Thin film metal sensors in fusion bonded glass chips for high-pressure microfluidics

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Abstract

High-pressure microfluidics offers fast analyses of thermodynamic parameters for compressed process solvents. However, microfluidic platforms handling highly compressible supercritical CO₂ are difficult to control, and on-chip sensing would offer added control of the devices. Therefore, there is a need to integrate sensors into highly pressure tolerant glass chips. In this paper, thin film Pt sensors were embedded in shallow etched trenches in a glass wafer that was bonded with another glass wafer having microfluidic channels. The devices having sensors integrated into the flow channels sustained pressures up to 220 bar, typical for the operation of supercritical CO₂. No leakage from the devices could be found. Integrated temperature sensors were capable of measuring local decompression cooling effects and integrated calorimetric sensors measured flow velocities over the range 0.5–13.8 mm/s. By this, a better control of high-pressure microfluidic platforms has been achieved.
1. Introduction

Having a viscosity comparable to gases and a liquid-like density, supercritical CO$_2$ is a benign, non-toxic and compressible solvent for several types of organic compounds. It is used for a multitude of applications, e.g., high-value products in the food industry [1], purification and analytics in the pharmaceutical industry [2] and, particle formation and loading of pharmaceutical compounds to carrier substrates [3-5].

Use of the fluid requires knowledge of multicomponent thermodynamic parameters like solubility. Conventional methods [6] used to determine high-pressure phase equilibriums are generally labour intensive, time-consuming and demand special safety precaution. Therefore, high-pressure microfluidics has emerged as a potentially fast and efficient alternative for the screening of such parameters [7]. But to increase its applicability, better control is needed of the highly variable fluids inside the chips. In this paper, the integration of sensor elements into high-pressure microfluidic channels is explored for applications with supercritical CO$_2$.

Thin film sensors and electrodes in microfluidic chips offer a large degree of added functionality. For low-pressure applications, this has been exemplified by systems for detection and control, e.g., systems that are impedance based [8] or electrochemical [9, 10, 12], and as microplasma sources [11]. With electrodes in the chips having direct access to the fluids, systems can have both a small footprint and a low dead volume.

Several research groups have demonstrated glass chips with integrated metal thin films [10-11, 13-15]. One fabrication method that we have followed involves etching a trench for the metal layer followed by a heat treatment at 625 °C [14]. However, such systems have only been demonstrated at low pressures. As one part of this paper, we investigate the conditions necessary to fulfill for chips with embedded Pt thin films to withstand the pressures needed for supercritical CO$_2$.

A supercritical fluid shows large variations in a number of physical qualities such as density, viscosity, thermal conductivity, and heat capacity, around its critical point, which for CO$_2$ is at 31.1 °C and 73.9 bar (7.39 MPa) [16]. The density must be controlled, as it is a key parameter affecting other properties, like the solubility of compounds. Further, CO$_2$ is a compressible fluid that can, during decompression, be subject to cooling by both the Joule-Thompson effect and by liquid-vapor phase transformations. For microfluidic platforms having flow restrictors and complex designs; CO$_2$ will be in a non-equilibrated state. Also, at near-critical conditions, a heterogeneous gas-liquid flow may add to the complexity [17-18]. To control such a fluid systems, it is necessary to have precise control of the temperature. The large surface to volume ratio in microchips allows for good heat transfer and common approaches for external temperature control includes having chips mounted on heated Cu blocks [19], immersed in oil [20] or mounted on resistive heat stages [21]. For all such systems, the temperature is measured externally and globally. Consequently, by adding local sensing capability, much-needed insight regarding the actual state of the fluid in the chip can be given.

This paper presents two high-pressure chips. To expose the cooling effects connected to decompression, a pressure drop chip with temperature sensors is used. The other chip is a calorimetric flow sensor. This sensor principle has been widely explored in the field of MEMS and works by measuring the temperature increase of a fluid as it passes over a heater [22-25]. While the response of such sensor principle is primarily dependent on flow velocity, other factors, including the fluid density and heat capacity, also affect the response [26]. By these two chips, it is explored how internal integrated temperature and flow velocity sensing can be achieved in
transparent microfluidic chips at the high pressures required for supercritical CO₂.

