

# Magnetically Enhanced Microflow Cytometer for Bead- and Cell-based Immunoaffinity Measurements in Whole Blood Samples



Scientific thesis for the attainment of the academic degree Master of Science (M.Sc.) of the Department of Electrical and Computer Engineering at the Technical University of Munich.

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**Submitted on** December 4<sup>th</sup>, 2020 at Munich

# Contents

1	Abs	ostract 4				
2	The	ory				
	2.1	Microfluidics				
		2.1.1	Flow Field inside Microchannels	5		
		2.1.2	Particles in Microfluidics	7		
	2.2	Surfac	e Chemistry	9		
		2.2.1	Surface Oxidation Methods	9		
		2.2.2	Silane Chemistry	12		
		2.2.3	Carbodiimide Crosslinker Chemistry	13		
		2.2.4	Microscopic Particle Surface Physics	15		
		2.2.5	The Biotin-Avidin-System	15		
	2.3	Magne	etoresistive Sensing	15		
		2.3.1	Sensing Principle	15		
3	Mate	erials a	nd Methods	16		
	3.1	Microf	luidics and Sensor Fabrication	16		
		3.1.1	Development of Layout	16		
		3.1.2	Patterning of Photoresist	16		
		3.1.3	Soft Lithography	17		
		3.1.4	Bonding of Microfluidic	17		
		3.1.5	Electrical Circuit	18		
		3.1.6	Electronic Readout	18		
	3.2	Magne	etic Beadometry	19		
		3.2.1	Standard Parameters	19		
		3.2.2	Concentration Measurement	19		
		3.2.3	Whole Blood Bead Spiking	19		
		3.2.4	Bead Capture Assay	19		
		3.2.5	Optical Particle Tracking	19		
	3.3	Surfac	e Bio-Functionalization	19		
	3.4	Tensio	metry	19		
		3.4.1	Surface Activation	19		
		342	Chemical Surface Functionalization	20		

		3.4.3	Surface Bioconjugation	20		
		3.4.4	Particle Functionalization	20		
4	Res	ults		21		
	4.1	Virtual	Prototyping of Cell Signals	21		
		4.1.1	Single Cell Signal	21		
		4.1.2	Cell Aggregates	21		
	4.2	Refere	nce Bead Surface Functionalization	21		
		4.2.1	Amine-Surface Biotinylation	21		
		4.2.2	Carboxy-Surface Biotinylation	25		
	4.3	Conce	ntration Measurements in MRCyte	25		
		4.3.1	Count Stability	25		
		4.3.2	Calibration of Flow Field	25		
		4.3.3	Differential Counting Setup	25		
	4.4	Proteir	Immobilization On The Microfluidic Channel Bottom	27		
		4.4.1	Physisorption	27		
		4.4.2	Covalent Attachment	28		
5	Disc	ussion		30		
6	Outl	ook		31		
List of Abbreviations						
List of Figures						
Lis	List of Tables					
Bil	Bibliography					
Sta	tatement					

# 1. Abstract

# 2. Theory

The main measurement principle by a giant magneto resistance (GMR)-Sensor has been already described and characterized exhaustively by Helou [1], Reisbeck [2] and Brenner [3]. Therefore, this theoretical part will focus on (bio-)physical aspects of a cell rolling motion inside a microfluidic channel and surface modification chemistry.

#### 2.1. Microfluidics

The main experiments of this work were carried out in microfluidic environments, which exhibit favorable properties compared to common turbulent systems. From a fluid-mechanical standpoint, shrinking the scales makes interfacial as well as electrokinetic phenomena much more significant, and reduces the importance of pressure and gravity.[4] However, electodynamics, chemistry and fluid dynamics are incetricably intertwined, so that fluid flow can create electric fields (and vice versa), with a degree of coupling driven by the surface chemistry. Many of the resulting phenomena arise or can explained by Cauchy-Momentum equation (eq. 2.3) and the resulting Navier-Stokes equation for incompressible fluids (eq. 2.4).

$$\frac{\partial}{\partial t} \iiint \rho d\mathbf{V} = -\iint \rho \mathbf{u} \cdot \vec{\mathbf{n}} d\mathbf{A}$$
 (2.1)

$$\nabla \cdot \mathbf{u} = 0 \tag{2.2}$$

$$\rho \frac{\partial \mathbf{u}}{\partial t} + \rho \mathbf{u} \cdot \nabla \mathbf{u} = \nabla \cdot \boldsymbol{\tau} + \sum_{i} \mathbf{f}_{i}$$
 (2.3)

$$\rho \frac{\partial \mathbf{u}}{\partial t} + \rho \mathbf{u} \cdot \nabla \mathbf{u} = -\nabla p + \eta \nabla^2 \mathbf{u} + \sum_{i} \mathbf{f}_{i}$$
Transient Convection Pressure Viscous Body Forces
$$(2.4)$$

conservation of mass, momentum reynolds number

#### 2.1.1. Flow Field inside Microchannels

The foremost characteristic of a microchannel is the laminar flow behavior, which causes deterministic pathlines. Mathematically, this is described by the reynolds number, which compares the intertia to shear forces. If it results below a certain threshold of 2000,

laminar flow can be assumed. This holds true for the utilized microfluidic with the dimensions  $12\,000\,\mu\text{m} \times 700\,\mu\text{m} \times 150\,\mu\text{m}$  (I x w x h) and aequous buffer solutions, where the channel width was used as characteristic length l. Hence, simplifications of the Navier-Stokes equation can be applied to our system.

