

## **Nanocytometer for smart analysis of peripheral blood and acute myeloid leukemia: a pilot study**

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1 **Nanocytometer for smart analysis of peripheral blood and acute myeloid leukemia:**  
2 **a pilot study**

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23

24 **Abstract:** We realize an ultra-compact nanocytometer for real-time impedimetric detection and classification of  
25 subpopulations of living cells. Nanoscopic nanowires in a microfluidic channel act as nanocapacitors and measure  
26 in real time the change of the amplitude and phase of the output voltage and, thus, the electrical properties of living  
27 cells. We perform the cell classification in the human peripheral blood (PBMC), and demonstrate for the first time  
28 the possibility to discriminate monocytes and *subpopulations* of lymphocytes in a label-free format. Further, we  
29 demonstrate that the PBMC of acute myeloid leukemia and healthy samples grant the label free identification of  
30 the disease. Using the algorithm based on machine learning, we generated *specific data patterns* to discriminate  
31 healthy donors and leukemia patients. Such solution has the potential to improve the traditional diagnostics  
32 approaches with respect to the overall cost and time effort, in a label free format, and restrictions of the complex  
33 data analysis.

34 **Keywords:** impedance cytometer, nanosensor, POC diagnostics, PBMCs, acute myeloid leukemia (AML),  
35 machine learning for data treatment

36

37 Healthcare of tomorrow will be dramatically affected by global processes that take place today, like societal  
38 shifts<sup>1</sup>, technological and digital revolution<sup>2,3</sup>. One of the main challenges within the healthcare sector is to establish  
39 new patient-care standards, based on *e.g.* new drug administering<sup>4</sup>, novel ultrasensitive diagnostics integrated into  
40 the gadgets<sup>5</sup>, to provide maximally personalized tests and doctor advices<sup>6</sup>. Medical data for patients will double  
41 every 73 days by 2020<sup>7</sup>. Taking into account the trends towards personalization in medicine, patient related data  
42 can reach millions of gigabytes during the lifetime<sup>8</sup>. To make this information serving its aim to improve the quality  
43 of care while controlling the costs, these data have to be analyzed using *conventional* and *unconventional*  
44 *algorithms*, involving elements of machine learning. This strategy helps to fully access and interpret information  
45 on demand using *e.g.* modern gadgets, connected to a cloud. Thus, artificial intelligence is now rapidly entering the  
46 medical sector. Ideal proof of concept realization of diagnostic devices combining the new technological trends  
47 with the novel data treatment protocols would be a nanoscaled sensor device, for *e.g.* cancer diagnostics,  
48 accompanied with algorithms involving machine learning elements to distinguish proper trends within the large  
49 amount of noisy data points. The development of such systems is currently in the emerging phase<sup>9,10</sup>, due to the  
50 number of existing technological challenges, *e.g.* reaching the stable performance of nanosensors as well as its  
51 current disintegration with the IT sector.

52 The primary goal of the current work is to show that all prerequisites for the development of a *nanobiosensor*  
53 *system combined* with a *smart analytical algorithm* to interpret the results can be achieved.

54 Leukemia is one of the common forms of blood cancer, affecting the production of white blood cells<sup>11</sup>, diagnosed  
55 in 352,000 people and caused 256,000 deaths worldwide in 2014<sup>12</sup>. Acute myeloid leukemia (AML) is the most  
56 frequent type in adults, with around 30% of all detected leukemia cases and relatively low five-year survival rate  
57 of 20-30%, strongly dependent on the age of the patient<sup>13,14</sup>. Diagnosis of AML is multidimensional<sup>15-19</sup>, including  
58 examination of blood by flow cytometry. More specifically, optical flow cytometry makes a big impact in blood  
59 cancer diagnostics<sup>20,21</sup> and evaluation of the immune response of the patient<sup>22,23</sup> via analysis of peripheral blood  
60 mononuclear cells (PBMCs)<sup>24-26</sup>. For a complete qualitative and quantitative detection of blood cancer<sup>26</sup> in PBMCs,  
61 the main immune cell subpopulations have to be distinguished, exploiting the *clusters of differentiation* (CD)  
62 responsible for cell surface marker expressions. Conventional flow cytometers rely on the use of specific molecular  
63 labels, *e.g.* monoclonal antibodies against cell surface markers<sup>27</sup>. Finally, a combination of this and above  
64 mentioned techniques rises the diagnostics costs of cancer up to hundreds of dollars per person<sup>28</sup>.

65 On-chip integrated nanodevices have emerged as a new generation of biodetectors<sup>29-38</sup>. A promising approach  
66 relies on measuring electrical signals, *e.g.* impedance. For the latter, *static* (electrical impedance spectroscopy  
67 (EIS)<sup>39-43</sup>) and *dynamic* (impedance cytometry<sup>44-47</sup>) modes of impedance detection are proposed. The latter one is  
68 performed at fixed frequency and is used to increase an analytic and information processing throughput. From the  
69 conceptual introduction of micro-Coulter counters<sup>44,48</sup>, impedance cytometry has evolved and strengthened its  
70 impact in biological contexts for single cell detection<sup>45,46</sup>, investigations of erythrocytes<sup>49,50</sup>, eukaryotes<sup>51,52</sup> and  
71 protozoa<sup>53</sup> (Figure 1 A).

72 The scientific community accepts that scaling down of the sensor dimensions boosts the sensitivity of common  
73 detection techniques. Despite the tremendous success of *e.g.* nanoscaled bio-FETs<sup>54</sup>, the sensitivity issue of  
74 impedance detectors and its possible improvement via cross-scale integration of the nanostructures, have not yet  
75 been addressed. All current realizations of impedance dynamic sensors are characterized by macro- to micro-  
76 dimensions, employing metal microscopic electrodes in a fluidic channel<sup>49,50,55,56</sup>. Proof-of-concept realizations of  
77 such devices are limited to detection of inorganic particles and isolated and purified/treated eukaryotic cells<sup>56-58</sup>  
78 with very few examples demonstrating the realistic systems, *e.g.* purified or diluted blood<sup>59</sup>, typically used for  
79 clinical diagnostics.

