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1 A novel approach for quantitative measurements of preferential 2 evaporation of fuel by means of two-tracer laser induced 3 fluorescence

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6 Abstract

7 This paper presents a novel approach for the measurement of vapor fuel concentration at engine relevant
8 conditions for the investigation of preferential evaporation in multicomponent fuel GDI sprays.

9 The result is achieved by combining in synergy two different tools: i) two-tracer laser induced fluorescence
10 (2T-LIF), a laser diagnostics capable of investigating qualitatively the preferential evaporation by the
11 simultaneous employ of two fluorescent tracers and, ii) a numerical model capable of relating the tracers
12 concentration to the vapor fuel composition.

13 In order to make this approach reliable for future spray application, this paper presents in details the
14 methodology, including the definition of the surrogate fuel and the tracers. Moreover, results of a fuel
15 evaporation experiment in a simplified environment are presented to validate the numerical model and to
16 assess the uncertainties associated to the technique in spray applications.

17 The evaporation experiment includes parametric variations of the effect of system pressure (1-5-10 bar),
18 and ethanol concentration (0%, 20% and 85% vol.), exploring problematic relevant to real GDI engines.

19 The experimental results provided also fundamental understanding of the preferential evaporation
20 process.

21 Ethanol is observed to form an azeotrope with lighter compounds segregating the evaporation of the
22 heaviest compounds at the end of the evaporation process.

23 Finally, the systematic error related to the calculation of vapor phase composition related to the
24 thermodynamic model approximations was assessed for GDI relevant application. The results indicated a
25 very low impact on the final results and discrepancies lower than 5%.

26 **Keywords:** *GDI, Preferential evaporation, Multicomponent fuel, 2-tracer LIF*

27 **Highlights:**

- 28 1) Novel approach to measure preferential evaporation in GDI sprays
- 29 2) Fuel vapor composition can be assessed experimentally basing on 2T-LIF
- 30 3) A thermodynamic surrogate for commercial gasoline is found
- 31 4) Preferential evaporation is studied at different ethanol content and evaporation pressures

32 **1 Introduction**

33 A highly competitive market and stringent regulations on the pollutant emissions lead engine
34 manufacturers to push to the limit the fundamental research on the in-cylinder related processes [1].

35 During the last two decades, direct fuel injection strategies (DI) have become of common use in
36 compression ignition engines (CI) and have begun to be a reference also for spark igniting engines (SI)
37 [2,3].

38 DI, in general, enhances the control on fuel mixing and combustion and enables the implementation of
39 new combustion modes, e.g. LTC and multi injections in CI engines [4,5] and stratified combustion in SI
40 engines [6]. DI involves several complex physical phenomena including liquid atomization, multiphase flow
41 and multicomponent fuel evaporation [3,7].

42 A real fuel, which is composed by hundreds of compounds of different volatility, can evaporate in a
43 specific fashion normally referred as preferential evaporation. Preferential evaporation takes place when
44 the evaporation process is considered to be slow compared to the mass diffusion of the components
45 within the droplet. This phenomenon causes the lighter compounds to evaporate at different timing than
46 the heavier compounds, and, depending on the mixing process, heterogeneities in the resulting mixture
47 composition might be found. [Different studies on GDI engines indicated preferential evaporation as a
48 possible source of heterogeneities in the composition of the resulting air-fuel mixture, by testing cases
49 typical of early or late injection phasing \[8–10\].](#) The consequence of preferential evaporation can also be
50 reflected on the combustion process [11], these heterogeneities might result in un-expected / un-wanted
51 effects such as abnormal combustions, or misfires. Few authors carried out experimental studies on GDI
52 sprays evaporation process. In particular, Itani et al. [10] showed experimentally how preferential
53 evaporation leads eventually to heterogeneities in the composition of a vaporized GDI spray under
54 condition typical of stratified combustion strategies.

55 Other authors investigated the effects of preferential evaporation computationally. Zigan et al. [9,12]
56 demonstrated how the evaporation of the droplets is strongly affected by preferential evaporation, and the
57 multicomponent fluid phase change has to be carefully taken into account to have correct predictions of
58 the final fuel vapor distribution [9].

59 The introduction in the market of new oxygenated fuels also represents a new challenge for engine
60 manufacturers. One of the reasons is that the different physical properties of these new fuels [13] yield
61 modifications in the final mixture which are not correctly understood [14]. Storch et al indicated that,
62 physical properties of ethanol blended fuels (E20 and E85) can cause important modifications in the spray
63 formation and in the combustion behavior [15].

64 Currently, CFD models used for engine development neglects this phenomenon due to the complexity of
65 the physics involved and the lack of quantitative validation data. The experimental investigations available
66 in the literature show only qualitative measurements. To in-cylinder enhance CFD simulations
67 development it is of primary importance to develop quantitative diagnostics that can provide validation
68 data.

69 The 2T-LIF diagnostic is a technique that has been used in previous studies to investigate preferential
70 evaporation. Krämer et al. employed four different fluorescent tracers (acetone, toluene, p-xylene and 3-
71 pentanone) to characterize 3 different volatility compounds of gasoline (low, medium and high) [16]. In
72 their work, they obtained information about the different fuel volatility class in a port fuel engine case by
73 imaging, on different experiments, different couples of tracers. Their results finally indicated
74 heterogeneities in the tracers' distributions and served as a *qualitative* evidence of heterogeneities in the
75 fuel vapor composition [10]. However, the current need of engine manufacturers in developing predictive
76 CFD models highlights the need of developing a technique to provide validation quantitative data about
77 this important phenomenon, which are currently not available in the literature.

78
79 In this work, a novel approach to perform quantitative measurements of preferential evaporation in GDI
80 sprays at real engine conditions is proposed by combining experimental and computational
81 methodologies: an optical diagnostic, 2-Tracer Laser Induced Fluorescence (2T-LIF), and a
82 thermodynamic model for the multi-component fuel evaporation.

83 The goal of this paper is not to presents experiments on GDI sprays, but it brings important contributions
84 to prove the reliability of the technique for future GDI applications. In particular:

- 85 1. It proposes a methodology to obtaine quantitative information of preferential evaporation in GDI
86 sprays. This methodology relies on the combination of 2T-LIF results and a thermodynamic

87 model based on Predictive Soave-Redlich-Kwong state equation. This approach enable to link
88 the tracer concentration measurements to the vapor fuel composition.

89 2. It provides validation data for the thermodynamic model, by performing evaporation experiments
90 in a simplified environment (a bubbler cell). These experiments, a part from providing validation
91 data, help understanding the effect of boundary conditions on preferential evaporation. In
92 particular, system pressure was varied matching values relevant for GDI engine application (1, 5
93 and 10 bar). Also, fuel ethanol content effect was tested (0, 20 and 85 % (vol.)) with the double
94 objective of investigating real world present and future fuels and observing the impact on the fuel
95 evaporation of the addition of a compound with different evaporating properties (ethanol) [17].

96 3. Finally, the accuracy of the methodology in measuring the vapor fuel composition is assessed.
97 More specifically, the impact of the discrepancies between thermodynamic model results and
98 experimental measurements on the estimation of the fuel vapor composition is presented. This
99 analysis is considered a fair estimation of the accuracy of the technique in GDI applications.

100 This paper is divided in five sections including the present introduction. One is devoted to the description
101 of the approach, indicating the fuel surrogate composition and the tracers to be used, presenting the
102 details of the thermodynamic models, and the calculations performed to compute the fuel vapor
103 composition. The following section focuses on the description of the experimental setup of the evaporation
104 experiment. The fourth section presents the evaporation experiment results and compares them to the
105 results provided computationally by the thermodynamic model. Also an assessment of the accuracy of the
106 quantitative 2T-LIF is performed on the bases of a deviation analysis. The main conclusions of the paper
107 are then summarized in the final section.

108 **2 Quantitative interpretation of 2T-LIF results**

109 **2.1 Global methodology**

110 2T-LIF is an optical diagnostic capable of investigating the preferential evaporation phenomena in fuel
111 vapor. The technique relies on the simultaneous imaging of the fluorescence signal emitted by two tracers

112 of different volatility added to a surrogate, optically transparent fuel. Recent studies demonstrated its
113 applicability to GDI sprays at engine relevant conditions [18]. Similarly to 1-tracer LIF techniques, a laser
114 sheet is used to cause the fluorescence of the tracers. However, in this case, the light emissions are
115 collected by two cameras which, by means of specific spectral filters, provide 2D information about the
116 fluorescent signal of each tracer separately. As for 1-tracer LIF, by correctly taking into account the photo-
117 physics of the tracer, it is possible to establish a relationship between the fluorescent signal and the local
118 mass fraction of the tracer [19,20].

