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MASTER THESIS

POLYMER PEN LITHOGRAPHY
FOR BIOACTIVE SURFACE FUNCTIONALIZATION

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SUMMARY

Polymer Pen Lithography (PPL) which is a hybrid of Dip-Pen Nanolithography (DPN) and soft lithography technique namely Microcontact Printing (µCP) has been tested for its pros and cons in patterning functional phospholipids under varying instances using high resolution characterization techniques. Although, having the advantage of simultaneously printing at several points of the substrate covering area on the order of millimeter by employing a Polydimethylsiloxane (PDMS) stamp however PPL is yet to make a mark to prove its ability to print pure lipids without admixing with carrier lipids which is not possible by cantilever-based scanning probe lithography techniques such as its parent technique DPN. The detailed analysis reveals a theoretically expected result of phospholipid layers stacked over the base Self Assembled Monolayer (SAM) of the phospholipid which is a direct result of adsorption of the Phosphate head over the hydrophilic glass or Silicon Dioxide substrate which assembles the monolayer constituting the hydrophilic head and hydrophobic tail in a regularly arranged manner. A statistical method is proposed which not only quantifies the stacked phospholipid layers in water, mimicking its natural state being the main constituent of cell membrane in blood plasma, but also has the potential to quantify the Scanning Probe Microscopy (SPM) probe which is used to characterize the sample such that the probe can be said to have certain factor which represent the destruction caused to the particular phospholipid feature while traversing over the sample in water. Moreover, the net overall charge of the phospholipid is observed to affect the topographic image due to the nature of characterization method used therefore a printing methodology to enhance the contrast is proposed to achieve optimum and reliable results. Finally, the difference between the feature is observed in different operating regimes based on the deformation and hence the contact area of the PDMS stamp which is observed to dictate the characteristics of the deposited feature.
PERSONAL CONTRIBUTION

At the time of publication of the thesis report the author is enrolled at Politecnico Di Torino (POLITO), Corso Duca degli Abruzzi, 24, 10129 Torino, Italy in an MS Nanotechnologies for ICTs program offered by Department of Electronics and Telecommunications (DET) Corso Castelfidardo, 39, 10129 Torino, Italy. The thesis is carried out under the supervision of Prof. Fabrizio Giorgis who is the Vice Head of Department of Applied Science and Technology (DISAT), Corso Duca degli Abruzzi, 24, 10129 Torino, Italy.

The experimental work of the thesis under POLITO’s ‘Thesis in a Company’ category is carried out at Institute of Nanotechnology (INT), Karlsruhe Institute of Technology (KIT) Hermann-von-Helmholtz-Platz 1, D-76344 Eggenstein-Leopoldshafen, Germany under the external supervision of the group leader of DPN and related techniques’ PD Dr. Dr. Michael Hirtz. All the experimental work and remarks are conducted by the author led by the lab supervisor at guest institute Dr. Uwe Bog. The only exception is for the SICM characterization and evaluation which is carried out at Institute of Physics, University of Münster, D-48149 Münster, Germany and Center for Nanotechnology (CeNTech), D-48149 Münster, Germany by Dr. Joo Hyoung Kim supervised by Prof. Kristina Riehemann and Prof. Harald Fuchs with the SICM software and hardware provided by Dr. Goo-Eun Jung of Park Systems Corp., KANC 4F, Gwanggyo-ro 109, Suwon 16229, Korea.
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1. Introduction

PPL (Polymer Pen Lithography) has been able to find applications in numerous fields to date but its primary use still lies in the deposition of bio-inks which have vast uses in biochemistry and cell biology. PPL is cantilever-free scanning probe lithography technique which has been shown [6] to obtain adjustable sized, high throughput, multiplexed patterns allowing deposition of different inks with a reasonably high compactness (area efficiency) over the substrate. It has been combined with thiol–acrylate photopolymerization chemistry for the creation of brush-polymer microarrays over large areas and nanometer-scale control over print position and size, therefore this technique of combination of localized photochemistry and functional-group tolerant chemistry finds application in diverse problems in biology, materials chemistry, and organic electronics [12].

PPL is basically an additive serial writing lithography method. Conventional lithography methods find application in many fields such as in the development of ICs, MEMS, video displays and projectors, data storage disks, biochips, miniaturized sensors, microfluidic devices, fine tunable bandgap photonic devices and diffraction devices. PPL stands out due to its low-cost, cantilever free design and high-throughput scanning probe lithography scheme that utilizes a soft elastomeric tip array as opposed to hard Si-based cantilever to deliver inks to a substrate surface in a direct write manner, the ink delivery is time and force dependent which determines feature size at either nanometer, micrometer, or macroscopic scales using the same tip array. PPL has been demonstrated to be capable of printing the complete range of DPN (Dip Pen Nanolithography) printable inks except for thermal DPN inks [13]. Additionally, a new group of dry inks which cannot be printed using DPN such as high gel–liquid phase transition temperature inks and non-soluble inks can also be printed using PPL due to the µCP (Micro Contact Printing) like “stamping” mode of transport hence inks neither need to be liquid nor need to be soluble in any carrier [13].

The bio-ink that has been employed for the PDMS stamp in deposition over glass substrate for this case is phospholipids. A generic phospholipid constitutes a hydrophilic phosphate group head and a hydrophobic fatty acid tails with a glycerol molecule in between. Phospholipids are the main constituents of cell membranes and tend to deposit with micro contact printing or DPN etc. by formation of SAM (Self Assembled Monolayers) with the hydrophilic head adsorbing over
the hydrophilic substrate. SAM are also known to rearrange themselves in an orderly manner. Lipid bilayers tend to form because of the amphiphilic characteristic of the lipids. Depending on the diffusion which in turns depend on the dwell time, chamber humidity and temperature of the deposition technique number of stack of lipid bilayers may stack up over the hanging hydrophobic tail of the SAM. Being the major part of the plasma membrane of living cells researchers are usually interested in observing the behavior of the lipid immersed in physiological buffer so that it can be used to model how lipid would behave in blood plasma which constitutes 90% water.

Scanning ion conductance microscopy (SICM) and atomic force microscopy (AFM) are the primary techniques used hereafter to characterize the phospholipids and determine their behavior in liquid. As the AFM is inherently destructive in nature a statistical method is to be proposed which not only quantifies the stacked phospholipid layers in water, mimicking its natural state being the main constituent of cell membrane in blood plasma, but also if possible measure the probe characteristics which is used to characterize the sample such that the probe can be said to have certain factor which represent the destruction caused to the particular phospholipid feature while traversing over the sample in liquid. Moreover, correlation of the overall charge of the phospholipid needs to be seen as to how it affects the topographic image due to the nature of characterization method used therefore a printing methodology to enhance the contrast is required to be proposed to achieve optimum and reliable results. Finally, the effect on the feature needs to be observed with high resolution for different deposition regimes based on the deformation and hence the contact area of the PDMS stamp which is known to dictate the characteristics of the deposited feature.
2. Methodology

2.1 Derivation of the printing methodology

PPL is the primary techniques used in this study to deposit lipid features over the substrate. In the last few years printing technologies, e.g. flexography, soft lithography, screen, gravure and inkjet printing gained the attention of manufacturing industries due to their low-cost, high volume and high-throughput production of electronic components or devices which are lightweight and small, thin and flexible, inexpensive and disposable. Printing methods are in general additive processes unlike the traditional lithographic techniques in the semiconductor and MEMS industry such as photolithography which are inherently subtractive. Figure 2-1 displays how the additive process is different in the sense that the desired pattern is directly implemented over the substrate whereas for the traditional case the material is deposited over the substrate and then the undesired part is selectively removed from the top of the substrate and sometimes also the bulk of the material as in the case of for example bulk micromachining in MEMS.

![Figure 2-1 Comparison between traditional subtractive patterning vs additive processes](image)

In the recent past there have been numerous efforts to develop a molecular printing technique having high throughput, high spatial positioning precision/resolution, high density integration and higher control over the feature size and shape. PPL brings all these advantages together in a...
single technique. PPL can be thought of as being derived from the combination of micro contact printing and dip pen nanolithography. The remaining Section shows how it combines the advantages of the two techniques to come up with all the advantages.

2.1.1 Micro Contact Printing

Micro Contact Printing is a soft lithography method of printing over a substrate. Soft lithography is principally a non-photolithographic strategy based on self-assembly and replica molding. It is a convenient, effective and low-cost method for manufacturing micro- and nanostructures by an elastomeric stamp with patterned relief structures on its surface which is used to generate micro and nano patterns and structures [10]. Industrially for micro- and nanostructures, photolithography is dominating but soft lithography is pretty much the close second. In soft lithography generally, polydimethylsiloxane (PDMS) is used as the elastomer due to low glass transition temperature. As in cast molding, at room temperature prepolymer being liquid can be shaped up using a Silicon Master of complementary shape, after the crosslinking/curing agent is mixed at raised temperature the PDMS hardens and the Si Master can be easily peeled off the structure thereafter. The complementary Si Master is fabricated using microfabrication techniques such as electron beam lithography for precision in getting the desired arbitrary shape. Figure 2-2 displays the stepwise preparation of the PDMS stamp.