80 Here, we present a nanosensor system, combining an ultra-compact impedance flow cytometer to analyze  
81 complex cell compositions with a software, based on conventional machine-learning algorithms, to interpret the  
82 measured data via exploiting the classification of cell subpopulations and respective clusters of differentiation (see  
83 Figure 1 B). Utilizing the term “nanocytometer”<sup>60</sup>, we work with a nanosensor that employs the interdigitated pairs  
84 of gold nanoelectrodes to reach the substantial increase of the sensitivity<sup>36,61</sup>, compared to the micron structures.  
85 We study untreated human PBMCs from healthy volunteers (Figure 1, C and D) and AML patients, and demonstrate  
86 significant differences in data patterns of healthy PBMC and AML samples (see Table S1 in **Supporting**). Thanks  
87 to the enhanced sensitivity of the device, we show the discrimination of the cells subpopulations in a label free  
88 format, *e.g.* B-, T, NK cells and myeloblasts that before was possible only using fluorescent biomarkers. The  
89 software processes the output voltage and phase signals measured by the detector in a multistep manner, followed  
90 by a final data clustering using the k-means algorithm. Fabrication of gold nanowire arrays is summarized in Figure  
91 2, A.I-III and C, and detailed in **Materials and Methods in Supporting Information**. The resulting cytometer  
92 devices possess 1 (sensor area  $\sim 46 \mu\text{m}^2$ ), 6 ( $\sim 506 \mu\text{m}^2$ ) and 18 pairs of gold nanowires ( $\sim 1610 \mu\text{m}^2$ , see calculations  
93 in **Supporting Information**) with the width of about 100 nm each. To optimize the sensor geometry, COMSOL  
94 simulations were carried out to reach the situation of a homogeneous electric field between the nanowires. This  
95 electric field is also enhanced (**Supporting Information S1-S3**), compared to the geometry without nanowires.  
96 The optimal nanowire configuration was found at a distance of 2  $\mu\text{m}$  from the nanowire tips to the opposing  
97 microelectrode pad, with a pitch about 1  $\mu\text{m}$  (Figure 2 B, and **Supporting Information S2**). In order to demonstrate  
98 the effect of a 2 $\mu\text{m}$  silica particle on the spatial distribution of the electric field and its enhancement near the  
99 nanowires, simulations were carried out in *yz*- and *xz*-planes. (Figure 2 B (*yz*-plane) and **Supporting Information**  
100 **S4**). Detailed comparison of the geometry and sensitivity characteristics of the nanocytometer with reported  
101 impedance sensor devices is provided in the Table S2 in **Supporting information**.

102 Next, a PDMS-based 3D flow-focusing system (Figure 2 C), confining the analyte in the middle and bottom of  
103 the channel (height 15  $\mu\text{m}$ , width 200  $\mu\text{m}$ ) close to the sensor (Figure 2 D, **Supporting Figure S5** for efficiency of  
104 the hydrodynamic focusing), was realized. Measurements were carried out with a lock-in amplifier (eLockIn205/2,  
105 Anfatec) for a direct readout of the signal. Flow rates were actively manipulated (0.1  $\mu\text{l}/\text{min}$  – 2.5  $\mu\text{l}/\text{min}$ ) using a  
106 syringe pump (neMESYS 290N, Cetoni) for injecting a sample solution (particles and cells solution, as well as

107 vertical and lateral focusing streams (100  $\mu\text{M}$ , KCl). The chip was measured under the microscope (Axiovert200,  
108 Carl Zeiss Microscopy) for complementary observations. With respect to the following analysis of *e.g.* peripheral  
109 blood, measurements were typically performed with the average cell rate of around 3-5 cells/s at 0.5  $\mu\text{l}/\text{min}$  (see  
110 **Supporting information S6**). The electrical characterization was carried out in both direct (DC) and alternating  
111 current (AC) modes to evaluate the equivalent circuit of the system and is summarized in **Supporting Information**  
112 **S7** and **S8**. The sensing device (*e.g.* 18 NW) exhibits a capacitive behavior with a characteristic butterfly shape in  
113 DC voltammetry (**Supporting Information S7**), also confirmed by a Nyquist diagram. Living cells, that cross the  
114 sensing area, cause a local alteration of the dielectric properties of the medium around the nanocapacitor, causing  
115 an instantaneous modulation of the equivalent circuit and its complex impedance.