119 The objective of this paper is to define a methodology to provide quantitative data of fuel preferential
120 evaporation in realistic GDI conditions.

121 To this end, a computational thermodynamic model is coupled to the 2T-LIF to link the tracer
122 concentration to the local fuel composition. In this section, the methodology is described from a
123 macroscopic point of view presenting the reasons behind the methodology proposed.

124

125 In order to provide a graphic description of the approach proposed, a diagram is presented in Figure 1.

126 As a first step, to carry out 2T-LIF is necessary to choose carefully the surrogate fuel and the fluorescent
127 tracers:

- 128 • *The surrogate fuel* has to be, on the one hand, optically transparent to assure that only the
129 tracers' fluorescence is collected; on the other, the distillation curve has to be as close as possible
130 to the one of a commercial gasoline.
- 131 • *The tracers* at the same time should have suitable fluorescence properties, in particular to have
132 well separated emission spectra and they have to be representative of different distilled fraction of
133 the fuel (e.g. one should be representative of the heavy compounds and the other of the light
134 compounds).

135 Once the fuel and the tracers are correctly selected, 2T-LIF can provide separately each tracer's local
136 mass fraction, and also the tracers' concentration ratio (T_R). The key point of this approach is the employ
137 of a *thermodynamic model* which is used to calculate the phase change process of a multicomponent fuel
138 taking into account the co-evaporation between compounds of different volatility.

139 However, to link the 2T-LIF results to the fuel vapor composition, thanks to the thermodynamic model,
140 some hypotheses about the droplet evaporations are needed, and will be presented later in the paper.

141

142 In this paper, the thermodynamic model is applied to fulfill three different tasks:

- 143 1. To define the gasoline surrogate composition needed for the optical experiments. The comparison
144 between the distillation curve provided by the model and the one provided by the experiments will
145 also provide a first preliminary validation of the thermodynamic model.
- 146 2. For its validation: an experiment is put in place to produce the evaporation of a multicomponent
147 fuel sample in well controlled conditions. The slow evaporation process characterizing this
148 experiment, stresses the preferential evaporation phenomenon: the comparison between the
149 experimental results and the thermodynamic model predictions represents the main validation of
150 the thermodynamic model and constitute a main outcome of this paper.
- 151 3. For the assessment of the uncertainties related to the vapor composition measurements in GDI
152 sprays: in this case, the typical tracers concentration range found in previous GDI measurements
153 [10] is used as an input in the model to assess the related vapor fuel compositions. The
154 discrepancies between model and experiments observed during the validation described in the
155 previous point are introduced to assess the impact of these differences on the fuel vapor
156 composition estimation.

157 However, it is important to underline that for each of the three applications of the thermodynamic model,
158 different hypotheses are needed to couple the model to the environment. These hypotheses, which have
159 a significant impact mostly on the second and the third case, will be detailed later on in the paper.

160 **2.2 Thermodynamic model**

161 In this section, the thermodynamic model employed to describe the phase behavior of the multicomponent
162 fuel is presented together with the numerical scheme employed for the calculations.

163 The thermodynamic model used is based on the Predictive Soave-Redlich-Kwong (PSRK) state equation.

164 The PSRK EoS is an enhancement of the classical SRK cubic state equation [21]. This approach was

165 proposed by Holderbaund and Gmehling in 1991 and it showed to be adapted to accurately describe
166 complex system, such as strongly non-ideal mixtures (i.e. ethanol blended fuels).

167
168 In this model, the pressure P is connected to volume v and temperature T by the classic expression of
169 SRK equation:

$$P = \frac{RT}{v-b} - \frac{a}{v(v+b)} \quad (1.)$$

170 Where a is a parameter associated to attractive interactions and b is co-volume which is the smallest
171 accessible volume. R is the universal gas constant. Equation (1) is directly used for pure compound if
172 parameters a and b are defined. For a mixture containing different components, the parameters of
173 equation of state can be calculated using the parameters of each pure component combined by the
174 following mixing rules.

$$\frac{a}{bRT} = \sum_i x_i \frac{a_i}{b_i RT} + \frac{1}{q_1} \left[\frac{G^{ex}}{RT} + \sum_i x_i \ln \left(\frac{b}{b_i} \right) \right] \quad (2.)$$

175

$$b = \sum_i x_i b_i \quad (3.)$$

176
177 Where x_i is the molar fraction of component i . The co-volume of the mixture is directly a linear combination
178 of the co-volume of each compounds and the attractive term is calculated from a mixing rules based on
179 Gibbs free energy (G^{ex}). q_1 is a parameter function of the equation of state and its value is equal to -
180 0.64663. The Gibbs free energy used in the $PSRK$ equation of state is the UNIFAC model which is based
181 on a group contribution method and gives a predictive character to this equation.

182
183 The pure component parameters a_i and b_i are only dependent on critical properties and specific properties
184 of the species according the following expressions:

185

$$a_i = 0.42748 \frac{R^2 T_{ci}^2}{P_{ci}} \alpha(T) \quad (4.)$$

186

$$b_i = 0.086641 \frac{RT_{ci}}{P_{ci}} \quad (5.)$$

187 Where T_{ci} and P_{ci} are respectively the critical temperature and the critical pressure of component i .

188 The function $\alpha(T)$ is given by:

189

$$\alpha(T) = T_r^{C_3(C_2-1)} \exp[C_1(1 - T_r)]^{C_2C_3} \quad (6.)$$

190 Where T_r is the reduced temperature (T/T_{ci}). C_1 , C_2 and C_3 are specific parameters which are determined
191 empirically in order to well fit the vapor pressure of pure compound.

192 If the pure compounds are well defined, the equation should be used in a predictive way. The equation of
193 state is used to calculate the phase behavior of the fluid using equilibrium calculation (flash).

194 **2.3 Fuel, tracers and filters selection**

195 For the correct application of the 2T-LIF the fuel composition as well as the tracers employed has to be
196 carefully selected. The requirement of the technique and the choices performed are detailed in the
197 following sections.

198 **2.3.1 Fuel surrogate definition**

199 As mentioned in the introduction, the fuel employed has to be transparent to the laser wavelength and
200 non-fluorescent: commercial gasoline cannot be employed and a specific fuel surrogate is needed. Also,
201 the surrogate fuel should be representative of a real gasoline evaporation having a similar distillation curve
202 in order to maintain the focus of the study on real engine application. Therefore, a specific fuel physical
203 surrogate has to be defined.

204 A simplified discrete model for petroleum-derived fuel is proposed to permit to track experimentally and
205 instantaneously multiple classes of volatilities.

206 In our approach the surrogate is constructed with saturated hydrocarbons in order to ensure its
207 transparency to UV laser excitation. The compounds are chosen and their proportions are tuned in order

208 to obtain a mixture that reproduces the experimental distillation curve of real fuel (ASTM standards [22]).
 209 Figure 2 shows the comparison of the commercial gasoline distillation curve and the distillation curve
 210 obtained for the surrogate fuel, including modeling and experimental (ASTM distillation characterization of
 211 the formulated surrogate fuel mixture) results. The comparison has two important outputs: i) the surrogate
 212 fuel well represents the evaporation behavior of the commercial gasoline and ii) the thermodynamic model
 213 well predicts the evaporation behavior of the surrogate fuel and provides a first validation of the
 214 thermodynamic model employed. In Table 1, the composition of the gasoline surrogate is displayed
 215 together with the main physical properties of the different components. For reference ethanol properties
 216 have been added to the table.

217 **Table 1. Composition of the gasoline surrogate and physical properties of each single component at reference conditions**
 218 **(T = 25 °C and p = 1 bar).**

	Gasoline surrogate			Ethanol
	n-pentane	iso-octane	n-undecane	
Chemical Formula	C ₅ H ₁₂	C ₈ H ₁₈	C ₁₁ H ₂₄	C ₂ H ₆ O
% vol.	36	46	18	0/20/85
Density [kg/m ³]	621.7	690.3	737.6	789.1
Normal Boiling Point [°C]	36.1	99.2	195.9	78.9
Enthalpy of vaporization [kJ/kg]	365.1	308.4	364.2	838

219

220 **2.3.2 Tracers selection**

221 The tracers to be employed for the 2T-LIF experiments are selected taking into account two main aspects:
 222 *Physical properties.* The tracers have to be representative of different distilled fractions of the target fuel.
 223 In particular, one has to evaporate together with the light compounds of the fuel and the other together
 224 with the heavier ones. Boiling temperature is an important parameter to evaluate the volatility of a
 225 compound. However, when a tracer is dissolved in a blend its volatility also depends on the other
 226 compounds of the mixture. Han and Steeper et al. [23,24] showed that the sole boiling temperature of
 227 each tracer is not sufficient to choose the tracer and the co-evaporation with the other components of the

228 blend has to be taken carefully into account. The tracers-blend co-evaporation was one of the primary
229 parameter for the tracer selection and it has been checked using the thermodynamic model.

