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Integrating a micro-mixing mechanism and on-line thermal processing for the large-scale ejection of polymeric liquid threads for producing ultrafine fibers

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Micro/nanofibers are structures that nowadays have a wide range of cutting-edge applications including energy generation and storage devices, smart textiles, cell growth, and tissue engineering. These fibrous materials are mostly produced from polymer solutions spun, under laminar flow conditions, into nanofibers by external forces. However, the turbulent interaction of gas-liquid interfaces offers an innovative approach for the high-throughput production of nanofibers. Here, we present Flow Blurring (FB), a solely pneumatic approach for the massive production of liquid threads of polymer solutions, which relies on a micro-mixing mechanism that triggers a turbulent motion capable of fragmenting a viscous flow. The as-ejected threads are subsequently processed thermally, on-line in a single-step, thus producing micro/nanofibers that form mats. The method operates with relatively large liquid flow rates, equivalent of a high production rate, and is thus suitable for industrial production of engineered nanomaterials. In this work, we used solutions of poly(vinyl alcohol) (PVA) to study its ejection and fragmentation dynamics through computational fluid dynamics (CFD) simulations. In addition, the physics underlying the regulation of the liquid flow rate in FB atomizers are proposed. Fibers with typical diameters in the range 400-800 nm were produced by online heating of the liquid threads. Liquid ejection experiments were performed under different operating conditions thus verifying the capability of the method for synthesizing submicrometer-sized fibers with high uniformity and production rates suitable for scaling up.

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Introduction

Micro and nanofibers are ubiquitous structures in cutting-edge applications including electrodes for energy generation and storage devices, smart textiles, cell growth, tissue engineering, coatings, and catalysts.¹ Many of those fibrous materials are produced from polymer solutions,²-⁴ which may contain additives such as nanoparticles, nanotubes, or soluble precursors.³,⁵-10 The polymer matrix either serves as a template for micro/nanofiber fabrication or is the material of interest itself. The applications of such polymer fibers are determined not only by the structure of the polymer's repeating building block or their physicochemical surface properties but also by their dimensions, such as their length and diameter.

Polymer micro- or nanofibers are generally obtained from the drying of a liquid filament drawn from a bulk by external forces as it occurs in techniques such as meltblowing,11 electrospinning, 3,12-18 or straining flow spinning. 19 The industrial demand for polymeric fibrous materials has prompted intense research to develop efficient and scalable fabrication methods to produce tailored materials.11 This quest has often resulted in the merging of two or more techniques to take advantage of their strengths. Thus, for instance, a centrifugal electrospinning method was presented that diverges from the traditional electrospinning in that it utilizes a type of rotating, bell-shaped ejector²⁰ instead of the typical, single needle-type. Likewise, a centrifugal multispinning technique was proposed, which consisted of a cylindrical reservoir with orifices on lateral, main surface from which the solution was ejected towards a rotating mesh.21 These methods are based on the ejection and subsequent collection of thin liquid threads, where the interplay of surface tension forces, viscous forces, and other external forces, such as electrical, are evident and critical to achieve rather thin and long fibrous materials. There should be therefore an energy input to overcome unfavorable forces, surface tension in this case, acting on the threads. Therein, it is consequently required the liquid to be electrically conductive to

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respond to external stimuli effected by the field from the collector electrode. Furthermore, from an engineering view-point the key features a method should fulfil to be implemented in practical applications are: simplicity and yet robustness, versatility to be used with a wide range of materials, consistency in its feeding system, and capability for large-scale production. In that sense, liquid atomization techniques offer a stable source of precursors and are widely used in materials synthesis.²²

Here, we present the coupling of a micro-mixing-based liquid atomization technique, for generating threads of polymer solutions, with an on-line thermal treatment to achieve high-volume production of ultrafine polymer fibers. The term micro-mixing refers to an intricate intermixing phenomenon occurring at small length scales, typically within the micrometer range, resulting in the mixing of both fluid phases. The approach is based on a pneumatic ejection of polymer solutions at high liquid flow rates, 23-25 of the order of tens of milliliters per hour, much higher than those used by standard techniques $(\ll 1 \text{ mL h}^{-1})^{26,27}$ followed by thermal treatment with a tubular furnace, in a single-step. Our method uses a so-called Flow Blurring (FB) atomizer^{23,25,28-30,} to fragment a liquid stream with a gas current, thus generating multiple, thin liquid threads that elongate as they move forward^{25,31,32} and are dried in-flight producing micro/nanofiber mats. Because the method functions with relatively low or medium temperatures it allows for the processing of materials which may undergo thermal degradation when exposed to higher temperatures.

A fundamental and unique feature of the method presented herein is its capability of mixing at micrometer length scales both fluid phases, liquid and gas. Such vigorous micro-mixing is triggered by a radial implosion of the gas into the liquid stream which causes the formation of bubbles within it.²³ As the bubbles recirculate in the liquid interior and move forward with the main current the fragment the liquid bulk thus originating the threads.^{23,29,33–35} Thus, this approach is suitable for the processing of viscous, polymer solutions. Although flow blurring atomization has been performed with water and other Newtonian liquids, its use for production of micro/nanofibers remains largely unexplored.

Furthermore, we used computational fluid dynamics (CFD) simulations as guidance to study the evolution of polymeric solution's ejection patterns and fragmentation, followed by fibers' production under different ambient conditions. Herein we used poly(vinyl alcohol) (PVA) as the test polymer due to its ease to dissolve in common solvents such as water or water/ethanol mixture, and because of its immense range of applications. Furthermore, we delve into the physics underlying the critical ejection process, which dictates the inception of liquid threads leading to ultrafine fibers.

2. Experimental methods

2.1 Materials and preparation of solutions

Poly(vinyl alcohol) (PVA, Mowiol 40–88, CAS: 9002-89-5) of average molecular weight of 205 000 g mol⁻¹ was purchased from Sigma-Aldrich and used as received, without further

purification. Water (Wasserlab, Type II, Analytical grade) and ethanol (PanReac, 99.5% v/v, CAS: 64-17-5) were used as solvents. The solutions were prepared by adding a previously weighted mass of PVA powder to a water/ethanol mixture (water mass fraction, $\chi_{\rm w}=0.35$) to result in a PVA concentration of 6 wt%. Subsequently, the solution was stirred using a hot plate (Corning PC-420D) until it became homogeneous and transparent. Mild heating was applied during the stirring process to facilitate polymer dissolution. To ensure uniform heating during stirring, the solution was submerged in a water bath. Once the stirring process was finished, the solution was allowed to cool to room temperature before using it in atomization experiments.