![Figure 2-2 Graphic display of the steps involved in preparation of a PDMS stamp from Si Master](image)
PDMS stamp properties of being chemically inactive, having a less value i.e. 21.6 dyn/cm of interfacial free energy, being non-hydroscopic, having high thermal stability in air, being transparent and being durable ensure its use in the current soft lithography world. However, there are some drawbacks which hinders soft lithography from becoming the dominant means of industrial fabrications, these include post-cure shrinking displayed at the bottom of figure 2-2, swelling by organic solvents, lesser accuracy due to elasticity and expansion and lastly the range of aspect ratio needs to be taken care of due to softness as it may lead to defects as a result of deformation [10]. Microcontact printing utilizes the PDMS stamp to pattern self-assembled monolayers (SAMs) [11] by contacting the surface of the substrate. Self-assembly is the spontaneous rearrangement of the subunits of the deposited ink, which is usually a ‘bioink’ such as protein, DNA and cell membrane, onto the substrate such that the whole structure is in the lowest possible form energetically. Other than monolayers self-assembly, microcontact printing is also used in self-assembly in two or three dimensions as well. The most popular microcontact printing technique which is relevant to this study is the one in which a planar stamp is used to print on a planar surface, however either of the stamp or the substrate need not necessarily be planar in general.

Soft lithography has many advantages such as being convenient, inexpensive, accessible to chemists, biologists, and material scientists, having basis in self-assembly which tends to minimize certain types of defects, most soft lithographic processes are additive and minimize waste of materials, readily adapted to rapid prototyping for feature sizes >20 mm, isotropic mechanical deformation of PDMS mold or stamp provides routes to complex patterns, having no diffraction limit features as small as 30 nm have been fabricated, nonplanar surfaces (lenses, optical fibers, and capillaries) can be used as substrates, generation and replication of three-dimensional topologies or structures are possible, optical transparency of the mask allows through-mask registration and processing, good control over surface chemistry, very useful for interfacial engineering, a broad range of materials can be used: functional polymers, sol–gel materials, colloidal materials, suspensions, solutions of salts, and precursors to carbon materials, glasses, and ceramics being applicable to manufacturing: production of indistinguishable copies at low cost, applicable in patterning large areas. The disadvantages include patterns in the stamp or mold may distorting due to the deformation (pairing, sagging, swelling, and shrinking) of the elastomer used, difficulty in achieving accurate registration with elastomers (<1 mm), doubtable compatibility with current integrated circuit processes and
materials, defect levels higher than for photolithography, µCP works well with only a limited range of surfaces [10].

2.1.2 Dip-Pen Nanolithography

DPN was invented in 1999 by the Mirkin Group, and it can be used to deposit molecules and materials on surfaces with sub-50 nm resolution [3]. It uses an AFM tip to selectively place different types of molecules at specific sites within a nano-scale pattern without using an intermediate such as resist or stamp. It has the ability to pattern with a wide variety of ‘inks’ and works best with ‘bioinks’ that tend to form SAMs as shown in figure 2-3 where the molecules are seen to adsorb to the surface of Au substrate by the ink forming a meniscus controlled by environment humidity and temperature which in turn effect the resolution of the technique. Therefore, it is a cantilever-based serial writing technique which has displayed been displayed to be compatible with many inks, from small organic molecules to organic and biological polymers, and from colloidal particles to metal ions and sols [1]. DPN is a particularly attractive tool for patterning biological and soft organic structures onto surfaces in ambient or inert environments without exposing them to ionizing UV or electron-beam radiation and without risking cross-contamination. Amongst many research activities utilizing the method all around the world some highlights include the use of DPN for in situ studies of surface reactivity and exchange chemistry, patterning biomolecular micro- and nanoarrays, building tailored chemical surfaces for studying and controlling biorecognition processes from the molecular to cellular level, generating chemical templates for the controlled orthogonal assembly of materials on surfaces and the use of DPN as a rapid prototyping tool for generating hard nanostructures using chemical etching on a length-scale comparable, or even superior, to that obtainable with e-beam lithography [1].

Figure 2-3 AFM tip traversing over the substrate leaving the traces of SAM behind [3]
materials on surfaces and the use of DPN as a rapid prototyping tool for generating hard nanostructures using chemical etching on a length-scale comparable, or even superior, to that obtainable with e-beam lithography [1]. In short, the combination of resolution, registration, and direct-write capability offered by DPN distinguishes it from any alternative conventional lithographic strategy and makes DPN a promising tool for patterning soft organic and biological nanostructures

2.1.3 Polymer Pen Lithography

PPL was reported [6] as a low-cost, high-throughput scanning probe lithography method that uses a soft elastomeric tip array, rather than tips mounted on individual cantilevers, to deliver inks to a surface in a “directwrite” manner. Polymer pen lithography merges the feature size control of DPN with the large-area capability of microcontact printing. Because ink delivery is time and force dependent, features on the nanometer, micrometer, and macroscopic length scales can be formed with the same tip array due to the piezo electric control typical of SPM methods. PPL PDMS elastomeric stamps may contain many thousands of pyramid-shaped tips prepared with a silicon master as shown in Figure 2-4, obtained by conventional photolithography followed by wet chemical etching, where the pyramids of the stamp are connected from the usually square base with a backing thin layer of PDMS of micrometric thickness (usually 50-100 microns) followed by the glass support, both of which play a vital role in improving the homogeneity of the PPL arrays on for example 3-inch (76.2mm) wafer as shown in Figure 2-5 [2].

Figure 2-4 A schematic illustration of the polymer pen lithography setup [2]
In PPL, similar to cantilever-based SPM lithography techniques the array tips are sharp and delivers ink upon making contact with the substrate at only the point of contact. The point of contact can be visually observed under a camera while printing as the transparent polymer pen array is seen to be elastically deformed upon contact resulting in an increased light reflection from the tip of the pyramids. This observation also comes in handy to position the array or in other words the stamp in parallel to the substrate in order to have homogenous distribution of ink as required in most of the applications where intentional feature size gradient as used in some studies [14] is not required. The number of pyramid pens can be in millions with their quantity depending on the tip to tip spacing in between the pens, the dimension of the base of the pens and eventually since the base of the pens consume finite area it also depends on the maximum capability of the equipment to hold a stamp of given area.

PPL has been used with varying strategies for the printing of functional phospholipid patterns that provide tunable feature size and feature density gradients over surface areas of several square millimeters by controlling the printing parameters, having shown two operate in two transfer modes of either of its pre cursor techniques’ [14]. Each of these modes leads to different feature morphologies as by increasing the force applied to the elastomeric pens the tip–surface contact area increases enhancing the ink delivery rate, a switch between a DPN and μCP transfer mode can thus be triggered. This results in a range of deposition properties of the ink feature from quasi-DPN to quasi-μCP which can be fine-tuned by the piezoelectric controller. Moreover, PPL has been demonstrated to possess the capability to do multiplexing [4], which is the integration of
more than one ink in an interdigitated microscale pattern and is still a challenge for microcontact printing. On the other hand, there is a strong demand for interdigitated patterns of more than one protein on subcellular to cellular length scales in the lower micrometer range in biological experiments.

2.2 Nature of the ink
The bio-ink that has been employed for the PDMS stamp in deposition over glass substrate for this case is phospholipids. A generic phospholipid constitutes a hydrophilic phosphate group head and a hydrophobic fatty acid tails with a glycerol molecule in between. Phospholipids are the main constituents of cell membranes and tend to deposit with micro contact printing or DPN etc. by formation of SAM (Self Assembled Monolayers) with the hydrophilic head adsorbing over the hydrophilic substrate. SAM are also known to rearrange themselves in an orderly manner. Lipid bilayers tend to form because of the amphiphilic characteristic of the lipids. Depending on the diffusion which in turns depend on the dwell time, chamber humidity and temperature of the deposition technique number of stack of lipid bilayers may stack up over the hanging hydrophobic tail of the SAM as shown in figure 2-6 [6]. Being the major part of the plasma membrane of living cells researches are usually focused on observing the behavior of the lipid immersed in physiological buffer so that it can be used to model how lipid would behave in blood plasma which constitutes 90% water.

With the aim of better controlling the resulting resolution and feature size especially the height or number of layers on top of SAM as given in the figure above, analysis and modeling of the dependence of lipid features (area, height and volume) directly depend on dwell time and relative humidity [7]. The dwell time as it dictates the time allowed for diffusion of the ink over to the
substrate and humidity as in the quasi-DPN mode it dictates the size of meniscus formed leading expressing the amount and resolution of lipid transfer. Feature shape is usually controlled by the substrate surface energy. Generally, a short dwell time growth is controlled by meniscus diffusion while at long dwell times surface diffusion is the dominant factor. The critical point for the switch of regime depends on the humidity for a given dwell time.

