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Table of Content

2	1 In	troduction	5
3	1.1	Motivation	5
4	1.2	Review summary	7
5	1.3	Global oil and gas reserves	8
6	1.4	Temporal and geographical trends in gas flaring	9
7	2 O	il and Gas Production	15
8	2.1	Exploration and exploitation	15
9	2.2	Composition of natural gas	16
10	2.3	Thermodynamic properties of varying fuel gas compositions	18
11	3 O	verview of the gas flaring processes	19
12	3.1	Gas flaring emission and its environmental impact	20
13	3.2	Emissions measurements around real-world gas flaring sites	23
14	3.3	Types of Flares	26
15	3.3	3.1 Steam- Assisted Flare	27
16	3.3	3.2 Air-Assisted Flare	
17	3.3	3.3 Pressure-Assisted Flare	
18	3.3	3.4 Non-Assisted Flare	
19	4 Es	stimating emissions from gas flaring	31
20	4.1	Determining the flame regime	31
21	4.2	Emission factors (EF) for gas flaring emissions	
22	4.3	Soot emission from gas flaring	35
23	4.4	Scaling soot emission from lab-based studies	
24	4.5	Soot modelling	41
25	4.6	Gas flaring emissions in global models and inventories	43
26	5 C	onclusion	44
27	6 G	lossary	47
28	7 R	eferences	48
29			

32 Gas flaring and resultant air pollution: A review focusing on Black Carbon

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41 Abstract

Gas flaring is a prominent source of VOCs, CO, CO₂, SO₂, PAH, NO_X and soot (black 42 carbon), all of which are important pollutants which interact, directly and indirectly, in the 43 Earth's climatic processes. Globally, over 130 billion cubic metres of gas are flared annually. 44 45 We review the contribution of gas flaring to air pollution on local, regional and global scales, with special emphasis on black carbon (BC, "soot"). The temporal and spatial characteristics 46 47 of gas flaring distinguishes it from mobile combustion sources (transport), while the openflame nature of gas flaring distinguishes it from industrial point-sources; the high 48 49 temperature, flame control, and spatial compactness distinguishes gas flaring from both biomass burning and domestic fuel-use. All of these distinguishing factors influence the 50 quantity and characteristics of BC production from gas flaring, so that it is important to 51 consider this source separately in emissions inventories and environmental field studies. 52 Estimate of the yield of pollutants from gas flaring have, to date, paid little or no attention to 53 the emission of BC with the assumption often being made that flaring produces a smokeless 54 flame. In gas flares, soot yield is known to depend on a number of factors, and there is a need 55

56 to develop emission estimates and modelling frameworks that take these factors into consideration. Hence, emission inventories, especially of the soot yield from gas flaring 57 should give adequate consideration to the variation of fuel gas composition, and to 58 59 combustion characteristics, which are strong determinants of the nature and quantity of pollutants emitted. The buoyant nature of gas flaring plume, often at temperatures in the 60 range of 2000 K, coupled with the height of the stack enables some of the pollutants to escape 61 further into the free troposphere aiding their long-range transport, which is often not well-62 captured by model studies. 63

64

65 Capsule Abstract

The review identified gaps in the estimation of emissions from the gas flaring process and
argues for explicit recognition of gas flaring emissions in emission inventories and global
models.

69

70 Keywords

Gas flaring; Oil and gas; Black Carbon; Incomplete combustion; Hydrocarbon; Emission
factors
facto

78

79	Variables	used a	and	their	meanings
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80 u_e exit velocity of the flue gas (m s	of the flue gas $(m s^{-1})$
---	------------------------------

81 s the air-fuel ratio

 ρ_e, ρ_{∞} the fuel gas and ambient densities respectively (kg m⁻³)

- ΔT_f mean peak flame temperature rise, K (taken as the difference between the adiabatic
- 85 flame temperature and the ambient temperature)

 d_e burner diameter (m)

- v_o kinematic viscosity of the gas-air mixture (m² s⁻¹)
- L characteristic flame length (m)
- $\dot{m_o}$ mass flux of the flue gas at the burner exit (kgm⁻²s⁻¹)
- f_s stoichiometric mixture fraction
- ϕ equivalence ratio
- T_{∞} ambient temperature

- -

99 **1 Introduction**

100 1.1 Motivation

Humans need energy to drive their technology: and hence, make life pleasurable and worth living. Different forms of energy are in use and new ones are developed in order to meet the increasing needs of society (MacKay, 2008). This quest for dependable, affordable and environmentally benign energy sources has occurred throughout human history; for the last century or so, crude oil has been the focus of world energy. In 2011, crude oil was estimated to provide 52.8 % of the world's energy (13113 Mtoe); with oil and natural gas accounting for 31.5 % and 21.3 % respectively (IEA, 2013).

Human reliance on oil and gas as an energy source is not without its attendant impact on the
environment. During production, detrimental impacts on the environment (air, water and soil)
include: oil spills and leakages; venting; sludge disposal; and flaring (Almanza et al., 2012;
Osuji and Adesiyan, 2005; Osuji and Onojake, 2004; Sonibare et al., 2010). The postproduction impact of oil and gas on the environment is also a major source of concern, but is
not the subject of this review.

114 Gas flaring is often a routine daily activity in oil fields around the world, particularly in oilrich regions of the world where the infrastructure to capture, store and utilise the gas 115 produced is not available. Flaring is most often associated with Nigeria and the Russian 116 Arctic, but it does still occur in more developed economies: the North Dakota Bakken shale 117 region, for instance (see, e.g., http://www.eia.gov/todayinenergy/detail.cfm?id=23752, last 118 119 accessed on 29 January 2016). Despite several calls by international bodies such as the World Bank's Global Gas Flaring Reduction (GGFR) initiative, the volume of gas flared globally 120 appears to have plateaued at around 130 billion cubic metres (bcm) since 2008, or may even 121 have increased (see section 1.4, below). According to GGFR (2013), there was an increase of 122 2 bcm in the volume of gas flared in 2011 compared to the previous year. The latest in this 123

series of initiatives by the World Bank - Zero Routine Flaring by 2030 - was launched by the
Secretary-General of the United Nations, Ban Ki-Moon and World Bank's President, Jim
Yong Kim, in May, 2015 (see <u>http://www.worldbank.org/en/programs/zero-routine-flaring-</u>
<u>by-2030</u>, last accessed on January 29, 2016).

Gas flaring, a prominent source of black carbon (BC) and other pollutants, has been ignored 128 or underestimated in emission inventories and models, as a result of which models are 129 struggling to predict measurements of BC in regions of intense gas flaring. The intensity of 130 gas flaring and specifics of atmospheric transport can combine to enhance the role of gas 131 flaring emissions over very large areas (e.g., the Arctic) (Stohl et al., 2013). Presently, 132 133 treatment of emissions from gas flaring is rather rudimentary in most global emission inventories. As at 2015, to the best of our knowledge, only two global pollutants inventories 134 explicitly accounted for emissions from gas flaring (see section 4.5). 135

Gas flaring is classified as a miscellaneous BC-rich source, a group which includes aviation 136 137 and shipping which together contribute about 9 % to global BC emission (Bond et al., 2013). 138 Gas flaring is a very different type of combustion compared to other BC sources in this category; gas flaring is characterised by either fuel-regulated or oxidant-regulated open-fire 139 140 (see below) resulting in flames that can be 8-10 metres in length, with flame temperature as high as 2000K. Gas flaring is a year-round activity in most of the intensive flaring regions of 141 the world, and so differs from transport sources in being stationary and differs from other 142 stationary sources (e.g. cooking and biomass burning) by being relatively constant in time. 143 The GAINS (Greenhouse gas Air pollution INteractions and Synergies) model estimated that 144 gas flaring contributes about 4 % of total anthropogenic BC emissions, majority of which are 145 from the leading gas-flaring nations; Russia, Nigeria, and countries in the Middle East (Bond 146 et al., 2013; Stohl et al., 2013). Gas flaring is estimated to contribute 260 Gg to global BC 147 148 estimates annually (Bond et al., 2013), while in Russia, it is estimated to have the largest 149 contribution of 36.2 % to anthropogenic BC emission (Huang et al., 2015). From a 3-year
150 model simulation, more than 40% of annual mean BC near the surface in the Arctic is
151 estimated to be contribution from gas flaring (Stohl et al., 2013).

152 Considering the small number of nations that still flare gas, a contribution of 4 % to global 153 BC represents a significant skew in the global apportionment of BC emissions. On a regional 154 scale, the contribution of this 'overlooked' source of ambient aerosol loading is likely to be 155 significant.

156 1.2 Review summary

For this review, we have collated previous studies (1984-2015) on gas flaring and its 157 contributions to ambient aerosol loadings. This review is the first to cover virtually every 158 aspects of gas flaring (process, trends, chemistry, flame dynamics and environmental impact). 159 The review starts with a brief assessment of the level and distribution of oil and gas reserves 160 around continents of the world. Compositional variation of the natural gas flared and their 161 thermodynamic properties are discussed because they are likely to have significant impact on 162 pollutant emission rates and overall amount. Next, we provide a discussion of the temporal 163 164 and geographical trends in gas flaring, including brief comments about how weather conditions in regions where gas flaring is common will impact near-field dispersion and long-165 166 range transport. This is followed by a broad discussion of the gas flaring process itself; highlighting how engineering and technology decisions impact on the emission of air 167 pollutants. Finally, several techniques used to estimate emissions from gas flaring are 168 discussed. Throughout this review, special attention is paid to soot (i.e. carbonaceous aerosol 169 predominantly composed of BC) emission from gas flaring because of the now known 170 contribution of BC to global warming and the apparent neglect of the contribution of gas 171 172 flaring to ambient air aerosol loadings in inventories and global models.

173 **1.3** Global oil and gas reserves

World-proven natural gas and oil reserves at the end of 2012 stood at 187.3 and 1668.9 trillion cubic metres (tcm) respectively, sufficient to meet 55.7 years of global production (BP, 2013). The distribution of these reserves among regions of the world is shown in Figures 1(a) and (b). The natural gas collected during the exploration of crude oil from the Earth's crust can be a very good source of fuel; transported in pipes for industrial or domestic use and also recycled back into the processing operation (Davoudi et al., 2013; Elvidge et al., 2009).



