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Roles of gas in capillary filling of nanoslits

Fabien Chauvet, Sandrine Geoffroy, Abdelkrim Hamoumi, Marc Prat and Pierre Joseph

Control and understanding of flows inside fabricated nanochannels is rich in potential applications, but nanoscale physics of fluids remains to be clarified even for the simple case of spontaneous capillary filling. This paper reports an experimental and modelling investigation of the role of gas on the capillary filling kinetics slowdown in nanoslits (depth going from 20 nm to 400 nm) compared to Washburn's prediction. First, the role of gas through the usually observed trapped bubbles during a nanoslits capillary filling is analysed thanks to experiments realized with water, ethanol and silicone oil in silicon-glass nanochannels. Bubbles are trapped only when slit depth is below a liquid-dependent threshold. This is interpreted as possible contact line pinning strength varying with wettability. Stagnant trapped bubbles lifetime is investigated for the three liquids used. Experimental results show that bubbles are first compressed because of the increasing local liquid pressure. Once the gas bubble pressure is sufficiently high, gas dissolution induces the final bubble collapse. Influence of the bubbles' presence on the capillary filling kinetics is analysed by estimating viscous resistance induced by the bubbles using an effective medium approach (Brinkman approximation). Surprisingly, the bubbles' presence is found to have a very minor effect on nanoslits capillary filling kinetics. Second, the transient gas pressure profile between the advancing meniscus and the channel exit is computed numerically taking into account gas compressibility. A non-negligible over-pressure ahead of the meniscus is found for nano-scale slit capillary filling. Considering the possible presence of precursor films, reducing cross-section for gas flow, leads to a capillary filling kinetics slowdown comparable to the ones measured experimentally.

A. Introduction

Flows at small scales are ubiquitous, from porous media to blood circulation inside microvessels; and they have raised scientific interest for a long time. From the 90's, the use of microfabrication technologies—initially developed for microelectronics—to manipulate the liquid inside deterministic networks of channels, has led to the overwhelming development of microfluidics. Motivation for this growth is mainly practical, since the field is strongly driven by the potential applications of labs-on-chip: miniaturization leads to smaller amount of sample, high-throughput and automated analysis, reduction of time and cost, key features for analytical chemistry, or even medicine (point of care diagnostics for instance). In spite of this applied finality, the possibility to build efficient, innovative microfluidic applications has not only originated from technological progresses, but also frequently from fundamental studies on fluid mechanics and flow physics, leading the way to novel or improved functions: diffusiophoresis in salt gradients for enhanced mixing, reduced friction on superhydrophobic surfaces, biomolecule concentration due to salt effects at a micro-nanochannel interface.

This last example involves fluid confined in nanometer-sized channels: nanofluidics is an emerging topic that appeared very recently and is strongly linked to nanotechnologies. It has also a strong potential for applications (let us cite perspectives for water salt removal, DNA elongation, or even energy conversion), but is very closely linked to fundamental phenomena. Indeed, nanoscale fluid physics is particularly rich and still only partially explored: hydrodynamic boundary condition, enhanced flows inside carbon nanotubes, proton titration by acidic walls, limit of validity of continuous description are some of the basic nanofluidic open fundamental questions.

In this context, we address in this paper the simple case of capillary invasion of nanochannels. This process, also baptized spontaneous imbibition in the context of porous media, is related to very different practical situations: in trees, in aging of concrete, for separation in zeolites, for oil recovery; which is an additional motivation to this study.

Almost one century ago, Lucas and Washburn described the dynamics of a liquid entering a tube by capillarity: meniscus velocity is driven by a balance between viscous friction in the
liquid within the already wetted part of the channel, and a driving force due to Laplace pressure at the liquid front. For the simple case of a rectangular tube with low aspect ratio (width \( w \gg \) depth \( d \)), the simplest description based on an increasing hydrodynamic resistance balancing a constant capillary driving force leads to a meniscus position \( h \) versus time \( t \) that can be written:

\[
h^2(t) = At \quad \text{with} \quad A = \frac{\gamma a \cos \theta}{3 \mu_i},
\]

where \( \gamma \) is the liquid surface tension, \( \theta \) is the contact angle and \( \mu_i \) is the liquid viscosity.

At the macro or even micrometer scale, the current description of spontaneous imbibition is quite accurate, with different refinements accounting for inertia, \(^{21,22}\) dynamic contact angle, \(^{23}\) or the transient velocity profile and pressure field at the entrance being considered, \(^{24}\) or including viscous dissipation in the gas phase. \(^{25}\) In porous media, observed deviations from the standard law \(^{26}\) have been analyzed as resulting from pore size distribution, but the analysis is complex as demonstrated recently; \(^{27}\) and the case of water-ethanol mixtures in model nanoporous packed beads shows non-trivial dynamics. \(^{28}\)

At the molecular scale, many numerical results have been derived by using molecular dynamics for situations related to imbibition: dynamic contact angle, \(^{29}\) capillary flows of \( x \)-alcohol or water in nanopores, \(^{30,31}\) evaporation of fluid bridges in slit-like pores, \(^{32}\) role of slip boundary condition on the capillary filling in carbon nanotubes, \(^{33}\) influence of a prewetting monolayer inducing a strong slip. \(^{34}\)

Nevertheless, as already evoked, there are still questions at dimensions below about 100 nm. A number of experimental works have recently been performed with silicon-glass nanoslits (low aspect ratio nanochannels), with depth in the 5–500 nm range, and widths of the order of 1–10 \( \mu \)m defined by photolithography. \(^{35}\) Fabrication of devices differ in their details: they use either silicon wet or dry etching, \(^{36,37}\) a spacer silicon oxide layer, \(^{38}\) a sacrificial core \(^{39}\) or two steps silicon oxidation. \(^{40}\) van Delft et al. have even included mirrors within devices, acting as a Fabry–Perot interferometer, for contrast enhancement. \(^{41,42}\)

There is no full consensus on the measured filling kinetics, but most studies report a semi-quantitative agreement with Washburn law: time progression of the meniscus follows a square-root law, but with a slowdown with respect to the macroscopic prediction. \(^{15,33-35,43-45}\) Fig. 1 gives an overview of the reported measurements. They are expressed as the ratio of the experimental measured factor \( A_m \) to the predicted one \( A \) in eqn (1), as a function of the channel depth \( d \). Dispersion of data is quite significant but most results deviate from the macroscopic prediction for depths below about 100 nm. These filling speeds that are significantly lower than classical continuum theory are observed for all the liquids investigated. The slowdown factor increases with size reduction, up to about 50% for tens of nanometers depth.