2.2 Density, surface tension, and viscosity measurements

The density of the PVA solution was obtained by weighting a known volume of solution and was found to be 885.3 kg m⁻³. The surface tension of the polymeric solution in air was measured with a KSV Instruments' contact angle meter (model CAM100) set up in a pendant drop configuration for static measurements in the range 0-180°. The instrument is equipped with a FireWire video camera module with a resolution of 640 imes480 pixels and with a light-emitting diode, monochromatic, light source. The objective lens provided with the camera is telecentric with a 55 mm focal length. The instrument's software applies a curve fitting using the Young-Laplace equation to calculate surface tension, which in this case was found to be 0.0324 N m⁻¹. The solution viscosity was characterized using a TA Instruments rheometer (Discovery HR-3) with cone-plate configuration with a gap of 52 µm. The cone angle and the plate diameter were 2.009° and 60 mm, respectively. Temperature control is achieved using a Peltier plate equipped in the instrument. The zero-shear viscosity, μ_0 , was taken as the viscosity at the lowest shear rate measured (25 Pa s). The viscosity measurements were performed at room temperature and ambient relative humidity (in the range of 50-60%).

2.3 Experimental setup for microfiber production

Fig. 1a depicts the experimental setup consisting of a tubular furnace (Carbolite Gero TF1-1200) equipped with a ceramic tube and a FB nebulizer (Ingeniatrics Tecnologías, Seville, Spain). A cylindrical, metallic mesh, formed by galvanized wires (Fig. 1b), was placed in the interior of the ceramic tube to collect the fibrous material and provide it with structural support. The tubular furnace has a maximum operating temperature of 1200 °C. Nevertheless, in our experiments the operating temperature ranged from 150 °C to 600 °C. Such range corresponds to the nominal furnace temperature (T_f) , which is higher than the real temperature (T_r) in the interior of the ceramic tube. Two temperature profiles along the lube length are presented in Fig. 1c to highlight such difference and to quantify T_r in the tube. As shown in the graph, there is a zone with a length of 20 cm in the central part of the furnace where the temperature is nearly-constant. The temperature gradient from the tube's center to its extremes is caused because of both ends are open (to allow for a free gas flow) and not thermally insulated.

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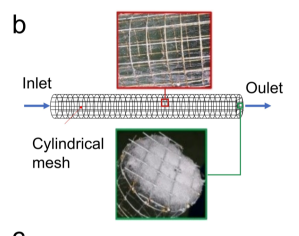
Tubular furnace

Ceramic tube

Compressed air (ΔP_l) Liquid container

Atomizer

Gas flow



meter

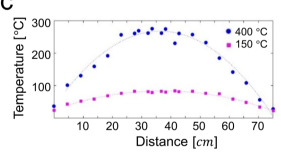


Fig. 1 (a) Experimental setup, (b) sketch of metallic mesh for fiber collection, and (c) furnace temperature profile.

The ceramic tube had inner and outer diameters of 7.4 cm and 8.4 cm respectively, and a length of 80 cm.

The Flow Blurring atomizer consists of two concentric tubes, with the inner one carrying the liquid and the outer one carrying the gas. It is characterized by its nominal orifice diameter $D=1200~\mu m$ and gap $H=250~\mu m$, thus resulting in a $\Phi=H/D=0.21$ (Fig. 2a). It has two inlets, one for the 6 wt% PVA solution and the other for the gas (air) flow. The air overpressure ($\Delta P_{\rm g}$) was measured with a digital manometer (Digitron 2026P) to deliver a specific gas flow rate ($Q_{\rm g}$) measured by a digital flow meter (Red-y compact series, Vögtlin Instruments) as indicated

in Fig. 1. The polymer solution was supplied pneumatically by feeding pressurized air into a hermetic, liquid container. The liquid flows through a polyamide tube with outer diameter 4 mm, inner diameter $D_{\rm i}=2.5$ mm, and length L=0.88 m. The liquid flow rate, $Q_{\rm l}$, was thus controlled by adjusting the overpressure in the liquid container, $\Delta P_{\rm l}$, together with the gas overpressure $\Delta P_{\rm g}$. The FB atomizer's unique geometry causes the gas flow to collapse radially inward towards the liquid stream coming from the liquid feeding tube at the atomizer discharge zone (Fig. 2a). As a result, a stagnation region is formed at the discharge of the liquid feeding tube, where the pressure exceeds that of the outer ambient environment $P_{\rm a}$ equal to a fraction of $\Delta P_{\rm g}$.

Regarding the experimental procedure, the FB atomizer was positioned at the furnace entrance. As a result, the polymer filaments, once ejected, passed through the ceramic tube where the applied heat facilitated rapid solvent evaporation. Depending on the liquid flow rate and furnace temperature, the filaments could either fully dry in-flight, resulting in fiber formation, or undergo partial drying. In the former case, solid fibers would impact both the mesh's wall, which runs parallel to the main flow direction, and the end wall, perpendicular to the flow direction. Conversely, if the filaments only partially dried, wetted fibers may impact both walls of the mesh, completing their drying process after landing. Once the atomization experiment concluded, the mesh was carefully removed from the furnace to prevent potential thermal degradation of the fibers. When detaching the fibrous material from the mesh collector, it is crucial to differentiate between material deposited on the mesh's lateral wall and that deposited on the crosssection of the outlet zone (refer to Fig. 1b). All experiments were conducted under atmospheric conditions at an average ambient temperature of 25 °C and relative humidity of 45%.

2.4 CFD simulations

2.4.1 Mesh geometry. CFD simulations have been carried out to further understand the physics of the micro-mixing phenomenon occurring in the interior of the flow blurring atomizer. The geometry used for the simulations was selected based on the specifications of the manufacturer and on an image of a 3D tomography image of a typical FB atomizer. The resulting mesh is depicted in Fig. 2b and was generated using a structured mesh approach. In these simulations, the atomizer is characterized by an inner diameter D = 0.7 mm and the geometrical parameter $\Phi = 0.14$. This is to ensure that the studies capture the essence of the phenomenon considering that the upper bound limit for the micro-mixing to take place is $\Phi = 0.25$. Otherwise, the operation regime may shift from the micro-mixing in the FB regime towards the so-called Flow Focusing regime.²³ Also, notice that the discharge zone, where the filaments develop, was modelled large enough not only to capture the ejections, but also to observe possible interactions among themselves.

A 3D structured mesh composed of 2 454 027 tetrahedrons was implemented for simulations. Thin elements were positioned in regions where the micro-mixing and the ejections

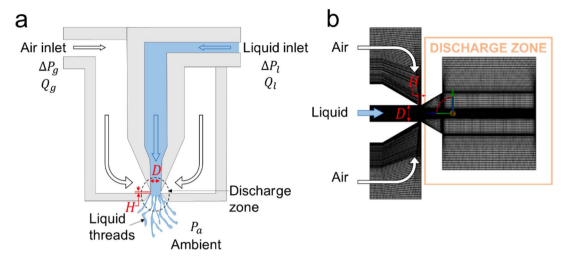


Fig. 2 (a) Sketch of a 2D cross-sectional view of FB atomizer; $D = 700 \, \mu m$ and $H = 100 \, \mu m$; (b) structured mesh for CFD simulations.