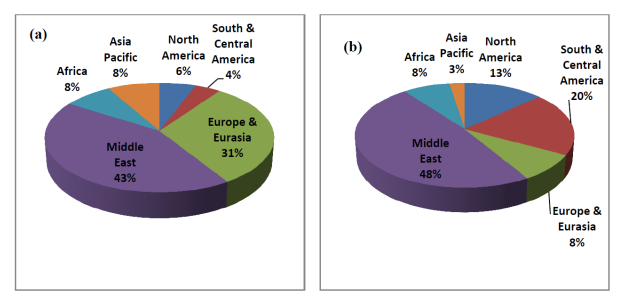


Figure 1: (a) The distribution of world natural gas reserves (b): The distribution of world oil
reserves (adapted from BP (2013).

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In developing countries and oil-rich regions where the technology, infrastructure, and market structure to put all of the natural gas to meaningful use are not available or inadequate, the excess natural gas then becomes a waste stream and is flared or vented. Gas flaring has been termed 'gross waste' by the World Bank's initiative against gas flaring (Global gas flaring reduction: GGFR), because flaring represents direct injection of fossil carbon into the atmosphere without capture and utilization of the heat produced by combustion.

191 **1.4** Temporal and geographical trends in gas flaring

The all-time peak of volume of gas flared, 172 bcm, was in 2005 (Elvidge et al., 2009). 192 Figure 2(a) shows the quantity of natural gas produced by the top 10 oil producing nations of 193 the world between 2000 and 2011, and for comparison, Figure 2(b) shows the estimated 194 quantity of gas flared by these major oil producing nations during the same period of time. 195 The estimated volume of gas flared globally in 2012 is also shown in Figure 2(b): the 2012 196 data is obtained from Elvidge et al. (2015). Data for the plots in figure 2(a) and (b) are from 197 IEA (2012) and GGFR (2012), respectively. Between 2008 and 2011, there was no 198 significant decrease in the amount of natural gas flared. In fact, there was a slight increase 199 200 between 2010 and 2011. According to (BP, 2015), the production excludes the quantity flared 201 or recycled. Therefore, total fossil fuel extracted and ultimately released to the environment is the sum of the production and quantity flared. 202

Figure 3 gives the temporal variation of the fraction of total gas extracted that is flared 203 between 2000 and 2011. Nigeria, Libya and Kazakhstan still flare a sizeable amount of their 204 total production. As at 2011, Nigeria and Libya flared about 30 % of their total gas extracted 205 while Kazakhstan flares about 15 %. The 30 % (15.2 bcm) flared by Nigeria is more than 206 207 twice Libya's total gas extracted (6.3 bcm) for the same period. Russia and Nigeria together account for about 35 % of the gas flared globally (Elvidge et al., 2009). Although the fraction 208 of gas flared is decreasing for most countries and for the largest emitters, several countries 209 210 show flat or increasing fractions of gas flared. The estimated quantity of natural gas flared in the US and Canada, as shown in Figure 2b, are just for the off-shore flaring which is 211 presumably responsible for the very low quantities recorded. After 2006, there is a factor of 212 22 and 10 increase in flaring from USA and Canada, respectively, resulting, presumably, 213 from increased exploitation of unconventional hydrocarbon reservoirs ('fracking'). 214

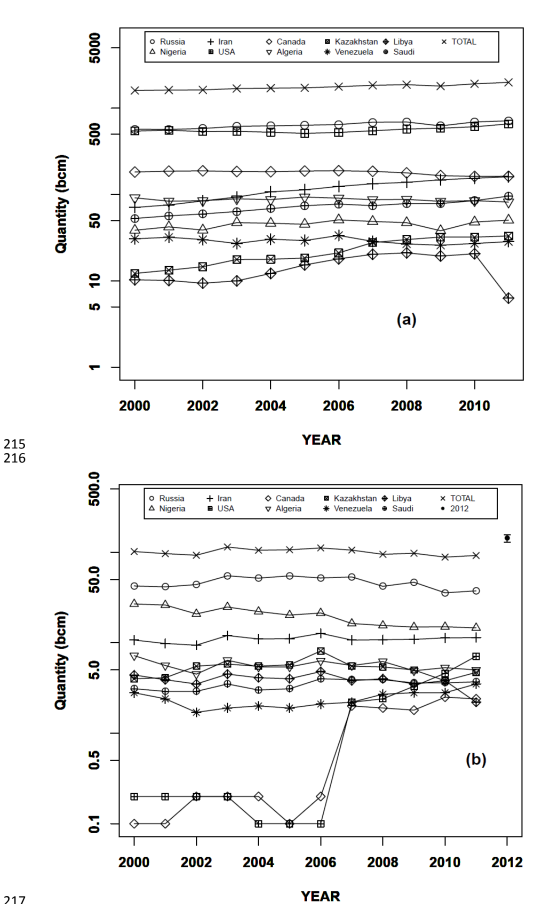




Figure 2: Trend of natural gas (a) production and (b) flaring in major oil producing nations 218 between 2000 and 2011 (adapted from IEA (2012) and (GGFR, 2012) respectively). 219

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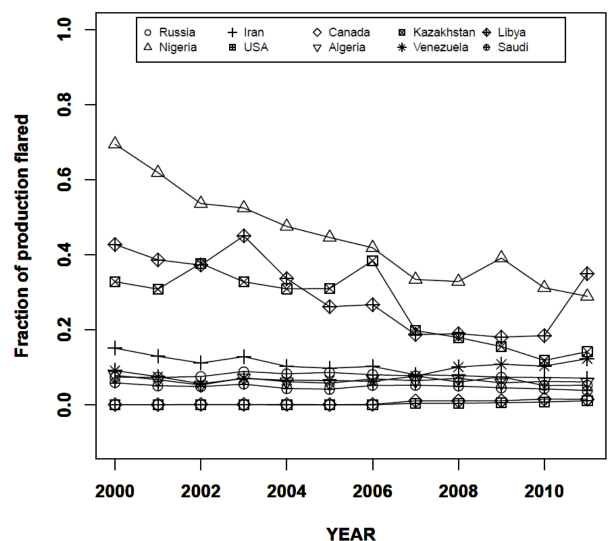


Figure 3: Trend of quantity flared compared to total production (data from Figure 2)
Table 1 gives the estimates, from satellite data, of the quantity of gas flaring by the top 20 gas

flaring nations of the world in 2008 and a brief summary of the climatic conditions of theregion based on the Köppen climate classification (see, e.g., Holden (2005))

Figure 4, reproduced, with permission, from Casadio et al. (2012), shows the global geographic distribution of gas flaring sites obtained by remote sensing techniques. In Africa, the flaring spots are clustered around the North - Algeria, Libya and Egypt - as well as around the Atlantic Ocean especially the Niger Delta area of Nigeria. In Europe, they are around Russia and Kazakhstan, and Iran, Iraq, Kuwait, and Saudi Arabia in the Middle East. Satellite

- remote sensing not only gives a general picture of the spatial distribution of flares, but also a
- 234 gross estimation globally at national levels.

Rank	Country	Gas Flared (bcm)	Regional Climate	Comments*
1	Russia	40.5	(Dfa, Dfb,	peri-Arctic emissions; pole-ward
			Dwa, Dwb),	atmospheric flow around Tibetan
			(Dfc,Dfd,	anticyclone in northern hemisphere
			Dwc, Dwd)	winter
			and ET	
2	Nigeria	15.1	Am and Aw	Tropical monsoon and trade-wind littoral and tropical wet and dry
				climate resulting from West Africa monsoon winds that changes direction with season; climatic condition is controlled by trade
_				wind and movement of the ITCZ
3	Iran	10.4	Bwh	Tropical and subtropical desert climate characterised by large
				diurnal temperature range. Deep
				turbulent boundary layer during th
				day; shallow stable boundary laye
				at night. Large-scale subsidence
				(descending branch of the Hadley circulation) above the boundary
				layer.
4	Iraq	7.0	Bsh	Mid-latitude steppe and desert
·	nuq	7.0	Dom	climate characterised by semiarid annual rainfall distribution
5	Algeria	5.5	Bwh	As for Iran
6	Kazakhstan	5.2	Bwk	Mid-latitude arid wet and dry
0	Razakiistaii	5.2	DWK	climate
7	Libya	3.8	Bwh	As for Iran
8	Saudi	3.5	Bwh	As for Iran
	Arabia			
9	Angola	3.1	Cwa	Humid subtropical climate;
	U			equatorward and poleward
				circulation during winter and
				summer respectively cause change
				in the movement of air masses from
				the cold polar and warm tropics
				within this climate.
10	Qatar	3.0	Bwh	As for Iran
11	Uzbekistan	2.7	Csa	Mediterranean climate; it is
				controlled by the variation between
				subtropical high in summer and

Table 1: Year 2008 estimated gas flared by top 20 gas flaring nations with an estimated error
 of ±2.11bcm.

				polar westerlies in winter
12	Mexico	2.6	Af, Aw, Bsh,	Complex climatic condition; two
			Bsk, Bwh,	tropical, two dry and three
			Cwa, Cwb,	temperate climates
			Cfa, Cfb	-
13	Venezuela	2.6	Aw	Tropical wet-dry climate
14	Indonesia	2.3	Aw, Am	As for Nigeria (see comment above)
15	USA	2.3	Bsk, Bsh,	Mediterranean/dry summer
			Bwk, Csa,	subtropical climate characterised by
			Csb, Af	moist winter and hot dry summer;
				subtropical anticyclones are key
				factor that control this climate
16	China	2.3	Cfa, Cwa	Humid subtropical climate; during
				winter the climate is influenced by
				the Siberian cold and during
				summer there is an inflow warm of
				air from the west
17	Oman	1.9	Bwh	As for Iran
18	Malaysia	1.9	Af	Tropical rainforest characterised by
				constant high temperatures and
				evenly distributed high
				precipitation; controlled by
				movement of the ITCZ and rising
				air along trade wind coast. Strongly
				affected by el Nino Southern
				Oscillation
19	Canada	1.8	Dfb, Dfd, Dsc,	As for Russia (see above)
			Af	
20	Kuwait	1.8	Bwh	As for Iran
	TOTAL	119.3		

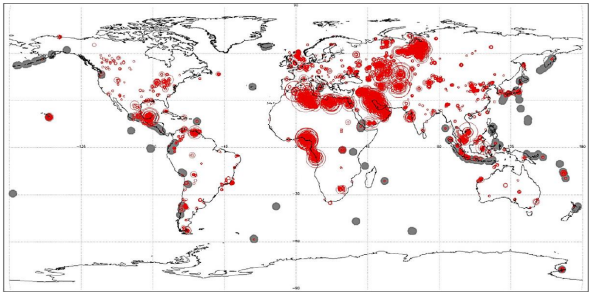
Source: Quantity of gas flared is obtained from Elvidge et al. (2009). *See Holden (2005) 237 238 239 Elvidge et al. (2015), using data collected by National Aeronautics and Space 240 241 Administration/National Oceanic and Atmospheric Administration Visible Infrared Imaging 242 Radiometer Suite (VIIRS), identified more than 7000 active flare globally in 2012: the bulk of which were found in the upstream sector of the oil and gas industries. In 2012, the 243 estimated volume of gas flared globally is 143±13.6 bcm (Elvidge et al., 2015). Compared to 244 119.3±2.11 bcm estimated for 2008 (Elvidge et al., 2009), this is an increment of about 20 % 245 in the central tendencies of the estimates for both years and is well outside the combined 246 uncertainty bounds of both estimates. An increment of about 20 % over four years, stands in 247 stark contrast to the decrease anticipated as a result of the World Bank's Global Gas Flaring 248

Reduction (GGFR) initiative. A new dimension to the problem is the inclusion of three new
countries – India, Egypt and Turkmenistan - in the list of top 20 gas flaring countries.