A number of specific studies and possible physical explanations have been proposed; they are briefly summarized below.

\( \dagger \) Nanotubes or nanopores are closer to nanochannels with aspect ratio of order unity; flows inside such structures are still highly debated, and their study goes beyond the scope of this paper.

The possible slowdown effect of dynamic contact angle has been suggested as a reduced capillary ‘motor’ in the process. \(^{38,42}\) However, a few arguments seem to disprove this effect. First, the governing parameter is the capillary number that compares viscosity to capillary stress: \( Ca = \mu_i U h/t \), where \( U \) is a typical velocity. For nanochannels, spontaneous imbibition, \( Ca \) can be written from eqn (1): \( Ca = a/6h(t) \). From usual models and measurements, \(^{44}\) a dynamic contact angle differing from the static one is usually obtained for a minimum capillary number \( Ca \sim 10^{-4} \), which holds only for distances lower than about 200 \( \mu \)m in a 20 nm deep nanoslit. It suggests that a hypothetical deviation due to this effect would only impact the very beginning of filling. Similarly, since dynamic contact angle depends on velocity, which strongly varies during capillary imbibition, the square root progression, experimentally observed by most groups, should not hold. Furthermore, at a given position after the meniscus \( h_0 \), \( Ca \) decreases when the depth of the channel \( a \) is reduced, so a dynamic contact angle slowdown effect should decrease with narrower channels.

Oh et al. have used water–ethanol mixtures with different proportions, for 6 nm and 16 nm deep channels. \(^{45}\) They conclude that specific adsorption or chemical selectivity effects are not relevant, and show that a simple additional resistance due to roughness is not sufficient to account for the experimental dynamics.

Haneveld et al. have worked on dimensions down to about 5 nm; they have realized a very accurate determination of channel’s depth, and measure effects notably smaller than most other studies. \(^{37}\) They propose, as a possible explanation of slowdown, the presence of immobilized liquid layers (with dimensions comparable to the molecular size). It can be related to the layering of the first liquid molecules, observed by surface force apparatus, for a variety of solvents. \(^{46}\)

Electroviscous effect, that can be viewed as an electroosmosis in the flow potential (which itself originates from the charged diffuse layer close to surfaces immersed in an electrolyte), has also been proposed to account for an enhanced hydrodynamic resistance. This point has been tackled both by experiments, with
studies on influence of the salt concentration on the meniscus dynamics, and by successive theoretical analyses. The effect is maximized when the depth of the channel is comparable to the screening length (extension of the diffuse layer). Nevertheless, models show that on a quantitative point of view, electroviscous slow-down is too low to explain the experimental results.

We can also note that the enormous negative pressure inside devices can deform the channels by elastocapillarity. The dynamics resulting from this fluid–structure coupling have been studied in detail by van Honschoten et al., and lead to an overall accelerated dynamics.

The transport by precursor films that can develop ahead of the meniscus, as evidenced in the simulations by Chibbaro et al., can also be a source of altered imbibition. It could, for instance, suck a part of the liquid, as suggested by Haneveld et al., to account for their experiments; this effect has been studied by a lattice-gas model. It can also modify the effective boundary condition, as shown by a numerical study.

An important point is that, even though many of the previously cited experimental studies have observed the formation of enclosed gas bubbles during the filling process, very few have assessed the influence of this phenomenon on the filling kinetics. Han et al. observed that for an ethanol–water mixture, air trapping disappears for the ethanol fraction above 40%, for 27 nm deep nanochannels. To our knowledge, the only published work on the influence of those bubbles on meniscus dynamics is by Thamdrup et al.; they measure how single bubble trapping leads to an instantaneous jump of the meniscus, but corresponds to an immediate decrease in the filling speed.

Phan et al. have measured fillings kinetics within dead-end nanochannels, a geometry which amplifies the role of the gaseous phase. Experimental results agree with a macroscopic hydrodynamic model accounting for gas compression and dissolution. However, the measured slowdown in the kinetics (about 50% for 45 nm deep channels) is slightly stronger than for identical open devices.

This collection of studies shows that we are still lacking a complete description of phenomena occurring during nanochannels spontaneous imbibition. In particular, the role of gas and liquid/gas/wall interfaces appears to be complex and has only hardly been investigated until now. In this paper, we present experimental results on capillary filling, with special focus on the trapping of bubbles, their dynamics, and their influence on the imbibition kinetics. We also discuss the role of gas present ahead of the meniscus: we develop a theoretical model accounting for gas compression during filling, demonstrating a significant pressurization. We take this effect into account to determine the meniscus dynamics, and extend this model to a situation with fully developed wetting films.