230 *Fluorescent emissions.* To enable a spectral separation of the signals emitted, the two tracers should emit
231 at different wavelengths with a minimum crosstalk. In this way, using a proper filtering, it is possible to
232 collect separately the fluorescence of each tracer and therefore to perform simultaneous measurements. It
233 is also important to avoid the co-existence of fluorescence species which interact among each other. In
234 particular, excited-state energy transfer between aromatic and ketones species have been reported to
235 interfere with signal quantification [25].

236
237 After a careful screening of the available information in the literature, the tracers selected for this study are
238 difluorobenzene (*DFB*) and 1-methylnaphthalene (*1MN*). The *DFB* represents the light compounds of the
239 tested fuel and the *1MN* the heavy compound of the fuel. The physical properties of the two tracers are
240 presented in Table 2 and their effective representativeness of the respective volatility classes will be
241 demonstrated by the experiments carried out in this work. It is important to note that *1MN* boiling point is
242 slightly higher than the maximum of the distillation curve of gasoline (c.f. Figure 2). In the results section it
243 will be proved that when the *1MN* is added in very low concentrations (~0.15%), the tracer co-evaporates
244 together with the heavier fractions of the Gasoline.

245
246 These two tracers features a good chemical stability and a very high quantum yield also at high
247 temperatures (e.g. 900K) [20,26–29]. Their emission spectrum is well separated enabling independent
248 filtering of the two signals (Figure 3) limiting the cross-talk. The high quantum yield of both tracers provide
249 sufficient fluorescent signal also at very low concentration as shown by Itani et al. (~0.15% (mass) [10]).
250 From a thermodynamic point of view, this aspect enables to not modify physical properties of the mixture.

251
252 The two tracers have been widely characterized in previous studies investigating the effect of the
253 temperature and pressure on fluorescent emissions and it is available in the literature [18,30,31]. The
254 photo-physics related to this process is described in details in [26]. This information is therefore available
255 for future quantitative interpretation of the 2T-LIF signal.

256 Despite their importance for future GDI application of the diagnostics, these features are not relevant to
 257 the present paper, in which the LIF measurement are performed at constant ambient temperature and
 258 pressure following a specific in-situ calibration. Therefore detailed information is referred to the cited
 259 bibliography.

260 The suitable thermodynamic characteristics together with the optimal fluorescent characteristics
 261 (spectrally well separated emissions and low temperature dependence) make of these tracers good
 262 candidates for GDI applications. In this work, the validation of the thermodynamic model will be therefore
 263 carried out blending the surrogate fuel presented in the previous section with 1MN and DFB.

264 **Table 2. Physical properties of the tracers. The properties are presented at standard reference conditions ($T = 25\text{ }^{\circ}\text{C}$, $p =$
 265 p_{atm})**

	DFB	1MN
Representative of	Lighter fuel compounds	Heavier fuel compounds
Chemical Formula	$\text{C}_6\text{H}_4\text{F}_2$	$\text{C}_{11}\text{H}_{10}$
Normal Boiling Point [$^{\circ}\text{C}$]	88.8	244.7
Density [kg/m^3]	1164.272	1016.43
Enthalpy of vaporization [kJ/kg]	312.97	405.15

266 **2.4 Fuel vapor composition measurement in a GDI spray**

267 The 2T-LIF provides quantitative information on local tracer mass fraction. As consequence of that, the
 268 tracers' ratio (T_R) is calculated by applying the definition below:

$$T_R = \frac{Y_{1MN}}{Y_{DFB}}, \quad (7.)$$

269 being Y_{1MN} and Y_{DFB} the mass fraction of respectively 1MN and DFB.

270 T_R has been used in previous studies to indicate qualitatively heterogeneities in the vapor fuel composition
 271 [10]. In this work, the thermodynamic model enables to establish a relationship between the ratio
 272 experimentally measured, and the fuel vapor composition.

273 To link the 2T-LIF data to the fuel vapor composition two possibilities approaches can be identified. The
 274 first, if the LIF-based imaging data are to be compared with CFD simulations, it may be possible to include
 275 the tracers in the simulation's evaporation model, which would then allow a direct comparison to the
 276 experimental data to evaluate the accuracy of that particular evaporation model. The downside of this
 277 approach is that a model may differ in the accuracy with which it predicts major fuel components vs. sub-
 278 percent trace species. On the other hand another simplified approach is proposed in this paper which
 279 need to make some assumptions about the fuel evaporation process. The vapor fuel composition
 280 calculation proposed is based on the following hypotheses:

- 281 1. It is assumed that the evaporation at the liquid-vapor interface is at equilibrium
- 282 2. The evaporation process is considered to be slow compared to the mass diffusion of the
 283 components within the droplet, and therefore the liquid phase is considered well mixed.

284 As a consequence of that, different scenarios are assumed depending on the value of T_R .

- 285 • $T_R = 1$. In this case, no preferential evaporation is observed and the composition is considered
 286 equal to the original fuel composition.
- 287 • $T_R < 1$. The high volatility tracer (DFB) prevails in the mixture. This mixture is considered to be
 288 originated by the first evaporated fractions of a droplet, which completed its evaporation
 289 elsewhere. The composition of this mixture is calculated by finding the value of evaporated mass
 290 fraction (M_{evap}) at which the ratio of the integral of the distillation curve of the tracers' is equal to
 291 $T_R, M_{evap,0}$:

$$292 \frac{\int_0^{M_{Evap,0}} y_{1MF} dM_{Evap}}{\int_0^{M_{Evap,0}} y_{DFB} dM_{Evap}} = T_r$$

293 Where, y_{1MF} and y_{DFB} indicate the distilled fractions of 1MF and DFB.

294 The mass fraction of each component Y_i is therefore obtained by integrating the distillation curves
 295 of each component i between 0 and $M_{evap,0}$

$$296 Y_i = \int_0^{M_{Evap,0}} y_i dM_{Evap}$$

297 • $T_R > 1$. The low volatility tracer (1MN) prevails in the mixture. The mixture is considered to be
 298 originated by the evaporation of a droplet which lost its lighter fractions elsewhere. In this case,
 299 $M_{evap,0}$ is found by calculating the integral of the distillation curve between $M_{evap,0}$ and 1:

$$300 \frac{\int_{M_{Evap,0}}^1 y_{1MF} dM_{Evap}}{\int_{M_{Evap,0}}^1 y_{DFB} dM_{Evap}} = T_r$$

301 In the same fashion, the mass fraction of each component Y_i is obtained by integrating the
 302 distillation curves of each component i between $M_{evap,0}$ and 1

$$303 Y_i = \int_{M_{Evap,0}}^1 y_i dM_{Evap}$$

304 In first approximation these hypotheses are applicable for a GDI spray, where the relatively low chamber
 305 temperature and gas density causes a slow evaporation process [32].

306 The relationship between T_R and vapor composition obtained with these hypotheses and on the results of
 307 the thermodynamic model is presented in Figure 4. The results refer to the composition of the gasoline
 308 surrogate described in Table 1 at atmospheric pressure for a tracer concentration of 0.15 % (in mass).
 309 This map indicates on the x-axis the tracers' concentration ratio, and, on the y-axis, the mass composition
 310 of the vapor. Each color represents a different compound of the surrogate, while the tracers, which are
 311 used in a very low concentration, are neglected in the map.

312 Also, the composition map describes a range of variation of T_R obtained by considering a complete
 313 theoretical evaporation process. However, the tracers' ratio range measurable in the reality depends on
 314 the sensitivity of the setup and of the cameras employed. For reference, based on the observations
 315 presented by Itani et al. [10,18] a "practical range" is identified and indicated on Figure 4 by considering
 316 the maximum T_R variations measured in their experiments.

317 **3 Experimental setup**

318 An experimental apparatus was put in place to provide phase equilibrium of tracers and validation data for
319 the thermodynamic model, with the objective of providing fundamental understanding of preferential
320 evaporation in multicomponent fuel and to validate the thermodynamic model.

321 The experimental setup involves a heated bubbler to evaporate gradually and in a finely controlled way a
322 certain volume of fuel; a N₂ dilution system to reduce the partial pressure of the fuel vapor extracted and
323 avoid condensation along the connecting pipeline; an optical cell, where the tracers' concentration in the
324 diluted fuel vapor is measured by means of laser induced fluorescence.