The geographical location of the sources of gas flaring emissions (as monitored from space, 251 252 see Figure 4), the atmospheric behaviour of emitted pollutants, and the pollutant "matrix" from other sources into which emission are made, determine to a large extent the effect gas 253 flaring emissions will have on regional pollution loadings and on climate. BC emissions in 254 the peri-Arctic and West African Monsoon (WAM) regions have the potential to interact with 255 regional radiative energy budgets and atmospheric circulations, leading to impacts on their 256 257 respective regional climates. Gas flaring emissions in the tropics, especially the WAM and South Asian Monsoon (SAM), could have significant regional impact as a result of the 258 259 intense convective activities and cloud formation in these regions.

260 Monsoon circulations are characterised by large-scale seasonal reversals of wind regimes. 261 Regions often referred to as 'monsoonal' include tropical and near-tropical regions which experience a summer rainfall maximum and most of these regions have a double rainfall 262 263 maximum (Barry and Chorley, 2009). The annual climatic regime over West Africa has many similarities to that of South Asia. Both are characterised by surface air-flow which is 264 determined by the position of the leading edge of a monsoon trough. Winds are south-265 westerly to the south of the trough and north-easterly to the North. The lack of a large 266 mountain range in the north of West Africa strongly enhances the northward advance of the 267 268 WAM compared to its South Asian counterpart. The position of the leading edge of the WAM trough may oscillate greatly from day-to-day through several degrees of latitude 269 (Barry and Chorley, 2009). In the WAM, deep convection occurs in organised systems 270 271 known as Mesoscale Convective Systems (MCS) (Mari et al., 2011; Mathon and Laurent, 2001). Deep convection in the tropics associated with the ITCZ is responsible for intense 272

- mixing, venting of the atmospheric boundary layer, and long-range transport of aerosol (Law
 et al., 2010; Reeves et al., 2010; Sultan and Janicot, 2003).
- 275



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Figure 4: Flaring hot spot sites (1991-2009) as monitored from space are indicated as red spots while grey spots represent position of active volcanoes during the same period Casadio et al. (2012).

280 2 Oil and Gas Production

281 **2.1 Exploration and exploitation**

Oil exploration can be a very complex and capital intensive process as oil deposits are often 282 located in reservoirs buried far into the Earth. These locations can be in very remote and 283 inhospitable parts of the Earth and can be either on- or off-shore. The multi-staged process of 284 exploration, exploitation and processing of crude oil in its raw form from the Earth's crust 285 can be broadly divided into upstream, midstream and downstream. Environmental 286 287 contaminants are expelled into the ambient environment - soil, air and water - at different 288 stages of the process. Crude oil is found in reservoirs which also contain gas. This gas is known as "associated natural gas", and it is separated from the crude oil at a Flow Station. 289 290 Natural gas includes both "non-associated" gas originating from fields producing only gaseous hydrocarbons, and "associated" gas produced in association with crude oil (IEA, 291

292 2012). Natural gas comprises mainly hydrocarbons, predominantly short-chained alkanes. At the separation stage, some of the natural gas is captured for domestic and industrial use while 293 the rest is disposed of, usually by *flaring* in open flames. Below, we will use the term 'fuel 294 295 gas' to refer to natural gas that is flared. The quantity of contaminants expelled at this stage of processing depends on the technology employed, quantity of crude oil processed, flare 296 geometry and design, prevailing meteorological conditions and the composition of the flared 297 298 gas (E & P Forum, 1994; Ismail and Umukoro, 2014; Johnson and Coderre, 2011; Obanijesu et al., 2009; Ouf et al., 2008; Sonibare and Akeredolu, 2004; Talebi et al., 2014). 299

Oil and gas are produced in many wells in different parts of the world at rates varying from 50 m^3 to 700 m^3 per day. As a result of the diverse nature of the geological features of the area where these explorations take place, the composition of oil and gas varies from one station to another.

304 2.2 Composition of natural gas

The composition of natural gas from 10 stations from around the world is given in Table 2. Fuel gas is a combination of C_1 to C_{7+} hydrocarbons which are predominantly alkanes. A typical fuel gas sample contains CH_4 , C_2H_6 , C_3H_8 , $n-C_4H_{10}$, $i-C_4H_{10}$, $n-C_5H_{12}$, $i-C_5H_{12}$, C_6H_{14} , C_7H_{16} , H_2S , CO_2 and N_2 , where 'n' and 'i' stand for 'normal' i.e., straight chained, 'iso' or branched-chained alkanes, respectively. The separation of gas and liquid is not perfect at the Flow Stations and as such trace amounts of liquid can occur in the gas stream, enhancing the abundance of higher molecular weight alkanes in the fuel gas.

Composition					Flow S	tations					Lab-l	oased
	1	2	3	4	5	6	7	8	9	10	Α	В
CH ₄	74.3	79.85	56.9	55.5	82.23	78.41	68.14	68.42	72.32	69.58	85.24	74.54
C_2H_6	14.0	11.54	21.2	18.0	2.38	5.68	14.22	7.65	2.41	0.25	7.06	12.17
C ₃ H ₈	5.8	2.25	6.0	9.8	4.24	0.23	10.27	11.27	6.24	12.54	3.11	5.37
nC_4H_{10}	2.0	2.58	3.7	4.5	0.94	0.70	3.23	4.39	8.12	2.35	1.44	2.49
iC ₄ H ₁₀	-	0.14	-	-	5.12	4.12	2.38	4.42	5.12	5.12	-	-
nC_5H_{12}	0.9	3.24	1.6	1.6	2.25	9.12	0.75	0.94	3.14	5.20	-	-
iC ₅ H ₁₂	-	-	-	-	2.14	0.25	1.01	1.55	2.48	2.54	-	-
C ₆ H ₁₄	-	0.14	-	-	0.25	0.23	-	0.18	0.15	1.97	-	-
N_2	2.9	0.1	-	0.2	-	0.05	-	0.16	-	0.24	1.24	2.15
CO ₂	-	0.16	7.1	8.9	0.45	1.21	-	1.02	-	0.21	1.91	3.28
H_2S	0.1	-	3.5	1.5	-	-	-	-	0.02	-	-	-
*С:Н	0.2659	0.2715	0.2569	0.2602	0.2730	0.2751	0.2860	0.2852	0.2893	0.2924	0.2541	0.2570
*Molar mass (g mol ⁻¹)	21.4	21.4	25.8	26.9	22.9	24.7	24.3	25.8	27.0	28.6	19.2	21.5
*Molar mass (g mol ⁻¹) without CO ₂ and N ₂	20.5	21.3	22.7	22.9	22.7	24.1	24.3	25.3	27.0	28.4	18.0	19.4
*HHV (MJ m ⁻³)	44.8	46.4	47.8	48.7	49.1	52.0	52.2	54.2	57.2	60.3	39.8	42.5

Table 2: Composition (in mole %) and Some Properties of Fuel Gas from Field Stations and Literature. 312

2 and 5 – 10 are adapted from Sonibare and Akeredolu (2004), 1, 3 and 4 from Ismail and Umukoro (2014) and lab-based (A&B) from McEwen and Johnson (2012) 313 314

Compositions 1, 3 and 4 are from Saudi Arabia, Kuwait and Iraq respectively while 2 and 5 - 10 are from different flow stations in Nigeria.

315 C:H = mass-weighted carbon to hydrogen ratio

316 * evaluated in this work

317 2.3 Thermodynamic properties of varying fuel gas compositions

The higher heating value (HHV) is the total enthalpy of the complete combustion reaction for 318 the gas mixture, plus the heat of condensation of the water produced during combustion; 319 320 lower heating value (LHV) is the total enthalpy of the complete combustion when water remains present in its gaseous form only (Flagan and Seinfeld, 2012). The Volumetric 321 heating value (VHV), measured in kJ mol⁻¹, usually refer to the HHV unless otherwise stated. 322 In gas flaring, HHV, calculable using data from standard thermodynamic tables, defines the 323 total amount of energy available to provide buoyancy to the flare. Buoyancy is an important 324 325 parameter in determining the dispersion of flare emissions (Beychok, 1994; Leahey and Davies, 1984). The heat content of a fuel gas depends on its molar mass and by extension the 326 density of the fuel. Figure 5 presents the best fit line relating HHV to the molar mass of the 327 fuel gases from the Flow Stations given in Table 2. This plot shows the extent of dependence 328 of HHV on the molar mass, and hence, density of the fuel gas. These properties of the fuel 329 gas, varies with the fuel composition. Note that the HHV values for the laboratory flares in 330 Table 1 are lower that the HHV for the Flow Station fuel gases. 331

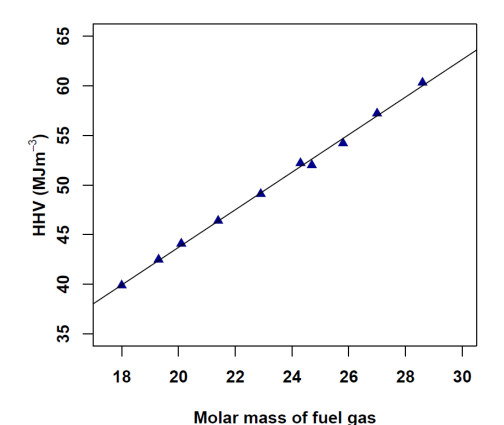


Figure 5: Higher heating value (HHV) as a function of molar mass of fuel gas for the Flow
Station data reported in Table 1.

