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1	Estimating the flammability of vapours above refinery waste
2	water laden with hydrocarbons mixtures
3	
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24 Abstract

25

26 In this study, the likelihood of fire hazards attributed to the vaporization of hydrocarbon 27 components derived from refinery wastewater drainage systems was assessed. Liquid samples 28 containing mixtures of hydrocarbon products and water were collected from a refinery drainage 29 system and subjected to a distillation process to separate oil and water. The oil-liquid phase was 30 analyzed using Gas Chromatography Mass Spectrometry (GC-MS) to examine the composition 31 of the sample. Hydrocarbon compounds ranging from C_9 to C_{16} were detected. Mole fractions of 32 28 selected components in the liquid phase were obtained from the GC-FID data and used to 33 calculate mole fractions of components in the gas phase via modified Raoult's law. Lower 34 Flammability Limits (*LFL*s) and Upper Flammability Limits (*UFL*s) for individual components 35 were calculated using a stoichiometric concentration method, while the LFL and UFL values for 36 the mixture (LFL_{mix} and UFL_{mix}) were calculated using the Le Chatelier equation. The LFL_{mix} 37 and UFL_{mix} values were used to construct a flammability diagram and subsequently used to 38 determine the flammability of the mixture. The findings of this study may assist in minimizing 39 fire hazards associated with presence of hydrocarbon vapours derived from refinery wastewater 40 streams.

41

42 Keywords: Gas chromatography; compositions; Lower and Upper Flammability Limits;
43 flammability diagram.

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- 45

47 **1. Introduction**

48

49 Refineries are complex systems of numerous chemical process operations that refine crude 50 oil into desired products such as Liquefied Petroleum Gas (LPG), petrochemical naphtha, motor 51 gasoline, kerosene, diesel and other products. Petroleum refineries can consume high quantities 52 of water for operational usage relative to other industrial and domestic users within a 53 geographical region. Consequently, these refineries can generate large volumes of wastewaters 54 containing various petroleum hydrocarbons, heavy metals, sulfur and ammonia at concentrations 55 that typically require treatment (Al-Haddad et al., 2007) prior to final discharge. With respect to 56 hydrocarbon components, highly flammable compounds such as benzene, toluene, ethylbenzene 57 and xylenes may be present and they pose a significant threat of fire hazard. Over time, the 58 mixture of water and hydrocarbon in drainage systems at certain conditions will naturally 59 separate and form distinct liquid phases, based on density and polarity of the material (EPA, 60 1998). Hydrocarbon compounds from these drainage flows can vary widely in composition from 61 day-to-day due to operational activities such as storage of waste liquids from drains, equipment 62 cleaning and spills. Some of these compounds can evaporate and turn into vapour at ambient 63 temperature and atmospheric pressure, thus creating potentially flammable mixtures in air. The 64 presence of flammable mixtures exposes drainage systems to the possibility of fire and explosion. 65 66

66 The composition of hydrocarbon compounds can be determined and ignitable liquids can be 67 identified (Newman et al, 1997) using comprehensive gas chromatography (GC) systems such as 68 gas chromatography–isotope ratio mass spectrometry (GC–IRMS) and gas chromatography-mass 69 spectrometry (GC-MS). GC analysis can determine the presence, type and concentrations of

70 contaminated hydrocarbons in wastewater (Senn and Johnson, 1987). However, GC analysis of 71 sample compositions is not sufficient to predict the flammability of hydrocarbon mixtures. In this 72 regard, a flammability diagram method can be suitably used for this purpose (Mashuga and 73 Crowl, 1999). Flammability diagrams generally show the "area" of flammability in mixtures of 74 fuel, oxygen and an inert gas. Mixtures of the three gases are usually depicted in a triangular 75 diagram, also known as a Ternary plot. For more detailed information on the ignitability of 76 mixtures and spark ignition, readers are directed to reviews by Mullins (1955), Lewis and von 77 Elbe (1987), and Mullins and Penner (1959).

78 Vapour-air mixtures ignite and burn only over a well-specified range of compositions. The 79 mixture will not burn when the composition is lower than the lower flammable limit (LFL), that 80 is, the mixture is too lean for combustion. The mixture is also not combustible when its 81 composition is too rich, that is, when it is above the upper flammable limit (UFL). As such, a 82 mixture is flammable only when its composition is between LFL and UFL. These flammability 83 limits can be measured experimentally, though they can still be determined without experimental 84 data (Crowl and Louvar, 2002). There are several available methods, databases and software 85 packages that provide sufficient flammability information for various hydrocarbons and they can 86 be sourced from Coward and Jones (1952), Zabetakis (1965), Sax (1984), Kuchta (1985), Ohtani 87 et al. (1994), Brooks and Crowl (2007a) and DIPPR (2010).