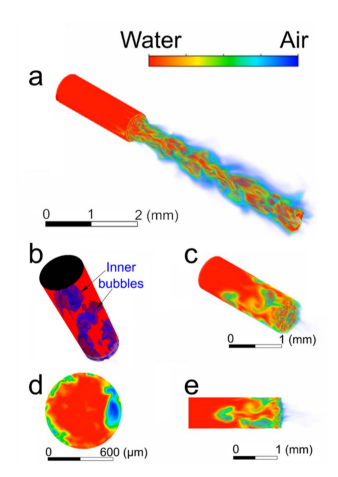


Fig. 3 Images of CFD simulations of water atomization. (a) So-called turbulent flow focusing with $\Delta P_g = 200$ kPa. Flow blurring with $\Delta P_g = 350$ kPa: (b) bubbly flow in the interior of the liquid; (c) perspective view of the FB regime, (d) cross-sectional view just upstream of the outlet of the liquid feeding tube; (e) lateral view of the FB regime. The color scale represents the volume fraction of both phases, red: α_l of water =1, and blue: α_g of air =1.

occur. Conversely, the size of elements at positions away from those regions of interest is considerably larger to decrease the computational cost without incurring in gross solution errors. The mesh quality was assessed using two the so-called orthogonal quality and skewness. Both of these parameters exceeded 0.9 and were lower than 0.5, respectively, in the regions of high interest, namely the micro-mixing zone and the discharge zone. These values indicate a highly reliable mesh in those critical areas. The simulation was conducted for a sufficient duration to allow the phenomenon to reach a steady state, in which the underlying rules governing the micro-mixing and ejection phenomena appeared to remain consistent over time, considering the inherently chaotic nature of the phenomenon. It is noteworthy to mention that we have extensive experience observing the ejection phenomena with ultra-high-speed cameras. That valuable knowledge is beneficial in understanding simulations' results, and in turn, it could be helpful to avoid excessive mesh refinements that would otherwise consume computational resources.

2.4.2 Modelling physics. The Navier–Stokes continuity and moment equations were used to study the dynamics of flow blurring atomization of polymer solutions, which are represented by the following expressions:

$$\frac{\partial \rho}{\partial t} + \nabla \cdot (\rho \mathbf{u}) = 0 \tag{1}$$

$$\rho \left[\frac{\partial \boldsymbol{u}}{\partial t} + \boldsymbol{u} \cdot \nabla \boldsymbol{u} \right] = -\nabla p + \nabla \cdot \left[\mu \left(\nabla \boldsymbol{u} + \nabla \boldsymbol{u}^T \right) \right] + \delta \left(\zeta - \zeta_f \right) \sigma (\nabla \cdot \boldsymbol{n}) \boldsymbol{n}$$

Here, u = (u, v, w) denotes the velocity field where u, v and w are the velocity components in the x, y and z directions, respectively; t denotes time; and p is the pressure field. The third term on the right-hand side of eqn (2) refers to the interface zone. This term is a delta-type function whose value is non-zero only at the interface between the gas and liquid, that is, when $\zeta = \zeta_f$, thus allowing to represent the effect of the curvature at the interface, normal to the surface of the two fluids. Additionally,

Paper

PVA Air $\Delta t = 55 \mu s$ 1

2

3

6

LIQUID

7

8

9

Fig. 4 Sequential frames of CFD simulations of PVA solution atomized in the FB regime. $\Delta P_{l} = 200$ kPa, $\Delta P_{g} = 300$ kPa. The atomizer 2D geometry is depicted in inset 4. The color bar represents α of both phases, red: α_{l} of PVA solution = 1, and blue: α_{g} of air = 1.

the gravity effect has been neglected because the Bond number $\text{Bo} = \rho_1 g D^2 / \sigma$ in our system is below unity, where ρ_1 and σ are the density of the liquid and its surface tension respectively.

The solution of the problem has been tackled using the commercial CFD software, ANSYS Fluent®. The PISO scheme is used to solve the pressure-velocity coupling. Within the spatial discretization, the Green-Gauss Node Based Model is used and the PRESTO! method has been selected for the resolution of the pressure terms. Additionally, a second-order upwind scheme has been selected for the momentum resolution and the transient problem has been formulated following a Bounded Second Order Implicit Scheme. High order term relaxation has been used to achieve a faster convergence and VOF solution stability controls was activated to avoid oscillations and

divergence phenomena during training. Furthermore, a variable time step was selected based on the Courant–Friedrichs–Lewy (CFL) number, which controls the convergence comparing the temporal and spatial feedforward. CFL number was fixed at a value of 2, giving a timestep of the order of 10^{-8} seconds. The inlet conditions were specified as *Pressure Inlet* for both the gaseous and liquid phases. The discharge zone was defined as a *Pressure Outlet* at ambient, atmospheric pressure and no suction conditions. Finally, when simulating FB atomization of polymeric solutions, a power-law viscosity model was implemented to consider the shear-rate dependant viscosity.

The multiphase system was modelled using the volume-offluid (VOF) framework, which is a surface-tracking technique applied to a fixed Eulerian mesh. It was designed to study the

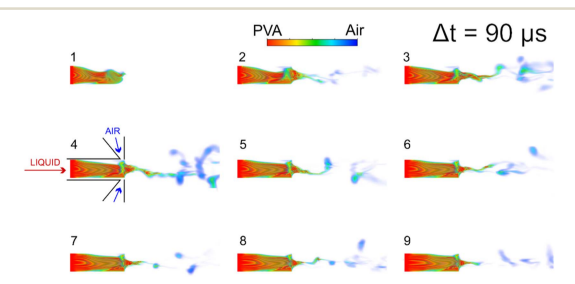


Fig. 5 Sequential frames of CFD simulations of PVA solution atomized in the FB regime. $\Delta P_{l} = 200$ kPa, $\Delta P_{g} = 330$ kPa. The atomizer 2D geometry is depicted in inset 4. The color bar represents α of both phases, red: α_{l} of PVA solution = 1, and blue: α_{g} of air = 1.

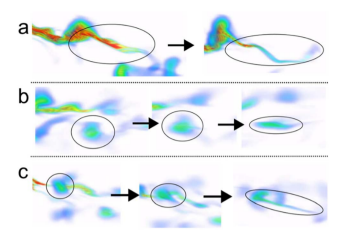


Fig. 6 Transformation mechanisms during ejection of polymer solution threads. (a) elongation, (b) droplet to filament, and (c) ejection. Red indicates $\alpha_l = 1$, and blue indicates $\alpha_q = 1$.