**B. Filling kinetics experiments**

**Materials and methods**

Nanochannels are realized by standard clean-room fabrication, as described in our previous work. Lateral dimension of the channels are designed on a photomask: width is varied between 3 μm and 10 μm, and length between 200 μm and 5 mm, as shown in Fig. 2(a) and (c). Nanochannels are etched in silicon by reactive ion etching in an ICP omega 201 machine, with conditions adapted to slow smooth etching: CF$_4$ at 10 mtorr pressure, with a flow rate equal to 20 sccm, coil power 50 W and bias power 25 W at 13.56 MHz. The etching rate is around 10 nm min$^{-1}$ and the obtained peak-to-peak roughness, measured by Atomic Force Microscopy (AFM), is below 1 nm on a 2.5 μm line (see Fig. 2(a)). Microchannels (10 μm deep, 200 μm wide) are also dry etched; they provide a fast and reproducible arrival of the liquid during the capillary filling experiments. Holes are drilled by sand blasting at microchannels ends.

Sealing of the silicon wafer to a glass substrate is ensured by anodic bonding after piranha cleaning (5 min in 50% H$_2$O$_2$/50% H$_2$SO$_4$ followed by 20 min rinsing in deionized water) and 30 min dehydration in an oven at 200°C. Bonding parameters (temperature $T = 370$ °C, voltage $V = 250$ V, limiting current 4 mA) are chosen to minimize channel collapse, as previously reported for very low aspect ratio devices. Glass roughness is a bit higher than that of the silicon, but remains of the order of 1 nanometer on a few microns (see Fig. 2(a)). Contact angles are measured just before the bonding step by drop deposition method (Digidrop apparatus). Full wetting is obtained for the different liquids, so the contact angle is supposed to be lower than 5°, the minimum angle that can reasonably be measured by this method. Channel depth is varied between 20 nm and 400 nm.
It is measured by a calibrated AFM. The values are consistent with the ones obtained on test zones of width much larger than those of the nanochannels by a mechanical profilometer and an optical interferometry profilometer, with a maximum deviation of a few percent.

Filling kinetics experiments are realized on an inverted microscope (Zeiss Axio Observer D1), with 10X objective, and a high speed camera (Fastcam SA3). Imbibition is characterized for deionized water (clean-room quality, 18 MΩ cm), ethanol, isopropanol (electronic grade), and silicone oil (Siliconöl M5, Carl Roth). Standard reflection illumination with a halogen lamp gives a good contrast between dry and wetted zone for depths down to 20 nm. Devices are stored in an oven at 200 °C for 30 min just before the experiments. The chip is inserted in a Teflon holder and liquid arrival is realized by deposition of a 5 µL droplet. The temperature is measured for each experiment; its value is in the range 21–26 °C. Relative hygrometry is around 45%.

Filling speed results

A typical snapshot of filling experiments is reported in the inset of Fig. 3(a), obtained for water in a 66.5 nm deep channel.

We observe trapping of bubbles at the liquid front; this point will be discussed in detail in the next section. Meniscus position is extracted by image analysis. The method we found to allow the best precision is to average image intensity on the channel width and threshold the obtained profile (filled part appears darker than the non-wetted zone); it enables us to determine the first arrival of liquid. Fig. 3(a) shows a typical result, also obtained for water in a 66.5 nm deep channel. Square of the meniscus position is reported versus time, together with prediction from Washburn law (eqn (1)). As reported in other studies, we measured a semi-qualitative agreement with the macroscopic prediction: the meniscus follows a square root law, but with a slightly reduced kinetics. The relative slowdown shows no dependence on channel width (from 3 to 10 µm) and length (from 200 µm to 5 mm), within a few percent. The typical reproducibility observed in our experiments is of the order of 10% for the value of A (defined in eqn (1)). This is consistent with estimation of the error bar originating from uncertainty on experimental parameters (channel depths, contact angle, image analysis, temperature and its influence on the viscosity).

Fig. 3(a) Typical filling kinetics, shown in the case of deionized water in a 66.5 nm deep nanochannel. Continuous line is the macroscopic Washburn prediction. Measured value of \(A_w\) is 20% lower than the theoretical one. (b) Superposition of our data on the kinetics data extracted from the literature.\(^{31,37-40,125}\)

Fig. 3(b) summarizes our determination of the ratio experimental/predicted coefficient \(A\) as function of the channel depth. Similarly to most other works, the slowdown effect we measure increases with confinement, up to around 25% deviation for 20 nm nanoslits for all the liquids tested.\(^{†}\)

Since this dynamics is hypothesized to be related to the presence of observed bubbles, we describe in the following section their trapping and time evolution.

C. Behavior of trapped bubbles

Threshold for bubble trapping

Fig. 4 shows the typical appearance of the phase distribution behind the meniscus, when it is located at a distance of order 750 µm from the nanochannel entrance, for three different liquids (water, ethanol and silicone oil), and four depths from 162 nm down to 42 nm.

First, a threshold depth \(a_t\) is clearly visible below which bubbles are trapped for the three liquids. Secondly, the value of \(a_t\) depends on the liquid: when reducing channel depth, bubbles are first trapped for water (85.7 < \(a_t\) < 162 nm), then for ethanol (66.5 < \(a_t\) < 85.7 nm), and eventually for silicone oil with very few bubbles observed only in the narrowest channels (42 nm deep). Note that 85.7 nm is probably very close to the trapping threshold for ethanol: it can be seen on the corresponding zoom on the spatiotemporal diagram of Fig. 5. Images for \(a = 20\) nm are not shown because of the poor contrast between liquid and gas at this nanoslit depth. Nevertheless, after careful analysis of the images, bubbles are observed with all the liquids for \(a = 20\) nm.

The precise physical mechanism responsible for bubble trapping is still unclear, but our observations suggest that it could be

Fig. 4 Snapshots taken during capillary filling of water, ethanol and one silicone oil for several nanoslit depths. For all the images, the distance between liquid front and the nanoslit entrance is \(\sim 750\) µm. Liquid appears as dark and gas (air) as white (flow from left to right).