116 Next, we compare three above fabricated nanocytometers to the reference nanowire-free microelectrodes  
117 (distance between microelectrodes - 50  $\mu\text{m}$ , width of pads - 35  $\mu\text{m}$ ) to investigate the enhancement of the sensitivity  
118 and the signal dispersion, depending on the sensor dimensions. We used silica colloidal particles and peripheral  
119 blood samples as reference objects. First, a single 10  $\mu\text{m}$  bead was placed onto the sensing area of all types of  
120 devices by micropipetting to investigate the EIS signature between 50 Hz and 20 MHz. Dielectric particles deform  
121 the semi-circle in the Nyquist diagram (Figure 3, A-B) via adding a particle related *serial RC-element* (accounting  
122 for the particles resistance and capacitance), connected in parallel to the initial RC-circuit. Based on Maxwell model  
123 for dielectric mixtures, the effect of particles and cells on the impedance signal is described using single shell  
124 models. Considering RC-like properties of the sensor, a particle or cell adds its capacitances and resistances of the  
125 membrane and cytosol. In the simplified model, cell membrane conductance and cytosol capacitance are ignored,  
126 resulting in a parallel addition to the RC-circuit of an in-series cytosol resistance and membrane capacitance<sup>62</sup>. We  
127 observe that devices with nanofabricated electrodes, possessing a single pair of nanowires, revealed stronger  
128 modulation of the amplitude and phase signals than both multiwire and microelectrode-based sensors (factor of 23  
129 for microelectrode geometry) (see Figure 3 A and B). Thus, enhancement of the electric field between the nanowires  
130 boosts the sensitivity of devices towards micro-objects, *e.g.* colloids and living cells. This statement is confirmed  
131 by cytometer-like measurements of 10  $\mu\text{m}$  large silica particles in 0.1x phosphate buffered saline (PBS), performed  
132 at 100 kHz using devices with 1, 6 and 18 nanowire pairs (Figure 3 C, D). Here, the solution of particles is injected  
133 into a microchannel, focused by streams of 100  $\mu\text{M}$  KCl and guided towards the sensors. In the following, the  
134 change of the amplitude  $\Delta V_{\text{out}}$  and the phase  $\Delta\text{Phase}$  of the output signal compared to the background, when a  
135 particle (or cell) is crossing the active area of the sensor, is evaluated for each detection event. The results are  
136 presented as clusters, depicting  $\Delta\text{Phase}$  (*y* axis) versus the  $\Delta V_{\text{out}}$  (*x* axis) of the output signal (**Supporting**  
137 **Information S9-S11** for details). Such representation of detection events allows us to compare not only output  
138 *signal modulations* but also the *dispersion* of the signal, measured by different devices. The devices with the  
139 smallest area reveal highest signal deviation, but they are prone to higher signal dispersion due to the stronger  
140 influence of the spatial location of the particle with respect to the electrodes (Figure 3, D). The sensitivity of the

141 device in arbitrary units and with respect to *resistance and capacitance changes per particle* is calculated for  
142 different sensor dimensions (**Supporting Table S2 and S12-S13**).

143 This result is a direct *fundamental consequence of the nanoscopic scaling effect* that makes a great impact in the  
144 field of nanobiosensorics. Indeed, the miniaturization of the detector size down to the dimensions of the analyte,  
145 boosts its sensitivity on one hand<sup>62</sup>, but unavoidably leads to an increase of the signal to noise ratio. We further  
146 apply a nanoscaled cytometer with 6 pairs of interdigitated nanowire electrodes for analysis of blood and  
147 diagnostics of AML by identifying human PBMC subpopulations with particular interest to classify the  
148 subpopulations of the cells within PBMCs in label-free format (*e.g.* monocytes, T-cells, B-cells, NK-cells<sup>63,64</sup>,  
149 myeloblasts). PBMCs of healthy donors are represented by subpopulations of peripheral cellular blood components  
150 exhibiting a round nucleus and visible granules<sup>65</sup>, consisting of monocytes (CD14) and lymphocytes which can be  
151 additionally divided into T cells (CD3<sup>+</sup>), B cells (CD19/CD20) and natural killer (NK) (CD16/CD56) cells<sup>63</sup>. In  
152 turn, the peripheral blood smear from the AML patients is highly probed with undifferentiated myeloid progenitor  
153 cells, the myeloblasts (CD34<sup>+</sup>/CD123).

154 First, we realize the measurements of PBMCs in order to determine the specific *calibration pattern*, peculiar for  
155 the impedance nanocytometer. The fresh human blood from healthy male donor and AML patient was purified  
156 using standardized Ficoll protocol (ratio 1:1) and resuspended in PBS buffer for measurements. This unified  
157 protocol has been applied to all further measurements, including impedance and conventional cytometry. Further,  
158 the traditional FACS technique was employed to sort the labeled subpopulations of PBMCs into separate vials (*i.e.*  
159 monocytes, B-, T-, NK- cells for healthy donor, and myeloblasts for AML positive patient) for *calibration*  
160 measurements (**Supporting Information S14**). The output potential  $\Delta V_{\text{out}}$  and phase shift  $\Delta\text{Phase}$  were acquired  
161 by applying a sinusoidal reference signal with an amplitude of 0.5 V and a frequency 500 kHz.

162 Next, we placed the microfluidic chip under the fluorescent microscope to perform parallel impedimetric and  
163 microscopy measurements. We did the calibration for healthy samples injecting each cells subpopulations one-by-  
164 one (1 - T-cells, 2 - NK cells, 3 - B cells, 4 - Monocytes), and *repeated them in random order sequence* (see 5 -  
165 NK-cells, 6 - T-cells, 7 - B-cells, respectively), to prove the fact of the signal differentiation and absence of drift  
166 in the system (Figure 4 A, different colors for coding each cell subpopulation). This data sequence resulted in the  
167 clusters of differentiations for each of the detected subpopulations (Figure 4, B-G). Measurements of myeloblasts  
168 are performed in similar manner and are summarized in Figure 4 L-N. Resulting cloud of the myeloblasts data is  
169 plotted in Figure 4 M (red circles) and converged with the whole data pattern of labeled PBMC of the AML patient  
170 (black circles) measured by the nanocytometer for localization of the subpopulation of malignant cells. Analysis of  
171 the whole datasets determines a *calibration pattern* (Figure 4 H - healthy and M, N - AML, **Supporting S15**).  
172 Afterwards, both *labeled* and *unlabeled* PBMC mixtures of a healthy volunteer (Figure 4 J and K, respectively) and  
173 AML patient #2 (Figure 4 M and N) were matched to compare with the aforementioned *calibrations* to fine-tune  
174 the thresholds for data clustering. Raw samples and calibration patterns match well at the level of the pattern shape,  
175 while normalization is needed to compare between labeled/unlabeled samples (**Supporting S16**). Normalization of

176 the AML data plot in the range [0, 1] enables to match the data patterns of myeloblasts (red circles), unlabeled  
177 samples of AML (black circles) and healthy PBMC (green circles, Figure 4 N). Interestingly, analysis of healthy  
178 PBMC (green) and myeloblasts (red) shows *additivity* of both patterns. Thus, we believe that the isolated labeled  
179 PBMC subpopulations of healthy and AML positive patients can serve as a valid guideline for impedimetric  
180 measurements of unlabeled PBMC samples.