Table 1 Characteristic velocity and liquid vein diameter calculated with scaling laws

Case	$\Delta P_{\rm g}$ (kPa)	$\rho_{\rm l} ({\rm kg \; m^{-3}})$	$Q_{\rm l} \left({ m L} \ { m h}^{-1} \right)$	U^* (m s ⁻¹)	<i>L</i> * (μm)
1 2	300	885.3	3.041	18.408	214.225
	330	885.3	1.615	19.307	152.430

position of the interface between immiscible fluids. VOF is a multiscale approach with a wide range of applications including the prediction of jet breakup, the motion of bubbles in a liquid, and the steady or transient tracking of liquid–gas interfaces. In this work, the VOF framework was implemented to study the evolution of the interface between the polymer solution and air as a result of their mixing at the micrometer length scale. The tracking of the interface between the two fluid phases was accomplished by the solution of a continuity equation for the volume fraction of the liquid. This equation has the following form,

$$\frac{1}{\rho_{l}} \left[\frac{\partial}{\partial t} (\alpha_{l} \rho_{l}) + \nabla \cdot (\alpha_{l} \rho_{l} \boldsymbol{u}) \right] = \dot{m}_{a \to l} - \dot{m}_{l \to a}, \tag{3}$$

where $\dot{m}_{l \to a}$ is the mass transfer rate from the liquid to the gas phase, $\dot{m}_{a \to 1}$ is the mass transfer rate from the gas to the liquid phase, and α_l is the volume fraction of the liquid phase. The volume fraction of the gaseous phase (α_g) was computed with $\alpha_l + \alpha_g = 1$. The density (ρ) and the viscosity (μ) of the fluid mixture are assumed to depend on the volume fraction as:

$$\rho = \alpha_{\rm g} \rho_{\rm g} + \alpha_{\rm l} \rho_{\rm l},$$

$$\mu = \alpha_{\rm g}\mu_{\rm g} + \alpha_{\rm l}\mu_{\rm l},$$

where μ_g is the viscosity of the gaseous phase.

To cope with the turbulent flow and the viscous effects taking part in the formation of inner bubbles and subsequent filament ejections^{23,24,32,33} the LES (large eddy simulation) model was selected. In LES, large eddies are resolved directly using the Navier–Stokes equations, while small eddies are resolved applying a Reynolds-Averaged-Navier–Stokes (RANS) model. The main feature of the LES model is the ability to spatially filter small-scale phenomena, which is performed by the following expression:

$$\overline{\phi}(x) = \int_{F} \phi(x') G(x, x') dx'$$

where *F* denotes the fluid domain, *G* is the filter function that determines the scale of the resolved eddies, $\phi(x')$ is the actual value of the flow variable, and $\phi(x)$ is the filtered flow variable.³⁵ The subgrid-scale stresses are computed from:

$$\tau_{ij} - \frac{1}{3}\tau_{kk}\delta_{ij} = -2\mu_t\overline{S}_{ij}$$

where \bar{S}_{ij} is the rate-of-strain for the resolved scale and μ_t is the subgrid-scale turbulent viscosity, which was modelled using the Smagorinsky–Lilly model,

$$\mu_{
m t} =
ho L_{
m s}^{\,\,2} \sqrt{2 \overline{S}_{
m ij} \overline{S}_{
m ij}}$$

where L_s is the mixing length for subgrid scales, which is computed with $L_s = \min(\kappa \times d, C_s \times \Delta)$ where κ is the von Kármán constant (typically 0.4), d is the distance to the closest wall, C_s is the Smagorinsky–Lilly constant and Δ is the local grid scale. C_s is used to adjust the magnitude of subgrid-scale turbulence, which was determined to be 0.23 for

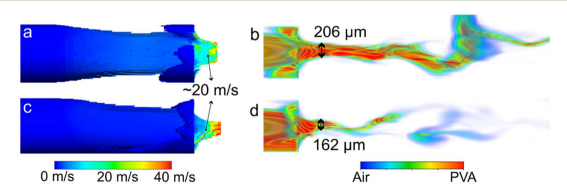


Fig. 7 Characteristic values obtained from simulations for PVA with $\Delta P_{l} = 200$ kPa and $\Delta P_{g} = 300$ kPa (a and b); and $\Delta P_{g} = 330$ kPa (c and d). Images on the left side are velocity contour plots and those on the right side are volume fraction contour plots.

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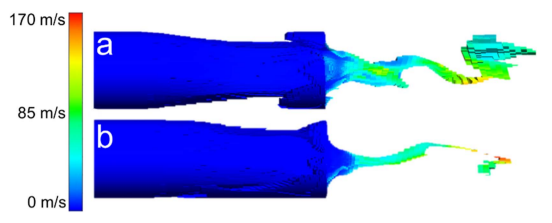


Fig. 8 Velocity fields values obtained from simulations for PVA with $\Delta P_l = 200$ kPa and ΔP_q of (a) 300 kPa; (b) 330 kPa.

Table 2 Summary of experimental conditions

	$\Delta P_{\rm g}$ (bar)	$Q_{\rm g}$ (L min ⁻¹)	$\Delta P_{\rm l}$ (bar)	$Q_{\rm l} ({\rm mL} {\rm h}^{-1})$	$T_{\mathrm{f}}\left(^{\circ}\mathrm{C}\right)$	$T_{\rm r}$ (°C)
Set 1	1.30	20	1.1	68.18	300	195
	0.82	15	0.71	50.37		
	1.80	25	1.43	38.17		
	2.29	30	1.82	34.96		
Set 2	0.82	15	0.71	50.37	150	75
					300	195
					450	315
					600	435

homogeneous isotropic turbulence within the inertial subrange. However, it was observed that this value resulted in excessive damping of large-scale fluctuations in the presence of mean shear and in transitional flows near solid boundaries, and it needs to be reduced in such regions. A value of approximately 0.1 yields optimal results for a broad range of flows, and this value is the default option in ANSYS Fluent®, and thus it was employed in this study too.^{35,37}

Results and discussion

3.1 Computational results

3.1.1 CFD model validation. FB atomization of lowviscosity, Newtonian liquids has been studied extensively both theoretically and experimentally by us and others. 23,29,30,38 From the seminal work of Gañán-Calvo, 23,30,38 several authors have reported direct observations of the formation of bubbles in the interior of the liquid feeding tube, in the surrounding of its tip, which is considered the main feature of this type of liquid atomization. 25,33-35,39,40 Such knowledge of the technique as well as our experience identifying various underlying phenomena^{23,29,31,38,41} allow us to use simulations as a solid tool to interrogate more complex systems. Therefore, to validate our computational approach for polymer solutions, firstly, simulations of FB atomization of water were carried out. In these set of simulations, the liquid overpressure (ΔP_1) was set at 200 kPa while the gas overpressure (ΔP_g) was varied to study its effect on

the atomization dynamics. Here, we present two cases: $\Delta P_{\rm g} = 200$ kPa and $\Delta P_{\rm g} = 350$ kPa.

