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L. García-Fernández, P. Fruton, Henri Bataller, J. M Ortiz de Zárate, Fabrizio Croccolo. Coupled non-equilibrium fluctuations in a polymeric ternary mixture. European Physical Journal E: Soft matter and biological physics, EDP Sciences: EPJ, 2019, 42 (11), 10.1140/epje/i2019-11889-4. hal-02473051

HAL Id: hal-02473051 https://hal.archives-ouvertes.fr/hal-02473051

Submitted on 12 Feb 2020 $\,$

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- Coupled non-equilibrium fluctuations in a polymeric ternary mixture 1 L. García-Fernández^{1,2}, P. Fruton¹, H. Bataller¹, J.M. Ortiz de Zárate³ and F. Croccolo¹ 2 3 ¹Laboratoire des Fluides Complexes et leurs Réservoirs – IPRA, UMR5150, E2S-Univ Pau & 4 5 Pays Adour/CNRS/Total, 1 Allée du Parc Montaury,64600, Anglet, France. ²Centre National d'Études Spatiales (CNES),2, Place Maurice Quentin, 75001 Paris, France. 6 7 ³Departamento de Estructura de la Materia, Física Térmica y Electrónica, Facultad de 8 Ciencias Físicas, Universidad Complutense de Madrid. Plaza de las Ciencias 1, 28040 9 Madrid, Spain.
- 10

11 Abstract

We investigate by dynamic shadowgraphy the non-equilibrium fluctuations at the steady state 12 of a thermodiffusion experiment in a polymeric ternary mixture of polystyrene-toluene-n-13 hexane. The structure function of the refractive index reveals the existence of quite different 14 15 decay times, thus requiring the analysis of a wide range of correlation times. This is related to the simultaneous presence of three distinct decay modes corresponding to (from fastest to 16 slowest) relaxation of temperature fluctuations, of the concentration fluctuations of the mixed 17 solvent, and of the concentration fluctuations of the polymer in the binary solvent. An 18 investigation of the decay times at the corresponding diffusive regimes provides a measurement 19 of the thermal diffusivity and the two eigenvalues of the mass diffusion matrix of the ternary 20 mixture. Similar experiments were performed in the past but here, to suppress the confinement 21 effect and obtain a more direct comparison with the theory, a thicker sample is studied. 22 23 Moreover, also a faster camera is used allowing the experimental observation of faster modes, like the propagative ones. The experimental values of the decay times are eventually compared 24 with those predicted by different available theories. Finally, we present a more complete 25 theoretical model to describe the non-equilibrium fluctuations in the bulk of a ternary mixture 26 at the steady state of a thermodiffusion experiment. 27

Keywords: Multicomponent mixtures, Polymer, Thermodiffusion, Shadowgraphy, Nonequilibrium, Fluctuations

30

31 **1. Introduction**

The majority of the fluids encountered in nature and industry are multicomponent mixtures. 32 The knowledge of their transport properties is crucial for many different applications from both 33 scientific and technological points of view. The efficiency of exploitation of the crude oil wells, 34 as well as the environmental alternative focused on the CO₂ storage in deep brine aquifers, are 35 36 some recent examples showing the interest of studying the transport properties of multicomponent mixtures subjected to non-equilibrium conditions [1, 2]. Complex fluids in 37 non-equilibrium conditions exhibit giant fluctuations of the thermodynamic variables, the so-38 39 called non-equilibrium fluctuations (NEFs) [3-5]. By analysing NEFs through light scattering techniques, the transport properties of the fluid can be determined both at atmospheric [6] and 40 at high pressure [7]. 41

The non-equilibrium condition can be induced, for example, by applying a temperature gradient 42 to a multicomponent fluid mixture, thus inducing a composition gradient within the fluid by 43 44 means of the thermodiffusion, or Soret effect [8-10]. This transport process can be investigated by optical techniques and particularly by light scattering thanks to its ability to visualize NEFs 45 without altering the intrinsic properties of the fluid. In this work, dynamic shadowgraphy has 46 47 been adopted [11-13] to study the fluctuations of the refractive index as generated by the NEFs 48 of the thermodynamic variables, like temperature and concentrations. By shadowgraphy, a large range of fluctuations sizes λ or, conversely, wave numbers $q = 2\pi/\lambda$, can be investigated at 49 the same time. This approach, combined with statistical analysis, represents a powerful 50 characterization technique able to provide simultaneous reliable measurements of different 51

transport properties, like mass diffusion coefficient or thermal diffusivity as well as Soret
coefficient [6, 14, 15].

A suitable characterization of transport processes in complex mixtures requires a deep 54 understanding of simpler fluids. So far, only binary mixtures [10, 16] have been extensively 55 characterized. The extension of theories and experiments from binary to ternary mixtures 56 requires further development due to the intrinsic and significant increase of difficulty with the 57 number of components of the mixture. Currently, a great effort is devoted to investigate the 58 transport phenomena in ternary mixtures, as performed in the present study. Thermodiffusion 59 experiments in ternary mixtures are performed on ground, by using different optical techniques 60 61 [10, 14, 17, 18], or in microgravity in order to avoid both convection and sedimentation. The thermodynamic characterization of ternary mixtures is one of the objectives of the following 62 ESA projects: Diffusion Coefficient Measurements in ternary mIXtures (DCMIX) [18-23], 63 64 Soret Coefficients in Crude Oil (SCCO) [1, 24-27] and Giant Fluctuations [28], as further elaborated in a recent colloquium paper [29]. 65

As a benchmark ternary mixture, the tetraline, isobutylbenzene and n-dodecane sample has been 66 studied by digital interferometry on ground and in microgravity within the DCMIX experiments 67 [18, 19, 30-38]. In addition, Bataller et al. [17] investigated this mixture on ground by one-68 69 wavelength shadowgraphy. Since no other alternative was available at that time, Bataller et al. [17] compared their experimental results with the theory of concentration NEFs developed for 70 ternary mixtures in the absence of gravity [39]. The two concentration modes related to the 71 presence of the two independent components could not be distinguished experimentally, and 72 only an average of the two eigenvalues of the mass diffusion matrix could be provided. This 73 74 result is expected, as the mixture is composed of similar-size molecules, and consequently the two eigenvalues of the diffusion coefficient matrix are very similar to each other. The two 75 eigenvalues \hat{D}_1 and \hat{D}_2 of the diffusion matrix **D** need to be different enough to distinguish the 76

two concentration modes. An equivalent way to set this statement is to define a dimensionless 77 number $Dr = \hat{D}_2/\hat{D}_1$ (Dr for Diffusion eigenvalue Ratio and \hat{D}_1 representing the slowest 78 diffusion mode) and require it to satisfy the relation $Dr \gg 1$. A ternary mixture containing a 79 polymer dissolved in a binary mixed solvent is a good candidate for such measurement, due to 80 81 the very different molecular size of its components and, correspondingly, a value of $Dr \approx 10$. This was experimentally proved for the diluted polymeric ternary mixture of polystyrene-82 toluene-n-hexane, as reported in [14], where a layer of L = 2 mm thick was investigated by 83 84 using one-wavelength shadowgraphy at a single image acquisition frequency of 28 Hz. By adopting the theoretical model of ternary mixtures in the absence of gravity, the dynamics of 85 the NEFs could be analysed for large wave numbers [14]. From such analysis, it was only 86 possible to determine the two eigenvalues of the mass diffusion matrix and the thermal 87 diffusivity. Afterwards, the diffusion matrix was simplified by neglecting the off-diagonal 88 elements and the result was utilised for obtaining an analytical relationship between the two 89 Soret coefficients, without an independent measurement of them. More recently [40], we have 90 further developed the theory of NEFs to include the effect of gravity, but not confinement so 91 92 that a direct comparison with the data of Ref. [14] is impossible since fluid layer thickness of only 2 mm makes confinement effects important, as evidenced in related literature for binary 93 mixtures [41, 42]. 94