$$Re = \frac{2\rho|\overline{u}|l}{\eta} \tag{2.5}$$

The step from the Cauchy momentum equation to the Navier-Stokes equation is complex and harbors several sources of error. First, an incompressible newtonian fluid as well as channel is assumed. The used water suspensions can be approximated with negligible compressibility, which is not true for the real case. Also, for blood or other shear-thinning fluids some deviations are prone for high errors. This happens due to the fact that the surface stress tensor  $(\tau)$  is decomposed into pressure and viscous contributions as shown in the equations 2.6. Then, the divergence relation of the respective viscous stress (eq. 2.7) does not hold for non-uniform viscosity  $\eta$ .

$$\tau = \tau_{viscous} + \tau_{pressure} = 2\eta \epsilon - p\mathbf{I}_{3\times 3}$$
 (2.6)

$$\nabla \cdot \boldsymbol{\tau}_{viscous} = \nabla \cdot 2\eta \epsilon = \nabla \cdot \eta \nabla \mathbf{u} \stackrel{only \ if \ \eta}{=} \eta \nabla^2 \mathbf{u}$$
(2.7)

Second, the channel height varies in reality as a result of fabrication inaccuracies. In the model case of a flow through a rectangular channel, no analytical solution of the Navier-Stokes equation exists, but a Fourier Series expansion if channel width is larger than channel height. [5] The equation 2.8 shows that height deviations can have prominent influence on a channel velocity simulation as it is proportional to  $h^2$ . Further, the flow rate (which is the velocity integral over the channel cross section) depends even on  $h^3$ .

$$u_x(y,z) = \frac{4h^2 \Delta p}{\pi^3 \eta l} \sum_{n,odd}^{\infty} \frac{1}{n^3} \left( 1 - \frac{\cosh(n\pi \frac{y}{h})}{\cosh(n\pi \frac{w}{2h})} \right) \sin(n\pi \frac{z}{h})$$
(2.8)

Third, the transient term (eq. 2.4) was neglected in all simulations, but a connected syringe pump possesses a slow rise time (Fig. 1a) and a remaining "pulsation error" in steady state (Fig. 1b). In effect, another error adds to the simulation, which is only valid after several ten seconds of the last flow rate change.

For later studies in a matlab model, the flow velocity and shear stress computations

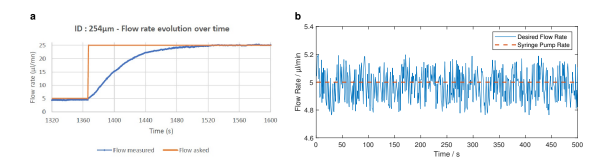


Figure 1: Syringe Pump error sources
Set flow rate: \_\_\_\_\_\_, Real Flow Rate: \_\_\_\_\_\_ a, Transient step answer of a syringe pump through a microtube with 254 μm inner diameter. b, Steady state flow rate error around the desired 5 μL min<sup>-1</sup> dispensing rate. A sinusoidal behaviour caused by the microstepping can be observed. [6]

were carried out with the error sources considered.

#### 2.1.2. Particles in Microfluidics

Stokes Drag Force Gravity Electro-static interaction Magnetic Force Friction Interface-Forces

## 2.2. Surface Chemistry

Introducing biological samples, such as plasma or whole blood, into microsystems needs more consideration of surface modification compared with buffered samples of adjusted pH containing cells or polymeric beads. Blood-material contact most often initiates surface-mediated reactions that lead to cell activation, blood clotting or biofilm formation. Therefore, most contact faces are passivized with chemically and biologically inert materials or even composed entirely from it. In any use case, where the sensor surface has to be functionalized with biomolecules, the surface inertness then requires specialized methods for permanent and reproducible adhesion.[7]

Molecules can be immobilized through various mechanisms on surfaces to achieve a biological or chemical functionality. The most simple is physisorption. Here, a biomolecule is bonded only by weak elektrostatic, van-der-Waals or dipole-dipole interaction with a adsorption enthalpy below 50 kJ mol<sup>-1</sup>. In contrast, this yields fast reaction rates, because no activation energy has to be overcome. Although a large number of molecules can be captured with this method, several drawbacks have been identified. [8], [9] For example, immobilized receptors can start to desorb or change their position, which in turn reduces sensitivity or causes false-positive results. [10], [11]