2. Background

Metal layers in bond interfaces

Models have been developed to describe how wafer bonding is influenced by surface variations [27]. With protrusions, gaps will form, figure 1. By such a model, wafers will bond, and conform to the gap, if the elastic strain energy required is less than the lowering in surface energy as the wafers adhere [28]. With small enough particles, elastomechanical instability allows for embedment, resulting in a gap of length 2R, around the particle that will be in the same order of length as the diameter, 2h, of the particle, figure 1 a. However, as soon as the particle is larger than a critical size, the gap size will be in the order of the thickness of the wafer. Specifically, when embedded in-between two 1.1 mm thick borosilicate glass wafers, with Young’s modulus of 64 GPa, Poisson’s ratio of 0.2 [29], and a surface energy in the range of 80 to 160 mJ/m², a rigid particle has a critical height in the range of 0.18 to 0.26 µm (following Eq. 6 in [27]). However, as soon as the particle is larger than its critical value, the gap radius will grow to 3 mm, or more (combining Eqs. 5 and 6 in [27]).

![Figure 1](image-url)

Figure 1 Schematic cross-sections: In a, a particle in the bond interface produces a gap. In b, a flat metal layer deposited on top of a wafer in the bond interface produces a similar gap. In c, the metal layer is instead deposited in an etched trench.

For a flat metal layer on one of the wafers, the critical height of the film will be lower, figure 1 b. Due to the sinusoidal-shaped gap, the metal layer thickness, d, needs to be less than 2h in order to give the same gap size. The metal layer width, w, will determine the contact points, and hence the d/h ratio. By forming an etched trench and lowering the metal layers into the substrate, the problem can be evaded, figure 1 c. This has been a successful approach for low-pressure applications [10-11, 13-14]. Gaps, u, formed along the edges by using such methods are then dependent on the dimensional mismatch between the trench and the metal, giving rise to possible voids around the metal. If these voids can be kept small, a high-pressure endurance of the chips should be possible; also when having many interconnects on a single chip.

The pressure tolerance of chips is dependent on several factors, including the geometry and wafer bonding scheme [30, 31]. For semi-circular channels, the largest stresses are generally seen along the bond interface. By modeling pressurized channels using finite element analysis, the stress along the channel can be calculated from the fracture pressure, and a comparable, geometry
independent measurement of strength can be made.

Flow of CO₂ in microfluidics

CO₂ is highly compressible in the region around the critical point and the fluid flow to and from the microfluidic chip depend on several variables. Given an equilibrated system as the experimental setup sketched in figure 2, the pressure difference, caused by a change in volume between two pistons, is driving the flow.

Figure 2 Experimental setup for chip characterization showing the forward moving piston, the chip, and the backpressure piston. System parameters at these three positions are shown. Two capillaries (orange) of 10 cm connect the chip to stainless steel tubing (gray) having the inner diameter given in µm. Cooling (light blue) is applied to the forward piston and the tubing before the chip.

Keeping the pressure difference between the two pistons small allows the system to be close to equilibrium. A decrease in volume of the forward moving piston corresponds to a volumetric flow rate of Q_f. However, this is not the same volumetric flow rate as in the chip. Hence, it is convenient to describe the piston flow in terms of mass flow, \( \dot{m} \). While maintaining pressure, \( P_f \), and temperature, \( T_f \), in the piston, \( \dot{m} \) can be calculated as

\[
\dot{m} = Q_f \rho_f
\]

with the density, \( \rho_f \), derived from \( P_f, T_f \), and an equation of state [16] valid under these conditions. The second piston can be set to regulate a backpressure, \( P_b \). The difference between \( P_f \) and \( P_b \) comes from the resulting pressure drop between the two pistons, \( \Delta P_{fb} \), while maintaining a flow between the pistons. Excluding the chip, the capillaries are the main source of flow restriction and \( \Delta P_{fb} \) can be estimated by considering the density \( \rho_c \) at the pressure, \( P_c \), and temperature, \( T_c \), found in and around the chip, and then applying the Hagen Poiseuille’s equation,
\[ \Delta P_{fb} = \frac{128 \mu L \dot{m}}{\pi D_H \rho_c} \] (2)

where \( L \) is channel length, \( \mu \) is the viscosity [32] and \( D_H \) is the hydraulic diameter, calculated from the perimeter and cross section of the channel. Though strictly valid for incompressible flow, it has been used with a good experimental agreement in other works [7]. For chips without pronounced flow restrictors, \( P_s, P_b, \) and \( P_c \) can be regarded as approximately equal.