In contrast, if the surface of the substrate is hydrophobic such as graphene the lipid formation instead of figure 2-6 would rather be similar to figure 2-7 [8] as the hydrophobic tail will be attached to the substrate instead of the hydrophilic head as in the previous case.

*Figure 2-7* A typical lipid deposition over hydrophobic substrate such as graphene [8]
3. Results and Discussion

Knowledge about surface charge of certain biological objects gives key information for understanding their structures, functions, and their behavior in a wide range of metabolisms. Recently, surface charge mapping methods within physiological condition based on various Scanning Ion Conductance Microscopy (SICM) measurement schemes have been developed and suggested. Although they have shown great capability for computing surface charge density and understanding biological events in electrostatic perspective from various samples ranging from lipid patches to live cells, there were little attempts to present well-defined, periodic, reproducible “standard sample” for these measurement methods, which is essential to achieve general applicability and calibration capability for whole scanning probe microscopy (SPM) measurement schemes. Here the study shows surface charge mapping associated with amplitude-based bias-modulated (BM-) SICM for various types of standard samples printed on glass substrates, which have different surface charge densities. Thereafter AFM images of the lipids have been used as the control, and it as found that BM-SICM mode has shown good stability, image quality, and reproducibility. It is highly expected that armed with our “standard samples”, SICM-based surface charge mapping method would have a greater momentum for its standardization and enhancement of its reliability.

Electrostatic force is the most accountable and fundamental physical interaction (among 4 fundamental physical interactions) between biological entities especially at their molecular level [19]. Thus, knowledge on surface charge distribution of biological substances is essential to understand their physicochemical behaviors inside a physiological buffer. However, despite of its importance with this information, measuring surface charge density of some sample within physiological conditions (ionic strength in the order of 100 mM where most of metabolic activities happen) has been quite challenging due to the presence of very tight Debye layer (less than 1 nm) formed in the vicinity of sample surface. Though Zeta potential measurement is the most widely used method for probing surface charge information for small (micro to nano) materials [20,21], this has its own limitations, firstly it usually assumes that the material under study should be colloidal particles, secondly the measured values largely fluctuate at physiological conditions (i.e. high ionic strength), and finally applying this method to ‘real’ biological systems such as live cell membranes is challenging, where they are usually supported
by certain substrates or scaffolds in more complex dimensions than 0-dimensional particles. From these aspects, surface charge mapping based on the scanning probe microscope (SPM) offers much attractions for measuring surface charge density, first because this value has basically 2-dimensional feature (C/m²) as SPM scans this 2D area, and second because SPM is operable within physiological conditions. There are noted attempts using atomic force microscope (AFM), the most widely used type of SPM, by surface functionalization of tip with some charged molecules [22]. But for this case too, experiments were done in a moderate ionic strength (~1 mM, hundred-fold dilute compared to physiological condition), and quantitative analysis was not available. Recently, two groups have shown capability for quantitative surface charge mapping method within physiological conditions based on scanning ion conductance microscopy (SICM) [23-28]. Even though these group use different measurement mode of SICM (Unwin et al. used a spectroscopy-based hopping mode, while Dong et al. exploited the conventional DC-SICM raster scanning mode), the basic idea is the same, that the local conductivity measured by pipette tip does not just indicate typical tip to sample distance, but also reflects the local ion concentration on the sample spot, where the tip is approached. At there, the counter-ions are relatively rich and co-ions are less than bulk ion concentration where the tip is distant (distant so that no counter-ions are absorbed or repelled by the tip) from Debye layer surrounding the sample. Studies have shown great applicability of this method in a wide range of samples, from lipid patches to live cells [23-28]. However, despite of this great deal of works on surface charge mapping studies within physiological conditions, very little attention has been paid to the development of “standard sample” for this method, which is essential to achieve general applicability and calibration capability for whole scanning probe microscopy (SPM) measurement methods. The samples which are called “standard samples” for the surface charge mapping method should show a periodic structure, whose sizes are well matched to typical scanning dimensions of SICM, and at the same time a well-defined electrostatic surface charge distribution, possibly coinciding with their topographic features, for ease of recognition. To show these features, we have studied polymer-pen-lithographied (PPL) lipid features, whose substrates are glasses (with negative charge from expressed hydroxyl group or silanol group on the surface here also within aqueous solution with neutral pH) for both cases. From these two groups of samples, we could see different pipette-sample interactions according to their surface charge distribution, showing that these structures could serve as good standard samples for the surface charge mapping method application. An AFM control is used to appreciate where the SCIM signal shows variation with the AFM lipid topography due to surface charge density of the lipid when doing the surface charge mapping.
Finally, a statistical method is proposed which not only quantifies the stacked phospholipid layers in physiological buffer, mimicking its natural state being the main constituent of cell membrane in blood plasma with ~90% water, but also has the potential to quantify the Scanning Probe Microscopy (SPM) probe which is used to characterize the sample such that the probe can be said to have certain factor which represent the destruction caused to the particular phospholipid feature while traversing over the sample in water.

### 3.1 Slightly Charged Phospholipid Inks

For the first batch of PPL a safe approach was taken to ensure that deposition of the lipid definitely takes place and at the same time is easily detectable under fluorescence microscope. There were two variants of inks used namely ink α1 and ink β1 both of which were slightly negatively charged with detailed composition provided in Section 3.1.1 and Section 3.1.2.

Even though ink α1 samples showed a nice pattern under fluorescence microscope in the lab at KIT primarily because of bright fluorescence of Rhodamine, the SICM characterization done at CeNTech, Munster was not very successful for sample which used ∞A probably because DOPC was not printed along as much or maybe washed away in the KCl solution which immersed the sample for the SICM traversal.

For the second type of samples though the features were observed because of higher percentage of primary lipid in ink β1 which did not wash away as much with the immersion in KCl solution. SICM results are shown in figure 3-1 which were obtained at 1 kHz, an input bias of 50mV while the sample was immersed in 0.15M KCl solution with a pH value of 7. The sample was traversed over a scan size of 15µm×15µm. The micro-pipette current was adjusted to a value of 800 pA and 1% setpoint value.

![Figure 3-1 SICM images at DC offset of -330 mV & 330 mV of ink β1 with cuts drawn at same place for comparison.](image-url)
In the DC mode polarity dependent profile, the SICM image comparison at two different DC offsets of -330 mV and +330 mV is drawn but a sharp contrast was not observed for 0 V offset. The red colored graph in figure 3-1 represents the height profile of the lipid at positive DC offset along the x-direction of the cut. On the other hand, the green colored graph represents the height profile of lipid ink at negative DC offset along the x-direction of the cut at the same place where the cut for the positive DC offset was made. It can be observed that a height of ~2nm is observed at the positive DC offset in contrast to a height of ~1nm at negative DC offset. Moreover, the glass in both cases seemed to have a negative charge translating to a negative height as per the working principle of SICM, this implied that the glass has a negative charge according to the calibration, parameter and environment (KCl immersion) in which the measurement of SICM was done. Additionally, the group at KIT which was not directly doing the measurement including myself had realized that the scan size of SICM is too small to be able to scan several features at once as maximum image size is 15µm×15µm.

3.1.1 Ink α1
The first type of samples was prepared with an ink containing 90 mol% of a very fluid lipid namely DOPC (1,2-dioleoyl-sn-glycero-3-phosphocholine), with structure as shown in Figure 3-2, which is well known to easily deposit over glass substrate was used. It was admixed with 10 mol% of Rhodamine ((lissamine rhodamine B sulfonyl) (ammonium salt)), with structure as shown in Figure 3-3, which is known to have a very bright red fluorescence even in low concentrations and hence easy to detect under a fluorescence microscope even if the deposited features are so small that they are not visible under a bright field optical microscope.

![Figure 3-2 Structural Formula of the employed DOPC [29]](image)

As may be observed in Figure 3-2 that DOPC has no overall net charge whereas from the structural formula of Rhodamine in Figure 3-3 it is clear that it has an overall net charge of -1. A predominantly neutral ink with 90% of neutral lipid and 10% of negatively charged lipid tends to be slightly negative and therefore it is expected that it would amount to a reasonable contrast when characterized under SICM to observe the natural topography of lipid in liquid.
Figure 3-3 Structural Formula of the employed Rhodamine [29]

For PPL machine a PDMS stamp of 100×50, where the former count signifies the center to center distance in between two consecutive pyramids in μm and the latter count signifies the base length and width of the pyramids’ square base in μm, prepared from its negative Si master fabricated with Electron Beam Lithography (EBL) was mounted. The stamp was then used by the NLP 2000 default software with the help of the precise piezoelectric control to generate ink patterns with the center to center closest proximity distance of 20μm. To be on a safe side and further ensure that the printing really took place at the micro-scale the samples were observed under upright fluorescence microscope as shown in Figure 3-4.

