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# Predicting the evaporation rate of stationary droplets with the VOF methodology for a wide range of ambient temperature conditions

4

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13

#### 14 Abstract

15 This paper presents CFD predictions for the evaporation of nearly spherical suspended droplets 16 for ambient temperatures in the range 0.56 up to 1.62 of the critical fuel temperature, under 17 atmospheric pressures. The model solves the Navier-Stokes equations along with the energy 18 conservation equation and the species transport equations; the Volume of Fluid (VOF) 19 methodology has been utilized to capture the liquid-gas interface using an adaptive local grid 20 refinement technique aiming to minimize the computational cost and achieve high resolution at 21 the liquid-gas interface region. A local evaporation rate model independent of the interface 22 shape is further utilized by using the local vapor concentration gradient on the droplet-gas 23 interface and assuming saturation thermodynamic conditions. The model results are compared 24 against experimental data for suspended droplet evaporation at ambient air cross flow including 25 single- and multi-component droplets as well as experiments for non-convective conditions. It is 26 proved that the detailed evaporation process under atmospheric pressure conditions can be 27 accurately predicted for the wide range of ambient temperature conditions investigated.

28

#### 29 Keywords: VOF, evaporation, multi-component, high temperature

#### 31 1 INTRODUCTION

32 Droplet evaporation is an important phenomenon realized in several engineering and physical 33 processes; it has been addressed in several textbooks, see selectively [1-4] and review articles 34 [5-8]. Numerical modelling of droplet evaporation started with the idealized evaporation of a 35 spherical, isolated, motionless and constant temperature droplet in an inert gas environment. 36 Under these assumptions, Godsave [9] and Spalding [10] derived the widely known " $d^2$ -law" 37 which predicts that the squared diameter of the droplet reduces linearly with time. Following 38 this approach, these strict assumptions were relaxed by taking into consideration the relative 39 velocity between the air and the droplet, the transient droplet heating, the Stefan flow effects, 40 the evaporation of multicomponent droplets and other secondary phenomena. The transient droplet heating was accounted with the Infinite Conductivity Model (ICM) [11] and the spatial 41 42 temperature distribution inside the droplet with the Finite Conductivity Model (FCM) [12], the 43 Effective Conductivity Model (ECM) [13] and the parabolic temperature profile model [14]. 44 Regarding the Stefan flow effects which arise from the radial vapor motion, the models of 45 Abramzon & Sirignano [13] and Yao et al. [15] are widely used, while there is a large variety of 46 available heat/mass transfer correlations to account for the relative droplet-gas motion [7]. The 47 performance of the aforementioned models was assessed in comparative studies such as in [16-48 19].

49 Modelling of multicomponent droplet evaporation has been also a challenging task since most 50 of the fuels are mixtures of many hydrocarbon components. For a bi-component mixture, the 51 preferential evaporation behavior obeys the distillation behavior of the two components, in 52 which the most volatile species evaporates first followed by the vaporization of the less volatile 53 component. The properties of the mixture depend, in addition to the local temperature, to the 54 concentration. Thus, additional equations need to be solved, accounting for the variation of the 55 species concentration in the liquid phase. Similar to the modelling of the thermal behavior of the 56 droplet, the temporal evolution of the species concentration is simulated by using 0-D or 1-D 57 models, usually known as the Infinite Diffusivity Models (IDM) [20-22] and the Finite 58 Diffusivity Models (FDM) [23-26], respectively.

The aforementioned simplified approaches are suitable for Lagrangian spray models predicting the evolution of sprays plumes in combustion systems, comprised by a large number of polydispersed size droplets, as they do not require a lot of computational resources. On the other hand, their applicability is limited to spherical or near spherical droplets; moreover, they cannot give insight of the complex 3-D fluid and thermal transport processes taking place during

64 evaporation. Such models require solution of the Navier-Stokes equations along with the energy 65 and species transport equations both for the liquid and gaseous phases simultaneously. The 66 earliest CFD works on droplet evaporation were those of [27-29] who examined the evaporation 67 of single-component spherical droplets and derived useful correlations for the Nusselt and 68 Sherwood numbers, followed later by [30-34]. These works have revealed the importance of the 69 spatial distribution of the velocity, temperature and vapor concentration field and they can be 70 considered as benchmark cases for the validation of 0-D and 1-D models. Later, they were 71 extended to multicomponent fuels in [35-38] and to high pressure conditions in [39]. A common 72 feature of the aforementioned CFD works is that they use a grid fitted to the liquid-gas 73 interface; this provides an accurate representation of the flow field and allows for the 74 determination of the local momentum-heat-mass transfer rates, but they are limited to 2D 75 axisymmetric flows with no or low surface deformation as in [40]. On the other hand, the 76 Volume of Fluid (VOF) and Level Set (LS) methodologies among others can capture the 3D 77 droplet deformation (and even breakup), but with a less accurate prediction of the flow field at 78 the interface region. Recent works on droplet evaporation with the VOF methodology include 79 those of [41-44]. In these works a local evaporation model was used; Schlottke & Weigand [41] 80 used a virtual mass averaged velocity to calculate the source term of the continuity equation at 81 the interface region which improved the model performance. Furthermore, Strotos et al. [42] 82 and Banerjee [43] performed parametric studies for bi-component droplet evaporation while 83 Ghata & Shaw [44] studied the vaporization of fiber-supported droplets and included the effect 84 of thermocapillary stresses at the interface.