\(†\) We have investigated different fluids: water (which is an electrolyte), ethanol and isopropanol (also polar but organic liquids) and a silicone oil (made of small polydimethylsiloxane chains). Even though they do not constitute an exhaustive set, they represent quite different systems concerning molecular liquid–liquid, liquid–gas, and liquid–surface interactions. Similar slowdown magnitude is observed for all liquids tested. It may thus be argued that specific surface effects seem to play a minor role, even though being able to confirm this point would require a dedicated study. In this work, we focus on effects which seem to be more generic than liquid–surface interactions to explain the experimentally measured slowdowns: trapped bubbles and gas pressurization.
related to contact line pinning by chemical or topological defects. As a matter of fact, even though we measure a zero contact angle just before bonding the devices at a macroscopic scale, wetting on silicon oxide surface (glass or silicon native oxide) is better for silicone oil (the spreading parameter is higher) than for ethanol and for water, so that water (and to a lower extent ethanol) should be more subject to pinning than the oil. The fact that such pinning has a stronger effect (and leads to the trapping of bubbles) when the confinement increases is qualitatively in agreement with fluid mechanics studies in macroscopic Hele-Shaw cells with chemical or topological heterogeneities.

Let us also note that, even in the absence of bubbles, pinning effects could contribute to the slowdown, and might even be interpreted as a dynamic contact angle (but with a very different scaling with velocity).

As can be seen in Fig. 4, bubble density is higher just behind the meniscus than close to channel entrance. A first explanation is that trapping occurs more sparsely at the beginning of the filling process, when the meniscus is moving faster. We believe this is related to a contact line pinning that is more significant for low velocities, as seen in the already cited Hele-Shaw cell studies. Nevertheless, bubble time evolution is probably the main cause for this spatial distribution; and we discuss their dynamics hereafter.

**Bubble dynamics**

For the cases investigated here, once a bubble is trapped it is not convected by the liquid flow but it stays at its formation place. This appears in spatiotemporal diagrams of a single nanoslit, see Fig. 5. These spatiotemporal diagrams consist of successive snapshots of one nanochannel which are stacked from the top (initial instant - empty nanochannel) to the bottom (filled nanochannel). Snapshots are separated in time by a constant time interval. Here, we focus on bubbles' temporal evolution. Bubbles created lastly have longer lifetime, as can be particularly well seen in Fig. 5 for water in \( a = 85.7 \) nm and ethanol in \( a = 66.5 \) nm. Furthermore, the lifetime seems to depend on the initial bubble size (small bubbles disappear more rapidly).

We can expect that, as analyzed in ref. 57, bubble volume decrease is due to both compression and dissolution of the air in the liquid. Compression is due to the increase in the liquid pressure as the distance between the front and the bubble location increases, see Fig. 6.

The time evolution of a single bubble volume is estimated from the images. Spatiotemporal diagrams are first binarized in order to be able to 'count' white pixels corresponding to the projected area of the highly crushed bubble chosen (liquid-gas interface can be reasonably considered as straight here). Multiplying this area by the depth \( a \) provides an estimate of the bubble volume, \( V_b \), accurate enough for the need of this study (uncertainty on \( V_b \) is quite high here due to the optical resolution along the slits widths). Several measured bubble volume temporal evolution at locations \( x_b \) are plotted as a function of time in Fig. 7. Measurement are presented for \( a = 85.7 \) nm for water (Fig. 7(a)), and \( a = 66.5 \) nm for ethanol and silicone oil (Fig. 7(b) and (c)), depths for which our experimental data give access to the full bubble lifetime. To estimate the bubbles compression effect, time evolution of the initial bubble volume trapped, \( V_{b0} \), is computed assuming an isothermal compression:
Fig. 6 Illustration of the increase in the liquid pressure and in gas bubble as the distance between the front and the bubble location increases. The dashed line represents liquid pressure profile at the instant when the bubble is trapped (meniscus at \( h = x_b \)). The continuous line represents liquid pressure profile at a later time.

\[
V_b = \frac{p_b}{\rho_b} V_{0b},
\]

where \( p_b \) is the pressure in the bubble. It depends on the local liquid pressure \( p(x_b) \) through Laplace's law:

\[
p_b = p(x_b) + \frac{2\gamma}{a} \cos \theta.
\]

Assuming a linear liquid pressure profile between \( x = 0 \) and \( x = h(t) \) leads to:

\[
p_b = p_a + \frac{(h(t) - x_b)}{a} \frac{12\mu_p \frac{dh}{dt}}{a^2}.
\]

Computations of \( V_b \) are performed using values of \( h(t) \) and \( \frac{dh}{dt} \) computed from the measured values of the coefficient \( A_m \) and \( h(t) = A_m t \). Time evolutions of \( V_b \) computed from this compression model, are shown in Fig. 7.

As can be seen from Fig. 7(a), \( V_b \) computed from isothermal gas bubble compression coincides well with experimental measurements for the case of water, except for \( x_b = 290 \mu m \) when \( t > 0.5 \) s. At long times, compression model predicts a positive value for \( V_b \) (limited maximal compression ratio) and then cannot predict the observed bubble disappearance. Indeed, this final collapse is due to gas dissolution in water. Two main processes can then be distinguished for the bubble volume decrease: compression followed by dissolution. Dissolution begins to act efficiently once the dissolution flux \( J \) (kg m\(^{-2}\) s\(^{-1}\)) is sufficiently large; it can be expressed as:

\[
J = \frac{D}{\delta} (c - 0),
\]

where \( D \) is the diffusion coefficient of the gas in the liquid, \( \delta \) is a certain transfer length scale and \( c \) is the gas concentration in the liquid at the liquid–gas interface. Gas concentration \( c \) depends on \( p_b \) through Henry's law: \( c = \frac{p_b}{k_{H}} \), where \( k_{H} \) is the Henry's constant, which depends on the liquid and gas under consideration. Eqn (5) becomes:

\[
J = \frac{D p_b}{\delta k_{H}}.
\]