In the present study, we perform a new shadowgraph experiment on a similar ternary mixture, but in this case a thicker sample with vertical extension of L = 5 mm is analysed in order to avoid the confinement effect, and be able to compare the experimental results with the theory of NEFs including gravity [40]. The present measurements are also carried out with a faster camera, allowing a more accurate characterization of thermal fluctuations and, thus, a better separation of them from concentration ones. Additionally, a new procedure of image acquisition combined with the concatenation of the structure functions allows us to investigate a wider

range of correlation times, as required for a complete analysis of this mixture that has a wide 102 range of decay times. The experimental results are thus compared for the first time with the 103 theoretical model developed for ternary mixtures in the presence of gravity [40], as well as with 104 105 the recently developed theory that includes the coupling of velocity, thermal and solutal NEFs in the case of binary mixtures [43]. These comparisons suggest that a further development of 106 the theory is required, in particular to include the coupling between the different NEFs in the 107 108 case of the ternary mixture. As part of the contents of this paper we present such a development and compare with the experimental data. 109

110

111 **2.** Experimental procedure

112 2.1 Polymeric solution preparation and characterization

113 The sample is prepared by dissolving 2 wt% of polystyrene (PS, $M_{\omega} = 4730 \ g/mol$, DIN-114 Poly(styrene) 4730 DA – PDI 1.03 by PSS-polymer) in 49 wt% toluene (Sigma-Aldrich, 115 24,451-1, 99.8%) and 49 wt% n-hexane (Sigma-Aldrich, 13,938-6, >99%) at ambient 116 temperature by using a magnetic stirrer. The thermophysical properties of the mixture, such as 117 density, viscosity and both thermal and mass expansion coefficients are determined.

118 The kinematic viscosity ν is measured at 25°C by a capillary viscometer (Ubbelohde 119 SCHOTT). The thermal and mass expansion coefficients, α and β_i , respectively, are defined as 120 follows:

121
$$\alpha = -\frac{1}{\rho} \left(\frac{\partial \rho}{\partial T} \right)_{\omega, p}, \tag{1}$$

122
$$\beta_i = \frac{1}{\rho} \left(\frac{\partial \rho}{\partial \omega_i} \right)_{T,p},$$
 (2)

where ρ is the mixture density, *T* is the temperature and ω_i is the concentration of component *i* in mass fraction. Following a common notation, components are ordered according to their (pure) density, hence, in this study 1 corresponds to PS, 2 corresponds to toluene and 3corresponds to n-hexane.

The coefficients α and β_i are indirectly determined from measurements of the mixture density 127 performed at different conditions through a Density Meter (ANTON PAAR, DMA 5000). For 128 the thermal expansion coefficient, the density is measured at different temperatures (from 23 to 129 27°C), while keeping the concentrations constant at $\omega_1 = 0.02$ and $\omega_2 = 0.49$. The mass 130 expansion coefficients are obtained after measuring the density at different concentrations while 131 keeping the temperature constant at 25°C. To obtain the mass expansion coefficient of the 132 polymer in the mixed solvent, β_1 , the concentration of the polystyrene is changed from $\omega_1 =$ 133 0.000 to $\omega_1 = 0.035$, keeping $\omega_2 = \omega_3$. For the mass expansion coefficient of the mixed 134 135 solvent, β_2 , the composition of the mixed solvent is modified from $\omega_2 = 0.47$ to $\omega_2 = 0.51$, keeping $\omega_1 = 0.02$ constant. Measured values of ρ , ν , α and β_i are summarized in Table 1. The 136 values of ρ , ν , α and β_2 are consistent with those reported in [44], determined for the equi-137 massic toluene-n-hexane binary mixture at $T_{mean} = 25 \ ^{\circ}C$. The measurements of viscosity and 138 139 density are repeated at least three times for each sample. The values reported in Table 1 are calculated as the average of the three measurements and the corresponding uncertainties are the 140 standard deviations. The parameter α is determined through Eq. 1, i.e. dividing the slope of the 141 density vs. temperature by the density of the sample at the nominal temperature. Similarly, β_i 142 are calculated through Eq. 2, i.e. dividing the slope of the density vs. concentration by the 143 density of the sample at the nominal concentration. 144

145 146

Table 1: Thermophysical properties of the polystyrene-toluene-n-hexane mixture

Parameter	Value
ρ	$(750.0 \pm 0.4) \times 10^{-3} \text{ g/cm}^3$
ν	$(5.24 \pm 0.01) \times 10^{-3} \text{ cm}^2/\text{s}$

α	$(9.1 \pm 0.7) \times 10^{-4}/\mathrm{K}$
eta_1	$(3.1 \pm 0.1) \times 10^{-1}$
β_2	$(2.6 \pm 0.1) \times 10^{-1}$

147

148 2.2 Thermodiffusion experiment

The experimental set-up is quite similar to the one used in previous studies and a more detailed 149 150 description can be found elsewhere [6, 7]. Before filling our thermodiffusion cell, its interior is evacuated by using a vacuum pump for more than four hours in order to evaporate all the 151 residual fluids. Subsequently, the mixture is injected into the cell. The sample is sandwiched 152 between two square sapphire windows of 8 mm thick, mechanically separated to provide a 153 vertical thickness of the fluid layer of $L = (5.0 \pm 0.1) mm$. Two thermoelectric (TEC) Peltier 154 devices are used to apply the temperature gradient to the fluid mixture. Both TECs are 155 independently connected to two proportional-integral-derivative (PID) temperature controllers 156 (Wavelength Electronics, MODEL LFI - 3751), being able to set the temperatures of the two 157 sides of the fluid slab with an absolute accuracy of 0.01 K and a relative RMS stability of 1 mK 158 159 over 24 hours. The temperature gradient is applied in the z direction, and the sample is always 160 heated from above in order to avoid convection, since both Soret coefficients are expected to 161 be positive [44-46]. When applying a temperature gradient, the two TECs actually act as heat pumps in the same direction, thus a heat exchanger flushed with water coming from a thermostat 162 (HUBER, ministate 125) removes the excess heat from the external side of the Peltier elements. 163 The thermodiffusion experiments are performed by imposing temperature differences of $\Delta T =$ 164 165 10, 15 and 20 K over the vertical extension of the sample cell. After imposing the temperature gradient, and before starting the image acquisition through the shadowgraph setup, it is 166 necessary that the sample reaches the stationary state where the Soret effect is completely 167 balanced by Fickean diffusion. In our experimental conditions, the diffusion time $\tau = L^2/D$ is 168

of the order of $8.3 \times 10^4 s$, i.e. about 23 hours. This value is obtained from the diffusion coefficient of the polymer in a typical solvent, like toluene $D \approx 3 \times 10^{-6} cm^2/s$ [45].

171 Density fluctuations $\delta\rho$ generated by both temperature fluctuations δT and concentration 172 fluctuations $\delta\omega_i$ give rise to refractive index fluctuations $\delta n = \delta T(\partial n/\partial T) +$ 173 $\sum_i \delta\omega_i (\partial n/\partial \omega_i)$, which can be detected by optical techniques, as described in the literature 174 [11, 13, 47]. In general, velocity fluctuations are not supposed to induce fluctuations of the 175 density or of the refractive index.