In the first case, despite the geometrical constrain Φ < 0.25, which dictates whether the micro-mixing occurs, the gas overpressure is not sufficiently high so as to implode into the liquid feeding tube. As a result, a gas cavity and inner bubbles are not formed and thus the micro-mixing is not triggered. As depicted in Fig. 3a, in this case, a surface-disturbed liquid jet is ejected, which corresponds to the regime known as Turbulent Flow Focusing (TFF³0). Under these conditions, a nearly axial liquid vein (red color) surrounded by a liquid mist composed of droplets is ejected. Also, the conical shape of the ejection is remarkable thus representing the dynamics of TFF reported theoretically by us.³0

By increasing the gas overpressure to 350 kPa, the flow blurring regime is achieved (Fig. 3b-e). Particularly, the perspective view of Fig. 3b depicts inner bubbles (blue color) in the interior of the tube. In this case, the ΔP_g is sufficiently high to implode into the liquid feeding tube and produce the inner gas cavity, thus triggering the micro-mixing mechanism. Furthermore, a cross-sectional view (Fig. 3d) at a position just upstream of the liquid feeding tube's outlet evidences the presence of numerous inner bubbles not only in the vicinity of the walls, but also in positions close to the central axis of the tube. The bubbles introduce a turbulence component into the liquid, which causes the mixing at micrometer length scales that leads to the formation of a polydisperse aerosol. The formation of bubbles in the atomizer interior is the main characteristic of the flow blurring regime as reported by us and others. 23,30,34,35

Rather than a liquid vein surrounded by droplets as in the TFF (Fig. 3a), in this case, the outcome is a fine aerosol. In this situation, the development of a boundary layer appears to be responsible for the formation of a thin air layer between water and the tube wall.²⁴ Note that the simulation captured the recirculation of the gas flux observed by us and other authors,^{42,43} which is indicated by the presence of a bubble (mainly green color) in the center of the liquid feeding tube (Fig. 3e). The qualitative results presented here agree with the physical characteristics of flow blurring atomization process

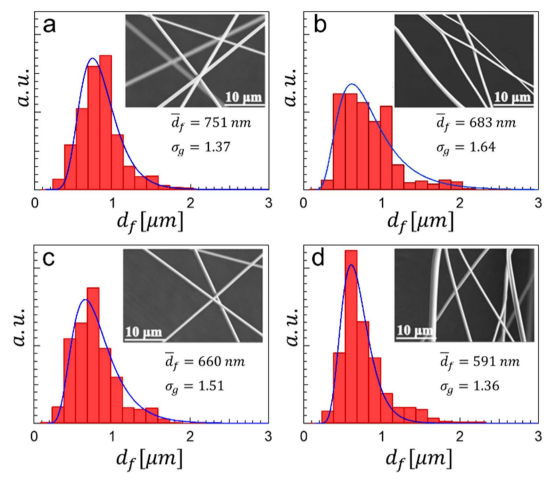


Fig. 9 Fiber diameter distribution for varying Q_l [mL h⁻¹] of (a) 68.18, (b) 50.37, (c) 38.17, and (d) 34.96. \bar{d}_f is the geometric mean diameter and σ_g is the geometric standard deviation.

that are reported in the literature. 23,24,32 It is important to emphasize that such bubble can be observed in both Fig. 3c and d, as they represent the same timestep from perspective and lateral views, respectively. The phenomenon is similar to that of Fig. 3b, where all internal bubbles are distinctly highlighted in blue. It is noteworthy to mention that in these simulations the bubbles do not consist entirely of air in the majority of cases. This is evident in Fig. 3d, where a lateral bubble exhibits a predominantly blue color (indicating a predominantly airfilled composition), while other greenish regions likely represent smaller bubbles. Owing to limitations in mesh resolution, the precise definition of bubble edges is not achieved, resulting in the simulation depicting these areas as regions with an approximate air-to-liquid volumetric fraction ranging from 40% to 60%. In spite of that, the simulations capture the essence of the phenomenon, and we thus conclude that the model is adequate to proceed with simulations of flow blurring atomization of polymer solutions as well as to obtain valid qualitative and quantitative results.

3.1.2 Simulations of PVA solutions' atomization in the flow blurring regime. For the simulation of flow blurring atomization of a 6 wt% PVA solution in water and ethanol (water mass fraction $\chi_{\rm w}=0.35$) a power-law viscosity model was implemented to take into account the shear-rate dependant viscosity

that characterizes the non-Newtonian behavior of the polymer solution. The approach is similar to that of water (the mesh is the same): the PVA overpressure, $\Delta P_{\rm l}$ was fixed at 200 kPa, while the gas overpressure was varied. Here, we present results for two cases: $\Delta P_{\rm g} = 300$ kPa (Fig. 4) and $\Delta P_{\rm g} = 330$ kPa (Fig. 5).

There exist key differences between atomization of PVA and that of water. For instance, rather than a gas cavity, in the PVA cases the air stream envelops the liquid jet and causes the diameter of liquid stream to reduce as it proceeds forward, out of the liquid feeding tube. Furthermore, a thin liquid column is ejected in both PVA cases, which downstream breaks up producing liquid filaments. This behavior was indeed observed by Ramos-Escobar *et al.*²⁴ and Marín-Brenes *et al.*,⁴⁴ using PVA and poly(ethylene oxide) solutions.

Comparison of simulations of both PVA cases evidences the effect of $\Delta P_{\rm g}$ or decreasing $Q_{\rm l}$. For lower $\Delta P_{\rm g}$, the ejected liquid column appears to have a larger characteristic diameter and travels further before it breaks up. The sequential frames of Fig. 4 depict a continuous liquid vein that appears to move in a chaotic fashion driven by the turbulence, and filament fragmentation is not observed. Conversely, rising the gas overpressure $\Delta P_{\rm g}$ leads to an ejection of a thinner liquid column, which rapidly breaks up producing filament-like ejections in a chaotic and turbulent fashion, as shown in the frames of

b a.u. $\overline{d}_f = 534 \, nm$ $\overline{d}_f = 683 \, nm$ = 1.61 = 1.64 $d_f[\mu m]^2$ 3 $d_f[\mu m]^2$ 3 d C a.u. $\overline{d}_f = 767 \, nm$ $\overline{d}_f = 774 \, nm$ = 1.69= 1.70 $d_f[\mu m]$ $d_f[\mu m]$ 900 e 800 700 600 500 400

Fig. 10 Diameter distribution for different T_r : (a) 75 °C, (b) 195 °C, (c) 315 °C, and (d) 435 °C. d_f represents the geometric mean diameter and σ_g is the geometric standard deviation. (e) Fibers' geometric mean diameter as a function of real furnace temperature, T_r (the green dotted line is to guide the eye only).

100 200 300 400 500

 T_r (°C)

Fig. 5, particularly from frame 5 onwards. Rising $\Delta P_{\rm g}$ equals to increasing the available energy to induce liquid filament thinning and its subsequent break up. In FB atomization, $\Delta P_{\rm g}$ in combination with $\Delta P_{\rm l}$ regulate the liquid flow rate, which implies that, in general, for a given liquid overpressure a higher gas overpressure results in a lower $Q_{\rm l}$. These simulations represent well direct observations reported by Modesto-López et al., ³² Ramos-Escobar et al., ²⁴ and Marín-Brenes et al. ⁴⁴ and confirm that for polymer solutions, in the FB regime, a lower liquid flow rate produces several, dispersed filaments, while increasing the liquid flow rate produces a continuum structure.

As the liquid structures move downstream, they undergo diverse transformations led mainly by inertio-capillary phenomena. Such mechanisms, described by Ramos-Escobar and others distinguish between those involving a single fluid entity and those related to two or more liquid structures interactions. Several of those transformations are present in our CFD simulations herein. Fig. 6a depicts the so-called elongation phenomenon where a relatively thick liquid filament is elongated by means of inertia leading to a thinner

structure. The transition from droplet to a filament due to inertia-capillary forces is clearly depicted in Fig. 6b similar to that captured by the high-speed recording of Ramos-Escobar *et al.*²⁴ Furthermore, Fig. 6c captures the collision of an upcoming filament with a droplet leading to a new ejection, which later develops into a filament with two small spheres at both ends (see Fig. 5 of ref. 31).

