survival (PFS; time from first dose to disease progression or death), OS (time from first dose
 to death), incidence and severity of AEs and significant laboratory abnormalities, AMG 337
 exposure and dose intensity, and pharmacokinetics.

5 Assessments

Radiologic tumor assessments (computed tomography or magnetic resonance imaging) per
RECIST version 1.1 were conducted at screening, during week 8 (±3 days), and every 8
weeks thereafter until week 32. After week 32, assessments were conducted every 12
weeks until study end.

10 Adverse events and serious AEs were monitored throughout the study. Patients underwent 11 a safety follow-up visit 30 (+3) days after the final administration unless the decision to 12 discontinue treatment was made >30 days after the last AMG 337 dose or the patient was 13 hospitalized at the time of the follow-up visit. In these instances, follow-up was conducted at 14 the first available opportunity. Patients were contacted every 3 months (±14 days) after the 15 safety follow-up visit or last response follow-up, whichever was later, until the final analysis 16 or the last active patient had died, whichever occurred first. AEs were graded according to 17 the Common Terminology Criteria for Adverse Events version 4.

18 Pharmacokinetics

Approximately 20 patients at selected sites participated in intensive pharmacokinetic
assessments. For these assessments, samples were collected predose and 0.5, 1.5, 3, and
6 hours postdose on cycle 1, day 1; predose on cycle 1, day 2; predose on cycle 1, day 15;
predose and 0.5, 1.5, 3, and 6 hours postdose on cycle 1, day 28; predose on cycle 2, day
1; predose on day 1 of cycles 3, 5, 7, and 9; every 12 weeks thereafter; and at safety followup.

- 1 All patients participated in general pharmacokinetic assessments. Samples for
- 2 pharmacokinetics were collected predose on days 1 and 15 of cycle 1; on day 1 of cycles 2,
- 3 3, 5, 7, and 9; every 12 weeks thereafter; and at safety follow-up. Samples were also taken
- 4 3 hours postdose on day 1 of cycles 1, 3, and 5.
- 5 Pharmacokinetic parameters were estimated by noncompartmental analysis of AMG 337
- 6 using Phoenix WinNonlin v.6.4 software (Centara; Princeton, NJ) on individual plasma
- 7 concentrations. The following parameters were estimated: maximum concentration (C_{max}),
- 8 time to C_{max} (t_{max}), area under the plasma concentration-time curve from 0 to 24 hours
- 9 (AUC₀₋₂₄), and accumulation ratio (AR), calculated as AUC on day 28 divided by AUC on
- 10 day 1.

11 Biomarker Analysis

- 12 *MET* gene amplification status to determine study eligibility was assessed in a single central
- 13 laboratory by IQFISH (Dako North America, an Agilent Technology Company, Carpinteria,
- 14 CA). *MET* amplification was defined as a *MET/CEN-7* ratio ≥2.0. In exploratory analyses,
- 15 *MET* gene copy number was evaluated. Biomarker assessments were conducted on
- 16 archival tumor tissue.

17 Statistical Analysis

No formal hypothesis testing was planned. The study focus was the estimation of the 18 19 magnitude of treatment effect as assessed by ORR in Cohort 1. The point estimate of ORR 20 and the corresponding exact binomial two-sided 95% CI were generated. The planned 21 sample size was approximately 100 for Cohort 1 and approximately 40 for Cohort 2. With 22 the planned sample size, the ORR could be estimated with a standard error not greater than 23 5%; the half-width of the 95% CI for the estimated ORR would be no more than 10%. 24 Assuming an observed ORR of 50%, the lower bound of the 95% CI for the estimated ORR 25 would exclude values <40%.

1 The full and safety analysis sets included all patients who received ≥1 AMG 337 dose. 2 Response analyses included all patients with measurable disease who received ≥1 3 AMG 337 dose. Pharmacokinetic analyses included all patients from the safety analysis set 4 with evaluable blood samples. All analyses were descriptive and focused on the estimation 5 of the magnitude of treatment effect. Descriptive statistics were provided for safety and efficacy endpoints. Safety summaries were provided for all G/GEJ/E patients and overall. 6 7 The number and percentage of patients with a best overall response of CR, PR, stable 8 disease, progressive disease, noncomplete response/nonprogressive disease were 9 determined. The stable disease classification required patients to have a response of stable 10 disease ≥6 weeks after the date of the first dose of AMG 337. ORR was calculated along 11 with the corresponding exact 95% CI using the Clopper-Pearson method (23). For time-to-12 event variables, the Kaplan-Meier estimates and corresponding two-sided 95% CI for the 13 median were determined, and survival plots were prepared.

