# *MET* deletion is a frequent event in gastric/gastroesophageal junction/esophageal cancer: a cross-sectional analysis of gene status and signal distribution in 1,580 patients

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**Background:** *MET* gene aberrations are found in several human cancers including gastric, ovarian and lung. In a large multinational cohort of patients with gastric/gastroesophageal junction/esophageal (G/GEJ/E) adenocarcinoma we assessed the MET status with respect to amplification and deletion and correlate the results with the phenotypical gene signal distribution pattern.

**Methods:** Tissue specimens from 1,580 patients were analyzed using a novel fluorescence in situ hybridization (FISH) assay employing a *MET*/CEN-7 IQFISH Probe Mix. *MET* amplification and deletions were defined as a *MET*/CEN-7 ratio  $\geq$ 2.0 and a *MET*/CEN-7 ratio <0.8, respectively. Furthermore, the link between the *MET* gene status and the phenotypical signal distribution was investigated.

**Results:** The prevalence of *MET* amplification and deletions was found to be 7.2% and 8.7%, respectively. Significant differences were observed with regard to geographic regions and sex. The Asian population had the highest percentage of *MET* amplification (9.4%) and the lowest percentage of deletions (3.2%). *MET* deletions was found more frequently among males (10.1%) compared to females (5.3%) and in esophagus (17.6%) compared to the stomach (5.7%). More than 50% of the patients who harbored *MET* gene amplification had a heterogeneous distribution of the FISH signals. Patients with a focal signal distribution were solely to be found among the *MET* amplified population. *MET* deletion were mainly observed in the group of patients with a homogenous signal distribution.

**Conclusions:** The screening data from this cross-sectional study showed that *MET* deletion and amplification are frequent events in G/GEJ/E cancer, which are linked to different phenotypical signal distribution patterns. The role of *MET* deletion in relation to tumor development is not fully understood but it is likely to play a role in the oncogenic transformation of the cells.

**Keywords:** Mesenchymal epithelial transition factor gene (*MET*); fluorescence in situ hybridization (FISH); gastric cancer; amplification; deletion

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## Introduction

The mesenchymal epithelial transition factor gene (MET) is a proto-oncogene located on chromosome 7q31 that encodes a transmembrane receptor with intrinsic tyrosine kinase activity known as Met (or cellular-MET, c-Met) (1). This receptor is also called the hepatocyte growth factor receptor (HGFR) after its ligand; hepatocyte growth factor (HGF) (1-3). Synthesis of HGF by neighboring cells and binding to Met results in tyrosine phosphorylation and activation of the Met receptor. During non-tumorigenic conditions, the paracrine activation of Met has key roles in embryogenesis, wound healing and organ regeneration (4). In cancer patients, aberrant Met signaling can result in suppression of apoptosis and cell proliferation, motility, migration, and invasion, which likely arises from different genetic alterations or dysregulation of MET (3-6). These genetic alterations also include amplification of MET, which has been found in several human cancers, including gastric, ovarian, lung, breast, renal and more (6).

In studies of gastric/gastroesophageal junction/ esophageal (G/GEJ/E) cancer, MET amplification has been reported with a prevalence in the range of 1.5% to 30.5%, depending on the study and the analytical method used (7-18). Using the in situ hybridization (ISH) techniques, such as fluorescence in situ hybridization (FISH), silver in situ hybridization (SISH), or chromogenic in situ hybridization (CISH), a prevalence in the range of 1.5% to 8.3% has been reported (7-15). MET amplification can also be detected using real-time polymerase chain reaction (qPCR); however, the prevalence seems to be somewhat higher than with the different ISH techniques with published data in the range of 3.7% to 30.5% (16-19). The results from these two analytical methods are not directly comparable due to differences in the way the assays detect gene amplification. The qPCR-based assays identify a gain in gene copy number regardless of the underlying cause and are thus unable to discriminate gene amplification from polysomy (6). This discrimination seems to be important as true MET amplification is more likely to lead to oncogenic MET addiction than polysomy (20). Furthermore, in a recent comparison of MET amplifications in solid tumors by in situ and extraction-based methods, large discrepancies was found compared with extraction-based methods such as PCR and NGS (21). Another factor contributing to the variability is the differences in the cut-off selected for the individual assays, thus comparing prevalence data for MET gene amplification across different study populations can

be challenging (9). Recently, in a review by Guo et al., it is likewise diligently described how MET amplification may vary with the technique or assay used (20). Even, comparing results from different ISH assays can be difficult due to the differences in the scoring algorithm used (7-15). When it comes to deletion of the MET gene in patients with G/GEJ/E cancer, far less is known both with respect to the prevalence as well as how this aberration influences the disease processes (22).