325 The fuel is evaporated gradually in a heated cell where N₂ is bubbled through the liquid phase. The N₂ +
326 vapor fuel extracted from the cell flows through a second cell at constant temperature where the 2T-LIF
327 technique is applied. The measurement can provide the relationship between the total evaporated mass
328 and the vapor tracers' concentration during the whole evaporation process. This relationship will be the
329 reference to be used for the validation of the thermodynamic model.

330 The details of the setup and of the data analysis are presented in the following section.

331 **3.1 Experimental apparatus**

332 The experimental apparatus is presented schematically in Figure 5 and it is composed of three main parts:

- 333 1. *Evaporation cell.* The aim of the evaporation cell is to cause the evaporation of the volume of the
334 tested fuel in a controlled and repetitive way. The cell is partially filled with a volume of the studied
335 fuel V_0 (~20 ml). A bubbler feeds a constant flow rate of nitrogen (\dot{m}_1) through the cell to ensure
336 the formation of saturated fuel vapor in the upper part of the chamber where the outlet is placed. A
337 set of heating resistances heats homogeneously the evaporation cell, while a control system sets
338 the heating power to increase gradually and in a controlled way the temperature of the cell. The
339 volume of liquid fuel is monitored thanks to an optical access to the evaporation chamber that
340 enables to measure the height H of remaining fuel; the corresponding mass is then calculated

341 taking into account the density corresponding to the instantaneous temperature of the fuel (T_1)
342 which is measured directly through a thermocouple immersed in the liquid fuel.

343 2. *Heated pipeline.* The N_2 , saturated with fuel vapor, exits the evaporation chamber, and passes
344 through a heated pipeline connecting the evaporation cell to the imaging cell. To prevent the re-
345 condensation of the fuel vapor along the pipe walls, the pipe is heated at a temperature (T_{pipe})
346 higher than T_1 . Moreover, a second constant stream of heated nitrogen \dot{m}_2 is fed, being the mass
347 flow rate \dot{m}_2 a multiple of \dot{m}_1 ($\dot{m}_2 = n \dot{m}_1$) to reduce the concentration of fuel vapor. The value of
348 n is approximately 20. The dilution has the two following advantages: i) it further reduces the
349 possibility of fuel condensation by decreasing the vapor partial pressure and ii) it increases the
350 mass flow rate through the imaging chamber improving the time response of the system and
351 therefore the accuracy of the measurement.

352 3. *Imaging cell.* The imaging cell is the part of the experimental apparatus devoted to the application
353 of the 2T-LIF. As for the connecting pipe, condensation of the fuel has to be prevented, and
354 therefore the whole cell is heated. Since the fluorescent emissions of the tracers are affected by
355 the temperature, the LIF cell temperature has to be kept constant and homogeneous during the
356 experiments, and its value was constantly monitored ($T_3 = 365^\circ\text{C}$). The optical access to the cell is
357 provided by three quartz windows (36 mm diameter): two of them are aligned to the laser beam
358 and a third orthogonal window is devoted to the LIF signal collection.

359 All the system is designed to perform tests up to a pressure of 15 bar. The nitrogen flow (\dot{m}_1 and \dot{m}_2) is
360 controlled by means of two automatic flow regulators upstream of the system, while the pressure of the
361 system (p_{sys}) is controlled by manually adjusting the aperture of a valve downstream of the LIF cell
362 (Figure 5). The complete details of the optical setup and of the data processing are presented in the
363 Appendix.

364 As explained in the appendix, the results are normalized for the total mass of the tracers in the fuel, and
365 considering the constant boundary conditions in the imaging cell, the results are independent on
366 fluorescence properties of the mixtures.

367

368 Figure 6 presents a sample of the results obtained by the analysis. The results present the tracers'
369 distilled fraction at each step of the fuel evaporation, indicating on the abscissa the Evaporated Mass
370 Fraction (*EMF*). In accordance to the physical properties of the tracers, *DFB* starts evaporating at the
371 beginning of the evaporation process and continues until the *EMF* is approximately 0.8. The *1MN*, with
372 higher boiling point, starts evaporating for *EMF* > 0.7 and it continues until the end of the process. Figure
373 6 also shows the comparison between three repetitions of the test: the results of the methodology appear
374 precise and robust, and in the remnant of the paper only the average of the three repetitions will be
375 presented. It is worth underlining that for all the tests performed a certain dispersion of the results was
376 observed at the beginning and at the end of the evaporation. The authors consider that data gathered at
377 the beginning and at the end of the experiment are less accurate due to different reasons. At the
378 beginning of the experiments the nitrogen flow is activated in the bubbler and in the dilution line: this
379 process might take few minutes before reaching the set-point values, especially at higher test pressures
380 and therefore the related data could be slightly biased in this period. On the other hand, at the end of the
381 experiments the measure of the remaining volume becomes more difficult due to the increasing impact
382 that the reading uncertainty ($\pm 20\text{mm}^3$) has on the total volume measured. For this reason for *EMF* < 0.1
383 and *EMF* > 0.9 interval of the evaporation diagrams are normally not considered for the comparisons and
384 the results will be considered only in the 10-90% mass fraction interval.

385 **3.2 Test matrix**

386 The experimental campaign has the double objective of investigating the effect of parametric variations on
387 preferential evaporation phenomenon and to provide data to validate the thermodynamic model.

388 The effect of the evaporation pressure was tested carrying out the tests at 1 bar (atmospheric), 5 and 10
389 bar. The fuel composition effect was also investigated by adding progressively ethanol to the surrogate
390 fuel. Apart from the standard gasoline surrogate defined in Table 1, two further surrogate/ethanol blends
391 were tested featuring 20 and 85% in volume of ethanol. In the remnant of the paper standard gasoline
392 surrogate will be referred as E00, while the 20% and 85% ethanol blends will be referred as E20 and E85.

393 **4 Experimental results and model validation**

394 In this section, the experimental results are presented and compared to the results provided by the
395 thermodynamic model. The effects of parametric variations in system pressure and ethanol content are
396 discussed.

397 **4.1 Gasoline surrogate at atmospheric pressure**

398 Figure 7 presents the results obtained for the evaporation of gasoline surrogate (E00) at atmospheric
399 pressure. Experimental and modeling results indicate a significant separation between the evaporation of
400 the two tracers: *DFB* evaporates together with the lighter fraction of the surrogate and then gradually its
401 concentration in the vapor decreases. The *DFB* fluorescent intensity (and therefore *DFB* concentration)
402 becomes lower than the noise level at approximately 80% of the evaporated fuel mass. On the contrary,
403 *1MN* fluorescence is detected approximately when the EMF exceeds 0.6 and it evaporates together with
404 the heavier fuel distilled fraction.

405 This first result confirms that the tracers chosen are capable of representing the different volatility class of
406 the surrogate fuel, with *DFB* being representative of the light compounds, while *1MN* is representative of
407 the heavier ones.

408 When the tracers evaporation curves are compared to the ones provided by the model, the evaporation
409 behavior of the two tracers is globally well represented by the model predictions along the whole
410 evaporation process. In particular, the co-evaporation process of *1MN* is well represented: even though
411 the boiling point of this tracer is higher than the one of the heaviest component of the surrogate, it starts
412 evaporating well before the end of the evaporation process.

413 **4.2 Effect of ambient pressure**

414 Figure 8 presents the experimental results (left hand side) and model predictions (right hand side) at
415 different ambient pressures for the gasoline surrogate (*E00*). Despite the increase in p_{sys} the comparison
416 indicates that the preferential evaporation phenomenon remains substantially similar: the *DFB* evaporates

417 evenly until 80% of the evaporated mass fraction and the *1MN* curve is characterized by a prominent peak
418 rising at $EMF > 0.9$. More in details, both, experiments and model, show qualitatively similar variation in the
419 evaporation profiles when p_{sys} is increased: a flattening of the *DFB* evaporation profile and its persistence
420 also during the last phase of the fuel evaporation. Also, a progressive sharpening of the *1MN* peak at the
421 end of the evaporation is observed. Although some discrepancies can be observed between model and
422 experimental results, it can be concluded that, globally, it captures the changes in the shape of the
423 evaporation of the tracers.

424 Figure 9 shows more results from the thermodynamic model, indicating how the system pressure affects
425 the evaporation of the different components of the surrogate. In particular it can be observed that it has a
426 non-negligible impact in homogenizing the evaporation of the components. E.g. n-pentane persists in the
427 liquid up to the end of the evaporation and n-undecane starts evaporating at a low rate since the
428 beginning of the process. In a practical way, this result indicates that increasing the system pressure can
429 potentially reduce the preferential evaporation effect.

430 **4.3 Effect of Ethanol**

431 The effect of the ethanol addition on the tracers evaporation is presented in Figure 10 showing for each
432 ethanol blend the results obtained at ambient pressure (left column) and at 10 bar (right column).

