Velocity and size measurement of droplets from an ultrasonic spray coater using photon correlation spectroscopy and turbidimetry

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Abstract: We have developed a combination of light scattering techniques to study and characterize droplets of an ultrasonic spray coater in flight. For this economically relevant spray coater, there is so far no reliable technique to systematically adjust the experimental parameters. We have combined photon correlation spectroscopy and turbidimetry to determine the size and speed of the droplets depending on parameters of the printing process as shroud gas pressure, flow rate, and atomizing power. Our method will allow us to predetermine these parameters to control the properties of the coated films as, e.g., thickness from tens of nanometers to micrometers.

1. INTRODUCTION

In industrial production, especially in the electronic industry, the preparation of thin films with reliable parameters as thickness or roughness with spray coating techniques is indispensable [1–5]. Ultrasonic spray coating (USSC) has been introduced for a range of active layers in electronics as organic thin film transistors [3], organic solar cells and photodiodes, and electroluminescent devices [6–8], as well as for electrodes and transistors [3,6]. Having reliable models [9] and methods to predict vital parameters to achieve reproducible and controlled films is of adamant importance. The droplet size can, e.g., be adjusted by regulating the ultrasonic frequency. Because of the small size of the droplets with narrow dispersion in size (less than 20% standard deviation according to [10]), it is possible to coat very thin homogeneous layers down to 10 nm thickness [11], which is unique compared with other spray coating techniques. This opens up the possibility to deposit functional coatings [12] on large and even three-dimensional (3D) surfaces [13]. Another important benefit is that the atomization occurs thanks to high frequency vibrations and no high velocity ejection though a small aperture. This allows the droplets to be deposited with a lower kinetic energy. Since no small aperture is required, the nozzle orifice will be less easily clogged with impurities or solid particles. Lang [14] was one of the first to find a relation between some spray parameters and the droplet size. With the formula of Lang [14], the droplet size can be predicted. For example, spraying water with the impact nozzle of Sonotek at a frequency of 120 kHz, droplets of 18 μm are expected. The relationship found was only correct when the liquid phase viscosity and the volumetric liquid flow rate do not affect the droplet size. Kim et al. [15] and Rajan and Pandit [16] have tried to predict the drop size more precisely.

The existing theories do not cover the question of how the droplet size, velocity, and concentration evolve during the flight from the ultrasonically generated droplet to the substrate but only focus on the droplet size when it arrives at the substrate. It is, however, important to understand the change of size, velocity, and concentration during the flight to predict the influence of the process parameters on the layer formation.

To answer these questions, a measuring technique needs to be developed that can determine the droplet characteristics during flight. For inkjet printing, this led to dimensionless numbers that perfectly describe the ink formulation suitable for printing [17]. However, inkjet printing is jetting only one droplet at the same time, and therefore measuring the droplet characteristics is relatively simple. Ultrasonic atomization creates thousands of droplets at the same time. This makes measuring the properties of the droplets as size, distribution in size, and velocity during the flight a complicated task. Applying a high-speed camera to measure the droplet characteristics during their flight sounds like a valuable solution. Measuring a 3D cone filled with a fog of moving droplets is, however, not that trivial with a standard high-speed camera. To follow the particles, frame rates of 20,000 are needed due to the speed of the particle; this reduces the number of pixels available. Here a compromise had to be found between the number of pixels that represent a droplet and
the field of view to track the droplets for the droplet velocity. For our camera and setup, this led to 4 pixels for a 20 μm droplet. The error in size is therefore considerable.

We made a survey, in which existing techniques could be applied [18], to study the parameters [19] of USSC. As optical methods, we found Fraunhofer diffraction [20], phase doppler anemometry [21], interferometric laser imaging [22], rainbow refractometry [23], and two-phase structured laser illumination planar imaging [24] as possible candidates. The mentioned techniques could be tried on ultrasonic-generated sprays, but this is outside the scope of our work. Our aim is to be able to measure the droplet velocity and size, to make an easy-to-handle setup. We want to avoid the use of dyes in the inks, as is necessary for some techniques above, since this leads to perturbations.

We found that a combination of photon correlation spectroscopy (PCS) [also known as dynamic light scattering (DLS)] to obtain the velocity of the droplets and turbidimetry to obtain their size is applicable to characterize ultrasonic generated droplets. For the PCS, in our case, the droplets are the scattering centers instead of particle in solution as one is used to. Due to the high difference in refractive index as well as density between liquid and surrounding gas, the droplets lead to a large enough scattering signal. For particle in solution, the size of the particle can be obtained from the Brownian motion of these particles. In the case of sprays, there is no considerable Brownian motion; thus, only the velocity can be determined by PCS and the size has to be obtained from turbidimetry. The droplet size was determined by turbidimetry, subsequently after PCS but within the same coating step [18–20]. This requires only the addition of a simple detector to monitor the light intensity passing through the sample.

