Intercomparison of detection and quantification methods for methane emissions from
 the natural gas distribution network in Hamburg, Germany

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16 Abstract:

17 In August and September 2020, three different measurement methods for quantifying methane 18 (CH<sub>4</sub>) emission from leaks in urban gas distribution networks were applied and compared in 19 Hamburg, Germany: the "mobile", "tracer release" and "suction" methods. 20 The mobile and tracer release methods determine emission rates to the atmosphere from 21 measurements of CH<sub>4</sub> mole fractions in the ambient air, and the tracer release method also 22 includes measurement of a gaseous tracer. The suction method determines emission rates by 23 pumping air out of the ground using soil probes that are placed above the suspected leak 24 location. The quantitative intercomparison of the emission rates from the three methods at a 25 small number of locations is challenging because of limitations of the different methods at different types of leak locations. 26

27 The mobile method was designed to rapidly quantify the average or total emission rate of many 28 gas leaks in a city, but it yields a large emission rate uncertainty for individual leak locations. 29 Emission rates determined for individual leak locations with the tracer release technique are 30 more precise because the simultaneous measurement of the tracer released at a known rate at 31 the emission source eliminates many of the uncertainties encountered with the mobile method. 32 Nevertheless, care must be taken to properly collocate the tracer release and the leak emission 33 points to avoid biases in emission rate estimates. The suction method could not be completed 34 or applied at locations with widespread subsurface CH<sub>4</sub> accumulation, or due to safety 35 measures, and this sampling bias may be associated with a bias towards leak locations with low 36 emission rates. The leak locations where the suction method could not be applied were the biggest emitters as confirmed by the emission rate quantifications using mobile and tracer 37 38 methods and an engineering method based on leak's diameter, pipeline overpressure and depth 39 at which the pipeline is buried. The corresponding sampling bias for the suction technique led 40 to a low bias in derived emission rates in this study. It is important that future studies using the 41 suction method account for any leaks not quantifiable with this method in order to avoid biases, 42 especially when used to inform emission inventories.

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### 48

# **1** Introduction

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50 Natural gas combustion has a lower carbon footprint than combustion of other fossil fuel 51 sources for the same thermal output (EIA, 2021). However, fugitive methane (CH<sub>4</sub>) emissions 52 can significantly turn the balance in terms of climate impact (Alvarez et al., 2012) because the 53 global warming potential of CH<sub>4</sub> over a 20-year time scale is 84 times higher than that of carbon 54 dioxide ( $CO_2$ ) (Myhre et al., 2013). The atmospheric abundance of  $CH_4$  has increased about 55 2.5-fold since the pre-industrial era (Bousquet et al., 2006). Following a short period of stable 56 levels after the year 2000, atmospheric CH<sub>4</sub> has continued to increase since 2006. Worden et al (2017) concluded that about 50 to 80% of the post-2006 increase originated from fossil 57 58 sources and Jackson et al. (2020) attributed the accelerated increase of 6 - 13 ppb yr<sup>-1</sup> from 59 2014 to 2017 (Nisbet et al., 2019), equally to the emission increase from fossil and agriculture 60 sectors.

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Gas distribution networks in cities are subject to maintenance programs by the operators to 62 detect and fix leakages that occur, as CH<sub>4</sub> is an incendiary gas and can be explosive at 63 64 concentrations between 4 and 16% in ambient air (DVGW, 2022). Since the safe operation of 65 the distribution network and leak repair is the primary objective of this maintenance, 66 quantification of emissions from leakages is rarely performed. The absence of regulations on 67 CH<sub>4</sub> emissions is another reason why leak rates are not routinely quantified, however CH<sub>4</sub> 68 emissions from the energy sector needs to be addressed properly within the EU CH<sub>4</sub> strategy 69 by 2050 (EC, 2020). Nevertheless, from the perspective of climate change and possible mitigation options, it is important that emissions from gas leakages are (i) quickly detected and 70 71 fixed and (ii) well quantified. Weller et al. (2020) and Alvarez et al. (2018) respectively 72 reported 5 and 1.6 times higher CH<sub>4</sub> emissions from leaks in the US gas distribution network 73 based on such observations compared to the national inventory reports.

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75 Leaks from buried pipelines can be due to corrosion or failure/defects in joints or materials 76 (EPA, 1996). When a leak occurs on a buried urban gas pipeline, the gas will generally 77 accumulate in the air space below the surface and then find its path to the atmosphere through 78 a single or several surface outlets. The outlets can be either unpaved soil surfaces, cracks in the 79 road or pavements, or associated with different types of cavities (manholes, communication 80 covers, rain drains, etc.). The major outlet is generally the one with the highest overall 81 permeability for gas released from the buried natural gas pipeline. On the way from the leak 82 location on a buried pipeline to the atmosphere through outlets, CH<sub>4</sub> may be oxidized by 83 methanotrophs in the soil and/or merge with CH<sub>4</sub> from other sources, e.g. biogenic CH<sub>4</sub> 84 emissions from sewage system.

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86 Routine leak surveys in Germany are conducted by walking with handheld CH<sub>4</sub> sensors above buried pipelines, referred to as the carpet method (DVGW, 2019). The success of leak detection 87 with the carpet method depends primarily on soil permeability (Ulrich et al., 2019), which is 88 89 influenced by soil moisture, texture, soil organic content and the location of the groundwater 90 table (Wiesner et al., 2016). Based on risk of explosion, gas leaks are classified into four types: 91 A1, A2, B and C (DVGW, 2019). This classification is based on the accumulation of CH<sub>4</sub> in 92 cavities (e.g. manholes, rain drains, etc.) or buildings and the distance of gas leaks to buildings 93 and cavities. If natural gas leaks into buildings or cavities, the leak classifies as A1, and it must 94 be repaired immediately to minimize explosion risk. If the gas leak has a distance up to 1 m to 95 buildings and does not fill cavities, it is classified as A2, and it must be fixed within a week. If 96 the distance is between 1 to 4 m to buildings, the leak is classified as B and the repair time 97 window is three months, and if the distance is more than 4 m then, the leak is considered as C

category and can be fixed according to the scheduled repair plan. There are 6,500 km of low
 pressure and 250 km of medium pressure gas pipelines in Hamburg which are monitored

100 every 4 years with the carpet method based on the national regulations in Germany. Gas

- 100 every 4 years with the carpet method based on the national regulations in Germany. Gas 101 leaks in cities are not quantified and thus also not a parameter affecting the course of action.
- 101 *Moreover, high pressure pipelines are monitored on annual basis with additional helicopter-*
- 103 based measurement platform.
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105 In recent years, mobile measurement methods using vehicles with fast and high-precision laser 106 instrumentation have been established for leak detection and emission quantification in 107 numerous cities (Fernandez et al., 2022; Defratyka et al., 2021; Luetschwager et al., 2021; 108 Keyes et al., 2020; Maazallahi et al., 2020; Ars et al., 2020; Weller et al., 2018; von Fischer et 109 al., 2017; Jackson et al., 2014). In-situ measurements of atmospheric CH<sub>4</sub> from mobile vehicles 110 are used to pinpoint and quantify CH<sub>4</sub> emission sources at street level in urban areas. The mobile method was calibrated using above-ground controlled release experiments, in which 111 112 known amounts of CH<sub>4</sub> were released from gas cylinders (Weller et al., 2019). Simultaneous 113 measurements of carbon dioxide  $(CO_2)$  and ethane  $(C_2H_6)$  can provide valuable additional information for attributing CH<sub>4</sub> sources (Maazallahi et al., 2020). A characteristic of the 114 115 resulting emissions distribution from gas distribution grids in cities is the existence of a few leak locations with very high leak rates, up to 100 L min<sup>-1</sup>, resulting in a right-skewed leak 116 117 emission rate distribution (Weller et al., 2020). Usually about 10% of the leaks are responsible 118 for between 30% to 70% of the emissions (Weller et al., 2019; Maazallahi et al., 2020). 119 Therefore, the CH<sub>4</sub> emission from the gas distribution system can be reduced very effectively 120 if the largest leaks can be found and fixed quickly, thus augmenting the routine leak detection 121 (carpet method) and repair programs with the mobile method.

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123 The tracer dispersion method is another method to quantify CH<sub>4</sub> emissions from point and area 124 sources. In this method, a tracer gas is released at a known rate close to the outlet of the gas leak, and both tracer and target gas concentrations are measured downwind. From these 125 126 measurements and the known tracer gas release rate, the target gas emission rate can be 127 determined with an uncertainty of  $\pm 15\%$  (Lamb et al., 1995) or less than 20% (Fredenslund et 128 al., 2019). Lamb et al. (2015) applied the tracer method to quantify leaks from urban 129 underground pipelines where they reported moderate agreement ( $\pm$  50%) to excellent 130 agreement ( $\pm$  5%) between the tracer and high-flow sampler method.

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132 Another approach to quantify underground leak rates from buried gas pipelines is the so-called 133 suction method. In this method air is pumped out of the ground at a known rate via probes 134 surrounding the underground leaks until an equilibrium CH<sub>4</sub> mixing ratio is reached in air out-135 flow, from which the CH<sub>4</sub> leak rate can be calculated. In Germany, this approach is applied to 136 a limited number of leak locations, which do not have to be repaired immediately or within 1 week. Suction measurements normally find leak rates that are  $< 2 \text{ Lmin}^{-1}$  (E.ON, personal 137 138 communication, 2020). The reported uncertainty range of this method is  $\pm$  10% based on 23 139 measurements in the 1990s (E.ON, personal communication, 2020). The discrepancy between 140 these rather low leak rates compared to leak rates inferred with the mobile method calls for 141 further investigation, since the suction method is also employed to derive network-wide 142 emission factors for the German country-wide gas distribution network (Federal Environment 143 Agency, 2020).

