Novel Principle for 2D Molecular Self-Assembly: Self-Intermixed Monolayer Phases of Sub-Phthalocyanine and C$_{60}$ on Ag(111)

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Abstract

In this thesis, a novel route to highly perfect bi-molecular self-assembly on a metallic surface is presented. Two kinds of organic molecules, the polar Chloro-[subphthalocyaninato]-boron(III) (SubPc) and the polarizable C_{60}-Buckminster fullerene, were co-deposited onto atomically clean Ag(111) under UHV conditions. Depending on the relative surface coverage of the two species, different well-ordered intermixed monolayers are formed: At a mixing ration SubPc/C_{60} = 2:3, an anisotropic stripe structure consisting of C_{60}-chains with an inter-chain periodicity of 3.3 nm is found. A mixing ration SubPc/C_{60} = 1:1 leads to a 2D hexagonal pattern with periodic vacancies. At room-temperature, the structures spontaneously emerge by self-organization and are stable at the same. The mixed structures show interesting and unusual features. One of the ordered bi-molecular overlayers, the 3:2-stripe-phase, exists in two chiral forms, although the individual molecules themselves are achiral. Different phases can co-exist simultaneously, and also a 2D molecular gas phase can be observed. The appearance of the molecular stripe domains in STM measurements strongly depends on the applied bias-voltage. The structural parameters and the schematic binary “phase-diagram” of this system are deduced from detailed room-temperature Scanning Tunneling Microscopy (STM) studies. The underlying interactions and the relevant properties of the molecules are discussed, based on STM, XPS and UPS measurements and on numerical simulations. The pattern formation is further treated under thermodynamic aspects of a two-component mixture. A counter-example to the intermixed system is further presented: silated Perylen and C_{60} molecules, co-adsorbed on clean Ag(100), do not intermix but lead to a 2D segregation.

These novel Self-Intermixed Monolayer Phases (SIMP) are different from previously known self-assembled molecular monolayers in that they form intermixed patterns at room temperature on uniform, unreconstructed atomically clean terraces. These self-organized patterns, as well as the additionally shown conformational changes of Porphyrin molecules, inspire to be incorporated in building blocks of possible molecular electronic devices.
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Abbreviations

1D, 2D, 3D One-, two, three-dimensional
AFM Atomic force microscope
BE Binding energy
C_{\infty} C_{\infty} Buckminster fullerene
Cu-TBPP Cu-tetra-[3,5 di-ter-butyl-phenyl] porphyrin
DOS Density of states
E_F Fermi-energy
ESCA Electron spectroscopy for chemical analysis
hcp Hexagonal close-packed
HOMO Highest occupied molecular orbital
KE Kinetic energy
LB Langmuir-Blodgett
LUMO Lowest unoccupied molecular orbital
LEED Low-energy electron diffraction
MBE Molecular beam epitaxy
ML Monolayer
OMBE Organic molecular beam epitaxy
Perylen N, N', N''-Tetra(trimethylsilyl)-4,9-diamino-perylenchinon-3,10-diimin
PES Photoelectron spectroscopy
Pc Phthalocyanine
SAM Self-assembled monolayers
STM Scanning tunneling microscope
STS Scanning tunneling spectroscopy
RT Room-temperature
SIMP Self-intermixed monolayer phase
SubPc Chloro[subphthalocyaninato]boron(III)
UHV Ultra-high vacuum
UPS Ultraviolet photoelectron spectroscopy
XPS X-Ray photoelectron spectroscopy
1. Introduction: Motivation and Outline

In the natural world, molecules are used for many purposes. Using molecule-based materials for electronics, sensing, and optoelectronics is a new endeavor, called molecular electronics, and the subject of riveting new research [1–4] and substantial popular press interest. A convenient definition of molecular electronics by M.A. Ratner [5] is the set of electronic behaviors in molecule-containing structures that are dependent upon the characteristic molecular organization of space. Early work focused on some visionary [6] or speculative [7] ideas and mechanistic suggestions, showing in the early 70ies that it is theoretically possible to use a single molecule as a rectifier. Some truly pioneering measurements of charge transfer across molecular adlayers [8] were done in 1971 and S. Roth [9] demonstrated an experiment in 1995 with an donor-acceptor molecule within a Langmuir-Blodgett film that behaved like a diode.

There is currently great interest in the adsorption of large organic molecules on inorganic surfaces. Partly, this activity is due to the increasing use of organic semiconductors for technological purposes. Organic light emitting diodes [10–12] and thin films transistors [13, 14] may serve as examples in this context. A recent break-through was achieved by designing logic circuits with field-effect transistors on the basis of single carbon nanotubes [15]. At the same time, a fundamental interest arises to extend the notions formed by myriad investigations of small molecule adsorption to the physisorption of larger adsorbates, which involve a higher degree of functionalization and can be designed by concepts of Supramolecular Chemistry [16–18]. An important class of molecular device functionality is based upon reversible changes of molecular conformation. One can argue that conformational changes in molecules represent the ultimate limit of nanomechanics.

The advent of self-assembly methodologies [19, 20] for efficient arrangement of a large number of molecules and the development of scanning probe techniques [21–24], both to prepare molecular nanostructures and to characterize their physical and electrical properties, made prototype molecular electronic devices into a reality.

The field of molecular electronics seeks to use individual molecules to perform functions in electronic circuitry now performed by semiconductor devices [25]. Individual molecules are hundreds of times smaller than the smallest planar features conceivably attainable by semiconductor technology. Because it is the area taken up by each electronic element that matters, electronic devices constructed from molecules will be hundreds of times smaller than their semiconductor-based counterparts. Moreover, individual molecules are easily made exactly the same by the billions and trillions. The dramatic reduction in size, and the sheer enormity of numbers in manufacture are the principle benefits offered by the field of molecular electronics.

In order to produce ultra-small structures which provide some functionality to our decimeter world, two key problems need to be addressed: First these devices need to be manufac-
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tured by efficient (parallel or self-assembled) techniques, and second, they need to be interconnected and “wired” at least to the micrometer level which is accessible by state-of-the-art technology.

The long-term goal is to provide moderate computational power and high-density memory in an extreme small, low-power format, which will not require multibillion-Euro fabrication facilities. The conventional CMOS technique may stall at 10 nm, depending on the fabrication economy, because the production has to be extremely accurate. This is meant by the often quoted “breakdown of Moore’s law”. In the mid 1960s, Gordon Moore, then working as a Director of R+D for Fairchild Semiconductor, came out with his famous quote to the press. He stated that he expected transistor density per Integrated Circuit to double every 18 - 24 months (“Moore’s Law”, see figure 1.1). It seems as if Gordon Moore’s vision, as the co-founder of Intel Corporation in 1968, was right up to now: The new Northwood core Pentium 4 processors have more than 55 million transistors. But not only the afore mentioned economic reasons, but also fundamental physical principles like leaking tunneling currents between two narrow structures could lead to a conversion of the production methods towards molecular electronics.

![Figure 1.1: The number of transistors integrated into Integrated Circuits is growing exponentially since the seventies following the prediction of Gordon Moore in 1968. For its continuity, however, new technologies might be needed. Data from [26]](image)

As a highly interdisciplinary field, progress depends on successful interactions across the boundaries of traditional subject areas. Molecular electronics encompasses biology, chemistry, computing, electronics and physics, each subject with its own methodology and jargon. Most importantly, there is a huge potential for research and discovery, which has motivated National and International Research Programs to collaborate with Multi-National Corporations in one or the other field. The close collaboration of scientists and technologists from many different fields propels nanoscience and nanotechnology.

During this PhD thesis, a novel principle of self-assembly was investigated which leads to nanostructures that have the potential to serve as building blocks for molecular electronic devices. The bi-molecular system Subphthalocyanine (SubPc) and C₆₀, co-deposited
on the clean Ag(111) surface, self-organizes into specific nanometer-size structures. The relative surface coverage of the two species determines the geometry of the observed patterns, which could in principle serve as the basis for storage devices at room-temperature. The conformational switching of single Porphyrin molecules gives similar ideas towards storage applications.

The outline of this thesis is as follows: Chapter 2 describes the ultra-high vacuum system which has been used to investigate single molecules at atomically clean surfaces. The preparation of the samples is described, including the substrate fabrication and the molecular deposition. The main analyzing tools, the Scanning Tunneling Microscope (STM) and the Electron Spectroscopy for Chemical Analysis (ESCA), are briefly explained. A short overview over the computational methods closes this chapter. In Chapter 3, the binary Self-Intermixed Monolayer Phases (SIMP) are discussed. It starts with the description of both individual components, SubPc and C₆₀, and then shows the various phases with their specific compositions. The adsorption geometry of all phases with their properties like induced chirality are further discussed. The chapter ends with the schematic room-temperature “phase diagram”. A detailed discussion of the underlying intermixing mechanism is given in Chapter 4, followed by a general consideration of thermodynamics of mixtures. It is not clear a priori whether two components mix or segregate. The phase behavior here is studied under thermodynamic aspects. Also given is a counter-example by a segregating binary system which does not lead to exciting intermixed structures. Chapter 5 gives a few examples of potential applications of the created patterns, including ideas for a continuation of this work. Appendix A presents an example of the attractive field of molecular conformation and its relevance in a larger context.
2. Instrumental Methods and Set-up

In this chapter, the experimental methods and the Nanolab at the University of Basel are described. In section 2.1, the UHV system is presented. In section 2.2, details on the Ag(111) and Ag(100) substrate preparation are given. Section 2.3 describes the molecular deposition system. The main tool used during this thesis is the scanning tunneling microscope (STM), described in section 2.4. Chemical analysis of the sample surface and measurements of the valence band were done by photoelectron spectroscopy, discussed briefly in section 2.5. Finally, an introduction to computational methods to investigate quantum molecular problems is given in section 2.6.

2.1. UHV System

Ultra high vacuum (UHV) is indispensable to study atomic processes at surfaces because it extremely reduces the density of contaminants. Under UHV condition with $10^{-11}$ mbar pressure, the number of molecules in the vacuum is so low that the mean free path length is enlarged from the order of $10^{-8}$ m at atmospheric pressure to the order of km [27]. The number of collisions at the sample surface therefore is reduced to a minimum and enables to study processes on atomically clean surfaces during several hours or days.

The Nanolab, see figure 2.1 and 2.2, consists of several individually pumped chambers which are separated by gate-valves. The chambers contain the tools for the various steps of sample preparation, molecular and metal deposition, analysis and sample transfer to the outer world [28–33]. All steps were performed in-situ. The partition into different chambers assures that the individual processes do not interfere with each other. An ingenious transport system allows the samples to reach all chambers.

2.2. Substrates

Well-defined substrates serve as support for the molecular layers. For practical uses, either single crystals with a certain orientation, see figure 2.3, or perfect grown metal films can be used. For this study, Ag(111) and Ag(100) films (see figure 2.3) were produced in-situ. After analyzing the metallic films, they were used for many experiments with molecules.

Ag(111) Preparation

The Ag(111) films with a surface lattice constant of 0.289 nm were grown hetero-epitaxially on air-cleaved Muscovite 2M-1 mica substrates with predeposited Au pads. After introduction to UHV, the mica substrate was first annealed at 840 K for 2h in order to obtain sharp
and clear hexagonal LEED spots. The silver layer was then deposited with an electron beam evaporator to a thickness of 120 nm, at a rate of 1 nm/sec and a pressure of $6 \cdot 10^{-8}$ mbar. The temperature of the mica during deposition was held at 570 K. Silver is reported to grow with atomically smooth surfaces [35] at this temperature, consisting of large step-free terraces. The Au from the contact pads is not visible in XPS measurements after the Ag deposition.