335 3 Overview of the gas flaring processes

Gas flaring is carried out with the aim to convert its hydrocarbon content, especially methane, to products that are less hazardous to the immediate vicinity of the flare site. Gas flaring is classified as a stationary combustion source for the purpose of air pollution regulation (USEPA, 2008). The combustion process involves the rapid oxidation of the fuel gas with the release of heat, gaseous and particulate pollutants, whose nature and quantity depend on the amount and composition of gas fuel burned, the combustion characteristics as well as the flare geometry and design (Ouf et al., 2008; Torres et al., 2012a).

Gas flaring may be categorized as *emergency*, *process* and *production* flaring depending on the basis of the flaring (Johnson and Coderre, 2011). *Emergency flaring* is unplanned: it is carried out at large facilities for safety purposes for a short duration of time. During

emergency flaring, a large volume of gas is disposed of quickly and, hence, the flow rate of 346 the fuel gas is very high. *Process flaring* is an intermittent disposal of unwanted gas that may 347 last for a few hours or a couple of days at often low flow rate. It occurs during well-testing as 348 349 well as start-up and shutdown of process units. *Production flaring* may occur continuously for years as long as the oil is being explored and exploited. The flow rate can be very high at 350 particular times especially during the initial development of a gas well (Johnson and Coderre, 351 2011). As a result of the length of time involved, which can be years, and the flow rate of the 352 gas flared, production flaring is the major process of concern for regional and global 353 354 pollution, including interaction with climate.

355 3.1 Gas flaring emission and its environmental impact

Gas flaring is the process of disposing of gas (referred to as fuel gas in this review), by 356 combustion in an open flame in the open atmosphere, using a burner tip designed specifically 357 for that purpose, in the course of routine oil and gas production operations (McEwen and 358 359 Johnson, 2012; OGP, 2000; Stone et al., 1992). Gas flaring is a source of greenhouse gases, precursor gases, VOCs, polycyclic aromatic hydrocarbon (PAH) and particulate matter (PM) 360 361 in the form of soot (Ana et al., 2012; Johnson et al., 2013; McEwen and Johnson, 2012; 362 USEPA, 2011; USEPA, 2012). These pollutants have been identified to have serious detrimental impact on animals, vegetation and human (Burney and Ramanathan, 2014; Dung 363 et al., 2008; Pope III et al., 2002; USEPA, 2010). The soot emitted from the combustion of 364 365 natural gas is predominantly black carbon (BC) (Johnson et al., 2013; Smith and Chughtai, 1995). 366

BC is a strong climate forcer and plays a prominent role in the nature of the Earth's climate because of its ability to absorb solar radiation, and hence, to result in a changed vertical gradient of warming by incoming solar radiation (Ramana et al., 2010; Ramanathan and Carmichael, 2008). It also affects cloud processes as well as decreasing surface albedo on ice 371 and snow causing them to melt faster (IPCC, 2007). It can be short-lived in the atmosphere, as it is removed from the atmosphere by dry and wet deposition (although insoluble in water, 372 it is wettable, particularly when 'aged' by atmospheric oxidation). When mixed with other 373 aerosol components in the atmosphere, BC can affect the climate for longer as its residence 374 time is increased in that state (Bond et al., 2013; Flanner et al., 2007; Quinn et al., 2008; 375 Ramana et al., 2010). Within current estimated uncertainties, BC is the second highest 376 contributor to global warming just after CO₂ (IPCC, 2007), and also the main light-absorbing 377 component of atmospheric aerosols (Bond et al., 2013; Chung and Seinfeld, 2005; Jacobson, 378 379 2002; Ramanathan and Carmichael, 2008; Seinfeld, 2008).

The radiative forcing capacity of a cloud of aerosol particles depends on the ratio of the BC to other components in the cloud (Ramana et al., 2010). The ubiquitous nature of BC coupled with its several effects on the Earth's climate makes the study of its sources and emission rates important. If adequate measures are put in place to reduce BC emission, short-term reduction of radiative forcing can be achieved in the Arctic and other oil rich regions of the world (Arctic Council, 2013; Feichter and Stier, 2012; Tripathi et al., 2005).

Studies have shown that the contributions from oil and gas processing, especially gas flares, 386 to air pollution have been severely underestimated (Edwards et al., 2013; Johansson et al., 387 2014; Schultz, 2014; Stohl et al., 2013). This is arguably due to the fact that emission factors 388 389 (EF) often used to estimate the emissions of pollutants from the oil and gas sector, especially 390 gas flaring, are too general and independent of site specifics such as fuel composition and combustion characteristics. There is a pressing research need for more measurements and 391 development of EFs estimates that vary with the fuel gas composition and combustion 392 393 characteristics.

At the local scale, ground-level measurements within one kilometre downwind of the flaring
sites have indicated an elevated concentration of particulate and gaseous pollutants (Edwards)

et al., 2013; Obanijesu et al., 2009; Sonibare et al., 2010). Likewise, on a regional scale,
flight campaign during ACCESS (Arctic Climate Change Economy and Society) around oil
and gas installations in Heidrun, Norway reported elevated concentrations of NO, SO₂ and
CO in the lower troposphere (Roiger et al., 2015).

NO_X (NO+NO₂), SO₂, CO₂, CO and unburned hydrocarbon are the major pollutant 400 constituents of gas flaring plumes (USEPA, 1995). The low amount of the nitrogen and 401 sulphur content of the fuel gas notwithstanding, the NO_X and SO₂ emission from gas flaring 402 remain significant because the ambient background level of both pollutants are usually low. 403 Hence, gas flaring can substantially enhance the local concentrations of these pollutants. 404 405 Complete combustion is often not achieved in most flaring conditions (Leahey et al., 2001). During incomplete combustion, methane (CH₄) and other unburned components of the fuel 406 gas are given off and some fuel components are partially oxidized to CO and soot rather than 407 408 completely oxidised to CO₂ (RTI, 2011; Strosher, 2000; Villasenor et al., 2003). Equations 1a and b, respectively, give simplified equations for the 'ideal' complete and partial oxidation of 409 410 the fuel gas. In equation (1b), 'rC' denotes soot production, ' vO_2 ' excess oxygen in the case of fuel-lean combustion and C_aH_b are PAH and other semi-volatile organics resulting from the 411 pyrolysis of hydrocarbons in the fuel gas. The value of m in equation (1b) determines a fuel-412 413 lean (over-fired; m > x + y/4) or fuel-rich (under-fired; m < x + y/4) combustion process. The level of combustion (oxidation) of the fuel gas is dependent on a number of factors: the 414 nature of the flame during combustion, the level of mixing of the gas and air in the reaction 415 zone, the amount of oxidant (oxygen) available, the VHV of the fuel gas and the prevailing 416 condition of the ambient wind (Stone et al., 1992). Flare design and geometry are also key 417 determinants of the level of combustion of the fuel gas. 418

419
$$C_x H_y + \left(x + \frac{y}{4}\right) O_2 \to x C O_2 + \frac{y}{2} H_2 O$$
(1a)

420
$$C_x H_y + m O_2 \rightarrow pCO_2 + qCO + rC + vO_2 + fH_2O + \sum_{a,b} d_{a,b} C_a H_b$$
 (1b)

421 For 'ideal' complete combustion, the oxidation of the hydrocarbon yields carbon dioxide and 422 water only (equation 1a), while the oxidation of the sulphur (as H₂S) and nitrogen content of the fuel gas gives SO_2 and NO_X respectively. In gas flaring, NO_X is produced by thermal 423 424 cracking of the nitrogen content of the fuel gas and entrained atmospheric nitrogen. The 425 amount of carbon-containing emission (CO₂, CO, C_aH_b, and BC) given off depends on the molar mass of non-CO₂ carbon per mole of the fuel gas (cf. Table 2). Smoking of a gas flare 426 does not necessarily imply that the combustion process is highly inefficient; because a small 427 428 amount of soot can absorb and scatter perceptible amounts of visible light, a flare with a combustion efficiency of 99 % could still smoke visibly (Castineira and Edgar, 2006). 429

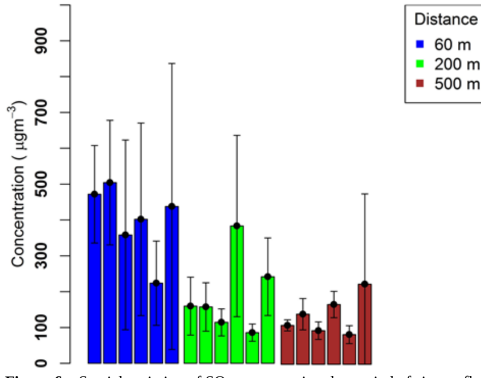
430 **3.2** Emissions measurements around real-world gas flaring sites

Few data of pollutants measurement around gas flaring sites are publicly available. Data from
the ACCESS aircraft campaign experiment in the Arctic (Norway) and a couple of groundbased in-situ measurements in Nigeria and the US show significant contributions from gas
flaring to ambient air concentrations of these pollutants.

In-situ ground measurements of air pollutants around typical oil and gas facilities where
varying degrees of gas flaring take place have been undertaken in the US (Edwards et al.,
2013; Edwards et al., 2014; Johansson et al., 2014), Mexico (Villasenor et al., 2003), Norway
(Roiger et al., 2015) and, Nigeria (Ana et al., 2012; Nwaichi and Uzazobona, 2011;
Obanijesu et al., 2009; Sonibare et al., 2010). Continuous noise levels higher than the WHO
limit of 70 dB were also observed around some gas flaring sites in Nigeria (Abdulkareem and
Odigure, 2006; Avwiri and Nte, 2004).

A 4-month sampling of three air pollutants (SO₂, CO and NO₂) around six flow stations in the
Niger Delta area of Nigeria was undertaken by Obanijesu et al. (2009) and Sonibare et al.

444 (2010). The measurements were made at 60, 200 and 500 m downwind of the flaring sites. Although to a varying degree depending on the capacity of the station, gas flaring is a 445 prominent daily activity within the stations. Mean pollutants measurements around the six 446 447 flow stations are shown in Figures 6(a), (b) and (c). The variation bar on the bar-plots shows the standard deviation of the measurements over the four-month period. The nature and 448 extent of dispersion of pollutants from a stationary source depend on the local meteorology 449 and topography of the area. As shown in Figures 6(a)-(c), the trend of the measurements from 450 the flaring sites are similar to observations from dispersion model studies where 451 452 concentrations of pollutants decreases exponentially with distance from the source (Hodgson et al., 2007). Site 4 is the only place where a significant deviation from this trend, especially 453 for CO and NO₂, was observed, suggesting the likelihood of contributions from other 454 455 source(s).