The corresponding flow velocity in the chips, \( v \), it is given by

\[ v = \frac{\dot{m}}{A \rho_c} \] (3)

where \( A \) is the channel cross section in the chip.

3. Materials and method

Design

Embedded electrodes are deposited on a wafer and then sealed by bonding another wafer with microfluidic channels on top. As such procedure must be compatible with the bonding method used to create strong high-pressure resistant chips, it limits the material choices. Pt shows a high degree of corrosion resistance and is compatible with both HNO\(_3\) and KOH [33] at the temperatures, 65-85 °C, used in the bonding scheme. Pt is also resistant to the thermal treatments needed after bonding. By depositing 100 nm thick and 10 \( \mu \)m wide Pt layers, it is possible to make resistive elements in the k\( \Omega \) range, having a footprint of 110 by 100 \( \mu \)m, thus fitting well within a fluid channel. The interconnects to the sensing elements, connected in a 4-point configuration, and heater, should be wide in order to give a low resistance, but at the same time narrow not to lower the high-pressure strength of the chips since the bond strength between metal and the bonded glass wafer probably will be the weakest link.

Two different kinds of demonstrator chips are made in this paper; a pressure drop chip and a flow sensor chip, and schematic drawings of these chips are found in figure 3. The outer dimensions of both chip designs were 15 by 5 mm. Both of the designs have semi-circular channels of 160 \( \mu \)m depth and 360 \( \mu \)m width, and are open to the sides of the chip, thereby forming inlets that give access to outer diameter 150 \( \mu \)m silica capillaries, forming a fluid interface.
Drawings of the pressure drop chip (top) and flow sensor chip (bottom). Widths of chips, channels and conductors are given in µm. The sensing elements and the interconnects over the channel interface are 10 µm wide. For the flow sensing elements, the temperature sensor closest to the flow is denoted T₁ and the one further away is denoted T₂.

The ratio between channels and bonded areas is 0.93 and 0.86 for the flow sensor and pressure restrictor, respectively. On the pressure drop chip, a small 5.4 µm deep and 60 µm wide restrictor channel is in-between the wide fluid inlet and outlet channels. At the ends of the restrictor, the channel width is larger, 660 µm. By this design, a significant pressure drop is formed in the chip when flowing CO₂, much more than the $\Delta P_{fb}$ of 0.24 bar of the surrounding system. Using Eq. (2) and $D_H$ calculated from the perimeter and cross section of the channel, $\Delta P_{fb}$ increases up to 11 bar. The pressure drop will cause CO₂ to expand, resulting in cooling. To investigate this behavior, thin film temperature elements are positioned right next to both the inlet and the outlet of the restrictor.

The other design, the flow sensor chip, consists of a straight channel of the same dimensions as the inlets and holds a flow sensor in its center. The sensor consists of a heating element and two temperature elements on either side of the heater in the flow direction. All temperature elements, in both designs, have 4 conductors leading to each sensor element, and by a 4-point measurement scheme, this allows for the localized measurement of resistance in the channel.

Fabrication

The devices were manufactured by structuring two glass wafers, having metal conductors on one side and etched channels on the other. The process scheme includes wet etching using HF, fusion
bonding and a thermal treatment at 625 °C for 6 h. In this section, the focus is on the fabrication of the electrode wafers and the bonding scheme.