According to eqn (6), \( J \) is proportional to \( D/\delta \) which will be assumed constant here. This hypothesis is justified because much higher variations are expected for \( p_b/k_{H} \) (\( p_b \) starts from \( p_a \) and
tends to \( p_b + 2 \gamma/a \) than for \( D\alpha \). Eqn (6) indicates that the dissolution flux is proportional to \( p_b \). Dissolution is then continuously enhanced during the filling process. In the following, we focus on the transition from compression-dominated to dissolution-dominated bubble evolution. It is characterized by measuring the gas bubble pressure \( p_{bd} \) from which experimental point \( V_b(t) \) deviates from the compression model in Fig. 7. Then, \( p_{bd} \) can be simply deduced graphically from a bubble volume ratio computation (see eqn (2)): \( p_{bd} = p_b(V_{bd}/V_b) \). As can be seen from the curve corresponding to \( x_b = 290 \mu m \), in Fig. 7(a), \( V_b \) decreases by a factor of about 10 before diverging from the compression model, leading to \( p_{bd} = 10 p_b \). The curve corresponding to \( x_b = 788 \mu m \) is entirely described by the compression model. We can then verify that the dissolution pressure \( p_{bd} \) is not reached in this case. Indeed a maximal value of \( 5p_b \) for \( p_b \) can be easily deduced from this plot. Therefore, for the latter case, bubble volume decrease is mainly due to mechanical compression since the dissolution process is not expected to be significant as long as \( p_b \) is lower than the dissolution pressure \( p_{bd} = 10 p_b \), previously deduced. The coherence of this analysis allows validating the proposed phenomenology to explain bubbles collapse: bubbles are first mechanically compressed due to the increase in local liquid pressure (see Fig. 6); \( J \) (which is proportional to \( p_b \)) increases during this stage until bubble volume decrease is mainly due to dissolution when \( p_b = p_{bd} \).

Compression process for the bubble located at \( x_b = 788 \mu m \) (Fig. 7(a)) takes more time because the bubble is further from the channel entrance than the bubble located at \( x_b = 290 \mu m \) for which \( h - x_b \) increases more rapidly (see eqn (3) and Fig. 6).

In the following, we focus on the case of ethanol and silicone oil (Fig. 7(b) and (c)) for which the experimental points \( V_b(t) \) follow the compression model curves only at the beginning for the ethanol (Fig. 7(b)) and not at all for the silicone oil (Fig. 7(c)). Given the previous analysis, these behaviors suggest that bubble volume decrease by dissolution for ethanol and silicone oil starts from lower values of \( p_{bd} \) than for water. In other terms, a lower bubble compression is needed to reach the sufficiently large values of \( J \) that mark the start of the dissolution stage. This can be understood considering Henry's coefficient values. For simplicity here, air bubble is considered as pure nitrogen. The values of Henry's coefficient \( k_H \) for nitrogen are extracted from Battino et al.\(^4\) This gives \( 1.0 \times 10^7 \) Pa m\(^3\) kg\(^{-1}\) and \( 3.6 \times 10^7 \) Pa m\(^3\) kg\(^{-1}\) for water and ethanol respectively. The \( k_{H4} \) for nitrogen in the silicone oil used in our experiments has not been found. We use the value \( 5.2 \times 10^7 \) Pa m\(^3\) kg\(^{-1}\) given in the same reference for a quite similar oil (octamethylcyclotetrasiloxane). Since \( J \) is inversely proportional to \( k_{H4} \), the value of \( J \) needed to observe a significant dissolution process is reached more rapidly (or at lower \( p_b \)) for ethanol than for water (2 orders of magnitude in \( k_{H4} \)) and even faster for silicone oil (3 orders of magnitude in \( k_{H4} \) between water and oil).

Finally, this analysis shows that bubble lifetime strongly depends on the solubility of the gas in the filling liquid. First, if the gas is highly soluble, it is expected that bubble lifetime depends only a little on filling kinetics (as for silicone oil in our study). Second, for moderately soluble gas, bubble volume decrease is first due to mechanical compression during which \( p_b \) increases at a rate depending on \( x_b \) and finally reaches \( p_{bd} \) from when dissolution induces bubble final collapse. It is interesting to note that we recover the compression–dissolution process already encountered in capillary filling of closed end nanochannels.\(^{97}\)

D. Discussion: roles of gas compression on the imbibition process

Influence of trapped bubbles on kinetics

It can be first hypothesized that the presence of stagnant bubbles in the liquid generates an increase in the overall hydrodynamic resistance in the invaded region and thus a possible slowdown of the capillary filling. However, the continuous creation of bubbles also induces an increase of the liquid front velocity because the volume to fill by the liquid is lowered. It is thus surmised that the competition between these two processes acts on the filling kinetics and this is analyzed in what follows.

One way to model the viscous resistance induced by the presence of bubbles in the narrow nanoslits is to use the Brinkman approximation.\(^{62,63}\) It allows estimating the effective permeability \( K \) of a slit consisting of a square periodic array of circular plots bounded by two parallel walls, see Fig. 8.

Note that such a way to model bubble viscous resistance would fail in the presence of bubbles occupying almost the entire channel width (that would lead to highly resistant corner flows\(^66\)). Nevertheless, high magnification bubbles snapshots in Fig. 5 for \( a = 66.5 \) nm show that bubble size is significantly inferior to channel width and there are no such corner films: liquid around bubbles occupies several channel depths. Bubble induced resistance should be much lower and can be estimated as a function of their sizes and gap spacing, as discussed in the following.