The limiting oxygen concentration (LOC) is defined as the minimum oxygen concentration in a mixture of fuel, air and inert gas that will propagate flame and is expressed in volume percent of oxygen (Zlochower and Green, 2009). In essence, LOC varies with pressure and temperature and depends on the type of inert (non-flammable) gas present. A reaction can not generate enough energy to heat the entire gas mixture required for the self-propagation of the

93	flame if the oxygen concentration is below the LOC (Crowl and Louvar, 2002). As such, the
94	LOC is a useful parameter in terms of fire hazard prevention since explosions and fires can be
95	prevented by reducing the oxygen concentration regardless of fuel concentration. This concept is
96	the basis for a common procedure called <i>inerting</i> used in safety engineering (Crowl and Louvar
97	2002; ASTM 2009). The LOC can be measured experimentally using a flammability apparatus
98	(British and European Standard, 2007; ASTM 2008a; ASTM 2008b) or it can be found from
99	different resources (NFPA 1994; CHEMSAFE, 2007). An example of experimental LFL and
100	LOC determination can be found from a study conducted by Brooks and Crowl (2007b), in
101	which they experimentally measured the LFL, LOC and the maximum safe solvent concentration
102	(MSSC) for ethanol and acetonitrile above aqueous solutions. If experimental and/or literature
103	data are not available, the LOC can be estimated using the stoichiometric combustion reaction
104	and the LFL. This procedure works relatively well for many different hydrocarbons (Siwek,
105	1996; Crowl and Louvar, 2002).
106	The objectives of this study were to estimate flammability limits and the LOC of the
107	hydrocarbon vapour mixture above refinery wastewater and subsequently determine the
108	flammability of the mixture. This was achieved by incorporating thermodynamics with process
109	safety concepts. Findings from this study can be used to investigate the root cause of fire
110	incidents in drainage systems due to the presence of flammable hydrocarbon vapour mixtures.
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116 **2.** Methodology

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- 118 2.1.1 Liquid Phase
- 119

136

120 Liquid samples were collected from drainage lines in a refinery located in Malaysia. 121 Residual water was removed from the samples using a simple distillation technique. The 122 composition of the organic-phase liquid was analysed using both GC-MS (for identification) and 123 GC-FID (Flame Ionisation Detection) (for quantitation). GC-MS was performed with a 124 Shimadzu QP5050 GCMS using the following settings: Electron impact ionization, electron 125 energy 70 eV and scan range 40 - 500 amu at 1 scan/s. The carrier gas (Helium 99.999%) flow 126 rate was set to 1.5 ml/min with column inlet pressure 54.8 kPa and linear velocity 36.10 cm/sec. 127 The end of the column was directly introduced into the ion source of a mass selective detector 128 operated in an electron impact ionization mode. Samples were injected into a HP5 fused silica 129 (5% phenyl polysilphenylene-siloxane) capillary column BPX5 (30 m length; 0.25 mm i.d.; 0.25 130 μ m film thickness) and the oven temperature was held at 55° C for 2 min. It was then increased to 131 300°C at 5°C/min and thereafter held for 40 minutes. The components of the liquid sample were 132 identified by comparing their mass spectra with the NIST Mass Spectral Database. Pure samples 133 of a selected number of compounds were analysed using the same GCMS conditions in order to 134 verify the match from the database. 135 Quantitative analysis of the peaks was performed with a Shimadzu GC2010 with an FID using

137 The mass fraction of each component in the liquid phase can be determined from GC-FID data138 (Eq. 1):

the same column and temperature parameters as for the GC-MS analysis.

139
$$\chi_i = \frac{A_i}{A_T} \tag{1}$$

140 where, χ_i is the mass fraction of component *i*, A_i is the peak area of component *i*, and A_T is the 141 peak area of all components.

142

143 The mass fraction is converted to mole fraction using Eq. (2):

144
$$x_i = \frac{\chi_i / M_i}{\sum \chi_i / M_i}$$
(2)

145 where, x_i is the mole fraction in liquid phase of component *i*, and M_i is the molecular weight of 146 component *i*.