181 Note that the discrimination between PBMC cells according to their dielectric properties has been predicted  
182 around two decades ago<sup>66,67</sup>. Natural reason is that the membrane surface of immune cells is not even, and its  
183 textures is related to the cells function<sup>68-70</sup>. Still, discrimination of *unlabeled lymphocytes at single cell level* was  
184 not demonstrated by now. We attribute successful discrimination of the lymphocyte cells in this work to the  
185 essentially increased sensitivity of the nanoscopic cytometer device (**Table S2**, PBMC measured by different  
186 sensors in **Supporting Information S16**).

187 Next, we strengthen the classification of the PBMCs by proportion analysis of all measured cells. All together 5  
188 samples from healthy volunteers were studied (4 male and 1 female, age 25-35 years, **Supporting Table S1**).  
189 Calibration patterns are used for determination of the clusters of monocytes and subpopulations of the lymphocytes  
190 within the solution (Figure 4 H-K, Figure 5 A (inset), F, and **Supporting Figures S16, S17**). Thus, the  
191 subpopulations of T cells (62.31%, purple), B cells (31.34%, green) and NK cells (7.34%, red) could be  
192 distinguished (Figure 5 A, inset). These percentages are in agreement with the proportions of cells within healthy  
193 human PBMCs predetermined via FACS (**Supporting Information S17**), deviating within 1-3% only (table in  
194 Figure 5 D).

195 Finally, for analysis of AML positive cases, all together 3 patients (2x female, 1x male) were tested, using a small  
196 sample volumes, compared to regular assays in the clinical practice<sup>71,72</sup> (~5  $\mu$ l). Blood from AML Patient#1  
197 (female) and a healthy donor (female) was taken at the same day for comparison (Figure 5 A, and inset in A).  
198 Further, 2 samples were analyzed additionally (**Supporting Figure S14** for FACS). Sensors with 18 pairs of  
199 nanowires (AML Patient #1) and 6 pairs (AML Patient #2 and #3) were utilized for these studies. All AML data  
200 were analyzed manually and compared to the calibration (Figure 4M) and healthy reference, measured earlier. An  
201 additional large data cluster was identified in all samples (black circles, Figure 5 A). We attribute it to the  
202 myeloblasts that account for 34.16% ( AML #1), 60,07% (AML #2) and 54,96% (AML #3) (Figure 5 D, and  
203 **Supporting Information S18, S19**). Results are in agreement with the proportions of myeloblasts cells, provided  
204 by flow cytometer analysis (**Supporting Information S14**), and are comparable to the data provided by World  
205 Health Organization (WHO)<sup>75</sup>. Further details on cell proportions and merged AML#2-AML#3 data are given in  
206 **Supporting S18**.

207 AML#1 raw data were additionally analyzed with the developed software for classification of cell subpopulations  
208 for comparison<sup>73,74</sup>. Algorithm for clustering of the of PBMC cells was divided into four subparts: signals baseline  
209 estimation, baseline subtraction, interquartile range analysis for peak detection and coupled peaks clusterization.



210 **(Supporting Information, Methods)**. Additional data cluster in the scatter analysis was identified with excellent  
211 precision, which can assist in the *pattern based disease diagnostics* (see Figure 5, B-C).

212 In conclusion, we demonstrate an ultra-compact impedance flow nanocytometer combined with software  
213 employing the conventional machine-learning algorithms. We successfully apply this system for the discriminative  
214 analysis of healthy PBMC (B-, T-, NK-cells and monocytes), as well as discrimination of PBMCs of leukemia  
215 patients, using extremely low sample volumes in a short time. The developed platform can contribute to the modern  
216 clinical diagnostics assays as a miniaturized, reusable, easy in operation tool, with an option of an autonomous  
217 analysis. Due to small dimensions of each individual sensor, many detectors can be integrated on a chip that paves  
218 a way towards a new type of miniaturized bio-analytics. Namely, the cytometer measurements format coupled to  
219 the smart data treatment opens a route towards the realization of a platform for the rapid *detection* and *recognition*  
220 of a broad spectrum of *e.g.* blood-related (**Supporting S20**) and immune system diseases. The task of the software  
221 is in the utilization of learning algorithms to train the network for the recognition of *multiple data patterns*,  
222 indicating different diseases, and used for the diagnostics of multiple patients. As the data complexity increases  
223 dramatically in this case, we envision the evolution of the signal treatment methods, *e.g.* towards deep learning  
224 approaches<sup>78</sup>.

225

## 226 **ASSOCIATED CONTENT**

227 **Supporting Information available online:** Supporting information contains an overview of datasets of healthy  
228 and AML-diagnosed donors, comparison of sensing areas in state-of-the-art impedance cytometry geometries,  
229 COMSOL simulations on sensor prototyping and electric field between optimized nanowire structure, electric field  
230 behavior in the presence of a particle, 3D Hydrodynamic Focusing Calculations, PBMC detection with various flow  
231 rates, electrical Characterization in AC and DC, sensitivity towards several numbers of analytes, settling time of a  
232 10µm particle on the sensor, scatter plot pairing and calculation, coefficients of variation (CV) for various sensing  
233 areas, phase shift and reference curve for resistance and capacitance shift, resistance and capacitance changes per  
234 particle, FACS analysis on subpopulations of healthy PBMCs and AML-diagnosed PBMCs, PBMC sub-population  
235 classification, PBMC detection with various sensing sizes and frequencies, manual analysis of PBMCs of healthy  
236 and AML-diagnosed donors, full-length Experiment, probing SiO<sub>2</sub> and PBMC mixture, signal modulation using  
237 silica microspheres of several diameters, detection Area calculation, detailed material and methods, code for data  
238 analysis.