Several studies in patients with G/GEJ/E cancer have reported that MET amplification is associated with an unfavorable clinical outcome, which has been shown from data generated both with the ISH and qPCR assays (8,10,16,17,19). The association between MET amplification and a poor disease prognosis has likewise been confirmed in a meta-analysis (23). Beside the disease prognostic characteristics, it has been suggested that MET amplification potentially possesses predictive properties in relation to Met-targeted therapy and, thereby could act as a companion or complementary diagnostic in relation to the tyrosine kinase inhibitors under development for treatment of G/GEJ/E cancer and other indications (13,15,24-26). Available data suggest that treatment plans targeting both Met and Her2 (human epidermal growth factor receptor 2 gene product) may be beneficial. For example, patients with HER2 amplified trastuzumab-resistant esophageal cancer showed that the effect of therapy with afatinib correlated with MET co-amplification (27). Furthermore, resistance towards afatinib in a gastric cancer cell line was reversed by MET knockdown indicating MET amplification as a resistance factor towards afatinib treatment (28).

Recent studies in patients with G/GEJ/E adenocarcinoma have suggested a link between gene amplification and a heterogeneous signal distribution pattern. This link has been shown for both the HER2 and MET genes (9,29). Here, we report data from a large cross-sectional study on MET gene aberrations in patients with G/GEJ/E adenocarcinoma analyzed with a FISH assay using the formamide-free, fast IQFISH hybridization buffer (30,31). The purpose of this exploratory study was to investigate the prevalence of MET amplification and deletions in tissue specimens from a large cohort of patients with G/GEJ/E adenocarcinoma and correlate these findings with the phenotypical gene signal distribution pattern. We present the following article in accordance with the STREGA reporting checklist (available at http://dx.doi.org/10.21037/ atm-20-4081).

#### Methods

#### **Patients/specimens**

The study included 1,580 formalin-fixed and paraffinembedded (FFPE) G/GEJ/E adenocarcinoma specimens consecutively collected from the screening population of an international multi-center phase II trial with the Met tyrosine kinase inhibitor AMG337 (Amgen) (25). The limited demographic and clinical data were collected in relation with the MET eligibility testing for inclusion in the clinical phase II study with AMG337. The G/GEJ/E specimens were cut in sections of 4 µm and mounted on glass slides. Apart from data on the MET gene status and the signal distribution pattern no other demographic and clinicopathological data were available except sex, age and tumor site. The study was conducted according to the Declaration of Helsinki (as revised in 2013), and informed consent was received from patients prior to testing of the biopsy specimens. Prior to study initiation the protocol was reviewed and approved by Institutional Review Boards/Ethics Committees in the respective countries. The study was conducted under a US Investigational New Drug Application and Investigational Device Exemption according to the International Conference on Harmonization (ICH) Good Clinical Practice (GCP) guideline (ClinicalTrials.gov ID: NCT02016534/EudraCT Number: 2013-001277-24). The study was conducted at 97 clinical sites worldwide. Evaluation of the stained slides was performed at one central laboratory in the USA by a technologist and subsequently independently reviewed by a pathologist.