148 2.1.2 Vapour Phase

149

150 It was necessary to estimate the concentrations of components in the gas phase, which also 151 contributed to the flammability of the mixture. Modified Raoult's law was used to estimate the 152 amount of liquid vaporized at ambient temperature; Eq. (3) was used to calculate the mole 153 fraction in the gas phase.

154
$$\gamma_i x_i P_i^{sat} = \varphi_i y_i P_t$$
 (3)

155 where γ_i is the activity coefficient for component *i*, x_i is the mole fraction of in the liquid

156 phase, P_i^{sat} is the vapour pressure of compound *i* as a pure component, φ_i is the fugacity

157 coefficient for component *i*, y_i is the mole fraction of component *i* in the vapour phase, and P_i is

158 the total pressure. Theoretically, the activity coefficient, γ_i , for an ideal solution is equal to 1.

159 However, since the mixture is non-ideal (real), the activity coefficient was calculated using the

160 UNIversal Functional Activity Coefficient (UNIFAC) method. The UNIFAC method expresses

161 the activity coefficient as the sum of a combinatorial part, $\ln \gamma_i^c$ and a residual part, $\ln \gamma_i^R$ (Eq.

162 4) (Fredenslund et al., 1975):

163
$$\ln \gamma_i = \ln \gamma_i^C + \ln \gamma_i^R$$
(4)

164 $\ln \gamma_i^C$ is given by Eq. (5):

165
$$\ln \gamma_i^C = \ln \frac{\Phi_i}{x_i} + \frac{z}{2} q_i \cdot \ln \frac{\theta_i}{\Phi_i} + l_i + \frac{\Phi_i}{x_i} \sum_{j=1}^M x_j l_j$$
 (5)

166
$$l_j \equiv \frac{z}{2} (r_j - q_j) - (r_j - 1)$$
 (6)

167 Where z is the average number of nearest neighbours around a group in solution (constant value

168 is used: z = 10). The segment fraction, Φ_i , and surface area fraction, θ_i , are defined, respectively

169 by Eqs. (7) and (8):

170
$$\Phi_i = \frac{r_i x_i}{\sum_{j=1}^M r_j x_j}$$
(7)

171
$$\theta_i = \frac{q_i x_i}{\sum_{k=1}^{M} q_j x_j}$$
(8)

172 The molecular volume, r_j , is defined by the sum of its constituent group given by Eq. (9):

173
$$r_j = \sum_{k=1}^{N} v_k^j \cdot R_k$$
 (9)

174 Where v_k^j is the number of k groups in molecule j, and R_k is the volume of group k. The

molecular surface area, q_j , is the sum of the individual group areas in the molecules given by Eq. (10):

177
$$q_{j} = \sum_{k=1}^{N} v_{k}^{j} . Q_{k}$$
(10)

178 where Q_k is the group surface area.

179 R_k and Q_k are obtained from the van der Waals group volumes and surface areas.

180 $\ln \gamma_i^R$ was calculated from Eq. (11):

181
$$\ln \gamma_i^R = \sum_{k=l}^N v_k^i \left[ln \Gamma_k - ln \Gamma_k^i \right]$$
(11)

where Γ_k^i is the group residual activity coefficient of group *k* in a reference solution containing only molecules of type *i*. Γ_k is the group residual activity coefficient in the solution. The coefficients Γ_k^i and Γ_k are related to the composition and temperature according to Eq. (12):

185
$$\ln\Gamma_{k} = Q_{k} \left\{ I - ln \left(\sum_{m=1}^{N} \Theta_{m} \cdot \psi_{mk} \right) - \sum_{m=1}^{N} \left[\frac{\Theta_{m} \cdot \psi_{km}}{\sum_{n=1}^{N} \Theta_{n} \cdot \psi_{nm}} \right] \right\}$$
(12)

186 In Eq. (12), the group interaction parameter, ψ_{mk} is defined by Eq. (13):

$$187 \qquad \Psi_{mk} = e^{-\frac{a_{mk}}{T}}, \quad a_{mk} \neq a_{km}$$
(13)

188 where a_{mk} is the group interaction parameter between groups *n* and *m*. The surface contribution,

189 Θ_m and the mole fraction of the group, X_m are defined by Eqs. (14) and (15), respectively:

190
$$\Theta_m \equiv \frac{Q_m \cdot X_m}{\sum_{n=1}^N Q_n \cdot X_n}$$
(14)