Therfore, most functionalization approaches rely on chemisorption where molecules are covalently bound to a surface. Due to the higher activation energy barrier this bonding mechanism works slower in comparison to physisorption, though higher temperatures or catalysators can promote an equilibrium. One of the most well-known strategies to bring reproducible thin films on surfaces is the formation of self-assembled monolayers (SAMs) where a dense layer of single molecules with high internal order forms upon dipping into a surface-active substance. [12]

#### 2.2.1. Surface Oxidation Methods

To modify a surface with functional silanes, oxidized sites (-OH (hydroxyl) resp. Si-OH (silanol) groups) have to be present. In order to increase the presence of those reactive groups on differing substrates, various activation methods such as piranha, oxygen gas  $(O_2)$  - plasma treatment or an hydrofluoric acid (HF) dip can be chosen. [13]

Critical for any surface engineering is the internal structure and in consequence the binding energies of the surficial groups. The three mainly used substrates in this work,

glass, poly(dimethyl siloxane) (PDMS) and silicon nitride (Si<sub>3</sub>N<sub>4</sub>), contain highly conserved, homogeneous surfaces and are mostly well characterized. The surface of glass exhibits already silanol groups intrinsically and consequentially demands only a removal of impurities. PDMS and Si<sub>3</sub>N<sub>4</sub> however have different compositions as shown in Fig. 8 and 3 hence requiring a strong oxidation agents to completely exchange its interface. [14]–[16]

Figure 2: Different substrate surfaces: glass and PDMS Surface groups and internal structure of quartz glass (a) and PDMS (b). After an oxidation step, the methyl groups are changed to hydroxyl.

#### Piranha Solution

Piranha is an oxidizer composed of hydrogen peroxide ( $H_2O_2$ ) and sulfuric acid ( $H_2SO_4$ ), typically in volume ratios between 1:3 and 1:7. The effectiveness of piranha in removing organic residues and creating hydroxyl groups is induced by two distinct processes. In the first process, which is notably faster, hydrogen and oxygen are removed as units of water by the concentrated  $H_2SO_4$ . (Reaction 2.9) This occurs due to the thermodynamically very favorable reaction with an enthalpy of  $-880 \, \text{kJ} \, \text{mol}^{-1}$  and produces Caro's acid ( $H_2SO_5$ ), one of the strongest oxidants known. [17]

$$H_2SO_4 + H_2O_2 \longrightarrow H_2SO_5 + H_2O$$
 (2.9)

$$H_2SO_4 + H_2O_2 \longrightarrow HSO_4^- + H_3O^+ + O$$
 (2.10)

In another process the sulfuric acid boosts hydrogen peroxide from a mild oxidizer into the more aggressive atomic oxygen by the dehydration of  $H_2O_2$ . (Reaction 2.10) These two dehydration processes in the mixture result on the one hand in a highly corrosive nature against organic materials, particularly against the difficult to remove carbon. On the other hand, it is strongly acidic and oxidizing which in turn requires great care and substantial safety measures to prepare and use it harmlessly.

#### **Hydrofluoric Acid**

One of the used substrates in this work is Si<sub>3</sub>N<sub>4</sub> as passivation layer above magnetic sensors as it has a significant better diffusion barrier against water or sodium ions and is chemically very inert. [18] However, due to its complex crystal structure it is also difficult to modify by com-

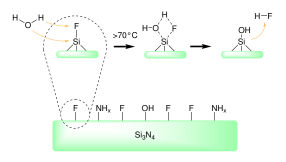


Figure 3: Proposed modification of  $Si_3N_4$  with HF

mon chemicals and the exact surface composition still subject to scientific discussion. [19] Apart from cleaning the surface with piranha, few other modification methods have been reported, but only one suitable for the direct generation of hydroxyl groups.

There, as depicted in 3, the reaction  $Si-OH + HF \longleftrightarrow Si-F + H_2O$  takes place reversibly due to the coincidence that Si-O and O-H as well as Si-F and H-F bonds have similar binding energies and hence the forward and reverse reactions a low activation energy. After Le Chatelier's principle, a depletion of HF in the bulk leads then to an increase in surficial hydroxyl groups. [20] In further works, it has been determined that an oxidation with a similar protocol based on aequous HF yields a variable Si-O-Si (siloxane) coverage with  $37 \pm 17 \%$  of a monolayer, which nevertheless can be used for stable, covalent attachment of silanes. Nominally the same surface coverages of silicon oxide and nitride surfaces could be achieved by ethoxy- and chlorosilanization. [21] As shown by [22], the subsequent surfaces exhibit beneficial biological properties and can be modified by further standard procedures.