Figure 3-4 Fluorescence microscopy image for ink α1
The result was positive, a homogenous pattern of circular lipids was observed with a feature diameter distribution given in table 3-1 as seen under a Texas Red filter with an exposure time of 400ms over a 20x zoom lens. The diameter had a mean of 2.839\(\mu m\) and a standard deviation of 0.835\(\mu m\).

![Histogram of feature size distribution](image)

**Table 3-1** Ink \(\alpha_1\) feature size distribution

The approach dot or the first point of contact can be observed to have a comparatively large feature size due to more dwell time and contacting more than once while defining the plane of print for the software to make the pattern.

### 3.1.2 Ink \(\beta_1\)

The second type of samples was prepared with an ink using a higher 95 mol% of DOPC having structure already shown in Figure 3-2 having known to be easy to deposit over glass substrate. It was admixed with 5 mol% of Dansyl (N-(5-dimethylamino-1-naphthalenesulfonyl) (ammonium salt)), with structure as shown in Figure 3-5, which is known to have a decent blue fluorescence hence easy to detect under a fluorescence microscope to aid in seeing admixed non-fluorescent lipids that may not be visible under a bright field optical microscope such as DOPC in our case.

![Structural formula of Dansyl](image)

**Figure 3-5** Structural Formula of the employed Dansyl [29]
Again, DOPC has no overall net charge apparent while from Figure 3-5 it is clear that Dansyl has an overall net charge of -1. A predominantly neutral ink with 95% of neutral lipid and 5% of negatively charged lipid tends to be slightly negative and therefore it is expected that it would amount to a reasonable contrast when characterized under SICM to observe the natural topography of lipid in liquid.

For PPL machine a similar PDMS stamp of 100×50, where the former count signifies the center to center distance in between two consecutive pyramids in μm and the latter count signifies the base length and width of the pyramids’ square base in μm, prepared from its negative Si master fabricated with Electron Beam Lithography (EBL) was mounted. Similar printing scheme of patterns with the center to center closest proximity distance of 20μm was employed. As a litmus test and ensure that the printing really took place at the micro-scale the samples were observed under upright fluorescence microscope with result shown in Figure 3-6.

![Figure 3-6](image)

Figure 3-6 Fluorescence microscopy image for ink β1 of Batch no. 1

A homogenous pattern of lipids can be observed with a feature diameter distribution given in Table 3-2 with a mean value of 10.84μm and standard deviation of 4.754μm under a DAPI filter with an exposure time of 400ms over a 20x zoom lens.
The approach dot or the first point of contact once more can be observed to have a comparatively large feature size due to more dwell time and contacting more than once while defining the plane of print for the software to make the pattern. Additionally, when compared to Figure 3-4 the average size of the features can be seen to be larger due to higher mol percentage of DOPC which, to reiterate, has such viscosity which allows more deposition of the lipids to the substrate.

### 3.1.3 Outlook

Having noticed that there was not an appreciable charge contrast of the deposited features and small scan size as apparent from SICM images, for the second batch prepared by PPL number of strategies were devised to observed better SICM characterization, firstly it would best to have a higher concentration of charged lipids than 5% or 10% as done previously in the respective inks so that a high charge contrast is achieved which would result in a better characterization. Secondly, as DOPC was seen to nicely deposit it could still be used as the primary constituent in terms of concentration in the inks to act as the carrier lipid for the charged lipids which may not have as nice deposition properties. Thirdly, the fluorescence lipid concentration percentage needs to be decreased to avoid false positives as seen in the previous case for ink α1 where a nice fluorescence microscopy image with bright red features was observed but nothing appreciable was observed in SICM images. Fourthly, the features need to be closely packed so that the scan size window of SICM can incorporate more than one feature. Lastly, it would be even better to prepare a sample which is a multiplex of positively charged lipid features and negatively charged lipid features so that the charge contrast is even greater.

**Table 3-2** Ink β1 feature size distribution

![Graph showing feature size distribution](image)
and importantly the negative charge tendency of the glass substrate can be either confirmed or negated.

3.2 Moderately Charged Phospholipid Inks

In the light of the decisions made thus far from the last Section, yet again, there were two variants of inks used namely ink α2 which was moderately positively charged and ink β2 which had a moderate-high negative charge with detailed composition provided in Section 3.2.1 and Section 3.2.2. The configuration details of the samples prepared by the multiplexing of both ink α2 and ink β2 are provided in Section 3.2.3

![SICM images of α2 samples at three different voltage biases of +340 mV, -340 mV and zero volts](image)

Figure 3-7 SICM images of α2 samples at three different voltage biases of +340 mV, -340 mV and zero volts

For the first type of samples which had moderately positively charged ink α2 SICM results are shown in Figure 3-7 at three different voltage biases. While the image at the positive bias does not show any convincing feature topography except for a printing impression, there is clearly some whiteness in the image which would usually represent the positive height deviation of possibly a lipid. However, for this case white patches have a size ~500nm contrary to the expected range in micrometers. Moreover, they don’t seem to have the periodic pattern which is the inherent characteristics of the features as per the employed printing scheme with detailed description given in Section 3.2.1. Therefore, the white patches are probably artifacts which must not be confused for lipid features. All the images were obtained at 1 kHz while the sample was immersed in 0.15M KCl solution with a pH value of 7. The sample was traversed over a scan size of 15µm×15µm except for the multiplexed image in which a scale of 10µm×10µm was used. The micro-pipette current was adjusted to a value of 800 pA and 1% setpoint value.
The second type of sample with moderately-high negatively charged ink β2, unfortunately it did not give reasonable results except for noise for all biases that were tried with it. After observing Figure 3-8 it would be safe to assume that the negatively charged lipids are camouflaged in the negatively charged glass substrate which was doubted from results of the batch with slightly charged inks in Section 3.1.

![Figure 3-8 SICM images of β2 samples at three different voltage biases of +340 mV, -340 mV and zero volts](image)

The images with which most expectations were associated because of multiplexing of both moderately charged positive ink α2 and moderately-high charged negative ink β2 definitely shows white patches as may be observed in Figure 3-9 given below.

![Figure 3-9 SICM images of α2 and β2 samples at three different voltage biases of 0 mV, -305 mV and -305 mV](image)
With the decreased 10µm×10µm scale it can be appreciated that the white patches have a size ~500nm contrary to the expected range in micrometers. Moreover, they don’t seem to have the periodic pattern which is the inherent characteristics of the features as per the employed printing scheme with detailed description given in Section 3.2.3. Therefore, the white patches are probably artifacts which must not be confused for lipid features. The negative bias was observed to enhance the role of artifacts in the image.

3.2.1 **Ink a2**

This type of samples was prepared with an ink containing 59.95 mol% of DOPC with structure as shown in Figure 3-2, which was previously used and found to deposit well over the glass substrate. It was admixed with 39.95 mol% DOTAP (1,2-dioleoyl-3-trimethylammonium-propane (chloride salt)), with structure as shown in Figure 3-10. For fluorescence purposes 0.1 mol% of Rhodamine was used which is order of magnitude lower than the previous cases to avoid getting false positive in SICM characterization and since it has a very bright red fluorescence it glows even in low concentrations and hence easily detectable under a fluorescence microscope.

![Figure 3-10 Structural Formula of the employed DOTAP](image)

As maybe observed in Figure 3-2 that DOPC has no overall net charge whereas from the structural formula of DOTAP in Figure 3-10 it is obvious that it has an overall net charge of +1. An almost 60% neutral ink with about 40% of positive lipid tends to be moderately positively charged, consequently it is expected that it would amount to a reasonable contrast with a previously known to be slightly negative glass substrate when characterized under SICM to observe the natural topography of lipid in liquid.

For PPL machine a PDMS stamp of 100×50, where the former count signifies the center to center distance in between two consecutive pyramids in µm and the latter count signifies the base length and width of the pyramids’ square base in µm, prepared from its negative Si master
fabricated with Electron Beam Lithography (EBL) was mounted as done previously. The stamp was then used by the NLP 2000 default software with the help of the precise piezoelectric control to generate ink patterns with the center to center closest proximity distance of 20µm which look like as shown in Figure 3-11 when observed under upright fluorescence microscope.

A somewhat homogenous pattern of lipids with feature diameter distribution given in table 3-3 was observed with a lens of 10x giving a mean value of 6.897µm and a standard deviation of 1.904µm under a Texas Red filter with an exposure time of 10s.
The approach dot or the first point of contact can be observed to have a comparatively large feature size due to more dwell time and contacting more than once while defining the plane of print for the software to make the pattern.