Before every new molecular experiment, the Ag(111) surface was cleaned by standard UHV techniques (repeated $Ar^+$ ion etching by 600 eV ions and post annealing to 573 K) which gave a well-ordered LEED pattern.¹

**Ag(100) Preparation**

Atomically clean Ag(100) surfaces were also used as substrates. In a first step, a GaAs(100) sample from a commercial wafer was introduced into UHV via the Fast Entry Air Lock. It then was heated to 850 K ($p < 1 \cdot 10^{-8}$ mbar) until sharp LEED spots were visible and indicated a clean, well-ordered crystallographic structure without contamination. Then a 1 nm thick Fe seed layer was deposited by MBE at 380 K with a rate of $0.01 \text{nm/sec}$ ($p \approx 3 \cdot 10^{-8}$ mbar). After that, a 150 nm thick Ag layer was epitaxially deposited at a constant rate of 0.1 nm/sec and 380 K sample temperature ($p \approx 1 \cdot 10^{-8}$ mbar). Finally, the Ag(100) sample was annealed at 620 K for 2 h and then analyzed in-situ by LEED, XPS and STM.

¹On the films, in contrast to single crystals, rings in the LEED pattern indicate different domains.
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![Image of Nanolab]

Figure 2.2: 3D model of the Nanolab exposing the manifold features of the system. (Technical drawing by A. Heuri)

The films are characterized by atomically flat areas and large terraces (hundreds of nm), which grow in different domains.

Before every new molecular experiment, the Ag(100) surface was cleaned by standard UHV techniques (repeated Ar⁺ ion etching by 600 eV ions and post annealing to 573 K) which gave a well-ordered LEED pattern.

### 2.3. **Organic Molecular Beam Epitaxy (OMBE)**

Ultrapure (99.99%) molecules are thermally deposited (by heating to 570–650 K) from tantalum crucibles at a rate of 0.5 to 1 ML min⁻¹ onto atomically clean substrates that are kept at room-temperature. A chimney on the button heaters collimates the sublimating molecules into a narrow solid angle. The sublimators are outgased for a long time in order to avoid impurities while depositing the films. The coverage, typically in sub-monolayer range, is measured with a water cooled microbalance at an accuracy of some percent of a
monolayer (ML). The film thickness can later be calibrated by core-level intensities in XPS measurements and coverage estimation in STM images. The pressure during the molecule-sublimation can rise up to $\approx 2 \cdot 10^{-8}$ mbar.

2.4. Scanning Tunneling Microscopy (STM)

The Scanning Tunneling Microscopy (STM), invented by Binnig and Rohrer in 1981 [22] (receiving Nobel Prize in 1986), delivers pictures of solid surfaces with atomic resolution. A direct real-space image of a surface is obtained by moving a tiny metal tip across a sample surface and recording the electron tunnel current between tip and sample as a function of the position [21, 36-39]. This finally makes it a powerful tool for us to study the orientational configurations of molecules grown on solid surfaces.

Tunneling is a genuine quantum mechanical effect in which electrons from one conductor penetrate through a classically impenetrable potential barrier – in the present case, the vacuum – into a second conductor [21, 36-40], see figure 2.4. The phenomenon arises from the “leaking out” of the respective wave functions into the vacuum and their overlap within classically forbidden regions. This overlap is significant only for atomic-scale distances and, in the first-order perturbation theory, the tunnel current, $I_T$, when a bias voltage $V_{bias}$ is applied between the two electrodes, is expressed as

$$I_T = (4\pi^2e/h) \sum_{s,t} |M_{st}|^2 [f(E_t) - f(E_s)] \delta(E_l + |e|V_{bias} - E_s)$$  \hspace{1cm} (2.1)

where $f(E) = [1 + e^{-E/\vartheta}]^{-1}$ is the Fermi function, $M_{st}$ is the tunneling matrix element between states $\psi_s$ and $\psi_t$ of the left and right electrodes respectively (tip and sample), calculated independently, $E_s$ is the energy of $\psi_s$ relative to the left-electrode Fermi level, and $E_t$ is the energy of the $\psi_t$ relative to the right-electrode Fermi level. The height of the potential barrier is the work function $\phi$. For most purposes, the Fermi functions can be replaced by their zero-temperature values, i.e. unit step functions. In the limits of small voltages $V_{bias} \ll \phi$, this expression then further simplifies to
2. Instrumental Methods and Set-up

\[ I_T = \left(4\pi^2 e^2 V_{\text{bias}} / h\right) \sum_{i,t} |M_{st}|^2 \delta(E_s - E_F) \delta(E_t - E_F). \] (2.2)

Bardeen [41] showed that, under certain assumptions, the matrix element in equation (2.2) can be expressed as

\[ M_{st} = \left(h / 8\pi^2 m \right) \int dS(\psi^*_s \nabla \psi_t - \psi^*_t \nabla \psi_s) \] (2.3)

where \( m \) is the electron mass and the integral is over any surface lying entirely within the barrier region and separating the two half-spaces. Now the ideal STM tip would consist of a mathematical point source of current, whose position we denote \( \mathbf{r}_T \). In that case, equation (2.2) for the current at a tip-sample separation distance \( d \) and small voltage \( V_{\text{bias}} \) would reduce to [42, 43]

\[ I_T \propto \sum_s |\psi_s(\mathbf{r}_T)|^2 V_{\text{bias}} \delta(E_s - E_F) \equiv \rho_{x,y}(\mathbf{r}_T, E_F) V_{\text{bias}} e^{-2\kappa d}, \quad \kappa = \sqrt{8m\pi^2 \phi / h} \] (2.4)

Thus the ideal STM would simply measure \( \rho_{x,y}(\mathbf{r}_T, E_F) \), the local density of states (DOS) at \( E_F \). The exponential decay of the signal leads to the unique resolution in tip-sample separation. Typically the current decays about one order of magnitude per Å. It is important to see how far this interpretation can be applied for more realistic models of the tip. Tersoff and Hamann [42, 43] showed that equation (2.4) remains valid, regardless of tip size, as long as the tunneling matrix elements can be adequately approximated with those for an s-wave tip wave function [38]. The tip position, \( \mathbf{r}_T \), must then be interpreted as the effective center of curvature of the tip, i.e. the origin of the s-wave which best approximates the tip wave functions.

All images in this thesis were recorded using the constant current mode (“topographic” mode). In this mode, a feedback loop system is used to force the tip via a piezoelectric driver to be always at such a distance to the sample surface (\( \approx 1\,\text{nm} \)) that the tunneling current flowing between these two electrodes remains constant. By recording the voltage, which has to be applied to the piezoelectric driver in order to keep the tunneling current constant, i.e. recording the height of the tip \( z(x,y) \) as a function of the position, a “topographic” image can be obtained in this mode. This mode can be used for surfaces, which are not necessarily flat on the atomic scale, thus for stepped surfaces or adsorbate covered surfaces. The topographic height of the tip movement can be obtained directly, provided the sensitivity of the piezoelectric driver elements is known. A disadvantage of the constant current mode is the finite response time of the feedback loop, which limits the scan speed. A schematic illustration of the constant current mode is shown in figure 2.5. Another mode of operation, which was not used for this study, is the constant height mode. In this mode the tip is scanned at constant height over the sample surface while the feedback loop is slowed or turned off completely. The variations in the tunneling current, which are recorded as a function of location, then contain the height information. An advantage of this mode is the
faster scan rate. This mode, however, is only applicable to atomically flat surfaces, thus not suited for molecular imaging.

For metals the local DOS \( \rho_{x,y}(r_T, E_F) \) is typically in good agreement with the topology. For adsorbates, however, the topology and the image obtained in constant current mode can differ completely due to a modified electronic situation. CO on Cu(211), for example, can appear as a protrusion or depression depending on the arrangement of the adsorbates and the presence of a CO molecule at the tip [44]. Similarly, the contrast of larger organic molecules is known to depend on its specific substituents [45]. The molecular corrugation also can be reversed upon tunneling within a specific range of the bias voltage [46]. A recent review [47] discusses the STM-contrast on adsorbates.

All measurements were done in-situ by a home-built room-temperature STM, using an electro-chemically etched tungsten tip which was cleaned in vacuum by electron bombardment heating. The voltage \( V_{bias} \) corresponds to the sample bias with respect to the tip, see figure 2.4. Here, tunneling from the tip into the unoccupied states of the sample was applied. For molecules this means tunneling into the LUMO. In the images, the slow scan direction is from bottom to top. The line scans were recorded in forward (left to right) and backward direction, with identical tunneling parameters. The architectural and electronic specifications of the STM used in the Nanolab are described in [28]. Details of the Eddy-current damping and the in-situ tip-change system can be found elsewhere [48].

2.5. Photoelectron Spectroscopy (PES)

Photoemission is nowadays one of the most widely used experimental methods in order to obtain important information of matter such as chemical composition, geometrical or electronic structure [49]. The phenomenon was discovered in 1887 by Hertz [50] and had become controversial for more than twenty years. In 1905, Einstein finally explained the
2. **Instrumental Methods and Set-up**

observation of Hertz by inclusion of the quantum nature of light [51], leading him to receive the Nobel Prize in 1922.

The principle of a photoemission process is sketched in figure 2.6. When matter is irradiated with photons, electrons may be excited to higher states, and if the excited electrons have an energy exceeding the work function, they can be emitted from the material. These electrons are called *photo-electrons* and their kinetic energy in vacuum, $E_k$, is given by the relation, (energy conservation)

\[
E_k = h\nu - E_B - \phi
\]  

where $h\nu$ is the photon energy, $\phi$ is the work function of the solid and $E_B$ is the binding energy of the initial electronic state. The binding energy is referenced by the Fermi level, $E_F$. Depending on the photon energy used, information on core levels (X-ray) or valence levels (ultraviolet, UV) of the sample electronic state is obtained. While the photoemission itself is described by Einstein, there have been many detailed theoretical studies on the process [49]. A semi-empirical approach [52], so-called three-step model, has been proven extremely useful in describing the photoemission from as solid. Unlike STM, this is a non-local method, but it allows quantitative analysis of the chemical composition and environment on macroscopic scale.

Photoemission measurements were performed with an ESCALAB MK II from V.G. Scientific Ltd., GB, which is equipped with a hemispherical analyzer, aligned with a fixed angle to the incident photon beam. The angular acceptance can be controlled by an adjustable aperture. All photoelectrons come from within an area of about 4 mm by 4 mm. The energy

![Figure 2.5: Principle operation of the STM. (Schematic: distances and sizes are not to scale.) The tunneling current $I_T$ which flows between the tip and the atomically flat and conducting sample when a constant bias voltage $V_{bias}$ is applied, decays exponentially with the tip-sample separation. The tip motion in the scans with constant current mode, controlled by a feedback loop and piezoelectric elements, is indicated over a surface step.](image-url)
scale is referenced towards the Fermi level $E_F$ of a bulk Ag sample.