The permeability of a square periodic array of cylinders of infinite length, \( K_{sp} \), can be deduced from the work of Sangani and Acrivos.\(^{68}\)

\[
K_{sp}(e) = \frac{W^2}{2f(e)},
\]

where \( W \) is the center to center distance between two plots (bubbles) and \( f(e) \) is a coefficient depending on the surface

Fig. 8 Sketch of the bubbles arrangement (gray plots) in a nanoslit for the estimation of the viscous resistance induced by their presence.
fraction of the plots (bubbles) $\epsilon = \pi r_p^2 W^2$ where $r_p$ is the plot radius (see Fig. 8). Discrete values of $f(\epsilon)$ are provided in ref. 65 and for the needs of the present study they are linearly interpolated. According to the Brinkman approximation, the effective permeability of a slit consisting of a square periodic array of plots bounded by two parallel walls with a gap spacing of $a$ can be estimated by:

$$K(\epsilon) = K_p(\epsilon) \left(1 - \frac{\tanh(Da)}{Da}\right)$$

(8)

where $Da = a/(2K_p(\epsilon))^{1/2}$. The liquid flow rate $q$ can then be linked to the pressure gradient along the slit using the Darcy’s law:

$$\frac{q}{aw} = \frac{K(\epsilon)}{\mu} \frac{d\rho}{dx}$$

(9)

We will first assume that the volume of the bubbles stays constant during the filling. Viscous resistance due to the presence of bubbles is then overestimated. A line of plots is selected in the square periodic array of plots to account for the bubbles positioning observed in the experiments (see Fig. 8). Therefore, we set $W = w$, where $w$ is the slit width, and we vary the bubble radius $r_b$. The flow rate is related to meniscus velocity taking into account that only the fraction $(1 - \epsilon)$ of total area is to fill by the liquid:

$$q = (1 - \epsilon)aw \frac{dh}{dt}$$

(10)

Assuming a pressure gradient of $2\gamma/ha$ and combining eqn (9) and (10) we obtain the Washburn’s law corrected for the presence of bubbles:

$$h(t)^2 = A \frac{12K(\epsilon)}{(1 - \epsilon)a} l = A_1 t,$$

(11)

where $A = \gamma \cos \theta/(3\mu)$ is the Washburn’s law classical prefactor (eqn (1)). In the following, the prefactor in eqn (11), $A_1$, is compared to $A$ in Fig. 9(a). Surprisingly, this modelling leads to the result that capillary filling should be faster than Washburn’s prediction for the narrowest nanoslits and slower for the less narrow ones. This is due to the counter-intuitive hydrodynamic behaviour of the very narrow nanoslits: when $a$ is very low, the effective permeability (eqn (8)) does not depend on the presence of the bubbles. This is shown in Fig. 9(b). At these very small scales the effective permeability $K(\epsilon)$ is actually very close to the Hele-Shaw permeability $K(0) = a^2/12$ as already shown by Tsay et al.\(^\text{40}\) Therefore, $A_1/A \sim 1/(1 - \epsilon)$ is necessarily higher than 1 and in this case the effect of the reduced volume to fill is more important than the viscous resistance induced by bubbles. For less narrow nanoslits, for $a > 100$ nm in Fig. 9(b), $K(\epsilon)$ is lower than the permeability of a slit free of bubbles and depends on the bubbles’ size. In this case, viscous resistance induced by bubbles is sufficiently large to overcompensate the effect of the reduced volume to fill. Nevertheless, such a slowdown is not expected in our experiments because no bubbles are observed for $a > 100$ nm.

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\(^\text{40}\) This holds except if bubbles are extremely close and almost obstruct the channel, this is the case for the plot for $2\gamma/\epsilon w = 0.95$ in Fig. 9(b). Such a bubble density is unrealistic compared to our observations and furthermore analysis should include corner flows modelling.

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**Fig. 9** (a) Ratio of Washburn prefactor corrected for the presence of bubbles to the classical Washburn prefactor as a function of the slit aperture for several bubbles sizes and a slit width of 3 μm. (b) Ratio between $K(\epsilon)$ and the permeability of a Hele-Shaw slit $K(0)$ for the same set of parameters as for (a).

For $a < 100$ nm, $K(\epsilon)$ should then always be $= K(0)$ and bubbles formation should lead to a faster filling kinetics.

The effect of the reduced volume to fill is overestimated in this analysis because, as it has been shown above, bubbles collapse after a certain time. Therefore, depending on the bubbles lifetimes, the effect of the reduced volume to fill is partially compensated by the bubbles collapses. If bubbles lifetime is very short, the overall effect should be null and then the filling kinetics should be the Washburn’s one. Capillary filling experiments realized with silicone oil are close to this situation (air bubbles dissolve rapidly in silicone oil) but we still measure a filling slower than the Washburn’s one. If a bubble lifetime is long, the filling kinetics is then expected to be faster but here again we measure a filling slower than the Washburn’s one, as it is the case for experiments with water.

Finally, these elements suggest that capillary filling kinetics slowdown at nanoscale cannot be explained by additional viscous resistance due to the bubbles. It is then expected to be due to other processes. The next section focuses on the possible but so far unexplored role of gas compressibility during the imbibition.