176 In our optical setup, a super-luminous diode (Superlum, SLD-MS-261-MP2-SM) with a wavelength of $\lambda = (675 \pm 13)$ nm acts as the light source, illuminating the sample parallel to 177 the temperature gradient. The light is provided out of a monomode optical fibre and the beam 178 is collimated by an achromatic doublet lens with focal length f = 150 mm and placed at its 179 focal distance from the fibre output. After the collimating lens, a linear polariser is positioned 180 181 before the sample and a second linear polariser (analyser) after it, providing a control of the light intensity independent of the integration time of the detector. Finally, the light beam 182 183 impinges onto a scientific-CMOS camera (Hamamatsu Digital Camera C13440, ORCA - Flash 4.0) whose detector size is s = 1.33 cm, placed in the near field, i.e. at a distance z =184 (200 ± 5) mm from the sample midplane. The s-CMOS camera allows a 'fast' image 185 acquisition frequency up to 100Hz at full frame (2048×2048 pixels), thus enabling a fruitful 186 characterization of the temperature fluctuations in the observable wave number range. The 187 shadowgraph signal is the result of the interference of the transmitted beam with the ones 188 scattered by the sample refractive index fluctuations. In our experiment series of N = 2500189 images of 2048 × 2048 pixels, corresponding to $q_{min} = 2\pi/s = 4.72/cm$, are recorded for 190 three different frequencies of f = 100, 10, 1 Hz, corresponding to minimum correlation times 191 of $dt_{min} = 0.01, 0.1, 1 s$. The recorded images are eventually analysed by the Differential 192 193 Dynamic Algorithm (DDA) briefly described in the following section [48].

194

195 *2.3 Dynamic shadowgraphy*

The DDA analysis consists of the following steps: images are recorded at different times as bi-196 dimensional intensity maps $I(\vec{x}, t)$ and are 2D-Fourier transformed and normalized to get 197 $i(\vec{q},t) = I(\vec{q},t)/I(0,t)$. Then, differences between pairs of normalised Fast Fourier 198 Transforms (FFTs) are calculated and their square moduli are determined $|\Delta i(\vec{q}, t, dt)|^2 =$ 199 $|i(\vec{q},t) - i(\vec{q},t+dt)|^2$. These quantities are computed for all possible delay times dt, 200 multiples of the dt_{min} set by the acquisition frequency of the detector. Finally, the results are 201 averaged first over time $\langle |\Delta i(\vec{q}, dt)|^2 \rangle = \langle |\Delta i(\vec{q}, t, dt)|^2 \rangle_t$ and second over the modulus of the 202 wave vector \vec{q} , $\langle |\Delta i(q, dt)|^2 \rangle = \langle |\Delta i(\vec{q}, dt)|^2 \rangle_{|\vec{q}|}$. The resultant function is the so-called 203 Structure Function (SF) $\langle |\Delta i(q, dt)|^2 \rangle$. All these calculations are performed by a custom 204 205 software running on a graphic card and taking advantage of the massive parallelization of the Graphic Processing Unit (GPU), thus requiring about one hour of computational time for each 206 207 experiment, i.e. the three series of 2500 images of 2048×2048 pixels [49].

208

209 2.4 Concatenation of the experimental structure function

As mentioned above, in the experiments reported in this study, sets of images of 2048×2048 210 pixels have been recorded at three different acquisition frequencies at the steady state of the 211 thermodiffusion process. Then, the experimental SF is determined by analysing each stack of 212 images acquired at a specific frequency. Examples of $\langle |\Delta i(q, dt)|^2 \rangle$ are shown in Fig. 1, as a 213 function of the wave number (Fig. 1a, 1c, 1e) and as a function of the correlation time dt for 214 different wave numbers (Fig. 1b, 1d, 1f). The SFs obtained at three different frequencies are 215 216 then merged in order to form a single concatenated SF, or c-SF (Fig. 1g), that covers a large range of correlation times of about four orders of magnitude, as shown in Fig. 1h. 217

As stated above, the c-SF is the result of merging the SFs determined at three different 218 frequencies (from left to right in Fig. 1h): the first points correspond to the data acquired at 100 219 Hz (from Fig. 1b, dark purple squares at short dt), the second ones to 10 Hz (from Fig. 1d, light 220 cyan circles at intermediate dt) and the third ones to 1 Hz (from Fig. 1f, dark orange triangles 221 at large dt). The concatenation procedure is made by simply selecting the relevant part of the 222 SFs as shown in Fig. 1. About the 10% of the delay times of each data set are used for the 223 224 concatenation, disregarding those that are already included in the slower frequency slot. This means that from the fastest acquisition set only delay times in the range 0.01-2.5 s are taken, 225 from the intermediate one the selected range is 2.5-25 s and from the slowest one 25-250 s. The 226 concatenation procedure has been already applied in our precedent work [43], but here it is 227 described in more detail. We point out that this scheme becomes useful whenever it is necessary 228 to investigate NEFs with quite different decay times and that it is similar to the multi-tau 229 230 approach utilised in dynamic light scattering [50].

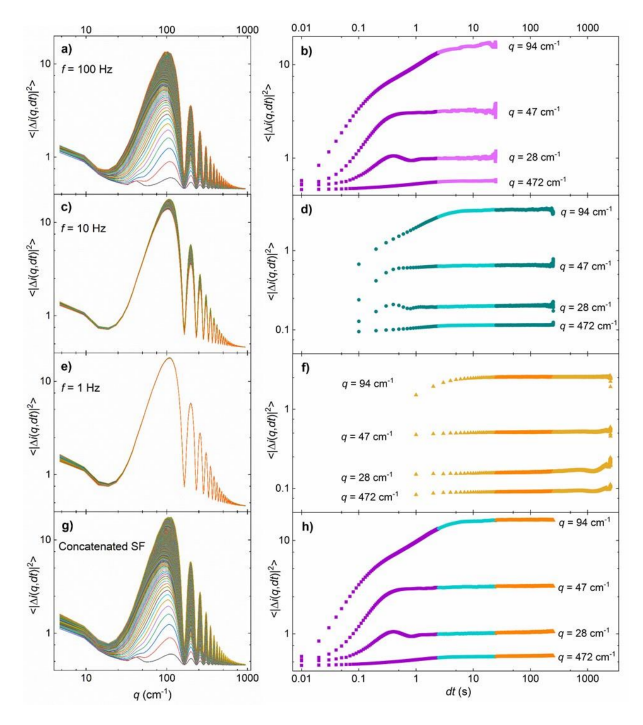


Figure 1: Structure function of the thermodiffusion experiment carried out at temperature
difference of 20 K for a series of images of 2048 × 2048 pixels at 100, 10 and 1 Hz, as a
function of the wave number (a, c, e, respectively) and as a function of correlation time (b,
d, f, respectively). Concatenated structure function as a function of the wave number (g)
and as a function of time (h). Purple squares, cyan circles and orange triangles correspond
to the data acquired at 100, 10 and 1 Hz, respectively.

238

239 2.5 Analysis of experimental signals

After the DDA and the concatenation process one gets the c-SF; it represents the raw experimental signals, obtained only from the acquired image series. For the analysis of these concatenated $\langle |\Delta i(q, dt)|^2 \rangle$, the physical optics theory of shadowgraph proposes the following expression [12, 48]:

244
$$\langle |\Delta i(q,dt)|^2 \rangle = 2\{T(q)S(q)[1 - I_{SF}(q,dt)] + B(q)\},$$
 (3)

where T(q) is the shadowgraph optical transfer function [11], S(q) is the static power spectrum, B(q) is the background and $I_{SF}(q, dt)$ is the intermediate scattering function, that usually is described as a sum of exponential decays: $I_{SF}(q, dt) = \sum_{i} a_i exp(-dt/\tau_i(q))$, where a_i are the amplitudes of the different modes (with $\sum_{i} a_i = 1$) and $\tau_i(q)$ the wave-number dependent relaxation times.