#### Oxygen Plasma

Apart from wet chemistry methods, the exposure of a surface to oxygen plasma yields hydroxyl groups as well. In a plasma chamber, a low-pressure gas is irradiated by kHz to MHz waves to excite and ionize its atoms. In consequence, the UV-radiation emitted by the gas can photolyse typical organic bonds and remove surface contaminations. Additionally, reactive oxygen species such as  $O_2^+$ ,  $O_2^-$ ,  $O_3$  or O either oxidize the surface as well or bind dissociated components with low vapor pressure. During an evacuation in the process, these molecules are removed from the chamber intrinsically. [23]

#### 2.2.2. Silane Chemistry

By the use of silane chemistry a surface is rendered organofunctional with alkoxysilane molecules. Since glass, silicon, alumina, titania, and quartz surfaces, as well as other metal oxide interfaces, are rich in hydroxyl groups, silanes are particularly useful for modifying these materials. [24]

The general formula for a silane coupling agent (Fig. 4) typically shows the two classes of functionality. X is a hydrolyzable group typically alkoxy, acyloxy, halogen or amine.

Following hydrolysis, a reactive silanol group is formed, which can condense with other silanol groups to form siloxane linkages. (Fig. 5) Stable condensation products are also formed with other oxides such as those of aluminum, zirconium, tin, titanium, and nickel. Less stable bonds are formed with oxides of boron, iron, and carbon, whereas alkali metal oxides and carbonates do not form stable bonds with siloxanes at all. The R group (Fig. 4) is a nonhydrolyzable organic radical that may posses a functionality that imparts desired characteristics. One

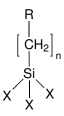


Figure 4: Trialkoxysilane Structure of a typical trialkoxysilane, X: hydrolyzable group, R: non-hydrolyzable organic radical, n: methylene chain-length

of the more common silanes is (3-aminopropyl)triethoxysilane (APTES), where the X group consists of an  $-O-CH_2-CH_3$  (ethoxy) group, the organic rest R is substituted by an  $-NH_2$  (amine) and the  $3-CH_2-$  (methylene) groups alter n to 3. [25]

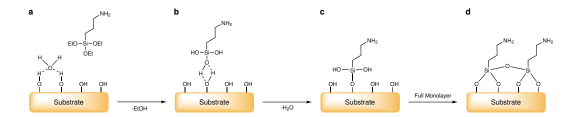


Figure 5: APTES Modification of an oxidized surface

a Before the condensation reaction, the oxidized surface forms hydrogen bonds with water molecules. The silane molecules are in the bulk solution. **b** The hydrolyzed silanol group adsorbs onto the surface and forms hydrogen bridges with it. **c** In a condensation reaction, under the loss of water, a covalent bond to the surface forms. **d** After the SAM assembly the surface is saturated with a covalent-bound, crosslinked silane film. [26]

The final result of reacting an organosilane with a substrate ranges from altering the wetting or adhesion characteristics of the substrate, utilizing the substrate to catalyze chemical transformation at the heterogeneous interface, ordering the interfacial region, and modifying its partition characteristics. Significantly, it includes the ability to effect a

covalent bond between organic and inorganic materials. Especially in optical or biological sensors, silane modifications open a broad range of applications.

However, the silanization reactions bear a few drawbacks which are often neglected. For instance, silane chemistry is strongly temperature and pH-dependent. [27], [28] Further, in a process to build SAMs out of APTES, the reaction has to be catalyzed by water. But already small changes in the water content cause dramatic deviations in layer thickness. [29] Additionally, silanes can crosslink to themselves through possible side reactions. (Fig. 5 D) [30]

#### 2.2.3. Carbodiimide Crosslinker Chemistry

The in previous manner produced amine-terminated films by APTES form the basis of many reactions and open the possibility to various applications, such as the direct attachment of biofunctional molecules by carbodiimide crosslinking chemistry.[31] Here, -COOH (carboxyl) groups are modified by 1-ethyl-3-(3-dimethylaminopropyl)carbodimide (EDC) and N-hydroxysuccinimide (NHS) to form a stable secondary  $R_1-CONH-R_2$  (carboxamide) bond with any primary amine.

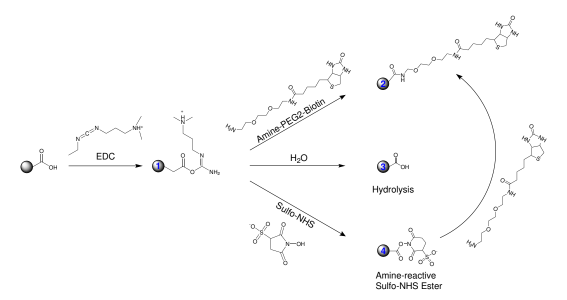


Figure 6: Carboxyl bead modification with EDC/NHS

The carboxy groups bead are activated with EDC to an active O-acylisourea intermediate. This can then either be nucleophilicly attacked by a primary amine of the amine- $PEG_2$ -biotin reactant or - due to its instability - hydrolzed back to a regenerated carboxyl surface. A present NHS-ester can also displace the O-acylisourea to form a considerably more stable intermediate which then itself reacts with any primary amine.