1 **RESULTS**

2 Patients

Between February 14, 2014, and May 16, 2016 (data cutoff), 2101 patients from 34 study 3 4 centers were screened; 132 (6%) patients had MET-amplification, and 60 patients were 5 enrolled (Fig. 1). Forty-five patients with measurable G/GEJ/E adenocarcinoma were 6 enrolled in Cohort 1; 10 patients with nonmeasurable G/GEJ/E adenocarcinoma were 7 enrolled in Cohort 2A; one patient with measurable G/GEJ/E adenocarcinoma who had 8 received prior MET antibody therapy was enrolled in Cohort 2B; and four patients with non-9 small-cell lung cancer (NSCLC) were enrolled in Cohort 2C. Five patients were HER2-10 positive/amplified (Cohort 1, n=4; Cohort 2A, n=1).

Most patients were male (72%) and white (64%); median (range) age was 62 (25–85) years (**Table 1**). Fifty-six patients (97%) had metastatic disease, and 57 (98%) had received at least one prior line of therapy (1 prior line, 29%; 2 prior lines, 29%; and >2 prior lines 40%). Seventy-two percent of patients did not respond to the first line of chemotherapy, 67% of patients did not respond to any line of chemotherapy, 66% had prior curative surgery for their cancer, and 78% had prior radiotherapy for the current malignancy. Thirty-nine patients (67%) had an ECOG performance status of 1.

18 Of the 60 patients enrolled, 58 (97%) received ≥1 AMG 337 dose and were included in the 19 efficacy and safety analyses; two patients (3%; 1 each from Cohorts 2A and 2C) did not 20 receive AMG 337. Forty-five (78%) had ≥1 dose reduction or dose withheld, most because 21 of toxicity (59%). At data cutoff, 57 (95%) had discontinued treatment (disease progression, 22 57%; AEs, 17%; patient request, 8%; other, 8%; death, 3%; noncompliance, 2%); one from 23 Cohort 2A remained on study. Median (95% CI) time to treatment discontinuation was 2.6 24 (1.9–3.6) months. Reasons for study discontinuation included death (68%), administrative 25 decision (17%), consent withdrawal (8%), and loss to follow-up (3%).

Enrollment in this and all AMG 337 studies was stopped and regulatory agencies were
notified when a protocol-permitted review of this study found an ORR that was lower than
expected based on preliminary data from the AMG 337 first-in-human study (22). As of July
2014, the first-in-human study had shown responses in 8 of 13 (62%) patients with *MET*amplified G/GEJ/E adenocarcinoma (1 CR, duration of 141 weeks; 7 PRs, duration up to 85
weeks), suggesting that the response rate in this study, which only enrolled patients with *MET*-amplified tumors, would be high.

8 Efficacy

9 The maximum change in the sum of longest diameter (SLD) of target lesions for patients in 10 Cohort 1 is shown in Fig. 2A. Twelve patients had maximum percentage reductions >30%, 11 and seven patients had increases in SLD of their target lesion. Eight patients in Cohort 1 12 achieved a best response of PR, for an ORR (95% CI) of 18% (8%-32%) in that cohort. Median (range) TTR was 7.6 (7.0-16.1) weeks, and median (95% CI) DOR was 6.0 13 14 (3.7-16.7) months in Cohort 1. Of those who achieved a PR, seven (88%) had disease 15 progression, and one (13%) was censored. Sixteen patients in Cohort 1 experienced a best 16 response of stable disease (defined as neither sufficient target lesion shrinkage to be 17 classified as PR nor sufficient increase to be classified as progression; Table 2); no 18 responses were observed in the patients with G/GEJ/E adenocarcinoma in Cohorts 2A or 2B 19 or in the patients with NSCLC in Cohort 2C.