Borosilicate glass (Borofloat 33, Schott) 1.1 mm thick and 4" diameter, were first cleaned in a Megasonic cleaning bath (PCT Systems Inc.) for 1.5 min with a solution of 25 wt.% KOH at 65 °C. This was followed by immersion in 69 wt.% HNO₃ at 80 °C for 10 min. The electrodes, deposited on one of the wafers, were made from Pt using a lift-off process. First, a 1 min 1000 W O₂ plasma cleaning process was done (Plasma Processor 300, Tepla) which was followed by the addition of an adhesion promoting primer, HMDS, in a priming oven (Star 2000, Star). 1 μm resist (1813, Shipley) was then added by spinning and baking at 115 °C for 1 min on a hotplate. The resist was exposed and then treated in an image reversal processing oven (58TA, Yield Engineering Systems Inc) and flood exposed. To form trenches, the wafers were etched in buffered oxide etch (BOE 1:7, J.T.Baker) to an aimed depth of 100 nm. Following rinsing and spin-drying, the depth of the trench was checked using profilometry (DektakXT, Bruker). The metal film was deposited using sputtering (K675XD, Emitech) with first a 10 nm thick Ti adhesion layer followed by a 100 nm thick Pt layer. After deposition, the resist was lifted off in an ultrasonic bath with acetone for 10 min. This was followed by rinsing in IPA and O₂ plasma cleaning for 10 min at 1000 W.

The other wafer, containing the fluid channels, was wet etched with HF using a mask made from HDMS primer, spray resist (AZ9260, Micro Chemicals), and a 1 μm thick sputtered molybdenum masking layer. For the deep and shallow channels, 49 wt.% and 12.3 wt.% HF was used, respectively. The full fabrication method of the fluid channels follows [30].

The bonding was performed in ambient air at atmospheric pressure, and alignment of the wafers was done manually over a stereoscope (SMZ800, Nikon) by first adding three 0.5 cm wide tape insertions (SWT 10, Nitto) between the wafer edges and then pressing firmly on the center, thereby starting the bonding process. As this locks the wafers in place, the tape was removed and all of the wafer area bonds. Bonded wafers were placed on SiN covered dummy wafers in a vertical oven (Micro TF-6, Koyo Lindberg) and were heat-treated at 625 °C for 6 h with a ramp up and ramp down at 1 °C/min. No weights were used on the wafers. After heat treatment, the wafers were diced using a dicing saw (DAD 361, Disco). By such action, both fluid inlets and metal interconnects are exposed on the sides of the chips. To better expose the electrode edges before making the electrical connections, the electrodes were lightly polished with SiC paper (600 grit).

Chip assembly

75 μm inner diameter and 150 μm outer diameter capillaries were manually inserted 2-2.5 mm into the inlets on the sides of the chips and glued using 2-component epoxy glue (Araldite Rapid, Huntsman Advanced Materials). To allow for transmission lightning, the chips were then placed
on 100 μm thick cover glass slabs. When the external temperature of chips needed to be raised, resistive Kanthal wirings were used. To achieve good thermal conductivity, the wirings were wound around AlN substrates and connected to the chips with thermal grease. As the chips required both electrical connections and a high-pressure fluid interface, a mounting fixture was made. It consisted of two printed circuit boards, PCBs, with centered holes. The cover glass with the chip was mounted in between the PCBs, figure 4. To gain electrical access to the chips, 200 μm thick Cu wires were on one end soldered onto pads on the top PCB, and on the other end glued to the sides of the chip using electrically conductive 2-component epoxy glue (CW2400, CircuitWorks). A flat cable connected to the top PCB provided electrical access to the instrumentation.

Experimental Setup

To characterize the chips, a high-pressure test system was used that consisted of two high-pressure piston pumps (100 DM, ISCO Teledyne), cooling baths and a high-speed camera mounted on a microscope. The forward moving piston, figure 2, contained CO₂ (99.9995%, AGA). The tubing and forward moving piston were cooled by a compressor (RK-20, Lauda) connected to a 20 L cooling bath held at 4.5 °C. The large thermal buffer was used for increased stability. The chips, mounted on a fixture with electrical and fluid interfaces, where held about 5 cm above this cooling bath and a high brightness LED was mounted just above the water line. To measure the temperature of the forward moving piston, Tₚ, a Pt100 sensor was used in 4-point configuration together with a data acquisition control unit (34970A, Agilent). The high-speed camera (Miro 320, Phantom Vision) was mounted on a stereoscope (SMZ800, Nikon) above the chip fixture, forming a high-speed transmission light setup.

The electric resistance of the integrated temperature elements was measured using another data acquisition control unit (9219, National Instruments) in 4-point configuration and the data was recorded using LabView. The heating element of the flow sensor was operated using a current controlled power supply (QL335TP, TTi).

