Increasing and decreasing droplets velocity in micro channels

L. Prat, F. Sarrazin, J. Tasseli\(^1\), A. Marty\(^1\)

Institut National Polytechnique de Toulouse
CNRS – Laboratoire de Génie Chimique – UMR 5503
BP 1301, 5 rue Paulin Talabot, 31106 Toulouse Cedex 1, France.
mel : Laurent.Prat@ensiacet.fr   tel : (+33)534615252   fax : (+33)534615253

\(^1\) CNRS - Laboratoire d’Analyse et d’Automatique des Systèmes
7 avenue du Colonel Roche - 31077 Toulouse Cedex 4, France.

Abstract: This paper deals with a specific aspect of non miscible liquid-liquid systems in microfluidic. For Chemical Engineering applications, the main constraints of functioning lies in the droplets velocity and frequency. Furthermore, the material used and the composition of the fluids is often imposed by the chemistry of the system (material resistance, fluids composition) and there is no possibility of adding other compound (surfactants for example). A technique under evaluation is presented: by using secondary channels and pumps, it is possible to increase or decrease at will the droplets velocity after they have been generated. Some experimental results are presented and discussed, including the possible limits of such an approach.

Keywords: microdroplets, flows manipulation, synchronisation, fine chemicals processes development
1. Introduction

The problem of controlling the flow patterns formed by two phase flows in microchannels has attracted a lot of attention. The parameters influencing these patterns are numerous; it involves the nature of the two fluids (including possible use of surfactants), the geometry and configuration of the injection, as well as the chip surface characteristics, etc. Indeed, the physics in such systems is largely dominated by interfacial effects. All this makes the problem difficult to describe in a simple way.

The first way of influencing the flow regimes (and the droplets sizes), is to vary the inlet fluids flow rates or their characteristics. These studies often used the Weber number (which compares inertial effects to surface tension). Gañán Calvo (Gañán Calvo, 1998) used a cross-type junction; Anna (Anna et al., 2004) demonstrated the existence of a wide range of droplet break up regimes in a flow focusing geometry. They studied the shear and the viscosity influences. The T-junction and the flow focusing have been used to form drops (Umihanowar et al., 2000; Thorsen et al., 2001; Nisisako et al., 2002; Dreyfus et al., 2003; Tan et al., 2003).

Several authors have noted the importance of the fluid interaction with the walls, for the flow pattern formation. Generally, the continuous phase completely wets the walls while the dispersed phase must be non-wetting (Nisisako et al., 2002; Tice et al., 2003; Dreyfus et al., 2003). The wetting property of liquids on the substrate can be changed by using surfactants or by surface treatments. As an example, the surface properties of PDMS can be changed by exposing it to UV light or by oxygen plasma treatment (Efimenko et al., 2002). These methods cause chain scissions in the polymer and therefore change the wetting properties of the surface. Hydrophobic microchannels (glass, silicon) can be created by coating the walls (silanization, for example (Brzoska et al., 1994)).

In Chemical Engineering, the ability of accurately controlling the flow characteristics of droplets produced in microfluidic systems without changing the flow rates and the natures of the respective phases is of great interest. For example, for a data acquisition objective, the main constraints to choose the velocity and frequency of the generated droplets are linked to the measurement tools. When determining kinetics of reaction using Raman or IR-absorption, the presence of the continuous phase can hide the product signals (inside the droplets). It is then necessary to synchronize the droplets with the analytical detector frequency. Others examples where such synchronization is useful can be found in micro-PIV measurements. In reactive systems, a very good synchronization is also needed when reaction occurs after the coalescence of droplets, each droplet containing one of the reagents (Sarrazin et al., 2005) or for laser induced reactions.

Then, in such applications of micro droplets (production or data acquisition), many constraints lead the implementation of chemical reactions in microfluidic systems:

- the physico-chemical properties of the reagents phase cannot be changed without changing the chemistry of the system;
- for acquisition purpose, the analytical means often lead to constraint the flows regime. Indeed, for inline measurements (ombroscopic, PIV, UV or Raman measurements), the pseudo stationary hypothesis is very useful in regards to analytical signal. The very large number of flowing droplets allows the use of statistic methods and allows integrating the acquisition signal during large periods;
- for mixing consideration, the velocity of the droplet in the microchannels is also constrained in certain values.