85 The present work examines the droplet evaporation using the solution of the Navier-Stokes 86 equations along with the energy conservation equation, the transport of species and the VOF 87 methodology for tracking the droplet interphase. The validation study examines single- and 88 multi-component droplets for a wide range of ambient temperature conditions under 89 atmospheric pressure. Relative to previous relevant CFD studies with the VOF methodology 90 resolving the complete fluid transport processes during vaporization, including the work of the 91 authors presented in [42], this is one of the first studies (at least to the authors' knowledge) that 92 examines droplet evaporation at temperature ambient conditions above the critical fuel 93 temperature and validates against experimental data the model for a wide range of temperatures. 94 The following sections include the description of the numerical model, the cases simulated, 95 followed by the model predictions for low and high ambient temperature conditions. The 96 conclusions of this work are summarized at the end.

#### 98 2 MATHEMATICAL MODEL

#### 99 2.1 Flow field equations

100 The numerical model involves the solution of the Navier-Stokes equations along with the 101 solution of the transport equation of the liquid phase. The model is an extension of the one used 102 in [45-47] including phase change. In addition, the numerical model employs the Volume of 103 Fluid (VOF) methodology [48] for capturing the droplet liquid-gas interphase. In this 104 methodology, the liquid phase is identified by the liquid volume fraction  $a_l$  (defined as the 105 liquid volume in a cell divided by the cell volume) and its transport is given by:

106 
$$\frac{1}{\rho_l} \left[ \frac{\partial (a_l \rho_l)}{\partial t} + \nabla \cdot (a_l \rho_l \vec{u}) = -\sum_{k=1}^{Nsp} \dot{m}_{evap,k}^{\prime\prime\prime} \right]$$
(1)

107 where  $\rho_l$  is the liquid phase density,  $\vec{u}$  is the velocity vector and  $\vec{m}_{evap,k}^{\prime\prime\prime}$  (kg/s/m<sup>3</sup>) is the 108 volumetric evaporation rate each of the *Nsp* species composing the liquid phase. The equation 109 (1) can be used for an arbitrary number of phases *Nph* (here *Nph*=2), while the corresponding 110 volume fraction of the gaseous phase can be obtained from the equality  $a_g=1$ - $a_l$ . The momentum 111 equations for both phases are written in the form:

112 
$$\frac{\partial(\rho\vec{u})}{\partial t} + \nabla \cdot (\rho\vec{u}\vec{u}) = -\nabla p + \nabla \cdot [\mu(\nabla\vec{u} + \nabla\vec{u}^T)] + \rho\vec{g} + \vec{F}_{vol}$$
(2)

113 where  $\vec{F}_{vol}$  is the volumetric force due to surface tension. This is calculated based on the 114 Continuum Surface Force (CSF) approach of Brackbill et al. [49] and it is equal to:

115 
$$\vec{F}_{vol} = \sigma \kappa \vec{n} \frac{\rho}{\frac{1}{2}(\rho_l + \rho_g)}$$
,  $\vec{n} = \nabla a$ ,  $\kappa = -\nabla \cdot \left(\frac{\vec{n}}{|\vec{n}|}\right)$  (3)

116 where  $\sigma$  is the surface tension coefficient,  $\vec{n}$  is the normal vector at the interface and  $\kappa$  is the 117 curvature of the interface. Alternatively, the surface tension forces can be included in the 118 momentum equations by using the Continuum Surface Stress (CSS) model by Lafaurie et al. 119 [50] which inherently includes the effects of the variation of the surface tension coefficient 120 along the interface. Equation (4) represents the CSS model, where **I** is the unit tensor and  $\otimes$  is 121 the tensor product of two vectors.