As the chips needed to be observed with the transmission light and microscope, a large cooled heat stage could not be used because it would block the light path. Instead, most of the heat needed to dissipate through the side of the chip, into the fixture. Because of this, during the flow sensor measurements, Tₚ varies between 17.7 and 23.5 °C for the subcritical condition, and between 33.0 and 46.8 °C for supercritical conditions as a result of the heat generated by the high brightness LEDs. For the pressure restrictor measurements, Tₚ varied between 15.9 and 16.2 °C. This is not a problem as the internal sensors in the chips directly measure Tₑ.

Geometric characterization

The metallization layers were characterized using scanning electron microscope, SEM, (1550, Zeiss), white light interferometry (Wyko NT1100, Veeco) and optical microscopy (AX70, Olympus). Access to the metal layers exposed to the insides of the channel was achieved by grinding a topside opening to the channels. Cross-sections of the metal layers that were embedded between wafers were done by grinding and etching in 12.3 wt.% HF for about 10 s. To avoid charging, samples used for SEM were coated with an approximately 6 nm thick sputtered Pd/Au film.

Measurements

Fracture tests were done using CO₂ that first was pressurized to 80 bar. Capillaries were glued to
one of the two inlets of each chip and connected to the forward moving pump. The other inlets were plugged with glue. During the pressure tests, the chips were put on cover glass slides without any clamping and no electrical connections were made. The connected capillary was 10 cm long and 150 µm in outer diameter and exerted a minimal mechanical force on the samples. By increasing the pressure at a rate of 0.5 bar/s until fracture, the fracture pressure was recorded. Chips were held at room temperature during these tests and a high-speed camera was used to follow the fracture behavior. To translate fracture pressure into fracture stress, the channel geometry of the flow sensor was modeled using finite element analysis (Comsol Multiphysics 5.2, Comsol). Leakage measurements were done with water and an external flow sensor (Nanoflow Sensor, IDEX) that was connected to the chips having their outlets plugged. The external flow sensor has a lower detection limit of 1.5 nL/min. Fluorescent dye has been capable of visualizing nanochannels [34-35]. To see if any leakage could be detected through the possible voids along the metal, samples were filled with an aqueous solution containing 14 g/L fluorescein (Sigma-Aldrich), both by means of capillary action and by pressurization to 90 bar. Following depressurization and flushing of excess fluorescein in the channels using water, the chips were observed using a fluorescent microscope (TE2000-U, Nikon).

To calibrate the sensors, the mounted chips and fixtures were put in a convection oven held at 50 °C (400, Memmert) together with a K-type thermocouple which was used as a temperature reference. The oven was turned off and allowed to cool during 5 h, giving correlated temperatures down to room temperature. The resistance change was linearly fitted against the temperature decrease. The temperature sensor noise was recorded while operating the chips without a pressurized fluid and taken as 2 times the standard deviation.

The pressure drop chip, used to study local temperature variations, was used for two experiments. In the first experiment, the backpressure piston was set to a constant pressure mode and the forward moving piston was set to constant volumetric flow mode. By such action, Pb was set to 50 or 57 bar and Qa was varied over the range 10 to 100 µL/min. The chip temperature, Tc, was measured using the internal temperature sensor located at the chip restrictor inlet. The decompression cooling effect over the restrictor, \( \Delta T \), was measured as the absolute difference between the inlet and outlet temperature sensors. In a second experiment, the Tc was recorded as a function of time and the forward moving piston was instead operated in constant pressure mode with the backpressure piston held at a fixed volume. Variations in Tc were recorded as Pf was changed in steps.

For measurements with flow sensor chips, the backpressure piston was again set to hold constant pressure and the forward moving piston was set to constant volumetric flow mode. As a measurement of Tc, the average temperature of the two sensor elements, denoted T1 and T2 in figure 3, was used for calculating a corresponding flow velocity v for each measurement. As no large \( \Delta P_{fb} \) is present; Pb \( \approx \) Pc \( \approx \) Pf and (1) and (3) together with Tc, Pb and Tf can be used to describe v. Two measurement series were conducted, with Tc held either under or over the critical temperature of 31.1 °C. For subcritical conditions, Pb was set to 80, 100 or 120 bar. For supercritical conditions, Pb was set to either 80 or 100 bar. The heating element was used at 1, 3 or 5 mA, producing a power of 0.41, 1.23 and 2.05 W, respectively.