For X-Ray Photoemission Spectroscopy (XPS), photo-energies $>100$ eV allow the measurement of the binding energy of core-levels states, chemical composition of surfaces and the study of chemical shifts which depends on the environment of the elements. XPS was done by using the Mg-K$_\alpha$ line with a photon energy of 1253.9 eV. The probing depth of XPS is in the order of 20-40 Å [53]. The molecular coverage of the sample was obtained by comparing the area of the Cls peak with the double peak of Ag3d [54].

For UV Photoemission Spectroscopy (UPS), photo-energies $<100$ eV allow the study of valence bands and occupied molecular orbitals. Nonmonochromated He-I ($h\nu = 21.2$ eV) radiation was used for UPS measurements. Since UPS has a very low information depth (typically 5-10 Å, due to inelastic scattering [53]), we here focus on the contact region between the molecules and between the molecules and the substrate.

![Diagram](image)

**Figure 2.6:** Principle of Photoelectron Spectroscopy. The irradiated matter emits electrons with kinetic energy $E_K$ if the photon energy $h\nu > E_B + \phi$ and is thereby mapping the occupied energy levels of the solid.
2. Instrumental Methods and Set-up

2.6. Introduction to Molecular Modeling

Molecular modeling is the science (and art) of representing molecular structures numerically and simulating their behavior with the equations of quantum and classical physics. Computational chemistry programs allow scientists to generate and present molecular data including geometries, energies, electronic properties, spectroscopic properties and bulk properties.

The starting point for many computer assisted molecular studies is generally a two dimensional drawing of a compound of interest. These diagrams can range from "back-of-the-envelope" sketches to electronically stored connection tables in which one defines the types of atoms in the molecule, their hybridization states and intramolecular bonds.

Connection tables are easily stored and searched electronically. However, they must be transformed into three dimensional representations of chemical structure to study chemical properties. Chemists use the mathematical descriptions of the rules of physical chemistry which are contained in quantum mechanics and molecular mechanics to accomplish this task.

In its purest form, quantum theory uses well known physical constants such as the velocity of light, values for the masses and charges of nuclear particles and differential equations to directly calculate molecular properties and geometries [55]. This formalism is referred to as ab-initio (from first principles) quantum mechanics.

The equation from which molecular properties can be derived is the Schrödinger equation, $H\psi = E\psi$, where $E$ is energy of the system relative to the situation in which all atomic particles are separated to infinite distances, $\psi$ is the wavefunction which defines the Cartesian and spin coordinates of the atomic particles and $H$ is the Hamiltonian operator which includes terms for both, potential and kinetic energy. Unfortunately, the Schrödinger equation can be solved only for very small molecules such as hydrogen and helium. Approximations must be introduced in order to extend the utility of the method to polyatomic systems.

The first approximation attempts to differentiate nuclei and electrons. It assumes that nuclei are much heavier than electrons and move much more slowly so that molecular systems can be viewed as electrons moving in a field of fixed nuclei (the Born-Oppenheimer approximation). Solutions to the Schrödinger equation using this assumption lead to values of effective electronic energy which are dependent on relative nuclear coordinates.

The second approximation allows the wavefunction $\psi$ to be represented as the product of one-electron (or spin) orbitals which are described by a set of basis functions. This formalism is called Linear Combination of Atomic Orbitals theory. Once the molecular orbitals have been derived, the orbital coefficients (which define the energy of the system) are calculated. Hartree-Fock theory is used to accomplish this goal. Hartree-Fock assumes that the energy of a set of molecular orbitals can be derived from the basis set functions describing the orbitals by a set of adjustable coefficients which are used to minimize the energy of the system.

Walter Kohn showed in 1964/65 that the energy of a quantum-mechanical system is uniquely determined by its electron density. This quantity is more easily handled than the complicated wave-function in the Schrödinger equation. Kohn also provided a method which made it possible to set up equations whose solution give the system's electron density and energy. This method, called Density Functional Theory has become widely used in chemistry.
since, because of its simplicity, it can be applied to fairly large molecules [56]. It provides a good accuracy as long as the electron correlation is small.

In general, ab-initio methods are able to reproduce laboratory measurements for properties such as the heat of formation, ionization potential, UV/Visible spectra and molecular geometry. Ab-initio quantum methods compute a number of solutions to a large number of equations [57]. John A. Pople (Noble Prize Chemistry 1998) has developed this computational method. The methods were made available to researchers through e.g. the Gaussian-98 ab-initio program. While recent publications have reported calculations on large molecules [58], the methods are generally limited to compounds containing between ten and twenty atoms due to the amount of computer time required for each calculation and the large amount of disk space needed to store intermediate data files. Physical/theoretical chemists have developed alternative approaches to computing structures and properties by simplifying portions of the calculation to circumvent these limitations. These methods are collectively named semi-empirical quantum methods.

Semi-empirical methods utilize approaches which are similar to ab-initio methods, but several approximations are introduced to simplify the calculations [59]. Rather than performing a full analysis on all electrons within the molecule, some electron interactions are ignored. These methods include the Hückel approach for aromatic compounds (in which the outer electrons in conjugated systems are treated, but the inner (or core) electrons are ignored, fixed geometry).

While semi-empirical methods require less computer resources than ab-initio methods, they are still compute intensive. In general, calculations are routinely performed on compounds which contain up to 100 atoms. The chief drawback of the method is that its application is limited to systems for which appropriate parameters have been developed. The computational requirements for quantum mechanical approaches on molecules consisting of several thousands of atoms (proteins, drug candidates) render these methods unusable for routine analysis. Thus, a further simplification in the way molecular geometries and their associated properties are computed is required. This approach is the Molecular Mechanics or Force Field method.

Rather than utilizing quantum physics, the method relies on the laws of classical Newtonian physics and experimentally derived parameters to calculate geometry as a function of steric energy. In contrast to ab-initio methods, molecular mechanics is used to compute molecular properties which do not depend on electronic effects (geometry, rotational barriers or vibrational spectra). Since the calculations are fast and efficient, molecular mechanics can be used to examine systems containing thousands of atoms. However, unlike ab-initio methods, molecular mechanics relies on experimentally derived parameters so that calculations on new molecular structures may be misleading.

On SubPc, AM1 semi-empirical calculations as well as ab-initio density functional calculations using the B3LYP exchange-correlation functional at the 6-31Gd level [60, 61] were performed, complementing earlier work [62–64]. The charge distribution, electrostatic potential, dipole moment and all molecular orbitals including the HOMO and LUMO are calculated. On Perylen, also ab-initio density functional calculations using the B3LYP exchange-correlation functional at the 6-31Gd level [60, 61] were done to calculate the same properties.
3. Self-Assembly of Phthalocyanines and $C_{60}$ on Ag(111)

In this chapter, a novel route to highly perfect molecular self-assembly through competing interactions of a bi-molecular system on a metallic surface is presented. After some introductory remarks on the topic of self-assembly in section 3.1, both molecules of the binary mixture are presented separately in section 3.2. Then the novel two-component phases are described: Depending on the relative surface coverage of the two species, Subphthalocyanine and $C_{60}$, well-ordered intermixed monolayers consisting of pseudo-linear $C_{60}$-chains (section 3.3) or 2D hexagonal patterns with periodic vacancies (section 3.4) are formed. The structural parameters and schematic binary “phase-diagram” of this system, which are deduced from detailed room-temperature STM studies, are discussed in section 3.5.

3.1. Molecular Self-Assembly and 2D Layering

The term *self-assembly* was coined by G. M. Whitesides [65] and covers a broad range of examples from different fields:

> "Molecular self-assembly is the spontaneous association of molecules under equilibrium conditions into stable, structurally well-defined aggregates joined by non-covalent bonds. Molecular self-assembly is ubiquitous in biological systems and underlies the formation of a wide variety of complex biological structures."

In a recent talk [66], he further characterized this phenomenon: In self-assembling processes, individual objects act quasi-individual and the results contain complex effects which cannot easily be understand (comparable to traffic, stock exchange, thunderstorm). Molecular self-assembly is a powerful method for assembling atomically precise material and devices. Biological organisms are composed of molecular building blocks, such as nucleic acids, proteins, and phospholipids, and are equipped to assemble these components into extremely well-organized structures – namely themselves. Self-organized systems are in the focus of nanotechnology research because of their potential use in the parallel "bottom-up" creation of functional supramolecular structures [67–69]. Self-assembling systems [65] will be required in order to efficiently produce nanoscale molecular electronic devices ("Let Nature do the job") and to create patterned molecular layers for a broad range of applications such as molecular data storage.

Molecular self-assembly has been shown to depend on intermolecular and on molecule-substrate interactions [70–72]. When molecules are deposited onto clean metallic surfaces, they can show a rich variety of phenomena [73]. The present interaction determine the growth, ordering and diffusion of the molecules on the substrate. In the experiments presented here, the molecules are bound to the substrate [74] and do not desorb to the vacuum. But due to the small corrugation of the surface potential, they can to some extent
3.1. Molecular Self-Assembly and 2D Layering

diffuse within a 2D confinement, as was exemplified in [75, 76]. This diffusive mobility ensures that lateral intermolecular interactions can play a significant role in adsorbate ordering. Such noncovalent intermolecular interactions are critical to an understanding of a wide range of disciplines from molecular biology to materials science [77]. In crystal engineering, supramolecular synthesis utilizes hydrogen bonding and other interactions including halogen-halogen, electrostatic interactions, and weak electron donor-acceptor complexation to organize molecules within a crystal [78–80]. When the concepts of 3D supramolecular synthesis are applied to the 2D case of physisorbed molecules on metal surfaces, the adsorbate-substrate interaction plays a role of equal importance with that of the intermolecular interactions [81]. The weak lateral forces exerted by the surface upon physisorbed molecules, and the image charges that occur in metal substrates, allow the weak intermolecular forces to play a significant role in the formation of long-range order in the adsorbed phase. An example of weak intermolecular interactions combined with physisorption comes from the STM study of DNA base molecules on Cu(111) [82]. DNA bases form superstructures on Cu(111) surfaces through intermolecular hydrogen bonding and their planar orientation on the surface. The relative inertness of Cu(111) allows the molecules to diffuse over the surface to spontaneously self-assemble into small clusters and chains. Further, the Langmuir–Blodgett (LB) technique relies on the weak Van der Waals interaction of long hydrophobic molecules on top of a water surface [83–85], whereas the formation of Self-Assembled Monolayers (SAM) is driven by the strong selective bonding of a head group (mostly thiols) to a metal (mostly Au(111)) or a semiconductor surface [86]. The SAMs are a well-established technique to produce 2D molecular crystals with variable properties by specially designing the molecular length and end-groups. Very recently, Yokoyama and co-workers [68] have observed similar small clusters and chains on Au(111) at 63 K. In this study, cyanophenyl-substituted porphyrins were found to form three-molecule clusters in the case of single substitution, four-molecule clusters for asymmetric double substitution, and branching chains for symmetrical double substitution. While these reports on DNA bases and substituted porphyrins are encouraging for 1D aggregation, they do not represent practical examples of 2D supramolecular structures. There are also reports [87] of highly ordered 2D structures at the solution-graphite interface where solvent and solute adopt well-defined bimolecular structures. Strongly ordering chemisorbed systems such as CO/NO + C₆H₆ (benzene) adsorbed on Pt, Rh, Ni and Pd [88–95] or CO and NH₃ (ammonia) on Ru(001) [96] or CO/NO and C₂H₅ (ethylidene) on Rh(111) [97] are not appropriate models for the non-covalent interactions envisaged here because both species undergo significant bonding with the used metal substrates.