122 
$$\vec{F}_{vol} = \nabla \cdot \left[ \sigma \left( |\vec{n}| \mathbf{I} - \frac{\vec{n} \otimes \vec{n}}{|\vec{n}|} \right) \right] , \ \vec{n} = \nabla a$$
(4)

123

124 Additionally, the transport equations for energy, vapor and liquid species concentration are 125 solved:

126 
$$\frac{\partial(\rho E)}{\partial t} + \nabla \cdot \left[\vec{u}(\rho E + p)\right] = \nabla \cdot \left[k\nabla T - \sum_{q=1}^{Nph} \left(a_q \sum_{k=1}^{Nsp} h_{q,k} \vec{J}_{q,k}\right)\right] - \sum_{k=1}^{Nsp} \dot{m}_{evap,k}^{\prime\prime\prime} L_k$$
(5)

127 
$$\frac{\partial (a_q \rho_q Y_{q,k})}{\partial t} + \nabla \cdot \left( a_q \rho_q Y_{q,k} \vec{u} \right) = \nabla \cdot \left( -a_q \vec{J}_{q,k} \right) + S_{Y,q}$$
(6)

128 
$$\vec{J}_{q,k} = -\rho_q D_{q,k,m} \nabla Y_{q,k}$$
, q=gas or liquid (7)

where  $\vec{J}_{q,k}$  is the dilute approximation of the diffusion flux of species k at the mixture of phase 129 q (neglecting the Soret effect),  $D_{q,k,m}$  is the diffusion coefficient of k<sup>th</sup> species in the mixture of 130 phase q (either gas or liquid). Equation (6) is solved in the gas phase (q=gas) for all the vapor 131 species (k=1÷Nsp) with a source term equal to  $S_{Y,q} = \dot{m}_{evap,k}^{\prime\prime\prime}$ . For the case of multicomponent 132 133 droplets, the same equation is also solved in the liquid phase (q=liquid) for all liquid species with a source term equal to  $S_{Y,q} = -\dot{m}_{evap,k}^{\prime\prime\prime}$ , except of the main liquid species (k=Nsp), since 134 135 its concentration can be computed from the summation of the liquid species concentrations, 136 which must equal to one.

The properties of the pure components are assumed to be a function of temperature according to [51, 52] and they are updated at every iteration during the numerical solution. The mixture properties (either in the gas or the liquid phase for the case of multicomponent evaporation) follow a mass average rule based on the concentration, while the gas phase density is obtained by assuming incompressible ideal gas.

142 The numerical settings used for the simulations are: Laminar flow, explicit VOF solution with 143 the Geometric reconstruction scheme [53], Second Order Upwind (SOU) discretization [54] for 144 the momentum and energy equations, First Order Upwind (FOU) for the species transport 145 equations, Body Force Weighted (BFW) pressure interpolation scheme [55], pressure-velocity 146 coupling with the PISO algorithm [56], variable timestep with Courant number *C*=0.25 both for 147 the interface tracking and the whole computational domain (global Courant number).

148 The commercial CFD code ANSYS FLUENT v14.5 [55] has been utilized for the solution of 149 the flow equations with a number of user defined functions (UDF) for resolving the above 150 processes; more specifically, UDFs have been used for applying the automatic local grid 151 refinement technique and the interface sharpening algorithm [47] required to minimize the 152 interface diffusion which is promoted by the evaporation source terms. Moreover, the 153 evaporation rate model described in the next section is also implemented using UDF together 154 with the functions addressing the fluid, vapor and gas properties as function of local temperature 155 [51, 52].

#### 157 **2.2 Evaporation rate model**

The evaporation rate is calculated as in [42, 43, 57, 58] by applying the simplified Fick's law atthe liquid-gas interface cells:

160 
$$\dot{m}_{evap,k}^{\prime\prime} = \rho_g D_{g,k,m} \left(\frac{dY_{g,k}}{dn}\right)_{surf}$$
, k=1÷Nsp (8)

161 The evaporation source term  $\dot{m}_{evap,k}^{\prime\prime\prime}$  is obtained by multiplying equation (8) with 162  $A_{l,cell}/V_{cell} = |\nabla a|$ . The evaporation rate of equation (8) is independent of the interface shape 163 and does not require any reference length or velocity. Although it is based on a diffusion law, 164 the flow conditions are taken into account through the solution of the Navier-Stokes equations 165 which advect the vapor and modify the local concentration gradient. Besides this simplified 166 Fick's law, a more physically correct expression was also examined which takes into 167 consideration the Stefan flow:

168 
$$\dot{m}_{evap,k}^{\prime\prime} = \frac{\rho_g D_{g,k,m} \left(\frac{dY_{g,k}}{dn}\right)_{surf}}{1 - Y_{g,k,sat}}$$
(9)

Equation (9) has been used in Schlottke & Weigand [41] among others and it is valid only for single-component evaporation. For multi-component evaporation, the exact expression for the evaporation rate was given in Renksizbulut & Bussmann [38] but it was not examined here since it is more complicated and computationally expensive. For low evaporation rates and when the droplet temperature is not close to the boiling temperature (i.e. small Spalding numbers,  $B_M <<1$ ), both expressions are valid.