### 3.2.2 Ink β2

The second type of samples was prepared with an ink constituting the same 59.95 mol% of DOPC as for α2 to act as the carrier lipid which ensures printing of the ink. It was admixed with 39.05 mol% of DGS-NTA (1,2-dioleoyl-sn-glycero-3-[(N-(5-amino-1-carboxypentyl)iminodiacetic acid)succinyl]), with structure as shown in Figure 3-12. The remaining 1 mol% was for PE CF (1,2-dioleoyl-sn-glycero-3-phosphoethanolamine-N-(carboxyfluorescein) (ammonium salt)) commonly known as fluorescein with structure displayed in Figure 3-13 which is expected to have a decent green fluorescence aiding detection under a florescence microscope and since it is not as bright as Rhodamine so one order of magnitude higher molar percentage of fluorescence lipid is used compared to α2.
DOPC as observed already has no overall net charge but from Figure 3-12 DGS-NTA has an overall net charge of -3. Although only 1 mol% but fluorescein can be considered to have -2 net charge as well. A 60% neutral ink with about 39% of highly negatively charged lipid and 1 mol% of slightly negatively charged lipid can be thought to be moderately-high negatively charged and therefore it is expected that it would amount to a reasonable contrast when characterized under SICM to observe the natural topography of lipid in liquid even if the substrate is slightly negatively charged glass.

For PPL machine a as in all previous case a PDMS stamp of 100×50, where the former count signifies the center to center distance in between two consecutive pyramids in µm and the latter count signifies the base length and width of the pyramids’ square base in µm, prepared from its negative Si master fabricated with Electron Beam Lithography (EBL) was mounted. Similar printing scheme of patterns with the center to center closest proximity distance of 20µm was employed. As a litmus test and ensure that the printing really took place at the micro-scale the samples were observed under upright fluorescence microscope with result shown in Figure 3-14.
A homogenous pattern of lipids can be observed at 10x hardware zoom having a mean diameter of 4.642\(\mu\)m and a standard deviation of 1.754\(\mu\)m under a FITC filter with an exposure time of 20s. The feature diameter distribution for the sample type is given in table 3-4.
3.2.3 Sample with multiplexing of ink $\alpha_2$ and ink $\beta_2$

This type of sample was prepared with a multiplexing of overall positively charged ink $\alpha_2$ (details in Section 3.2.1) as well as overall negatively charged ink $\beta_2$ (details in Section 3.2.2) within a close proximity to have a high contrast and both types of lipids within the limited scan size of SICM.

To print in this multiplexed manner with PPL machine a PDMS stamp with four instead of one Section of 100×50, where the former count signifies the center to center distance in between two consecutive pyramids in $\mu$m and the latter count signifies the base length and width of the pyramids’ square base in $\mu$m, prepared from its negative Si master fabricated with Electron Beam Lithography (EBL) was used. The stamp was then used by the NLP 2000 default software with the help of the precise piezoelectric control to generate ink patterns with the center to center closest proximity distance of 20$\mu$m for each ink. As it can be perceived from samples with solely ink $\alpha_2$ (details in Section 3.2.1) and solely ink $\beta_2$ (details in Section 3.2.2) that the feature sizes were of radius 2$\mu$m and 5$\mu$m respectively which is not so large as to cause mixing of both the inks henceforth with 10$\mu$m axial offset in both axis inks were multiplexed. Its fluorescence image can be observed in Figure 3-15 taken from an upright fluorescence microscope.
Figure 3-15 Fluorescence microscopy image for samples with ink $\alpha_2$ multiplexed with ink $\beta_2$

A somewhat similar homogenous pattern of both inks was observed under a lens of 10x showing a mean feature diameter of 3.628 $\mu$m and standard deviation of 1.487 $\mu$m with a superposition of images taken under a Texas Red filter and FITC filter at an exposure time of 15s for each filter. The detailed feature diameter distribution is given in table 3-5 below.

Table 3-5 Feature size distribution of samples with multiplexing of ink $\alpha_2$ and ink $\beta_2$
The approach dot or the first point of contact can be observed to have a comparatively large feature size due to more dwell time and contacting more than once while defining the plane of print with the software to make the pattern for both ink α2 and ink β2.

### 3.2.4 Outlook

There were many takeaways from the moderately charged ink characterization results obtained to have an improved iteration thereafter. Consequently, for the third batch prepared by PPL number of strategies were devised to observed better SICM characterization results, firstly it was thought that since the lipids are printing very well at half concentration of carrier lipid namely DOPC it would best to have an even higher concentration of charged lipids than ~40% as done previously in the respective inks for a high charge contrast. Secondly, as features were not seen well under SICM for the highly anticipated multiplexed version of ink α2 and ink β2 it was proposed to even closely pack the features to observe inside the scan size window of SICM. Lastly, it was projected that multiplexing of inks to have a negative and positive ink side by side was not that good of an idea, in fact a highly positively charged ink would stand out better over the negatively charged glass substrate as per the calibration of SICM done at CeNTech, Munster.

### 3.3 Highly Charged Phospholipid Inks

For the third batch of PPL a highly positively charged lipid ink which was used to draw a high contrast with the negatively charge glass substrate. The ink viz. ink γ with detailed composition provided in Section 3.3.1 was used for this purpose. The ink γ samples showed a nice pattern under fluorescence microscope in the lab at KIT without intermixing of the features for 10µm closest proximity distance.

Figure 3-16 shows the results obtained with BM-SICM with positive DC offset and Figure 3-17 shows the results obtained with BM-SICM with negative DC offset where brightness corresponds to the vertical height of the features. The patches were observed to have a periodic pattern to confirm that they indeed represent lipids.
With negative offset the features were observed with much less noise as in Figure 3-17 as compared to results with positive offset.

### 3.3.1 Ink γ

The samples were prepared with an ink containing 99 mol% of MVL5 (N1-[2-((1S)-1-[(3-aminopropyl)amino]-4-[di(3-amino-propyl)amino]butylcarboxamido)ethyl]-3,4-di[oleyloxy]-benzamide) with structure as shown in Figure 3-18, which was found to deposit over the substrate by trial and error method. It was admixed with 1 mol% of Rhodamine ((lissamine rhodamine B
sulfonyl) (ammonium salt)), with structure already given in Figure 3-3, which was used for its bright red fluorescence and hence easy detection under a fluorescence microscope so that features not visible under a bright field optical microscope can be seen.

![Structural Formula of MVL5](image)

**Figure 3-18** Structural Formula of the employed MVL5 [29]

As maybe observed in the figure above MVL5 can be observed to have a high overall net charge of +5 whereas from the structural formula of Rhodamine in Figure 3-3 it is clear that it has a slight overall negative net charge of -1 in addition of being in low concentration. A predominantly neutral ink with 90% of highly positively charged lipid and 1% of slightly negatively charged lipid tends to be overall highly positively charged and therefore it is expected that it would amount to a reasonable contrast with negative inherent charge over the glass substrate when characterized under SICM to observe the natural topography of lipid in liquid.

For PPL machine a PDMS stamp of 100×50, where the former count signifies the center to center distance in between two consecutive pyramids in µm and the latter count signifies the base length and width of the pyramids’ square base in µm, prepared from its negative Si master fabricated with Electron Beam Lithography (EBL) was mounted. The stamp was then used by the NLP 2000 default software with the help of the precise piezoelectric control to generate ink patterns with the center to center closest proximity distance of 10µm to cater for the small scan size of SICM. To be on a safe side and further ensure that the printing really took place at the micro-scale the samples were observed under upright fluorescence microscope as shown in Figure 3-19.
The ink was seen to be deposited well as a homogenous pattern of lipids with mean diameter of 4.616µm and a standard deviation of 2.684µm under a Texas Red filter with an optical zoom of 10x. The detailed feature diameter distribution is given in table 3-6 below.

The approach dot or the first point of contact can be observed to have a comparatively large feature size especially due to suitable viscosity of MVL5, more dwell time and contacting more than once while defining the plane of print for the software to make the pattern.
3.4 Lipid known to have good adherence to glass substrate

For the AFM topographical analysis of lipids an ink without fluorescence was employed, therefore it was necessary to ensure that the constituent of the ink was previously shown to print perfectly. DOPC with structural formula shown in Figure 3-2 in pure form without admixing any other type of lipid was used as profile for its dip pen nanolithography deposition with respect to the dwell time is already known from a study [16] as displayed in Figure 3-20.

![Figure 3-20 DOPC height profile against DPN contact dwell time over substrate [16]](image)

As the data is only applicable to DPN therefore it is only valid if the PPL stamp is barely touching the substrate as proven in a previous study [14] by which PPL has been shown to operate in 3 different regimes from DPN to hybrid to eventually μCP type mode of operation with increasing contact area of stamp with the substrate as given in Figure 3-21.
An intentional gradient was introduced to observe how the lipid topography differs in both modes of operation. A dwell time of 10 sec taking into account the data given Figure 3-20 was used to safely ensure a definite printing of the lipid over the substrate for detailed SPM characterization thereafter. The chamber had an environment humidity of 25% at room temperature and pressure. For characterization a clear distinction between approach dots or pattern dots was not made and there is a possibility that the feature in question may be from either of the two forms.