191
$$X_{m} \equiv \frac{\sum_{j=1}^{M} v_{m}^{j} \cdot x_{j}}{\sum_{j=1}^{M} \sum_{n=1}^{N} v_{n}^{j} \cdot x_{j}}$$
(15)

192 The fugacity coefficient, φ_i , for each component in the mixture was determined using the Peng-

193 Robinson method (Eq. (16) (Peng and Robinson, 1976):

194
$$ln\varphi_{i} = \frac{b_{k}}{b}(Z-I) - ln(Z-B) - \frac{A}{2\sqrt{2B}} \times \left(\frac{2\sum_{i}^{N} x_{i}a_{ik}}{a} - \frac{b_{k}}{b}\right) ln\left(\frac{Z+2.414B}{Z-2.414B}\right)$$
(16)

where φ_i is the fugacity coefficient, *b* is van der Waals co-volume defined by Eq. (17), *Z* is the compressibility factor defined by Eq. (18), *B* is a constant defined by Eq. (19), *A* is a constant defined by Eq. (20), x_i is the mole fraction, and *a* is the attraction parameter defined by Eq. (21):

$$199 b = \sum_{i}^{N} x_i b_i (17)$$

$$200 \qquad Z = \frac{Pv}{RT} \tag{18}$$

$$201 \qquad B = \frac{bP}{RT} \tag{19}$$

$$202 \qquad A = \frac{aP}{R^2 T^2} \tag{20}$$

203
$$a = \sum_{i}^{N} \sum_{j}^{N} x_{i} x_{j} a_{ij}$$
(21)

where *P* is the pressure defined by Eq. (22), *T* is the temperature, *v* is the molar volume, *R* is the universal gas constant, a_{ij} is defined by Eq. (23), and *i*, *j*, *k* are component identifications.

206
$$P = \frac{RT}{v-b} - \frac{a(T)}{v^2 + 2bv - b^2}$$
(22)

207
$$a_{ij} = (1 - \delta_{ij}) a_i^{1/2} a_j^{1/2}$$
 (23)

208 where, δ_{ij} is an empirically determined binary interaction coefficient for components *i* and *j*.

209 Applying Eq. (24) at the critical point, we obtain:

210
$$a(T_c) = 0.45724 \frac{(RT_c)^2}{P_c}$$
 (24)

211
$$b(T_c) = 0.07780 \frac{RT_c}{P_c}$$
 (25)

212 $Z_c = 0.307$

213 At temperatures other than the critical:

214
$$a(T) = a(T_c).a(T_r,\omega)$$
(26)

$$215 \qquad b(T) = b(T_c) \tag{27}$$

216 where $a(T_r, \omega)$ is a dimensionless function of reduced temperature and acentric factor, ω and

- 217 equals unity at the critical temperature.
- 218 The total pressure for the mixture can be calculated from Eq. (28):

219
$$P_t = \sum \gamma_i x_i p_i^{sat}$$
(28)

The vapour pressures of the components were calculated using the classic (Eq. 29) and extendedAntoine equations (Eq. 30):

222
$$\log_{10} P^{sat} = A - \frac{B}{C+T}$$
 (29)

223
$$\log_{10} P^{sat} = A - \frac{B}{T} + C \log_{10} T + DT + ET^2$$
 (30)

where

225 *P*^{sat} is the vapour pressure (mmHg) while *A*, *B*, *C*, and *D* are the component-specific constants.

These constants were obtained from several sources (Wichterle and Linek, 1971; Yaws, 1992;

227 Dykyj et al., 1999; Yaws et al., 2009; www.chemspider.com).

228 2.2.1 *LFL and UFL*

For some situations, it may be necessary to estimate the flammability limits without

230 experimental studies. Jones (1938) found that for many hydrocarbon vapours, the LFL and UFL

231 were functions of the stoichiometric concentration of fuel (C_{st}) (Eq. 31 and Eq. 32):

232
$$LFL = 0.55C_{st}$$
 (31)

233
$$LFL = 3.5C_{st}$$
 (33)

234 The stoichiometric concentration, C_{st} for organic compounds was determined using the general

combustion reaction (Eq. 33).