239

## 240 **AUTHOR INFORMATION**

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243 **Author Contributions**

244 J.S, S.K. conducted experiments and simulations under supervision of L.B and G.C. E.A., M.B., A.F., and M.B.  
245 L.G.D. and M.R., J.M.M. and K.S. contributed with biological samples from their institutes. L.B, and G.C oversaw  
246 the research in their groups. The manuscript was written by L.B and J.S with input from E.A, L.G.D and M.R, S.K.  
247 G.M, developed the software for data analysis. All authors co-wrote the paper and agree to its contents.

248 **Notes**

249 The authors declare no competing financial interest.

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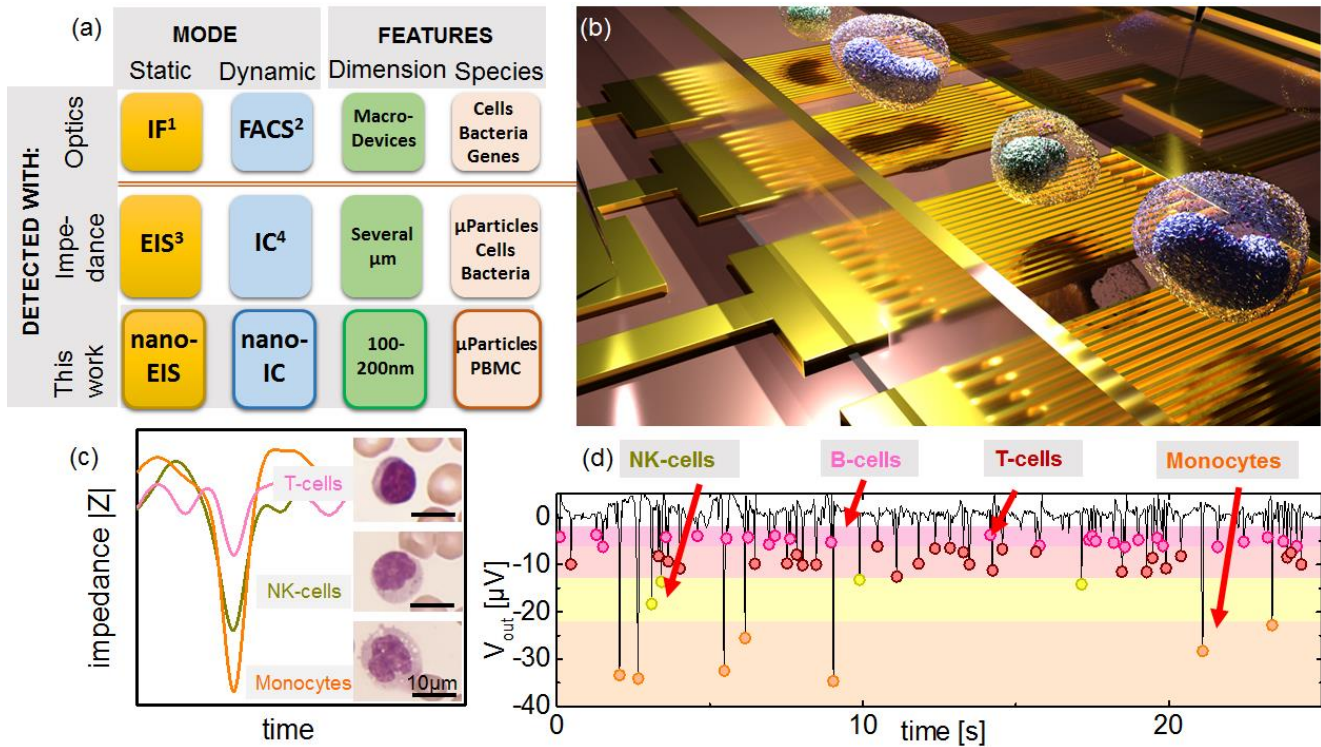
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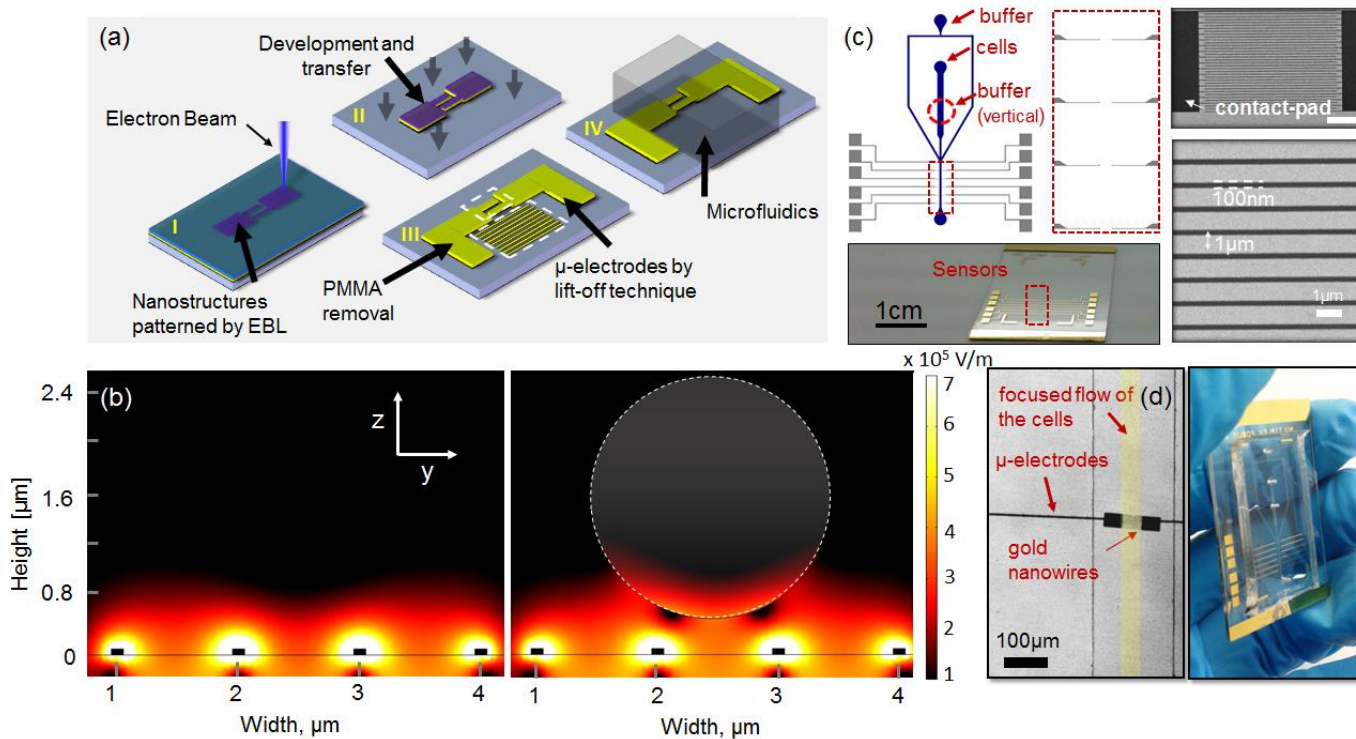
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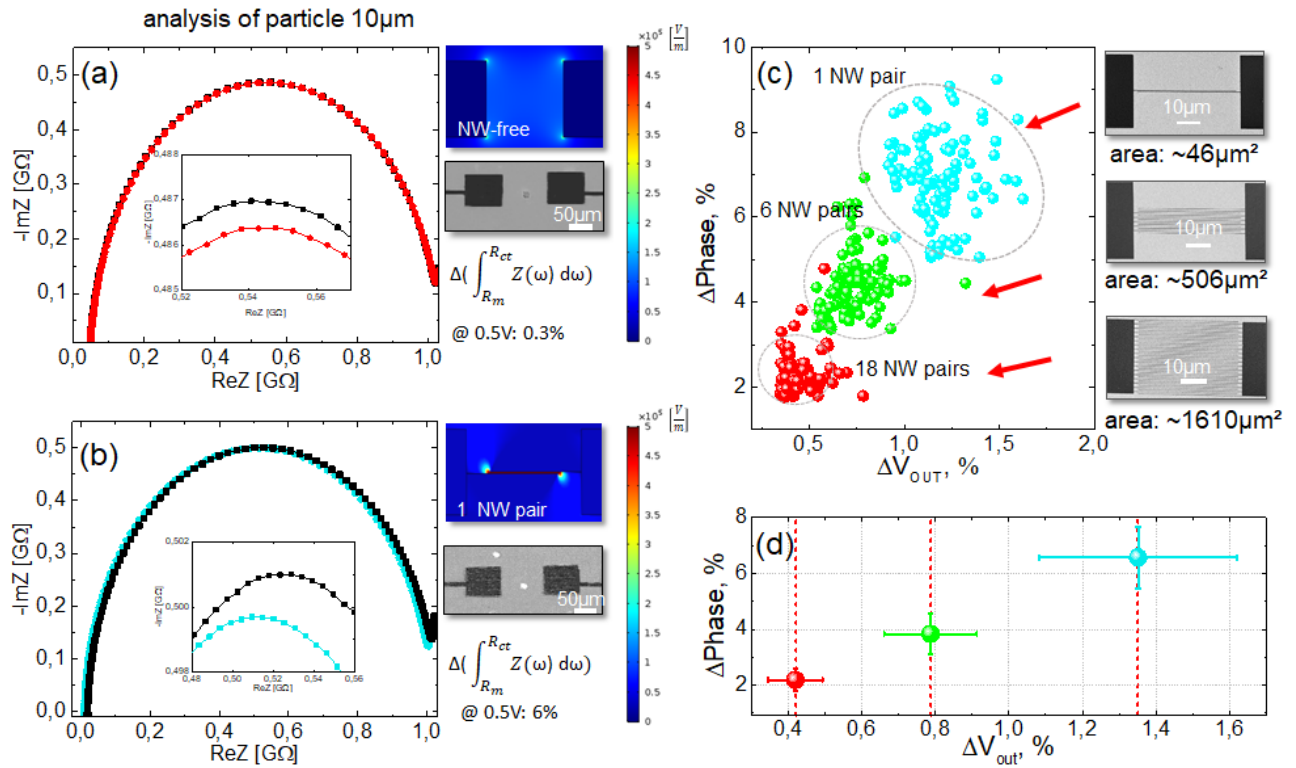
## Figures