The general reaction mechanism is depicted in Fig. 6 for the example of a microbead surface, but it can equivalently be applied to any other modified surface or molecule. The initial carboxyl group is esterified by EDC to an active o-acylisourea intermediate and leaves rapidly upon nucleophilic attack of an amine with release of an iso-urea byproduct. A zero-length amide linkage is formed. (Fig. 6, 1->4) Sulfhydryl and hydroxyl groups also will react with such active esters, but the products of such reactions, thioesters and esters, are relatively unstable compared to an carboxamide bond. (Fig. 6, 1)

However, this reactive complex is slow to react with amines and can hydrolyze in aqueous solutions, having a rate constant measured in seconds. If the target amine does not find the active carboxyl before it hydrolyzes (Fig. 6, 3), the desired coupling cannot occur. This is especially a problem when the target molecule is in low concentration compared to water, as in the case of protein molecules. Notwithstanding, forming a NHS ester intermediate from the reaction of the hydroxyl group on NHS with the EDC active-ester complex increases the resultant amide bond formation remarkably. (Fig. 6, 3->4) [32]

Another critical point in carbodiimide chemistry is the solubility of the compounds. EDC, NHS and N-hydroxysulfosuccinimide (sulfo-NHS) are soluble in aqueous and organic solvents. Nevertheless, activation with non-sulfonate NHS decreases water-solubility of the modified carboxylate molecule, while activation with sulfo-NHS preserves or increases its water-solubility by virtue of the charged sulfonate group. [33]

## 2.2.4. Microscopic Particle Surface Physics

# 2.2.5. The Biotin-Avidin-System

## 2.3. Magnetoresistive Sensing

Short intro over MRCyte Foto of setup with arrows to necessary parts Microscope Stages PEEK holder Helmholtz coils Kepco MFLI DAQ

## 2.3.1. Sensing Principle

Loss because of reduced velocity and magnetic drag

Different produced GMR stacks Wheatstone Bridge setup Magnet alignment

# 3. Materials and Methods

#### 3.1. Microfluidics and Sensor Fabrication

#### 3.1.1. Development of Layout

#### 3.1.2. Patterning of Photoresist

3" (100) silicon wafers (Si-Mat) were dehumidified in a drying oven (UN30, Memmert) for 2 h at 150 ℃ to 180 ℃. Then, immediately after they reached room temperature, they were placed centered inside a wafer spinner (WS-650-23B, Laurell Technologies). For the desired layer thicknesses 2 mL to 3 mL SU8-30XX (Microchem) were poured carefully onto the center of the wafer and the following program was carried out:

- 1.  $500 \text{ rpm for } 10 \text{ s at } 100 \text{ rpm s}^{-1}$
- 2.  $3000 \text{ rpm for } 30 \text{ s at } 300 \text{ rpm s}^{-1}$
- 3. Ramp down at  $300 \text{ rpm s}^{-1}$

Upon finish, the wafer was gripped outermost with wafer tweezers and soft-baked on a hot plate (super nuova+, Thermo Scientific) for 5 min at 65 ℃ and at least 10 min at 90 ℃. The optimal duration was determined if the gently touched resist did not stick to the tweezers. To prevent cracks in the resist caused by a fast temperature change, the wafer was cooled on the hotplate to room temperature. Such processed wafers were stored for a maximum of 4 weeks in a light-tight storage box.

To pattern the resist, the i-Line of a laser lithograph (Dilase 250, Kloe) was used. In preparation of the writing layout a AutoCADz \*.dxf-file with only one layer of polylines was imported to the program "Kloe Design", converted to contours and subsequently to polygons. For the filling a spot-size equivalent to the minimal structure resolution (as measured in Hicsanmaz [34]) and an overlap of at least 50 % was chosen. The writing trajectories were displayed for a last control before the export to ensure only closed contours. Finally the contour and filling were exported into separate files.

Both files were loaded in this order into the "Kloe Dilase" program. Also the preprocessed wafer was placed inside the laser writer and attached to the vacuumed stage. With the integrated camera the global zero was set to the wafer center by finding the horizontal or vertical edges and adding/subtracting the radius of the wafer (3"  $\approx \varnothing$  76.2 mm) intensities, writing speeds

#### 3.1.3. Soft Lithography

The fabricated wafer was placed the center of a 90 cm petri dish. A PDMS mold was created by vigorous mixing of the pre-polymer base with its curing agent (Sygard 184, Dowsil) in a ratio of 10:1 (w/w). For 3" wafers, thin channels were casted from 15 g, normal channels from 20 g PDMS in the petri dish. Gas bubbles were removed from the mixture in a desiccator for 20 min at 2 hPa, and the clear PDMS was cured in an oven (Um, Memmert) for 1 h at 60 ℃. After curing, the PDMS mold was released from the petri dish carefully, taken off the wafer and stored in a clean petri dish upon further processing.