Here, a novel route to self-assembly of highly perfect binary molecular monolayers on a metallic surface through competing non-covalent interactions is presented. These novel “Self-Intermixed Monolayer Phases” (SIMP) are different from previously known self-assembled molecular monolayers in that they form intermixed patterns at room temperature on uniform, unreconstructed atomically clean terraces. Unlike in SAM and LB films, the intermixed molecules do not segregate [98–100]. In contrast to former two-component experiments [81], the individual loosely-packed components are easily discriminable because of their specific symmetries. Therefore, experiments e.g. towards single molecular storage application are straightforward.
3. Self-Assembly of Phthalocyanines and C$_{60}$ on Ag(111)

3.2. Single Component Phases: Phthalocyanines and C$_{60}$ on Ag(111)

Figure 3.1.: Structure of Chloro[subphthalocyaninato]boron(III) (SubPc). Colors specify different elements: C, dark gray; Cl, green; H, white; N, blue; B, pink, hidden by the Cl. The height is approximately 0.5 nm, the diameter ca. 1.3 nm.

The growth and self-organization of pure monolayer islands of the individual components sublimed onto Ag(111) have been studied previously by STM [33, 75]. The first component, Chloro-[subphthalocyaninato]-boron(III) (SubPc) [101], is a polar molecule with a characteristic triangular symmetry (figure 3.1). It is the lowest phthalocyanine homologue with unusual electrical and optical properties [102]. These compounds are only known as boron derivatives. In contrast to planar four-fold symmetric phthalocyanines, here, the center metal atom is replaced by a boron-chlorine unit which binds three instead of four isoindol-residues. Therefore the SubPc molecule has a C$_{3v}$ symmetry. Due to the $sp^3$ bonded boron it is non-planar but nevertheless has an aromatic 14-$\pi$-electron system. The synthesis of SubPc was made by Meller and Ossko in 1972 [101] and 2 years later its structure was determined by Kitaibl [103] using X-ray diffraction. In the next 15–20 years great progress for the structural characterization of the SubPc molecules was not achieved even though a notable group of SubPc derivatives was synthesized [104, 105]. One of the most striking features of the X-ray crystal structure is the presence of pairs of SubPc molecules in Van der Waals contact [102]. In a secondary organization, the SubPc pairs form a sheet in which the concave faces point inward and the axial groups point outward toward the next sheet.

In STM images the SubPc molecules appear as three-lobed objects. Individual phenyl-rings can be recognized in STM images with submolecular resolution, see figure 3.2. SubPc molecules adsorbed on Ag(111) self-organize into a honey-comb pattern [33], see figure 3.3. Furthermore, subphthalocyanines can be organized at the supramolecular level in liquid crystals and LB films [102]. They are considered as useful molecules that could act as recording media for DVD-R applications [106] and could be employed for rewriting at short wavelength (purple color: The calculated HOMO-LUMO gap is 2.7 eV and fluorescence measurements in a dimethylformamide solution give a value of 2.2 eV [62]).

The second component is the well-known C$_{60}$ carbon fullerene$^1$ [107] (figure 3.4). Since
the discovery of a method for synthesis of \( C_{60} \) molecules in macroscopic quantities [108],

\[ \text{Also called Buckminsterfullerene, Buckyball, } C_{60} \text{ Fullerene, Carbon cluster (} C_{60} \text{), Fullene-60, Footballene, Footballene (} C_{60} \text{), Icosahedral } C_{60}, \text{ Soccerballene or [5,6]Fullerene } C_{60}. \]

Figure 3.2.: SubPc adsorbed on Ag(111). A single molecule is marked by a circle. The three sub-units of a molecule represent the phenyl-rings [62]. Scan range 21 nm x 9 nm. \( V_{\text{bias}} = 2 \text{ V}, I_t = 200 \text{ pA}. \)

Figure 3.3.: A monolayer of SubPc on Ag(111); The molecules are arranged in a hexagonal honeycomb pattern [75]. Scan range 56 nm x 56 nm. \( V_{\text{bias}} = 1.3 \text{ V}, I_t = 200 \text{ pA}. \)

Figure 3.4.: Structure of \( C_{60} \). The diameter of the carbon cage is approximately 7 Å.
great activity has been directed toward the understanding of the growth and structural properties of C₆₀ layers. Much interest has been focused on the interfacial properties of these layers in contact with various surfaces [109]. In particular, because of charge transfer across C₆₀-metal interfaces, structures composed on these interfaces have attracted considerable attention [110].

The carbon atoms of each C₆₀ molecule have a pentagon-hexagon arrangement. The hexagonal pattern is found in graphite which exhibits sp² bonding characteristics. However, to allow the molecule to curve around to form a sphere, pentagonal bonding is also present. This introduces some sp³ hybridization. One feature of this molecule is that all the valence electrons are involved in C–C bonding with no unpaired electrons left over giving a closed-shell electronic structure. In its bulk form, fullerite, C₆₀ molecules interact with each other via weak van der Waals forces. The energy gap between the lowest unoccupied molecular orbital (LUMO) and the highest occupied molecular orbital (HOMO) is 1.9 eV. C₆₀ therefore is an intrinsic semiconductor. As for other semiconductors, the electrical characteristics may be altered by doping with atoms of different species. Due to the electrophilic (the LUMO is three fold degenerate) nature of C₆₀, much attention has been given to doping fullerene crystals to form n-type semiconductors, metals or superconductors [111].

The threefold degenerated LUMO is shown in figure 4.1 in section 4.1. In the room-temperature STM images, however, the fullerenes appear as spheres and no details from the intramolecular structure are visible. This is due to the vibrational modes of the molecules at room-temperature, which are fast compared to the scanning speed [112] (see also section 4.4.1).

The ultra-pure C₆₀ with a density of 1.65 g/cm³ [113] were sublimed at approximately 650 K. The growth and self-organization of C₆₀ on Ag(111) have been studied previously by STM [112, 114, 115]. C₆₀ sublimed onto Ag(111) form close-packed islands with p6mm symmetry which are fluid to be rotated ±30° against the Ag lattice [116], see left side of figure 3.6 and figure 3.7. It is a commensurate $(2\sqrt{3} \times 2\sqrt{3})R \pm 30°$ structure and is the most stable phase of C₆₀ adsorbed on Ag(111) [115]. The lattice mismatch of these C₆₀(111) islands is -0.2% at room-temperature and all molecules are adsorbed at hollow sites [116].

### 3.3. Mixture of C₆₀ and SubPc: 3:2-Stripe Phase

C₆₀ and SubPc co-deposited onto the atomically clean Ag(111) substrate are observed to self-organize into periodic 2D patterns at room-temperature. In typical STM images (figure 3.5), individual molecules within ordered patterns can be clearly identified by their characteristic triangular or round shapes, which match those determined in previous STM studies [62, 75, 112, 116]. The highly anisotropic arrangement of the molecules on the right hand side of figure 3.6 consists of zigzag rows of SubPc molecules (green) separated by quasi-linear chains of C₆₀ molecules (yellow). Within these rows, the C₆₀ molecules are grouped into aligned triplets, where the center molecule is sandwiched between two SubPc molecules. Remarkably, in this self-assembled pattern, a significant fraction of the surface between the molecules remains uncovered (appears dark in figure 3.5 and figure 3.6). In this phase, the molecules intermix at a fixed ratio of SubPc:C₆₀ = 2:3. Alternating C₆₀ chains and SubPc
zigzag rows form a 2D periodic “stripe pattern” which belongs to the planar space group \( p2 \). The stripes are parallel to the close-packed \( C_{60} \) rows of an island of the hexagonal phase of \( C_{60} \) on Ag(111) [116], which appears on the left side of figure 3.6. These close-packed rows of the pure \( C_{60} \) layer are parallel to the [TT2] direction of the substrate and can thus be used to identify the orientation of the stripe pattern. Such coexisting \( C_{60} \) islands are formed next to the mixed phase when the \( C_{60} \) content of the deposited film exceeds a critical ratio. The \( C_{60} \) within the hexagonal phase occupy highly-symmetric threefold-hollow sites [116] of the underlying Ag(111) substrate, and thus the molecules in the mixed layer have a definite registry with respect to the substrate lattice i.e. they adapt to the atomic corrugation of the surface potential. Nevertheless, the observed self-intermixing assembly requires a sufficient mobility of each component at 293 K. Previous authors have reported significant room-temperature diffusion of \( C_{60} \) on some metals. On Ag(111) this diffusion is manifested via the growth of islands from step edges [112]. For SubPc, the room-temperature coexistence of ordered islands with patches of mobile molecules in a 2D “lattice gas”-phase has been reported recently [75]; a similar 2D gas-phase is observed (see figure 3.10 in section 3.4.2). The mobility of the molecules is further supported by time lapsed-images in the vicinity of vacancy defects, as exemplified by the split appearance of the \( C_{60} \) molecule marked red (at the top of figure 3.6). The STM data confirm the perfection of the atomic and molecular arrangement of the substrate and of the grown binary layers: virtually no substitutional defects or additional adsorbates appear on the extended ordered terraces. This is a prerequisite to recognize details of the complex self-organization.

Clearly, the intermolecular interactions between SubPc and \( C_{60} \) are producing an entirely new 2D crystalline structure that is energetically favored over the patterns resulting from either parent component.

![Figure 3.5: Self-intermixed monolayer on Ag(111) and its molecular components. Artificially colored Scanning Tunneling Microscopy image of the basic unit of the molecular stripe pattern with superimposed schematic contours (SubPc green, \( C_{60} \) yellow). Individual components can be identified by size and shape. Scan range 4.3 nm x 3.2 nm. \( V_{bias} = 1.3 \) V, \( I_t = 20 \) pA.](image-url)
3. **Self-Assembly of Phthalocyanines and C_{60} on Ag(111)**

3.3.1. **Adsorption Geometry of the Stripe Pattern**

A model of the registry is given in figure 3.7. The model is based on a thorough analysis of STM images with different phases. The known hexagonal close-packet phase of pure C_{60} molecules and the honeycomb pattern of pure SubPc molecules are used as references. Images with two different phases were surveyed for angles and distances. The stripes grow in

![Figure 3.6: Self-intermixed monolayer on Ag(111): Artificially colored STM image of a monolayer of co-adsorbed SubPc and C_{60} on Ag(111) (imaged area, 17nm x 25nm, V_{bias} = 1.3 V, I_{t} = 20 pA). The self-intermixed stripe phase is shown on the right with alternating C_{60} chains (yellow) and SubPc rows (green). The bare substrate areas appear dark. The distance between adjacent parallel C_{60} rows is 3.3 nm. Up to 38 parallel stripes with a length of 100 nm have been observed. The monoclinic unit cell of p2 symmetry contains three C_{60} and two SubPc molecules. One C_{60} molecule, colored in red near the top edge of the C_{60} island, jumped to an adjacent site between two horizontal scans. A pure C_{60} island coexists on the left hand side.](image)
3.3 Mixture of C\textsubscript{60} and SubPc: 3:2-Stripe Phase