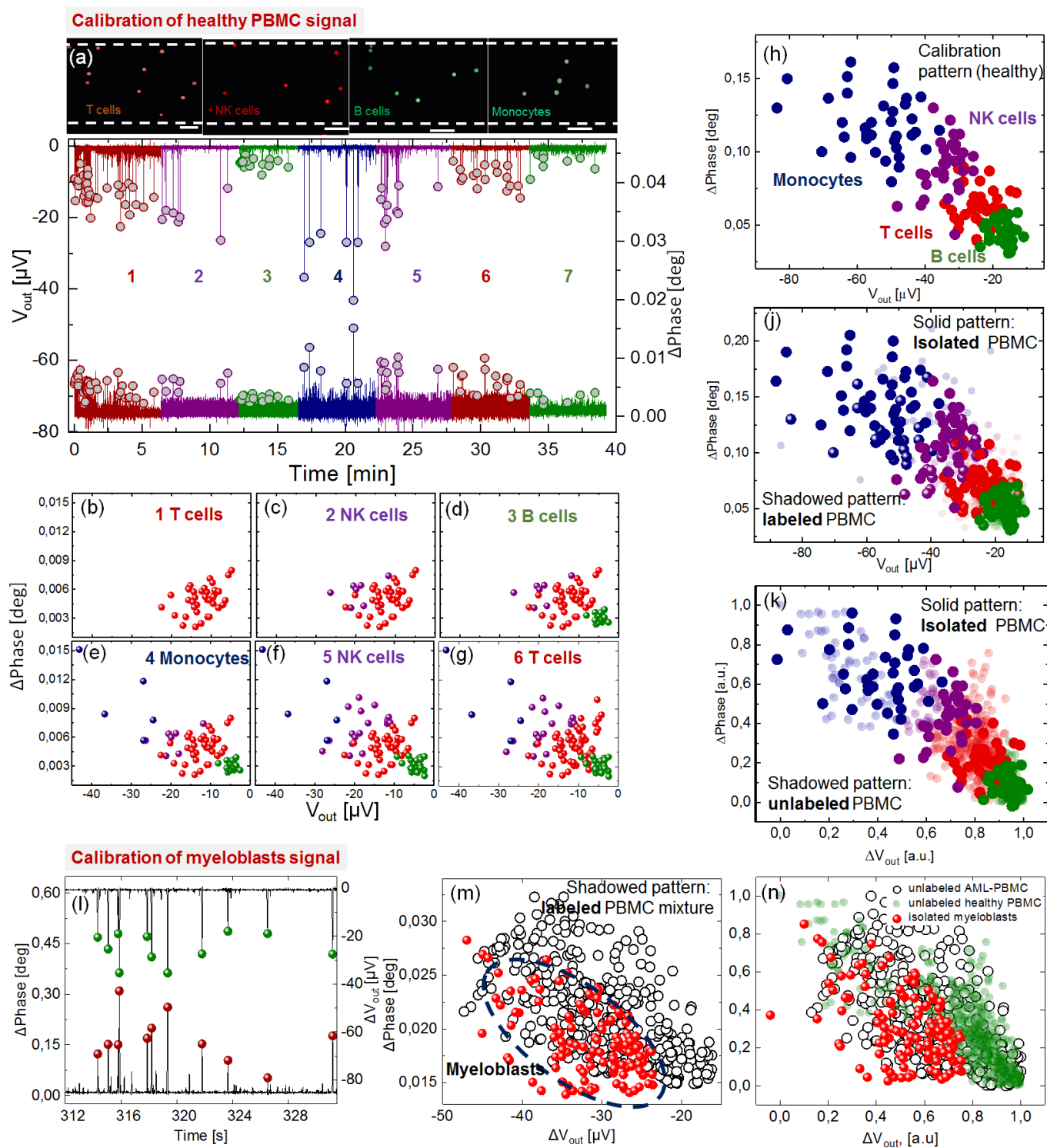
**Figure 1:** Conceptual figure describing the developed sensor platform. **(a)** Contribution of the nano-sensor platform to previously reported and state of the art techniques, *i.e.* immunofluorescence (IF)<sup>80</sup>, fluorescence-activated cell sorting (FACS)<sup>81</sup>, electrical impedance spectroscopy (EIS)<sup>47</sup> and impedance cytometry (IC)<sup>64</sup>. **(b)** Schematically illustration of PBMC detection by nano-impedance cytometry **(c)** Comparison of signal magnitudes of PBMCs with different diameters. **(d)** Real-time output response of the sensor with complex mixture of PBMCs.



**Figure 2:** Fabrication and integration of the nanoscaled sensor chip. **(a)** Nano-impedance cytometer fabrication process. **(b)** COMSOL Multiphysics simulation of the electric field perturbation in presence of a dielectric microparticle in the microfluidic channel. **(c)** Layout of the nano-sensor array of 6 independent accessible electrode pairs approaching the contact pads of the EBL-patterned design. The main channel of the microfluidic geometry is placed to incorporate the electrodes. **(d)** 3D hydrodynamic focusing technique allowing analyte guidance in the middle and at the bottom of the channel.