To corroborate our results, we used a high-speed camera for particle image velocimetry/shadowgraphy to see we are in the right order of magnitude of the size and velocity.

2. METHODS AND THEORY

A. Ultrasonic Spray Nozzle

Schematic setup of an ultrasonic spray nozzle is shown in Fig. 1 with its characteristic parts and parameters.

Ultrasonic nozzles [14] produce standing waves in the ink on the atomizing surface. This is the result of mechanical vibrations produced by piezoelectric transducers (Fig. 1) in the nozzle. The nozzle, powered by an ultrasonic generator delivers an electric signal [parameters: power (W), amplitude A, and frequency $f_c$ (Hz)] [21] to the cooled transducers. The piezoelectric transducer and the nozzle tip transfer the electric signal to mechanical vibrations ($A_m \propto A_c$, frequency $f_m = f_c$ [22]). Typically used frequencies $f_m$ are located between 25 kHz to 180 kHz. The dimensions of the nozzle are designed to be in resonance in the operating frequency. If the amplitude of the standing waves is high enough, small droplets will break off (atomization effect) [14]. In general, higher frequency nozzles produce smaller droplets [21]. The droplet size is also influenced by adjusting the nozzle height (usually 20–90 mm) since the solvent evaporates (partly) during flight. After atomization, the droplets do not contain any kinetic energy. Therefore, a so-called shroud gas (e.g., nitrogen or argon) is used to give the droplets kinetic energy and lead them to the substrate. A higher shroud pressure (psi) will result in droplets with higher velocity. By adjusting the volumetric ink flow rate (ml/min) of the liquid feed, the number of droplets can be strongly influenced. These parameters directly influence the droplet size [23], velocity, and spray angle ($\alpha$) of the atomized solution. This directly affects the formed layer, especially its thickness.

B. Photon Correlation Spectroscopy

PCS is a well-seasoned DLS technique [24–26] for the determination of particle sizes via their Brownian motion, e.g., in polymer analytics [27]. In the standard experiments, one obtains the intensity–intensity or homodyne autocorrelation function,

$$g_2(q, t) = \langle |E(q, t_0)|^2 |E(q, t_0 + t)|^2 \rangle \iff g_2(q, t) = \langle I(q, t_0) I(q, t_0 + t) \rangle,$$

(1)

with $E$ being the field, $I$ being the intensity of the scattered light, and $q$ being the magnitude of the scattering vector. A representation of the correlation function in terms of a distribution of exponentials, is the so-called Kohlrausch–Williams–Watts function \([KWW, \text{Eq. (2)}] [28,29]\), with $\beta$ the stretching parameter describing the distribution [30] and $A$ describing its amplitude. The KWW is in the case of monodisperse particles in solution, a pure exponential with $\tau$ the characteristic time of this exponential function:

$$g_2(q, t) = A \exp \left( -\frac{t}{\tau} \right)^\beta; \quad \beta = [0, 1].$$

(2)

This homodyne correlation function is connected to the field–field or heterodyne autocorrelation function $g_1(q, t)$ via the Siegert relation [26],

$$g_2(q, t) = C \left[ 1 + f g_1(q, t) \right]^2.$$

(3)

Following [26], $g_1(q, t)$ can experimentally be obtained by adding a local oscillator, i.e., direct laser light into the detector. The dominating term in the expansion of the mathematical representation of such a heterodyne experiment is $g_1(q, t)$ [26]. An effect to note is that in the case of multiple relaxation processes in the case of $g_2(q, t)$ these are products of (stretched) exponentials while in the case of $g_1(q, t)$ these are a sum. The main advantage of heterodyne light scattering is that for low
scattering signals, the signal to noise of the correlation functions is enhanced.

Compared to standard PCS experiments, there are two questions to address in relation to our experiment:

(a) Is the comparatively low number of scattering droplets problematic? This is not the case: PCS is an interferometric technique in the sense that one observes the produced speckle field in the far field and its dynamic change. This speckle field is independent of the number of scattering moieties. Modern PCS instruments utilize as in our case single mode detection—we observe only the change of a single speckle since every speckle represents a different electrodynamic mode [31]. This maximizes the amplitude called contrast in PCS and can be used to minimize systematic noise [32].