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Hendrick et al. (2016) used surface flux chamber measurements carried out between 2012 and
2014 to estimate gas leak rates from 100 leak locations in the Boston area that were detected

- 147 using mobile measurements (n = 45) in 2011 from Phillips et al. (2013) and additional locations 148 from later mobile surveys (n = 55). They reported CH<sub>4</sub> emission rates from gas leaks ranging 149 from 0.003 g min<sup>-1</sup> to 16 g min<sup>-1</sup>, corresponding to roughly 0.0 - 24.4 L min<sup>-1</sup>. They also 150 reported that their estimate using chamber measurements underestimated total CH<sub>4</sub> emissions, 151 likely because the chambers didn't capture the total CH<sub>4</sub> emitted from the leak. This is similar
- to the enclosure measurements results from Weller et al. (2018).
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154 The flow through a hole in a pipeline can also be calculated theoretically and empirically from 155 the physical properties of the hole, mainly the ratio of hole to pipeline diameter and the 156 overpressure in the pipeline. There are three different engineering model types to estimate 157 emissions from gas leaks: the hole model, the rupture model and modified models to bridge the 158 gap between hole and rupture models (Hu et al., 2020; Moloudi and Esfahani, 2014; Yuhua et 159 al., 2002; Arnaldos et al., 1998). These types of models are either to estimate leak strength from 160 a pipeline in open space or a buried pipeline. A leak on a buried pipeline has higher surrounding resistance depending on soil conditions compared to a situation where the pipeline is in open 161 162 space. Such models have been used to quantify emissions from holes in pipelines in open space 163 (Hou et al., 2020; Manda and Morshed, 2017; Moloudi and Esfahani, 2014; Mahgerefteh, Oke 164 and Atti, 2005; Yuhua et al., 2003; Kayser and Shambaugh, 1991) but also from buried 165 pipelines (Liu et al., 2021; Ebrahimi-Moghadam et al., 2018; Okamoto and Gomi, 2011; Yan, 166 Dong and Li, 2015). Cho et al. (2021) introduced a model, which takes into account soil 167 properties including absolute and relative permeability and porosity, the underground spread of the leak, surface CH<sub>4</sub> mole fractions and depth of the buried pipeline based on experiments 168 169 with a controlled release rate. This model was calibrated based on release rates ranging from 1.3 g min<sup>-1</sup> to 5.7 g min<sup>-1</sup>, corresponding to roughly 2.0 - 8.7 L min<sup>-1</sup>. 170

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172 In this study, we present results from measurements with the mobile, the tracer release and the 173 suction methods in Hamburg, Germany, in August and September 2020. We present the 174 quantitative emission estimates as well as a qualitative intercomparison of the three methods, 175 in particular related to the applicability and the strengths and weaknesses of the different 176 methods at different leak locations. We investigate differences between the leaks detected from 177 mobile measurements and leak locations reported from the routine leak detection surveys 178 performed by the local gas utility (hereinafter LDC (Local Distribution Company)). Finally, 179 we discuss implications of our study for national emission inventories.

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## **2** Materials and Methods

#### 182 **2.1 Campaign preparation and general overview**

As a preparation for the intercomparison campaign, all partners contributed to the preparation of an "intercomparison matrix" where the characteristics and deployment details of the different methods were specified. This matrix is provided in section S.1 of the Supplemental Information (SI). The matrix includes descriptions related to the identification of gas leaks, the quantification of gas leaks, adjustments of the method to the intercomparison exercise and upscaling. It also laid out an initial plan for the intercomparison in terms of identification of suitable locations and deployment of the different methods.

- 190 According to this plan (Fig. 1), we first applied the mobile method to identify potential gas leak
- 191 locations, namely leak indications (LIs). When the mobile method had detected one or more
- 192 emission outlets (See Sect. S.2 in SI) and classified them as a potential gas leak location, the
- 193 carpet method was applied to confirm the leak and determine the confine leak location. Some
- additional locations that had previously been identified by the carpet method (leak categories
- 195 B and C) were added to the list of target locations.

196 Following leak detection, the mobile quantification method (multiple transects) was applied on 197 all the locations and the tracer and suction methods were applied at the confirmed leak 198 locations, and with some restrictions regarding safety and method capacities. The release 199 location for the tracer quantification method was confirmed based on surface screening using 200 a handled methane analyzer. For comparison of the mobile and tracer release methods with the suction and hole methods we assumed that (i) a steady state between pipeline leakage under-201 202 ground CH<sub>4</sub> accumulation and emission to the atmosphere had been reached (Kirchgessner et al., 1997) and (ii) methanotrophs and methanogens have negligible impact on quantification of 203 204 gas leak emissions. Thus, the total emission rate of all outlets in the vicinity of a leak location 205 is equal to the natural gas emission rate from the pipeline leak. We will discuss implications of 206 the above assumptions for selected cases. After leak repair, the LDC reported leak hole sizes, pipeline diameters and pipeline operational pressures, allowing leak rate estimation with the 207 208 hole method.



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Figure 1 – Flowchart of application of leak detection methods (blue colors) and quantification methods (red colors) followed by repair actions and intercomparison of

212 the detection and quantification methods

#### 213 2.2 Measurements setups

#### 214 2.2.1 Mobile measurement setup

215 Onboard the measurement vehicle (VW Transporter) we operated two cavity ring-down 216 spectrometers (CRDS), model G2301 and model G4302 (Picarro, Santa Clara, California, USA). The G2301 measures CH<sub>4</sub>, CO<sub>2</sub> and water vapor (H<sub>2</sub>O) at a flow rate of  $\approx 0.2$  L min<sup>-1</sup> 217 and 0.3 Hz frequency. The G4302 has a flow rate of  $\approx 2.2$  L min<sup>-1</sup> and sampling frequency of 218 219 about 1 Hz for CH<sub>4</sub>, C<sub>2</sub>H<sub>6</sub> and H<sub>2</sub>O. The air intake for both instruments was from the same 220 tubing attached to the front bumper. This setup allowed us to directly compare the 221 enhancements observed from the two instruments during surveys. The G4302, which is in a 222 shape of a backpack, was also used in attribution of outlets emissions in walking surveys to 223 check presence of  $C_2H_6$  in emission outlets.

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#### 225 2.2.2 Tracer release measurement setup

The tracer release method was applied by releasing acetylene ( $C_2H_2$ ) at the emission outlet 226 227 identified by the mobile leak detection and confirmed by the carpet method. The tracer gas 228 was released at the main emission outlet, which was confirmed by surface screening using a handheld CH<sub>4</sub> analyzer. Tracer release rates between 1.3 and 2.6 L min<sup>-1</sup> from a gas cylinder. 229 230 A Picarro CRDS, G2203 instrument was used to measure CH4 and C2H2 mole fractions 231 continuously with  $\approx 0.3$  Hz frequency. The instrument was installed in a measurement vehicle 232 (VW Caddy), and air was sampled from the atmosphere through an inlet on the roof about 2m 233 above ground. The tracer method was applied either in static mode, where air was sampled in 234 one or a few locations downwind from the outlets and tracer release locations (n = 11) or mobile 235 mode (n = 5), where the plumes were transected while measuring concentrations of CH<sub>4</sub> and 236 C<sub>2</sub>H<sub>2</sub>. The choice of mode depended on the site conditions including road accessibility and 237 wind direction. The tracer release setup including instrumentation used as well as mobile mode 238 is described in detail in Mønster et al (2014), and the principle of the static mode is described 239 in Fredenslund et al (2010).

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#### 241 2.2.3 Suction measurement setup

In the suction method, 12 probes were used to insert in the soil around the confirmed gas leak location by the LDC. The probes are connected to a pump to extract accumulated subsurface CH<sub>4</sub> from the leak. CH<sub>4</sub> mole fraction at the outflow is measured with a Flame Ionization Detector (*MEEM*, 2018).

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#### 247 2.2.4 Carpet method setup

Leak detection experts from the LDC operate a methane detector (Sewerin instruments, Gütersloh, Germany) on a rolling device, where a plastic cover (the carpet) moves over the ground and provides a loose seal to the surrounding atmosphere, facilitating preferential analysis of air emanating from the surface right below the carpet. The instrument gives an acoustic signal when a high  $CH_4$  from a potential leak has been detected. The instrument can detect  $C_2H_6$  with a gas chromatograph, which take about couple of minutes per outlet location.

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#### 255 **2.3 Detection, confirmation and attribution of emissions at gas leak locations**

#### 256 **2.3.1 Mobile detection of possible leak location**

For leak detection with the mobile method, we first evaluated  $CH_4$ ,  $C_2H_6$  and  $CO_2$  signals during mobile surveys. If (i)  $CH_4$  and  $C_2H_6$  signals were observed with a ratio of less than 10% with no  $CO_2$  signal or (ii)  $CH_4$  was observed (< 500 ppb enhancement on G4302) with no  $C_2H_6$ and  $CO_2$  signals, then we parked the mobile measurement car, detached the G4302 analyzer from the system and searched for gas outlets on foot with the G4302. This detailed search for

outlets was performed to (i) confirm the presence of both  $CH_4$  and  $C_2H_6$  signals (ii) map the

spatial spread of outlets and (iii) spatially constrain the possible gas leak location. The reported possible gas leak locations from the mobile method were then reported to the LDC for confirmation and localization of the leak with the carpet method and subsequent underground measurements.

#### 268 2.3.2 Attribution of leak indication signals from mobile measurements

269 To attribute an observed leak indication (LI) from mobile measurements to a source category, 270 namely fossil, microbial and combustion, we used CO<sub>2</sub> and C<sub>2</sub>H<sub>6</sub> signals, which were 271 continuously measured along with  $CH_4$ . We quantitatively evaluated  $C_2:C_1$  ratios (%) when (i) 272 the CH<sub>4</sub> enhancements were larger than 0.5 ppm (ii)  $C_2H_6$  enhancements were also larger than 273 15 ppb and (iii) the determination coefficient ( $\mathbb{R}^2$ ) of the linear regression between CH<sub>4</sub> and 274 C<sub>2</sub>H<sub>6</sub> was larger than 0.7. If CH<sub>4</sub> signals in mobile measurements were associated with CO<sub>2</sub> 275 and high  $C_2H_6$  mole fractions ( $C_2:C_1 > 10\%$ ), we attributed those emissions to combustion (Maazallahi et al., 2020). When we repeatedly observed CH<sub>4</sub> enhancements, no CO<sub>2</sub> 276 277 enhancements and C<sub>2</sub>:C<sub>1</sub> ratios between 1 and 10%, or we observed persistent CH<sub>4</sub> signals in 278 several passes we did further on-foot inspection of the outlets. If the emissions from the outlets clearly pointed to a fossil origin based on the CH<sub>4</sub> and C<sub>2</sub>H<sub>6</sub> signals, we labeled the locations 279 280 as potential gas leak locations and reported them to the LDC for confirmation. We only considered a location as a gas leak for further investigation if the LDC confirmed the existence 281 282 of a gas leak.