175 The key parameter in the implementation of the evaporation rate model is the calculation of the 176 concentration gradient magnitude  $(dY_g/dn)_{surf}$ . The interfacial cells are assumed to be saturated and the vapor saturation concentration  $Y_{sat}$  is calculated as a function of the cell temperature 177 178 (and liquid composition when multicomponent evaporation is examined). An algorithm is used 179 to identify the closest neighbor cell (subscript  $\infty$ ) to the direction of the normal vector of the interface  $\vec{n}$ ; this is illustrated in Fig. 1. Denoting as  $\vec{d}$  the vector connecting the two cells, two 180 181 options exist for the calculation of the concentration gradient (see details in Fig. 1) in non-182 orthogonal unstructured grids. The first one is the simplest but less accurate when the vectors  $\vec{n}$ 183 and d are not aligned. The second one is more accurate since it accounts for the angle between 184 the two vectors; it works for all types of grid and it is the default method in the present model.



Fig. 1: Illustration of the two different methods employed for the estimation of the vapor concentration gradient at the liquid-gas interface. The 2<sup>nd</sup> method has been finally adopted in the present work.

Finally, the vapor diffusion coefficient  $D_{g,k,m}$  of the k<sup>th</sup> species in the gas mixture, is obtained by using the dilute approximation (Eq. 10), in which  $X_{g,k}$  is the mole fraction of species k in the gas phase and  $D_{g,i,j}$  is the binary diffusion coefficient of species i in species j; the latter is a function of temperature:

194 
$$D_{g,k,m} = \frac{1 - X_{g,k}}{\sum_{j,j \neq i} (X_{g,j}/D_{g,i,j})}$$
 (10)

195

#### **3 DESCRIPTION OF THE TEST CASES SIMULATED**

197 The mathematical model used is validated against 3 sets of experimental data for suspended droplet 198 evaporation for: (a) convective air flow conditions (experimental data of Daïf et al. [59]), (b) the 199 data of Wong & Lin [60] and (c) under microgravity conditions in a quiescent gas environment 200 (experimental data of Nomura et al. [61]). The cases examined are presented in Table 1; they all 201 refer to atmospheric ambient pressure. As it can be seen, they cover a wide range of conditions 202 including single- and multi-component evaporation, sub- and super-critical gas temperature 203 conditions, low and moderate *Re* numbers, as also species with a large volatility difference (n-204 heptane and n-decane). More specifically, cases 1-3 correspond to the experimental data of Daïf et al. [59] who examined droplet evaporation at low temperature conditions and moderate Re 205

numbers for droplets consisting of n-heptane (case 1), n-decane (case 2) and a mixture of the two with 74% n-heptane (case 3). Case 4 corresponds to the experimental data of Wong & Lin [60] who examined the evaporation of n-decane droplets in a high temperature gas environment and low Re number flow. Cases 5-8 correspond to the experimental data of Nomura et al. [61], who investigated suspended n-heptane droplets in high temperature ambient conditions. Each set of the aforementioned experimental data is examined in a separate section in the present paper, in which the experimental and the numerical setup are presented in detail.

213

ed

215

case	species	$d_0$	$T_{d,0}$	$T_\infty$	$\mathcal{U}_{\infty}$	Re <sub>0</sub>	ref
		[mm]	[K]	[K]	[m/s]		
1	C7	1.052	300	356	3.20	159	[59]
2	C10	1.386	315	348	3.10	210	[59]
3	C7-C10	1.334	294	348	3.10	203	[59]
	74-26%						
4	C10	1.961	315	1000	1.0	17	[60]
5	C7	0.700	300	471	~0	~0	[61]
6	C7	0.700	300	555	~0	~0	[61]
7	C7	0.700	300	648	~0	~0	[61]
8	C7	0.700	300	741	~0	~0	[61]

216

#### 218 4 RESULTS & DISCUSSION

# 4.1 Evaporation at low temperature conditions. Experimental data of Daïf et al. [59]

In this section the numerical model is validated against the experimental data reported in Daïf et al. [59] for cases 1-3 of Table 1. These refer to subcritical ambient temperature conditions and include the evaporation of single- and multicomponent droplets at moderate *Re* numbers. In [59] the droplets were held in suspension at the spherical head of a capillary tube (400 and 200µm respectively), which was placed perpendicularly to the gas flow. They used a video camera to record the droplet diameter and an infrared thermo-graphic system to measure the droplet's surface temperature.