The liquid ejection's velocity in the vicinity of the liquid discharge region (U^*) and the as-ejected liquid vein's diameter (L^*) strongly influence the evolvement of the liquid filaments. A scaling analysis was used to propose an equation for U^* based on operation parameters:²⁴

$$U^* \sim \left(\frac{\Delta P_{\rm g}}{\rho_{\rm l}}\right)^{1/2} \tag{4}$$

Additionally, in the atomizer interior the Reynolds number was assumed to be Re \gg 1 and Bo \ll 1, which makes it reasonable to neglect gravity effects. Furthermore, in the current case, in which either the liquid flow rate or the viscosity are large enough to prevent the formation of the gas cavity, the characteristic diameter of the liquid vein L^* is assumed to be²⁴

$$L^* \sim \left(\frac{Q_1}{U^*}\right)^{1/2} \tag{5}$$

The values of U^* and L^* calculated with eqn (5) and (6) based on the simulations' conditions are summarized in Table 1. The equivalent values obtained directly through the simulations are indicated in Fig. 7. It is noteworthy to mention that the values obtained in the simulations are of the same order of magnitude as those obtained with the scaling law proposed in Ramos-Escobar et al.24 Both results, from the simulations and from the scaling law, fit very well and thus describe the ejection phenomena with the micro-mixing mechanism accurately. Looking at the characteristic diameters, the trend represented by the scaling law indicates that increasing the liquid flow rate produces thicker liquid veins. This phenomenon is also shown in the CFD simulations and can be observed directly in the color plots (Fig. 7). Although the frames shown in Fig. 7 were chosen arbitrarily, the orders of magnitude were verified in several frames to add confidence to the results.

The liquid ejection velocities away from the exit orifice are shown in Fig. 8. Notice the high acceleration that the liquid vein experiences in such region, increasing its velocity from nearly 10 m s $^{-1}$ to 100 m s $^{-1}$ in a distance no longer than 5 mm. Furthermore, the order of magnitude of the velocities represented here are consistent with those reported in Modesto-López *et al.* 32 measured from ultra-high speed videos which were of the order of 80–100 m s $^{-1}$.

In order to obtain quantitative data of the filaments' diameter downstream of the ejection orifice, where the liquid jet has already evolved, measurements were performed using the software ImageJ. ⁴⁵ Approximately 50% of all measurements were in the range of 40–90 μ m while the remaining 50% were in the range of 90–180 μ m. Therefore, roughly, a characteristic

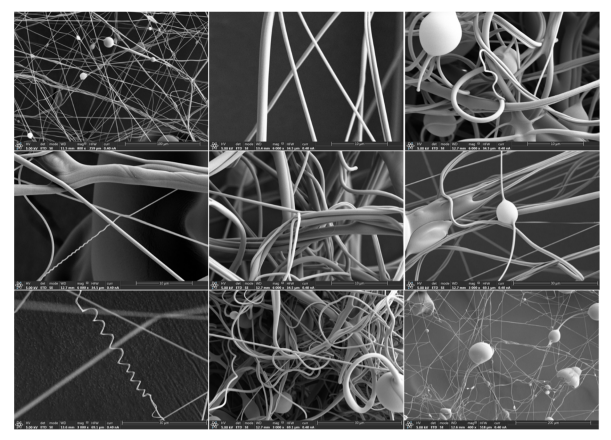


Fig. 11 SEM images of PVA fibers collected on the lateral mesh

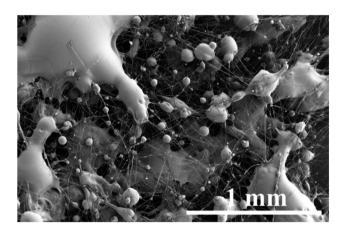


Fig. 12 Material collected on the transversal zone of the cylindrical mesh.

diameter of the filaments at a 5 mm distance from the outlet orifice would be 100 $\mu m. \,$

3.2 Liquid flow rate and driving pressures in the flow blurring atomizer

The regulation of Q_1 is critical in achieving a good control of the liquid threads' diameters. Given the relatively large values of Q_1 in our experiments, the use of volumetric means as syringe pumps was not a practical solution. Thus, we opted for

a continuous pressure differential system. However, determining such a differential to calculate the liquid flow rate is not trivial for FB atomizers because of the particular fashion in which the liquid ejection takes places, that is, through a radial implosion of the gas. As a result, a stagnation region is formed at the discharge of the liquid feeding tube, where the pressure exceeds that of the outer ambient environment $P_{\rm a}$ equal to a fraction of $\Delta P_{\rm g}$. In the following analyses, the physics underlying the liquid flow rate regulation are thoroughly described. First, the discharge coefficient is defined as:

$$C_{
m d}=rac{Q_{
m g}}{Q_{
m g,t}},$$

where $Q_{\rm g,t}$ is the theoretical volumetric gas flow rate, calculated assuming isentropic conditions and flat velocity profile at the discharge, and $Q_{\rm g}$ is the experimentally measured volumetric gas flow rate (measured at standard conditions). It is normally coincident with the discharge coefficient of a circular outlet ($C_{\rm d} \cong 0.8$, e.g. see ref. 46). The latter coefficient will also be calculated from experimental measurements. The discharge coefficient of the atomizer used herein satisfies the experimental fitting:

$$C_{\rm d} = 0.8 - 1.8 \,{\rm e}^{-1.5 \left(\frac{\Delta P_{\rm g}}{P_{\rm a}} + 1\right)},$$

which is consistent with other studies of gas discharge coefficients from small orifices. This fitting yields accurate results with errors below 0.5% along the experimental ranges of $\Delta P_{\rm g}$ used in this study.