250 A simple observation of Fig. 1h, without any calculation, shows that, depending on the wave number range, up to three different decay times can be distinguished in the c-SF. In some cases, 251 252 these decay times differ by more than one order of magnitude. In addition, and also depending on the wave number range, one can observe in Fig. 1h damped oscillations in $\langle |\Delta i(q, dt)|^2 \rangle$. 253 This oscillating (propagating) phenomenon means that at least two of the relaxation times $\tau_i(q)$ 254 255 present in the $I_{SF}(q, dt)$ of Eq. 3 (and their associated amplitudes) form a pair of complex conjugate numbers. We conclude that a complex scenario of NEFs with different decay times 256 257 is present in the case of a ternary mixture with large Le and Dr numbers, and a key question is the selection of the number of modes in the $I_{SF}(q, dt)$ of Eq. 3 and whether they are purely 258 diffusive (monotonically decaying) or propagating (oscillating). A rational choice needs to be 259 guided by the available theoretical models for NEFs in ternary mixtures, which we review next. 260

261

262 **3.** Theoretical framework

263 3.1 Thermodiffusion in ternary mixtures

In a ternary mixture, the mass concentrations of the three components are related by: $\omega_1 + \omega_2 + \omega_3 = 1$, which means that only two concentrations, e.g. ω_1 and ω_2 , are independent. The corresponding diffusion fluxes J_1 and J_2 in the centre-of-mass frame of reference have thermodiffusion and Fickean components, as expressed by [51]:

268
$$\begin{pmatrix} J_1 \\ J_2 \end{pmatrix} = -\rho \begin{bmatrix} \begin{pmatrix} D_{11} & D_{12} \\ D_{21} & D_{22} \end{pmatrix} \begin{pmatrix} \nabla \omega_1 \\ \nabla \omega_2 \end{pmatrix} + \begin{pmatrix} \omega_1 (1 - \omega_1) & -\omega_1 \omega_2 \\ -\omega_1 \omega_2 & \omega_2 (1 - \omega_2) \end{pmatrix} \begin{pmatrix} D_{T1} \\ D_{T2} \end{pmatrix} \nabla T \end{bmatrix},$$
(4)

where ρ is the density of the mixture, D_{ij} are the components of the centre-of-mass diffusion matrix, $\nabla \omega_i$ are the concentration gradients, D_{T_i} are the invariant thermodiffusion coefficients and ∇T is the applied temperature gradient.

At the stationary state, the diffusion fluxes become zero and, following Ortiz de Zárate [51], the Soret coefficients S_{T_i} can be expressed by:

274
$$\begin{pmatrix} \omega_1(1-\omega_1) & -\omega_1\omega_2\\ -\omega_1\omega_2 & \omega_2(1-\omega_2) \end{pmatrix} \begin{pmatrix} S_{T1}\\ S_{T2} \end{pmatrix} \nabla T = - \begin{pmatrix} \nabla \omega_1\\ \nabla \omega_2 \end{pmatrix},$$
(5)

The introduction of the matrix concentration pre-factors in Eqs. 4 and 5 forces the thermodiffusion coefficients D_{T_i} and the Soret coefficients S_{T_i} to display the same frameinvariance properties of the single D_T or S_T of a binary mixture, that is, they are numerically the same irrespective of whether mass fractions or mole fractions are used to express the composition [51]. Here, for simplicity, we display the equations only in the mass fraction framework.

Fluctuating Hydrodynamics (FHD) has been recently extended to ternary mixtures first considering equilibrium fluctuations [52] and then non-equilibrium fluctuations [39, 40]. The first NE theory [39] neglected the effects of gravity and confinement. In that case, there is no mixing between the decay rates of the four hydrodynamic modes (fluctuations of velocity, temperature and two concentrations) which continue to decay with their well-known equilibrium values given by kinematic viscosity, thermal diffusivity and the two eigenvalues 287 \widehat{D}_i of the diffusion matrix **D**. These \widehat{D}_i , are always real and positive, and can be related to the 288 four components D_{ij} of Eq. 4 by [39, 40]:

289
$$\widehat{D}_{1,2} = \frac{1}{2} \Big[D_{11} + D_{22} \mp \sqrt{(D_{11} - D_{22})^2 - 4D_{12}D_{21}} \Big].$$
(6)

In accordance with the aforementioned common notation (i.e. 1 for PS, 2 for toluene and 3 for n-hexane), $\hat{D}_1 < \hat{D}_2$ since \hat{D}_1 represents the slowest diffusion mode essentially related to the diffusion of the polystyrene in the binary solvent.

Since refractive index does not depend on fluid velocity, only three of the four hydrodynamic modes are directly observable by optical techniques. Hence, within the approximations of Ref. [39] (no gravity and no confinement) we conclude that for a thermodiffusion experiment in a ternary mixture, a suitable expression for the $I_{SF}(q, dt)$ can be provided by the sum of three exponential decays given by two concentration modes, plus one mode for temperature fluctuations:

299
$$I_{SF}(q, dt) = a_1 \exp[-dt/\tau_1(q)] + a_2 \exp[-dt/\tau_2(q)] + (1 - a_1 - a_2) \exp[-dt/\tau_3(q)],$$

300 (7)

301 where a_i are the normalised amplitudes of the different modes and $\tau_i(q)$ the corresponding 302 relaxation times.

303

304 3.2 Concentration NEFs in ternary mixtures including gravity effects

For NEFs, first the theory was developed in the absence of gravity force [39]; while gravity effects were incorporated by Martínez Pancorbo et al. [40]. Both of these papers adopted the so-called large Lewis number approximation that uncouples temperature fluctuations from concentration fluctuations. Next, we briefly summarize the main results of Ref. [40] and recall the expressions useful for the experimental data analysis. The presence of gravity causes a mix between the two concentrations NEFs. The corresponding decay times can be described as $\tau_i(q) = 1/[\gamma_i(q)q^2]$, where the diffusivities $\gamma_i(q)$ in dimensionless form, are given by the following equation [40]:

313
$$\gamma_{i}(\tilde{q}) = \frac{1}{2} \left(\widehat{D}_{1} + \widehat{D}_{2} - \frac{Ra_{i}\widehat{D}_{i}}{\tilde{q}^{4}} \right) \times \left[1 \mp \sqrt{1 - \frac{\frac{4\widehat{D}_{1}\widehat{D}_{2}}{\left(\widehat{D}_{1} + \widehat{D}_{2}\right)^{2}} \left[1 - \frac{Ra_{s,ter}}{\tilde{q}^{4}} \right]}{\left[1 - \frac{Ra_{i}\widehat{D}_{i}}{\tilde{q}^{4}(\widehat{D}_{1} + \widehat{D}_{2})} \right]^{2}} \right], \tag{8}$$

where $\tilde{q} = qL$ is the dimensionless wave number, and Ra_i are the two solutal Rayleigh numbers defined for ternary mixtures by:

316
$$Ra_i = \frac{gL^4}{\nu \hat{D}_i} \beta'_i \nabla \omega'_i, \qquad (9)$$

317
$$Ra_i\widehat{D}_i = Ra_1\widehat{D}_1 + Ra_2\widehat{D}_2, \tag{10}$$

318
$$Ra_{s,ter} = Ra_1 + Ra_2,$$
 (11)

In Eq. 9, the primes indicate the concentrations (mass fractions) in which the diffusion matrixis diagonal, for a generic diffusion matrix [53]:

321
$$\binom{\omega'_1}{\omega'_2} = \begin{pmatrix} 1 & \frac{D_{22} - \hat{D}_2}{D_{21}} \\ \frac{D_{11} - \hat{D}_1}{D_{12}} & 1 \end{pmatrix} \binom{\omega_1}{\omega_2}.$$
 (12)

The use of "diagonal" concentrations simplifies the theory but, as we shall see, somewhat complicates the interpretation of experimental results. In general, they differ from the measurable mass fractions.