To evaluate the temperature difference, \( \Delta T \), response with v, data was fitted to a nonlinear model of the form,

\[
\Delta T = a(1 - e^{-bv}) + c ,
\]

where a, b and c are constants. To define a statistically relevant working range of the flow sensor
at a specific condition, the detection limit is determined using the root-mean-square error, $s_{\text{rmse}}$, of each fitting curve, and it is taken as $2s_{\text{rmse}}$. Given a resolution, $\Delta v$, in this work set to 1 mm/s, the upper limit of the working range is calculated from the minimal sensitivity, $\frac{\Delta T}{\Delta v}$, required to produce a detectable change of $\Delta T$, i.e. $2s_{\text{rmse}}$.

4. Results

Electrode characterization

The heater and temperature sensors were integrated into the channels of the chip, with sensor elements exposed to the fluid, figure 5. The electrodes were embedded by letting the metal fill an etched trench in the channel lid, figure 6.

Figure 5 The flow sensor (top) and pressure drop (bottom) device. The smaller sensing area has 10 $\mu$m wide electrodes.
Figure 6 The Pt metal of an electrode in the channel is seen to the left. The glass is seen to the right, with the transition in the middle. The view is tilted 70° in the direction perpendicular to the metal and glass transition.

Figure 7 shows a height representation of the exposed part of the nominally 10 µm wide electrodes in the channels after bonding and thermal treatment. The metal surface had a surface roughness of 4 nm and larger hills and valleys. Due to these variations, the metal was at most 35 nm above the glass plane. As this surface topography is inherent to the deposited film, these variations are also present at the bond interface. This implies the presence of gaps with a shape similar to figure 1b and a height \( d \) in the range of 35 nm. A trench between the metal and glass was formed, having an average depth of 20 nm and width of about 2.5 µm. This trench can also be seen in the tilted view in figure 6.

![Figure 7](image)

**Figure 7** Height representations, (a), of a 10 µm wide embedded electrode exposed to the channel inside. The profilometry curve in (b) has valleys on each border of the metallization.

After bonding, the enclosed metallization had no apparent voids or delamination surrounding it when cross-sections of the metal layers were examined using SEM and the glass seems to conform around the metal layer. This is fundamental for leak resistance at high pressure.

When operating the chips with CO\(_2\) at 90 bar, no leakage was seen when observing the chips under the stereoscope. Most importantly and conclusive, no fluorescence could be seen either on top or along the metal pathways when the chips were filled with the fluorescent solution. Also, when testing for leakage using the external flow sensor, no flow was registered to keep the pressurization of the chips. This implies that any leakage was below the detection limit of 1.5 nL/min. The fracture tests showed that fracture occurred at pressures in the range of 80 to 220 bar. The pressure drop and flow sensor chips fractured at pressures between 80 to 173 bar \((n = 6)\) and 190 to 220 bar \((n = 4)\), respectively. For these measurements, the average fracture pressures were 126 and 201 bar, while the standard deviations were 12 and 35 bar, respectively. Modeling of the flow sensor chip showed that 201 bar corresponds to an average fracture stress of 73 MPa. As a comparison, other studies with glass chips with similar geometries but without integrated electrodes showed fracture stresses between 129 and 254 MPa [30]. Both types of chips were operated continuously at 90 bar for more than one week.

By investigation of the fracture surfaces, it was seen that the crack formed during fracture originates from the bond interface edge of the channels. The crack deviates from this interface plane, forming a fracture surface with a topographic height difference from the bond plane of up to 30 µm. But, in the proximity the channels, the direction of the fracture deviates so that the
fracture goes along and exposes the metallization layer, showing that the metallization affects the crack growth.