# MET IQFISH Testing

FISH staining using the *MET*/CEN-7 IQFISH Probe Mix (For Investigational Use Only (IUO), Dako/Agilent Technologies) and reagents in the Histology FISH Accessory Kit (Dako/Agilent Technologies) was performed according to the IQFISH staining procedure (9). Following mounting of the FFPE specimens on positively charged glass slides, heat pre-treatment was carried out in a microwave oven. Then pepsin digestion was done at 37 °C followed by ethanol dehydration with subsequent drying prior to probe application. The Texas Red-labeled DNA probe (*MET*) and the fluorescein-labeled PNA probe (CEN-7) were co-denatured with the tissue specimen for 10 minutes at 66 °C followed by hybridization at 45 °C for 90 minutes using a Hybridizer (Dako/Agilent Technologies). Following hybridization, excess probe was washed away with Stringent Wash Buffer at 63 °C for 10 minutes. Then the slides were dehydrated in ethanol baths, dried at room temperature and, finally, the slides were mounted in DAPIcontaining Fluorescence Mounting Medium with glass coverslips.

MET FISH stained slides were inspected using a fluorescence microscope equipped with 20x, 40x and 100x objectives and enumeration was performed at the largest magnification. Ratios of MET/CEN-7 from the invasive tumor area were calculated by counting of signals from 20 nuclei. Based on this ratio, specimens were characterized as amplified (MET/CEN-7  $\geq$ 2.0), non-amplified (MET/ CEN-7 between  $\geq 0.8$  and 2.0), or deletion (MET/CEN-7 <0.8). Specimens having a ratio between 1.8 and 2.2 were considered borderline cases and signals from additional 20 nuclei were counted and the final ratio and MET status was determined from the 40 nuclei. The cut-off level for MET deletion was selected based on previous observations for other genes using FISH assays (32-34). Signals in normal cells within the sample were used to verify staining quality of individual specimens, as these cells are expected to have a normal diploid status with two MET gene signals and two CEN-7 signals in every nucleus. Before the MET FISH assay was used in the study, it underwent an extensive analytical validation (9).

# MET signal distribution

In addition to the assessment of MET gene status (amplified/ non-amplified/deletion), the MET signal distribution pattern was evaluated. In these tumors, MET gene signals can preferentially have a homogeneous or heterogeneous distribution pattern (9). The homogeneous signal distribution is observed when most tumor cells in the tumor specimen are equally amplified or equally non-amplified. The heterogeneous signal distribution is observed when tumor cells having amplified and non-amplified status are intermingled across the tumor. Specimens with the heterogeneous signal distribution are further divided into two categories: (I) When MET amplified tumor cells are grouped together, the signal distribution is categorized as focal, and, (II) when MET amplified tumor cells are interspersed in tissue areas of cells exhibiting low or normal MET/CEN-7 ratio, the signal distribution is defined as mosaic (9). A similar phenotypical signal distribution pattern has been described for HER2, another gene known to be amplified in a subset of patients with G/GEJ/E cancer (29).

 Table 1 Demographic and tumor characteristics of the G/GEJ/E cancer patients screened for *MET* amplification (N=1,580)

Characteristics	N (%)	
Age, years		
Mean (SD)	60.3 (12.1)	
Median (Range)	62.0 (19-96)	
Regions		
Asia <sup>1</sup>	277 (17.5)	
Europe/Australia <sup>2</sup>	1,142 (72.3)	
North America <sup>3</sup>	154 (9.7)	
South America <sup>4</sup>	7 (0.4)	
Sex		
Male	1,132 (71.6)	
Female	448 (28.4)	
Tumor site		
Gastric	981 (62.1)	
Gastroesophageal Junction	169 (10.7)	
Esophageal	233 (14.7)	
Metastatic	198 (12.5)	

<sup>1</sup>Republic of Korea; <sup>2</sup>Australia, Belgium, Czech Republic, France, Germany, Greece, Hungary, Italy, Poland, Russian Federation, Spain, United Kingdom; <sup>3</sup>Canada, United States of America; <sup>4</sup>Chile, Peru; G/GEJ/E, gastric/gastroesophageal junction/ esophageal; *MET*, mesenchymal epithelial transition factor gene.