283 If at a particular location, we observed several CH<sub>4</sub> maxima, for example from different outlets,

- we considered the "strongest" outlet as the main emission point. The "strongest" emission point refers to a point where we observed the highest CH<sub>4</sub> mole fraction when the G4302 intake inlet was put at a distance of  $\approx 2 - 5$  cm above the surface or outlet. When several emission outlets
- with similar mole fractions were found, we considered the spatial average of the coordinates as the main emission point. The tracer method then released  $C_2H_2$  at the main outlet emission point.
- The LDC reported a C<sub>2</sub>:C<sub>1</sub> ratio of 3.0% (96.20 ± 0.02 mol % CH<sub>4</sub> and  $2.88 \pm 0.00$  mol % C<sub>2</sub>H<sub>6</sub>, GNH personal communication) for the gas composition in the grid for the period of August and September 2020 in Hamburg. This ratio was reported 3.5% (95.09 mol % CH<sub>4</sub> and 3.37 mol %, GNH personal communication) in April 2020.
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#### 295 **2.3.3 LDC leak detection and confirmation**

296 Since the pipeline locations are known to the LDC, the method can be applied precisely above 297 the pipelines, including visible cracks and cavity outlets in the close vicinity, increasing the 298 possibility of leak detection. Once the carpet method detects a CH<sub>4</sub> source, a second 299 measurement is performed above the location with the highest signal, where air is accumulated 300 and analyzed for the presence of C<sub>2</sub>H<sub>6</sub>. The C<sub>2</sub>H<sub>6</sub> detection in the carpet method is not online with higher detection threshold and in batch mode (gas chromatography), which takes time, 5 301 302 -10 minutes per location. If sufficiently high CH<sub>4</sub> and C<sub>2</sub>H<sub>6</sub> levels are found, the leak is 303 categorized in one of safety categories of A1, A2, B or C. 304

#### 305 2.3.4 Precise underground leak localization

When a leak has been confirmed with the carpet method, a precise localization of the leak is performed by drilling holes about 20-40 cm into the ground along the pipeline track and measuring the sub-surface  $CH_4$  concentration. The location with the maximum sub-surface reading is assigned the most likely leak location where the repair teams open the road and attempt repair of the leak. The final exact leak location is reported after opening ground for the repair reactions. Mostly the locations reported from the carpet method matches the locations 312 reported from the leak repair team, which depends on the transport pathways of emission 313 undersurface and surface coverage.

314

#### 315 **2.4 Emission quantification**

#### 2.4.1 Mobile measurements quantifications 316

After the detection of the target locations, we performed additional transects at these locations 317 318 on different days. We accepted a mobile measurement transect of a leak location for further 319 analysis if (i) the GPS signals of transects were logged correctly along the street track and (ii) 320 at least one of the two instruments, G2301 (for quantification and attribution) and / or G4302 321 (for attribution), were running during the transect and (iii) the transect track included at least 322 one GPS coordinate less than 50 m from the leak location. The start and end point of the 323 accepted transects were determined as the locations where the driving tracks intersected with a 324 circle with radius of 100 m centered at the gas leak location reported by the LDC, or a reported 325 outlet location from the mobile method, for the locations where the LDC did not confirm a 326 leak. The segments between the start and end points were evaluated one by one (See an example 327 in Sect. S.4.1 in SI) to determine various parameters, e.g., the maximum  $CH_4$  enhancements, 328 plume area, driving speed, distance to the actual leak locations, etc. The plume area is the 329 integral of the CH<sub>4</sub> enhancements above background along the driving track from the location 330 where the  $CH_4$  enhancement exceeds > 10 ppb until the location where it falls again below the 331 10 ppb threshold.

332 Gas leak quantification from mobile measurements is based on an empirical equation derived 333 from controlled release experiments reported by von Fischer et al., (2017) and reevaluated in 334 Weller et al., (2019) (Eq. 1).

335

 $Q = \exp((\overline{Ln(C_{max})} + 0.988) / 0.817)$ б Eq. 1 337

338 In Eq. 1,  $C_{max}$  is the maximum CH<sub>4</sub> enhancement (ppm) observed during each transect next to 339 the leak location. The maximum CH<sub>4</sub> enhancement should be more than 10% above CH<sub>4</sub> background level to be considered for the quantification algorithm. The emission rate is 340 denoted by Q and it is in L min<sup>-1</sup>.  $\overline{Ln(C_{max})}$  is the mean of the logarithm of the maximum 341 342 mole fraction enhancements for all accepted transects.

343 The standard quantification method only uses transects where CH<sub>4</sub> enhancements are more 344 than 10% or  $\approx$  200 ppb above background level. This 10% enhancement threshold corresponds to about 0.5 L min<sup>-1</sup> emission rate in Eq. 1. Thus,  $\approx 0.5$  L min<sup>-1</sup> is the minimum emission rate 345 that can be quantified with Eq. 1 and leaks with smaller emission rates are ignored by design 346 of the method. Below we investigate the effect of relaxing the enhancement threshold. The 347 application of the tracer release technique in mobile mode allowed us to use the known  $C_2H_2$ 348 349 release rate and the measured C<sub>2</sub>H<sub>2</sub> plumes to independently validate the mobile approach, 350 including the effect of the enhancement threshold. We also investigated the effect of distance 351 between CH<sub>4</sub> maxima to gas leak locations, which is not a parameter in Eq. 1.

352 The uncertainty of the emission rate for each location in the mobile method was calculated 353 using standard error and t-factor (95% confidence) for the locations with at least three CH4 354 enhancements greater than the 10% threshold.

In addition to evaluating the maximum CH<sub>4</sub> enhancement from each transect we also derived 355 356 the plume area (mixing ratio times distance and in unit of ppm m) for comparison between the 357 instruments. In principle, the plume area should provide a more robust quantification of an 358 ambient CH<sub>4</sub> plume than the maximum enhancement: When a plume spreads out, individual 359 realizations of the plume can be sharper and higher, or wider and lower, depending on meteorological conditions, but the plume area should be less affected. In addition, when an 360 361 instantaneous plume is sampled with two instruments with different gas flow rates, instruments with a lower flow rate will be affected by mixing of air in the measurement cell. This will lead
to a lower maximum enhancement but a wider peak, and thus the peak area should lead to a
better comparison between the instruments.

#### 366 **2.4.2 Tracer measurements quantifications**

The tracer method uses Eq. 2a to quantify CH<sub>4</sub> emissions in mobile mode (integral over space dimension) and Eq. 2b in the static mode (integral over time dimension). Parameters relevant for the evaluation with the tracer method are provided in Sect. S.4.2.

370 
$$Q_{CH_4} = Q_{C_2H_2} \cdot \frac{\int_{start}^{end} C_{CH_4} dx}{\int_{start}^{end} C_{C_2H_2} dx} \cdot \frac{MW_{CH_4}}{MW_{C_2H_2}}$$
Eq. 2a

371 
$$Q_{CH_4} = Q_{C_2H_2} \cdot \frac{\int_{start}^{c_{H_4}} C_{CH_4} dt}{\int_{start}^{end} C_{C_2H_2} dt} \cdot \frac{MW_{CH_4}}{MW_{C_2H_2}}$$
 Eq. 2b

Here C is the mole fraction (ppm) and MW is the molecular weight of the species, 16 g mol<sup>-1</sup> 373 for CH<sub>4</sub> and 26 g mol<sup>-1</sup> for C<sub>2</sub>H<sub>2</sub>.  $Q_{CH_4}$  is the CH<sub>4</sub> emission rate estimate for CH<sub>4</sub> (g s<sup>-1</sup>) and  $Q_{C_2H_2}$  is the controlled release rate of C<sub>2</sub>H<sub>2</sub> (g s<sup>-1</sup>). The C<sub>2</sub>H<sub>2</sub> flow rate was controlled and 374 375 measured with a flow controller (Brooks Sho-Rate). In addition, the mass of C<sub>2</sub>H<sub>2</sub> released at 376 377 each location was measured by weighing the release cylinder before and after the tracer release 378 with a precise scale (KERN DE60K5A). The change in mass was then converted to a mass flow rate using the release time. To convert the emission rate from mass ( $g s^{-1}$ ) to volume (L 379 380 min<sup>-1</sup>) we used normal temperature and pressure (NTP) conditions, T = 293.15 K, p = 1.01325381 bar. The locations of tracer release (C<sub>2</sub>H<sub>2</sub>) at the confirmed gas locations were determined with 382 the combined information from the mobile and the carpet methods.

383 The tracer gas can also be used to pinpoint and confirm the emission source location. Prior to 384 quantification, it is important that the emission outlet is located for proper tracer release (see 385 Fig. 1) and source simulation and that other potential interfering emission sources can be ruled 386 out. This is secured by performance of upwind and downwind CH<sub>4</sub> mole fraction screening. During transecting of the CH<sub>4</sub> and tracer plumes, the two plumes should match, if this is not 387 388 the case, the tracer release should be relocated until a proper plume match is obtained. If an 389 emission source consists of multiple outlets, the combined emission from all outlets can be 390 measured by releasing the tracer at the main outlet and increasing the measuring distance until 391 one confined overlapping plume of CH<sub>4</sub> and tracer gas is obtained. If the distance cannot be 392 increased to access limitations, tracer should be released at each single emission outlet. 393

#### 394 **2.4.3 Suction measurements quantifications**

The quantification of a leak with suction method is possible after pumping accumulated air out of soil and reaching CH<sub>4</sub> mole fraction equilibrium in the outflow. With the equilibrium CH<sub>4</sub> reached and the known pumping rate through the probes, it is then possible to calculate emission rate (See Sect. S.4.3 in SI).

399

365

#### 400 **2.4.4 Hole method, based on leak and pipeline properties**

The LDC reported the physical properties of gas leaks and pipeline conditions. These include leak area, pipeline diameter and pipeline operational pressure. In order to get an estimate of the upper physical limits of gas leakage through a hole with the given properties, we used the empirical model by Liu et al., (2021), which was designed to quantify emissions from buried natural gas pipelines to estimate emission rates from the leaks (Eq. 3), hereinafter "hole" 406 method.

407

- 408  $Q = 0.567 \cdot [(h + 139.592)^{-0.1} 0.542] \cdot d^{1.5} \cdot p^{0.7}$  Eq. 3
- 409

Here, Q is the gas leak rate in m<sup>3</sup> h<sup>-1</sup> (at standard atmospheric conditions and converted to 410 NTP), h is the depth of the buried pipeline in cm, d is the gas leak hole diameter in mm and p 411 412 is the pipeline overpressure in kPa. We used 150 cm as pipeline depth for all the locations in 413 Hamburg to estimate emission rate. We note that the model that we employed is for buried 414 pipelines not pipelines in open space, and emission estimates for the gas leak emission rate in 415 open space would be even higher (See Sect. 4.4 in SI). Ebrahimi-Moghadam et al. (2018) 416 showed that CH<sub>4</sub> emission from a pipeline hole area can be between 7 to 10 times higher in 417 open space relative to the subsurface conditions.