**Influence of gas pressurization ahead of the invasion front**

One possible explanation of the slower invasion observed in nanochannels might lie in the effect of the gas flow in the part of the channel occupied by the gas phase. This issue was addressed in Hultmark et al.\(^\text{25}\) where it was shown that the viscous resistance associated with the gas flow could indeed explain the deviation from classical theory. However, the gas compressibility was not taken into account in Hultmark et al.\(^\text{25}\) as a result the effect of the gas phase was significant for extremely long systems only. In contrast with Hultmark et al.\(^\text{25}\) the gas compressibility is taken into account in what follows and it is shown that a significant pressurization of the gas occurs during the imbibition process. The effect of the pressurization effect on the imbibition dynamics is then discussed. Before, one can give a simple argument explaining why the gas pressurization effect should be incorporated in the analysis of the gas pressure in the nanoslit. As shown below, the gas pressure is governed by a diffusion equation whose diffusion coefficient is given by $D_g = \mu_g/\epsilon w^2$ where $\mu_g$ is the gas dynamic viscosity, $a$ the slit aperture and $\rho_g$ is the gas
pressure. As a result, a characteristic time of the pressure
"diffusion" is $t_p = L^2/D_p$, where $L$ is the channel length. With $L = 1$
mm, $a = 20$ nm and $p_g = 1$ bar for example, this gives $t_p \approx 300$ s, which is much greater than the imbibition time ($\approx 2$ s). As
a result, quasi-steady pressure profiles cannot be assumed in the
gas phase and the transient effect due to the gas compressibility
should be taken into account.

The situation under study is sketched in Fig. 10. The equation
governing the pressure in the gas phase is obtained by combining the
gas mass balance equation,

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

(12)

the momentum balance equation, which reduces to the lubrica-
tion approximation with only one non-zero average velocity
component, the one in the direction $x$ of the invasion

$$u_x = -\frac{a_g^2 \frac{\partial p_g}{\partial x}}{12 \mu_g}$$

(13)

where $a_g$ is the channel aperture in the gas phase (this consider-
ation of the aperture for the gas phase as possibly different from
the aperture of the channel will be made clear later), the ideal gas
law

$$p_g = \frac{M}{R T} (p_a + p_g)$$

(14)

where $R$ is the ideal gas constant, $T$ is the temperature, $p_a + p_g$ is
the total gas pressure; $p_g$ is the pressure at the exit of the nano-
channel (thus the atmospheric pressure) and $p_a$ is the deviation
from the atmospheric pressure.

This leads to the following non-linear transient diffusion
equation,

$$\frac{\partial p_g}{\partial t} = \frac{\partial}{\partial x} \left( \left(p_g + p_a\right) \frac{a_g^2 \frac{\partial p_g}{\partial x}}{12 \mu_g} \right)$$

(15)

which must be solved subject to the following boundary
conditions

$$p_g = 0 \text{ at } x = L$$

(16)

$$\frac{\partial h}{\partial t} = -\frac{a_g^2 \frac{\partial p_g}{\partial x}}{12 \mu_g} \text{ at } x = h(t)$$

(17)

and the initial condition $p_g = 0$ everywhere in the slit at $t = 0$.

The imbibition dynamics, i.e. $h(t)$, depends in turn on the gas
pressure at the imbibition front. Taking into account the inertial
effects occurring at the very beginning of the process so as to
avoid the velocity singularity at $t = 0$ of the Washburn model, the
equation governing the evolution of the front position $h$ (see
Fig. 10) can be expressed as (details of the derivation will be given
elsewhere):

$$\frac{d^2 h^2}{dt^2} = \frac{1}{\ell_0} \left[ \frac{h_0^2}{\ell_0} \cos \theta \left( \frac{a_1}{a_m} - \frac{p_g(t)}{p_o} \right) - \frac{dh^2}{dt} \right]$$

(18)

where:

$$p_o = \frac{\gamma \cos \theta}{a_1}; \quad \ell_0 = \frac{\rho a_1^2}{12 \mu_o}; \quad h_0 = \frac{\sqrt{\rho_0 \gamma a_1^3}}{6 \mu_0}$$

(19)

$\gamma$ is the surface tension, $\theta$ the contact angle ($\theta = 0$ throughout
the article), $\mu_o$ is the liquid dynamic viscosity and $a_1$ is the channel
aperture in the invaded region ($a_1 = a$ where $a$ is the aperture of
the slit). Note that as sketched in Fig. 10, we have the possibility
to consider a radius of curvature of the meniscus different from
the classical value $a/2\cos \theta$. This radius is denoted by $a/2$. The
initial condition associated with eqn (18) is $h = 0$ at $t = 0$.

From the above equations, it is clear that the imbibition
dynamics and the evolution of the pressure in the gas phase are
coupled. Furthermore, it can be noted that the pressure diffusion
equation, eqn (15), is non-linear and should be solved over a
shrinking domain as the front moves into the slit. It seems
difficult to avoid a numerical approach to the above problem.
The details of the method of solution will be presented
elsewhere and can be summarized as follows. To solve eqn (15)–
(17) governing the gas pressure, we first introduce a new coor-
dinate system $X/ = 1 - x/L)(1 - h(t)/L), t = t$ allowing us to
express the problem in a computational domain constant in size.

The trick is the same as the one used in Camassel et al.46 where
the interested reader can find further details. Then the equation
in the fixed domain is solved numerically using the commercial
simulation software COMSOL Multiphysics. Eqn (18) is solved
using a variable order solver based on the numerical differen-
tiation formulas (NDFs) available with Matlab. To obtain a fully
converged solution, one must iterate between the solution for the
gas pressure and the solution of eqn (18) governing the front
dynamics. For the present article, we have used a simplified
procedure in three steps. The first step consists in solving eqn (18)
with $p_g(t) = 0$, which thus gives the classical invasion dynamics.
In the second step, we have computed the gas pressure $p_g(t)$
assuming the classical invasion dynamics, i.e. using $h(t)$ as
computed in step #1. The third step consists in solving again
eqn (18) using the solution $p_g(t)$ obtained in step #2. This
simplified procedure is sufficient to get insight into the possible
impact of gas pressurization on the imbibition dynamics.

Furthermore, we have distinguished two main situations
depending on the possible presence of precursor liquid films
along the slit walls in advance of the bulk meniscus. We begin
with the simplest case where the walls ahead of the front are dry.

Case #1: dry walls in the gas region. In this case, the problem is
solved assuming $a_1 = a_m = a$. Examples of results for pure
water as working fluid are presented in Fig. 11.