#### 3.1.4. Bonding of Microfluidic

Under laminar flow, crosslinked molds were cut into pieces with the respecting single microfluidic ( $\mu$ F) with a razor blade. Holes for in- and outlet were punched through the containing channels with a biopsy puncher (ID 0.5 mm, WellTech). The substrates and  $\mu$ Fs were sonicated in acetone and deionized water (diH<sub>2</sub>O) for 5 min and dried with filtered nitrogen gas (N<sub>2</sub>) completely. For the bonding of PDMS to various substrates different protocols have been established:

#### **PDMS Glueing**

Here, a micron-height layer of uncured PDMS was used as an adhesive layer between  $\mu$ F and substrate. Approx. 3 mL were poured onto a 3" wafer and spun down for 5 min at  $6000 \, \mathrm{min^{-1}}$ . The microchannel was placed on the substrate by visual control of a stereo microscope (SMZ800, Nikon) with 8-fold magnification. Subsequently, the bonding process could be finished by a 1 h bake at 60 °C or over-night at room temperature.

#### **Plasma Bonding**

Due to the chemical nature of glass (or more generally oxides) and PDMS, the respective parts can be activated by the exposure to a controlled oxygen plasma which generates additional silanol (Si-OH) groups on their surfaces and removes impurities at the same time. Bringing the activated surfaces in contact triggers the formation of covalent bonds almost immediately. First, the acetone-wiped substrates and the microchannels were centered inside the plasma cleaner (Zepto, Diener). Second, vacuum was applied to a final pressure <0.2 hPa. Third, the chamber was flushed with pure O<sub>2</sub> until a chamber pressure from 0.7 hPa to 0.8 hPa had been stabilized. Fourth, the plasma process was executed with 30 W (Power-Poti: 100) for 45 s to 60 s (Time-Poti: 15-20). Upon

finish, the chamber was flushed for 5 s and ventilated. Immediately, the corresponding workpieces were brought into contact and pressed together gently. To ensure a durable bond, the assembled workpieces were baked for 1 h at 60  $^{\circ}$ C.

$$Here goes the mass flow equation$$
 (3.1)

#### **Reversible Bonding**

To bond the  $\mu$ F to a substrate reversibly and without residues, the channel can be brought into contact with the bottom part without any adhesinon agent. For low-pressure as well as vacuum driven flows, this method is preferrable due to its time and work efficiency.

#### 3.1.5. Electrical Circuit

Ground PCB Stacked PCBs with spacer

#### 3.1.6. Electronic Readout

test,test

#### **Hysteresis Alignment**

test,test

#### Single GMR

test,test

#### **Dual GMR**

one MFLI supplies both at same freugency. Aux Trigger tested, but no advantage.

### 3.2. Magnetic Beadometry

- 3.2.1. Standard Parameters
- 3.2.2. Concentration Measurement
- 3.2.3. Whole Blood Bead Spiking
- 3.2.4. Bead Capture Assay
- 3.2.5. Optical Particle Tracking
- 3.3. Surface Bio-Functionalization

## 3.4. Tensiometry

#### 3.4.1. Surface Activation

To functionalize any silicon containing surface with Si-OH groups which the utilized silane could interact with, multiple surface activation pathways were explored. First, substrates were cleaned in hydrochloric acid (HCI):methanol (MeOH) and  $H_2SO_4$  before they were immersed in boiling water. Second, surface silanol groups were achieved by piranha immersion and third by a HF dip.

For all methods, the following reagents were used:  $diH_2O$  (0.054  $\mu$ S, Merck MilliQ)), acetone (>99.9 %, VWR), ethanol (EtOH) (absolute, VWR), MeOH (99.8 %, VWR), acetic acid (AcOH) (glacial, VWR), HCl (37 %, Sigma-Aldrich), H<sub>2</sub>SO<sub>4</sub> (95 % to 98 %, VWR), H<sub>2</sub>O<sub>2</sub> (30 % (w/w), Sigma-Aldrich), HF (5 %, VWR)

#### **Work Safety Remarks**

Before the work with one of the acid solutions was carried out, serveral safety measures were implemented. As any diluted acid solution becomes very hot immediately due to the exothermic reaction, every container should be placed inside a cooled water or ice bath. Additionally, the beaker as well as concentrated acid flasks should be gripped firmly by a laboratory stand to avoid a tip over. As the reactivity of chemicals is highly temperature-dependent, the solutions was processed further when they had been cooled to  $\leq 70$  °C. It should be also noted that - as in every chemical reaction, but especially ones with H<sub>2</sub>SO<sub>4</sub> - the acid was always poured into the other reactant to avoid splashing and boiling.