(b) Is the size of the particles/droplets compared to the wavenumber $k$ of the incident light causing problems in determining size or velocity? For our experimental setup and our aim, this is not a problem. One has to take into account not only scattering but refraction and scattering within the droplets. It was found that this influences the results at small angles close to the forward direction [33–36]. We perform PCS at 90° scattering angle.

1. Droplet Velocity

If the particle respective of the droplets in a PCS experiment have a directed flow, than due to the Doppler shift, one observes, in the frequency domain in the spectral density $S(q, \omega)$, a frequency shifted peak. This is the position of the Doppler shift of the moving particles. However, in the case of the results at small angles close to the forward direction $\omega_{D}$, and our aim, this is not a problem. One has to take into account not only scattering but refraction and scattering within the droplets. It was found that this influences the results at small angles close to the forward direction [33–36]. We perform PCS at 90° scattering angle.

The light is scattered into an angle $\theta$, in our case 90°. The angle between the velocity vector $V$ and the scattering vector $q$ is defined as $\phi$. It should be noted that when $V \perp q$ the frequency or Doppler shift cannot be observed. To be able to measure, we tilted the beam of the shroud gas by 5° against the vertical, and now $q$ and $V$ are not perpendicular to each other anymore. For determining the propagation vector $k_i$, the Bragg condition can be used where $q = |q|$ is the magnitude of $q$.

$$q = 2k_i \sin \left( \frac{\theta}{2} \right) = \frac{4\pi n}{\lambda_i} \sin \left( \frac{\theta}{2} \right) \rightarrow k_i = \frac{2\pi n}{\lambda_i}.$$  

where $n$ is the refractive index of the measured medium. The incident light wavelength is represented as $\lambda_i$. If Eqs. (5) and (6) are combined, the droplet velocity can be extracted,

$$V = \frac{q \lambda_i}{2\pi \cos(\phi) \sin \left( \frac{\theta}{2} \right)}.$$  

In the final Eq. (8), $\omega(q)$ is replaced with the frequency ($f(q)$). The velocity of the droplet can be calculated by having one variable parameter $f$. The frequency is defined by fitting the correlation function [27] obtained through PCS as will be shown later,

1. Turbidimetry

The droplet size can be determined by performing one additional turbidimetry measurement [39]. In this experiment, the amount of transmitted light is measured and placed in relation to the droplet diameter followed by the Mie theory. For turbidimetry, the advantage of determining size and concentration at the same time using several wavelengths is not possible straightforward, as only one of the parameters can be determined independently. The Mie scattering at diameters above 5 μm show too little difference (Fig. 2) using different wavelengths.

1. Droplet Size

The size of droplets in suspension can be determined by measuring the turbidity $\tau_{\lambda_0}$. Turbidimetry measures the damping of a light beam traveling through the spray caused by the absorption and scattering of light by the droplets/particles,

$$\tau_{\lambda_0} = \frac{1}{L} \ln \left( \frac{I_0}{I} \right).$$  

$I_0$ and $I$ are representing the intensities of the incident and weakened light beams. The optical path length through the measured medium is $L$. The amount of absorption and scattering is related to the concentration and size of the droplets. Therefore, turbidity can be related to the droplet size. For droplets with monodisperse spherical diameter, $D$, a relation is defined [39],

$$\tau_{\lambda_0} = \phi N \frac{\pi D^2}{4} Q_{\text{ext}}.$$  

Absorption in a medium is related to the extinction coefficient \( Q_{\text{ext}} \). This depends on the incident light wavelength \( \lambda \) and droplet diameter \( D \). The concentration of the droplets is defined as \( \phi_N \). After combining Eqs. (9) and (10), \( D \) can be determined,

\[
D = \sqrt{\frac{-\ln \left( \frac{I}{I_0} \right)}{L \phi_N Q_{\text{ext}}}}. \tag{11}
\]

The extinction coefficient \( Q_{\text{ext}}(D) \) has an asymptote of two (Fig. 2). In our experiment, \( Q_{\text{ext}} = 2 \) is appropriate to calculate the droplet diameter because the expected average droplet size is 18 \( \mu \)m. In turbidimetry, the particle size and concentration can be determined independently by using a set of different wavelengths. In our experiment, there is only a negligible difference in extinction coefficient between two wavelengths for droplets between 10 and 100 \( \mu \)m. Based on this, only the droplet diameter can be determined with the turbidimetry measurement. The amount of multiple scattering is in our case limited and negligible, thanks to the light transmission higher than 95% [40] as shown from our experimental data in the inset in Fig. 2. As was pointed out in [40,41] only for transmission below 40% or vice versa obscuration/obfuscation 50%, a considerable influence of multiple scattering is expected. This high transmission is a direct consequence of the low density of droplets in this technique. This high transmission allows us to perform the high-speed camera observation across the spray cone, and the droplets in the focus of the camera are well resolved.