Fifty-four patients (Cohort 1, n=45; Cohort 2A, n=9) were included in the PFS and OS
analyses. Forty-five patients (83%) had a PFS event; median (95% CI) PFS was 3.4
(2.2-5.0) months (Fig. 2B). Thirty-six patients (66.7%) died; median (95% CI) OS was 7.9
(4.8-10.9) months (Fig. 2C).

24 Exposure

- Across all cohorts, median (range) number of treatment cycles completed was 3.0 (1-21),
- and duration of treatment was 2.2 (0-20) months. Forty-five patients (78%) had ≥1 dose

change or reduction, largely because of AEs (n=34; 59%). Median (range) actual dose
intensity was 297.8 (59-345) mg/d; relative dose intensity was 99% (20%-115%).

3 Adverse Events

4 Fifty-seven patients (98%) had ≥ 1 treatment-emergent AE (**Table 3**). The most frequently 5 reported AEs (\geq 30% of all patients) were headache (60%), nausea (38%), vomiting (38%), 6 abdominal pain (33%), decreased appetite (33%), and peripheral edema (33%). Forty-one 7 patients (71%) had grade \geq 3 AEs, and 34 (59%) had serious AEs. Ten patients (17%) had 8 AEs leading to AMG 337 discontinuation; these AEs were headache (n=2 patients) and 9 upper abdominal pain, increased blood bilirubin, cholangitis, fatigue, general physical health 10 deterioration, increased hepatic enzyme, edema, peripheral edema, and vomiting (n=1 11 patient each). Nine patients (16%) had fatal AEs; none were deemed treatment-related by 12 investigators. Overall, AEs of interest were reported for 90% of patients; the most frequent 13 was headache. Headache pain (worst level at onset) was evaluated on a scale from 1 (very 14 mild pain) to 10 (extreme pain) for 35 patients who had any postbaseline headache pain; 15 nine (26%) had scores \geq 6; the remaining (45%) had scores ranging from 1 to 5. AMG 337 is 16 a potent inhibitor of the adenosine transporter, which was considered the underlying cause of headache. Other AEs of interest were edema (57%), skin and subcutaneous disorders 17 (35%), and drug-related hepatic disorders (35%). 18

19 Pharmacokinetics

The pharmacokinetic analysis set comprised 467 plasma samples from 58 patients; 16 with G/GEJ/E tumors underwent intensive pharmacokinetic sampling and had sufficient data for analysis (Cohort 1, n=12; Cohort 2, n=4). Pharmacokinetics were similar between cohorts, with no large variation from days 1 to 28 (**Table 4**). Mean C_{max} ranged from 3080 to 4110 ng/mL; mean t_{max} was approximately 3 hours; mean AUC₀₋₂₄ ranged from 32,800 to 48,200 h·ng/mL, and accumulation was minimal: mean AR was 0.946 and 0.965 for Cohorts 1 and 2, respectively.

1 Biomarkers

- 2 A tumor *MET*/CEN-7 ratio ≥2.0 was a study eligibility criterion. Among the 58 patients
- 3 included in the analysis, the mean (range) *MET/CEN-7* ratio was 7.0 (2.0–20.4), and mean
- 4 (range) gene copy number was 16.4 (3.5–51.3). Among the 47 patients who were evaluable
- 5 for treatment response, the mean (range) *MET/CEN-7* ratio was 7.7 (2.4–12.0) among the 8
- 6 responders (17.0%) and 7.1 (2.0–20.4) among the 39 nonresponders (83.0%).

1 DISCUSSION

2 To our knowledge, this is the largest *MET*-amplification screen in G/GEJ/E cancer to date; 3 2101 patients with G/GEJ/E cancer were screened using an analytically validated IQFISH 4 assay. Previous MET pathway inhibitors assessed in clinical trials have focused on patients 5 with high levels of MET protein expression (24,25). In this study, we enrolled patients with 6 tumors that exhibited MET gene amplification, a relatively rare event, as determined by 7 *MET/CEN-7* ratio ≥2.0. *MET* amplification indicates pathway "addiction" and suggests that 8 MET inhibition could be beneficial in *MET*-amplified patients (16.20), a result supported by 9 animal models (24). MET inhibition resistance can be accomplished through activation of 10 other pathways (24). For example, activation of HER2 or epidermal growth factor receptor 11 pathways in MET-amplified GEJ tumor cell lines can overcome MET inhibition (24). This 12 resistance may partially explain why an antitumor response to AMG 337 was not observed in 13 more patients.