236
$$C_m H_x O_y + z O_2 \rightarrow m C O_2 + \left(\frac{x}{2}\right) H_2 O$$
 (33)

237 where z is equivalent moles O_2 /moles fuel which can be determined from Eq. (34):

238
$$z = m + \frac{x}{4} - \frac{y}{2}$$
 (34)

239 The stoichiometric concentration, C_{st} can be determined as a function of z by Eq. (35):

240
$$C_{st} = [\text{moles fuel/ (moles fuel + moles air)}] \times 100 = \frac{100}{[1 + (z/0.21)]}$$
 (35)

241 Substituting Eq. (34) into Eq. (35) and applying Eq. (31) and Eq. (32) yield Eqs. (36) and (37):

242
$$LFL(vol \%) = \frac{0.55(100)}{4.76m + 1.19x - 2.38y + 1}$$
 (36)

243
$$UFL(vol\%) = \frac{3.50(100)}{4.76m + 1.19x - 2.38y + 1}$$
 (37)

- 244 where *LFL* and *UFL* are the lower and upper flammable limits, respectively.
- 245 The LFL and UFL for mixtures (LFL_{mix} and UFL_{mix}) were calculated using the Le Chatelier
- equation (Eqs. 38 and 39) (Le Chatelier, 1891):

247
$$LFL_{mix}(vol\%) = \frac{1}{\sum(y_i/LFL_i)}$$
 (38)

248
$$UFL_{mix} (vol\%) = \frac{l}{\sum(y_i/UFL_i)}$$
(39)

where LFL_i and UFL_i are the lower and upper flammable limits, respectively, for component *i* in fuel and air while *n* represents the number of combustible species.

251

253

LOC can be estimated using the stoichiometry of the combustion reaction and the *LFL* (Crowl and Louvar, 2002). Eq. (40) can also be used to estimate LOC for a vapour mixture

256 (Zlochower and Green, 2009):

257
$$LOC_{mix} = \sum y_i R_i / \sum y_i / L_i^* = \sum y_i R_i / \sum y_i R_i / LOC_i$$
 (40)

$$258 \qquad L_i^* = LOC_i / R_i \tag{41}$$

259 where LOC_{mix} is the limiting oxygen concentration for the vapour mixture (vol%), R_i is the

- stoichiometric molar ratio of oxygen to compound i in the vapour phase, and LOC_i is the
- 261 limiting oxygen concentration for individual compound (vol%). Figure 1 shows the methodology
- 262 outline adopted in this study.
- 263
- 264
- 265
- 266
- 267

268 **3.** Results and Discussion

269

270 3.1 Compositions

271

272 The components were identified using a combination of the retention time, comparison of 273 mass spectrum derived from GC-MS analysis of the hydrocarbon mixture to a mass spectral 274 library, and comparison to known standards. Figure 2 shows the retention time and peak 275 abundance data for 77 components detected in the liquid phase via GC-MS analysis. Considering 276 the difficulties of relying solely on mass spectral database matching for the identification of 277 compounds that are very similar in molecular structure, it was felt prudent to take the following 278 precautions in deciding which of the 77 individual compounds observed in the gas 279 chromatogram should be used in calculations. Firstly, using pure compounds that were readily 280 available it was ascertained that peaks 3, 14, 41, 52, 75, and 77 were simple straight chain 281 hydrocarbons (Table S1 - electronic Supplementary material) and that the compound ID from the 282 NIST database matched with the compounds injected. These results confirmed that the output 283 from the mass spectral comparison was valid, particularly for compounds with higher than 95% 284 match. Subsequently, all compounds with a minimum 95% database match were considered to 285 have been correctly identified. As a check on this assumption it was noted that the database 286 identification of peaks 28, 42, 50, and 63 were for simple straight chain alkanes and produced a 287 homologous series with the positively identified compounds. Additionally, it was noted that the 288 difference in retention times between these compounds, running from nonane to hexadecane, 289 were consistent providing further evidence that these were compounds of a homologous 290 hydrocarbon series. There are 28 compounds (Table S1- also contains the properties relevant for

291 this study) that meet the above criteria and their order of elution based on their boiling points, 292 (i.e. those with higher boiling points elute later) is also consistent with expected results. These 293 selected 28 compounds cover all the major identified peaks (including numbers 42, 50, 63) and 294 account for a 56% of the total area in the chromatogram (as determined from GC-FID data). 295 The identification of individual components shows that the liquid sample contains a large 296 number of hydrocarbon components ranging from C_9 to C_{16} . The predominant hydrocarbon 297 groups are alkane and alkene with a relatively minor presence of arenes. The peak numbers 14, 298 28, 41, 52, 65, and 75 are assigned to decane, undecane, dodecane, tridecane, tetradecane, and 299 pentadecane, respectively, and these six components constitute more than 43% of the total 300 chromatogram area,. The mass fraction of each component was estimated by dividing the GC-301 FID peak area of each component by the total peak area. The mass fraction was then converted to 302 mole fraction according to Eq. (2) and the resulting distribution for the mole fraction for each 303 component x_i in the liquid phase is shown in Figure 3.