In addition, the gas and liquid overpressures, ΔP_g and ΔP_l , driving Q_g and Q_l within the ranges of interest for producing the desired output, result relatively close. That is because the FB atomizer's unique geometry causes the gas discharge to collapse radially inward towards the liquid stream coming from the liquid feeding tube, at the atomizer discharge. Such radial implosion produces a relatively stagnant region with comparable liquid and gas velocities at the discharge of the feeding tube with an absolute pressure P_{1g} equal to a fraction of the absolute gas pressure upstream of the atomizer (P_{og}) , i.e. $P_{1g} =$ $k_{\rm n1}P_{\rm og} = k_{\rm n1}(\Delta P_{\rm g} + P_{\rm a})$, with $0 < k_{\rm n1} < 1$. The gas is unsteadily forced inside the liquid tube, in the vicinity of its outlet, producing the characteristic recirculating liquid-gas region of the flow blurring technology. The upstream front of this recirculating region has an absolute pressure $P_{\text{oR}} = k_{\text{on}} P_{\text{og}}$ such that $P_{\text{oR}} > P_{\text{1g}}$, and therefore $0 < k_{\text{n1}} < k_{\text{on}} < 1$ since the liquid-gas biphasic flow must be accelerated towards the tube outlet. Thus, the total pressure driving the liquid column from the pressurized container is given by the formula:

$$\Delta P_{\rm Tl} = (\Delta P_{\rm l} + P_{\rm a}) - k_{\rm on} P_{\rm og}, \equiv P_{\rm ol} - k_{\rm on} P_{\rm og}$$

Both $k_{\rm on}$ and $k_{\rm n1}$ are expected to be slightly dependent on $P_{\rm og}$ owing to the nonlinear (turbulent) nature of the whole gas and liquid flow outlets from flow blurring atomizers. In our setup, the Reynolds number of the liquid flow in the liquid tubing between the pressurized container and the atomizer is smaller than unity, and therefore the liquid flow rate is finally given by the formula:

$$Q_{\rm l} = \frac{\pi D_{\rm i}^4}{128 \text{ µL}} \Delta P_{\rm Tl} = k_{\rm L}^{-1} \Delta P_{\rm Tl}. \tag{6}$$

Care is taken to keep the liquid level in the pressurized container at the same height as the atomizer outlet. However, the hydrodynamic resistance, $k_{\rm L}$, of our setup is small ($k_{\rm L}=3.82\times 10^{-3}$ bar min mL $^{-1}$), resulting in small pressures compared to both $\Delta P_{\rm l}$ and $\Delta P_{\rm g}$. Thus, experimental measurements for the calculation of $k_{\rm on}$ for the specific atomizer used will entail large errors using standard pressure sensors. This value is usually comparable to the coefficient of discharge $C_{\rm d}$ of the atomizer.

A way to assess the consistency of all those experimental measurements is to evaluate the value of an applied gas pressure $P_{\rm og}^*$, a critical pressure above which the value of $k_{\rm n1}$ becomes independent of $P_{\rm og}$: this occurs when the gas outlet becomes sonic. Naturally, the stagnation pressure of this sonic discharge must be approximately equal to $P_{\rm 1g}$. Since we use air (adiabatic constant $\gamma=1.4$) discharging at atmospheric pressure $P_{\rm a}=1.012$ bar, the critical pressure must be $P_{\rm 1g}^*=1.89P_{\rm a}$, and therefore the eventual constant value of $k_{\rm n1}^*=P_{\rm 1g}^*/P_{\rm og}^*$ must be:

$$k_{\rm nl}^* = 1.89 \frac{P_{\rm a}}{P_{\rm og}^*} < k_{\rm on}^* = \frac{P_{\rm ol} - k_{\rm L} Q_{\rm l}}{P_{\rm og}^*}$$
 (7)

The data in Table 2 allow a quite precise calculation of $P_{\text{og}}^* \cong 2.8$ bar above which k_{on}^* becomes constant and equal to $k_{\text{on}}^* = 0.862 \mp 0.005$. From eqn (7) above, one has $k_{\text{n1}}^* = 0.675 < k_{\text{on}}^*$, thus proving the consistency of measurements, where an expected pressure drop equal to $(k_{\text{on}}^* - k_{\text{n1}}^*) P_{\text{og}}$ will accelerate the liquid and push the liquid–gas biphasic flow across the recirculating region for $P_{\text{og}} > P_{\text{og}}^*$.

An interesting by-product of the above-described investigation is the description of the cascade of pressures occurring at the outlet of flow blurring atomizers, which, to the best of our knowledge, has not been attempted at this level of detail before.

3.3 Microfiber production

The performed experiments and their conditions are listed in Table 2. Two set of experiments were carried out; in the first set, the gas flow rate $Q_{\rm g}$ vas varied to study its effect on the final fiber diameter and structure. It is noteworthy to mention that a change of $Q_{\rm g}$ inherently implies that $Q_{\rm l}$ should be varied to adjust the atomization conditions to the flow blurring regime, in which the micro-mixing takes place. That is, $\Delta P_{\rm l}$ was maintained to a minimum admissible for the atomization process to occur.

In those experiments, $T_{\rm r}$ was kept constant at 195 °C ($T_{\rm f} = 300$ °C). In the second set of experiments, $Q_{\rm g}$ and $Q_{\rm l}$ were fixed at 15 L min⁻¹ and 50.37 mL h⁻¹, respectively, while varying $T_{\rm r}$ in the range from 75 °C to 435 °C. This second set of experiments aimed at elucidating the effect of solvent evaporation rate on the diameter and structure of the fibers. Those experiments combined with data from the CFD simulations allowed to estimate the in-flight stretching rate of the filaments.

3.3.1 Set 1: variation of liquid flow rate. The fiber diameter distribution as well as a representative SEM image of resulting fibers as a function of Q_1 are depicted in Fig. 9. The data indicate that rising the liquid flow rate causes an increase in fiber diameter, and such correlation is expressed as $\bar{d}_{\rm f} \sim Q_{\rm l}^{0.307}$. Recalling the scaling analysis, this tendency is consistent with the experimental results since the diameter of the ejected liquid vein grows with the liquid flow rate. Thus, it seems reasonable to think that as the liquid vein becomes thicker, the phenomenon propagates downstream leading to the production of thicker fibers. Furthermore, despite the relatively high liquid flow rates employed for atomization experiments, the fiber mean diameters are in the submicrometer regime, below 750 nm, and the mode diameter is of the order of 500 nm. The results thus indicate that the liquid flow rate is one of the main operating parameters influencing the size of both, the liquid filaments and the fibers.

3.3.2 Set 2: variation of furnace temperature. The fiber diameter distribution for varying furnace temperature T_r as well as a SEM image of their corresponding fibers are represented in Fig. 9. There is a noticeable effect of T_r on the size of the produced fibers. In this case, the solvent evaporation rate would

increase with $T_{\rm r}$, thereby reducing the time that the liquid threads exist as such. Therefore, they would be converted into fibers at a shorter time, and consequently they would not have sufficiently long time to elongate and thin.