To solve Eq. (11), there are two unknown parameters, namely the diameter and the concentration of the droplets. The other parameters are known. To obtain the droplet diameter, Eq. (12) is applied,

\[
\phi_N = \frac{\dot{V}_{\text{liquid}}}{V_{\text{sphere}} A V}. \tag{12}
\]

Here, \( \dot{V}_{\text{liquid}} \) is the volumetric flow rate of the syringe pump. The volume of the droplet \( V_{\text{sphere}} \) can be replaced by an expression including the diameter \( D \) assuming the droplets are perfectly spherical. The velocity \( V \) of the droplets will be defined by PCS. Equation (8) is only correct if one assumes that the velocity over the entire surface \( A \) is constant and perpendicular in the cone. Therefore, edge effects of the velocity are not considered in this work. The surface area \( A \) can be determined by using a simple camera to measure the angle of the cone. From Eqs. (11) and (12), the droplet diameter can be obtained.

3. MATERIALS AND SETUP

For the combination of heterodyne PCS and turbidimetry, an experiment was developed (Fig. 3). Droplets were created and sprayed from an ultrasonic nozzle (Sono-Tek, Impact). This nozzle was mounted on an XYZ-stage under an angle of 85 deg to the scattering plane.

A. Turbidimetry

A laser diode (Qioptiq/Excelitas, iFLEX2000, \( \lambda = 660 \) nm, 40 mW) was used as light source, after the laser, and a beam expander (X3) was placed. The beam expander is needed to reduce the energy density of the light on the detector to avoid saturation by maximizing the signal. The intensity of the expanded beam can be fine-tuned with an attenuator (neutral density filter). This light transmits through the spray of droplets and reaches the custom build detector (including photodiode 220D, OEC GmbH). In front of the detector, a pinhole is placed whose opening is adjusted to the size of the laser beam diameter. Therefore, only transmitted light enters the detector reducing the amount of stray light. Two flip mirrors made it possible to switch easily from turbidimetry to the PCS measurement (Fig. 3).

B. Photon Correlation Spectroscopy

Measurements were performed with a heterodyne setup. This means that elastic light and inelastic light are combined for self-beating. The heterodyne detection mode was chosen for its stability and better signal/noise ratio. Light from the blue laser diode (\( \lambda = 488 \) nm) is divided into two beams by using a parallel glass plate with a reflectance of 5%. The intensities of both beams can be regulated separately with attenuators (Fig. 3). The scattered light coming from the spray falls into a single mode fiber with a Y-beam splitter into two single-photon counting modules (Avalanche diode, Perkin Elmer, SPCM-AQR). This signal is correlated by the Multiple Tau Digital Correlator (ALV GmbH, ALV-7004) in pseudo cross correlation.

C. High-Speed Camera

A high-speed camera (Photron, Mini AX100 200 K-M-32 GB) is mounted together with a bi-telecentric objective (Thorlabs, X2, MVT23200) as a reference measurement system. Illumination of the view field by a white light source (SCHOTT, KL 2500 LCD) combined with a telecentric backlight illuminator (Techspec, 52 mm). A frame rate of 20,000 frames/second was needed to obtain the velocities with high enough precision. This leads to a reduction in the field of

![Fig. 2. Diameter of the droplets influences the extinction coefficient (\( Q_{\text{ext}} \)). This is modeled with two different wavelengths of the incident light: 488 nm and 660 nm for droplet size distribution with standard deviation of 20% for each particular median diameter (data generated with http://philiplaven.com/mieplot.htm). The inset shows the measured transmitted light intensity versus the horizontal place in the spray cone. In this experiment, the diameter of the cone at the place the transmission was measured was 34 mm.](image-url)
Fig. 3. Experimental setup for turbidimetry combined with PCS. Three measurement techniques are combined: heterodyne PCS to measure the droplet velocity, turbidimetry measuring the droplet size, and a high-speed camera to corroborate our results. The ultrasonic spray coater is symbolized by the scattering volume as a red dot and is not explicitly drawn for clarity.

view for the camera to attain this frame rate. Therefore, in our experiment, the resolution for a 20 µm particle is about 4 pixels.