418

## **3 Results**

419 420

#### 421 **3.1 Leak Detection**

422 15 possible leak locations were detected by the mobile method in the initial surveys, (labeled 423 as HH001 – HH015). At 13 out of these 15 locations, leaks were confirmed by the LDC, HH007 424 and HH012 locations were not confirmed as gas leak locations. In addition, the LDC identified 425 5 other leak locations (labeled as HH100 – HH104) that had not yet been fixed (category B and C). The overview of the measurements (detection and quantification) is provided in the SI (See 426 427 Sect. S.5 in SI). At some locations we also observed that vegetation was impacted negatively 428 by the presence of leaks in their vicinities, a known phenomenon as high levels of methane cause harmful anoxic conditions for the plant roots (See Sect. S.6 in SI). At several locations 429 430 the outlet identification was straightforward, because we only observed one outlet, but at 5 431 locations we observed numerous outlets spread over a large area. Figure 2 shows the spread of emission outlets at one of the locations (Fig. 2a), with correlations of  $CH_4$  and  $C_2H_6$  at the 432 433 "strongest" outlet (Fig 2b). Fig. 2c shows precise gas leak location practice of the LDC at one 434 of the other locations.



435

Figure 2 – (a) aerial image of location HH004 (© Google Maps). Yellow pins show surface
 emission outlet locations, and the red point shows the actual pipeline leak location

reported by the LDC; (b) correlation between CH4 and C<sub>2</sub>H<sub>6</sub> measured from a
telecommunication cover; (c) Map (not to scale) of drilled holes (blue dots) to locate the
pipeline gas leak at HH008. The red star shows the actual pipeline gas leak location as
indicated by the undersurface CH4 mole fractions (See Sect. S.3, Fig. S3)

442

#### 443 **3.2 Leak Quantification**

Table 1 shows the results of the leak emission rate quantifications from the four methods. All these locations were quantified by the mobile method, although for 6 of them the 10% enhancement threshold was not reached. 16 locations were quantified by the tracer release method and 8 by the suction method. A complete overview of key parameters for all measurements (detection and quantification) is provided in Sect. S.5.

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	ID	Leak quantification methods (L min <sup>-1</sup> )							Info. from the LDC			
		Mobile (measurements from G2301)				Suction		(	Pip	Lea	Leak cor	Pipe I
		Transect (s) w/ CH <sub>4</sub> Enh. > 10% threshold	Emission average	Emission range; 95% confidence	Tracer (L min <sup>-1</sup> )	Emission (L min <sup>-1</sup> )	Status	Hole (L min <sup>-1</sup> )	eline buried year	k size (cm <sup>2</sup> )	type; Safety nsiderations	line Size and Material <sup>#</sup>
Detected by mobile method	HH001	n = 1 (10%)	0.7	-	0.06	<1.8	INC	39	1935	2.5	С	DN80ST
	HH002	n = 5 (50%)	4.9	0.7 - 36.0	0.22	< 0.7	INC	45	1935	3.0	A2	DN80ST
	HH003	n = 6 (86%)	7.5	1.1 - 53.0	1.37	-	-	-	1963	-	A1	DN100ST
	HH004	n = 4 (100%)	7.8	1.8 - 34.5	5.33	-	-	-	1959	-	A1	DN80ST
	HH005+	n = 19 (51%)	1.8	0.9 – 3.6	0.21	-	-	-	1935	-	A2	DN80ST
	$\rm HH006^{*}$	n = 11 (39%)	1.2	0.8 - 1.8	0.02	0.3	CPLT	33	1934	0.5	В	DN80ST
	$\rm HH007^{\circ}$	n = 0 (0%)	-	-	-	-	-	-	-	-	-	-
	HH008	n = 6 (26%)	1.5	0.4 - 6.4	0.32	<1.3	INC	-	1934	-	С	DN80ST
	HH009×	n = 9 (38%)	3.9	1.5 – 9.8	4.86	<3	INC	-	1928	-	A1	DN80ST
	HH010	n = 3 (38%)	1.6	0.2 - 13.7	0.51	< 0.7	INC	-	1937	-	С	DN200ST
	HH011^×	n = 4 (50%)	1.9	0.2 - 18.6	0.37	-		150	1963	15	A1	DN300ST
	HH012°	n = 0 (0%)	-	-	_	-	-	-	-	-	-	-
	HH013^	n = 2 (40%)	1.8	-	-	-		65	1939	5	A1	DN80ST
	HH014	n = 24 (55%)	1.6	1.1 – 2.5	1.41	-	-	65	1950	5	A1	DN100ST
	HH015	n = 1 (50%)	1.0	-	0.38	< 0.9	INC	19	1935	1	A1	DN80ST
Reported by the LDC	HH100	n = 1 (13%)	0.7	-	0.14	-	-	-	1994	-	С	d225Pe
	HH101	n = 0 (0%)	-	-	0.07	< 0.7	INC	-	1960	-	С	DN80ST
	HH102	n = 0 (0%)	-	-	0.01	-	-	-	1928	-	С	DN125ST
	HH103	n = 0 (0%)	-	-	0.03	-	-	-	1963	-	В	DN150ST
	HH104	n = 0 (0%)	-	-	-	-	-	-	1930	-	С	DN100ST
451 <sup>+</sup> The LDC reported three leak locations, $\approx 30$ m distance between the two ends, for this												nis
	452 location: two leaks with area of 5 cm <sup>2</sup> and one leak with area of 1 cm <sup>2</sup>											

450 Table 1 – Results of gas leak quantification with different methods in Hamburg, Germany

location: two leaks with area of 5 cm<sup>2</sup> and one leak with area of 1 cm<sup>2</sup> \* Complete measurements for the suction method and used for averaging

^ Leak size reported as sum of total hole area of all the leaks on the pipeline

<sup>×</sup> Large difference between leak location and the tracer release location

<sup>°</sup> The LDC did not confirm a gas leak

457 458 <sup>#</sup>Pipeline materials, steel (ST) or Polyethylene (Pe), pipeline Diameter Nominal (DN), which is close to the inner pipeline diameter in mm

459

#### 460 **3.2.1 Mobile method**

461 The mobile method was applied at all the 20 locations (18 confirmed and 2 unconfirmed gas leak locations). At 14 (all confirmed gas leak locations) out of the 20 locations, CH<sub>4</sub> 462 463 enhancements above the 10% threshold were observed and could be evaluated with the 464 standard algorithm. The emission rate estimates for these 14 gas leak locations ranged from 0.7 465 to 7.8 L min<sup>-1</sup>. At the 6 other locations we didn't observe any CH<sub>4</sub> enhancements above the 10% threshold. When we lowered the enhancement threshold to 10 ppb, the emission rates 466 467 were 0.07 (HH007; not confirmed gas leak location), 0.1 (HH012; not confirmed gas leak location), 0.04 (HH101), 0.02 (HH102), 0.05 (HH103) and 0.02 L min<sup>-1</sup> (HH104). Of the 5 468 469 leak locations reported by the LDC, 4 did not show any enhancement maximum above the 10% 470 threshold, i.e., these locations would not have been identified with the default algorithm (Weller et al., 2018) and would thus not produce an emission estimate. 471

472 Fig. 2 shows a summary of all individual observed enhancement maxima with the G2301 473 analyzer from all transects with the mobile vehicle, which were used for the quantification of emission rates with Eq. 1. The figure illustrates the large spread in enhancement maxima for 474 475 multiple passes at each location, similar to Luetschwager et al (2019), leading to large 476 uncertainties in emission estimates of individual locations. Fig. 2 also shows the diversity of 477 the various locations, where at some locations most or all of the observed enhancement maxima 478 are above the 10% threshold (e.g. HH003 and HH004), at several locations none of the 479 enhancement maxima was above the threshold (e.g. HH101 and HH104) and at other locations 480 many transects showed enhancement maxima both above and below the threshold (e.g. HH006, 481 HH008, HH009, HH014).

482 As shown in Fig. 3, there is a wide range of CH<sub>4</sub> enhancement observations per location. This 483 depends on wind conditions, distance of the observed plume maximum to the emission outlet 484 location, the superposition of emissions from several outlets and likely other variables such as 485 soil water content. The mean relative uncertainty from the mean emission rate values for the 486 mobile method is  $\approx$  70% for lower and 400% fort the upper ends for the locations with at least 3 transects (n = 10) which pass the 10% enhancement threshold (significant signals) in this 487 study. The lower and upper ranges go down to 60% and 275% for the locations with at least 5 488 489 transects (n = 7) with significant CH<sub>4</sub> enhancements.



490

Figure 3 – CH<sub>4</sub> enhancement maxima from all individual transects for each location using
G2301. Red points show CH<sub>4</sub> enhancement maxima below the 10% threshold, green

493 points show CH<sub>4</sub> enhancement maxima above the 10% threshold, thus used for the 494 standard quantification. Blue circles show the  $\overline{\text{Ln}(C_{max})}$  of all the green points for each 495 location, and black triangles show the derived mean emission rate (based on all green 496 points) using Eq. 1 for the location with at least one green point (right y-axis).

## 497498 **3.2.2 Tracer method**

499 The tracer method performed emission rate quantification at 16 gas locations out of 20 locations. The derived emission rates range from 0.03 to 5.3 L min<sup>-1</sup> (Table 1). For 4 locations 500 the tracer method was not applied because (i) the emissions were not persistently observable 501 502 and the LDC also didn't confirm existence of gas leaks at these locations (n = 2; HH007 and 503 HH012) or (ii) the leak had already been repaired (n = 1; HH013) or (iii) no emission was 504 detectable during the visit of the tracer team (n = 1; HH104). For two of the locations (HH11 505 and HH09), where leaks were confirmed and the tracer method was successfully deployed, 506 later investigations during repair actions (see Fig. 1) showed that the surface emission outlets 507 were located far (15 to 60 m) from the actual gas pipeline leak location indicating underground 508 gas migration. It is evident from Table 2 that the tracer technique can also quantify very small emission rates, below the cut-off of the mobile technique of 0.5 L min<sup>-1</sup>. Emission rate 509 510 estimates derived from the tracer technique were in general lower than the ones derived from 511 the mobile technique, except for three sites where they were comparable (HH004, HH009 and 512 H014).