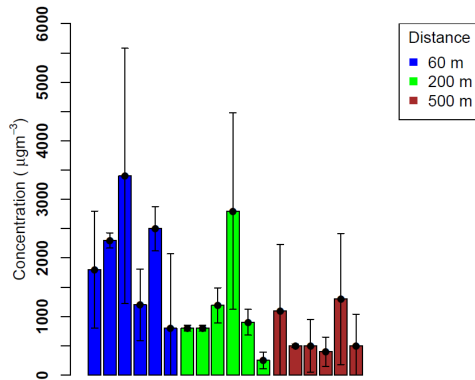


457 Figure 6a: Spatial variation of SO₂ concentration downwind of six gas flaring sites
458 (adapted from Obanijesu et al. (2009))

459

456

460



464

Figure 6b: Spatial variation of CO concentration downwind of six gas flaring sites (adapted from Sonibare et al. (2010))

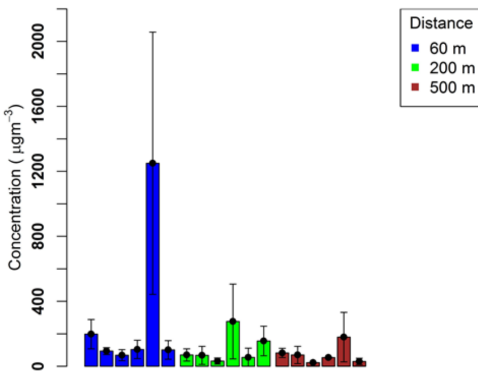


Figure 6c: Spatial variation of NO₂ concentration downwind of six gas flaring sites (adapted from Sonibare et al. (2010))

471 A summary of the few available in-situ ground measurements downwind of gas flaring sites

472 in other regions of the world are given in Tal	ble 3.
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BC (ng kg ⁻¹)	O ₃ (ppbv)	VOC (ppbv)	PAH (ng m ⁻³)	NO (ppbv)	NO ₂ (ppbv)	SO ₂ (ppbv)	CO (ppbv)	Ref
-	> 120	100 -350	-	> 3.5	> 7.5	-	> 80	Edwards et
								al. (2013)
>40	> 25	-	-	> 1.2	-	> 1.2	> 90	(Roiger et
								al., 2015)
-	-	-	0.34 -	-	-	-	-	(Ana et al.
			3.3×10^4					2012)

476

477 **3.3 Types of Flares**

Based on the design and operating condition of the system, flares can be categorised as air-478 assisted, steam-assisted, non-assisted and pressure-assisted. Flares are assisted primarily to 479 enhance the turbulence and mixing of the fuel gas and air in the combustion zone, so as to 480 suppress smoking of the resulting flame (Castineira and Edgar, 2006; Enviroware, 2012; 481 Torres et al., 2012a). The choice of assistance, therefore, affects flame chemistry, as 482 discussed below. Air and pressure-assisted flares are not as efficient as steam-assisted flares 483 in terms of the carbon conversion efficiency (CCE) (Castineira and Edgar, 2006). Complete 484 485 combustion of the fuel gas requires sufficient air for combustion and adequate mixing of the air and fuel gas. The efficiency of a gas flare at a given moment in time depends on the HHV 486 of fuel gas (see table 2), design of the burner, mixing of air and fuel gas in the combustion 487 zone, composition of the fuel gas, wind speed and direction, and ambient temperature and 488 pressure (Kostiuk et al., 2004; Stone et al., 1992; Torres et al., 2012b). 489

490 Equivalence ratio, ϕ , is a measure of the amount of oxygen available for the combustion of 491 the fuel gas (Flagan and Seinfeld, 2012; McAllister et al., 2011). It is defined as:

492 $\Phi = \frac{\left(\frac{A}{F}\right)_s}{\left(\frac{A}{F}\right)_a} \dots$	(2)
--	-----

493 where $\left(\frac{A}{F}\right)_{s}$ is the stoichiometric air-fuel mass ratio and $\left(\frac{A}{F}\right)_{a}$ is the actual air-fuel mass ratio. 494 $\phi < 1$ implies fuel-lean mixture, that is, more oxygen than is needed for the combustion of 495 the fuel is available; $\phi = 1$, a stoichiometric mixture, where the exact amount of oxygen 496 needed is made available, and $\phi > 1$, a fuel-rich mixture, less oxygen than is needed is 497 available for the combustion of the fuel. Burners used in gas flaring and the entire gas flaring 498 set-up are designed to produce a flame operating at $\phi \sim 1$ taking into account the economic 499 aspects such as cost effectiveness of applying a process like steam assistance.

500 3.3.1 Steam-Assisted Flare

Steam-assisted flare involves the introduction of steam jet into the combustion zone of the 501 flare to provide added momentum and turbulence to the fuel gas and air, enhancing mixing 502 and as such suppressing the tendency for smoke in the flare (Castineira and Edgar, 2006; 503 Müller-Dethlefs and Schlader, 1976; Stone et al., 1992). It is the most efficient assist given to 504 flare to suppress smoking during combustion because the steam affects flame chemistry as 505 well as mixing. The steam acts to break up long-chain hydrocarbons to smaller chains that 506 burn with less smoke (Castineira and Edgar, 2006; Fortner et al., 2012; Müller-Dethlefs and 507 Schlader, 1976; Torres et al., 2012a). Steam undergoes thermal dissociation in a flare flame 508 to give H and OH free radicals that react with carbon to give CH2- and -CHO radical 509 moieties. The steam-induced free radicals enhance the formation of C=O bonds rather than C-510 511 C bonds, promoting completeness of combustion. The steam can also react with intermediate products like CO, oxidizing it further to CO₂ (Castineira and Edgar, 2006; Müller-Dethlefs 512 and Schlader, 1976). 513

514

14
$$CO(g) + H_2O(g) \longrightarrow CO_2(g) + H_2(g)$$
 $(\Delta H = -41 \text{ kJ mol}^{-1})$ (3)

For a steam assisted flare, combustion efficiency starts to decrease when the steam-to-fuel gas ratio goes beyond a threshold which depends on the heat content of the fuel gas and the location at which the steam is injected into the combustion zone in the flare (USEPA, 2014).

In the US, a lower heating value (LHV) limit of 11.18 MJ/m³ is imposed on fuel gas in order 518 to be suitable for steam-assisted flares (Castineira and Edgar, 2006). Fuel gas with high heat 519 content promotes higher combustion efficiency in steam-assisted flares (McDaniel and 520 521 Tichenor, 1983; Torres et al., 2012b). Over-steaming results when too much steam is injected into the combustion zone; over-steaming is analogous to over-aeration in air-assisted flares. 522 Over-steaming causes a decrease in the flame temperature by serving as a heat sink. A 523 decrease in combustion efficiency of the flare results, along with, increased noise caused by 524 cavitation created within the flame (Castineira and Edgar, 2006; Stone et al., 1992). 525

The formation of NO_X in steam-assisted flares is reduced, compared to non-assisted flares, and further reduced at large values of equivalence ratio (ϕ) due to a drop in the flame temperature (Miyauchi et al., 1981; Müller-Dethlefs and Schlader, 1976). Steam-assisted flares are rather expensive to maintain, especially for large gas facilities, as a large-scale steam generator is required.

531 3.3.2 Air-Assisted Flare

In air-assisted flares, forced air from a low-pressure blower is used as an additional source of 532 momentum and turbulence to the fuel gas in the combustion zone and, hence, enhances the 533 mixing of the fuel gas and air in the zone (Castineira and Edgar, 2006; Stone et al., 1992; 534 535 Torres et al., 2012a). Air-assisted flaring involves the installation of an air blower that provides the forced air at the bottom of the stack. The major advantages of the air-assisted 536 flare are that it is less expensive to run, extends the life-span of the flare by cooling the tip of 537 538 the flare and is easier to maintain than other configurations (Castineira and Edgar, 2006). For an air-assisted flare, the combustion efficiency of the flare decreases linearly above a 539 threshold limit of the air assist to fuel-gas ratio, but the rate of decrease is slow for a fuel gas 540 with a higher VHV (Torres et al., 2012b). Incomplete combustion can occur when air-fuel 541

gas ratios go beyond the optimum value, to the extent that the flame may be put out as aresult of over-aeration (Castineira and Edgar, 2006).

544 3.3.3 Pressure-Assisted Flare

In pressure assisted flares, the fuel gas stream pressure is controlled by varying the volume 545 flow of the fuel gas, and used to enhance the mixing of the fuel gas and air in the combustion 546 zone. A high-pressure burner is used to promote atomization of any liquid hydrocarbon and 547 enhance the mixing of the fuel gas with air to bring about a complete or near-complete 548 combustion. Pressure assistance often requires significant amount of space in a remote area 549 550 because of the burners arrangement at ground level. Fuel gas exit velocity increases with pressure at the burner. Pressure-assisted flares usually have burners arranged on the ground 551 and as such must be located carefully within the oil and gas production plant (Enviroware, 552 553 2012; Stone et al., 1992).

554 3.3.4 Non-Assisted Flare

555 For non-assisted flares, no provision is given to provide momentum and enhanced mixing for the fuel gas and air. The method is often used for gases with low VHV, that is, fuel gas 556 having low C-to-H ratio (in alkanes, C-to-H ratio increases from a minimum value of 0.25 for 557 methane; see Table 2). The C-to-H ratio determines the smoking tendency of hydrocarbons, 558 with smokiness increasing with C-to-H (USEPA, 1995). Note that the experimental fuel gas 559 compositions listed in Table 2 are at or below the lowest C-to-H ratio of the Flow Station gas 560 compositions listed. Non-assisted flaring is used for gases that require smaller amounts of air 561 to undergo complete combustion (Enviroware, 2012; Stone et al., 1992). 562

Table 4 summarises the features of the different types of flares discussed in Section 3.3.