### 3.4.1 Completely decompressed PDMS pyramid

Figure 3-22 shows the feature profile of the lipid with PPL operating in the quasi-μCP mode as the contact area was high with a blue arrow shown in the figure for visual aid displaying the amount of decompression of the PDMS pyramid corresponding to the situation in question.
As may be observed in the figure above the contact area is around 2500 micrometer squared which is the total area of single PDMS pyramid of the PPL stamp implying a complete decompression of the pyramid. As the feature is circular it means that quite a lot of spreading already took place before withdrawing the stamp which means that this feature represents the approach dot as the image clearly shows the dwell time was high by the standards of DOPC. The height of the lipid along the cut can be observed to increase in discrete steps and then apparently decrease due to flattening of the image as per the axial cut. The apparent minimum in the middle of the cut is an artifact and this minimum is in fact representative of the maximum height of the feature. After the maximum, an almost mirror image of the first half can be observed. Over the self-assembled monolayer near the edges of the PDMS pyramids used there are visible lipid bilayers stacks one over another mounting up till the maximum in the middle. The asymmetric trench in the middle is a depression as clear reduction of lipid layers around depression can be observed which may be due to uneven movement while in contact or at the time of stamp lift-off. At the maximum, height is slightly more than 30nm from the axial cut, with reference of substrate, which amounts to around 10 lipid bilayers above the self-assembled monolayer as the height of a single lipid bilayer is around 3nm. This matches well with the observable AFM image where the discrete lipid steps can be counted to the same amount.
3.4.2 Partially decompressed PDMS pyramid

Figure 3-24 shows the feature profile of the lipid with PPL operating in the hybrid mode as the contact area was almost half of the base area of PDMS pyramid with a blue arrow shown in the figure for visual aid displaying the amount of decompression of the PDMS pyramid corresponding to the situation in question.

![Figure 3-24 DOPC print with intermediate decompression of the PDMS pyramid [14]](image)

As may be observed in the figure above the contact area is around 1250 micrometer squared which is half of the total base area of single PDMS pyramid of the PPL stamp implying an intermediate decompression of the pyramid. The height of the lipid along the cut can be observed to increase rapidly with discrete steps and then there is an apparent decrease to an almost steady value which again is an artifact as in the last case introduced by flattening. After the midpoint of steady value which in fact represents the maximum height of lipid, an almost mirror image of the first half can be observed. Over the wetting layer near the edges of the PDMS pyramids used as may be observed in the AFM image at the bottom without fingers, over it are more lipid bilayers stacks having fingers over the circumference. At the maximum, height is slightly more than 12nm from the axial cut, with reference of substrate, which amounts to around 4 lipid bilayers above the self-assembled monolayer as the height of a single lipid bilayer is around 3nm.
3.4.3 Decompressed PDMS pyramid tip

Figure 3-25 shows the feature profile of the lipid with PPL operating in the quasi-DPN mode as PDMS pyramid was touching the substrate with its tip decompressed with a blue arrow shown in the figure for visual aid displaying the decompression of the PDMS pyramid tip corresponding to the situation in question.

![Figure 3-25 DOPC print with decompression of the PDMS pyramid tip][14]

As may be observed in the figure above the contact area is orders of magnitude lower than the total base area of single PDMS pyramid of the PPL stamp implying a negligible decompression of the pyramid. The height of the lipid along the cut can be observed to not have sharp edges and a decreasing profile as in the previous cases but instead have an almost steady profile. This means that the artifacts are minimal in this case as the lipid height is not very high hence flattening of the image does not affect much. Not a lot of spreading of ink has taken place as may be evident from the image with feature being more rectangular than circular. A steady value of around 4.5nm is observed which amounts to around 2 lipid bilayers above the self-assembled monolayer as the height of a single lipid bilayer is around 3nm and that of SAM is around 1.5nm.
3.5 Pure Lipid ink deposition without carrier lipid over glass substrate

Having successfully deposited highly positively charged pure lipid MVL5 for SICM characterization opened the potential of PPL in depositing pure lipid inks without the carrier lipid DOPC which is not possible with DPN. An ink without fluorescence with pure DOPE lipid having structural formula shown in Figure 3-26 was employed. Unlike DOPC, the printing properties of pure DOPE using PPL over glass substrate were unknown. DOPC has been constantly used in this study as the carrier lipid known to deposit well over glass but DOPE alone was not used in this study anywhere before this. As in the previous case an intentional gradient like Figure 3-22 was introduced to observe DOPE’s printing in all the regimes over the glass substrate if it takes place in either of the quasi-DPN meniscus type transfer or quasi-µCP stamping type mode of operation as the contact area of stamp with the substrate is increased.

A dwell time of 5 sec to ensure sufficient contact of inked stamp to the substrate was used with chamber humidity of 25% at room temperature and pressure. Subsequently, it was taken for detailed AFM. Again, a clear distinction between approach dots or pattern dots was not made and there is a possibility that the feature in question may be from either of the two forms.

![Figure 3-26 Structural Formula of the employed DOPE [29]](image)

3.5.1 Completely decompressed PDMS pyramid

Figure 3-27 shows the feature profile of the lipid with PPL operating in the quasi-µCP mode as the contact area was high with a blue arrow shown in the figure for visual aid displaying the amount of decompression of the PDMS pyramid corresponding to the situation in question.
For this experiment a stamp with 25µm square base width was used. As may be observed in the figure above the contact area is around 625 micrometer squared which is the total area of single PDMS pyramid of the PPL stamp implying a complete decompression of the pyramid. The height of the lipid along the cut can be observed to display train of impulses in the left half at regular intervals and negligible deposition in the right half. Feature height slightly above 4nm was observed which amounts to around 2 lipid bilayers over the wetting monolayer as the height of a single lipid bilayer is around 3nm.

3.5.2 Partially decompressed PDMS pyramid
Figure 3-24 shows the feature profile of DOPE lipid with PPL expected to operate in a more hybrid mode closer to DPN meniscus type transfer as the contact area was almost half of the base area of PDMS pyramid with a blue arrow shown in the figure for visual aid displaying the amount of decompression of the PDMS pyramid corresponding to the situation in question. However, PPL was still found to deposit somewhat square features and not quasi-circular which means that the deposition still takes place in stamping mode which is why DOPE lipid transfers in the first place because without carrier lipid it is not known to deposit with DPN.
As may be observed in the figure above the contact area is around 400 micrometer squared which is two thirds the total base area of single PDMS pyramid of the PPL stamp implying an intermediate decompression of the pyramid. The height of the lipid is relatively uniform unlike the impulses in the last case. A uniform height of around 4.5nm which amounts to around 3 lipid monolayers as the height of a single lipid monolayer is around 1.5nm.

3.6 Proposed Lipid Quantification Method for future studies
The results in the previous subSections have given enough empirical data to find a pattern in the results and come up with a generic quantification method for lipids. A figurative display deposited lipid layers is given in Figure 3-29 from a study [18] which has then been labeled as suited for the derivation of the quantification method. The layer $L_0$ represents the height of a single layer at the bottom which is the self-assenbled monolayer. $L_0$ is then followed by a the first inverted lipid bilayer namely $L_1$, followed by the second inverted lipid bilayer $L_2$ and so on and so forth.
As a relatable example for instance Figure 3-30 shows 3-D image of the scenario given in Section 3.4.3 with a quasi-DPN mode operation of PPL patterning of DOPC lipid.

The lipid feature can be observed to have discrete heights as limited by the layer thickness which only comes in discrete value i.e. single lipid bilayer has a thickness of ~3nm. To quantify the amount of lipid deposited in discrete steps it is necessary to step-wise integrate the planes.
Therefore, the proposed method integrates for the factor $I_0$ the self-assembled monolayer which it takes as the reference normalization factor for the later layers. For the calculation of $I_0$ a $z$-axis limited volume integral is to be calculated with limits $z=0$ to $z=1.5\text{nm}$ which is the average thickness of a single lipid layer. Since one of the three axes is already defined the volume integral in this case would not be a triple integral but would be reduced to a double integral over $x$-$y$ axis. The $x$-$y$ axis limit of the substrate can be arbitrary as per the requirement of the study and dimensions of the substrate to be calculated. The following formulas are only valid if the substrate surface is perfect lying on the $x$-$y$ plane without any tilt.

$$I_0 = \iiint 1.5 \times 10^{-9} \, dxdy$$

OR

$$I_{n0} = 1$$

The formula for $I_0$ is essentially the integral of the layer $L_0$ as displayed in Figure 3-29 which would amount to a value on the order of $10^{-9}$ or similar so a parallel quantification scheme to make intuitive sense defines normalized integral $I_{n0}$ equal to 1 can be used to get a convenient relative values for the later layers. Similarly, $I_1$ can be calculated for the $z$ limits from $z=1.5\text{nm}$ to $z=4.5\text{nm}$ which would incorporate the first reversed lipid bilayer over the SAM (self-assembled monolayer).

$$I_1 = \iiint 3 \times 10^{-9} \, dxdy$$

OR

$$I_{n1} = I_1 / 2I_0$$

By this scheme $I_1$ would amount to the absolute integral of the layer $L_1$ as displayed in Figure 3-29 which would amount to a value on the order of $10^{-9}$ or similar so a parallel quantification scheme to make intuitive sense defines normalized integral $I_{n1}$ such that it would give a value in between zero to 1 which would intuitively represent the ratio of molecules of lipid in the self-assembled monolayer which also had another lipid on their top in layer $L_1$.