513

### 514 **3.2.3 Suction method**

Due to the time-consuming nature of the suction measurements, initially 10 gas leak locations 515 516 had been planned for deployment of the suction method in this campaign. The goal was to 517 cover a wide range of expected emission rates, as stated in the intercomparison matrix. The 518 suction method was applied at 8 gas leak locations (see Table 1) out of which the suction 519 quantification was complete (HH006) according to protocol where an equilibrium 520 concentration has to be reached. This was at HH006, with a derived emission rate of 0.3 L min<sup>-</sup> 521 <sup>1</sup>. At several of the locations where the mobile method had indicated high emission rates, 522 subsurface accumulation was widespread, and the suction method was either not deployed (n 523 = 3; HH003, HH04, HH011) or the measurements were incomplete (n = 7; HH001, HH002, 524 HH008, HH009, HH010, HH015 and HH101) because of either safety reasons or because the 525 suction team estimated that they would be unable to complete the measurements within a day. 526 For the 7 locations with incomplete suction measurements, the emission rates were reported 527 ranging from 0.7 to 3 L min<sup>-1</sup>. These can be regarded as upper limit estimates because suction 528 was not yet completed and CH<sub>4</sub> concentrations would have supposedly dropped further.

## 529530 **3.2.4 Hole method**

For 5 locations where the leak area of a single gas pipeline leak was reported, the corresponding emission rates are between 19 to 65 L min<sup>-1</sup>. For locations HH011 and HH013, the hole area was reported as the sum of several holes and the total hole area for these two locations resulted in an emission rate of 150 and 65 L min<sup>-1</sup>, respectively. The quantification from the hole method is higher than from the mobile, tracer and suction methods by at least an order of magnitude.

536

### 537 **3.3 Leak categories**

The 20 (18 confirmed + 2 not confirmed) locations can be divided into four main categories related to measurement challenges of the various methods. These categories may overlap.

- 540 (i) Large subsurface CH<sub>4</sub> accumulation
- 541 (ii) Insufficient CH<sub>4</sub> enhancements for mobile quantification
- 542 (iii) Large CH<sub>4</sub> enhancement variability for mobile quantification

543 (iv) Several outlets and / or leaks or atmospheric turbulence

In this section we present the overall results and discuss in detail one selected location for each
of these categories. The remaining locations (with similar characteristics) are presented in the
SI.

547

#### 548 3.3.1 Location type I – Large subsurface CH4 accumulation and multiple outlets

The spatial spread of surface emission outlet locations identified with the G4302 instrument as part of the mobile method provides an indicator for the extent of the subsurface accumulation of CH<sub>4</sub>. For 5 locations, emission outlets were found at great distance from each other, in order of tens of meters. The total emission of a gas leak is equal to the sum of emissions from all the surface outlets at a location, thus it is necessary to quantify each outlet separately to get the total emission.

- 555 HH011 (Fig. 4) is an example where very widespread CH<sub>4</sub> accumulation and migration was 556 observed. During the initial mobile gas leak detection, leaks were located at the intersection of
- 557 streets 1 and 2, close to a subsurface vent and a rain drain,  $\approx 2$  m far apart, (the yellow pin in 558 Fig. 4a) based on clear signals from these outlets and a sign next to the road indicating presence
- of gas pipelines. The vent showed a  $C_2:C_1$  ratio of 2% ( $\mathbb{R}^2$  of 0.8 and max CH<sub>4</sub> mole fraction
- of 31 ppm) and we observed C<sub>2</sub>:C<sub>1</sub> ratio of 2.8% with  $R^2$  of 0.96 and max CH<sub>4</sub> mole fraction
- 561 of  $\approx$  70 ppm from the rain drain, clearly indicating a large / dominant contribution from fossil
- 562 CH<sub>4</sub>. However, after quantifying the emission from these two leaks using the mobile and the
- tracer release methods, the LDC found the actual gas pipeline leak, during the repair actions,on the south side of the intersection, far from the vent and the rain drain, at the intersection of
- 565 street no. 3 and no. 2 indicating that the gas had travelled about 60 m underground. It is possible 566 that the leak resulted in several gas emission outlets, likely closer to the gas pipeline leak
- 567 location. The emission rate measured using the mobile method was  $1.6 \text{ L} \text{ min}^{-1}$  based on 5
- 568 plume transects and is likely underestimated because some emission outlets potentially were
- 569 not included in the performed plume transect. It should also be noted that the distance from the
- 570 gas pipeline leak location to the plume transect is larger than the distances applied during the 571 controlled release calibrations (average 15 m) (Weller et al., 2019).
- 572 The tracer was released at the vent and the rain drain and thus measured the combined emission
- 573 from these two outlets to be  $0.4 \text{ L} \text{ min}^{-1}$ . If the gas pipeline leak gave rise to multiple
- 574 unidentified surface emission outlets, the emission from the gas pipeline is underestimated. IN
- fact, Fig. 4b shows that a  $CH_4$  plume without  $C_2H_2$  was observed during the tracer release measurements at HH011, confirming that at least one other source of methane emission was
- 577 present nearby.
- 578 Based on the previous experience at locations with widespread subsurface accumulation it was
- 579 concluded that the suction method could not be applied at this location. The other case in this
- 580 category was HH009.



581 Figure 4 – aerial image of HH011 (© Google Maps). A gas leak location with widespread 582 583 undersurface CH4 accumulation. The yellow pin shows the assumed leak location and location of tracer release, which was very different from the actual leak location as 584 585 identified by the LDC (red circle). St. 1-4 are added to identify streets that are discussed 586 in the text. General wind direction during tracer release deployment is shown with an orange arrow. CH<sub>4</sub> (in blue) and C<sub>2</sub>H<sub>2</sub> (in red) levels measured at a plume transect. One 587 588 of the CH<sub>4</sub> plume is proportional to the C<sub>2</sub>H<sub>2</sub> plume while the other CH<sub>4</sub> plume lacks the 589 C<sub>2</sub>H<sub>2</sub> signals suggesting existence of at least another emission outlet.

590

591 The LDC reported the total area of several holes in the pipeline as  $15 \text{ cm}^2$  for HH011, which is 592 the largest leak size among all the locations. If we assume that there was one hole with this 593 size, then the emission rate estimated by Eq. 3 will be 150 L min<sup>-1</sup>, a hole of 5 cm<sup>2</sup> gives 594 emission rate of 65 L min<sup>-1</sup>. The pipeline for this location was DN300ST and has been in 595 operation since 1963.

596

#### 597 **3.3.2 Location type II – Insufficient CH4 enhancements for mobile quantification**

598 At HH101, on a narrow ( $\approx$  3 m wide) street, which had about 1 m wide bare soil pavement on 599 one side, the LDC reported a gas leak location based on their routine surveys. On both sides of the street there were about  $\approx 1.5$  m tall bushes and some trees. All three methods (mobile, tracer 600 601 and suction method) were deployed at this location. Gas emissions found their way to the atmosphere through cracks in the asphalt with  $C_2:C_1$  ratio of 2.5% ( $R^2$  of 0.93) with max CH<sub>4</sub> 602 603 mole fraction of  $\approx 25$  ppm. None of the CH<sub>4</sub> enhancement maxima observed during the mobile 604 surveys at this location were above the 10% enhancement threshold with the G2301 instrument, 605 thus this location would not be labeled as LI and no quantification would be reported from 606 mobile method as implemented in Weller et al (2019) and Maazallahi et al. (2020). The tracer 607 method was applied in static mode at a distance of  $\approx 15$  m and reported an emission rate of 0.1 L min<sup>-1</sup>, which is compatible with the emission strength being below the "detection limit" 608 609 defined by the 10% cut-off of the standard algorithm (0.5 L min<sup>-1</sup>). When the emission strength is evaluated using the CH<sub>4</sub> enhancements below the cut-off, the value is 0.04 L min<sup>-1</sup>. The 610

- 611 suction method was applied at this location but an equilibrium was not achieved after 9 hr, i.e.
- incomplete suction measurements, and an upper limit for the emission rate of  $\approx 0.7$  L min<sup>-1</sup> was 612
- reported. The fact that the suction measurement was incomplete at this location with a small 613
- 614 emission rate shows that subsurface accumulation can also be large for smaller leaks.
- Three of the leak locations in this study only showed one CH<sub>4</sub> enhancement above threshold. 615
- The 10% threshold is a constraint, which removes enhancements less than about 200 ppb. This 616
- 617 means for the locations where we only have one transect with CH<sub>4</sub> enhancements more than the 10% threshold, the minimum emission rate estimated is about 0.5 L min<sup>-1</sup>, no matter how 618
- many transects we had with CH<sub>4</sub> enhancements less than the 10% threshold. This situation was 619
- 620 observed for HH001, HH015 and HH100 (Fig. 5). In this case, the mobile method likely
- 621 overestimates the total leak rate, because only the maximum enhancement is used for 622 quantification. The tracer method reported low emission rates for these three sites 0.12 L min<sup>-</sup>
- 623 <sup>1</sup> on average (n = 6).
- For the two locations (HH007 and HH012) where the LDC didn't confirm gas leaks (despite 624
- periodic observation of C<sub>2</sub>H<sub>6</sub> at outlets during the mobile surveys) none of the transects showed 625 CH<sub>4</sub> enhancement maxima above the 10% threshold. At HH007, the outlet was through cracks
- 626
- 627 in the pavement but at HH012 the outlets were from manholes. At HH007 the outlet location 628 had shifted by about 2 m for two different days (4-week gap). We note that the correlation
- coefficients between  $CH_4$  and  $C_2H_6$  at these locations were between 0.4 and 0.6, so less than 629
- 630 0.7, which is the threshold correlation we accepted for the outlets. As a leak was not confirmed
- 631 for these locations, the tracer and suction methods were not applied.
- 632

#### 633 3.3.3 Location type III – Large CH<sub>4</sub> enhancement variability for mobile quantification

- For several locations, we observed a large variability of CH<sub>4</sub> enhancements from different 634 transects. One example is HH008, where only 6 of the 23 transects exceeded the 10% threshold, 635 i.e. the leak was only observed in about every 4<sup>th</sup> transect. The leak location of HH008 is an 636
- example where CH<sub>4</sub> enhancements from several transects cover a wide range. Based on the 6 637 transects, which showed enhancement maxima above the 10% threshold, a leak rate of 1.5 L 638 639 min<sup>-1</sup> is derived. This may be an overestimate since many transects with maxima below the 640 threshold were not considered. For this location the mobile tracer method was applied, which 641 resulted in a leak rate quantification of 0.3 L min<sup>-1</sup>.
- The suction method derived an upper emission estimate of 1.3 L min<sup>-1</sup> from incomplete 642 measurements at HH008. The LDC reported a C category leak for this location from a DN80ST 643 644 pipeline, which was installed in 1934.
- 645