The experimental setup (Fig. 3) was situated on a breadboard with active vibration insulation (Scientific Instruments GmbH, TableStableTS300) mounted in a light-tight blackbox (eliminating disturbing correlations caused by extraneous scattered light sources as room lighting, sunlight).

4. DATA TREATMENT

After obtaining the raw data from the experiments, it is necessary to process this data in order to obtain the required results. In this chapter, it is explained how to transform the obtained data to the needed parameters for the Eq. (8). Subsequently, the droplet velocity-diameter can be obtained.

A. Photon Correlation Spectroscopy

A digital multitau correlator (ALV GmbH, ALV-7004) is used for signal processing to obtain $g_1(q,t)$ (Fig. 4) or $g_2(q,t)$. The correlation function itself has an exponential decay with a shifted additional peak. This peak represents the Doppler shift. To obtain the velocity of the droplets, the angular frequency of this peak must be determined. Figure 4 shows that it is not possible to fit the correlation function properly and extract the angular frequency from the attenuated cosine, which is strongly damped.

Therefore, we made a fast Fourier transform (FFT, Fig. 5) of the data. Now the Doppler shift can be read out from the real part of the spectrum. Modern correlators like the one used have a quasi-logarithmic time base. A FFT needs a linear time base with the sampling rate to be double as the highest frequency in the data (Niquist theorem). Therefore, before applying the Fourier transformation, the data is interpolated linearly with a time base equivalent to the fastest lag time.

The shifted peak against the central Lorentzian represents the Doppler shift (Fig. 5). The fit was performed with the Levenberg–Marquart algorithm used in Origin (Originlab.com) nonlinear fitting routine. To reduce the complexity, only two Lorentzians are fitted in the interested area.

Fig. 4. Heterodyne correlation function $g_1(q,t)$ of a PCS measurement—the small peak at $3 \times 10^{-5}$ s contains information about the measured droplet velocity. $g_1(q,t)$ is fitted with Eq. (4a) [26] (function displayed in Table 1). The experimentally observed cosine is strongly damped. The inset shows an enlargement of the main feature. The fit function is displayed as a full (red) line.

Table 1. Fit Function Used (Fig. 4)$^a$

| Equation | $y = A_1 \exp\left[-\left(\frac{t}{\tau_1}\right)^{\beta}\right] + A_2 \exp\left[-\left(\frac{t}{\tau_2}\right)\right] \cos(\omega t) + \text{BG} |$
|-----------|--------------------------------------------------|

$^a$This is a combination of KWW Eqs. (2) and (4a).
Fig. 5. $S(q, t)$, the real part of the Fourier transformation of $g_1(q, t)$. The frequency spectrum is fitted with two Lorentzians according to Eq. (4b). The cumulative peak of the fit is displayed on top of the function as a full (red) line.

Table 2. Parameters of Fit with Two Lorentzians

| Parameter   | Value   |
|-------------|---------|
| FWHM1       | 1.75E5  |
| Amplitude1  | 1.57E8  |
| Peak2       | 5.74E5  |
| FWHM2       | 1.08E6  |
| Amplitude2  | 9.97E8  |
| baseline    | 0       |

Table 3. Spraying Parameters of the Series of Experiments in Fig. 6 with Increasing Shroud Pressure

| Shroud Pressure (psi) | Flow Rate (ml/min) | Atomizing Power (W) | Spray Height (mm) | Ink   |
|-----------------------|--------------------|---------------------|-------------------|-------|
| 0.2                   | 0.4                | 0.6                 | 0.8               | 1.0   |
| 1.0                   | 1.2                | 1.5                 |                   | Water |

Fig. 6. Fast Fourier transformation of the experimentally obtained $g_1(q, t)$. The peak of the second Lorentzian shifts to the right by increasing the shroud pressure resulting in a higher droplet velocity.

5. RESULTS AND DISCUSSION

To test our measurement system, we performed USSC experiments with water as ink. Here, the shroud pressure, volumetric flow rate of the ink, and atomizing power have been increased to study the droplet size and velocity. The results are compared with the data from the high-speed camera as well as with predictions from theory for the droplet diameter [16]. Theoretical calculations or initial experimental measurements for the droplet speed for USSC are not available in literature, indicating the innovative character of this research.