433 Model and experimental results indicate that the addition of ethanol has a non-negligible impact on the
434 evaporation of the fuel. In particular the variation in the shape of the *DFB* and *1MN* distillation curves
435 indicates that the evaporation of the light and heavy compounds happens in the three cases in a different
436 way.

437 For *E85* case the *1MN* evaporation is observed mainly during the last 5% of the fuel mass fraction
438 evaporation. Even though the measurements concerning the last 10% of evaporated fuel are
439 characterized by low accuracy, both, experiments and models well capture the modification in the
440 evaporation curves of *DFB* and *1MF* induced by the ethanol addition. The effect of ethanol is well
441 represented by the model, at least in a qualitative way. In particular, results indicate that increasing the
442 ethanol content causes *1MN* to evaporate progressively at higher values of *EMF*. However, the trend for
443 *DFB* is not as simple: while for *E20*, *DFB* evaporation is delayed when compared to *E00*, for *E85* *DFB*

444 evaporate at lower values of EMF. In Figure 11 more results from the thermodynamic model are
445 presented showing the vapor concentration predicted for each distilled fraction (the tracer composition is
446 neglected). These results show that the evaporation of the fuel changes radically when passing from E20
447 to E85. Consistently with its lower boiling temperature n-pentane is the first fraction to evaporate for the
448 three cases. Following the same logic, n-undecane is the last component to evaporate (boiling point 195
449 °C). On the contrary, iso-octane evaporates after ethanol for E20 and before in E85. This behavior can
450 be related to the formation of an azeotrope among ethanol forms with lighter compounds for E85 which
451 links together the evaporation of n-pentane, iso-octane and ethanol. This behavior is related to the physic-
452 chemical affinity between ethanol, iso-octane and n-pentane. Similar effects have been described in [13].
453 More in details, it can be observed as for E20 the azeotrope causes a similar evaporation among iso-
454 octane and ethanol up to the complete evaporation of the latter. After this point the ethanol an evident
455 change of behavior of iso-octane can be observed (c.f. red arrow in Figure 11). On the contrary, for E85,
456 due to the higher presence of ethanol, the iso-octane evaporates to completion before the ethanol,
457 changing radically the shape of its evaporation curve.

458 A closer look to Figure 11 and Figure 10 shows that the modification is the shape of the curved observed
459 for DFB follow quite closely the ones observed for iso-octane. This fact is also underlined by the
460 appearance of a peak on the DFB evaporation curve observed for E20 (c.f. red arrow in Figure 10). This
461 peak corresponds to the peak in iso-octane evaporation observed for iso-octane (c.f. red arrow in Figure
462 11). This aspect underlines how the thermodynamic model is important to understand the tracers' results:
463 in fact by taking only into account the boiling point, one might expect DFB to be equally representative of
464 Ethanol and iso-octane. These results, as others in the literature [16,18,32], show clearly that more
465 complex interactions need to be taken into account.

466 Figure 11 also provides another important indication about the lower volatility classes of the surrogate (n-
467 undecane). Probably due to the high difference in boiling point, n-undecane does not co-evaporate with
468 the azeotrope mentioned above, and it is indicate to evaporate to completion only when the last fraction of
469 ethanol is evaporated. In a more practical sense, an important ethanol blending in commercial gasolines
470 (e.g. E85), while homogenizing the evaporation of the light and medium volatility class of gasoline might
471 cause a higher segregation of the heavier compounds.

472 Ethanol, due to its higher evaporation enthalpy, is in general considered to delay sprays evaporation due
473 to its high latent heat [15]. However, the results observed in this section indicate that, if the well mixed
474 hypothesis of the liquid fuel is verified, as it is reasonable for the evaporation experiment, ethanol
475 evaporate well before the heavier volatility compounds, causing less co-evaporation of the heavier
476 fractions (e.g. n-undecane).

477 **4.4 Vapor composition calculation and accuracy estimation**

478 To provide a further insight about the accuracy of the methodology proposed the coupling between the
479 thermodynamic model and the results from the 2T-LIF measurements are analyzed investigating the
480 impact that the discrepancies between experimental and modeling results presented in section 4, have on
481 the global accuracy of the technique.

482 Instead of calculating the complete uncertainty analysis of the evaporation experiments, which analysis
483 would go beyond the scope of this section, the deviation between the model and the experimental results
484 has been considered as a first order estimation of the error of the model predictions. This deviation has
485 been employed to estimate how the error in the thermodynamic model prediction on tracers' evaporation,
486 propagates to the vapor fuel calculation. In this way: fixing the relationship between the vapor composition
487 and the fuel evaporated mass, the map presented in Figure 4 has been re-calculated in two different ways
488 1) using the T_R obtained by the model simulation (as done for Figure 4) and 2) using the T_R obtained
489 experimentally.

490 In Figure 12 the comparison is presented for the different fuels tested (from top to bottom E00, E20 and
491 E85) and for two p_{sys} (from left to right 1 and 10 bar). The comparison shows that for all the cases, the fuel
492 vapor concentration is obtained with accuracy also when using the experimental input. In the practical
493 range the difference is in general lower than 2 % and never above 5%. This means that despite the
494 discrepancies observed in the evaporation experiment between model and experimental results the
495 impact on the vapor fuel composition calculation are expected to be acceptable (<5%) in the relevant
496 range of measurements.

497 **5 Summary and Conclusions**

498 This paper presents a novel approach for the quantitative interpretation of 2T-LIF experiments, making of
499 it a powerful optical diagnostic to study preferential evaporation in GDI engines.

500 The methodology combines a state-of-the art optical diagnostic (2-tracers laser induced fluorescence) with
501 a numerical model, and enables to assess the effects of preferential evaporation in a multicomponent and
502 eventually to perform quantitative measurement of vapor-phase fuel composition. An optically transparent
503 surrogate for commercial gasoline has been found which replicates closely the distillation curve of the real
504 fuel.

505 This paper, a part from presenting the methodologies, presents the results of an experiment that, at the
506 same time, investigates the effect of boundary conditions on preferential evaporation process and
507 assesses the accuracy of the methodology proposed. The main outcomes are:

- 508 • An increase in system pressure increases the co-evaporation among the different volatility class.
- 509 • The addition of ethanol was observed to have a complex effect on fuel evaporation. When added
510 in lower concentrations (E20) it is observed to evaporate before isooctane as expected
511 considering the lower boiling point. However, for E85, where ethanol is the main component of the
512 blend it causes n-pentane and iso-octane to evaporate to completion before ethanol. At the same
513 time for E85 the evaporation of the n-undecane is segregated and the end of the evaporation
514 curve.
- 515 • Ethanol is observed to form an azeotrope with iso-octane and n-pentane due to their physical-
516 chemical affinity changing significantly their evaporation curves.
- 517 • The two tracers proposed, have been found to co-evaporate together with the light-medium
518 volatility class of gasoline (difluorobenzene) and the heavy ones (1-methylnaphtalene).

519 Finally, the impact of hypothetic errors of the thermodynamic model on the vapor fuel composition
520 calculation was assessed, by using as error indication the deviation between experimental and
521 modeling results. The analysis indicates that, the impact on the vapor fuel calculation for the
522 conditions relevant to engine application were acceptable (< 5%).

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527 and the tracer characterization data provided.

528 **Symbology**

529 1MN: 1-methylnaphtalene

530 2T-LIF: two-tracer laser induced fluorescence

531 a, b, a_i, b_i : constants

532 C_1, C_2, C_3 : constants

533 DFB: difluorobenzene

534 E00: gasoline surrogate

535 E20: gasoline surrogate + ethanol 20% (vol)

536 E85: gasoline surrogate + ethanol 85% (vol)

537 EMF: evaporated mass fraction

538 E_{laser} : laser pulse energy

539 F.S.: camera sensor full scale

540 GDI: gasoline direct injection

541 I_1, I_2 : average counts measured by the camera for the area A_1 and A_2

542 L_1, L_2 : Fluorescent intensity measured in the portion of the sensor A_1 and A_2

543 M_1, M_2 : quantity proportional to the mass of tracer passed through in the imaging cell during the
544 integration interval

545 N : average noise counts (measured in the area A_N)

546 p : pressure (generic)

547 p_{sys} : pressure of the experimental apparatus

548 R: universal gas constant
549 SRK: Soave-Redlich-Kwong.
550 T: temperature (generic)
551 T_1 : Temperature measured within the bubbler
552 T_3 : Temperature measured within the imaging cell
553 T_R : Tracers' concentration ratio
554 T_{pipe} : temperature of the connecting pipe
555 V_{fuel} : volume of the liquid fuel in the evaporation cell
556 v: specific volume (generic)

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628 **7 Appendix A**

629 In this section the details of the optical setup and data processing are presented.

630 **7.1 Optical setup**

631 The laser beam employed for the 2T-LIF was the fourth harmonic of the pump wave of an Nd:YAG laser
632 (266 nm). The maximum output of each laser beam was about 60 mJ and was modulated along the full
633 range from 0 to 100% of the total power to avoid camera saturation using a beam attenuator (*Altechna-*
634 *Watt Pilot*). The beam energy corresponding to each fluorescence image was measured and recorded
635 using a beam sampler reflecting 10% of the beam intensity to a high-speed power meter. The laser
636 fluence in the test section was always kept below 0.7 mJ/mm^2 assuring to remain in a linear fluorescence
637 regime [30,31].