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Experiment</th>
<th>Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>molecular spacing in pure C\textsubscript{60} HBC islands</td>
<td>1.0±0.1 nm</td>
<td>1 nm</td>
</tr>
<tr>
<td>inter-row spacing \perp stripes</td>
<td>3.3±0.3 nm</td>
<td>3.32 nm</td>
</tr>
<tr>
<td>SubPc-SubPc distance \parallel stripes</td>
<td>3.0±0.3 nm</td>
<td>3.0 nm</td>
</tr>
<tr>
<td>centerhole-centerhole distance in star pattern</td>
<td>5.7±0.5 nm</td>
<td>5.2 nm</td>
</tr>
<tr>
<td>(\angle) (C\textsubscript{60}-triplet, stripe-axis)</td>
<td>±8±1°</td>
<td>±8.2°</td>
</tr>
<tr>
<td>(\angle) between holes in star pattern (section 3.4)</td>
<td>60.5±1°</td>
<td>60°</td>
</tr>
<tr>
<td>(\angle) (holes in star pattern and stripes)</td>
<td>90±1°</td>
<td>90°</td>
</tr>
<tr>
<td>(\angle) (honey-comb axis, stripe axis)</td>
<td>84±1°</td>
<td>84.5°</td>
</tr>
<tr>
<td>(\angle) (C\textsubscript{60} HBC, C\textsubscript{60} HBC-center axis)</td>
<td>9±1°</td>
<td>10°</td>
</tr>
<tr>
<td>(\angle) (stripe domains)</td>
<td>60±1°</td>
<td>60°</td>
</tr>
<tr>
<td>(\angle) (honey-comb holes, star holes)</td>
<td>68±1°</td>
<td>65°</td>
</tr>
<tr>
<td>(\angle) (honey-comb holes, honey-comb holes)</td>
<td>60±1°</td>
<td>60°</td>
</tr>
<tr>
<td>(\angle) (honey-comb chiral domains) [33]</td>
<td>9±1°</td>
<td>9.42°</td>
</tr>
</tbody>
</table>

Table 3.1.: Comparison between the experimental data and the theoretical analysis of the registry model.

lots of domains that can differ in the angle and the phase, see figure 3.8. This gives further
insight to the substrate-overlayer relationship. The comparison of the measurements with
the model (figure 3.7 and 3.13) is given in the table 3.1. The results correlate perfectly,
although it is not possible to image the substrate simultaneous with the adsorbates and an
error has to be taken into account.\(^2\) The model perfectly fits in geometry (distances and
angles) within all phases (C\textsubscript{60} HBC, SubPc\textsubscript{honey-comb}, stripe-phase and star-phase (presented in
section 3.4)). Here, all SubPc molecules are located at identical adsorption sites (center
on-top) and have the same appearance in STM images.

Remarkably, in this model the three C\textsubscript{60} molecules within a triplet have not the same
adsorption site with respect to the underlying substrate: The corner fullerenes still are
adsorbed in hollow positions like in the pure island, whereas the center fullerene of a triplet
now is located on-top. This different adsorption-geometry may lead to different electronic
bonding between adsorbate and substrate and therefore to a different appearance in STM.
Quite often in STM studies of molecular films, it is possible to recognize different appearances
of molecules, i.e. to recognize molecules with a variation in the apparent height [114].
Since STM maps the local density of states (see equation (2.4)), it is controversial whether
the brightness difference is related either to a real difference in height (i.e. molecules are
pushed out) or to an electronic effect (i.e. a spatially non-homogeneous charge transfer).
Here\(^3\) on Ag(111), bright molecules can also have different bonds in equivalent adsorption
sites [112, 117]. In the case of Ag(100), see section 4.4.1, the different structures are not
due to geometric effects, but to non-homogeneous charge distribution and different chemical
bonding with the Ag substrate [117]. On Al(111), the observed non-equivalent adsorption

\(^2\) The STM was calibrated with a Si(111)\textsubscript{7x7} sample, but at least 5\% error in the lateral accuracy have to
be allowed.

\(^3\) and also in the case of Au(111)
3. *Self-Assembly of Phthalocyanines and C∞ on Ag(111)*

sites are due to the surface reconstruction [118], whereas on Au(110), the C∞ adsorption induces a cooperative self-assembly of Au atoms and C∞ [109].

![Diagram](image)

Figure 3.7.: Proposed model for the registry of the *stripe* phase on Ag(111). The positions and the orientations of the C∞ and SubPc molecules are depicted on the atomic silver lattice.

![Images](image)

Figure 3.8.: a) Rotational and b) translational domain shifts of the *stripe* phase in accordance with the hexagonal (111) substrate (60° in a), resp. 1.7 nm in b) which means a shift by 6 Ag sites). Scan range a) 56 nm x 48 nm, b) 23 nm x 21 nm.
3.3.2. Chiral Supra-Molecular Structures of Achiral Molecules

*Chiral* is the Greek expression for *hands* and was used the first time by Louis Pasteur 1847 to describe macroscopic crystals that are its mirror images.\(^4\) Chirality is a geometrical property that plays an important role in physics, biology and chemistry. Two chiral forms of the same molecule can have extremely different properties: The taste of an orange and a lemon arise from the same molecule but with different chirality. Likewise the taste of caraway and spearmint. Two different enantiomers (chiral forms) have identical physical properties (such as boiling and melting point, density, spectroscopic and chromatographic features) and are spectroscopically not distinguishable, but sometimes well with STM. Recent results give more insight into the mechanism of chiral recognition [119–121].

Two mirror-symmetric species of the *stripe* pattern exist: The C\(_{60}\)-triplets are turned out of the overall row axis by 8° either clockwise or counterclockwise. Therefore I can clearly distinguish between \(\lambda\) (left) and \(\rho\) (right) domains which differ by a mirror operation. Interestingly, within one domain I can only find one chiral form, that means the chirality is strictly correlated. The reason might be found in the growth-process of an island: The first grown stripe probably defines the chiral form, then the following accretion has to suit it, see also section 3.5. Here, homochiral chains are formed by achiral molecules. These superstructures show adsorption-induced chiral motifs, a phenomenon that is called “organizational chirality” [122].

![Mesoscopic chirality segregation into \(\lambda\) and \(\rho\) domains of the *stripe* pattern](image1)

Figure 3.9.: Mesoscopic chirality segregation into \(\lambda\) and \(\rho\) domains of the *stripe* pattern: Enantiomorph supramolecular structures are formed by a chiral packing of achiral molecules. The scan ranges are approximately 10 nm x 10 nm. In both scans \(V_{bias} \approx 1.3\,\text{V}, \, I_t \approx 20\,\text{pA}\).

3.4. Mixture of C\(_{60}\) and SubPc: Binary 1:1-Star Phase

A higher fraction of SubPc in the co-deposition process leads to a distinctly different pattern. Figure 3.10 shows the previous *stripe* phase (I) co-existing with a new hexagonal “*star phase*”

\(^4\)As a graduate student, he was studying two acid-crystals in the dregs of wine with identical chemical composition but different optical properties.
3. **Self-Assembly of Phthalocyanines and C₆₀ on Ag(111)**

(II). The triangular sub-unit (figures 3.11 and 3.12) of the hexagonal pattern consists of a C₆₀ trimer which is surrounded by three SubPc molecules pointing towards center holes. The mixing ratio of SubPc:C₆₀ in this pattern is 1:1. Between these two patterns (figure 3.10, I and II), irregular streaks of comparable height are clearly visible (figure 3.10, III). These streaks can be identified as mobile molecules in a 2D gas phase which diffuse faster than the imaging process, for more details see section 3.4.2, using the analysis recently applied to pure SubPc monolayers [75].

The self-assembled mixed monolayers observed here (*star* and *stripe* patterns) are distinctly different from previously known 2D molecular self-assembled or otherwise ordered binary layers. The ratio and relative positions of the components and the periodicity in the monolayer are maintained within large domains that are stable at room-temperature. Moreover, the spacing and orientation of the SubPc molecules suggests a special ordering mechanism. Therefore, I introduce the term “Self-Intermixed Monolayer Phases” (SIMP) for the novel co-adsorbed structures observed here.

![Figure 3.10: Self-Intermixed Monolayer Phases (SIMP): coexisting star and stripe patterns. This STM image shows two different SIMPs of SubPc and C₆₀ on Ag(111) (imaged area, 34 nm x 34 nm, Vbias = 1.9 V, It = 20 pA). On the right hand side, the 3:2-stripe SIMP (I) and on the left hand side, the hexagonal 1:1-star SIMP (II) is observed. Point defects are marked by arrows (top right: a vacancy defect is interrupting a C₆₀ row; left: a center hole is filled with a molecule). Apparently random tip excursions of single molecular height (III) are visible between the two ordered regions and are identified as mobile molecules in a 2D gas phase [75].](image-url)
3.4. Mixture of C_{60} and SubPc: Binary 1:1-Star Phase

Figure 3.11.: STM image of a hexagonal cell in the star SIMP, with superimposed schematic shapes. A C_{60} triplet is surrounded by three SubPc molecules. Through a vacancy in the center, the level of the Ag substrate is imaged. Scan range 11.5 nm x 11.5 nm, V_{bias} = 1.9 V, I_t = 20 pA.

Figure 3.12.: With a good STM tip, the intramolecular structure of the SubPc molecules can be resolved and their orientation within the pattern is unravelled. Scan range 17 nm x 12 nm, V_{bias} = 1.9 V, I_t = 20 pA.
3. Self-Assembly of Phthalocyanines and $C_\infty$ on Ag(111)

3.4.1. Adsorption Geometry of the Star Pattern

Based on the analysis presented in section 3.3.1 and table 3.1, the following registry model for the 1:1-star phase is proposed, see figure 3.13. It is a commensurable $(18 \times 18)R0^\circ - 2(3\text{SubPc} + 3C_\infty)$ structure. Note that the allocation to the exact adsorption sites and the molecular orientations remain rather speculative.

![Proposed model for the registry of the star phase on Ag(111). The center hole is large enough to offer space for an additional molecule (see left arrow in figure 3.10).](image)

3.4.2. 2D Molecular Gas-Phase

Between two SIMPs, often a 2D molecular gas-phase is found. In STM images, this can be recognized as streaks in the fast horizontal scan-direction, see top of figure 3.14. The line-scan, see bottom of figure 3.14, makes clear that the same single-molecular events can be found in the condensed phases and in the gas phase: the objects consistently have an apparent height of 0.5-0.6 nm and a width of 1-2 nm and can be identified as molecules. For a more detailed analysis of this so-called lattice-gas see [75]. The composition of the gas is not clear a priori, but could be explored by a variable temperature STM (e.g. freezing the movement or growing of a certain domain) or by a careful series of deposition experiments with one dominating component.
3.4. Mixture of $C_{60}$ and SubPc: Binary 1:1-Star Phase

Figure 3.14.: Molecular gas-phase: between the two well-ordered SIMPs, random tip excursions can be seen due to mobile molecules (top part). Scan range 27 nm x 15 nm, $V_{bias} = 1.9$ V, $I_t = 20$ pA (section from fig. 3.10). The line-scan ("height-profile", lower part) makes clear that the same objects can be found in the condensed phases and in the gas phase. The motion of the molecules in the gas-phase during the scanning-process, however, inhibits any correlation of the successive scan-lines.
3. **Self-Assembly of Phthalocyanines and C\(_\infty\) on Ag(111)**

<table>
<thead>
<tr>
<th>Phase</th>
<th>Super-structure</th>
<th>Matrix</th>
<th>Unit-cell [Å(^2)]</th>
<th>Ref.</th>
</tr>
</thead>
</table>
| Ag(111)| \(1 \times 1\) \(-(1\text{Ag})\)                  | \[
\begin{pmatrix}
1 & 0 \\
0 & 1
\end{pmatrix}
\] | 7.233                | Fig. 3.7 |
| C\(_\infty\) | \((2\sqrt{3} \times 2\sqrt{3})R\pm30^\circ -(1C_{\infty})\) | \[
\begin{pmatrix}
2 & 2 \\
-2 & 4
\end{pmatrix}
\] | 86.8                | Fig. 3.7 |
| SubPc  | \((\sqrt{111} \times \sqrt{111})R\mp4.71^\circ -(2\text{SubPc})\) | \[
\begin{pmatrix}
10 & 1 \\
-1 & 11
\end{pmatrix}
\] | 810.0                | [33] |
| star   | \((18 \times 18)R0^\circ -2(3\text{SubPc}+3C_{\infty})\) | \[
\begin{pmatrix}
18 & 0 \\
0 & 18
\end{pmatrix}
\] | 2343.5              | Fig. 3.13 |
| stripe | \((\sqrt{108} \times \sqrt{151})R30^\circ -(2\text{SubPc}+3C_{\infty})\) | \[
\begin{pmatrix}
6 & 6 \\
-9 & 14
\end{pmatrix}
\] | 998.1                | Fig. 3.7 |

Table 3.2.: Crystallographic properties of various phases.