#### **Plasma Activation**

Hier plasma protocolle raussuchen

#### **Hydrochloric-Sulfuric Acid Activation**

To degrease any glass or  $Si_3N_4$  surface, a protocol according to Dressick, Dulcey, Georger, *et al.* [35] was used. There, the surfaces were first sonicated in acetone and  $diH_2O$  for 5 min. Afterwards these were immersed in a 1:1 (v/v) solution of HCI:MeOH for >30 min, rinsed with  $diH_2O$  copiously and soaked in  $H_2SO_4$  for >30 min as well. Then, the samples were rinsed again in deionized water. To form silanol groups on the activated surface, the surfaces were finally immersed in >90 °C heated (SuperNuova+, Thermo Scientific)  $diH_2O$  for at least 2 h.

#### Piranha Activation

In this method, activation was carried out in a 1:5 (v/v) piranha solution at 70 ℃ for 30 min. After treatment, the samples were rinsed carefully with diH<sub>2</sub>O three times.

#### **Hydrofluoric Acid Activation**

#### 3.4.2. Chemical Surface Functionalization

Chemically activated surfaces were now coupled with APTES covalently. Therefore an aqueous silane solution was prepared from EtOH with volume fractions of 5% diH<sub>2</sub>O, 0.5% aqueous AcOH (pH 4.5) and 1% APTES in this order. The samples were soaked immediately after their activation in the silane solution. The reaction was carried out for 2 h to 4 h at >40%. At finish, all specimens were rinsed three times or sonicated for 5 min in absolute EtOH.

Then, the amine terminated surface modification was enhanced by a carbodiimide conjugation with Poly(acrylic) Acid (PAA). As above, a reaction consisting of 0.5 M 2-(N-morpholino)ethanesulfonic acid (MES) buffer with 1 mg mL<sup>-1</sup>, 6 mM EDC and 3 mM NHS was activated for 15 min on a magnetic stirrer. Subsequently, the prepared samples were immersed in the solution for 1 h on a rotation shaker (VWR). As final cleaning, the slides were rinsed or sonicated for 5 min in diH<sub>2</sub>O and stored in fresh diH<sub>2</sub>O at 4 °C up to 14 d upon further use.lib:chem:Anti-EpCAM-PAA

#### 3.4.3. Surface Bioconjugation

#### 3.4.4. Particle Functionalization

# 4. Results

test,test

## 4.1. Virtual Prototyping of Cell Signals

Signal Similarity For Cells With Varying Bead Coverages

Cross-Correlation between single dipole with sum magentic moment and surface covered with randomly distributed magnetic particles

#### 4.1.1. Single Cell Signal

#### 4.1.2. Cell Aggregates

# 4.2. Reference Bead Surface Functionalization

#### 4.2.1. Amine-Surface Biotinylation

Streptavidin-Atto488 reference calibration Anti-Biotin-PE working? BNF-Dextran-Streptavidin unspecific binding?

Figure 7: Amine bead modification with Sulfo-NHS-Biotin

An amine terminated bead is incubated with sulfo-NHS-Biotin to cover its surface by amide-Biotin. As byproduct the sulfo-NHS-ester 1-hydroxy-2,5-dioxopyrrolidine-3-sulfonate splits off.

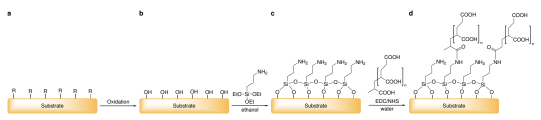


Figure 8: General process chain of chemical surface modification

Any substrate with various surface groups R (a) is oxidized to exhibit hydroxyl groups.(b). Then a silane SAM is attached (c) and subsequently modified by carbodiimide chemistry with PAA. (d)

#### **Magnetic Polystyrene Bead**

# Non-Magnetic Polystyrene Bead

## 4.2.2. Carboxy-Surface Biotinylation

# 4.3. Concentration Measurements in MRCyte

## 4.3.1. Count Stability

Measurement over 1h Measurement of Syringe Tubing Losses

4.3.2. Calibration of Flow Field

4.3.3. Differential Counting Setup Sensitivity Calibration Concentration Measurements

# 4.4. Protein Immobilization On The Microfluidic Channel Bottom

# 4.4.1. Physisorption

Quantification in Plate Reader Trial with Neutravidin + Sensor (Esthis Versuch)

#### 4.4.2. Covalent Attachment

# Plasma-Based Approach Water-Based Approach

Sonicate in Acetone and Water 5' 1:1 HCI:Methanol  $H_2SO_4$  Treat for 30 min in light boiling water

# 5. Discussion

test,test

Contact angle for silanization of surface methods more useful -> should be 1st approach for characterization

Anti-Biotin-PE working? BNF-Dextran-Streptavidin unspecific binding?