564

	Steam-assisted	Air-assisted	Pressure-assisted	Non-assisted
Method	Steam is introduced into the combustion zone to enhance mixing.	Air is introduced from a blower to enhance the mixing and turbulence of the fuel gas in the combustion zone	The vent pressure of the gas flow is used to enhance mixing at the tip of the flare burner	No assistance is given to the combustion process
Efficiency	Most efficient in terms of suppressing soot formation. Some of the CO formed can be oxidized to CO ₂	Less efficient than the steam assisted flare but relatively efficient that the other two types	Not as efficient as steam and air- assisted but can equally suppress sooting.	Only efficient for non- sooting combustion especially in light hydrocarbons
Benefits	Fuel with high heat value, and hence, high sooting propensity can be disposed of with relatively less soot	Prolongs the life span of the flare tip. Less expensive than steam-assisted and easy to maintain, hence, it is the most commonly used.	Enhance combustion efficiency when the gas flux pressure is sufficiently high enough without the additional cost of steam and air generation	Can be used for occasional emergency flaring of near smokeless gas
Relative size	They are often large flares as they include the steam generator and are usually employed in large gas facilities.	Not as large as the steam assisted.	May be of same size as air- assisted flare depending on the flow capacity of the facility	Often smaller in size compared to the other types
Shortcoming	Over-steaming can result in reduced efficiency of flare. It is also expensive to maintain on a large-scale	Over- aeration can also result in less efficiency. A limit of air assist to gas ratio must be maintained for effectiveness of the flare.	The fluctuation of gas flow pressure has a bearing consequence on the efficiency of the combustion. Requires large space in a remote area.	Cannot be used for dense fuels with high sooting propensity which are typical gas in oil and gas processing facilities

Table 4: Summary of the properties of flare types

568 **4** Estimating emissions from gas flaring

Emissions from a typical gas flare can be solids, liquids or gases. As a result of the 569 inaccessible nature of full-scale real-world gas flares, several techniques have been used to 570 quantify the amount of emissions from such flares. Such methods include measurement or 571 source monitoring often by lab-based, pilot-study-based or field-based study (Johnson et al., 572 2011; McEwen and Johnson, 2012), application of emission factors obtained from 573 measurements and scaling calculations (Giwa et al., 2014; Huang et al., 2015; Sonibare and 574 Akeredolu, 2004; Talebi et al., 2014; USEPA, 1995) and, simulations, often by computational 575 fluid dynamics (CFD) (Almanza et al., 2012). 576

In real-world flares, complete combustion cannot be achieved always and everywhere. Incomplete combustion of the fuel gas can be due to poor efficiency of the flare system, flame temperature (flame dynamical characteristics), insufficient oxygen resulting in poor stoichiometric air/fuel gas mixing ratio, the condition of the fuel gas in the combustion zone and prevailing ambient meteorological condition (Stone et al., 1992). Carbon monoxide (CO) can represent 24 - 80 % (on carbon a molar basis) of emissions for an incomplete combustion process (Torres et al., 2012b).

584 **4.1 Determining the flame regime**

It is important to define clearly the configuration of the fire (flare) as this is essential for an 585 adequate estimation of the yield and transport of pollutant species from the combustion 586 process. Flames can be classified along a spectrum ranging from turbulent diffusion flames 587 588 (of the kinds discussed above) to pool fires (e.g. tar-pool fire) based on the nature and dynamics of the fuel in the flame as well as the design of the burning process (Delichatsios, 589 1987, 1993a). In this review we are concerned with gas flares, which are classified as 590 turbulent jet-diffusion flames. They are so classified because of the high pressure associated 591 with the release of the fuel gas into the flame. 592

Jet-diffusion flames in the environment can be categorized based on the momentum flux 593 ratio, R, of the jet plume versus the horizontal momentum flux of the ambient wind (Huang 594 and Wang, 1999). Flares with high R (R>10) may be further categorized depending on 595 whether the flame characteristics are driven by the buoyancy of the hot plume or momentum 596 of the fuel gas (McEwen and Johnson, 2012). Both buoyancy and momentum are important 597 in determining the character of flares. A combination of several dimensionless parameters -598 599 Richardson number, Richardson ratio, fire Froude number, gas Froude number, and Reynolds number - have been used in studies to configure the regime of the flame in a jet-diffusion 600 601 flame. The Froude numbers measure the ratio of the inertia force on an element of the fluid (in this case, gas or fire) to the weight of (i.e. gravitational force acting on) the fluid element. 602 The fire Froude number, Fr_f , gas Froude number, Fr_g and Reynolds number, Re, have 603 proved to be useful dimensionless parameter to define the flame regime (Becker and Liang, 604 1982; Delichatsios, 1993a; Delichatsios, 1993b; Delichatsios, 1987; Sivathanu and Faeth, 605 1990). 606

607
$$Fr_g = \frac{u_e f_s^{3/2}}{(gd_e)^{1/2} (\frac{\rho_e}{\rho_\infty})^{1/4}} \dots$$
(4)

608
$$Fr_f = \frac{u_{efs}^{3/2}}{\left(\frac{\Delta T_f}{T_{\infty}}gd_e\right)^{1/2}\left(\frac{\rho_e}{\rho_{\infty}}\right)^{1/4}}$$
....(5)

$$609 \qquad Re_s = \frac{u_e d_e}{v_o} \tag{6}$$

610
$$f_s = \frac{1}{S+1}$$
.....(7)

In the definition of Fr_f in equation (5), $\left(\frac{\Delta T_f}{T_{\infty}}\right)g$ is the effective acceleration generated by individual hot eddies burning at the flame temperature (Delichatsios, 1987). Among the three dimensionless numbers defined by equations (4) - (6), the Reynolds number is used to determine the status of flow, either turbulent or laminar. The fire Froude number, Fr_{f} , is used to identify the dominant mechanism between buoyant-generated turbulence and momentumgenerated turbulence. In practice, Fr_{f} can be used to parameterise soot yield from turbulent diffusion flames (Delichatsios, 1993b; McEwen and Johnson, 2012).

618 **4.2** Emission factors (EF) for gas flaring emissions

The Emission factor (EF) of a pollutant is the amount of the pollutant released into the 619 atmosphere per unit activity or per unit raw material consumed. It can be obtained from 620 experimental measurements carried out on several sources which represent a particular 621 622 emission source type. For example, road transport emission factor can be compiled by measuring the amount of each pollutant (CO₂, CO, PM, NO_x) given off by cars (petrol and 623 diesel), heavy duty vehicles and motorbikes per litre of fuel burned for every km travelled 624 (Gertler et al., 1998; USEPA, 1995; Zhang and Morawska, 2002) under given driving 625 conditions. It is often expressed in $g m^{-3}$ (pollutant produced per unit volume of raw material 626 consumed), g kg⁻¹ (pollutant produced per unit mass of raw material consumed), g km⁻¹ 627 (pollutant produced per unit distance travelled). Emission factors have been compiled by 628 several agencies, which include the US Environmental Protection Agency (USEPA), the 629 European Environmental Agency (EEA), the United Kingdom Department for Environment, 630 Food and Rural Affairs (Defra), and GAINS (Greenhouse gas Air Pollution Interactions and 631 Synergies), for several source categories based on technical sessions of lab-based studies, 632 pilot studies or actual field measurements. Of these agencies, ECLIPSE and USEPA have 633 EFs specifically for emissions from gas flares: the GAINS emission factor for BC from gas 634 flaring is 1.6 g m⁻³: the equivalent USEPA value has four discrete values between 0 and 6.4 g 635 m^{-3} depending on the smokiness of the flame (see below). Stohl et al. (2013) in their study 636 used emission factor of 1.6 g m⁻³ obtained from ECLIPSE (Evaluating the Climate and Air 637

638 Quality Impacts of Short-lived Pollutants) emissions data set to simulation BC emissions in639 the Arctic.

It is difficult to carry out an accurate estimate of emission from gas flares directly from field measurements. Conventional experimental techniques are not suited due to the severe operating conditions that occur in the flaring process and the almost uninhabitable nature of the gas flaring area to both man and the field equipment during the process of gas flaring (Ismail and Umukoro, 2014; McDaniel and Tichenor, 1983; McEwen and Johnson, 2012; RTI, 2011; Talebi et al., 2014). As a result of the unsteady and opaque nature of gas flare flames, remote sensing provides only a partial answer to the difficulties of in-situ monitoring.

In Chapter 13 of USEPA's 5th edition of the compilation of air pollutant emission factors 647 648 known as AP-42, published in January 1995, emission factors (EF) for pollutant emitted from industrial flaring of waste gas were given; these were recently updated in 2014 (though still 649 in the draft stage) except those for soot. The EFs published in 1995 were based on a study 650 651 conducted by McDaniel and Tichenor (1983) aimed at determining combustion efficiency and hydrocarbon destruction efficiency for flares operated under different condition. The 652 recent updates of EFs in AP-42 give emission factors of 0.17 kg GJ^{-1} , 1.43 kg GJ^{-1} and 0 - 6.4 653 $g m^{-3}$ are given for CO, NOx and soot, respectively. 654

The fuel used in the pilot study to estimate EFs for industrial flare pollutants in the AP-42 compilation was predominantly propylene and inert diluents. As such, the EFs, especially for soot, might not be an adequate representation of a typical flare in the oil and gas industry with varying fuel gas composition (Table 2). For AP-42, the collection of soot at the experimental stage was not done by a conventionally accurate method. For instance, the particulate matter (soot) given off was not collected isokinetically in accordance with USEPA's method 5 for sampling particulate matter from stationary source (USEPA, 2000). An emission factor of 0.0

 $\mu g L^{-1}$ (microgram per litre of fuel gas at standard temperature and pressure) given for soot in 662 non-smoking flares of industrial flares in AP-42 is of limited utility and presumably denotes a 663 upper limit of 0.05 μ g L⁻¹, because the soot yield from flares is never likely to be precisely 664 zero. Even seemingly modest emission of soot (BC) can be significant, given what is now 665 known of the effects of BC on climate, weather and human health from findings in recent 666 studies (Bond et al., 2013; Ramanathan and Carmichael, 2008; Stocker et al., 2013; Tripathi 667 668 et al., 2005; Wang et al., 2014).

The mass/mole balancing technique for estimating the yield of pollutants from the 669 combustion of hydrocarbon is a widely used technique. Using this technique, Ismail and 670 Umukoro (2014) and Sonibare and Akeredolu (2004) estimated the yield for SO₂, NO_X, CO₂ 671 and CO from the combustion of fuel gas (hydrocarbon with inert diluents) at various levels of 672 combustion. Both studies did not account for unburned carbon (soot) even for their 'severely' 673 674 incomplete combustion process reactions. Ismail and Umukoro (2014) varied combustion efficiency (CE) and air available for combustion, and with CE as low as 0.5 (50 %) still did 675 not account for unburned carbon. 676

677 4.3

Soot emission from gas flaring

Soot, which is predominantly black carbon (BC), is a product of the incomplete combustion 678 679 of biomass, solid fuel and fossil fuel (Goldberg, 1985; Koch et al., 2009). Globally, fossil fuel 680 combustion is estimated to contribute 3 Mt of BC to the atmosphere annually (Bond et al., 2004). Annually, the contribution of gas flaring to global BC concentration is estimated to be 681 260 Gg (Bond et al., 2013), that is approximately 0.1% of the total contribution from fossil-682 fuel use, of which Russia is estimated to contribute 81.0 Gg (Huang et al., 2015). 683

The formation and quantification of soot from the combustion of hydrocarbon is a rather 684 complex thermo-chemical process that is not well understood, despite decades of research 685

(Castineira and Edgar, 2006; Haynes and Wagner, 1981; Johnson et al., 2011; Maricq, 2009).
Soot given off from the combustion of hydrocarbon is predominantly elemental carbon and
the amount given off depends on a number of factors, including the efficiency of the
combustion process and the fire dynamical characteristics.