14 Of the 2101 patients screened for eligibility for this study, including patients with G/GEJ/E 15 adenocarcinoma and NSCLC, only 132 (6%) had MET amplification, which is consistent with previously reported rates of 2% to 10% (16,18), yet this is a small percentage of the total 16 17 G/GEJ/E population. In this study, which enrolled 60 of those eligible patients and evaluated 18 AMG 337 as monotherapy, PRs as the best response were observed in eight patients with 19 G/GEJ/E tumors; no responses were observed in patients with NSCLC or in patients with 20 nonmeasurable gastric cancer who had previously received MET inhibitors. Biomarker 21 analysis did not uncover an association between the level of MET gene amplification and 22 response to AMG 337 treatment; however, the total number of responders in this analysis 23 was small.

Pharmacokinetics and rates/types of AEs were similar to those from previous AMG 337
studies (22); the most common treatment-emergent AEs were headache, vomiting, and
nausea. Headache is a common adverse reaction to adenosine receptor agonists/transport

1 inhibitors and may be reversed by adenosine antagonists such as caffeine (25,26).

AMG 337 pharmacokinetics was characterized by rapid absorption and no accumulation
over 28 days of dosing.

4 Results from preclinical studies and the phase 1 AMG 337 first-in-human study 5 (ClinicalTrials.gov; NCT01253707) indicated that tumors with MET amplification had 6 sensitivity to AMG 337 (21,22,27). However, an interim analysis of this study initiated when 7 30 patients had completed two 28-day cycles found response rates that were lower than 8 expected based on preliminary data from the AMG 337 phase 1 study. Responses had 9 been observed in 62% of patients with *MET*-amplified tumors in the phase 1 study; 10 responses were observed in only 13% of evaluable patients (3 of 24 patients with at least 1 11 postbaseline scan) in the analysis of this study that was available as of January 22, 2015. 12 Consequently, this study was terminated early, and enrollment in all AMG 337 trials was 13 discontinued. Reasons for the differences in response rates between the phase 1 study and 14 the phase 2 study are unclear. The number of patients in the phase 1 study was small (111 15 enrolled, 27 MET-amplified), the response rates in the final analysis of the phase 1 study 16 were lower (30%, 8 of 27 *MET*-amplified patients), and patients in the phase 1 study may 17 have been enriched for factors other than MET that are not currently understood. The phase 18 1 study enrolled patients with a broader range of tumor types; the phase 2 study included 19 patients who had received prior therapy for advanced disease (not just patients refractory to 20 standard treatment or for whom no standard therapy was available), and the proportion of 21 patients with metastatic disease was higher in the phase 2 study (97% vs 89%). Future 22 studies are necessary to determine which biomarker(s) would be predictive of response to 23 MET-targeted therapy, which signaling pathways contribute to resistance, and whether 24 combination therapy with a MET inhibitor and another targeted agent would show greater 25 efficacy than was observed here.

The MET inhibitors onartuzumab (a monovalent monoclonal antibody that binds the extracellular domain of MET, blocking interaction with the MET ligand HGF) and

1 rilotumumab (a monoclonal antibody that selectively targets HGF) have been examined in 2 MET-IHC-positive G/GEJ cancer (28,29). The phase 3 METGastric and RILOMET-1 studies demonstrated no PFS or OS benefit with either MET inhibitor in combination with 3 4 chemotherapy (28,29). The phase 2 YO28252 study of onartuzumab plus FOLFOX in 5 patients with metastatic GEJ or gastric adenocarcinoma reported a median PFS of 5.95 months for onartuzumab plus FOLFOX versus 6.80 months for placebo plus FOLFOX in all 6 7 patients, a median OS of 8.51 versus 8.48 months in the MET-positive subset, and an ORR 8 of 60.5% in the intent-to-treat population (30). In the present single-arm, phase 2 study of 9 AMG 337 as monotherapy in patients with MET-amplified solid tumors, median PFS and OS 10 were 3.4 and 7.9 months, respectively, and the ORR was 16% overall (18% in patients with 11 measurable MET-amplified G/GEJ/E adenocarcinoma [Cohort 1]).

