1.B.2.c - Venting and Flaring

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Category Code	Method				AD				EF						
1.B.2.c	T2			AS				CS							
Key Category	SO ₂	NOx	ΝНз	NMVOC	CO	ВС	Pb	Hg	Cd	Diox	PAH	нсв	TSP	PM ₁₀	PM ₂ 5
1.B.2.c	-/-	-/-	-	-/-	-/-	-/-	-	-/-	-	-	-	-	-/-	-/-	-/-

Method(s) applied							
D	Default						
T1	Tier 1 / Simple Methodology *						
T2	Tier 2*						
Т3	Tier 3 / Detailed Methodology *						
С	CORINAIR						
CS	Country Specific						
M	Model						
* as described in the EMEP/	EEA Emission Inventory Guidebook - 2019, in category chapters.						
(source for) Activity Data	a						
NS	National Statistics						
RS	Regional Statistics						
IS	International Statistics						
PS	Plant Specific						
As	Associations, business organisations						
Q	specific Questionnaires (or surveys)						
M	Model / Modelled						
С	Confidential						
(source for) Emission Fa	ctors						
D	Default (EMEP Guidebook)						
CS	Country Specific						
PS	Plant Specific						
M	Model / Modelled						
С	Confidential						

Pursuant to general requirements of the Technical Instructions on Air Quality Control TA Luft (2002), gases, steam, hydrogen and hydrogen sulphide released from pressure valves and venting equipment must be collected in a gas-collection system. Wherever possible, gases so collected are burned in process combustion. Where such use is not possible, the gases are piped to a flare. Flares used for flaring of such gases must fulfill at least the requirements for flares for combustion of gases from operational disruptions and from safety valves. For refineries and other types of plants in categories 1.B.2, flares are indispensable safety components. In crude-oil refining, excessive pressures can build up in process systems, for various reasons. Such excessive pressures have to be reduced via safety valves, to prevent tanks and pipelines from bursting. Safety valves release relevant products into pipelines that lead to flares. Flares carry out controlled burning of gases released via excessive pressures. When in place, flare-gas recovery systems liquify the majority of such gases and return them to refining processes or to refinery combustion systems. In the process, more than 99 % of the hydrocarbons in the gases are converted to CO₂ and H₂O. When a plant has such systems in operation, its flarehead will seldom show more than a small pilot flame.

	Unit	1990	1995	2000	2005	2010	2015	2018	2019
Flared natural gas	millions of m ³	36	33	36	18.7	12.1	10.5	10.6	15.6
Refined crude-oil quantity	millions of t	107	96.5	107.6	114.6	95.4	93.4	87.7	87

Flaring takes place in extraction and pumping systems and at refineries. In refineries, flaring operations are subdivided into regular operations and start-up / shut-down operations in connection with disruptions.

Source of emission factor	Substance	Unit	Value
Flaring emissions in natural gas extraction	NMVOC	kg/ 1000 m ³	0.005
Flaring emissions in natural gas extraction	NOx	kg/ 1000 m ³	1.269
Flaring emissions in natural gas extraction	SO ₂	kg/ 1000 m ³	8.885

Source of emission factor	Substa	nce l	Init		Value			
Flaring emissions in natural gas extraction CO			kg/ 1000		0.726			
Source of emission factor	Sub	stance	Unit	Value				
Flaring emissions at petroleum production	NO×		kg/t	0.008				
Flaring emissions at petroleum production facilities				kg/t	0.010			
Flaring emissions at petroleum production	CO		g/t	0.1				
Source of emission factor		Subst	ance	Unit	Valu	ıe		
Flaring emissions at refineries: normal flar	itions	NMV	OC.	kg/m³	0.00)4		
Flaring emissions at refineries: normal flaring opera			CC)	kg/m³	0.00)1	
Flaring emissions at refineries: normal flaring opera			ions SO ₂		kg/m³	0.00)3	
Flaring emissions at refineries: normal flaring opera			NC) _×	g/m³	0.4	-	
Source of emission factor	Sub			stance	Unit	Val	lue	
Flaring emissions at refineries: disruptions	g ope	rations	NMV	OC	kg/t	0.0	01	
Flaring emissions at refineries: disruptions	g ope	rations	СО		kg/t	0.0	01	
Flaring emissions at refineries: disruptions	g ope	rations	SO ₂		kg/t	0.0	07	
Flaring emissions at refineries: disruptions	ope	rations	NO×	·	kg/t	0.0	04	

The emission factors have been derived from the 2004 and 2008 emissions declarations Theloke et al. 2013 $^{1)}$. In 2019, they were updated for CH₄, N₂O, CO, NMVOC, NO_x and SO₂, on the basis of Bender & von Müller, 2019 $^{2)}$. Venting emissions are taken into account in category 1.B.2.b.iii. The SO₂ emissions are obtained from the activity data for the flared natural gas (Table 178) and an emission factor of 0.140 kg / 1,000 m³, a factor which takes account of an average H₂S content of 5 % by volume. The emission factors are determined on the basis of emissions reports, crude-oil-refining capacity and total capacity utilisation at German refineries. The guide for this work consists of the evaluation assessment of Theloke et al. (2013) $^{3)}$.

Recalculations

will be published later - in meantime please refer to chapter 8.1 "recalculations"

Planned improvements

Currently no improvements are planned.

References

^{1), 3)} Theloke, J., Kampffmeyer, T., Kugler, U., Friedrich, R., Schilling, S., Wolf, L., & Springwald, T. (2013). Ermittlung von Emissionsfaktoren und Aktivitätsraten im Bereich IPCC (1996) 1.B.2.a. i-vi - Diffuse Emissionen aus Mineralöl und Mineralölprodukten (Förderkennzeichen 360 16 033). Stuttgart.

²⁾ Bender, M., & von Müller, G. (2019). Konsolidierung der Treibhausgasemissionsberechnungen unter der 2. Verpflichtungsperiode des Kyoto-Protokolls und der neuen Klimaschutz-Berichterstattungs-pflichten an die EU (FKZ 3716 41 107 0).