#### 646 3.3.4 Location type IV – Several outlets and / or leaks or atmospheric turbulence

647 On a  $\approx$  5 m wide street, we detected two leaks about 80 m away from each other, HH001 and 648 HH002 (Fig. 5a). It was a cobblestone street and there were bushes and few trees planted, 649 mostly on one side of the street. The mobile method performed 10 transects at both locations 650 and all the transects were accepted for the evaluation. The tracer team could quantify both locations using static measurements. The suction team began to quantify HH002 and HH001, 651 652 but during quantification of HH001, there was a small accident (fire due to contact of drilling head with electric cable) and the leak had to be fixed immediately. The plumes on this street 653 were sufficiently separated to positively identify two different leaks on the same street. In 654 655 contrast, at location HH005, we observed several maxima for the same transect, but because 656 the maxima were close to each other, those were clustered together in the mobile measurement 657 algorithm (Fig. 5b). Later the LDC reported even three individual pipeline leaks on this street. 658 In another example (HH010), some transects showed several plume maxima although only one 659 emission outlet and later on only one gas pipeline leak was found (Fig. 5c). However, the 660 release of the tracer resulted in several matching CH<sub>4</sub> and tracer gas plumes confirming that 661 the emission indeed occurred form a single outlet and that the multiple plumes at this location 662 were due to inhomogeneous plume dispersion. This illustrates that the existence of several 663 maxima in one transect does not necessary correspond to presence of several leaks and/or 664 outlets, but it can also be related to a spatially heterogeneous/disturbed plume. This shows that 665 the signals from the mobile detection method are not sufficient to allow determining the 666 number of leaks at a location with several plume at a close distance from each other in a single 667 transect.

668



669

Figure 5 - several maxima observed during a single transect on one street showing different situations: two well isolated leaks with about 80 m distance from each other (a1 and a2, HH001 and HH002), three pipeline leaks close to each other with several emission outlets (b1 and b2, HH005) and one leak and one outlet but several CH4 enhancement maxima due to turbulence (c1 and c2, HH010), aerial images: © Google Maps.

675

After detection by mobile measurements, emissions out of the ground were detected at HH001 676 and HH002 with the G4302 backpack within 3 m distance from the gas pipeline leak locations, 677 678 which was later reported by the LDC. For the single transect with a maximum above the 10% 679 threshold observed with the mobile method, the derived emission rate at HH001 was 0.8 L min<sup>-</sup> 680  $^{1}$  (n = 1). For HH002, the derived emission estimate for the transects with maxima above the threshold is 5.2 L min<sup>-1</sup> (n = 5) from the mobile method. At HH002, individual derivation of 681 emission from separate CH<sub>4</sub> enhancement gives a wide range between 0.7 and 36.0 L min<sup>-1</sup> 682 683 (95% confidence) from the mobile method (see category III above). For HH001, the tracer method was applied in static mode at  $\approx 30$  m distance to the release point and  $\approx 40$  m far from 684 HH002. The derived emission rate for HH001 is 0.06 L min<sup>-1</sup> and for HH002 0.22 L min<sup>-1</sup> from 685 the tracer method. For HH001, after about 5 hr of pumping, the suction quantification had to 686 be stopped due to the incident described above. Based on the incomplete suction measurement 687 an upper limit for emission rate of  $\approx 1.8$  L min<sup>-1</sup> for HH01 was estimated. An emission estimate 688 689 of  $\approx 0.7$  L min<sup>-1</sup> was derived for HH002 from an incomplete suction measurement. The LDC

690 reported leak size of  $\approx 2.5 \text{ cm}^2$  for HH001 and for  $\approx 3 \text{ cm}^2$  for HH002 which then give emission 691 rate of 39 and 45 L min<sup>-1</sup> respectively from the hole method. For both locations, leaks were due 692 to pipeline corrosion.

693

#### 694 **3.4 Emission rates of different leak safety types**

The 18 confirmed gas leak locations that were investigated in the campaign were categorized 695 696 into the four safety categories, A1 (n = 7), A2 (n = 2), B (n = 2) and C (n = 7). The mobile 697 method quantified all the A1 and A2 leaks (n = 9) with an average emission rate of 3.6 L min<sup>-</sup> <sup>1</sup>. 5 out of 9 leaks in categories of B and C leaks were quantified with the mobile technique 698 699 including the 10% threshold with average emission rate of 1.1 L min<sup>-1</sup> (n = 5). Apart from one 700 location, which had to be fixed before the measurements, the tracer method quantified the A1 701 and A2 leaks (n = 8) and reported an average emission rate of 1.8 L min<sup>-1</sup>. The tracer method 702 also quantified all the B and C leaks (n = 9) with an average emission rate of 0.1 L min<sup>-1</sup>. 703 Mostly due to the safety and time constraints and medium to large underground accumulations 704 of CH<sub>4</sub>, the suction method could provide incomplete measurements at only 3 locations of A1 705 and A2 leaks with an average emission rate of  $1.5 \text{ Lmin}^{-1}$  (n = 3). The suction method measured at 5 out of 9 B and C locations, one of the measurements was complete and the others were 706 incomplete, with an average emission rate of 1.0 L min<sup>-1</sup> (n = 5). Although the number of 707 quantified leaks is limited, all the three methods show that the emission rates from category A1 708 709 and A2 leaks are higher than category B and C leaks (Fig. 6). This indicates that the site 710 selection bias of measurements for the suction method due to safety concerns (see qualifier 711 above), can lead to a bias in the emission rate in this method.





### **Figure 6 – Emission rate differences between different gas leak categories**

### 714 **4 Discussion**

715 **4.1 Leak detection methods** 

#### 716 **4.1.1 Leak location vs outlet location**

There is a difference between the location of the leak in the gas pipeline (leak location; See Sect. S.7 in SI) and the location where the gas is emitted to the atmosphere (outlet locations; See Sect. S.2 in SI). Furthermore, a single leak in the gas pipeline can result in multiple emission outlets at the surface. In this campaign we observed that in most cases (2 out of 18), the emission outlet at the surface occurred only a few m (sometimes < 1 m) from the location

of the leak in the gas pipeline. However, in one case, an emission outlet was detected about 60

m away from the leak location indicating significant underground gas accumulation andmigration (see Fig. 4).

725

#### 726 **4.1.2** Intercomparison of the gas leak detection methods

727 The mobile method detects atmospheric CH<sub>4</sub> enhancements while measuring continuously with 728 ppb precision from an inlet installed at the front bumper of the car while LDCs apply the carpet 729 method with an instrument precision at the ppm level. High precision for the carpet method is 730 not needed as the inlet to their instruments is connected to a carpet, which is attached to the 731 ground. The mobile method can cover larger areas in shorter times, but not all roads, walkways, 732 or other surface areas where pipelines are buried are accessible with a vehicle. The advantage 733 of the carpet method is that it can precisely follow the pipeline map, which also means that it 734 can locate leaks more precisely. The mobile method use a 10% threshold to neglect unreliable 735 gas leak sources, which sometimes results in neglecting actual signals from small leaks. Also 736 the mobile measurements do not detect all leaks due to the dependence on the wind direction 737 (only downwind sources leaks can be detected). Luetschwager et al. (2021) suggested that 5 to 738 8 plume transects give > 90% probability of gas leak detection at a given location, so if all the 739 streets in an urban area are covered 5 to 8 times, > 90% of the leaks can be detected by mobile 740 measurements.

Both the mobile and the carpet method use  $C_2H_6$  signals for distinguishing between fossil and microbial CH<sub>4</sub> emissions, and as for  $C_2H_6$ , the instrument used in the mobile method is more sensitive, and faster. In the carpet method, the laboratory analysis of  $C_2H_6$  is slow and with higher detection threshold compared to the mobile method, where  $C_2H_6$  is measured in realtime during the surveys, and also on foot from the emission outlet. The CRDS instrument provides real-time measurements of CH<sub>4</sub> and  $C_2H_6$  at 1 Hz frequency so checking various outlets at a possible gas leak location is faster.

- At 14 out of the 20 locations in this study, gas leaks were detected (CH<sub>4</sub> signals passing the 748 749 10% threshold) and quantified with the mobile method. However, we observed that 4 out of 5 750 locations reported by the LDC would not have been detected in mobile surveys without prior 751 information on existence of the leaks because the maximum enhancement was below the 752 mobile detection threshold. At the only location (HH100) from the list of the LDC, where 753 mobile method could quantify the emissions, the outlets were located on the road and the vehicle was driving on top of the outlet. For this location only one of the transects passed the 754 10% enhancement threshold, and the quantification for this location was  $\approx 0.7 \text{ L min}^{-1}$ , close 755 to the detection threshold of this method,  $\approx 0.5$  L min<sup>-1</sup>. One of the other locations, HH101, 756 757 reported by the LDC had similar surrounding conditions (e.g. presence of buildings, road 758 conditions, etc.) as the other leaks detected by the mobile method, but still the mobile method 759 was not able to detect a gas leak at this location without a priori information from the utility. 760 The quantifications made by the tracer method suggest that the emission rates of the locations 761 provided by the LDC were much lower than the locations detected by mobile measurements 762 (Table 2). The 10% threshold in the mobile method precludes the identification of small leaks 763  $(< 0.5 \text{ Lmin}^{-1})$ , which would only be identified by the carpet method.
- 764

### 765 **4.2 Signal attribution in mobile detection method**

#### 766 4.2.1 Attribution during mobile survey in car

During the mobile measurements we used two approaches to find correlation between  $CH_4$  and C<sub>2</sub>H<sub>6</sub>. When we compare the online measurements point by point, the probability of detecting a fossil signal is high, as only one single significant reading is sufficient to indicate a fossil signal. When we use the R<sup>2</sup> of the linear correlation between  $CH_4$  and  $C_2H_6$  enhancements above the cut-off, the attribution is more reliable. In a large dataset without a priori information on the existence of a gas leak at different locations, the correlation method is more trustworthy
 as the point-by-point method could be affected by instrument noise and/or spikes.