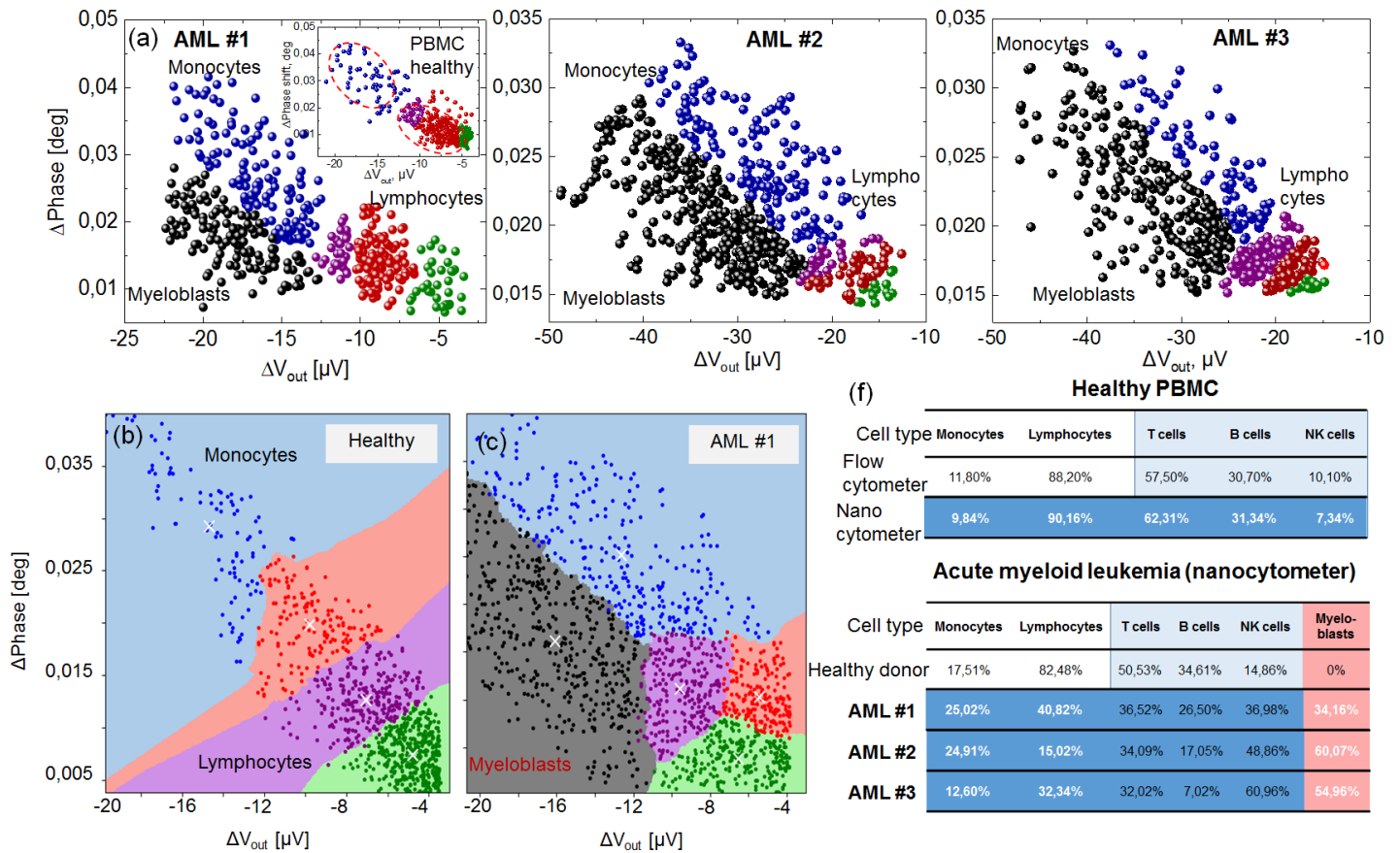
**Figure 3:** Comparison of the signal change with and w/o present micro-particle in the sensing area in static mode. While (a) only micro-electrodes grant a weak electric field and thus have a small signal change when a particle is present (0.26%), introduction of 16 interdigitating nanowires (b) and single nanowire pair. (c) cytometer mode summary: signal modulation while detecting 10  $\mu\text{m}$  particle, using different sensor dimensions. (d) Calculated change of device output signal in dependence of sensor dimensions, from (c).



**Figure 4:** Detection and classification of isolated PBMCs and formation of the calibration pattern. (a) detection of the main subpopulations of PBMCs one-by-one, *i.e.* lymphocytes and monocytes. (b-g) Formation of the pattern: plot of individually measured fluorescently labelled PBMC cells. (h) Exemplary calibration pattern. (j) Impedance cytometry of unlabeled PBMCs: matching of the labelled PBMC mixture with the calibration. The lymphocyte cluster is divided based on its subpopulations, namely NK-, T-, and B-cells. (k) Matching the

unlabelled PBMC mixture with the calibration pattern. The lymphocyte cluster is divided based on its subpopulations, namely NK-, T-, and B-cells. Panels (l)-(n): calibration of the signal for impedimetric detection of myeloblasts. (l) Detection of the labeled isolated blasts one-by-one in time domain; (m) formation of the data cloud and its localization within the pattern of peripheral blood of the AML positive donor; (n) matching the myeloblasts cluster (red) with the unlabeled PBMC of the AML positive donor (black open circles) and PBMC of healthy donor (gray circles).





Healthy PBMC					
Cell type	Monocytes	Lymphocytes	T cells	B cells	NK cells
Flow cytometer	11,80%	88,20%	57,50%	30,70%	10,10%
Nano cytometer	9,84%	90,16%	62,31%	31,34%	7,34%

Acute myeloid leukemia (nanocytometer)						
Cell type	Monocytes	Lymphocytes	T cells	B cells	NK cells	Myeloblasts
Healthy donor	17,51%	82,48%	50,53%	34,61%	14,86%	0%
AML #1	25,02%	40,82%	36,52%	26,50%	36,98%	34,16%
AML #2	24,91%	15,02%	34,09%	17,05%	48,86%	60,07%
AML #3	12,60%	32,34%	32,02%	7,02%	60,96%	54,96%

**Figure 5:** (a) Detection and characterization of PBMC of healthy human donors (see Inset) and AML patients#1-3. A new cluster is found in the AML samples caused by the presence of the myeloblast subfamily. (b) Impedance cytometry scatter plot of PBMCs of healthy donor (n=1000 cells) and (c) AML patient (n=1400 cells) calculated via the machine learning algorithm. (d) Overview of the individual cell counts, comparison to healthy patient. AML patients PBMCs shows a myeloblast percentage in the range 30-60%.



**TOC image:**

Ultra-compact nanocytometer for real-time impedimetric detection and classification of subpopulations of living cells in peripheral blood.