### 3.5. Schematic Room-Temperature Binary Phase Diagram

The multi-phase behavior of this binary system on Ag(111) has been studied as a function of composition in a series of deposition experiments. The results are summarized in the schematic “phase-diagram” (figure 3.15). From the top to the bottom, self-intermixed monolayers are shown with an decreasing SubPc:C\(_\infty\) mixing ratio from 1:0 to 0:1. In the first column, pure mono-molecular phases of SubPc and C\(_\infty\) are depicted at the top and the bottom. Homogeneous *star* and *stripe* SIMPs correspond to a mixing ratio of SubPc:C\(_\infty\) = 1:1 and 2:3, respectively, and are shown in the middle of the left column. Between these mixing ratios, coexistence of two adjacent phases is observed, as shown in the second column. Using this schematic 2D “phase-diagram”, a mixing ratio for deposition of the molecular components can be chosen to control the formation of selected 2D patterns [123].

Coadsorbate-induced ordering is a cooperative interaction between two different adsorbates which yields different ordered structures than either adsorbate forms by itself. The crystallographic properties of all superstructures from figure 3.15 are listed in table 3.2.

**Intermixing Activity**

The exact transition from one pattern to another upon additional deposition of molecules remains unsearchable, mainly because of technical reasons. The molecules are deposited in the OMBE chamber, whereas the observations are made in the STM-chamber, see figure 2.1. It is impossible to find the same surface location within nanometer precision and to control the mutation of the pattern after a sample-transfer of several meters. But there is a solution to get some aspects of the transition from one pattern to another sometimes. By preparing a sample with pure SubPc honey-comb pattern and then depositing a small amount of C\(_\infty\), I can get some “snapshots” of the transition process.\(^5\) The C\(_\infty\) molecules inter-diffuse

\(^5\)Because of slightly inhomogeneous deposition, there exist areas with a tiny amount of C\(_\infty\). Other areas with higher density do not contribute to the process, presumably because the step-potential prevents inter-step diffusion and the molecules cannot cross steps of the substrate [75, 124].
and form “channels”, where two C\textsubscript{60} molecules replace one SubPc molecule. Note that
the SubPc domain of both sides of the “channel” are phase-shifted and the C\textsubscript{60} molecules
start to form a stripe similar to the known geometry (here, however, two C\textsubscript{60} molecules are
sandwiched between two staggered SubPc molecules). The chirality of this new single stripe

Figure 3.15.: Schematic room-temperature “phase-diagram” of SubPc and C\textsubscript{60} with corre-
spanding STM images of self- intermixed monolayers (imaged area, 13 nm x
13 nm, 1.0 V < V\textsubscript{bias} < 1.9 V, 20 pA < I\textsubscript{i} < 100 pA, C\textsubscript{60} blue, SubPc green).
Left column: pure SIMPs displayed against the ratio SubPc:C\textsubscript{60}: 1:0, honey-
comb phase of pure SubPc; 1:1, self-intermixed monolayer star phase; 2:3, self-
intermixed monolayer stripe phase; 0:1, hexagonal phase of pure C\textsubscript{60}. Right
column: Coexisting phases in equilibrium with molecular 2D gas for ratios
SubPc:C\textsubscript{60} which are between the above-mentioned values.
3. Self-Assembly of Phthalocyanines and C₆₀ on Ag(111)

(see section 3.3.2) is most probably given by the chirality of the preexisting honeycomb pattern.⁶

Figure 3.16.: “Snapshot” of the transition from pure SubPc honeycomb pattern to an intermixed phase. The C₆₀ molecules inter-diffuse and form “channels”, where two C₆₀ molecules replace one SubPc. Scan range 19 nm x 17 nm. Vₘₐₛ = 2 V, Iₖ = 15 pA. This STM image is slightly filtered with a median filter.

⁶The honeycomb pattern can exist in two chiral domains [33].
4. Intermixing Mechanism

In order to understand this remarkable self-assembly behavior, possible molecule-molecule and molecule-substrate interactions have to be considered. These interactions and the molecular mobility determine nucleation, growth and ordering of molecular adsorbates. Upon deposition, these processes drive the system towards a minimum of the overall free energy. I think that the mechanism is based on a delicate balance of electrostatic and Van der Waals interactions. The most important underlying interactions and the relevant properties of the molecules [125] are discussed qualitatively in section 4.1. Further observations that explain the molecular interactions are given in section 4.2. A general consideration of thermodynamics of mixtures is treated in section 4.3. This chapter then is closed with a counter-example of the SIMPs: A bi-molecular system that does not intermix but that segregates into separate single-component phases is introduced in section 4.4.

4.1. A Delicate Balance of Electrostatic and Van der Waals Interactions

Neither SubPc nor $C_{60}$ form strong covalent bonds with the Ag(111) substrate, whereas it has for instance been observed for either molecule on Si(111) [62, 116]. Nevertheless, both molecules bind rather strongly to Ag(111): the $C_{60}$ monolayer desorbs only above 670 K [126, 127], whereas I have indications that the SubPc molecules decompose above 470 K. At room temperature, however, isolated molecules of both types diffuse and cannot be imaged by STM. This indicates that the corrugation of the molecule-substrate interactions is weak. In the following two paragraphs, the electronic properties and bonding of each component, $C_{60}$ and SubPc, are discussed first, and then the interactions governing the observed ordering in the SIMPs.

The first component, $C_{60}$, has a high polarizability, because its frontier $\pi$-orbitals extend around the carbon cage, see e.g. figure 4.1. Significant attractive Van der Waals interactions originate from mutually induced polarization fluctuation on the molecules [126, 128] and on the metal surface. An additional weak chemical bond to the Ag(111) substrate with ionic character arises from the charge transfer of approximately $\Delta q$ of 0.8 electron from the metallic substrate to each $C_{60}$ in the monolayer, estimated from photoemission spectra, see figure 4.13, and vibrational energy shifts [129]. This charge transfer implies an induced dipole moment pointing towards the substrate which in principle leads to a long-ranged, lateral repulsion between these parallel dipoles. The packing and aggregation of the $C_{60}$ ad-layer is determined by the balance between the Van der Waals attraction, the dipolar repulsion and the "steric" (Pauli) repulsion between contacting molecules and the substrate.\textsuperscript{1} The close
4. Intermixing Mechanism

match between intermolecular distances in the monolayer and in the fcc crystal of bulk C$_{60}$ (with space group $Fm\bar{3}m$) \cite{126} suggests that the long-range electrostatic repulsion between the adsorption-induced dipoles is of minor influence.

![Image](image)

Figure 4.1.: LUMO of C$_{60}$ [132].

For the second component, SubPc, AM1 semi-empirical calculations as well as ab-initio density functional calculations were performed using the B3LYP exchange-correlation functional at the 6-31Gd level \cite{61} complementing earlier work \cite{62, 63}. Both approaches indicate an excess of electronic charge on the electronegative atoms which surround the electron-deficient boron. This charge is compensated by an electron deficit which is mainly localized on the six central carbon atoms (figure 4.2).

This results in a calculated permanent axial dipole moment of 0.95 eÅ pointing away from the Cl, the magnitude of which compares well to a measured value of 1.1 eÅ and to the value of 1.2 eÅ predicted by AM1 calculations \cite{63}, see figure 4.3. Each SubPc molecule is bound to the substrate by a Cl–Ag interaction and by electrostatic attraction of its polar charge distribution with the induced counter-polarization of the metallic substrate. On the basis of XPS and UPS data \cite{33, 124} it is concluded that the SubPc is adsorbed intact with the Cl pointing towards the substrate. This is further confirmed by the observed destabilization of ordered 2D islands in STM images at reverse bias voltage $V_{bias}$ where the dipole moment is opposing the electric field in the tunneling junction. The permanent dipole moment of SubPc together with its parallel image in the metallic substrate gives rise to a noticeable repulsive lateral interaction between the SubPc molecules \cite{75}. On the other hand, SubPc is less polarizable than C$_{60}$, owing to its less extended frontier π-orbital, which are mainly localized around the central C-N-C-ring \cite{62}. Therefore, the Van der Waals attraction between adsorbed SubPc molecules is weaker. This picture is supported by the tendency of SubPc to form ordered structures with voids and with intermolecular separations \cite{75} of 1.79 nm which exceed the lateral extent of the molecule. The relative angular orientation of neighboring adsorbed SubPc molecules is determined by the details of the charge distribution: the dominant isotropic part of the electrostatic repulsion is reduced if the overall positive corner of one molecule points towards the face of its neighbor which exposes a negative charge around an N atom (see figure 4.2). Thus, for SubPc layers, unlike for C$_{60}$, a balance

\footnote{Because of a small charge deficiency between the single bonds and a charge excess on the double bonds, the intermolecular potential in addition contains a small bond-to-bond Coulomb energy dependent on the molecule orientation \cite{130}. This results in a variation of the domain energy by about 0.1 eV per molecule \cite{131}.}
between stronger electrostatic repulsion and weaker intermolecular attractions determines layer packing and ordering.

Figure 4.2.: Relaxed geometry and effective atomic charges (Mulliken population analysis) from a density functional computation on a single free SubPc molecule [125]. For the identification of atomic species, the color-code of figure 3.1 is used. The distribution of delocalized electronic states and partial charges determines the intermolecular and molecule-substrate interactions which presumably leads to the observed ordering as discussed in the text.