- We also used CO<sub>2</sub> signals and their correlation with CH<sub>4</sub> signals to investigate interference
- from combustion or microbial processes. For only 7 plumes at 6 locations, we detected
- correlations between  $CO_2$  and  $CH_4$ , which could indicate either oxidation of  $CH_4$  to  $CO_2$  or
- mixture of microbial  $CH_4$  emissions from e.g. the sewer system with the emissions from natural gas leaks. The number of these possible co-emissions is low compared to the number of total
- transects (only  $\approx 7\%$  of the plumes with CH<sub>4</sub> enhancements greater than 10%), thus such an
- admixture of microbial  $CH_4$  should not impact the quantification from mobile method significantly.
- 782

#### 783 **4.2.2 Plume attribution to emission outlets**

784 The outlet attribution was performed using the G4302 CRDS instrument which is portable like 785 a backpack. We checked the outlets (See Sect. S.2, Fig. S1) around the locations of interest and 786 evaluated the correlation between CH<sub>4</sub> and C<sub>2</sub>H<sub>6</sub> and the persistence of the emissions on 787 different days. In theory, it is possible to estimate contributions of fossil and microbial CH<sub>4</sub> in 788 a plume using the ethane signals during the mobile measurements with the vehicle and the 789 reference C<sub>2</sub>:C<sub>1</sub> ratio provided by the LDC. However, due to the low C<sub>2</sub>H<sub>6</sub> signals in ambient 790 air, it was not feasible to quantify the possible contribution of microbial methane emissions. 791 Nevertheless, the C<sub>2</sub>H<sub>6</sub> signals of the G4302 CRDS instrument were still very useful to identify 792 a location as a possible gas leak location or not. For all the 15 locations, which were initially 793 detected by the mobile method we observed detectable C<sub>2</sub>H<sub>6</sub> signals, including the two 794 locations which later were not confirmed as a gas leak location by the LDC. This suggests that 795 either the leak is at a greater distance and depending on the transport of the emission we 796 periodically can see the signals at the detected outlets or that there are sources that produce 797 both  $CH_4$  and  $C_2H_6$  in the vicinity of the location.

798

### 799 **4.3 Leak quantification methods**

#### 800 4.3.1 Mobile method

801 If the outlets are close to each other, we may observe several  $CH_4$  enhancements close to each 802 other or overlapping when a single transect is performed at a close distance. If we assume that 803 the number of  $CH_4$  maxima is equivalent to the number of real outlets that exist on a road and 804 only use the maximum enhancements from the most pronounced plume to calculate the 805 emission rate, the total emission will be underestimated with the mobile method.

Emission rate estimates with the mobile method from individual transects are associated with 806 807 high uncertainty, related to variabilities in either above-ground or under-ground conditions. For 808 example, an unfavorable wind direction (above ground condition) can result in missing a plume 809 from a gas leak. The mobile measurement van itself may also affect the measurement, e.g., by 810 creating pressure fluctuations. Luetschwager et al. (2021) showed that the quantifications from 811 the same leak in individual mobile transects can vary by more than an order of magnitude. In Hamburg, we found that the range can be even a factor 50 or 100 in exceptional cases (Table 812 813 2). This high variability illustrates that if we perform only one transect per location, the 814 estimated leak emission rate can result in high under / overestimation in emission estimate for the single location, as was also reported by Maazallahi et al. (2020). This large uncertainty for 815 816 individual locations is less severe when the results are extrapolated to the city-level, where the

817 sample size is also large, including over- and underestimates (Brandt et al., 2016).

In our previous study in Hamburg (Maazallahi et al., 2020) the overall average emission rate for all the LIs was estimated 3.4 L min<sup>-1</sup> LI<sup>-1</sup> (n = 145) while for the fossil-attributed locations it was 5.2 L min<sup>-1</sup> LI<sup>-1</sup> (n = 45; standard error of 3.1). This showed that the biggest

821 emitters were among the fossil categories. In the present study, the average emission rate from

mobile measurements for the gas leak locations is 2.7 L min<sup>-1</sup> LI<sup>-1</sup> (n = 14; standard error of 822 0.6). The higher average emission rate per fossil location in the first campaign may have been 823 caused by the fact that in that campaign only a smaller number of transects were performed per 824 825 location (on average 1.1 in the precious study versus 6.9 transects with  $CH_4 > 10\%$  threshold 826 per location in the present study). Luetschwager et al. (2021) stated that after 6 transects with CH<sub>4</sub> exceeding the 10% threshold per location the average overestimation of leak size estimates 827 828 will be less than 10%. In addition, the differences in sample size and locations in these two 829 studies (45 versus 14 locations in the first and second studies respectively) may partially explain the difference in average. This is because the probability of detecting large emitters, 830 831 which increase the average emission rate of all leaks, increases with sample size.

832 The two CH<sub>4</sub> sensors onboard the mobile van play specific roles in the detection and 833 quantification of leaks. CH<sub>4</sub> enhancements on the G2301 are 3.8 times lower than the G4302. 834 This is an artefact of the G2301, which smoothes the signal compared to the G4302 because of the slower pump and sampling rate (See Sect. S.8.1 in SI). On the other hand, this results in 835 836 more signals passing the 10% threshold on G4302. This then also leads to higher detection probabilities using G4302 (See Sect. S.8.2 in SI). Higher record of CH<sub>4</sub> enhancements then 837 838 also results in higher emission rate quantification using Eq. 1 (See Sect. S.8.3 in SI). We use 839 the G2301 for quantification, since this is the instrument that was also used for introduction of the mobile equation quantification in Weller et al. (2017). The quantification of the gas leak 840 841 locations using Eq. 1 depends only on the CH<sub>4</sub> enhancements. This gives about a factor 2 higher 842 emission rates from G4302 than from G2301 for the same plumes. When we evaluate the plume 843 areas from the two instruments, they are much closer to the 1:1 line (See Sect. S.8.3 in SI). This 844 agrees with findings from another study using two different in-situ instruments onboard a 845 mobile car (See Sect. S1.5, Fig. S6 from Ars et al. (2020)). They also found that the plume area 846 is closer to the 1:1 line in mobile measurements even if the air intakes are not at the same 847 location of the vehicle. This suggests that the plume area is a more robust parameter than 848 maximum enhancement for emission rate quantification and a leak rate quantification equation 849 using the plume area should be developed.

850 In general, the closer the air intake is to the emission point the higher the CH<sub>4</sub> mole fraction 851 reading is (See Sect. S.9 in SI), but when several outlets are present at one location it is not 852 possible to uniquely determine the distance to the emission point, and also determine which 853 plume belongs to which outlet. Eq. 1 from Weller et al. (2019) only uses the maximum CH<sub>4</sub> 854 enhancements above the 10% threshold from each pass. In their controlled release experiments the average distance between the leak and measurement was 15.75 m. Analysis of our results 855 (Table S4, Sect S.5 in SI) shows that higher maximum concentrations are encountered more 856 857 often when the distances of the transect to the leak location are small. For example, at HH002 858 the transect was very close to the main emission point, which likely leads to the substantially higher emission rate estimate derived from the mobile method (4.9 L min<sup>-1</sup>) compared to the 859 860 tracer method (0.22 L min<sup>-1</sup>). On the other hand, at HH011 the mobile method underestimates the emission rate (See Sect. 3.3.1), as at this location the measurement distance to the leak was 861 larger than reference distance of 15.75 m applied by Weller et al. (2019). This suggests that to 862 863 reduce the quantification error for individual leak locations, distance should also be included 864 in an improved transfer equation.

The effect of neglecting or retaining the transects with enhancement maxima below the 10% threshold was quantitatively investigated for 5 locations where the tracer team conducted mobile measurements (See Sect. S.10 in SI). These measurements were evaluated as "controlled release" experiments for  $C_2H_2$ , because the actual  $C_2H_2$  release rate is known, and measurements were made in mobile mode. The standard mobile quantification algorithm with the 10% threshold yields emission estimates that are in relatively good agreement with the released quantities, whereas the estimates are biased considerably low when measurements with maxima below the threshold are retained. This supports the use of the original method, which removes transects with an improper realization of the plume. Relating to section 4.5, it must be noted, however that in these measurements the distances of the  $C_2H_2$  maxima to the release points were between 30 to 45 m, thus larger than the normal distance of mobile CH<sub>4</sub> measurement to the emission outlets (from few meters up to 30 m).

877

#### 878 **4.3.2 Tracer method**

879 The tracer method is more labor intensive than the mobile method. However, the strength of 880 the method is the application of a tracer gas providing the plume dilution and avoiding the use 881 of atmospheric dispersion models and weather information. If the tracer release location does 882 not reflect the sum of all the outlet emissions at a gas leak location, or misses some of the 883 outlets, then the total emission quantification from the gas leaks will be underestimated. An 884 example of such a case is site HH011 in this study where the leak location in the gas pipeline (after quantification; see Fig. 1) was found to be located about 60 m upwind the targeted 885 886 emission outlet. During tracer quantification, an additional CH<sub>4</sub> plume (not defined by the 887 tracer gas) was observed indicating more than one emission outlet (Fig. 4). The confirmation 888 for this is the finding of gas leak location by the carpet method. The emission rate of the 889 targeted emission source (the vent and the drain) is thus not representing the combined 890 emission from the gas leak in the pipeline located 60 m upwind the emission source. Further 891 surface screening and leak detection would have been needed to identify and quantify all 892 emission outlets.

893

#### 894 **4.3.3 Suction method**

895 The suction method is the most labor-intensive quantification method. Following a strict, safety 896 first, protocol the gas utilities fix leaks in the A1 safety category immediately upon detection 897 and A2 leaks within a week. Given logistical constraints, the suction method therefore mainly 898 or exclusively quantifies B or C leaks (50% of confirmed gas leak location in this study). We 899 investigated whether such a site selection bias could lead to a bias in the average quantified 900 emission rate in the inventory report. In this study, we observed that the leaks detected from 901 the mobile methods were mostly in the A1 and A2 category and the biggest emitters (based on 902 the mobile and tracer release measurements) had soil CH<sub>4</sub> accumulation of a magnitude that 903 prevented successful application of the suction method. Further research is needed to identify 904 the physical mechanism(s) to explain the observed correlation between A1 and A2 leaks and 905 high emission rates. As a hypothesis, the presence of soil cavities associated with leak category 906 A1 may result in higher permeability, i.e. lower underground resistance, which then leads to 907 higher emission rate for the same pipeline hole size compared to locations with no cavity.