The diameter of soot particles emitted ranges between 10 – 200 nm and most commonly lies between 10 and 50 nm. Figure 7 shows a TEM micrograph of soot particle and agglomerates from acetylene flames. The very fine particle of soot are from seemingly 'non-sooting' flames while at the other extreme are those from heavily sooting flames (Flagan and Seinfeld, 2012; Glassman, 1996; Haynes and Wagner, 1981). Combustion of hydrocarbon components of the fuel gas has PAH and BC signatures which can be used as tracers for emissions from the flaring of fuel gas (Fortner et al., 2012; Maricq, 2009).

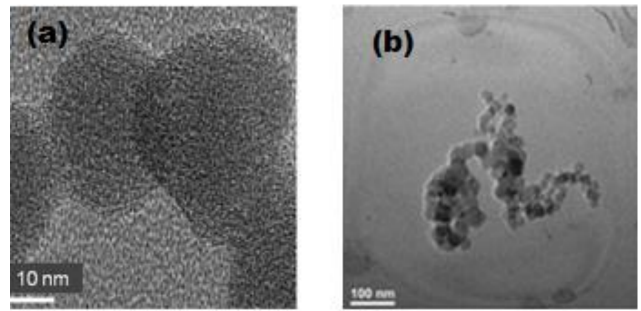


Figure 7: TEM micrograph of soot (a) microstructure (b) agglomerates (Tumolva et al.,
2010).

701 Soot is formed when the carbon particles are cooled below their ignition temperature and 702 there is a deficiency of oxygen (Stone et al., 1992). Considering the variation of the

703 composition of gas flared from one station to the other, EFs available for estimating emissions from gas flares are overly generalised. The fuel gas used in most of the studies to 704 estimate the EFs is either propylene or propane or a mixture of both with nitrogen added to 705 706 alter the heat content as well as the use of predominantly methane-based fuels which are known to be low-sooting. 707

Several attempts have been made to quantify and study the characteristics of soot yield from 708 709 gas flares using various approaches in lab-based, pilot-based and field-based studies as well as simulation techniques. Emission factors from some of these studies are given in Table 5. 710 711 Other emission factors not highlighted in this table are based on the USEPA's AP-42 emission factors. 712

- 713
- 714

Study (year) Emission factor (g m⁻³ **Emission rate** Type of study Fuel of fuel burned) $(g s^{-1})$ 80% propylene 20% propane USEPA (1995) $0.0, 0.9, 4.26.4^{a}$ Pilot study Johnson et al. 2.0 (±0.66) Field study associated (2011)natural gas Almanza et al. 0.025 - 0.22**CFD** simulation associated (2012)natural gas McEwen and 0.51 Lab-scale flare hydrocarbon Johnson (2012) (alkanes) Johnson et al. (2013)0.067 Field study **GAINS 2011** Modelling 1.6 associated IMP (2006)^b 3.37 Simulation natural gas associated CAPP (2007) 2.563 natural gas

Table 5: A Summary of Emission Factors/Emission rate for Soot from Industrial Flares 715

^a0.0 is for non-smoking flame, 0.9 for light smoking flares, 4.2 for averagely smoking flare 716 717 and 6.4 for *heavily smoking* flares

^b cited in (Almanza et al., 2012) 718

719

The studies by Johnson et al. (2011) and Johnson et al. (2013) gave emission rates ($g s^{-1}$) and 720 not emission factors $(g m^{-3})$ because both were actually field studies that quantified the soot 721

given off per unit time by estimating the travel (speed) of the soot in space using a chargecouple device camera (CCD) camera viewing a real field flare.

Emission of soot from the combustion of hydrocarbon varies with the VHV of the gas and, as 724 such, the emission factor for soot can be estimated from its VHV (McDaniel and Tichenor, 725 1983; RTI, 2011; Torres et al., 2012b). McEwen and Johnson (2012), using a lab-based 726 experiment, studied the BC emission from the combustion of hydrocarbon. The study varied 727 the VHV of the fuel gas and measured the soot yield for each combustion process. The VHV 728 used here is the higher heating value (HHV), or gross heating value. The relationship between 729 the two variables (soot yield and HHV) obtained from the study by McEwen and Johnson 730 731 (2012) is given in equation (8):

$$EF_{soot} = 0.0578(VHV) - 2.09$$

When the composition of the fuel gas is known, its VHV can be calculated from standard 733 thermochemical tables (cf. Table 2). It should, however, be noted that because the VHV 734 contains no information on flame dynamics, this relationship is only appropriate for the flame 735 dynamics conditions of the experiment. Although, equation (8) is a readily available 736 relationship to estimate soot emission from hydrocarbon combustion, its application is 737 738 restricted to a complete or near-complete combustion process. Inserting the estimated VHV into the relation developed by McEwen and Johnson (2012) gives an estimate of the soot 739 yield in $g m^{-3}$. 740

741 **4.4** Scaling soot emission from lab-based studies

Lab-based studies of emissions from flares are the most common and readily available method to estimate emissions from full-scale flares. However, considering the size (diameter) of the flare stack, flow rate of fuel gas, exit velocity, and the resultant buoyancy of full-scale flare, there is a need to scale up the emissions yield from lab-based studies to be representative of a full scale flare. In earlier studies, several dimensionless parameter have been considered for such scaling purpose; these include Richardson ratio, Ri_L (Becker and Liang, 1982), fire Froude number, Fr_f (McEwen and Johnson, 2012), and the first Damköhler ratio, Da_1 (Becker and Liang, 1982). Richardson ratio, Ri_L , is defined as the ratio of the buoyancy-generated turbulent kinetic energy (TKE) of the flame to the TKE of emitted gas jet at the exit:

752
$$Ri_L = \frac{gL^3}{(u_e d_e)^2} \left(\frac{\rho_\infty}{\rho_e}\right) \qquad (9)$$

Richardson ratio is the basis to assess the turbulent regime of the flame: when $Ri_L \ll 1$, 753 buoyancy-induced mixing between emitted gas jet and background air is much weaker than 754 jet-induced mixing, and consequently the flame is dominated by forced convection; when 755 $Ri_L \gg 1$, jet-induced mixing is much weaker than buoyancy-induced mixing and the flame is 756 dominated by natural convection. Fire Froude number, Fr_f , is defined in Equation (5) and can 757 be interpreted as the ratio of the jet's inertia to the buoyancy force acting on it. Fire Froude 758 number can be used to assess the dominating force to "stretch" the flame: when $Fr_f \gg 1$, the 759 jet's momentum is the dominating factor and when $Fr_f \ll 1$, the flame's buoyancy force is 760 761 the dominating factor. Comparing Fr_f against Ri_L , we understand that both are used to assess the dominating factor between jet-related quantity and buoyancy-related quantity, but there 762 are two differences: (1) the quantity is momentum (\propto velocity) for Fr_f , but TKE (\propto velocity 763 \times velocity) for Ri_L ; (2) the ratios between jet-related quantity and buoyancy-related quantity 764 are reciprocal: $Ri_L \propto$ (buoyancy-related quantity)/(jet-related quantity) whereas $Fr_f \propto$ (jet-765 related quantity)/(buoyancy -related quantity). Therefore, these two parameters are closely 766 related and mathematically their relationship should be $Ri_L \propto Fr_f^{-2}$. Precise relationship 767 between them can be derived from Equations (5) and (9): 768

769
$$Ri_{L} = \left(\frac{\rho_{\infty}}{\rho_{e}}\right)^{3/2} \left(\frac{L}{d_{e}}\right)^{3} \left(\frac{T_{\infty}}{\Delta T_{f}}\right) \cdot f_{s}^{-3} \cdot Fr_{f}^{-2} \qquad (10)$$

It is noted that $\left(\frac{L}{d_e}\right)^3$ can be interpreted as the volume expansion ratio of the flaring gas due to burning and $\left(\frac{T_{\infty}}{\Delta T_f}\right)^{-1}$ can be approximated as the temperature rising ratio of the flaring gas due to burning. Based on the gas law, these two are proportional to each other for an isobaric process (from the exit to the flame tip) which is a good assumption for gas flaring, i.e. $\left(\frac{L}{d_e}\right)^3 \propto \left(\frac{T_{\infty}}{\Delta T_f}\right)^{-1}$, or equivalently, $\left(\frac{L}{d_e}\right)^3 \left(\frac{T_{\infty}}{\Delta T_f}\right) \approx \text{const.}$ Therefore, Equation (10) confirms the relationship of $Ri_L \propto Fr_f^{-2}$. This suggests a strong dependence between adopting Ri_L and adopting Fr_f as the scaling parameter.

The first Damköhler ratio, Da_1 , however, is defined as the ratio of residence time of fuel in flame (τ_{res}) to chemical time of burning process (τ_{chem}):

779
$$Da_1 = \frac{residence\ time\ in\ flame,\ \tau_{res}}{chemical\ time,\ \tau_{chem}} = \frac{\left(\frac{L}{u_e}\right)}{\tau_{chem}}$$
(11)

780 It describes the extent of the oxidation process within the flame in relation to the oxidant's feed rate. For a large Da_1 (i.e. $\tau_{res} \gg \tau_{chem}$), the velocity fluctuating component does not 781 have much influence on the chemistry of the flame. The chemical reaction is able to proceed 782 to completion within the residence time in the combustion zone, resulting in intensive 783 chemical reaction and hot diffusion flame. For a small Da_1 (i.e. $\tau_{chem} \gg \tau_{res}$), turbulence 784 can significantly affect the chemistry and structure of the flame. The rate of chemical reaction 785 and hence, heat release may be affected, causing combustion product to be mixed with 786 reactants within a time interval shorter than the chemical reaction time (Liberman, 2010; 787 William, 1985). From the perspective of processes, Da_1 involves an extra dimension (i.e. 788 chemical processes) which is not reflected by either Ri_L or Fr_f . In principle, we should 789 790 consider Da_1 as one more scaling parameter.