# Statistical analyses

The statistical analyses were performed based on all screened subjects with an evaluable *MET* gene status assessment. Descriptive statistics for continuous variables are provided as mean, standard deviation, and median, while frequency and percent distributions were provided for non-continuous variables. To investigate *MET* gene status prevalence within each subgroup based on different baseline covariates, Chisquare tests were performed. P values generated from the analyses were mainly included as a descriptive measure suggesting comparative strength of association rather than a test of hypotheses. Accordingly, nominal P values are reported with no adjustment for multiplicity.

# Results

The results presented in this study are based on FISH data

from 1580 patients with G/GEJ/E adenocarcinoma screened for *MET* amplification in relation to an international multicentre phase II trial. The vast majority of the patients came from Europe and Australia (72.3%) but patients from Asia, North America and South America were also screened for the trial (*Table 1*). The median age of the patient population was 62.0 years with more than 71% being males. For 62.1% of the patients, the tumor was localized in the stomach and for the remaining part of the population the tumor sites were esophagus, GEJ or metastatic.

# MET gene status

Based on the screening of the 1,580 G/GEI/E cancer patients, the prevalence rate of MET amplification was found to be 7.2% (N=113) and 8.7% (N=138) for MET deletion. The MET/CEN-7 ratio showed a wide range with ratios ranging from 0.5 up to 39.2. Differences were observed regarding geographic regions, sex and tumor site, as shown in Table 2. The Asian population had the highest percentage of patients with MET amplified tumors (9.4%) and concurrently the lowest percentage of MET deletion (3.2%) (P<0.0001, Chi-square test). With regard to sex, MET deletion was more frequent among males (10.1%) compared to females (5.3%) (P=0.0082, Chi-square test). The tumor site seems to have less influence on the prevalence of MET amplification except for the metastatic tumors where 9.1% were found to be amplified, which is slightly higher than for the other tumor sites. For the patients with MET deletion, a prevalence of 17.6% was found when the tumor was located in esophagus compared to 5.7% in the stomach (P<0.0001, Chi-square test).

# MET signal distribution

In addition to the calculation of the *MET*/CEN-7 ratios and determination of gene status an assessment of the signal distribution pattern was performed. Data from this assessment was available from 1,579 of the 1,580 screened patients as shown in *Table 3*. In total, 6.3% of the screened population showed a heterogeneous signal distribution, either focal or mosaic with the mosaic distribution pattern slightly more prevalent (4.0%) than the focal distribution (2.3%) (P=0.0051, One proportion z-test). No major differences were observed regarding the distribution between the heterogeneous and the homogenous pattern when it comes to sex, tumor site, and geographic regions (*Table 3*).

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Table 2 WET screening results and gene status based on WET/CELV-7 Table (IV=1,500)				
	Amplified ( <i>MET</i> /CEN-7 ≥2.0), n (%)	Non-amplified ( <i>MET</i> /CEN-7 <2.0 and MET/CEN-7 ≥0.8), n (%)	Deletion ( <i>MET</i> /CEN-7 <0.8), n (%)	Total, n (%)
All	113 (7.2)	1,329 (84.1)	138 (8.7)	1,580 (100.0)
Regions				
Asia	26 (9.4)	242 (87.4)	9 (3.2)	277 (100.0)
Europe/Australia	75 (6.6)	953 (83.4)	114 (10.0)	1,142 (100.0)
North America	11 (7.1)	128 (83.2)	15 (9.7)	154 (100.0)
South America	1 (14.3)	6 (85.7)	0 (0.0)	7 (100.0)
Age, years				
Mean (SD)	59.1 (13.0)	60.0 (12.0)	64.0 (11.6)	60.3 (12.1)
Median (range)	59.0 (25–85)	62.0 (19–87)	65.0 (26–96)	62.0 (19–96)
Sex				
Male	76 (6.7)	942 (83.2)	114 (10.1)	1,132 (100.0)
Female	37 (8.3)	387 (86.4)	24 (5.3)	448 (100.0)
Tumor site				
Gastric	70 (7.1)	855 (87.2)	56 (5.7)	981 (100.0)
GEJ	11 (6.5)	137 (81.1)	21 (12.4)	169 (100.0)
Esophageal	14 (6.0)	178 (76.4)	41 (17.6)	233 (100.0)
Metastatic	18 (9.1)	160 (80.8)	20 (10.1)	198 (100.0)
MET/CEN-7 ratio				
Mean (SD)	7.1 (5.7)	1.1 (0.2)	0.7 (0.1)	
Median (range)	5.5 (2.0–39.2)	1.1 (0.8–<2.0)	0.7 (0.5–<0.8)	

Table 2 MET	screening results and	gene status based	on MET/CEN-7	ratio (N=1.580)
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MET, mesenchymal epithelial transition factor gene. CEN-7, centromere of chromosome 7; GEJ, gastroesophageal junction.