228 The computational domain, the boundary conditions and the grid used for these cases are shown 229 in Fig. 2a; they are similar to these used in Strotos et al. [42]. The flow was assumed to be 2D-230 axisymmetric since the Re number is below 400 for all cases examined and the presence of the 231 capillary tube is not expected to induce 3D effects. The spherical head of the suspender was 232 assumed to be an adiabatic impermeable wall and the liquid contact angle was set to 10 degrees 233 as in [42]. Regarding the initial conditions, uniform velocity, temperature and concentration 234 fields were assumed. The grid used is unstructured with 2 levels of adaptive local refinement; 235 this is based on the work of Theodorakakos & Bergeles [62] and implemented here as in 236 Malgarinos et al. [45]. A grid independent solution is achieved by using 2 levels of local grid 237 refinement as shown in Fig. 2b for the prediction of the dimensionless droplet volume temporal 238 evolution. This is a good compromise between accuracy and computational cost which is 239 approximately 1.7CPU-days for each second of physical time for the single-component cases 240 and 2.3CPU-days/sec for the bi-component evaporation case.



Fig. 2: (a) Computational domain and boundary conditions employed. The inset figure shows the grid topology around the liquid-gas interface for two levels of adaptive local grid refinement. (b) Temporal evolution of droplet's non-dimensional volume for the case 1 using one-, two- and three-levels of local grid refinement.

248 The temporal evolution of the non-dimensional droplet squared diameter and the mean droplet 249 temperature for the cases of Daïf et al. [59] are presented in Fig. 3a and Fig. 3b, respectively. 250 The predictions are in good agreement with the experimental data regarding the droplet size, 251 while the wet-bulb temperature of n-decane is overestimated by 4°C; this is within the accuracy of previously published studies for the same case [25, 42, 63, 64]. Single-component droplets 252 253 (cases 1 and 2) after an initial heat-up period follow the  $d^2$ -law and they reach the wet-bulb temperature corresponding to the experimental conditions. The two-component droplet (case 3) 254 255 exhibits preferential evaporation in which the more volatile n-heptane evaporates first and then 256 the less volatile n-decane follows. The relevant physical processes during the multicomponent 257 evaporation have been discussed in depth discussed in Strotos et al. [42, 65].



Fig. 3: Model predictions for the experimental data of Daïf et al. [59]. Temporal evolution of (a) droplet size and (b) mean droplet temperature. No experimental data were provided for the droplet temperature in case 1.

264 Initially, the spherical suspender lies at the centre of the droplet; under the action of 265 gravitational and aerodynamic forces the droplet moves either upwards or downwards as shown 266 in Fig. 4. The n-heptane droplet (case 1) is relatively light and moves upwards (opposite to the 267 direction of gravity in Fig. 4), while the droplets of the other two cases are heavier and they move initially downwards (in the direction of gravity in Fig. 4). As the droplet evaporates and 268 269 becomes lighter, the gravitational force become smaller compared to the aerodynamic one and 270 the droplet progressively moves to the other side of the suspender. After reaching an 271 equilibrium position, the droplets are held in touch with the suspender due to the adhesion 272 forces. Behind the droplet, a large recirculation zone appears and its length is plotted versus the 273 gas phase *Re* number in Fig. 5; the experimental data of Taneda [66] for a solid sphere are also 274 shown. The wake length increases with increasing Re number and it is slightly restricted 275 compared to the one formed under the same flow conditions behind a solid sphere. This is 276 attributed to the liquid phase recirculation which moves the flow separation point towards the 277 rear of the droplet and suppresses the length of the wake, as also reported previously in 278 Sirignano [2].



280

Fig. 4: Predicted transient droplet motion and flow field streamlines for the experiments of Daïf

282 et al. [59] (cases 1-3).





Fig. 5: Dependence of the wake recirculation length with the *Re* number for cases 1-3.

286

In Fig. 6 representative velocity, temperature and vapor fields are presented for the n-decane case at 0.25 and 20.0sec; the velocity field is depicted by the streamlines colored with the velocity magnitude. The spatial distribution of the temperature and the vapor field is influenced by the recirculation zones formed inside and behind the droplet, as also the presence of the solid suspender. Inside the droplet, a relatively cooler region is identified at the vortex core (as also observed in [29, 30]), while at the wake behind the droplet the temperature and the vapor concentration fields are similar between them and they are influenced by the gas phase vortex.

294



Fig. 6: Velocity streamlines, temperature and vapor concentration field for the n-decane droplet
(case 2).