791 **4.5 Soot modelling**

Mathematical modelling is a technique that has been used by scientist and engineers over the years to understand the relationship between sets of input and output parameters in a process, especially where the 'real world' process is often remote or grossly complicated to assess. The region of validity of such model outputs is often limited as several assumptions and constants are applied in the modelling to further simplify the process being studied.

Soot formation and oxidation in pre-mixed and non-premixed (diffusion) hydrocarbon flames 797 have been studied using several modelling techniques including computational fluid 798 dynamics (CFD). The main problem with mathematical modelling of turbulent combustion of 799 hydrocarbon is the problem of modelling turbulent flow and chemical kinetics and the 800 interaction between flow and chemical reaction (Magnussen and Hjertager, 1977). Kennedy 801 (1997) classified the models for soot formation as empirical correlations, semi-empirical 802 803 correlation models and models with detailed chemistry and physics of soot formation. The flame temperature, C:H ratio and number of carbon atoms in the fuel (hydrocarbon) are 804 important parameters considered to have strong influence on the sooting propensity of the 805 hydrocarbon (Harris et al., 1986). These parameters have been the basis of measurements 806 used in most empirical correlation models. 807

808 In their study to modelling soot formation and combustion, Magnussen and Hjertager (1977) assumed that soot is formed from a gaseous hydrocarbon in two stages (a) formation of 809 radical nuclei, and (b) soot particle formation from these nuclei. Applying expressions for the 810 rate of formation of radical nuclei and rate of soot particle formation expressions developed 811 by Tesner et al. (1971a); (Tesner et al., 1971b) in their model, tested on both pre-mixed and 812 813 diffusion flame, they predicted soot concentrations that are in close agreement with experimental data. They concluded that soot formed and was contained in eddies and burned 814 away during turbulence dissipation. 815

Moss et al. (1989), using a two-equation model for the evolution of soot volume fraction and number density, simulated the formation of soot. They included the influences of nucleation, surface growth and coagulation on the rate of soot formation. As given in equations 12(a) and (b), their model contains simplified expressions to quantify particle nucleation, growth and coagulation; using three empirical constants that are dependent on the fuel to control the rate of these processes. A major finding from their study is that soot volume fraction is proportional to the square of pressure.

823
$$\left\{\frac{d(\frac{n}{N_0})}{dt}\right\} = \alpha(\zeta) - \beta(\zeta)(\frac{n}{N_0})^2 \dots 12(a)$$

824
$$\rho_s \left\{ \frac{df_v}{dt} \right\} = \gamma(\zeta)n + \delta(\zeta) \qquad 12(b)$$

where ρ_s is the assumed density for solid carbon (1.8 x 10³ kgm⁻³), N_o is Avogadro's number and ζ is the mixture fraction. Rates of the processes are expected to be a function of the mixture fraction. The rates of the processes are defined explicitly in terms of the fuel density, ρ , temperature. T, and fuel mole fraction, X_c as described in equation 13:

831 coefficients $C_{\alpha,\beta,\gamma,\delta}$ and activation temperatures, T_{α} and T_{γ} are obtained from experimental 832 data. In equations 12(a), the first and second term on the left are nucleation and coagulation 833 processes, respectively. And, in equation 12(b), the first and second term on the left represent 834 the growth and nucleation, respectively.

Lautenberger et al. (2005) developed a CFD model to study the formation and oxidation of 835 soot. Since their model was to generate enough accurate predictions of soot emission 836 concentrations in order to estimate CFD simulations of fire radiation from turbulent flames, 837 838 considerations were given only to phenomena which were essential. Soot estimation in their model was based on a further simplified form of the two-equation model of Moss et al. 839 (1989). There are no fuel-specific constants in their estimations, but rather, they used the 840 841 laminar smoke point height to account for the sooting propensity of different fuels. The smoke point of a flame is its length just before the onset of the release of visible smoke. 842 843 Length of a flame is dependent on the extent of completeness of the combustion and heat content of the fuel gas (Beychok, 1994). 844

The use of CFD simulation soot has made it possible to make predictions about the relationships between the various processes involved in soot formation and oxidation in both pre-mixed and diffusion hydrocarbon flames as well as the sensitivity of the soot formation to some of the complex phenomena. However, the veracity of CFD simulation results in limited by the availability of field measurements for model evaluation.

850 4.6 Gas flaring emissions in global models and inventories

To the best of our knowledge, the only two emission datasets that explicitly includes gas flaring emissions are EDGAR (Emissions Database for Global Atmospheric Research) and ECLIPSE (Evaluating the Climate and Air Quality ImPacts of Short-livEd Pollutants) gridded emissions datasets.

In EDGAR v4.2, for gas venting and flaring, emissions were calculated for 1994 onwards using the amount of gas flared estimated from satellite observations of intensities of light from various gas flares. The estimated quantities of gas flared and emission factors obtained from either inventory guidance documents or confidential information were used to generate gridded annual emission datasets for different countries on a resolution of $0.1^{\circ} \times 0.1^{\circ}$. In 2005, the inclusion of new primary data sources for gas flaring in EDGAR v4.2 gives rise to a change of +75 % of EDGAR 4.1 value (130 Tg) in global CO₂ emission (European Commission, 2009).

ECLIPSE is provided by the IIASA (International Institute for Applied Systems Analysis). 863 Emission calculations for the historical years (2005 - 2010) were developed in a series of 864 regional and global projects. For gas flaring, emissions were calculated using data available 865 from NOAA, NASA and the World Bank collaborative work to estimate the volume of gas 866 flared globally. The volume of gas flared was estimated using NASA MODIS active fire 867 detection products (Elvidge et al., 2007; Elvidge et al., 2011). Emission factors and other 868 parameterizations were obtained from peer-reviewed data on emission performance of 869 various technologies. The calculation was performed with the IIASA GAINS model (Klimont 870 et al., 2013). The ECLIPSE v4 global emission dataset is available on a 0.5° x 0.5° lon-lat 871 resolution. 872

Stohl et al. (2013), performing a 3-year black carbon (BC) simulation in the Arctic, using a Lagrangian particle dispersion model FLEXPART driven by the ECLIPSE dataset, estimated that more than 40 % of mean surface BC surface concentration in the Arctic is attributable to gas flaring. Although, more studies on the sensitivity of global models to gas flaring emission is needed to give a clear and precise quantification of the contribution of gas flaring emission to global aerosol loadings, the change in global CO_2 level in EDGAR4.2 underscores the importance of an explicit inclusion of gas flaring emissions in global models.

880 **5** Conclusion

The World Bank has been at the fore-front of the campaign to reduce gas flaring through the public-private partnership project "Global Gas Flaring Reduction (GGFR)"

(http://www.worldbank.org/en/programs/gasflaringreduction). The "Zero Routine Flaring by 883 2030" initiative was launched in April 2015 by the World Bank, United Nations, 884 governments and oil companies. As at April 17 2015, a total of nine countries have agreed to 885 886 the "Zero Routine Flaring by 2030" initiative. At the time of writing, some major flaring nations have yet to sign up to the initiative. Indeed, some major flaring countries still struggle 887 to meet targets for gas flaring set in the late 1980s. A steep increase in flaring since 2006 has 888 889 been reported for the USA and Canada, associated, presumably, with the exploitation of unconventional hydrocarbon reservoirs. 890

Considering the wide spectrum, quantity and effects of pollutants emitted from gas flaring, coupled with the estimated quantity of gas flared globally, it is surprising that so little effort has been put into adequately understanding the yield of pollutants, especially BC, from the process in real world field situations. The wide variation of fuel gas compositions from flow stations around the world underpins the importance of developing strategies that take these compositions into consideration when estimating emissions.

The steep decrease in the fraction of total gas production flared between 2000 and 2006 seems to have stabilised between 2007 and 2010, and the fraction has even increased in 2011. The overall quantity of gas flared since 2000 has been steady between 93 and 110 bcm. An increase in total production since 2009 has brought about a corresponding increase in the quantity of gas flared. Incentives and stringent policies are not yet in place to encourage more companies and countries to partner with the World Bank in their "Zero Routine Flaring by 2030" initiative.

Elevated concentrations of BC, CO, H_2S , SO₂, NO, NO₂ and PAH measured around flaring sites (ground based and aircraft) is indicative of the detrimental impact of gas flaring on the environment. Clusters of gas flaring sites around the tropics and near-tropic regions of the

world, where there is the likelihood on enhanced atmospheric mixing of the emissions into
the lower and even mid-troposphere, coupled with the high temperature of the emitted plume
suggests the possibility of long-range transport of these pollutants.

Emission factors used for BC emission from gas flaring are inadequate to estimate emission 910 from a typical real-world gas flare as most of the fuels used in the studies for such emission 911 factors are not representative of fuel gas from most Flow Stations around the world. In the 912 studies that employed the mass/mole balancing technique to estimate pollutants from the 913 combustion of hydrocarbons, estimated EFs for CO₂ and CO have been given, but there is no 914 consideration of the amount of unburnt carbon given off. It should be noted that these studies 915 916 did not consider flame dynamics changes in their estimations. When estimating emissions from gas flares, there is the need to ascertain the nature (regime) of the combustion flame as 917 the flame nature and temperature plays pivotal role in determining the pollutants yield, and 918 919 this has not yet been routinely considered in dispersion modelling and global inventories.

Global models need to update the sources of BC to include gas flaring, especially in regions
prone to long-range transfer of gas flaring emission from leading gas flaring nations
including Russia, Nigeria and the Middle East.

923

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931			
932	6	G	lossary
933		1.	Heating value – the heat evolved per unit of a gas when it undergoes complete
934			combustion at standard temperature and pressure. It can be measured in unit mass,
935			unit mole or unit volume.
936		2.	Higher (gross) heating value (HHV) – The heating value when all of the water present
937			in the combustion product in gas phase condenses to liquid.
938		3.	Lower (net) heating value (LHV) - The heating value when all of the water present in
939			the combustion product remains in the gas phase.
940		4.	Volumetric heating value (VHV) – Total heat evolved per unit volume of a gas.
941		5.	Equivalent ratio, ϕ - This is the ratio of the stoichiometric air-fuel mass ratio to the
942			actual air-fuel mass ratio
943		6.	Mtoe – Million tonnes of oil equivalent
944		7.	Laminar smoke point - The length of flame from a gas jet just before the onset of the
945			release of visible smoke.
946			
947			
0.17			
948			
949			

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