Figure 4.3.: Electrostatic potential of the free SubPc molecule mapped onto the electron density surface [60]. Excess of electrons at the Cl in the center are colored red (negative electrostatic potential), more positively charged portion at the peripheral phenyl-rings blue (positive potential). This electron distribution is responsible for the axial dipole moment [60] and for the molecular interactions.
4. Intermixing Mechanism

For the mixed phases, I note that the packing density of SubPc molecules in the stripe and star phases is about 7% and 20% higher than in the honeycomb phase of pure SubPc even if the same area is assigned to each C_60 as in the pure hexagonal phase. In analogy with ordered structures of small co-adsorbed molecules [89–91], this suggests a higher binding energy. In the case of benzene co-adsorbed with CO or NO, where also different phases were observed [89–92], this has been attributed to increased attraction between antiparallel induced dipole moments which originate from the adsorption chemistry of the two species. In our case, stronger binding also partly arises from the interactions of the SubPc permanent dipole moment with the antiparallel induced dipole moment of the neighboring C_60 molecules. In addition, Van der Waals and anisotropic Coulomb interactions contribute. A summary of all the contribution is given in figure 4.4. But at the moment, it is not possible to deconvolute the individual contributions unambiguously. The proposed explanations are not final and a more detailed theoretical analysis is certainly desirable (e.g. numerical calculations including the substrate and/or several adsorbates). In particular the adsorption characteristics of the SubPc molecules, their mobility and appearance in STM images, as well as the detailed geometries, registries and relative energies of the observed ordered phases must still be quantitatively understood.

4.2. Discussion of Other Intermixing Models

There are many factors to consider when discussing this complex system. Unfortunately, even in works on simpler atomic systems [133], no conclusive rules could be derived about long-range electrostatic or local direct-interactions. The net effect in the redistribution of charge upon adsorption (and the modification of bonds) is very complicated and has to be recalculated in every single case. Further models can be proposed to explain the intermixing phenomenon as explained in section 4.1:

- Long-range interactions mediated by substrate surface states: On Ag(111), the corresponding wavelength of Friedel oscillations outranges the characteristic intermolecular spacing and strong surface-state – adsorbate coupling is ruled out, see section 4.2.1.

- Intrinsic dipole-dipole interaction: They are too weak, see section 4.2.2.

- Charge transfer between the molecules: Possible effect, could explain the intermolecular attraction, discussion in section 4.2.3.

- New chemical bonds between the components: UPS and XPS measurements (see section 4.2.4) show no chemical interaction between the components, but solely an inter-diffusion. The interface between the two components proves to be stable at room-temperature (STM).

---

2This result easily from following consideration (for numbers see table 3.2): One C_60 molecule in the single component phase needs 87 Å^2, one SubPc molecule in the pure honeycomb pattern needs 403 Å^2. In separate phases, these two molecules thus need 492 Å^2. In the star phase, however, 1 SubPc + 1 C_60 only need 391 Å^2. In the stripe phase, 2 SubPc + 3 C_60 need 908 Å^2, whereas the same molecules in single-component phases need 1071 Å^2. This defines the higher packing density of 7% and 20%.
4.2. Discussion of Other Intermixing Models

4.2.1. Long-Range Interactions Mediated by Substrate Surface States

Theoretical calculations on adsorbed atoms and molecules have shown that the interactions between adsorbates are primarily indirect (i.e., via the substrate wavefunctions) when the species are separated so that the overlap of adsorbate orbitals is negligible [95]. The substrate wavefunctions oscillate in sign as a function of the separation distance, and decrease (with \( r^{-2} \)) in magnitude with increasing separation distance [134]. When these interactions are found to be attractive at certain separation distances, ordered domains can form [135]. The possibility of such long-range interactions mediated by substrate surface states [136], however, can in my opinion be excluded because on Ag(111) the corresponding energy band edge is only 0.05 eV below the Fermi level [137], and the corresponding wavelength of Friedel

\[
\text{Attractive and repulsive contributions}
\]

Figure 4.4.: Summary of the forces in SIMPs: molecule-substrate and molecule-molecule interactions for both components separately and the mixed phase are shown. Attractive contributions are in blue, repulsive contributions in red.
oscillations ($\lambda = 3.8$ nm) considerably exceeds any of the observed nearest-neighbor spacing, see figure 3.7. Other anisotropic adsorbate systems show such a coupling [137, 138].

4.2.2. Intrinsic Dipole-Dipole Interaction

A calculation of the (permanent) dipole – (substrate induced) dipole interaction between SubPc and C$_{60}$ gives an attractive value of $+8.3$ meV for the antiparallel dipole moments [139]. At room-temperature, this is not enough to explain the SubPc–C$_{60}$ attraction. From the change in work-function, the dipolar interaction energy for example of the pairing CO+benzene on Rh(111) was estimated to be $+90$ meV [89]. In their study, this was enough to stabilize these chemisorbed molecules in an intermixed phase.

4.2.3. Intermolecular Charge Transfer: Apparent Height Analysis

Previous observed intermixed systems [81, 88–95, 97] are based on a typical electrophilic and electrophobic molecular pairing with a charge transfer between the molecules. The leading forces for stabilization in these donor–acceptor systems are then believed to be dipole–dipole interactions (induced antiparallel dipole moments, see previous section).

The analysis of the apparent height of the C$_{60}$ and SubPc molecules gives some indications of a charge transfer from the 14-$\pi$-electron system of the SubPc to the $\pi^*$-orbital of the C$_{60}$, which is well known as a strong electron acceptor. This supports the observed moderation and stability of the mixed monolayers at the presence of two adsorbates. However, other observations (section 4.2.4) do not support this charge transfer mechanism.

Since STM images represent a convolution of the electronic (LDOS) and topographic structure, a change in the apparent height can thus be due to a change in the physical height of the molecule, a change in the conductance of the molecule or both, see equation (2.4). If the geometry of the system is known, information of the electronic situation can be deduced and a charge transfer might become detectable.

To investigate this proposed intermixing mechanism, height-profiles of STM images have been accurately analyzed. Images with coexisting phases and identical scanning parameters have been chosen to compare the apparent molecular heights. By investigating all phases, the apparent height of the C$_{60}$ in all configurations was found, see figure 4.5.

An electron transfer from the SubPc to the C$_{60}$ should lead to a reduction of the C$_{60}$ height in STM images of mixed phases since electrons would be transferred from the SubPc HOMO to the threefold degenerated C$_{60}$ LUMO orbitals. The additional filling of the LUMO would then reduce the tunneling probability (compared to the already 0.8 substrate-electron filled C$_{60}$ LUMO in the pure fullerene layer). In STM images (with negative bias, tunneling into the LUMO, see figure 2.4), the C$_{60}$ molecules would then be observed with a smaller signal. Moreover, additional SubPc neighbors should enhance the effect.

\[^3\text{Interaction energy } w(r) = \frac{q_1q_2e^{-r}}{4\pi\varepsilon_0}. \text{ Supposed a permanent dipole moment for SubPc of } 0.95 \text{ eÅ and a } C_{60} \text{ charge transfer of } 0.75 \text{ e from the Ag(111)-substrate over a distance of } 0.2 \text{ nm and an inter-molecular separation-distance of } 4 \text{ lattice constants (see figure 3.7). This calculation does not include a charge transfer between the molecules which would intensify the dipolar interactions.}\]
4.2. Discussion of Other Intermixing Models

Figure 4.5.: Apparent height of C\textsubscript{60} molecules in different phases, imaged at 1.9 V bias voltage \(V_{bias}\). The variations are assumed to arise from different local environments modified by the SubPc molecules.

Indeed, the measurements show such a decrease of the C\textsubscript{60}-height as a function of the number of “donating” SubPc neighbors. The pure C\textsubscript{60} layer without any SubPc contact and no potential charge transfer shows the highest apparent height. Also the behavior of the C\textsubscript{60}\textsuperscript{stripe-center}, C\textsubscript{60}\textsuperscript{stripe-corner} and the C\textsubscript{60}\textsuperscript{star} molecules support the charge transfer theory from the SubPc to the C\textsubscript{60}. The C\textsubscript{60}\textsuperscript{stripe-corner} molecule is the largest of these three cases, which in this model is explained by the absence of any SubPc contact. The higher coordinated C\textsubscript{60}\textsuperscript{stripe-center} and C\textsubscript{60}\textsuperscript{star} molecules are smaller than the low coordinated C\textsubscript{60}\textsuperscript{stripe-corner} molecule which can be explained by the direct contact to SubPc neighbors. The small height-difference between the C\textsubscript{60}\textsuperscript{stripe-center} and C\textsubscript{60}\textsuperscript{star} molecules can be interpreted in the same arguments of charge transfer: Since the C\textsubscript{60}:SubPc mixing ratio is 3:2 in the stripe pattern and 1:1 in the star pattern, the C\textsubscript{60} in the latter pattern appears smaller because it is supplied with the charge from one SubPc, in the stripe pattern only from \(\frac{2}{3}\) of a SubPc. In conclusion, this analysis supports the SubPc→C\textsubscript{60} charge transfer theory.

There could be additional effects to explain the differences of the apparent heights, like the adsorption on different sites on the Ag(111) substrate \cite{114} (see figure 3.13 and 3.7), random blinking\textsuperscript{4} \cite{112} and substrate- or coadsorbate-induced conformational changes\textsuperscript{5} \cite{140}. Up to this point, I discussed the apparent height of the fullerenes at a certain bias voltage and I showed that this effect could be linked to a possible charge transfer between the molecules. The situation gets more complex when looking at different bias voltages.

**Bias-dependent Images** During imaging with the STM, the bias voltage \(V_{bias}\) can be varied. Only one polarity was applied (sample positive), because the SubPc islands are known to be stable only in this field orientation due to the molecules’ dipole moment \cite{33}, see also section 4.1. Depending on the applied bias voltage, the SubPc or the C\textsubscript{60} molecules

\textsuperscript{4}For an example see also section 4.4.

\textsuperscript{5}For an example see also appendix A.
4. **Intermixing Mechanism**

appear more pronounced. At low voltage (e.g. 0.5 V), only the fullerences can be imaged, see figure 4.6. This is consistent with observations on pure SubPc films [33]. That the C∞ molecules can be observed also at lower voltage is a hint for a metallic coupling induced by charge transfer from the Ag substrate, as already discussed in section 4.1 and further treated in different contexts in section 4.2.3. A detailed analysis of various STM images, recorded with the same tunneling current of 20 pA, shows that the apparent heights of both molecules becomes similar only at elevated bias voltage, see figure 4.7. The functional relation looks reasonable, but is not straightforward, since the appearance in STM is governed by the electronic structure of the adsorbed components. We are tunneling from the tip into the LUMOs. The electronic structure of free SubPc is known from computations [60], that of C∞ also from inverse PES experiments [141]. The two DOS show more details like the three peaks of C∞ (at 0.4, 1.5 and 2.8 eV [60]) or two peaks in the case of SubPc (at 0.6 and 2.26 eV [141]) and therefore should lead to a more complex behavior than the measurements and the fitted curve of figure 4.7. More data is required to fully understand the observed bias dependence and to correlate it to theoretical explanations. At this step I just state that the appearance of the *stripe* pattern strongly depends on the applied bias voltage.

![STM image of a stripe island recorded at different bias voltages](image)

Figure 4.6.: STM image of a stripe island recorded at different bias voltages \(V_{\text{bias}}\), indicated on the left side. The tunneling current \(I_t\) was kept constant at 20 pA. At low \(V_{\text{bias}}\), only the C∞ molecules remain visible. Due to the specific electronic structure of the molecules, the appearance of the pattern strongly depend on \(V_{\text{bias}}\). Scan range 34 nm x 51 nm.