12 The present study had several limitations. This was a single-arm study; thus, within-study 13 comparison of the response rate with standard of care was not possible. Additionally, early 14 termination likely influenced the final results. Although patients were enrolled based on MET 15 amplification status, testing of *MET* amplification was conducted using archival tumor tissue, 16 and the test result may not have been reflective of tumor status during the study. It is 17 possible that some tumors may have changed between the time archival tumor samples 18 were collected and the time patients were enrolled and treated or that other genomic 19 alterations in some tumors may have affected response to inhibition of the MET signaling 20 pathway. In the future, this may be addressable using novel diagnostic tools (eg, liquid 21 biopsy) to evaluate dynamic changes occurring during therapy (30).

In conclusion, this study demonstrated an ORR of 18% with AMG 337 monotherapy in
heavily pretreated patients with advanced *MET*-amplified G/GEJ/E adenocarcinoma and a
median duration of response of 6.0 months (Cohort 1). Although it is unlikely that
monotherapy would be beneficial in a large group of patients, it is possible that a select
group of patients could benefit from AMG 337 or that combination therapy strategies could
be useful; however, such approaches would require further study.

1 ACKNOWLEDGMENTS

- 2 The authors wish to acknowledge Meghan Johnson, PhD, Miranda Tradewell, PhD, and
- 3 James Balwit, MS, CMPP (Complete Healthcare Communications, LLC, an ICON plc
- 4 company, West Chester, PA), whose work was funded by Amgen Inc., and Micah Robinson,
- 5 PhD (Amgen Inc.) for assistance with writing this manuscript as well as Robert D. Loberg,
- 6 PhD (Amgen Inc.) for his work in developing and deploying the MET FISH assay. David
- 7 Cunningham is funded by the National Institute of Health Research Biomedical Research
- 8 Centre at the Royal Marsden and Institute of Cancer Research.

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- 21

1 DATA SHARING STATEMENT

2 There is a plan to share data. This may include de-identified individual patient data for 3 variables necessary to address the specific research question in an approved data-sharing 4 request; also related data dictionaries, study protocol, statistical analysis plan, informed 5 consent form, and/or clinical study report. Data sharing requests relating to data in this 6 manuscript will be considered after the publication date and 1) this product and indication (or 7 other new use) have been granted marketing authorization in both the US and Europe, or 2) 8 clinical development discontinues and the data will not be submitted to regulatory authorities. 9 There is no end date for eligibility to submit a data sharing request for these data. Qualified 10 researchers may submit a request containing the research objectives, the Amgen product(s) 11 and Amgen study/studies in scope, endpoints/outcomes of interest, statistical analysis plan, 12 data requirements, publication plan, and qualifications of the researcher(s). In general, 13 Amgen does not grant external requests for individual patient data for the purpose of re-14 evaluating safety and efficacy issues already addressed in the product labeling. A 15 committee of internal advisors reviews requests. If not approved, requests may be further 16 arbitrated by a Data Sharing Independent Review Panel. Requests that pose a potential 17 conflict of interest or an actual or potential competitive risk may be declined at Amgen's sole 18 discretion and without further arbitration. Upon approval, information necessary to address 19 the research question will be provided under the terms of a data sharing agreement. This 20 may include anonymized individual patient data and/or available supporting documents, 21 containing fragments of analysis code where provided in analysis specifications. Further 22 details are available at the following: http://www.amgen.com/datasharing.