Classically speaking it is impossible to have a lipid layer over air as there needs to be a continuous stack of lipids to support the lipids in the upper layer which is the principle this quantification methods develops upon. Therefore, normalized integral cannot possibly be greater than 1. In contrast, a value of zero would represent the complete absence of the first layer. A negative value is impossible to obtain as per the definition.
Similarly, $I_2$ can be calculated for the $z$ limits from $z=4.5$nm to $z=7.5$nm which would incorporate the second reversed lipid bilayer over the SAM.

$$ I_2 = \iint 3 \times 10^{-9}dxdy $$

OR

$$ I_{n2} = I_2 / 2I_0 $$

By this scheme $I_2$ would amount to the absolute integral of the layer $L_2$ as displayed in Figure 3-29 which would amount to a value on the order of $10^{-9}$ or similar so a parallel quantification scheme to make intuitive sense defines normalized integral $I_{n2}$ such that it would give a value in between zero to 1 which would intuitively represent the ratio of lipid molecules in layer $L_1$ which also had another lipid bilayer on their top in layer $L_2$.

Similarly $I_3$, so on and so forth later layers can be quantified by the generic formula for the $m$’th positive integer given below.

$$ I_m = \iint 3 \times 10^{-9}dxdy $$

OR

$$ I_{nm} = I_m / 2I_0 $$

To end with, this method not only has the capability of quantifying the deposited lipids over the substrate but can also decisively enumerate how much destructing a characterization technique such as SPM techniques i.e. AFM and SICM in this case can potentially be to the deposited lipids. By observing the change in integral values $I_m$ and normalized ratios $I_{nm}$ one can easily observe the worth of these formulas as a tool to compare different characterization methods for the suitability of samples as delicate as the lipids samples. Moreover, the change in characterization parameters e.g. voltage setpoint, DC offsets for AFM to ensure minimum destruction to the samples can be facilitated by means of this quantification method.

This method would particularly come in handy as the research group continues to do the AFM characterization as done in the previous Section but in physiological buffer unlike air as done in this study. This is because in physiological buffer the sample is prone to more destruction as compared to air apparent from the SICM experiments in this study too.
4. Experimental Setup

4.1 Ink
All phospholipids employed in our experiments were obtained from Avanti Polar Lipids, USA, and used as dissolved in chloroform. For the first batch DOPC (1,2-dioleoyl-sn-glycero-3-phosphocholine) was admixed with 10 mol% RHODAMINE (lissamine rhodamine B sulfonyl an ammonium salt) in the first experiment and admixed with 5 mol% DANSYL (1,2-dioleoyl-sn-glycero-3-phosphoethanolamine-N-(5-dimethylamino-1-naphthalenesulfonyl an ammonium salt) in the following experiment for detection with fluorescence microscope. For the second batch DOPC was admixed with 40 mol% DOTAP (1,2-dioleoyl-3-trimethylammonium-propane a chloride salt) and 0.1 mol% RHODAMINE in the first experiment whereas DOPC was admixed with 40 mol% DGS-NTA and 1 mol% PE CF (1,2-dioleoyl-sn-glycero-3-phosphoethanolamine-N-(carboxyfluorescein an ammonium salt) for the following experiment. We chose New Multivalent Cationic Lipid for siRNA Delivery MVL5 (N1-[2-((1S)-1-[((3-aminopropyl)amino]-4-[di(3-amino-propyl)amino]butylcarboxamido)ethyl]-3,4-di[oleyloxy]-benzamide) a highly positively charged lipid for the third batch. When hydrated with chamber humidity control, the ink allowed control of its diffusion and transport properties. Note that for the third batch MVL5 was admixed with 1 mol% of the RHODAMINE labeled phospholipids to facilitate detection of the lipid membranes by fluorescence microscopy. The admixing of inks were done inside the fume hood displayed in Figure 4-4.

4.2 PPL Stamp Fabrication
The PPL pen arrays were prepared using PDMS polymer solution as described in literature with some variation [14]. To reiterate it was prepared by mixing 3 g of vinyl- compound-rich prepolymer (ABCR, Karlsruhe), 20 μL of 2,4,6,8- tetramethyltetravinylecycloketatlosiloxane (Sigma-Aldrich, Germany) and a drop of platinum catalyst (platinum divinyltetramethyl disiloxane complex in xylene, Gelest, USA). The polymer solution was stirred in a falcon tube and put in a desiccator shown in Figure 4-1 at low pressure (~0.1 bar) for 15 min. to degas. An amount of 1 g (~30% methylhydrosiloxane)- dimethylsiloxane copolymer (ABCR, Karlsruhe) cross-linker was then mixed into the h-PDMS solution, stirred, degassed again, and poured onto the silicon master with the desired pyramidal array designed by Electron Beam Lithography as shown in Figure 4-2. Standard microscopic glass slides (VWR, Germany) were cut to pieces slightly bigger than the stamp sizes in the Si-master to ensure an easy peel off the next day using
a scalpel. The glass slides were carefully placed on the filled Si master and pressed down to fill the inverted pyramids of the Si-master homogeneously and to ensure that none of the air bubbles remain. The PDMS polymer was cured overnight on a hot plate as shown in Figure 4-3 at 70 °C. Next day the residual PDMS over the stamp was carefully removed from the side of glass and the stamp array was detached from the master by using a scalpel. Thereafter the excess PDMS which could possibly affect printing results was removed from the microscopic glass slide stamp.

![Desiccator used for the experiment](image1)

**Figure 4-1** Desiccator used for the experiment

![Si masters used for the experiment](image2)

**Figure 4-2** Si masters used for the experiment
4.3 Substrates

The glass substrates were prepared with slight variance from the method described [15] as glass (standard coverslips, VWR, Germany) were cleaned by ultrasonication submersed for 10 min in chloroform and then for an additional 10 min in isopropanol. Thereafter the coverslips were dipped into nanopure water (18.2 MΩ cm) several times and dried with nitrogen blow. All the aforementioned steps were conducted inside the fume hood with as displayed in Figure 4-4, along with the experimental setup for substrate preparation, for safety purposes. The coverslips were then used as substrate for the printing experiments.

For the AFM, Surface enhanced ellipsometric contrast (SEEC) glass substrates with $\lambda/4$ optical multilayered solutions as reported in [9] which keeps the polarization state of the incident light unchanged on naked substrate which greatly enhances the imaging contrast. It nevertheless controls the polarization state of light locally air or liquid where the deposition in our case of lipid takes place to aid label-free visualization of the sample with as high as an axial density in z-dimension as ~0.1 nm for the substrates used here manufactured by Nanolane.
4.4 Printing Process

Patterns were printed with an NLP 2000 system (NanoInk, USA) as reported previously [15]. The machine used for this study is shown in Figure 4-5. Before inking, stamps were treated with oxygen plasma (0.2 mbar, 100 W, 10 sccm O₂, 2 min, ATTO system, Diener electronics, Germany) to enhance hydrophilicity of the surface. For each experiment stamps were inked by spin-coating 2ul of respective inks (30 s at 3500 rpm) to spread the lipids uniformly by the spin coater on the right most edge of the fume hood in Figure 4-4. The polymer pen array was pasted on the holder of an NLP 2000 instrument (NanoInk, USA) by two-component epoxy given in Figure 4-6. The holder shown at the bottom right of Figure 4-7 is custom made in a workshop as by default NLP 2000 is used for SPT instead of PPL which is in fact the primary focus of this study, so a different holder was developed. A sacrificial cover slip is consumed for parallel alignment of stamp and substrate by leveling as the pen array was tilted with respect to the substrate with the internal controls of the printing setup as appropriate until sufficient balance is achieved. Ink was printed in such a way as to create an array of dots with minimum of 20μm horizontal and vertical distance apart for first 2 and last 2 batches, and 10μm horizontal and vertical distance apart for the third batch.
Figure 4-5 NLP 2000 setup used for patterning in all the experiments of the study

Figure 4-6 Two component UHU epoxy used to paste the stamp over the holder
Figure 4-7 Customized holder for PPL on bottom right and glass slides used as the base for stamp on the left-most side

4.5 Characterization Parameters

The Fluorescence Microscopy, Scanning Ion Conductance Microscopy (SICM) and Atomic Force Microscopy (AFM) in air was employed to characterize the phospholipids deposited over the substrate the details of which are given in the following sub-Sections.