299 In Fig. 7 the spatial distribution of the evaporation source term (in kg/m<sup>3</sup>s) at selected time 300 instances is presented for the n-decane droplet; the non-interpolated contours shown correspond 301 to the region between VOF values equal to 0.01 and 0.99. As it can be seen, there is a narrow 302 region at the interface (2-3 cells) in which the evaporation source term is distributed and it 303 maximizes near the 0.5 iso-VOF line, in which the quantity  $|\nabla a|$  maximizes too (see also the 304 inset figure at t=20 sec). The thickness of this region in length units depends on the grid density, 305 but it has a minor effect on the overall evaporation process, as shown in Fig. 2b. The 306 evaporation rate increases with time, since the droplet temperature also increases; it is 307 interesting to observe that the maximum evaporation rate is observed around 30-40° away from 308 the front stagnation point, which was also observed in Shih & Megaridis [31]. The minimum 309 evaporation rate is observed near the point of flow separation at approximately 130-140°; this 310 behavior was also observed in Haywood et al. [28]. The spatial distribution of the evaporation rate is not smooth and this is ought to the imperfect distribution of the VOF values over the 311 312 computational cells.



Fig. 7: Spatial distribution of the evaporation rate at different time instances for the n-decane
droplet (case 2).

318 One of the parameters examined was the effect of the thermo-capillary flow (Marangoni effect) 319 induced by the variation of the surface tension coefficient along the interface. This is achieved 320 by using the CSS formulation [50] instead of the CSF model [49] in order to calculate the 321 surface tension forces in the momentum equations. As it was also shown in Strotos et al. [42], 322 the temperature along the interface does not exhibit large variations and the Marangoni effect 323 does not affect the droplet evaporation for these specific cases. On the other hand, it seems to 324 create a kind of unsteadiness by increasing the fluctuations of the droplet shape but without a 325 noticeable effect on the liquid phase internal circulation; unsteadiness in thermo-capillary flows 326 were also observed in Shih & Megaridis [32]. The single-component cases 1-2 were also 327 examined by using the evaporation rate of equation (9). The results have shown that it has a 328 negligible effect on the prediction of the phenomenon due to the low evaporation rate and thus it 329 is not presented.

330

314

# 4.2 Evaporation at high temperature conditions. Experimental data of Wong & 332 Lin [60]

The 4<sup>th</sup> case of Table 1 corresponds to the experimental data of Wong & Lin [60] who examined the evaporation of n-decane droplets in a supercritical gas environment ( $T_{\infty}$ = 1000K,  $T_{cr,C7}$ =617.7K). The droplet was suspended at a ceramic shell attached to a 100µm glass filament; the droplet internal temperature distribution was measured by using thermocouples (25µm wire diameter and ~70µm bead diameter) placed at different positions inside the droplet. This set of experimental data has been widely used in validation studies for 0-D and 1-D models [16, 19, 63, 67, 68] as also in CFD studies by [30, 33], which assumed a spherical droplet shape and neglected the presence of the shell-like suspender. In the present study the presence of the suspender has been considered but assumed to be spherical since its exact geometry and dimensions were not reported in the relevant paper.

343 The model predictions for the droplet size and the mean droplet temperature are shown in Fig. 8 344 by using the two variations of the evaporation model described in section 2.2, i.e using equation 345 (8) and (9), respectively. The predictions for the droplet size are satisfactory for both models 346 with the second one having a slightly higher evaporation rate at the beginning and lower at 347 subsequent times. In both cases the droplet's size regression deviates significantly from the  $d^2$ -348 law due to the intense and long droplet heat-up period. During this period, the thermal 349 expansion of the droplet is more intense compared to the droplet evaporation and the droplet 350 size slightly increases; this phenomenon has been observed in [39, 69] among others. Regarding 351 the model predictions for the droplet temperature, the available experimental data for the droplet 352 temperature were obtained by placing three thermocouples at fixed positions ( $r/R_0=0, \pm 0.6$ ). 353 Such information was difficult to be obtained with the current numerical setup since the droplet 354 moves over the suspender and also deforms. So, in Fig. 8 the model prediction for the mean 355 droplet temperature is presented, which is closely related to the measured quantities. The 356 temperature prediction is in accordance with the experimental measurements but this time the 357 simplified Fick's model of eq (8) has a better performance predicting accurately the wet-bulb 358 temperature. The present results are within the accuracy of the results obtained with other 359 numerical methodologies mentioned at the beginning of this section. It has to be noted that at 360 the latter stages of evaporation, the interface diffusion problems associated with the VOF 361 methodology become more pronounced and are more intense for high evaporation rates; this 362 explains why the droplet temperature predicted with equation (9) further increases for t > 2.5 sec.

363



Fig. 8: Model predictions for the experimental data of Wong & Lin [60]. (a) temporal evolutionof droplet size and (b) mean droplet temperature.