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TABLES

Table 1. Patient Demographics and Disease Characteristics^a

	Cohort 1	Cohort 2A	Cohort 2B	Cohort 2C	All Patients
Characteristic	(n=45)	(n=9)	(n=1)	(n=3)	(N=58)
Sex					
Female	11 (24)	3 (33)	1 (100)	1 (33)	16 (28)
Male	34 (76)	6 (67)	Û	2 (67)	42 (72)
Median (range) age, years	62 (34–85)	58 (25–81)	59 (59–59)	64 (58–67)	62 (25-85)
Race	, , , , , , , , , , , , , , , , , , ,			, , , , , , , , , , , , , , , , , , ,	
White	27 (60)	6 (67)	1 (100)	3 (100)	37 (64)
Asian	17 (38)	3 (33)	O Í	Û	20 (35)
Other	1 (2)	Û	0	0	1 (2)
Ethnicity					
Hispanic/Latino	0	1 (11)	0	0	1 (2)
Not Hispanic/Latino	45 (100)	8 (89)	1 (100)	3 (100)	57 (98)
Region				()	
Asian	17 (38)	3 (33)	0	0	20 (35)
Europe/Australia	26 (58)	4 (44)	1 (100)	3 (100)	34 (59)
North America	2 (4)	2 (22)	Û	Û	4 (7)
ECOG performance status					
0	15 (33)	3 (33)	1 (100)	0	19 (33)
1	30 (67)	6 (67)	O Í	3 (100)	39 (67)
Disease stage at screening	(),				
Locally advanced	2 (4)	0	0	0	2 (3)
Metastatic disease	43 (96)	9 (100)	1 (100)	3 (100)	56 (97)
Primary tumor location	(),		(
Stomach	33 (73)	7 (78)	0	0	40 (69)
GEJ	6 (13)	1 (11)	1 (100)	0	8 (14)
Esophageal	5 (11)	1 (11)	0	0	6 (10)
Other	1 (2)	Û	0	3 (100)	4 (7)
Prior lines of therapy	· · /			· · ·	
0	0	1 (11)	0	0	1 (2)

1	12 (27)	4 (44)	1 (100)	0	17 (29)
2	14 (31)	2 (22)	0	1 (33)	17 (29)
>2	19 (42)	2 (22)	0	2 (67)	23 (40)
Median (range) MET/CEN-7 ratio	6.2 (2.0–20.4)	4.7 (2.1–14.7)	2.5 (2.5–2.5)	4.7 (2.7-8.6)	5.4 (2.0–20.4)

^aFull analysis set. All data are n (%) unless otherwise stated.

Table 2. Efficacy Analyses^a

	Cohort 1	Cohort 2A	Cohort 2B	Cohort 2C	All Patients
Efficacy, n (%)	(n=45) [°]	(n=10) ⁵	(n=1) ^o	(n=4) ⁵	(N=60) ^o
Response analysis set inclusion	45 (100)	0	1 (100)	3 (75)	49 (82)
Response analysis set exclusion ^a	0	10 (100)	0	1 (25)	11 (18)
No measurable tumor per RECIST	0	10 (100)	0	0	10 (17)
at baseline					
Did not receive AMG 337	0	1 (10)	0	1 (25)	2 (3)
Best response ^a					
CR	0	0	0	0	0
Partial response	8 (18)	0	0	0	8 (16)
Stable disease	16 (36)	N/A ^c	1 (100)	1 (33)	18 (37)
Non-CR/Non-PD	0	N/A ^c	0	0	0
PD	12 (27)	N/A ^c	0	1 (33)	13 (27)
Not assessed	9 (20)	N/A ^c	0	1 (33)	10 (20)
Objective response rate, % ^d	18	N/A	N/A	N/A	16
95% exact CI, %	8–32	N/A	N/A	N/A	7–30

PD=progressive disease; N/A=not applicable.

^aResponse analysis set; defined as all enrolled patients with measurable tumor per RECIST at baseline who received ≥1 dose of AMG 337. ^bAll enrolled patients.

^cNo enrolled patients from Cohort 2A met the criteria for inclusion in the response analysis set; however, among patients from Cohort 2A excluded from response analysis set, 1 patient experienced stable disease, 5 patients experienced non-CR/non-PD, and 2 patients experienced PD; the response assessment was not conducted in 1 patient.

^dResponses required confirmation.