In summary, in this model [40] the two concentration modes couple, while temperature fluctuations remain uncoupled, keeping their equilibrium decay rate unaltered. Since for typical values of Ra_i the two decay times of Eq. 8 are real numbers, this theory [40] suggests to still use Eq. 7 for the $I_{SF}(q, dt)$. We note that the notation of Eqs. 9-11 implies that, for a ternary mixture, two Lewis numbers (Le_1 and Le_2) can be defined as $Le_i = a_T/\hat{D}_i$.

330

331 *3.3 Coupling of concentration and temperature NEFs in binary mixtures*

In the simplest case of binary mixtures, most of the available literature on concentration and 332 333 temperature NEFs makes use of the simplification that the two modes are always decoupled. This is essentially true whenever the large Lewis number approximation is valid, that is the case 334 of ordinary liquid mixtures whose mass diffusion coefficient is much smaller than the thermal 335 diffusivity. However, the decoupling assumption fails to be valid for very large fluctuations, 336 i.e. for small wave numbers, as recently demonstrated both experimentally and theoretically 337 338 [43]. In particular, only the full coupling of velocity, temperature and concentration NEFs in the bulk fluid can explain the appearance of propagating modes giving rise to a $I_{SF}(q, dt)$ that 339 includes oscillations at very small wave numbers. This mechanism also induces a slowing-down 340 341 of the non-propagating concentration NEFs similar to that caused by the confinement effect 342 [41, 42].

This theory of full coupling between NE velocity, temperature and concentrations fluctuations has been published only for binary mixtures. We briefly recall here the main steps of the theoretical framework, while for details we address the reader to the literature [43].

In order to take into account a more complete description of the behaviour of NEFs within a thermodiffusion experiment, one should write the evolution equations of the fluctuations of velocity v_z , temperature and concentration of the single independent component:

349
$$\frac{\partial}{\partial t} \begin{bmatrix} \delta v_z(q,t) \\ \delta T(q,t) \\ \delta \omega(q,t) \end{bmatrix} = -\mathbf{G}(q) \begin{bmatrix} \delta v_z(q,t) \\ \delta T(q,t) \\ \delta \omega(q,t) \end{bmatrix},$$
(13)

350 where the inverse linear response matrix in its dimensionless form can be written as:

351
$$\mathbf{G}(\tilde{q}) = \begin{bmatrix} PrLe\tilde{q}^2 & -1 & 1\\ -Ra_s PrLe/\psi & Le\tilde{q}^2 & 0\\ Ra_s PrLe & \psi\tilde{q}^2 & \tilde{q}^2 \end{bmatrix},$$
(14)

In the previous equation the well-known dimensionless numbers appear, namely the Lewis number $Le = a_T/D$, where a_T is the thermal diffusivity; the Prandtl number $Pr = \nu/a_T$; the separation ratio $\psi = c_0(1 - c_0)S_T\beta/\alpha$; and the solutal Rayleigh number $Ra_S = [\beta g \nabla \omega/(\nu D)]L^4$.

The solution of Eqs. 13 and 14 requires the computation of the eigenvalues and eigenvectors of the matrix $G(\tilde{q})$:

358
$$\det[\mathbf{G}(\tilde{q}) - \lambda \mathbf{1}] = \left[\Gamma_{\nu_z}(\tilde{q}) - \lambda\right] [\Gamma_T(\tilde{q}) - \lambda] [\Gamma_\omega(\tilde{q}) - \lambda], \tag{15}$$

with NE fluctuations dimensionless decay times given by $\tau_i(\tilde{q}) = 1/[\Gamma_i(\tilde{q})]$. Although Eq. 15 can be solved analytically, the resulting expressions of the roots as a function of q are so complicated that, in practice, it is more useful to have a numerical solution for fixed values of the parameters Pr, Le, Ra_s and ψ , in the whole range of q.

363 Depending on the parameter values and on the wave number q, Eq. 15 may have a pair of 364 complex conjugate solutions, which typically happens at small q values. The resulting 365 $I_{SF}(q, dt)$ function in those cases shows a sinusoidal term:

366
$$I_{SF}(q, dt) = a_1 \exp[-dt/\tau_1(q)] + \left\{\frac{1-a_1}{\cos[\phi(q)]}\right\} \cos[\Omega(q)dt + \phi(q)] \exp[-dt/\tau_2(q)]$$

367 , (16)

368 where $\Omega(q)$ is the oscillation frequency and $\phi(q)$ a phase term.

369

370 *3.4 Coupling of concentration and temperature NEFs in ternary mixtures*

The next step in the development of FHD is to include all the couplings among the fluctuating modes for a ternary mixture. Of course, in that case one needs to consider the evolution of fluctuations in velocity δv_z , temperature and concentration of the two independent components:

374
$$\frac{\partial}{\partial t} \begin{bmatrix} \delta v_z(q,t) \\ \delta T(q,t) \\ \delta \omega_1(q,t) \\ \delta \omega_2(q,t) \end{bmatrix} = -\mathbf{G}_t(q) \begin{bmatrix} \delta v_z(q,t) \\ \delta T(q,t) \\ \delta \omega_1(q,t) \\ \delta \omega_2(q,t) \end{bmatrix},$$
(17)

where the inverse linear response matrix for "diagonal" concentrations can be written indimensionless form as:

377
$$\mathbf{G}_{t}(\tilde{q}) = \begin{bmatrix} PrLe_{t}\tilde{q}^{2} & -1 & 1 & 1\\ -PrLe_{t}^{2}Ra & Le_{t}\tilde{q}^{2} & 0 & 0\\ PrLe_{t}^{2}Ra\psi'_{1} & \psi'_{1}\tilde{q}^{2} & \tilde{q}^{2} & 0\\ PrLe_{t}^{2}Ra\psi'_{2} & Dr\psi'_{2}\tilde{q}^{2} & 0 & Dr\tilde{q}^{2} \end{bmatrix}.$$
(18)

To make dimensionless Eqs. 17 and 18, L is used as unit of length and L^2/\hat{D}_1 as unit of time; 378 consequently, $\tilde{q} = qL$ for wave numbers, and L/\hat{D}_1 is the unit of velocity. Hence, everything is 379 made dimensionless with respect to the slower diffusive mode. Other dimensionless parameters 380 in Eq. 18 are: one (unique) ternary Lewis number $Le_t = a_T / \hat{D}_1$, the Diffusion eigenvalue Ratio 381 $Dr = \hat{D}_2/\hat{D}_1$, the thermal Rayleigh number $Ra = -\alpha g L^4 \nabla T/(\nu a_T)$ and the "diagonal" 382 separation ratios $\psi'_i = D_{T,i}^{\omega'} \beta'_i / (\widehat{D}_i \alpha)$, with $D_{T,i}^{\omega'}$ being the (diagonal) thermodiffusion 383 coefficients as defined in Ref. [18]. We point out that this approach is different from the one 384 used in our previous publication [40] and reported in section 3.2, where two Lewis numbers 385 386 were introduced.