The link between MET gene status and signal distribution is shown in Table 4. For patients with MET amplified tumors, a total of 51.3% also had a heterogeneous signal distribution. This link was especially strong for the group of patients with a focal signal distribution pattern as no patients with focal heterogenous signal distribution were found among the MET non-amplified or deleted groups, as shown in Figure 1. The mosaic heterogeneous signal distribution was detected in 64 out of the 1,579 screened patients and found among all three MET gene status categories; amplified, non-amplified and deletion. The proportion of patients who had MET amplification and a mosaic signal distribution was 34.4% (Table 4). Figure 1 visualize the distribution of the MET/CEN-7 ratios in relation to the three gene signal distribution patterns. Patients who harbored tumors with MET deletion were

mainly found in the group with a homogenous signal distribution, as shown in *Table 4* and *Figure 1B*.

#### Discussion

So far, our knowledge on *MET* amplification in patients with G/GEJ/E cancer has been based on data from relatively small studies that have included up to a few hundred patients (7-19). Here, we report *MET* gene aberration data from 1,580 patients with G/GEJ/E adenocarcinoma analyzed with a validated *MET* FISH assay using the formamide-free, fast IQFISH hybridization buffer. To the best of our knowledge, it is the largest cohort reported so far. Compared to other analytical methods for detection of gene aberrations, such as qPCR and nextgeneration sequencing, the ISH technology offers the

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**Table 3** *MET* screening results and signal distribution pattern  $(N=1,579)^1$ 

-				
	Focal Heterogeneous, n (%)	Mosaic Heterogeneous, n (%)	Homogenous, n (%)	Total, n (%)
All	36 (2.3)	64 (4.0)	1,479 (93.7)	1,579 (100.0)
Regions				
Asia	11 (4.0)	6 (2.2)	259 (93.8)	276 (100.0)
Europe/Australia	23 (2.0)	55 (4.8)	1,064 (93.2)	1,142 (100.0)
North America	2 (1.3)	2 (1.3)	150 (97.4)	154 (100.0)
South America	0 (0.0)	1 (14.3)	6 (85.7)	7 (100.0)
Sex				
Male	25 (2.2)	45 (4.0)	1,061 (93.8)	1,131 (100.0)
Female	11 (2.5)	19 (4.2)	418 (93.3)	448 (100.0)
Tumor site				
Gastric	19 (1.9)	41 (4.2)	920 (93.9)	980 (100.0)
GEJ	3 (1.8)	5 (3.0)	161 (95.3)	169 (100.0)
Esophageal	7 (3.0)	12 (5.2)	214 (91.8)	233 (100.0)
Metastatic	7 (3.5)	7 (3.5)	184 (93.0)	198 (100.0)

<sup>1</sup>Data on signal distribution was missing for one patient. *MET*, mesenchymal epithelial transition factor gene; GEJ, gastroesophageal junction.

Table 4 MET ge	ene status based on META	/CEN-7 ratio and	signal distribution	pattern $(N=1,579)^1$
0				

	Amplified ( <i>MET</i> /CEN-7 ≥2.0), n (%)	Non-Amplified ( <i>MET</i> /CEN-7 <2.0 and MET/CEN-7 ≥0.8), n (%)	Deletion ( <i>MET</i> /CEN-7 <0.8) n (%)	Total, n (%)
All	113 (7.2)	1,328 (84.1)	138 (8.7)	1,579 (100.0)
Signal distribution				
Focal heterogeneous	36 (100.0)	0 (0.0)	0 (0.0)	36 (100.0)
Mosaic heterogeneous	22 (34.4)	40 (62.5)	2 (3.1)	64 (100.0)
Homogenous	55 (3.7)	1,288 (87.1)	136 (9.2)	1,479 (100.0)