Two remarkable results emerge from Fig. 11. First the pres-
surization of the gas phase is quite significant and reaches an
absolute value as high as 2.5 bar (1.5 bar above the exit pressure)
in the 20 nm slit for example. Second, this pressurization effect at
the meniscus lasts over a significant period of the imbibition
process. As can be seen from Fig. 11, the pressurization dynamics
is characterized by a plateau, whose relative duration increases as
the slit aperture decreases, followed by a period of relatively slow
pressure decay as the front further advances into the slit. The

![Fig. 10 Sketch of imbibition process in a slit.](image_url)
of the front. The numerical procedure is the same as for the dry walls except that the gas section is now reduced owing to the presence of the films. The working fluid is water as before. As a result of the reduced gas section, the gas pressurization effect increases as the liquid film thickness increases. As illustrated in Fig. 12, the effect is sufficient to slow down the imbibition dynamics significantly. For example, the case $a_{	ext{f}}/a_{0} = 0.25$ in Fig. 12 corresponds to films of thickness 7.5 nm along each wall of the slit.

Owing to the simplifications made in the modelling (no effect of films on capillary pressure jump, dynamics of film precursors ignored—they are supposed to be present at all times—possible effect of gas pressurization on film thickness ignored, possible slip effect at the film surfaces, etc.), further work is needed to assess the effect of precursor films on a more firm basis.

Note also that rarefaction effects in the gas phase were neglected on the ground that a significant pressurization was expected. However, estimates of Knudsen number for the pressures obtained in our simulations indicate that the rarefaction effect, which increases the apparent conductance of the gas flow and thus tends to diminish the pressurization effect, is not necessarily negligible. Taking into account the rarefaction effect will not change our results qualitatively, but will reduce the pressurization effect.

Nevertheless, the simple computation presented here offers a possible explanation to the slowdown in the imbibition dynamics observed in the experiments.

**E. Conclusions**

We discussed the phenomena arising during the capillary filling of narrow nanoslits, with gap of a few tens of nanometers. After a short literature survey, we described how experiments on capillary filling of microfabricated 20 to 400 nm deep nanoslits shed light on processes involved in nanoscale flows. As reported previously by several authors, experiments show a slowdown compared to macroscopic prediction and the deviation increases with confinement. We presented possible origins of this phenomenon. In particular we discussed in detail the role of gas compressibility effect cannot be neglected in nanoslits. Bubble formation, collapse and their effect on viscous resistance was analysed experimentally for three different liquids: water, ethanol and silicone oil. Nanoslit aperture threshold, below which bubbles appear, was found to depend on the liquid used. The bubble formation mechanism is still unclear but seems to be related to contact line pinning by unavoidable nanoscopic heterogeneities, since the measured aperture threshold decreases for highly wetting liquids. Further work is planned for a more accurate description of this effect. Bubble collapse was shown to follow two stages: first, the bubble is compressed due to increase of the local liquid pressure that induces increase of gas bubble pressure; second, for sufficiently high gas bubble pressure,
dissolution leads to final bubble collapse. A simple model qualitatively explained why bubble lifetime strongly depends on the gas solubility in the liquid, as it is observed in experiments. The intuitive effect of viscous resistance induced by the presence of bubbles was estimated using an effective medium approach. Actually, this analysis showed that bubbles should not induce significant viscous resistance for nanoslit aperture <100 nm. This counter-intuitive purely hydrodynamic effect comes from the fact that viscous resistance is mainly due to the very low nanometric slit gap which dominates over any assembly of obstacles spaced by a micrometric distances in the slit. Therefore, nanoslit capillary filling kinetics slowdown does not seem to be due to the presence of bubbles in the filled part of the channel. However, the effect of successive contact line pinnings during the meniscus walk on the overall kinetics is not obvious and should be considered as a candidate for explaining the slowdown by a decrease of the imbibition driving force. Bubble formation seems to be controlled by contact line pinning, so that even though they do not increase the viscous resistance, their presence reveals meniscus pinning which may cause the slowdown.

Another effect that contributes to slow down the imbibition dynamics is related to the gas compressibility. We showed from a numerical model coupling the imbibition dynamics with the evolution of gas pressure in the nanoslit that the imbibition process is characterized by a significant pressurization of the gas phase. This effect was shown to last over most of the imbibition but is more marked in a first phase that can represent almost half of the total imbibition duration for the narrowest slit tested. However, this effect is sufficient to slow down the imbibition dynamics significantly only in the presence of prewetting liquid films, reducing the gas section in the region ahead of the imbibition front.

Those effect show that capillarity at small scale can be complex and follows quite counter-intuitive rules, as recently demonstrated by Willmott et al. who showed the spontaneous filling of hydrophobic micropipettes by water droplets.89

In spite of the insights brought by the present paper, the full understanding of capillary filling of nanoslits remains a subject rich in open questions such as finding the reason why bubbles are trapped; studying the influence of roughness (a recent study shows the interplay between shape and roughness for meniscus at small scales89); assessing the role of gas solubility.

Experiments on forced imbibition (with an external additional driving force) could be a clue to discriminate between possible mechanisms.

Other related situation are also interesting, such as filling in the presence of surfactants,71 or the case of channels with two dimensions in the nanometer range, closely related to reduced friction recently observed in carbon nanotubes.4,65

Nano-fluidics offers numerous other situations where novel effects can be expected, in particular when the channel dimensions become comparable with typical length scale: (i) molecular size for complex fluids (polymer melt or semi-dilute solutions),72 as well as individual molecules – see works on DNA elongation and sorting, first applications of nano-fluidics73,90; (ii) mean free path for gases (rarefaction effect);48 (iii) Debye screening length for water solutions, potentially enabling electrical charge-based filtering, sorting, concentrating, energy conversion.1,12,77,78

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Notes and references