As in the case of binary mixtures, the solution of Eq. 17 requires to numerically evaluate the dimensionless decay rates $\Gamma_i(\tilde{q})$ as the four eigenvalues of the matrix $\mathbf{G}_t(\tilde{q})$, namely:

389
$$\det[\mathbf{G}_{t}(\tilde{q}) - \lambda \mathbf{1}] = \left[\Gamma_{\nu_{z}}(\tilde{q}) - \lambda\right] \left[\Gamma_{T}(\tilde{q}) - \lambda\right] \left[\Gamma_{\omega_{1}}(\tilde{q}) - \lambda\right] \left[\Gamma_{\omega_{2}}(\tilde{q}) - \lambda\right].$$
(19)

A more detailed explanation of the full-coupling theory for ternary mixtures is outside the scope of the present publication and will be the subject of a future paper. We simply mention that, depending on q, complex solutions of Eq. 19 may exist. As was the case with binaries, for parameter values typical of ternary liquid mixtures, these propagating modes do appear as a mixing between the temperature and the viscous (velocity) modes. The consequence is an oscillatory $I_{SF}(q, dt)$ such as the one expressed in Eq. 16, but with an additional exponential term to represent the extra concentration mode.

397

398 4. Results

399

400 *4.1 Intermediate scattering function analysis*

401 As mentioned in section 2.5, for most wave numbers the $I_{SF}(q, dt)$ of Eq. 3 is usually defined as a sum of exponential decays. However, a preliminary analysis of the experimental SF is 402 403 required to check the suitability of this model. As it can be seen in the c-SF shown in Fig. 1h for q = 94/cm, three exponential decays can be detected, corresponding (from left to right) to 404 the relaxation of the temperature and concentration (faster and slower mass modes) NEFs. 405 Then, the $I_{SF}(q, dt)$ can be modelled for most wave numbers by a sum of three exponential 406 decays, as defined in section 3.1 by Eq. 7 for a system including one thermal mode plus two 407 408 solutal, well separated, modes. In the studied system, the corresponding three exponential decays represent the following three modes: long decay times for the two concentration modes, 409 slower for mass diffusion of the denser component (PS in the binary solvent), and faster for 410 mass diffusion of the two components of the solvent mixture (toluene and n-hexane); and the 411 shortest decay time corresponds to the thermal diffusivity of the overall mixture. 412

Nevertheless, for small wave numbers such as q = 28/cm, oscillations are detected in the c-413 414 SF (see Fig. 1h), indicating the presence of propagating modes [43]. Fig. 2 shows the oscillations in the experimental c-SF for several wave numbers in the range 19/cm < q <415 47/cm. These need to be taken into account by using a proper $I_{SF}(q, dt)$ function, such as the 416 one described in section 3.3 by Eq. 16. The latter equation is defined for a binary mixture, and 417 418 includes one solutal mode, i.e. the exponential decay, and a coupled thermo-viscous mode, i.e. the damped oscillation. For analysing ternary mixtures, one should supplement Eq. 16 with an 419 additional exponential term, representing the second solutal mode. However, after carrying out 420 a careful comparison of the corresponding fitting output, we have found that the addition of this 421 term gives more scattered results due to the increase in the number of fitting parameters. For 422 that reason, Eq. 16 has been used to fit the raw data in the wave number range where oscillations 423

are detected in the c-SF. The slower concentration mode is, thus, unobservable in this wavenumber range.

426

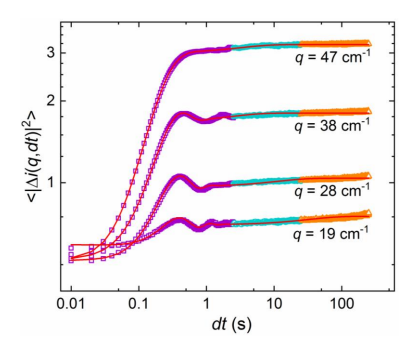


Figure 2: Concatenated structure function of the thermodiffusion experiment carried out at
temperature difference of 20 K for a series of images of 2048 × 2048 as a function of time
for different small wave numbers. Purple squares, cyan circles and orange triangles
correspond to the data acquired at 100, 10 and 1 Hz, respectively, and the red curves to the
fitting functions through Eqs. 3 and 16.

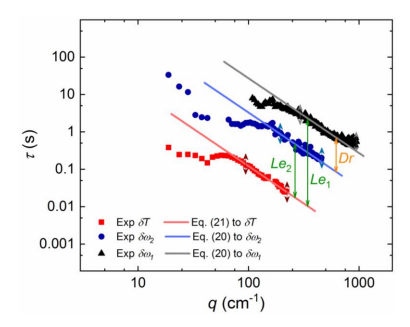
433

Generally, the quality of these fittings is good, as demonstrated in Fig. 2, with an average R^2 of about 0.99, which translates into a 3% mean relative uncertainty for the decay times determined by these fittings. However, both R^2 and the relative uncertainty of the decay times depend on the wave number. The best accuracy is obtained for intermediate wave numbers where the signal-to-noise ratio of the c-SF is the largest (see Fig. 1g). Decay times with a relative uncertainty larger than 10% have been disregarded for the further analysis.

440

441 *4.2 Decay times of the NEFs*

442 Figure 3 shows the decay times of the different fluctuating modes as resulting from the modelling of the experimental c-SF by using Eqs. 3 and Eq. 7 or Eq. 16 depending on the 443 analysed q. Three decay times are easily distinguished (from top to bottom): the slowest mode 444 (black symbols) describes the concentration mode of the polymer into the binary solvent; the 445 intermediate one (blue symbols) the concentration mode of the binary solvent mixture; and the 446 447 fastest one (red symbols) the temperature mode. These decay times cover a span of about three orders of magnitude, which is quite impressive for an imaging technique. This justifies the 448 necessity of performing measurements at different acquisition frequencies and of carrying out 449 450 the concatenation procedure adopted here. Besides decay times, when the Eq. 16 is used, the fitting procedure also gives the oscillation frequency $\Omega(q)$ of the propagating modes appearing 451 in Fig. 2 for wave numbers q < 47/cm. 452



453

454 Figure 3: Decay times of NEFs on the ternary mixture of PS-toluene-n-hexane in a
455 thermodiffusion experiment performed at temperature difference of 20 K, as a function of
456 the wave number q. Lines represent fitting functions through Eqs. 20-21 over the diffusive
457 wave number range, as indicated in the text.

458

The experimental decay times shown in Fig. 3 are to be discussed first in the diffusive regime only (for large enough wave numbers), where the three decay times are well-separated. As elucidated at the end of section 3.1, in this region of large wave numbers the effect of gravity is negligible for both concentration modes, and the decay times of the two concentration modes are given by the eigenvalues of the diffusion matrix:

$$\tau_i(q) = 1/(\widehat{D}_i q^2). \tag{20}$$

By fitting the experimental decay times in the diffusive regime (q > 290/cm) for the 465 concentration NEFs, c-NEFs, of component 1 and q > 190/cm for the c-NEFs of component 466 2) to Eq. 20, the eigenvalues of the diffusion matrix can be determined. The diffusion coefficient 467 \widehat{D}_1 obtained from the slowest concentration mode represents the mass diffusion eigenvalue of 468 the polymer in the binary solvent mixture. \widehat{D}_2 calculated from the intermediate decay times 469 470 (i.e. faster concentration mode) represents the mass diffusion eigenvalue of toluene in n-hexane (i.e. interdiffusion of the solvents in the solvent mixture). The resulting two eigenvalues \widehat{D}_1 and 471 \widehat{D}_2 are reported in Table 2. These values in Table 2 are calculated by averaging the fitting results 472 of the experiments performed at three temperature differences, 20, 15 and 10 K, and the 473 uncertainties are the corresponding standard deviations. These values are comparable with those 474 475 obtained in our previous study [14] for a similar polymeric mixture prepared with the same components of this study but a slightly different polymer molecular weight, and with those 476 477 reported in [43, 45] for the diluted binary mixture of PS-toluene and in [44, 46] for the binary 478 solvent mixture of toluene-n-hexane. Furthermore, the value of the Dr number can be determined as the ratio between the two eigenvalues, resulting in Dr = 8.4 (see Table 2). This 479 480 value totally justifies the selection of the polymeric mixture as a suitable sample for experimentally studying well-separated concentration modes. 481

482 In a similar way, in order to get information about the thermal diffusivity a_T of the ternary 483 mixture, one can try to fit the data points of the thermal mode with the theoretical predictions for thermal NEFs only. In the diffusive regime, the decay time of the thermal mode can beexpressed as [54]:

486
$$\tau_{th}(q) = 1/(a_T q^2),$$
 (21)

and the fitting of temperature NEFs for q > 90/cm in Fig. 3 provides the thermal diffusivity value reported in Table 2. This result is again comparable with the one obtained in previous studies for a similar polymeric mixture [14] as well as with that reported in [46] for the equimolar toluene-n-hexane binary solvent mixture.