638 The simultaneous detection of the fluorescent signal obtained by the two tracers was performed by means
639 of an intensified CCD camera (Princeton instrument – Pimax) fitted with a stereoscope or image-doubler.
640 The stereoscope is a commercial device (provided by LaVision) and it enables to project two times the
641 images “seen” by the two input channels on two different portion of the camera’s sensor. The sample
642 stereoscopic image presented in Figure 13 shows the LIF window, and a ruler placed within the chamber
643 along the laser beam path. Each of the stereoscope channels is fitted with a different filter as defined in
644 2.3.2, to collect separately the fluorescence signal of 1-methylnaphtalene ($370 \pm 18 \text{ nm}$) and di-
645 fluorbenzene ($280 \pm 10 \text{ nm}$). The filters enable to collect separately the signal from the two tracers with a
646 negligible cross talk (c.f. Figure 3)

647 **7.2 Test Procedure**

648 The surrogate fuel (already doped with the tracers and stored at about -10°C) is poured into the
649 evaporation cell which temperature is kept at -5°C thanks to a refrigeration system. After closing the
650 evaporation cell, the nitrogen start flowing through the bubbler and through the dilution system and at the
651 same time the 2T-LIF system starts its acquisition. The N_2 mass flow rate is kept constant by two mass

652 flow rate meter placed upstream of the circuit. The laser starts pulsing at its working frequency (10 Hz),
653 and the intensified LIF stereoscopic images are acquired at a reduced rate (0.2 Hz). For each LIF image
654 also the following data are recorded:

- 655 • Temperatures of the system T_1 , T_{pipe} , T_3 (see Figure 5).
- 656 • Fuel height in the evaporation cell H, converted into volume left V_{fuel} . A CCD camera is used to
657 monitor the level of the fuel in the evaporation cell. An automatic processing algorithm is
658 employed to obtain the corresponding fuel volume. In order to improve the accuracy of the volume
659 measurement, the nitrogen flow to the bubbler is stopped shortly before the acquisition of the
660 image (0.2s) for a short time (0.25 s). This interruption of the bubbler flow is not considered to
661 affect the nitrogen mass flow measurement since this process was periodic (0.2 Hz) and the
662 duration of the interruption was short relatively to the period.
- 663 • Laser pulse energy E_{laser} . The laser pulse energy is measured inserting a beam-sampler before
664 the test chamber that directs part of the laser beam towards a laser energy-meter.

665 The data is acquired until the complete evaporation of the fuel. In order to have a good measurement
666 precision, the fuel is evaporated gradually, and the whole process lasts 1-1.5h.

667 **7.3 Image processing and data analysis**

668 The images collected by the camera during the fuel evaporation were analyzed to establish a relationship
669 between the evaporation of the two tracers and the evaporation of the fuel.

670 Each LIF image was analyzed in the following way:

- 671 I. The signal recorded by the camera for each channel is evaluated by averaging the pixel counts in
672 the interrogation areas indicated in Figure 14 (I_1 and I_2), which correspond to the same part of the
673 chamber.
- 674
- 675
- 676 II. The average counts numbers corresponding to each interrogation area are transformed in a value
677 proportional to the photons reaching the sensor (L_1 and L_2), by taking into account the

678 background noise, evaluated for each image in an area (indicated as A_N in Figure 14) where no
679 fluorescence signal is observed (N).

$$L_1 = \frac{I_1 - N}{F.S. - N} , \quad L_2 = \frac{I_2 - N}{F.S. - N} \quad (8.)$$

680 being $F.S.$ the full scale of the sensor.

681
682 III. The LIF signals are normalized to the measured laser pulse energy E_{laser} . The values obtained
683 (L_{n1} and L_{n2}) are proportional to the tracer mass concentration in the visualization cell.

$$L_{n1} = \frac{L_1}{E_{laser}} , \quad L_{n2} = \frac{L_2}{E_{laser}} \quad (9.)$$

684 IV. As the gas flowing through the cell is mainly composed by nitrogen, the global mass flow through
685 the LIF cell can be assumed as constant. Therefore, the integral of $L_{1,n}$ and $L_{2,n}$ on time is
686 proportional to the tracer evaporated mass. This value is calculated as,

$$M_1 \propto \int_{t_1}^{t_2} L_{1,n} dt , \quad M_2 \propto \int_{t_1}^{t_2} L_{2,n} dt \quad (10.)$$

687 V. The mass of fuel evaporated at each time interval (determined through level measured in the
688 evaporating cell) can therefore be related to the tracer evaporated masses M_1 and M_2 . Once
689 normalized with reference to the total initial fuel mass, the curves shown in Figure 6 are obtained.
690 On the x-axis the fuel evaporated mass fraction, while on the y-axis the normalized distilled
691 fraction for each tracer is presented. The integral of each curve on the fuel evaporated mass
692 fraction is the unity.

693 It is important to underline that considering the constant pressure and temperature condition in the
694 imaging chamber, the signal calibration is based on the normalization for the total tracer mass blended in
695 the fuel. Therefore the results are not dependent on fluorescent properties of the tracers.

696 8 List of captions

697 Figure 1. Scheme of the workflow for the quantitative measurement of fuel vapor composition.

698 Figure 2. Distillation curve of commercial gasoline (Euro 5, Directive 2009/30/EC) and the surrogate chosen for this study.

699 The model results are plotted together with the experimental characterization.

700 Figure 3. Fluorescence spectra of *1MN* and *DFB* for a 266 nm wavelength excitation at 350 °C and atmospheric pressure.

701 The transmission of the filters employed in the evaporation experiment (described in the following section) are indicated by
702 the areas in light colors [18].

703 Figure 4. Example of composition map for E00 at 1bar. The practical variation range derived for the experimental

704 measurements of the GDI application is indicated by the red dashed lines.

705 Figure 5. Schematic of the experimental validation setup.

706 Figure 6. Sample output from the data processing. The relationship between the distilled fraction of each tracer and the fuel

707 evaporated mass fraction is presented. E00, $p_{sys} = 1 \text{ bar}$. The gray areas represents the regions of low experimental

708 accuracy.

709 Figure 7. Experimental (dashed lines) and modeling (solid lines) results. Tracers' distilled fraction during the evaporation of

710 the fuel. The gray areas represents the regions of low experimental accuracy. E00, $p_{sys} = 1 \text{ bar}$

711 Figure 8. Experimental (left) and modeling results (right). Tracers' distilled fraction during the evaporation of the fuel. Effect

712 of ambient pressure on the evaporation of E00.

713 Figure 9. Modeling results of the fuel compounds' evaporation of at different p_{sys} .

714 Figure 10. Effect of ethanol addition on the evaporation of the surrogate fuel. Experimental results and model predictions.

715 Figure 11. Evaporation of the different fuel compounds. E00 (left), E20 (center) and E85 (left) at ambient pressure.

716 Figure 12. Fuel vapor composition obtained using modeling (solid lines) and experimental results (dashed lines). From top

717 to bottom E00, E20 and E85. p_{sys} 1bar (left) and 10 bar (right). The practical range for the 2T-LIF application is indicated in

718 red dashed lines.

719 Figure 13. Sample image obtained using the image-doubler. The LIF cell window image from the two channels of the

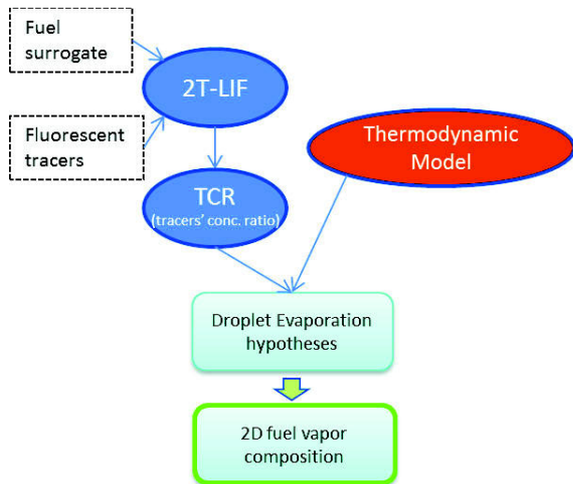
720 stereoscope.