Indeed, for $T_r \ge 300$ °C the fiber diameter appears to remain constant, implying that independently of how high T_r is, under such ejection conditions the maximum achievable $\bar{d}_{\rm f}$ is of the order of 800 nm, as depicted in Fig. 10e. In addition, as reported by us in previous works and as demonstrated in our simulations, liquid threads undergo different transformation mechanisms (ejections, elongation, rotation, etc.) in flight. One of the most frequent and, perhaps, one of the most important for producing thin fibers is the elongation process, which arises from an inertia-capillary mechanism. The elongation takes place as long as the liquid threads preserve their mobility, which is, at least partially, dependent on ambient temperature through solvent evaporation. Increasing temperature in the furnace leads to an acceleration of solvent evaporation and further solidification of the liquid filaments, thus stopping or retarding the elongation process and, consequently, preventing the reduction of the fiber diameter simultaneously decreasing the length of collected fibers. This phenomenon is of significant importance at lower temperatures and becomes less relevant at higher temperatures where, as shown in Fig. 10, the increment of geometric mean diameter was smaller. For instance, at $T_r =$ 315 °C, $\bar{d}_{\rm f} = 767$ nm and at $T_{\rm r} = 435$ °C, $\bar{d}_{\rm f} = 774$ nm, that is a difference of less than 1%. However, between $T_r = 75$ °C and T_r = 195 °C the fiber geometric mean diameter increased from 534 nm to 683 nm, approximately 22%. For the flow conditions and the geometry of the furnace and collector considered herein, there is a critical $T_{\rm r}\sim 300$ °C ($T_{\rm f}\sim 450$ °C) at which further temperature increase does not result in variation of fiber diameter in fibers collected on the lateral zone of the mesh. Since the solvent of the 6 wt% PVA solution was a mixture of water/ethanol (35/65 w/w), following Raoult's law, it is reasonable to assume that the boiling point of the solvent is approximately 80 °C (molar fraction of ethanol 0.42, molar fraction of water 0.58) and at the 75 °C of the furnace the filaments had longer time to elongate as they travel before the solvent evaporates completely. In addition, fibers produced at $T_r > 315$ °C exhibited calcination effects, as observed when the mesh was removed from the furnace. The melting point of PVA is approximately 200 °C, thus exposing the fibers to higher temperature may result in thermal degradation. Furthermore, increasing $T_{\rm r}$ above 315 °C not only has minor impact on the fibers' diameters, but it also leads to poor quality fibers, probably as a consequence of extremely fast evaporation resulting from both the high temperatures and high velocities (increasing dramatically heat transfer by convection).

Fig. 11 show several SEM images of fibers collected on the lateral mesh under varying conditions. The images show PVA fibers of varying diameters and it elucidates the versatility of the technique as facilitator of novel micro- and nanofibrous materials.

The SEM images shown in Fig. 11 represent fibers collected in the lateral zone of the cylindrical mesh. Nevertheless, a SEM image showing material collected on the transversal section, at the end of the cylindrical mesh, is depicted in Fig. 12. Albeit the presence of fibers is noticeably in this zone, the dominant structures are large spheroids and smaller quasi-perfect spheres, both significantly large compared with the dimensions of the fibers. Two facts are evident from such image, first, filaments elongate in flight before colliding on the mesh were they stop and dry thus leading to solid structures, and at the final stages of evaporation, surface tension forces take over forming the spheres. Second, a longer heating zone would be required to evaporate the solvent contained in the filaments thus increasing the fiber production rate, implying that liquid ejection technique does not limit the fiber production. Nevertheless, these type of beaded micro- and nanofibers find use in high-added value applications such as microencapsulation for drug delivery or as filter media.⁴⁷⁻⁵⁰

To estimate the elongation rate of filaments per unit distance of flight, a correlation between the data obtained using CFD simulations and the data collected with the experimental setup was established. First, a representative filament' initial diameter of $D_0=50~\mu \mathrm{m}$ for the ejection zone is selected. Furthermore, it will be assumed that the filament is a perfect cylinder and because the PVA concentration is 6 wt%, the fiber will be considered to constitute the 6% of the mass of the cylindrical filament volume, thus:

$$0.06 (V_{\rm fil} \rho_{\rm sol}) = 0.06 \left(\frac{\pi D_{\rm fil}^2}{4} L \rho_{\rm sol} \right) \sim \frac{\pi D_{\rm fib}^2}{4} L \rho_{\rm PVA}$$
 (8)

where $V_{\rm fil}$ is the volume of the initial filament, $D_{\rm fil}$ its diameter, $D_{\rm fib}$ the diameter of the collected fiber, L the length of the filament and fiber, and $\rho_{\rm sol}$ (885.3 kg m⁻³) and $\rho_{\rm PVA}$ (500 kg m⁻³) are the density of the solution and the density of bulk PVA, respectively. Therefore,

$$D_{\rm fil} \sim \frac{D_{\rm fib}}{\sqrt{0.06}} \left(\frac{\rho_{\rm PVA}}{\rho_{\rm sol}}\right)^{0.5} = 5.43 D_{\rm fib},$$
 (9)

Furthermore, fibers collected at 50 cm from the ejection point had a characteristic diameter of 0.8 µm thus implying $D_{\rm fil}\sim 4.35~\mu m$ and, according to our CFD simulations, the velocity of the filaments was of the order of $\nu_{\rm flight}=100~{\rm m~s^{-1}}.$ Therefore, a filament diameter reduction of 91% occurred in only 5 ms. If the initial liquid threads did not undergo branching, splitting or breakup, its length would increase more than two orders of magnitude.

4. Conclusions

This study presents a micro-mixing-based method for the large-scale production of submicrometer-sized polymer fibers. The method involves ejecting polymer solutions at high liquid flow rates using a Flow Blurring (FB) atomizer, followed by on-line thermal treatment, resulting in the production of micro/nanofiber mats in a single step. The atomizer plays a crucial role in fragmenting the liquid stream with a gas current to generate thin filaments. Computational Fluid Dynamics (CFD) simulations were employed to investigate the dynamics of polymer solution ejection and fragmentation, while

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experiments were conducted to examine the impact of gas pressure and furnace temperature on the final fiber diameter and structure.

The simulations demonstrated that the flow blurring regime was attained when the gas overpressure was sufficiently high to implode into the liquid feeding tube, creating an interior gas cavity and triggering the micro-mixing mechanism. Moreover, cross-sectional views obtained from the simulations revealed the presence of numerous inner bubbles not only near the walls but also in positions close to the central axis of the tube. By utilizing CFD simulations to study the dynamics of polymer solution ejection and fragmentation, a deeper understanding and control of the atomization process can be achieved, potentially leading to the development of more efficient and precise methods for polymer fiber production.

Poly(vinyl alcohol) (PVA) solutions were utilized in the fiber production experiments. In the initial set of experiments, the gas flow rate was varied while maintaining a constant liquid flow rate to examine its influence on the final fiber diameter and structure. The furnace was held at a nominal temperature of 300 °C. In the second set of experiments, the gas and liquid flow rates were fixed while the furnace temperature was varied from 150 °C to 600 °C. Increasing the liquid flow rate resulted in an increase in the fiber diameter, which can be expressed as a geometric mean diameter $(\bar{d}_{\rm f}) \sim Q_{\rm l}0.3$. Furthermore, the experiments carried out at lower furnace temperatures (150 °C and 300 °C) resulted in fibers with a diameter of approximately 600 nm, while those carried out at higher temperatures (450 °C and 600 °C) resulted in fibers with a diameter of nearly 770 nm. Such fact suggests that the solvent evaporation rate has a significant effect on the diameter and structure of the fibers.

The method described herein has the potential for efficient and scalable production of polymer fibers for a variety of cutting-edge applications. The use of relatively low or medium temperatures allows for the processing of materials which may undergo thermal degradation at higher temperatures, and the capability of functioning with high liquid flow rates, which can reach tens of milliliters per hour, allows for the production of large amounts of fibers in a short period of time.

Conflicts of interest

There are no conflicts to declare.

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