367

368 The predicted flow, temperature and vapor concentration field for the experimental case of [60] 369 are shown in Fig. 9. The gravitational forces are quite strong and the droplet moves downwards 370 to the direction of gravity. Since the *Re* number of the flow is below 20, there is no recirculation 371 in the gas phase downwind the droplet and the temperature and vapor concentration fields have 372 a slightly different topology compared to the cases 1-3 presented in the previous section 4.1. 373 Furthermore, the thickness of the temperature and concentration boundary layers seem to be 374 higher compared to those of cases 1-3 and this ought to the fact that the boundary layer thickness is proportional to  $(RePr)^{-1/2}$  [70] as also to the thickening of the boundary layer due to 375 376 the intense evaporation rate [19].

377 Inside the droplet, the liquid circulation is quite disturbed forming two unsteady vortices which 378 tend to homogenize the temperature. This is in contradiction with the experimental observations 379 of Wong & Lin [60] as also with the numerical solution of Megaridis [30]. The inability to 380 capture this phenomenon in the current work is probably ought to the parasitic velocities 381 induced by the VOF methodology [71, 72], which originate from the imbalance between surface 382 tension and pressure gradient forces. The case examined has a low Re number and the parasitic 383 velocities become important while they are further magnified due to the high density ratio 384 between the liquid and the gas phase, which exceeds 2000 in this case. Nevertheless, the overall 385 behavior of droplet evaporation is not affected by the spurious currents and the experimental 386 data are correctly predicted. In an effort to reduce the magnitude of these velocities, several 387 options were examined, such as reducing the computational time-step down to a Courant 388 number of 0.1, increasing the smoothing iterations of VOF function and using different 389 discretization schemes for the VOF; however, none of these methods could improve the predicted velocity field. The unphysical velocities could be potentially minimized by applying 390 391 the algorithm described in Francois et al. [72], but the pressure-velocity coupling in FLUENT is 392 hardcoded and cannot be modified as stated in [73]. The CSS surface tension scheme was also 393 examined, but the induced thermo-capillary flow along with the spurious velocities resulted in 394 high deformation of the droplet shape. Thus, the CSS simulations are not considered accurate 395 and they are not presented.



Fig. 9: Predicted flow field streamlines, temperature and vapor concentration field for the
experimental case of Wong & Lin [60].

# 4024.3Evaporation at high temperature conditions. Experimental data of Nomura403et al. [61]

404 Cases 5-8 of Table 1 correspond to the experimental data of Nomura et al. [61], who examined 405 the evaporation of suspended n-heptane droplets for a wide range of ambient temperature and 406 pressure conditions; these data have been widely used to validate 0-D and 1-D models (see [18, 407 39] among others). The experiments were conducted at microgravity conditions to prevent the 408 natural convection and the vapor was flowing radially as stated in Nomura et al. [61]. The 409 droplet diameters examined were in the range 0.6-0.8mm but the exact values were not reported. However, as the time axis of their plots was presented as  $t/d_0^2$  this normalization results to the 410 411 prediction of vaporization time independent on the selection of the droplet diameter. In the present work, a diameter of 0.7mm was selected as the mean among the experimental values. 412 413 Regarding the initial droplet temperature, a value of 300K was used. Simulations of the cases of 414 [61] were simplified by disabling the solution of the flow equations and accounting only for the 415 transient and the diffusion terms in the conservation equations. This selection is justified by the 416 fact that the velocities are quite low and the vapor flows radially [61]. Note that this is not 417 equivalent to a 1-D model which assumes spherical symmetry, since a 2-D computation domain 418 is used (Fig. 10) and the grid cells are not aligned with the liquid-gas interface (see inset figure 419 in Fig. 10). More importantly, the gas phase temperature and concentration are also solved, so 420 there is no need to estimate "film" gas properties near the interface, as also the Nusselt and

421 Sherwood numbers are not a-priori assumed, but become a part of the solution through the 422 predicted temperature and vapor concentration gradients. It has also to be noted that this 423 approach is not applicable in the experiments examined in the previous sections, since the gas 424 phase velocity plays a dominant role. The grid used is shown in Fig. 10 and it represents an 425 adiabatic box extended  $25d_0$  far from the droplet. The box dimensions are large enough so that 426 the average gas temperature remains practically constant.

427



428

Fig. 10: Numerical grid used for the experimental data of Nomura et al. [61].

The model results for the dimensionless squared droplet diameter are shown in Fig. 11a for the four different ambient temperatures examined. As it can be seen, the model results are satisfactory for a wide range of ambient temperatures even above the critical one  $(T_{cr,C7}=540.2K)$ . The droplet evaporation rate is shown in Fig. 11b. The evaporation rate increases at the initial stages of the evaporation due to droplet heat up and after reaching a maximum it decreases due to the reduction of the droplet surface area. This behavior is observed in all cases and it generally increases with increasing ambient temperature.