- 491 From the two diffusion eigenvalues and the thermal diffusivity, the Le_i numbers can be 492 determined, which are also listed in Table 2.
- 493

494 *Table 2: Diffusion coefficient eigenvalues, Lei and Dr numbers of the polymeric ternary*

495

Parameter	Value
\widehat{D}_1	$(3.8 \pm 0.1) \times 10^{-6} \mathrm{cm^2/s}$
\widehat{D}_2	$(3.1 \pm 0.1) \times 10^{-5} \mathrm{cm}^2/\mathrm{s}$
a_T	$(8.5 \pm 0.2) \times 10^{-4} \mathrm{cm^2/s}$
Le ₁	227 ± 12
Le ₂	27 ± 2
Dr	8.4 ± 0.6

mixture

496

497 **5. Discussion**

498 Comparison between experimental and theoretical decay times

As described in section 3.2, Martínez-Pancorbo et al. [40] presents the analysis of the fluctuations of two independent concentrations together with the effect of buoyancy, but considering neither the effect of confinement nor the presence of temperature and velocity

fluctuations. Consequently, the coupling between modes which is known to be important at 502 small wave numbers, as demonstrated for binary mixtures [43], is not taken into account. In 503 order to be insensitive to confinement effects the experiments reported here have been 504 performed in a thick cell with vertical dimension of L = 5 mm. In these conditions, the effect 505 506 of confinement should not be visible in the investigated wave number range. In fact, confinement effects appear only at dimensionless wave numbers $\tilde{q} = qL < 5$, corresponding 507 here to dimensional wave numbers q < 10/cm, at the very bottom of the investigated range of 508 509 *q* [41, 42].

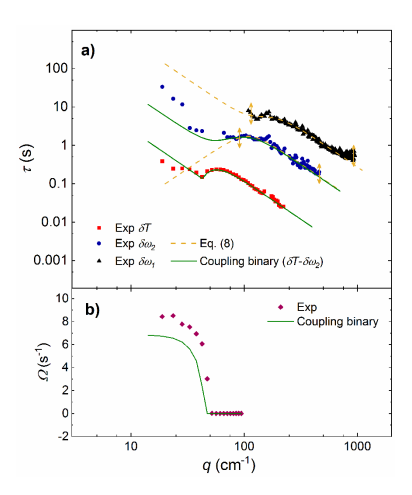
Regarding coupling effects, it is mandatory to consider them whenever oscillations are visible in the experimental c-SF at small q (see Fig. 2). The complete FHD theory, including the presence of velocity and temperature fluctuations and the various couplings among all modes, has been previously published by us [43] for the case of binary mixtures, as summarised in section 3.3. The results of [43] can be very useful for discussing the impact of the velocitytemperature coupling on the measured decay times.

516 In the following, we compare the experimental values of decay times at different wave numbers (obtained by fitting the c-SF through Eqs. 3 and 7 or 16), with the two published models [40, 517 518 43]. This allows to understand the impact of the different effects on the measured c-SF, and to evaluate if a more complete theoretical framework is needed or if on the contrary, the available 519 theories can be simply applied in different wave number ranges in order to extract the transport 520 properties of the investigated ternary mixture. As described in the previous section, at least the 521 522 diffusion coefficients can be safely determined by fitting decay times at large wave numbers, 523 where NEFs are dominated only by the diffusion phenomenon. The measurement of the Soret 524 coefficient is more complicated as it requires the analysis of the effect of gravity [6], or other 525 suitable analysis [55]. The simultaneous and independent characterization of the two Soret 526 coefficients of a ternary mixture requires a complete knowledge of the diffusion matrix

including the cross-diffusivities. This has not been achieved yet by Shadowgraphy and is the ultimate goal of our current research. Although we estimate that this can be done with onewavelength Shadowgraphy, a more precise information can be obtained with a two-wavelength version of the technique. The two-wavelength Shadowgraphy is currently being developed in our lab, also in relation to the future space experiments of the NEUF-DIX project [28].

Figure 4 shows the experimental decay times and the oscillation frequency obtained from the
c-SF (data points are the same as in Fig. 3), together with theoretical predictions provided either
by Eq. 8 (dashed yellow curves) or by numerically solving Eqs.14-15 (solid green curves).

535



536

Figure 4: a) Decay times and b) oscillation frequency of NEFs on the ternary mixture of PStoluene-n-hexane in a thermodiffusion experiment performed at temperature difference of
20 K, as a function of the wave number q. Experimental data are shown by points, while lines

represent the numerical calculations of Eqs. 8 (dashed lines) and the theory developed for
binary mixtures including the effect of propagating modes (solid lines).

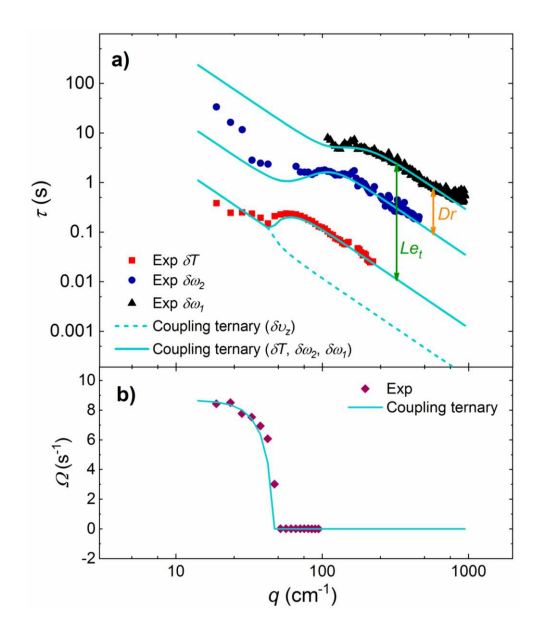
542

The experimental data are fitted to Eq. 8 by fixing the diffusion matrix eigenvalues (\hat{D}_1 and \hat{D}_2) 543 to the corresponding values presented in Table 2. The parameters obtained from this fitting to 544 Eq. 8 are: $Ra_i \hat{D}_i = (260 \pm 40) \ cm^2/s$, and $Ra_{s,ter} = (4.6 \pm 1.0) \times 10^7$. As visible in Fig. 4a, 545 the experimental results suggest that the available theory for the ternary mixture reported by 546 Martínez Pancorbo et al. [40] cannot describe the behaviour of c-NEFs in the present 547 experiment for the whole range of wave numbers. The theory provides suitable predictions for 548 intermediate and especially for large wave numbers, but fails in the limit of small q (in 549 particular for the fastest concentration mode) where the effect of the coupling with temperature 550 fluctuations, and the consequent appearance of propagating modes in the experimental structure 551 function become significant (see section 3.3) [43]. Hence, investigation of the coupling with 552 temperature fluctuations is required for a full understanding of the present experimental results. 553 554 We proceed into two steps: first we compare with the published theory [43] for the binary mixture case (one concentration mode) as reviewed in section 3.3; second we compare with the 555 theory for ternary mixtures (two concentration modes) as briefly presented here in section 3.4. 556 557 This allows appreciating the features that are genuinely related to the ternary mixture case.