721 Figure 14. Sample 2T-LIF image. The fluorescence signal obtained in the two channels along the laser beam path is

722 displayed. The signal evaluation areas are indicated by the green squares.

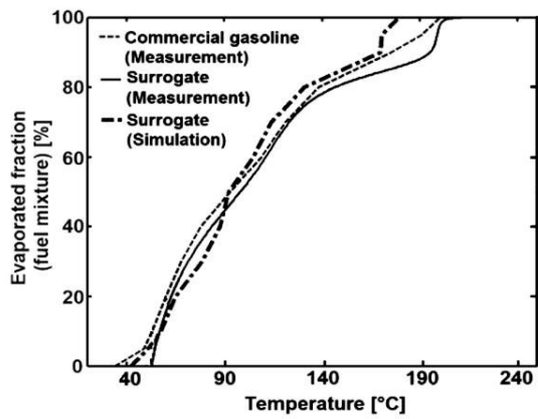
723

724 **9 List of figures**



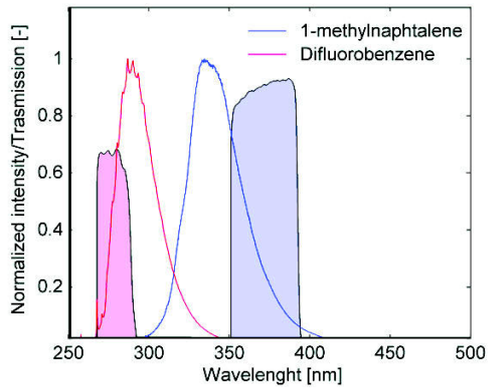
725
726 Figure 1

727



728
729 Figure 2

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731

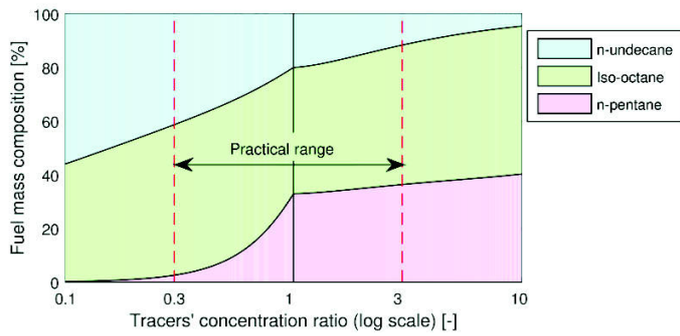
732 Figure 3

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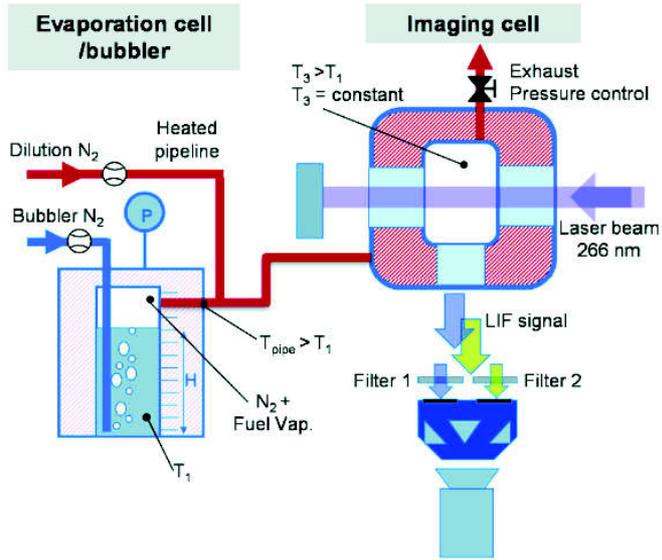
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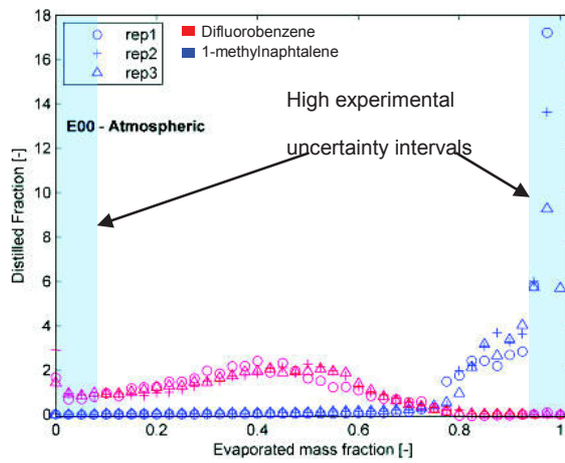
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738 Figure 4

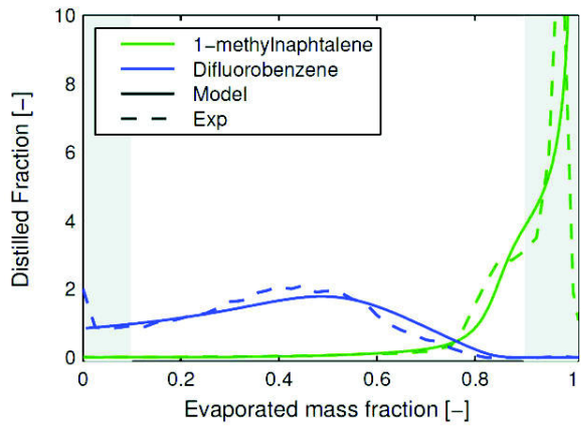
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740
 741 Figure 5
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745
 746 Figure 6
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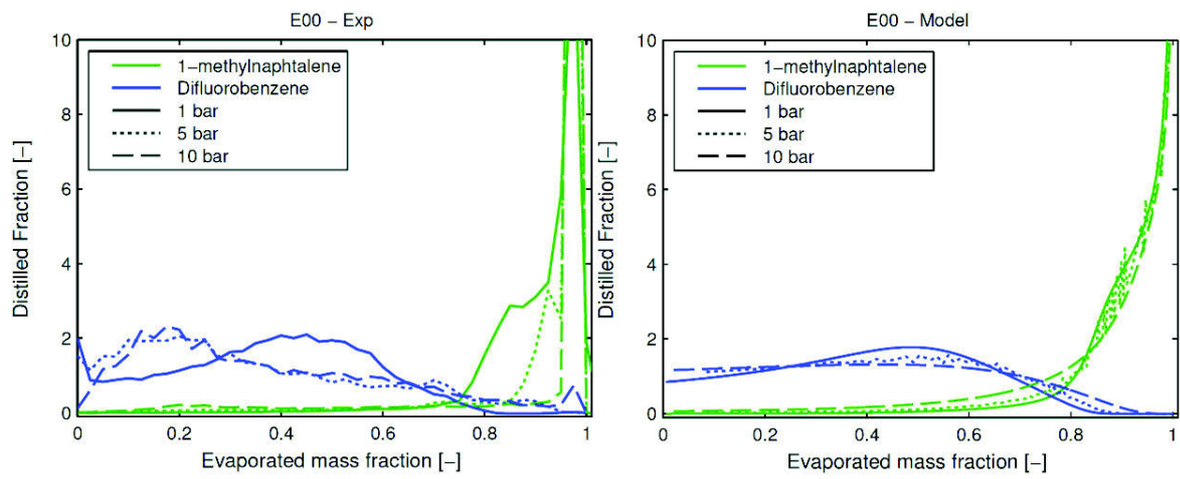


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751 Figure 7

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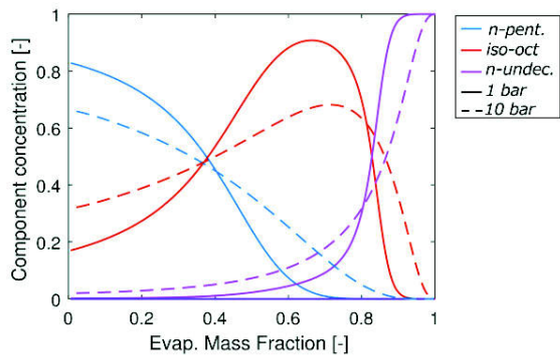