Table 3.	Treatment-Emerg	gent Adverse Ev	vents ^a

	Patients	
AE, n (%)	(N=58)	
All AFs	57 (98)	
$Grade > 3^{b} AF$	41 (71)	
Serious AF	34 (59)	
Serious treatment-related AF	12 (21)	
Fatal AF	9 (16)	
AEs of interest	52 (90)	
ΔFs reported in >10% of patients	()	
Headache	35 (60)	
Nausea	22 (38)	
Vomiting	22 (38)	
Abdominal pain	19 (33)	
Decreased appetite	19 (33)	
Peripheral edema	19 (33)	
Fatigue	13 (22)	
Asthenia	12 (21)	
Diarrhea	12 (21)́	
Hypoalbuminemia	11 (19)	
Back pain	10 (17)	
Constipation	10 (17)	
Dry skin	9 (16)	
Dyspepsia	9 (16)	
Edema	8 (14)	
Pruritus	8 (14)	
Pyrexia	8 (14)	
Upper abdominal pain	7 (12)	
ALT increased	7 (12)	
Dizziness	7 (12)	
Dyspnea	7 (12)	
Rash	7 (12)	
Ascites	6 (10)	
Hypotension	6 (10)	

ALT=alanine aminotransferase.

^aSafety analysis set.
 ^bPer Common Terminology Criteria for Adverse Events version 4.

	Cohort 1							Cohort 2						
		Day 1			Day 28				Day 1			Day 28		
Statistics	t _{max} , h (n=12)	C _{max} , ng/mL (n=12)	AUC₀₋₂₄, h⋅ng/mL (n=12)	t _{max} , h (n=8)	C _{max} , ng/mL (n=8)	AUC ₀₋₂₄ , h-ng/mL (n=7)	AR (n=7)	t _{max} , h (n=4)	C _{max} , ng/mL (n=4)	AUC₀₋₂₄, h⋅ng/mL (n=3)	t _{max} , h (n=3)	C _{max} , ng/mL (n=3)	AUC ₀₋₂₄ , h-ng/mL (n=3)	AR (n=3)
Mean (SD)	3.2 (1.4)	4110 (1850)	48,200 (22,000)	3.0 (1.4)	3260 (832)	36,800 (11,800)	0.946 (0.389)	3.0 (2.1)	3080 (535)	36,,000 (7530)	1.8 (1.3)	3100 (448)	32,800 (9020)	0.965 (0.404)
Median (range)	3.0 (1.5– 6.1)	3700 (1570– 7170)	42,200 (17,600– 94,300)	3.0 (1.5– 6.0)	3520 (2040– 4380)	39,100 (21,000– 54,900)	1.07 (0.328– 1.37)	2.3 (1.5– 6.0)	3160 (2410– 3570)	34,100 (27,900– 44,300)	1.9 (0.4– 3.0)	3330 (2580– 3380)	37,400 (22,400– 38,500)	1.13 (0.504– 1.26)
CV%	41.9	45.0	47.6	45.2	25.5	32.1	41.1	69.7	17.4	20.9	70.8	14.5	27.5	41.9

Table 4. Pharmacokinetics^a

CV=coefficient of variation.

^aPharmacokinetic analysis set.

FIGURES

Figure 1. Patient Disposition. Cohort 1: patients with *MET*-amplified G/GEJ/E adenocarcinoma with measurable disease per RECIST; Cohort 2A: patients with *MET*-amplified G/GEJ/E adenocarcinoma with nonmeasurable disease per RECIST; Cohort 2B: a patient with *MET*-amplified G/GEJ/E adenocarcinoma with measurable disease per RECIST; who had received prior MET antibody therapy; Cohort 2C: patients with non–small-cell lung cancer. *At data cutoff, May 16, 2016.

Figure 2. Percentage change in sum of longest diameter of target lesion(s) per RECIST (A), progression-free survival (B), and overall survival (C) for patients with G/GEJ/E carcinoma. The dashed line in panel A marks the median; error bars in panels B and C indicate 95% CI. NE=not evaluable. *Unconfirmed PR graded as SD.





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A Multicenter Phase 2 Study of AMG 337 in Patients With *MET* -Amplified Gastric/Gastroesophageal Junction/Esophageal Adenocarcinoma and Other Solid Tumors

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Clin Cancer Res Published OnlineFirst October 26, 2018.



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