<sup>1</sup>Data on signal distribution was missing for one patient. *MET*, mesenchymal epithelial transition factor gene; CEN-7, centromere of chromosome 7.

possibility to study the connection between genotype and phenotype in the tumor tissues. In the current study, this option was used to investigate the link between *MET* gene status and the phenotypical signal distribution pattern, which has been described previously for both the *HER2* and the *MET* gene (9,29). Determination of gene copy numbers by FISH is a well-established analysis method used for many years by pathologists and technologists to determine clinically relevant *HER2* levels in breast and gastric cancer. The current FISH ratio method in which *MET* loci and the centromeric region of chromosome 7 are detected using separate fluorescent labels has been thoroughly validated (9) and is completely analogous to determination of the HER2/CEN17 ratio also performed in gastric cancer tissue. Therefore, we have no reason to suspect a bias in the reported results should be introduced by the testing method.

The median age of the study population was 62.0 years with the typical sex distribution for G/GEJ/E cancer of approximately two third being males (35). Among the 1,580 screened patients, 113 cases were found to be amplified corresponding to a prevalence rate of 7.2%, which is within the range of what other studies have shown using different type of ISH assays (7-15). When comparing



**Figure 1** Dot plot of *MET*/CEN-7 ratios versus the signal distribution pattern for all patient specimens (N=1,579) (A). Data on signal distribution is missing for one patient. Dot plot of *MET*/CEN-7 ratios versus the signal distribution pattern for all patient specimens up to a *MET*/CEN-7 ratio of 3.0 (N=1,499) (B). MET, mesenchymal epithelial transition factor gene; CEN-7, centromere of chromosome 7.

the different geographical regions, MET amplification was significantly more prevalent in the Asian patient population compared to patients screened in other regions. No published studies have investigated this population difference previously, hence, it is difficult to confirm if a region/race difference exists with respect to the prevalence rate of MET amplification in G/GEJ/E cancer patients. However, a couple of studies conducted in China have shown relatively high prevalence rates; one using a FISH assay showed a prevalence of 8.3% and another study using a qPCR assay detected MET amplification in 30.5% of the patients (8,16). Again, it should be underlined that results from these two types of assays are not directly comparable but data from these studies could indicate a relatively high rate of MET amplification among Asian patients. No major difference was found in the prevalence rate for MET amplification between the three tumor sites (G/GEJ/E) except for the metastatic sites, which were slightly higher. The information in the literature on tumor site differences is limited, but a single published study with similar types of patients seems to confirm our findings (13).

A link between gene amplification and a heterogeneous signal distribution pattern was found in the study. More than 50% of the patients with *MET* amplified tumors had a heterogeneous gene signal distribution, either focal or mosaic. For one group of patients, this link was especially strong, as all patients having tumors with a focal signal distribution were among the 113 patients with *MET* amplification. The link between *MET* amplification and a

heterogeneous signal distribution pattern was also shown when the assay was validated (9), and a similar correlation has previously been reported among patients with HER2 amplified G/GEJ/E tumors (29). It is known that Met plays a key role in the malignant transformation especially in tumors with MET amplification, leading to an invasive growth with increased cell motility and dissociation as well as tissue infiltration and stimulation of angiogenesis (36). The fact that the focal heterogeneous signal distribution is ultimately linked to MET amplification status is surprising and it could indicate that MET amplification is a requirement for the appearance of focal heterogenous signal distribution. Previous studies in adenocarcinomas of the colon have indicated that HGF secreted from stromal cells in the tumor microenvironment activates Wnt- and betacatenin signaling, which is important for maintenance of cancer stem cells and de-differentiation of non-cancer stem cells into cancer stem cells (37). Since Met signaling has been shown to enhance tumor growth, stimulate cancer cell motility and increase invasiveness, it would be interesting in future studies to determine if focal areas with MET amplified tumor cells in adenocarcinomas of G/GEJ/E are clinically relevant entities that populate cancer stem cells or highly tumorigenic cells and drives disease progression. The link between the mosaic signal distribution and gene amplification was somewhat weaker and here, approximately 35% of the patients had MET amplified tumors.