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755 Figure 8

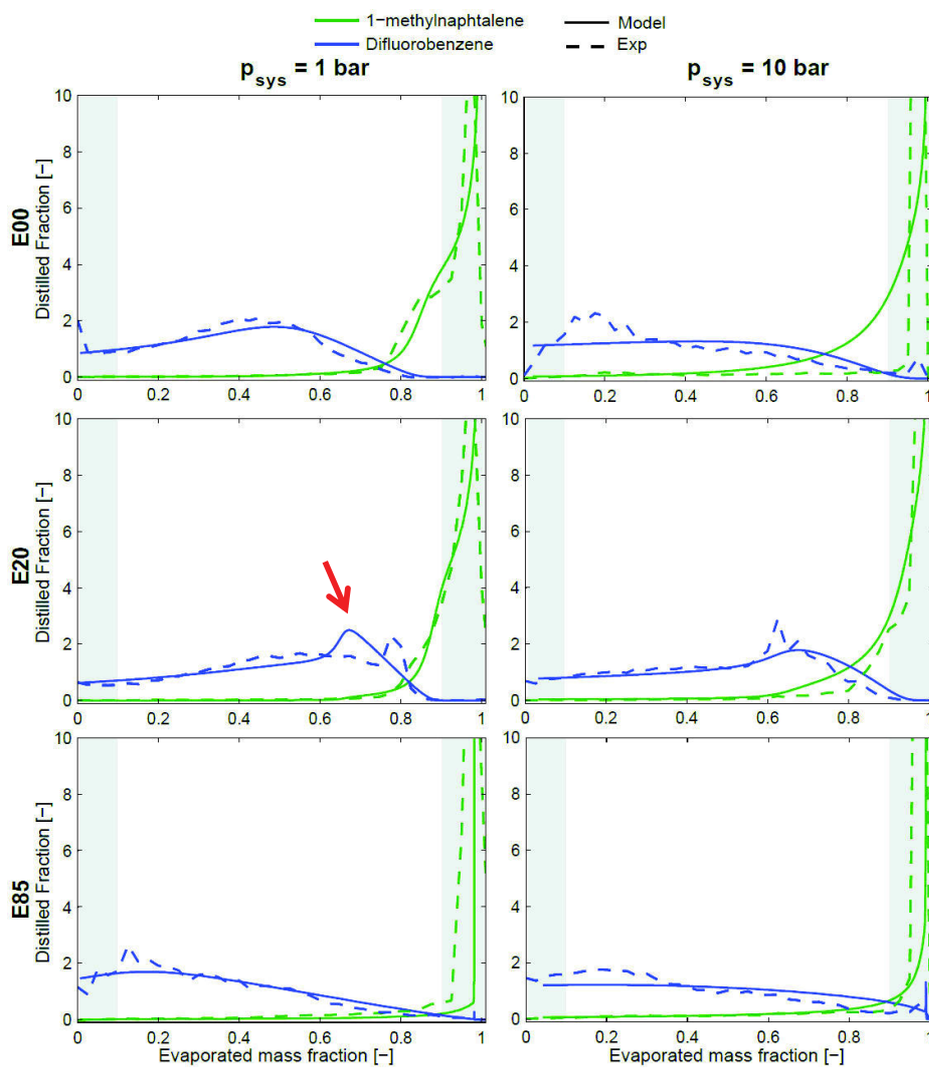
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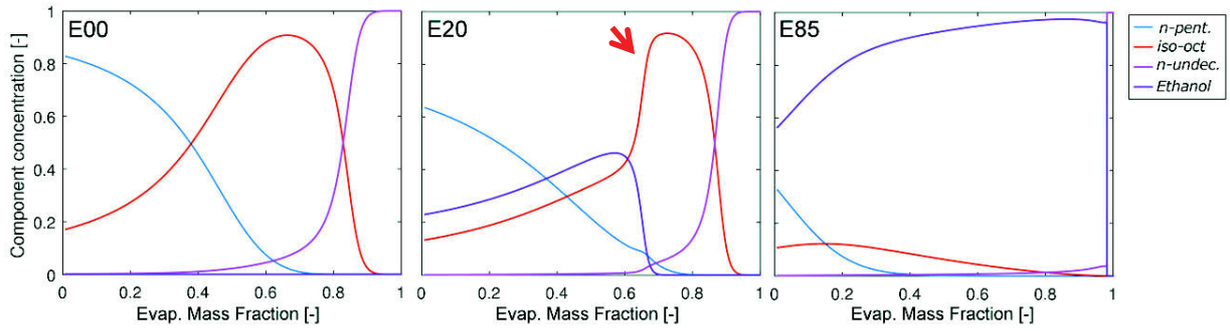
759 Figure 9



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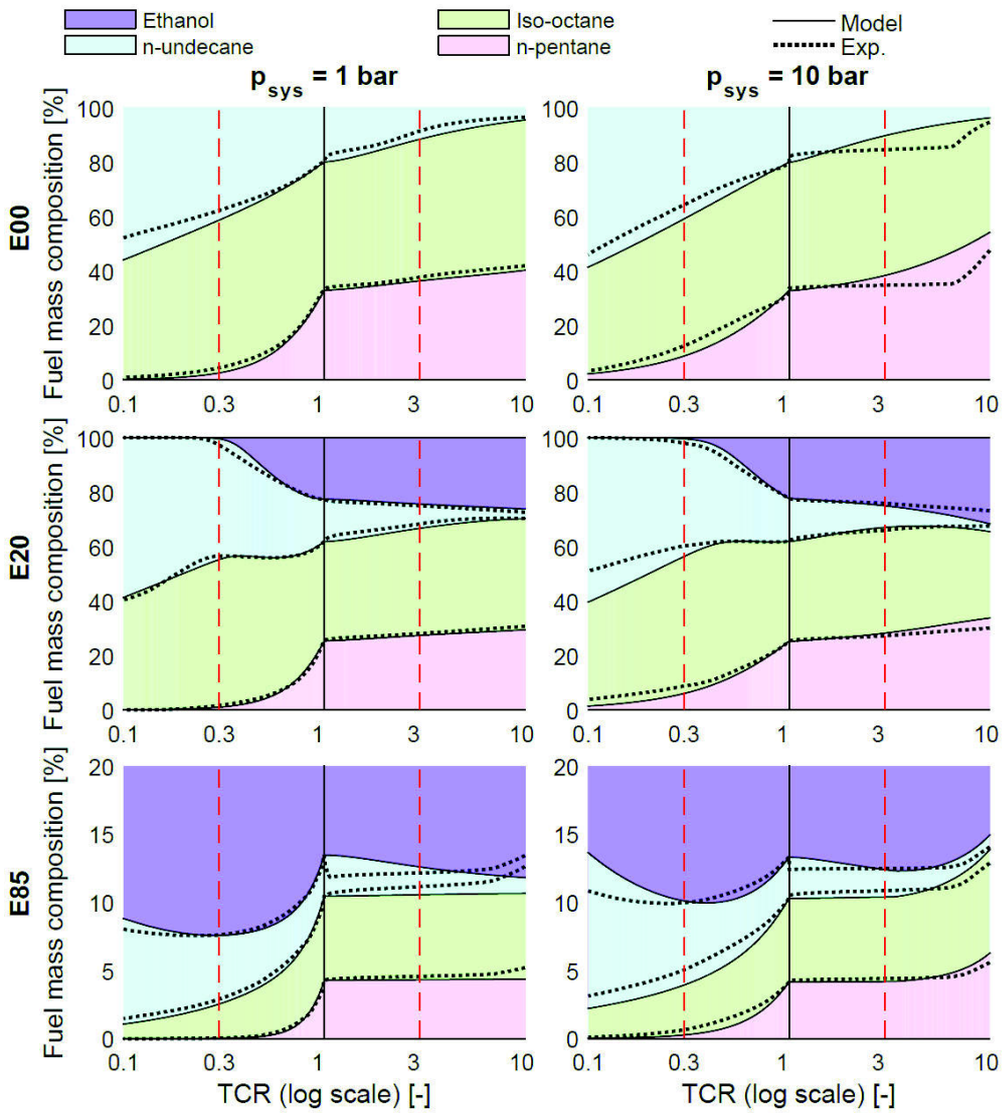
761 Figure 10

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764 Figure 11



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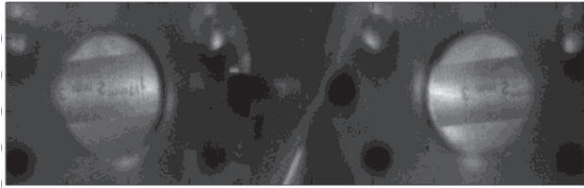
766 Figure 12

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768

Left channel

Right channel



769

770 Figure 13

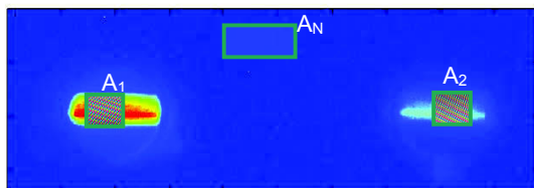
771

1-methylnapthalene

Difluorobenzene

470 ± 20 nm

285 ± 10 nm



772

773 Figure 14

774