- The suction method was intended to be deployed right before the repair actions. For some of these locations, the suction method was in operation for more than 10 hours, but due to the high soil CH<sub>4</sub> accumulation, the measurements were stopped and labeled as incomplete in this study.
- soil CH<sub>4</sub> accumulation, the measurements were stopped and labeled as incomplete in this study.
   For the other locations with high soil CH<sub>4</sub> accumulation, the suction method was not attempted,
- given the expectation (based on experience at the incomplete locations) that completion of
- 913 measurements for leak rate quantification at those locations was unlikely.
- 914

#### 915 **4.3.4 Hole method**

Based on the leak size, pipeline depth and overpressure, the average emission rate was estimated at 40 L min<sup>-1</sup> (n = 5). We note that these estimated should be regarded as upper limits since flow restrictions outside the pipe are not included. The emission range of individual gas leaks based on the hole method is between 19 to 150 L min<sup>-1</sup> for 1 cm<sup>2</sup> to 15 cm<sup>2</sup> hole sizes respectively, larger than any of the measurement-based quantification methods. This method

921 requires information about the overpressure of the gas pipeline, depth of buried pipeline and

922 size of a leak and it does not include the information about soil properties, which can impact 923 the emission rate.

923 924

#### 925 **4.3.5 Intercomparison of methods**

In this study, a reliable quantitative intercomparison of the three methods (mobile, tracer and suction methods) was attempted. A complete comparison of all three methods was possible at only one out of 20 locations (18 confirmed gas leak locations) because of the long time (>8-10 hrs) needed for full equilibrium of the suction method, whereby emission rates for 7 out of the 8 leaks quantified by the suction method were reported as maxima rather than absolute values (Table 1). At these 7 locations the emission was thus overestimated.

In total, the average CH<sub>4</sub> emissions from natural gas pipeline leaks for the same locations where we have quantifications from mobile and tracer methods (n = 13) are 2.8 and 1.2 L min<sup>-1</sup> respectively. The suction method could only be completed at one location. The average emission rate reported for all the locations from the suction method (high bias due to incomplete measurement) is 1.2 L min<sup>-1</sup> (n = 8).

- 937 The higher emission rates derived with the mobile method are in qualitative agreement with
- 938 previous studies. Weller et al. (2018) compared quantifications from the mobile measurements 939 described in von Fischer et al. (2017) with the tracer method and surface enclosure method in
- four US cities. They reported that mobile measurement estimates were  $\approx 2.3 \text{ L min}^{-1}$  greater
- than the tracer method mean estimates of  $\approx 3.2 \text{ L min}^{-1}$  (n = 59). This was attributed to the overestimation of small leaks (< 2.4 L min<sup>-1</sup>) in the mobile measurements method, which we
- have also discussed above for our dataset. In addition, performance of only a few transects at
- 944 individual locations also lead to systematically high biased emission rate estimates for higher 945 emission rates (Luetschwager et al., 2021). Indeed, at the locations where we only have one
- 946 transects with CH<sub>4</sub> enhancements above the 10% threshold, there is an overestimation from
- 947 mobile method compared to the tracer method. For example, at HH001 (n=1), HH015 (n=1)
- and HH100 (n=1) the mobile method estimated emissions of a factor 4 higher in comparison
   to the tracer method. The analysis of Luetschwager (2019) clearly shows that this high bias is
- 950 reduced when numerous transects are performed. Therefore, we carried out multiple transects
- 951 to reduce this systematic bias. We note that there are also large differences between the mobile 952 and tracer methods, e.g. HH002 and HH006. We suspect that the very short gas leak location
- 953 distance to the mobile driving transects can explain partially the difference. Moreover, 954 existence of another leak in the category of A1 at the HH006 location which had to be fixed
- 955 prior to the tracer method could explain the difference in emission rate magnitude at this
- location. Nevertheless, the limited number of transects and the 10% threshold can contribute
  to an overestimation of the average leak rate with the mobile method at an individual location.
  At the same time, however, the mobile method fails to detect leaks entirely when the leak outlet
- is located downwind of the mobile van. The fact that the mobile method misses downwind
  emissions constitutes a method specific factor towards biasing city-wide emissions low, which
  qualitatively counteracts the high bias above.

#### 962 **4.4** Possible suction method sampling bias with implications for emission inventories

963 Following our communications with the emission inventory experts (personal 964 communications with Christian Böttcher, 2022), we cannot fully reconstruct the methods that are used in the existing national inventory report to establish the emission factors due 965 966 to lack of transparency. However, the German environmental agency (UBA) is considering 967 to use the results of the recent large scale measurement campaign based on the suction 968 method (MEEM, 2018) in future publications of the national emission inventory in Germany 969 (Federal Environment Agency, 2021). The utilities choose leak locations for application of 970 the suction method where there are no safety concerns and/or immediate leak closure is

971 compulsory. This implies that this method is not applied at locations of the A1 category, which

- demand immediate repair (P. 27 in *MEEM*, 2018). Due to logistic constraints and the timeconsuming nature of the suction measurements, they are likely also not (or rarely) applied at
  locations in the A2 category, which require repair within a week. Thus, suction measurements
- have a location sampling bias towards leaks in the B and C category. This is supported by the
- 976 fact that the leak locations that were contributed by the LDC to the intercomparison campaign
- 977 were locations in the B and C category. This study investigated whether this location sampling
- 978 bias could result in an emission rate bias, which could contribute to the fact that the suction
- 979 method did not report leaks with emission rates as high as they have been reported by the 980 mobile method in this study or during previous measurements in the same city (Maazallahi et
- 981 al., 2020).
- 982 In this study, emission rates from A1 and A2 category leaks were larger compared to those
- from B and C category leaks (Figure 6). The emission rate differences vary by measurement
- 984 method: a factor 2 for the mobile method (n = 9 for A1&A2, n = 4 for B&C), a factor 11 for 985 the tracer method (n = 8 for A1&A2, n = 8 for B&C) and a factor 1.6 for the suction method
- 986 (n=3 for A1&A2, n = 5 B&C). For the mobile method, there is a clear separation between the
- A1&A2 versus the B&C categories. The highest emission estimate for the B&C group
  (HH010) is similar to the lowest emission rate estimate for the A1&A2 group (HH014).
  Furthermore, HH011 in the A1 category was very likely biased low because of the wrongly
- 990 assumed leak location.
- 991 For the tracer method, the difference between the two groups is largest, an order of magnitude, 992 and we know that emissions are underestimated at least at one location of the A1 category 993 (HH011). The uncertainty of the tracer method is much smaller than the difference between the 994 two groups. The tracer method also illustrates that 4 of the 5 leaks that were contributed by the 995 LDC to the intercomparison campaign were extremely small. If these would be representative 996 for locations where the suction method is usually applied, it would indeed indicate a severe 997 emission rate bias for the suction method, not because the measurements themselves are biased, 998 but because locations with low emission rates are targeted with this method. In the 999 intercomparison campaign, we attempted to apply the suction method also at locations of the 1000 A categories, but at 8 out of 9 locations from the A category, the suction measurements could 1001 not be applied for safety reasons, or suction could not be completed, because of the widespread subsurface accumulation (Table 2). At the other A location (HH014), the suction method could 1002 1003 not be applied as the ground had been already opened for the repair.
- 1004

# **5** Conclusion

1005 1006

1007 In summer 2020, we compared three gas leak rate quantification methods, namely the mobile, 1008 tracer, and suction methods, in Hamburg, Germany. While the mobile and tracer methods 1009 have been compared previously, this is the first peer-reviewed study that includes the suction 1010 method, although suction measurements could not be completed in one day at most locations. 1011 The mobile method can cover large areas in a short time, but some of the smaller leaks (< 0.51012 L min<sup>-1</sup>) are not identified as a gas leak location due to the 10% enhancement threshold in the 1013 standard mobile quantification algorithm. While the mobile method quantification algorithm is 1014 designed to accurately report city-level total gas distribution leak rates (i.e., considering a large 1015 sample size), it has large (known) uncertainties for individual leaks. The tracer method has a 1016 smaller uncertainty, but it is labor intensive in comparison to the mobile method. On average, 1017 CH<sub>4</sub> emissions from natural gas pipeline leaks were higher from mobile quantifications in 1018 comparison to tracer quantifications. For many locations, we encountered several outlets and 1019 with widespread underground gas accumulations. At one location, after deployment of the 1020 mobile and the tracer quantification and during the repair actions, it was found out that the 1021 actual leak in the gas pipeline was located  $\approx 60$  m away from the identified emission outlet 1022 indicating significant underground gas migration. It is possible that this leak had several 1023 emission outlets that were not identified and the emission quantified from the single outlet is 1024 thus not representative for the whole emission from this leak.

1025 The suction method has a low reported uncertainty, but it is even more labor and time intensive than the tracer method. Due to the time and effort needed to plan and execute the measurements, 1026 1027 the suction method is likely never applied in routine operation at A1 or A2 safety category 1028 leaks that mandate immediate or near-time repair. In our study, it was also not feasible to apply the suction method at locations with large subsurface CH<sub>4</sub> accumulations. Our results thus 1029 1030 indicate a systematic difference between A1 and A2 (high emissions) versus B and C (low 1031 emissions) category locations, and generally larger emission rates are inferred with the mobile 1032 and tracer methods for sites with widespread subsurface accumulation.

1033 This study did not allow a direct, quantitative comparison of emission rates estimated with all three different methods because of the inability to quantify the same leak locations with all 1034 1035 methods. However, this inability illuminates the importance of site selection for deriving representative emission factors based on empirical measurements. Specifically, the results 1036 1037 suggest that a significant emission rate bias could exist for measurements that are carried out 1038 with the suction method. Our results therefore stipulate that representative site selection 1039 includes sampling at all leak safety categories (MEEM, 2018). Otherwise, this could lead to a sampling and emission rate bias in the national inventory of gas leak CH<sub>4</sub> emission in Germany. 1040

1041

1042 Authors contributions: TR, HM and SS conceived and designed the study. TR coordinated 1043 the campaign in collaboration with DBI, Technical University of Denmark (DTU), 1044 Environmental Defense Fund (EDF), E.On and Gasnetz Hamburg (GNH) teams. HM carried out the mobile measurements, emission outlet attribution, performed the analyses of mobile 1045 1046 data and collectively with TR analyzed the intercomparison results. AD, CS and AMF performed the tracer method and reported the emission rates from the tracer dataset. 1047 HDvdG and TR made instruments and equipment available for the mobile method and CS 1048 1049 provided those for the tracer method. HM wrote the paper, and all co-authors supported the 1050 interpretation of the results and contributed to improving the paper.

1051

1052 **Competing interests**: The authors declare that they have no conflict of interest. 1053

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