In the current study, we also investigated the prevalence rate of *MET* deletion, defined as *MET*/CEN-7 <0.8, and

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surprisingly, this aberration was found in 138 patients corresponding to 8.7% of the screened population. Unexpectedly, this prevalence rate was higher than for MET amplified tumors and the MET deletions were almost exclusively found among tumors with a homogeneous signal distribution pattern. Data in the literature on MET tumor deletion is very limited and we have only been able to identify a single publication on the topic. This is from a study investigating MET gene aberrations in a group of patients with salivary gland cancer using a FISH assay and here, the prevalence of MET tumor deletion was found to be 7.7% (22). Furthermore, in this study MET amplification and MET deletion were clearly associated with patient age, male gender, increased tumor size, lymph node metastasis and high-grade malignancy. The implications of MET deletion in relation to tumor growth and survival is largely unknown but based on an experimental study using a liver cancer model, it has been suggested that loss of MET signaling increases cellular stress leading to an imbalance in redox homeostasis in the hepatocytes resulting in tumor progression (38). There seems to be a greater variability for the MET deletion data compared to what has been observed for MET amplification with respect to the different study variables. A significant sex difference was shown with a nearly doubling of the MET deletion rate in males (10.1%) compared to females (5.3%). Furthermore, a likewise significant difference in the prevalence of MET deletions was found regarding tumor site. Here, a prevalence of 5.7% was found for the stomach, increasing to 12.4% and 17.6% for the GEJ and esophagus, respectively.

The data obtained from this screening population does not give direct mechanistic information regarding activity of Met transcripts or proteins in these tumors. Met expression in the partially MET deleted tumors could be highly active due to changes in transcriptional regulation or due to activating mutations. Therefore, additional understanding of the mechanisms affected by Met during cancer progression could be relevant for new treatment options and optimal biomarker detection. In the current study access to demographic and clinicopathological information was limited, which prevented correlation of MET gene status with this type of information. Although this is a limitation of the study, we believe that observations on the MET amplification and deletion as well as the phenotypical gene signal distribution pattern are relevant, especially as they are based on data from a large population of patients with G/GEJ/E cancer.

In summary, the current study provided the largest dataset on the prevalence of *MET* amplification and

deletion in patients with G/GEJ/E cancer. The prevalence of MET amplification found in our population was 7.2%. Furthermore, the data confirmed an association between MET amplification and the heterogeneous signal distribution previously shown in relation to the validation of the MET IQFISH assay (9). Unexpectedly, our study showed a relatively high prevalence of MET gene deletion, which has not previously been described for G/GEJ/ E cancer. A total of 8.7% of the patients were found to have tumors with MET deletion and these were almost exclusively linked to a homogeneous signal distribution. Current knowledge of the role of MET deletion in relation to tumor development is limited, but since MET deletions have been observed in context of increased malignancy it may play a role in oncogenic transformation of the cells as has been described for MET amplifications.

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### Footnote

Reporting Checklist: The authors have completed the STREGA reporting checklist. Available at http://dx.doi. org/10.21037/atm-20-4081

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*Ethical Statement:* The authors are accountable for all aspects of the work in ensuring that questions related

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to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted according to the Declaration of Helsinki (as revised in 2013), and informed consent was received from patients prior to testing of the biopsy specimens. Prior to study initiation the protocol was reviewed and approved by Institutional Review Boards/Ethics Committees in the respective countries. The study was conducted under a US Investigational New Drug Application and Investigational Device Exemption according to the International Conference on Harmonization (ICH) Good Clinical Practice (GCP) guideline. The study was conducted at 97 clinical sites worldwide. ClinicalTrials. gov ID: NCT02016534 (https://clinicaltrials.gov/ct2/ show/NCT02016534). EudraCT Number: 2013-001277-24 (https://www.clinicaltrialsregister.eu/ctr-search/ search?query=2013-001277-24).

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