304

305 3.2 Mole Fraction in Vapour Phase

306

The activity and fugacity coefficients for all components in the vapour mixture are calculated and the values shown in Table S1. The average activity and fugacity coefficients for the vapour mixture were determined to be 1.02 and 0.88 respectively. Figure 4 shows the mole fraction for each component in the vapour phase, y_i based on combustible basis. Interestingly, the values of y_i for most components are small; possibly due to the low volatility of the heavy hydrocarbon components detected in the liquid phase. The low volatilities of the components were due to the low vapour pressure, which range from 5×10⁻³ mmHg to 6.74 mmHg. The total

314	mole 1	raction of the vapour mixture is 0.12 vol %, while the air content in the mixture is 99.88
315	vol%.	Therefore, the percentages of N_2 and O_2 in the mixture are approximated to be 79 vol%
316	and 21	vol%, respectively.
317		
318	3.3	LFL, UFL, and LOC

319

214

The calculated *LFL* and *UFL* as well as *LOC* values are listed in Table S1. The calculated *LFL* and *UFL* values are in agreement with those obtained from other databases and open literatures (Figures 5a and 5b). It should be noted that not all *LFL* and *UFL* values could be found in other database and published literatures and some of these values were not indicated in Table S1. The *LFL_{mix}* and *UFL_{mix}* are determined to be 0.72 vol% and 4.60 vol%, respectively while the *LOC_{mix}* value is 11.40 vol%.

326

327 3.4 Flammability Diagram

1 C C C I

328

329 A flammability diagram is a conventional method used to assess the flammability of mixture 330 of gases. The flammability diagram is represented by three axes, namely, (1) fuel (hydrocarbon 331 vapour mixture), (2) inert material, and (3) oxygen. In order to plot the flammability diagram, 332 concentrations of the fuel, oxygen, and inert material (in volume or mole %) are required. The air 333 line is plotted by taking the compositions of air from Table S1 (78.91 % nitrogen and 20.98 % oxygen). The intersection of the stoichiometric line with the oxygen axis is given by 100(z/1+z)334 (Crowl and Louvar, 2002). The LOC_{mix} line can be drawn by locating the LOC_{mix} value (11.40%) 335 336 on the air axis and then drawing a parallel line until it intersects with the stoichiometric line. To

337	construct the flammability zone, the values of LFL_{mix} and UFL_{mix} are required and they are					
338	located on the air line while the flammability zone is the area to the right of the air line. Figure 6					
339	represents the triangular flammability diagram for the hydrocarbon mixture. It can be clearly					
340	seen that the stoichiometric line crosses the flammable zone. Therefore, it can be inferred that the					
341	vapour mixture is flammable.					
342						
343	4. Conclusions					
344						
345	In this study, theoretical work including thermodynamic fundamentals and flammability					
346	calculations were applied to estimate the flammability limits of hydrocarbon vapours derived					
347	from refinery wastewater. Detailed inspection of the customised flammability diagram showed					
348	that the vapour mixture derived from the wastewater was flammable and may pose a potential					
349	fire hazard. As such, it is always prudent for refinery workers to exercise caution when handling					
350	wastewater laden with hydrocarbon residuals. Findings from this study afford a useful basis for					
351	safety officers and engineers from other refineries to assess the potential fire hazards associated					
352	with hydrocarbon vapours derived from wastewater streams.					
353						
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473	Figure	Captions

- **Fig.1.** Flowchart on the methodology used in this study.
- **Fig. 2**. GC analysis for the liquid sample.
- **Fig. 3**. Percentages of hydrocarbon mole fractions in the liquid phase.
- **Fig. 4**. Percentages of hydrocarbon mole fractions in the vapour phase.
- **Fig. 5a&b.** Comparison between calculated and literature values of *LFL and UFL*.
- **Fig. 6**. Triangular flammability diagram of the hydrocarbon mixture.







Fig. 3. Mole fractions of the selected 28 hydrocarbon components in the liquid phase.



Fig. 4. Mole fractions of the selected 28 hydrocarbon components in the vapour phase.





`Fig. 5a &b. Comparison between calculated and literature values of *LFL* and *UFL*.



Fig. 6. Triangular flammability diagram of the hydrocarbon mixture.