All these constraints imply that for a given fluid system, the frequency, velocity and flow rates of continuous and dispersed phases must be situated in a narrow “operating window”.
The present paper presents a feasibility study on a way to slow down or accelerate droplets by removing or increasing part of the continuous phase inside of the micro channels. In a first part, the geometry of the reactor is described, as well as the expected behaviour. In a second part, the droplets generation is studied in function of the flow rates. In the last part, the droplets slow down experiments are presented and analyzed with global approach.

2. Experimental device

The micro-channels are made out of two parts of polydimethylsiloxane (PDMS); one containing the channels is formed on wafers made by photolithography. The two parts are stuck together by contact after oxidation in plasma chamber. Thus, all the channels walls are in PDMS, which limits fluids wetting problems, like water sticking on glass. The channels section is rectangular, with a constant height equal to 60 micrometers and a width different according to the position: 100 µm at the T-junction and on the main channels, 20µm on the secondary channels. The channels precision has been verified by profilometer and is over 93%.

The fluids are stocked in syringes placed on syringe pumps and they are flushed at constant flow rates into polyethylene tubings connected to the micro-channels. The continuous phase is silicone oil (20 cSt viscosity), and the dispersed phase is water.

According to the high droplets velocities and the small channel lengths, the observation can be done only with a fast camera (NV1000) connected to a microscope (Nikon SMZ-10). Images acquisition rate varies between 600 and 1200 frames per second. Exposition time (about 400 microseconds) should be adapted so as to satisfy two conditions: it should be large enough to capture enough light for the contrasts distinction, and short enough to conserve sharp drops edges.

Figure 1 shows the typical geometry of the devices. They are composed of a T-junction used for droplets generation and of smaller in width T-junctions to fill-out or fill-in the continuous phase (oil) and then slow down (or accelerate) the droplets.

![Figure 1: geometry of the micro system.](image-url)
3. Observation, discussion and conclusion

In order to calibrate the presented system, the size of the droplets generated with a water flow rate of 100μL.h⁻¹ \( (F_{\text{H2O}}) \) has been studied while varying the oil flow rate \( (F_{\text{OIL}}) \). The objective in this short paper is not to study the generation system (with a Weber number approach for example). This information should be used to qualify the studied system. The pictures obtained are presented in figure 2.

![Figure 2: droplets generation function of oil flow rate](image)

The second series of experiments has consisted in filling out part of the continuous phase in a 3 consecutive 20μm T-junctions. Figure 3 shows two typical images recorded during the experiments and the evolution of the droplets velocity in function of \( F_{\text{OIL, out}} \). With this device, it is then possible to slow down the droplets and then gain a degree of freedom in the system.

![Figure 3: droplets velocity (mm.s⁻¹) in function of oil fill-out rate (μL.h⁻¹)](image)

One difficulty lies in the deformation of the droplets when passing in front of the secondary T-junctions part. As it can be seen in the pictures of figure 3, when the continuous phase is flowing out, the droplet is deformed and can break up.

This is illustrated in figure 4. The used geometry is a junction with an angle of 135° angle in respect with the main flow direction. The width of the secondary channels is 30μm and 100μm for the main channel. Function of the fill-out oil flow rate, the oil droplet is deformed when passing in front of the junction. When \( F_{\text{OIL, out}} \) reaches 125μL.h⁻¹, the droplets break up in a symmetrical way: two small parts in the secondary channels and the main part in the main channel.
This study shows that it is possible to slow down or accelerate droplets in non miscible liquid-liquid micro systems by filling in or out secondary channels with the continuous phase. The limit is reached when the droplets break up due to high shear stress at the secondary channels junction. In a Chemical Engineering point of view, these results are of great interest for numerous applications where there is no real possibility to act on the surface tension by modifying the composition of the liquid. Such applications can be found in analytical means implementation, in reactive systems or for separation purpose.

Of course, this study needs enhancement in order to predict the possibilities in term of operating domain of such a system. But the presented technique, by introducing a new operating variable, allows increasing the operating domain.

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