*Temperature sensing in the pressure drop chip*

Cooling as a consequence of flows through the restrictor on the pressure drop chip was seen, figure 8. The temperature at the inlet was 15.9 and 16.2 °C. With $P_b$ at 50 bar, a $\Delta T$ up to 5.1 °C was seen as well as a transition from liquid to a gas phase at the outlet, thus forming a multiphase system. With $P_b$ slightly higher, 57 bar, the gas phase disappeared and a lower $\Delta T$ was observed, around 0.6 °C. This difference was of similar levels also for higher pressures, and depressurization of $P_f$ from 90 to 70 bar caused a 1.8 °C drop of $T_c$ from 13.2 °C down to 11.4 °C with a rate of 0.2 °C/s. The temperature sensor noise was 0.007 °C.

*Flow sensing in CO$_2$*

Figure 9 shows $\Delta T$ as a function of flow rate $v$ for both sub- and supercritical conditions. Except for the measurement with 2.05 W at low temperature, subcritical conditions, no measurements at 80 bar could be fitted to a typical flow sensor response explaining the lack of curve fits for most of the 80 bar measurement series. The working ranges of the flow sensor at the specific conditions are presented in table 1. In the table, the resulting density range for the given condition is presented. Operating the sensor at 2.05 W yields the largest thermal gradient and the largest range, between 0.5 to 13.8 mm/s for a $\Delta v$ of 1 mm.

While the sensor noise of the flow sensing chip is similar to that of the pressure drop chip, the measurement variation, $2s_{\text{sensor}}$, is much larger, up to 0.057 °C for the conditions that could be used to fit a curve. This variation is affected by the fluid system and the varying properties of CO$_2$.

At low pressure, boiling occurs at the heater during operation, as exemplified in figure 10 where the pressure is 60 bar and the heating output is 2.05 W. The flow sensor is not working properly when boiling occurs.
Figure 8 Cooling effect, $\Delta T$, at the restrictor of the pressure drop chip as a function of $Q_r$. 
**Figure 9** Temperature difference as a function of flow velocity for subcritical (top) and supercritical (bottom) state. Fitting curves with corresponding range limits have been added.

**Figure 10** Boiling with bubble formation at the flow sensor heater running at 2.05 W. The pressure is 60 bar. The flow goes from right to left in the picture.
Table 1. Flow sensor range.

<table>
<thead>
<tr>
<th>$P_b, T_c^a$ (bar, °C)</th>
<th>Power (W)</th>
<th>$P_c (kg/m^3)^b$</th>
<th>Range (mm/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>120, 17.8-22.3</td>
<td>880-886</td>
<td>0.6-4.7</td>
<td>0.5-7.2</td>
</tr>
<tr>
<td>100, 18.2-22.9</td>
<td>854-861</td>
<td>0.1-1.2</td>
<td>0.5-8.1</td>
</tr>
<tr>
<td>80, 18.3-22.8</td>
<td>826-834</td>
<td>$^c$</td>
<td>$^c$</td>
</tr>
<tr>
<td>100, 36.3-46.2</td>
<td>537-680</td>
<td>0.8-3.5</td>
<td>0.9-3.2</td>
</tr>
<tr>
<td>80, 33.5-46.4</td>
<td>254-511</td>
<td>$^c$</td>
<td>$^c$</td>
</tr>
</tbody>
</table>

$^a$ Maximum and minimum values.  
$^b$ Calculated from $P_b$ and $T_c$.  
$^c$ Deviating fitting curve – No range

5. Discussion

Embedded metal layers

Insight in the conditions at bonding is given by figure 7. As the etched trench depth and the metallization thickness mismatch, the electrodes are up to 35 nm higher than the glass, thereby suggesting a situation similar to figure 1 b, where metal is present above the bond layer. While this height is less than the critical size for a spherical particle, the large width of the exposed metal could raise some concerns over the ability to bond. However, no problems have been seen during bonding and the chips can sustain high pressures. The mismatch between the etch trench and the metal deposition results in a 2.5 µm wide and 20 nm deep valleys surrounding the electrodes, but after bonding, this gap was no longer possible to detect in SEM.

While any voids on the nanoscale would allow fluids to flow [35-36], no evidence of leakage has been found either by using fluorescent dye or flow measurements. The lack of any leakage, visible gaps or voids, suggests a mechanism that seals the structure during the thermal treatment. Glass sinters at lower temperatures through viscous flow and a surface energy reduction [37]. SiO$_2$–B$_2$O$_3$–Al$_2$O$_3$ glass powders have been shown to sinter at their glass transition temperature, $T_g$, of 624 °C and have optimal sintering temperatures of 700 °C [38]. As the $T_g$ of borofloat glass is 625°C [29], both the temperature and time of the heat treatment as well as the dimensions of possible gaps satisfies the conditions for sintering.