4.5.1 Fluorescence Microscopy

Micro-graphs are taken with the DAPI, FITC and texas-red channels of an upright microscope for the first batch as shown in Figure 4-8.
For the second batch inverted microscope shown in Figure 4-9 with the Texas red filter was used for immediate confirmation of whether printing took place before going for detailed SICM characterization. The exposure time and zoom value is given in each Section as per the case.
For post image processing ImageJ was used to process the images from fluorescence microscopy as given in section where results are discussed. Auto thresholding was used to obtain the area of the lipid features which stood out due to fluorescence over the reference background substrate. Thereafter area of contours with circularity of greater than 0.1 was obtained and then their diameter was displayed as a histogram distribution.

4.5.2 SICM

SICM studies presented here were performed by an NX-Bio (Park Systems, Suwon, South Korea). SICM measurements were done in amplitude-based BM-SICM mode [17] provided by Park Systems both with raster scanning method, not with spectroscopic hopping scheme [23, 25-27]. Typical scanning speed for these two modes was selected to be 0.3 Hz. For amplitude-based BM-SICM mode applied AC voltage signal was with drive of 50 mV and 1 kHz frequency (where we can get the clearest topographic images), and its responsive amplitude of measured AC current signal was used as feedback value for topographic imaging of our standard samples [17]. Although McKelvey et al. found that phase signal from detected AC current was more sensitive for recognizing tip-to-sample distance [17], we cannot use this mode because of its saw-tooth like artifact in low frequency (~1 Hz) upon the monitored phase value with unidentified reason. 1 kHz of input AC voltage frequency was still enough for observing topographic change induced by surface charge from our standard samples according to the applied DC offset voltages. Control over this amplitude-based BM-SICM mode was done with SmartScan (ver. 1.0 Build 74 RTM8a, Park Systems, Suwon, South Korea. The SICM images were flattened and analyzed with the XEI software program (ver. 4.3.0.Build5, Park Systems).

At first, we have examined the reliability of our amplitude-based BM-SICM mode with a topographic standard sample in a raster-scanning manner. This standard sample was made with casted PDMS upon the calibration grating (TGZ1, NT-MDT, Moscow, Russia, 20 nm height with 3 um lateral period). BM-SICM mode produced clear topography for this sample (without applying any DC offset) as expected. Throughout the whole measurements with SICM, regardless of the mode under use, the setpoint was chosen to be 1% decrease from the bulk value of monitored physical quantities (AC amplitude for BM-SICM mode, After checking that this BM-SICM mode is trustful, we moved onto standard samples for surface charge mapping of PPL-lipid patches (positively or negatively charged as the case may be) both upon the glass substrates (negatively charged within neutral pH aqueous solution), which is designed to show well-defined electrostatic charge contrast from patterned region to the remaining glass.
Throughout whole measurements, no meaningful difference in relative amplitude noise level has been found at these opposite DC offset polarities. For the case of the lipid samples, typical thickness of patterned layer was in the range of 3–5 nm, indicating that pattern consists of just a few layers of phospholipids. Even though our lipid ink molecules have more positive charges per a head (+4).

Till now, we have shown amplitude-based BM-SICM studies on topographic change induced by surface charge from its standard samples made constituting lipids. Our lipid patterns meet all needs required for other established SPM measurement schemes, well-defined, periodic topography and surface charge in large area (larger than mm² scale). BM-SICM mode was found to be better at its stability, image quality, and reproducibility with samples imaged at a frequency of 1 kHz, an input bias of 50mV while the sample was immersed in 0.15M KCl solution with a pH value of 7. The sample was traversed over a scan size of 15µm×15µm. The pipette current was adjusted to a value of 800 pA and 1% setpoint value. The values were recorded at different DC offset for different batches as per the imaging requirement to obtain optimized image. Finally, it proves that this amplitude-based SICM mode could be used as a reliable and robust scheme for surface charge mapping studies within physiological conditions.

4.5.3 AFM

For a high-resolution and detailed characterization of the patterns of samples from fourth and fifth batch AFM was employed on a Dimension AFM system (Bruker, Germany) in tapping mode with HQ:NSC15/Al BS cantilevers (MikroMasch, USA) with 325 kHz nominal resonance frequency. All measurements were done in air under ambient conditions. The images were taken with PeakForce QNM mode. Image processing and export was carried out in the onboard software package. The parameters such as setpoint, scanning frequency etc. were adjusted such that the best possible image by causing the least possible destruction while traversing the sample occurred. These values differed for fourth and fifth batches that were characterized by AFM and even further differed depending on the features deposited in either quasi-DPN, hybrid or quasi-μCP mode. The case specific details of the parameters are given below.
DOPC features with Quasi-DPN mode:

![Figure 4-10 AFM parameters for DOPC features with Quasi-DPN mode](image)

DOPC features with hybrid mode:

![Figure 4-11 AFM parameters for DOPC features with hybrid mode](image)
DOPC features with Quasi-\(\mu\)CP mode:

Figure 4-12 AFM parameters for DOPC features with Quasi-\(\mu\)CP mode

DOPE features with Quasi-DPN mode:

Figure 4-13 AFM parameters for DOPE features with Quasi-DPN mode
DOPE features with Quasi-μCP mode:

Figure 4-14 AFM parameters for DOPE features with Quasi-μCP mode
5. Conclusion

PPL was found to successfully print phospholipids over glass substrate with the control to print in the operating mode of either of its precursor namely DPN and µCP. With the perfectly parallel levelling of the PPL stamp with the substrate almost equal precision to that of DPN was noticed as the stamp master manufactured by EBL is also very accurate in terms of distance in between the eventual PDMS pyramids used to deposit features. The adsorption of phospholipids head with glass substrate forming SAM stacked over on top by lipid bilayers was found to have step-like discrete patterns and understandably so because of the amphiphilic molecular structure of the lipids employed. A theoretical method is proposed which not only quantifies the stacked phospholipid layers in physiological solution, mimicking its natural state being the main constituent of cell membrane in blood plasma, but also has the potential to quantify the Scanning Probe Microscopy (SPM) probe which is used to characterize the sample such that the probe can be said to have certain factor which represent the destruction caused to the particular phospholipid feature while traversing over the sample in water.

As for the characterization results, SICM was found to give best results for phospholipids with highly positively charged headgroup as it gave rise to the best contrast in terms of charge with the glass substrate background. The AFM images clearly showed bigger features with larger contact area of PDMS pyramid touching the substrate at the time of deposition. This observation is consistent with the fluorescence imaging results from a previous study [14] and the printing of DOPE via PPL is a successful demonstration that PPL can be used to print lipids (pure) that cannot be written via DPN. The AFM studies revealed that this is because the PPL does a more µCP mode in this case (compared to the more DPN mode with DOPC), which is not obtainable in DPN.

Although the SICM measurements were done with samples immersed in a physiological buffer but all AFM results in this study were obtained with samples in air which is not reminiscent in terms of explaining lipids’ natural behavior e.g. being main constituent of plasma membrane in blood plasma. So, with the opening of gateways to precisely print such lipids with PPL which are inconceivable to otherwise write via DPN and research group at the hosting institute
continuing SPM characterization of lipids inside physiological buffers a lot of potential in terms of contributing to the field of science and medicine is expected in the near future.
6. References


## LIST OF ABBREVIATIONS

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Full Form</th>
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<tbody>
<tr>
<td>DPN</td>
<td>Dip pen nanolithography</td>
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<tr>
<td>PPL</td>
<td>Polymer pen lithography</td>
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<tr>
<td>AFM</td>
<td>Atomic force microscopy</td>
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<tr>
<td>SEM</td>
<td>Scanning electron microscopy</td>
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<tr>
<td>PE</td>
<td>Printed electronics</td>
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<td>PDMS</td>
<td>Poly-Dimethylsiloxane</td>
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<td>NMR</td>
<td>Nuclear magnetic resonance</td>
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<tr>
<td>ICs</td>
<td>Integrated Circuits</td>
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<tr>
<td>MEMS</td>
<td>Micro electromechanical systems</td>
</tr>
<tr>
<td>SPM</td>
<td>Scanning probe microscopy</td>
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<tr>
<td>μCP</td>
<td>Microcontact Printing</td>
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This thesis is dedicated to all humankind with a humble request to have mutual respect and of propagating kindness, love and tolerance.