As already mentioned, Fig. 4 shows, superposed on the experimental data, the theoretical predictions (green solid lines) of the theory [43] with one concentration mode. The theoretical curves have been obtained numerically (solving Eqs. 14 and 15) by keeping constant the Lewis number Le = 27 to the same value of Le_2 shown in Table 2, and varying the values of Pr and ψ to best fit the data. The solutal Rayleigh number $Ra_{S,2} = -(q_{S,2}^*L)^4$ is also kept constant at -5.8×10^6 , after evaluating the value of the solutal rolloff wave number $q_{S,2}^*$. This value represents the wave number where the fluctuation $\delta\omega_2$ behaviour changes from diffusive to

gravitational [6]. The resulting values for the free fitting parameters are the following: Pr =565 9.0 and $\psi = 0.9$. As it can be observed in Fig. 4a, the decay times provided by the numerical 566 solution agree very well with the experimental decay times of the temperature NEFs in the 567 568 whole range of wave numbers. However, for the diffusion mass mode, at the smallest wave number range, the agreement is not as good as for the temperature, and a slowing-down of the 569 570 experimental decay times is visible in comparison to the corresponding numerical solution (green solid line). The theoretical results of the oscillation frequency are likewise similar to the 571 572 experimental data (see Fig. 4b). This last comparison between the experimental and theoretical 573 decay times shows the importance of including temperature fluctuations in the theoretical analysis. 574

Next, we finally take into account the more complete theory of NEFs, which includes the coupling of thermal and viscous fluctuations to the two independent concentration modes present in a ternary mixture, as briefly described in section 3.4. The comparison between experimental and theoretical decay times is shown in Fig. 5. The effect of considering the two concentration modes and the associated couplings is evident by comparing Fig. 4 and 5.



- 580
- 581

Figure 5: a) Decay times and b) oscillation frequency of NEFs on the ternary mixture of PStoluene-n-hexane in a thermodiffusion experiment performed at temperature difference of 20 K, as a function of the wave number q. Experimental data are shown by points, while lines represent the theory developed for ternary mixtures including the effect of propagating modes.

The theoretical curves shown in Fig. 5 have been obtained numerically (solving Eqs. 18 and 19) by keeping constant $Le_t = 227$ and Dr = 8.4 (the Le_1 and Dr values shown in Table 2), and varying the values of Pr, ψ'_1 and ψ'_2 to best fit the data. Additionally, the term $PrLe_t^2Ra =$

 $\alpha g L^4 \nabla T / \widehat{D}_1^2$ appearing in the matrix $G_t(\widetilde{q})$ of Eq. 18 is also kept fix at 1.6×10^{11} , considering 590 for this number the α value of Table 1, the \widehat{D}_1 value of Table 2, and the applied temperature 591 gradient. The resulting values for the free fitting parameters are the following: $Pr = 10, \psi'_1 =$ 592 0.7 and $\psi'_2 = 0.4$. The theoretical results show that for large wave numbers $(q \ge 47 cm^{-1})$, four 593 real solutions are obtained, demonstrating that all the modes are decoupled in this wave number 594 range. The velocity mode cannot be observed experimentally, as it does not give rise to 595 refractive index fluctuations. For small wave numbers ($q < 47 cm^{-1}$), two real solutions and a 596 597 pair of complex conjugate ones are obtained numerically, indicating that velocity and temperature fluctuations are mixed at small q. This mixing induces the appearance of 598 propagating modes, observed in the experimental c-SF of Fig. 2 as oscillations at small q. 599

Applying the full theory of section 3.4, we found a reasonable agreement between the experimental and the theoretical data for the three modes, especially for large and intermediate wave numbers. However, we would like to point out that the numerical values of Pr, ψ'_1 and ψ'_2 must be taken into account with caution as the fitting procedure is not robust. The focus here is mainly on the qualitative understanding of the behaviour of the different modes in a ternary mixture when the coupling among them is included.

One relevant point is the slowing-down of the two concentration modes and their consequent diffusive-like behaviours at very small wave numbers (see Fig. 5a). In addition, it is worth noting the good agreement obtained for the frequency of the propagating mode, as can be seen in Fig. 5b. It clearly improves the results of the binary model shown in Fig. 4b.

Of course, the effect of confinement should be also included in the theory if one wants to
describe layers of ternary mixtures of any vertical thickness. Unfortunately, this additional
development requires more effort and no analytical solution can be obtained.

613

614 6. Conclusions

The concentration and temperature NEFs in a thermodiffusion experiment of a ternary mixture 615 616 composed of polystyrene, toluene and n-hexane are characterized by dynamic shadowgraphy. The decay times of the NEFs are determined and three modes are clearly distinguished. The 617 618 fastest decay times are associated to the temperature NEFs, the intermediate ones to the concentration NEFs of the molecular solvent and the slowest ones to the concentration NEFs 619 of the polymer. To encompass the very different decay times (i.e. large range of correlation 620 621 times), images are acquired at various frequencies, and the corresponding SFs are merged into the so-called c-SF by following the described concatenation procedure. 622

In order to exploit the obtained decay times, first, they are fitted with a simple diffusive model 623 at large wave numbers to obtain the diffusion eigenvalues \widehat{D}_1 and \widehat{D}_2 and the thermal diffusivity 624 a_T . Secondly, the theory developed for the two concentration modes of a ternary mixture in the 625 presence of gravity is compared with the experimental results for the first time. The 626 experimental decay times of the concentration modes at intermediate and large wave numbers 627 628 appear to be in good agreement with this theoretical model. However, for smaller wave numbers, a slowing-down of the experimental decay times of NEFs is detected with respect to 629 the predictions of that theory. Due to the large thickness of the thermodiffusion cell, 630 631 confinement effects cannot be considered as the origin of such discrepancy. In addition, evidence of propagating modes on the structure function suggests that the aforementioned 632 slowing-down could be originated by the coupling of the temperature and viscous modes to the 633 concentration ones. Comparison with the decay times from the theory developed to describe 634 such coupling in a binary mixture shows a better agreement for the thermal and the faster 635 636 concentration modes at intermediate wave numbers. This positive outcome lead us to further develop the theory of NEFs in the case of ternary mixtures, by including the coupling of 637 638 velocity, temperature and the two concentration modes under strong gravitational stabilization and for large size fluctuations (small wave numbers). A better qualitative agreement is foundbetween the new theoretical development and the experimental data points.

641

642 Acknowledgements

The authors of the University of Pau kindly thank the financial support from the Centre National
d'Etudes Spatiales (CNES) and from the funding partners of the Industrial Chair CO2ES: E2SUPPA, TOTAL, CNES and BRGM. L. García-Fernández gratefully acknowledges the CNES
for the post-doctoral research grant. The research at the Complutense University was supported
by grant ESP2017-83544-C3-2-P of the Spanish *Agencia Estatal de Investigación*.

648

649 Authors contribution statement

650

The experimental measurements and data analysis were performed by L. García-Fernández, H.
Bataller and F. Croccolo. P. Fruton contributed to the concatenation procedure of the Structure
Function. J.M. Ortiz de Zárate and L. García-Fernández focused on the new theoretical
development presented in section 3.4. All the authors contributed to the discussion of the results
and in writing the manuscript.

656

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