A. Increasing Shroud Pressure

1. Droplet Velocity

The frequency shift was studied increasing the shroud pressure (parameters Table 3). If the shroud pressure increases, so should the droplet velocity. By increasing the shroud pressure, the second peak shifts to the right (Fig. 6), meaning that the second peak is located at a higher frequency resulting in a bigger Doppler shift giving a higher measured droplet velocity [Eq. (8)]. This proves that PCS can detect velocity changes in ultrasonic sprays. Three independent series of measurements (Table 3) were performed. The obtained droplet velocities from the PCS and HSC correlate [Figs. 7(a) and 7(b)]. The results of the PCS have the same magnitude and trend as measured with the HSC. The droplet velocity increases with increasing shroud pressure. The spread and accuracy of the results are improved with the PCS compared to the HSC.

2. Droplet Diameter

Turbidimetry measurements were made as a series of three independent experiments (Table 3). Using the PCS obtained droplet velocity [Fig. 7(a)], the average diameter of the droplets is obtained (Fig. 8). Spraying water with the impact nozzle (120 kHz), the expected droplet size according to Lang [14] agreed with our turbidimetry measurement. The average diameter is $18 \pm 3 \mu m$ (Fig. 8) and is not influenced by the shroud pressure.
Fig. 7. Droplet speed versus increasing shroud pressure (pressure of the nitrogen shroud gas) from three independent runs; line is the average as linear fit. The results of (a) the PCS have the same slope within the error as (b) the HSC. Higher accuracy of PCS results compared to the HSC (slope: PCS, 2774 and HSC, 2683).

Fig. 8. Droplet diameter versus increasing shroud pressure (pressure of the nitrogen shroud gas) from three independent runs; line is the average as linear fit—no influence in the shroud pressure range studied (slope: $-0.03$).

B. Increasing the Volumetric Ink Flow Rate

1. Droplet Velocity

The effect of the volumetric ink flow rate on the droplet velocity was investigated with PCS by performing three independent series of measurements (Table 4). Decreasing droplet velocity is observed with increased volumetric ink flow rate [Fig. 9(a)]. The magnitude and trend did agree with the HSC [Fig. 9(b)].

2. Droplet Diameter

Turbidimetry measurements were made from three series of independent measurements (Table 4). Together with the obtained droplet velocity [Fig. 9(a)] and the turbidimetry measurement, the droplet diameter is obtained (Fig. 10). This diameter is dependent on the volumetric ink flow rate and did agree with the theoretical predictions according to Rajan and Pandit [16].

3. Increasing Atomizing Power

For completeness, we studied the influence of the atomizing power from 1 W to 4 W at a spray height of 40 mm, a flow rate of 1.5 ml/min, and a shroud pressure of 0.8 psi as well and found only moderate influence on the droplet velocity and a linear increase in droplet diameter.

6. CONCLUSION

By a combination of PCS and turbidimetry, the determination of droplet diameter-velocity relations and their dependence on flow rate, shroud pressure, and atomizing power of ultrasonically generated sprays was demonstrated. In our study, we show that this combination of PCS and turbidimetry is a valuable solution to measure the droplet characteristics. To the best of our knowledge, measurements on droplets with PCS and turbidimetry have not been explicitly demonstrated before.

The velocities measured with PCS are in agreement with the velocities measured with a high-speed camera. The measured droplet diameter confirms the theoretical predictions.

Table 4. Spraying Parameters of the Series of Experiments in Fig. 9 with Increasing Ink Flow Rate

| Shroud Pressure (psi) | Flow Rate (ml/min) | Atomizing Power (W) | Spray Height (mm) | Ink       |
|-----------------------|--------------------|---------------------|-------------------|-----------|
| 0.8                   | 1.0-2.0-3.0-4.0    | 2.5                 | 40                | Water     |

Fig. 9. Droplet speed versus increasing volumetric ink flow rate. (a) PCS, (b) HSC show the same trend and quantitative results; full (red) line is the average as linear fit (slope: PCS, $-147$ and HSC, $-180$).

Fig. 10. Droplet diameter versus volumetric ink flow rate from three independent runs; full (red) line is the average as a b-spline. The droplet diameter is increasing nonlinear as result of the increasing volumetric ink flow rate.
Our modified setup was even able to characterize the droplet velocity and diameter more accurately and cost-efficiently than using a high-speed camera where, due to the high frame rates needed, the spatial resolution is limited. This opens a way to map out the influences of the spray coat parameters on the quality of wetting related to the ink. This study extends the application and possibilities of PCS and turbidimetry to droplets and the limits of particle size. Therefore, a combination of PCS and turbidimetry is a powerful tool to characterize ultrasonic generated droplets.

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