Both demonstrator chips withstand the pressures needed for operation around the critical point of CO$_2$. Consistent with the larger geometry of the pressure drop chip, the average fracture pressure was 126 bar compared to the 201 bar achieved for the flow sensor chip. The simpler semi-circular channel geometry of the flow sensor allows for strength-comparisons using modeling to find the correlating fracture stress. The average fracture stress was 73 MPa which is about 40% less than the stresses seen for chips without electrodes, found in both our previous work [30] and the works of others [39-40]. This and the fact that the crack formation is irregular, following and exposing the metallization, suggest that metallization lowers the fracture strength and the integrated electrodes act as defects. The fracture strength in glass chips is a complex property, not only dependent on the stress, and crack initiating defects are affected by stress corrosion [30]. This makes the device strength also dependent on the time, temperature and environment. While the exact crack formation is unknown, it is prudent to keep both metal structures and channel dimensions small where the electrodes enter the pressurized channels, thus lowering both the
defect size and local stress.

\textit{Measurements of CO}_2 \textit{in a microfluidic system}

The chips clearly demonstrate the possibilities with integrated electrodes for added control of supercritical fluids in microfluidics. Using these systems, direct measurements of the inner wall temperature are made; enabling observations of both the cooling effect at pressure drops and flow velocity measurements up to 0.5-13.8 mm/s.

It can be confirmed that cooling effects are present while using a restrictor channel on the chip. The boiling point of CO\textsubscript{2} is 14.3 °C at 50 bar and 19.8 °C at 57 bar. Therefore, the detectable temperature changes of figure 8 at 57 bar represent liquid CO\textsubscript{2} expanding as it exits the resistor. Meanwhile, for the measurements at 50 bar, there is also the formation of a gas phase with a connected energy of evaporation. As this forms a non-equilibrated multiphase composition of both gas and liquid CO\textsubscript{2}, quantitative predictions of the energy output is hard. As multiphase flows are strongly dependent on inertial, viscous and surface tension forces, and with the high variability of such forces for near-critical CO\textsubscript{2}, getting full control is elusive [17]. Large temperature drops are troublesome for advanced fluid handling where it is common to work close to solubility limits. For several compounds of interest, a drop in temperature of a few degrees can give a supersaturated solution with the compound of interest, and particle formation can result in clogging. Localized temperature sensors as the ones presented here for high-pressure applications give an increased understanding and a precise process control, thereby allowing for optimizations for complex fluid operations.

The flow sensor has been shown to give reliable results at specific conditions. In general, when densities are high, \(\Delta T\) shows lower scattering. However, while operating the sensor at 80 bar and supercritical conditions, the large variation made it impossible to make a normal flow curve fit. This is because a calorimetric sensor response is connected to the heat capacity, \(C_p\). In figure 11, it can be seen that at pressures just above the critical point, \(C_p\) shows large variations around a maximum and then decreases with increasing pressure.

For best performance of the flow sensor, sensing should be done either at subcritical conditions or at higher pressures, where both \(C_p\) and density show less variation. As exemplified in this paper, it is, however, possible to extract flow data even at the highly variable conditions around the supercritical point. For an integrated microfluidic platform with several functional units, different temperature or pressure zones on the same chip can be required and conditions around the critical point may present. Due to the highly complex and often non-equilibrated nature of supercritical fluids, the thermal and fluid dynamic aspects of such integrated sensors should be further modeled together with experimental data to give valuable insights.
Pt electrodes have been integrated into high-pressure glass chips being able to sustain pressures and temperatures for supercritical CO₂. Localized cooling effects of up to 5 °C are seen in a pressure drop chip. Using the integrated, high-pressure tolerant, calorimetric flow sensor, flow sensing under supercritical conditions was done with a workable range up to 0.5-13.8 mm/s. The integration of thin film sensors adds an essential control to high-pressure microfluidic platforms.

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