# 6. Outlook

# List of Abbreviations

# Symbols

au - surface stress tensor
$\eta$ - dynamic viscosity
$\mu$ F - microfluidic
ho - density
$\sum_i \mathbf{f}_i$ - body forces
A
AAF - artificial Anti-Ferromagnet
AcOH - acetic acid
AFM - Anti-Ferromagnetism
amineNH <sub>2</sub>
APTES - (3-aminopropyl)triethoxysilane
C
carboxamide - R <sub>1</sub> - CONH - R <sub>2</sub>
carboxylCOOH
D
diH <sub>2</sub> O - deionized water
E
EDC - 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide
ethoxyO-CH <sub>2</sub> -CH <sub>3</sub>
EtOH - ethanol
F
FM - Ferrimagnetism
FWHM - full width at half maximum
G
GMR - giant magneto resistance
н

$H_2O_2$ - hydrogen peroxide
$H_2SO_5$ - Caro's acid
H <sub>2</sub> SO <sub>4</sub> - sulfuric acid
HCI - hydrochloric acid
HF - hydrofluoric acid
nydroxylOH
1
PA - isopropanol
M
MACS - MACS running buffer
MeOH - methanol
MES - 2-(N-morpholino)ethanesulfonic acid
methyleneCH <sub>2</sub>
MNP - magnetic nanoparticle
N
$N_2$ - nitrogen gas
NFM - non-ferro-magnetic
NHS - N-hydroxysuccinimide
0
$O_2$ - oxygen gas
P
PAA - Poly(acrylic) Acid
PBS - phosphate buffered saline
PCB - printed circuit board
PDMS - poly(dimethyl siloxane)
PM - Paramagnetism
S
SAM - self-assembled monolayer
Si <sub>3</sub> N <sub>4</sub> - silicon nitride
silanol - Si_OH

siloxane - Si-O-Si
SMA - styrene maleic anhydride
SPM - superparamagnetism
sulfo-NHS - N-hydroxysulfosuccinimide
U
u - flow field

# List of Figures

1	Syringe Pump error sources
	Set flow rate: —, Real Flow Rate: — a, Transient step answer of a
	syringe pump through a microtube with 254 $\mu m$ inner diameter. $\boldsymbol{b}$ , Steady
	state flow rate error around the desired $5\mu L\text{min}^{-1}$ dispensing rate. A
	sinusoidal behaviour caused by the microstepping can be observed. [6] 7
2	Different substrate surfaces: glass and PDMS
	Surface groups and internal structure of quartz glass (a) and PDMS (b).
	After an oxidation step, the methyl groups are changed to hydroxyl 10
3	Proposed modification of $\mathbf{Si}_{3}\mathbf{N}_{4}$ with HF
4	Trialkoxysilane
	Structure of a typical trialkoxysilane, X: hydrolyzable group, R: non-hydrolyzable
	organic radical, n: methylene chain-length
5	APTES Modification of an oxidized surface
	a Before the condensation reaction, the oxidized surface forms hydrogen
	bonds with water molecules. The silane molecules are in the bulk solu-
	tion. <b>b</b> The hydrolyzed silanol group adsorbs onto the surface and forms
	hydrogen bridges with it. ${\bf c}$ In a condensation reaction, under the loss of
	water, a covalent bond to the surface forms. <b>d</b> After the SAM assembly
	the surface is saturated with a covalent-bound, crosslinked silane film. [26] 12
6	Carboxyl bead modification with EDC/NHS
	The carboxy groups bead are activated with EDC to an active O-acylisourea
	intermediate. This can then either be nucleophilicly attacked by a primary
	amine of the amine-PEG2-biotin reactant or - due to its instability - hy-
	drolzed back to a regenerated carboxyl surface. A present NHS-ester
	can also displace the O-acylisourea to form a considerably more stable
	intermediate which then itself reacts with any primary amine

7	Amine bead modification with Sulfo-NHS-Biotin					
	An amine terminated bead is incubated with sulfo-NHS-Biotin to cover its					
	surface by amide-Biotin. As byproduct the sulfo-NHS-ester 1-hydroxy-					
	2,5-dioxopyrrolidine-3-sulfonate splits off	2				
8	General process chain of chemical surface modification					
	Any substrate with various surface groups R (a) is oxidized to exhibit					
	hydroxyl groups.( $\mathbf{b}$ ). Then a silane SAM is attached ( $\mathbf{c}$ ) and subsequently					
	modified by carbodiimide chemistry with PAA. ( <b>d</b> )	2				

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## Statement

I declare that I have authored this thesis independently, that I have not used other than the declared sources / resources, and that I have explicitly marked all material which has been quoted either literally or by content from the used sources.

Munich, December 4<sup>th</sup>, 2020, Signature