440 Fig. 11: Model predictions for the experimental data of Nomura et al. [61]. (a) temporal 441 evolution of droplet size and (b) evaporation rate.

443

#### 444 **5 CONCLUSIONS**

445 The Navier-Stokes equations coupled with the conservation equations for energy and species 446 concentration and the VOF methodology were used to study the evaporation of suspended 447 single- and multi-component droplets. The local evaporation rate was estimated by using the 448 Fick's diffusion law, which is independent of the interface shape and grid density. The model 449 performance was assessed by comparing the results against experimental data for single- and bi-450 component fuel droplets under sub- and super-critical ambient temperature conditions. 451 Prediction of the overall evaporation rate for a wide range of cases was satisfactory, while 452 prediction of the droplet's temperature was within the accuracy of other published works. For 453 the cases with Re of O(100) the recirculation zones at the gas and the liquid phase are correctly 454 predicted and agree with the numerical solutions from other researchers. For the case with Re of 455 O(10), the model could not capture the liquid phase internal recirculation zone since the 456 parasitic velocities induced by the VOF methodology were large compared to the flow velocity. 457 Overall, the numerical model can be considered as a reliable tool able to predict the lifetime and 458 temporal evolution of droplet evaporation in a wide range of ambient temperature conditions for 459 moderate and high Re numbers. In comparison to simplified (but also faster) 0-D and 1-D 460 models, the present model gives insight into the spatial distribution of the flow variables and 461 more importantly, it can be extended without any modification to predict the evaporation of 462 deformed droplets and droplets undergoing breakup; see [74]. Nevertheless, with the today's 463 computational resources, the present model cannot be used to simulate sprays comprised by a 464 large number of droplets.

465

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### **7 NOMENCLATURE**

Symbol	Quantity	SI Unit
Α	Surface area	m <sup>2</sup>
$B_M$	Spalding number $B_M = (Y_{g,sat} - Y_{g,\infty})/(1 - Y_{g,sat})$	-
Cp	Specific heat capacity	$J/(kg \cdot K)$
d	Diameter	m
đ	Vector connecting adjacent cells	m
D	Mass diffusion coefficient	m <sup>2</sup> /s
Ε	Specific internal energy	J/kg
F	Force	Ν
g	Gravity acceleration	m/s <sup>2</sup>
h	Specific enthalpy	J/kg
Ι	Unit tensor	-
$\vec{J}_{q,\mathbf{k}}$	Species flux	kg/m <sup>2</sup> s
k	Thermal conductivity coefficient	$W/(m \cdot K)$
L	Latent heat of vaporization	J/kg
<i>ṁ</i> ′′	Evaporation rate	kg/m <sup>2</sup> s
<i>ṁ</i> ′′′	Evaporation source term	kg/m <sup>3</sup> s
n	Normal coordinate to gas-liquid interface	-
Nph	Total number of phases	-
Nsp	Total number of liquid species	-
р	Pressure	N/m <sup>2</sup>

Pr	Prandtl number	-
R	Radius	m
r	Coordinate in radial direction	m
Re	Reynolds number $Re = \rho_{g,\infty} d_0 u_{\infty} / \mu_{g,\infty}$	-
S	Source term	
Т	Temperature	K
t	Time	S
и	Velocity	m/s
V	Volume	m <sup>3</sup>
<i>x,y,z</i>	Coordinates	m
Y	Mass concentration	kg/kg

# 474 Greek Symbols

Symbol	Quantity	SI Unit
α	Liquid volume fraction	-
К	Curvature	$m^{-1}$
μ	Viscosity	kg/(m·s)
ρ	Density	kg/m <sup>3</sup>
σ	Surface tension coefficient	N/m

475

## 476 Subscripts

Symbol	Quantity
0	Initial
$\infty$	Conditions far from the droplet

cell	Cell
cr	Critical
d	Droplet
evap	Evaporation
exp	Experimental
g	Gas phase
k	k <sup>th</sup> species of the mixture
l	Liquid phase
m	Mixture
mean	Mean
q	of phase q
sat	Saturated
surf	Surface interface
vap	Vapor

### **abbreviations**

C10	n-decane
C7	n-heptane
CSF	Continuum Surface Force
CSS	Continuum Surface Stress
ECM	Effective Conductivity Model
FCM	Finite Conductivity Model
FDM	Finite Diffusivity Model
ICM	Infinite Conductivity Model

	IDM	Infinite Diffusivity Model	
	VOF	Volume Of Fluid	
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480			
481	8 REFERENC	DES	
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