4.2.4. **Formation of a New Chemical Complex**

In the next few sections, I will show that the SubPc and C∞ molecules in the intermixed phases do **not** form a new chemical compound. The hypothesis was that the SubPc and
the $C_{60}$ molecules form building blocks of a new quantum state, e.g., a SubPc-$C_{60}$-SubPc complex in the stripe pattern. But potential associated molecule-molecule interactions and their changes in the electronic states are not seen in PES experiments or are too weak to be detected with our system. The photoelectron spectroscopy method is used to characterize the valence band and the $1s$ core level of the molecules in different combinations, as well as to determine the coverage of the molecular films.

**Coverage determination:** The coverage of the molecules was determined by comparing the $1s$ peak with the Ag$_3$d$_{3/2}$ peak. Thereby, 1 ML of $C_{60}$ means that the substrate is covered with 1 layer of fullerenes in the $2\sqrt{3} \times 2\sqrt{3}$ arrangement. 1 ML of SubPc stands for a complete layer of SubPc molecules in the honeycomb pattern. The obtained coverage for $C_{60}$ or for SubPc is proportional to the thickness obtained with the microbalance, as can be seen in figure 4.8. This is in no way surprising, but demonstrates that the molecules are confined to the 2D substrate surface and that they do not desorb after the deposition.

Furthermore, the value of the $1s$-area on a SIMP sample is simply the sum of the $C_{60}$ and SubPc signals, as they correlate perfectly with the sum of the microbalance values. Either molecule does not affect the other component and the first component (SubPc) remains at the surface even after the co-deposition of the second component (which finally is a weak form of “sputtering”). The deposition process therefore is additive, relating to section 4.3. In addition, these linear relations in figure 4.8 show that all measurements were done in the submonolayer range where no 2nd layer effects yet take place. Weaker bonded molecules of a second layer would influence the XPS signal [142] and lead to a kink in figure 4.8.

![Figure 4.7: The ratio of the apparent heights of $C_{60}$ and SubPc is shown as a function of the applied bias voltage $V_{bias}$. The tunneling current $I_t$ was kept constant at 20 pA. At elevated $V_{bias}$, the apparent heights of both molecules becomes the same, whereas at lower bias voltage, the $C_{60}$ dominate because their LUMO is partly occupied owing to a charge transfer from the substrate. The SubPc molecules are hardly visible at low $V_{bias}$, which is consistent with observations on pure SubPc films [33].](image-url)
4. Intermixing Mechanism

The absolute thickness calculation in units of monolayers is tricky because it uses some geometric factors describing the expected patterns. This can lead to large errors. However, a comparison to other UPS spectra [143] shows an excellent agreement.

![Graph](image)

Figure 4.8.: Comparison of the coverage measurement by the microbalance and by XPS. Both methods correlate well.

**C1s peak shows no new chemical component:** XPS measurements were first performed on pure C$_{60}$ thin films (θ < 1 ML). In the measured XPS spectrum (at the top of figure 4.10), only one single C1s peak at 284.9 eV relative to the Fermi level, could be resolved, corresponding to the interpretation that all carbon atoms of C$_{60}$ are equivalent. However, the overall line shape is slightly asymmetric, with a well-pronounced tail extending to the high-binding-energy side, as also was seen on other metals [144]. This indicates that fullerene adsorption leads to a metallic interlayer and breaks the molecular symmetry, thus promoting a variety of inequivalent C sites [144]. Studies done on thick films, in contrast, show a symmetric C1s peak [145]. A characteristic shift of the C1s peak as a function of the coverage is seen in figure 4.9. At a coverage of 0.4 ML, the shift is Δ = 0.35 eV. This shift can be explained by a reduced image screening for molecules in the thickest films [142, 146]. A controversial study [128] assigns this core level shift to a charge transfer because Scanning Tunnelling Spectroscopy (STS) experiments show the same shift as Photoelectron Spectroscopy experiments.

All these results are in good agreement with other fullerene studies. When comparing C$_{60}$ spectra to intermixed spectra, it is important that the coverage is considered because of its influence to the C1s peak position.

In addition, XPS measurements were done on pure SubPc and two-component films (spectra see figure 4.10). The C1s signal of the different films was analyzed. The chemical environment of the carbon atoms in C$_{60}$ and SubPc is different: the 60 carbon atoms of the

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Bulk fullerenes show a symmetric C1s peak because all C atoms are sitting in perfectly equivalent sites with each C atom being coordinated to two partner atoms through a double bond (hexagon side) and to the remaining C atom through a single bond (pentagon side), see section 3.2.

without the ability of photoelectron screening
fullerene are all bound to three carbon neighbors via two single bonds and one double bond (see figure 3.1). The 24 carbon atoms in the SubPc molecules are bonded in the following three configurations (see figure 3.1): 6 of them are connected to three carbon atoms, 12 of them to two carbon neighbors and a hydrogen atom and 6 of them to two nitrogen and one carbon atom.

The SubPc C1s spectra therefore show two peaks (see figure 4.10): at 285.1 ± 0.1 eV (C–C compound type) and at 286.75 ± 0.1 eV (C–N compound type), in accordance to [148]. The spectra of the C60 phase and the mixed layer (same figure) shown only one maximum at 284.9 ± 0.1 eV, respectively at 284.75 ± 0.1 eV (C–C compound type). In both C60 and mixed layer spectra, the peaks are slightly asymmetric with a tail extending to the high-binding-energy side. In the case of the pure fullerene this is attributed to the metallic interlayer (see above). In the mixed films, the asymmetric tail also covers the smaller C–N peak. Here, both types of molecules, each with different carbon environments, contribute to the measurement. It can be seen that all C–C peaks are at the same position within 0.15 eV. Thus, the carbon compound C1s peak in the mixed phase shows no drastic shift upon co-deposition. From these observations it can be deduced that the carbon atoms of the SubPc and the C60 molecules in the mixed phases still are in their original environment and the molecules are chemically not modified.

Valence band measurements show no new chemical component: Also in the UPS measurements, one can see that both molecules remain intact, just laying side-by-side, as is guessed from the STM images and XPS measurements. No new chemical compounds are believed to be formed.

Both SubPc and C60 films show up with their characteristic UPS spectra [33] (see figure 4.11). The UPS profile of the clean Ag(111) surface is dominated by the Ag 4d states spanning the 13-16 eV K.E. range (not shown here), while the featureless plateau located

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8 This residual shift probably arises from the coverage dependence of the individual peaks.

---

Figure 4.9.: The C1s peak of the C60 film shifts as a function of the coverage. The chemical shift, compared to bulk fullerenes, where the C1s peak shows up at 285.3 eV, points toward an ionic character of the adlayer–substrate interaction [113, 147].
4. Intermixing Mechanism

at higher K.E’s up to the Fermi edge are primarily due to the delocalized Ag_{5sp} states (see lowest curve). The small peak at 17 eV K.E. is attributed to a satellite of the d-bands by the He-I_{\beta} radiation with $h\nu = 23.08$ eV. The C_{60} (HOMOs at $\approx 18$ and $\approx 17.5$ eV K.E.) have a weak interaction with the low density Ag_{5sp} states of the substrate [143], so that the C_{60} molecular orbitals will retain their character [127]. The high density of states of the Ag d-band only start at higher binding energies. This advantage of Ag over other noble metals ensures a large energy window for the C_{60} derived states close to the Fermi level free from interference by the d-band. In addition, the Ag(111) surface is known to be stable with regard to reconstruction upon adsorption.

The C_{60} UPS spectrum is characterized by the manifold HOMO- and HOMO-n orbital related structures, in agreement with published results [147]. Figure 4.11 shows the region around the Fermi energy at different C_{60} coverage. The features at lower K.E. (not shown here) are mixtures of $\pi$ and $\sigma$ orbitals [149]. The two peaks at $\approx 18$ and $\approx 17.5$ eV K.E. stand for the HOMO and HOMO-1 and they are derived from $\pi$-states with an angular momentum $l$ equaling 5 and 4, respectively [150]. The HOMOs of the C_{60} are known to shift towards lower K.E. as a function of the coverage [147, 151]. When comparing to the mixed phase later, this shift has to be taken into account. The parallel shift of the HOMO and HOMO-1 as a function of the coverage is clearly visible, as summarized in figure 4.12. The simultaneous shifting is a clear sign for the ionic dominance of the Ag-C_{60} interaction, a so-called “rigid-band shifting” without covalent character. This shifting of the spectra can be explained by charge transfer [128]. Theoretical calculation of C_{60} and C_{70} support the rigid shift [145]. Another study [142] explains this shifting of the spectra by screening due to

![Figure 4.10: XPS spectra of the C1s peak of pure C_{60}, pure SubPc and a two-component film on Ag(111) measured with Mg-K_{\alpha} excitation. A chemical shift can be observed for the C–C and C–N carbon compound peaks of the SubPc film. The two different chemical environments of the carbon atoms in SubPc also lead to the asymmetry in the main peak of the mixed film. In all phases, however, the C–C peak-position is unchanged.](image)
the metallic first monolayer and the substrate. The valence band peaks at 13-16 eV kinetic energy related to the Ag(111) states are progressively reduced as C_{60} is deposited on the surface (not shown here). The progressive formation of C_{60} induced electronic levels can be observed. Spectra at sub-monolayer coverage are not affected by annealing to 500 K [143].

In pure Ag(111) samples, the Ag 5s\textit{p} background signal in the vicinity of the Fermi level is nearly zero. With small C_{60} coverage, the film gets a metallic character as can be seen in the extra-emission at the Fermi level in figure 4.13, top curve [144, 147, 151]. This is consistent with the Cls results which also confirm the ionic character of the adlayer-substrate bond (see figure 4.9) and the analysis of the apparent heights (see section 4.2.3). The new states at \( E_F \) reflect the tail of the \( \pi^* \)-derived LUMO orbital which is completely unoccupied in solid C_{60} and is sizeably shifted towards the Fermi level when the fullerene interacts with the Ag(111) substrate. This is clearly a strong confirmation of the ionic character of the C_{60}-substrate bond [112], the LUMO-derived extra-emission being the direct fingerprint of the charge transfer from the metal crystal towards the C_{60} molecule. Integration of this peak yields a charge transfer of about 0.75 electrons per C_{60} [127], consistent with vibrational energy shifts [129]. The term metallic here reflects the finite DOS at \( E_F \) as measured by STS or PES methods, but does not imperatively stand for a conductivity over large distances. For this purpose a lateral mobility of the charges is further required. The same formation of metallic-like interfacial states is also known from Perylenes (see 4.4.2) on Ag(111) [152].

The states at \( E_F \) effectively are new molecular states and not an artefact of the experimental method. It can be ruled out that the signal is a satellite (projection) of the C_{60} HOMO by the He-I\( \beta \) line (23.08 eV), although the positions offer this interpretation. From measurements on the pure Ag film I know that the He-I\( \beta \) signal is only 3\% of the He-I\( \alpha \) signal. The new states at \( E_F \), however, form 45\% of the LUMO. This is too intense to derive from the satellite line.

The SubPc spectra (see figure 4.13 and figure 4.14) shows characteristic features [33]: The HOMO is around 18.25 eV K.E., the same region where also the C_{60}-HOMO is located. Also the HOMO of the SubPc shifts to higher B.E. with increasing coverage, namely 0.4 eV from 0.4-8.4 ML, mainly attributed to charging effects of the molecular layer (more details can be found in [33]).

**UPS measurements on two-component layers:** We found that the valence spectra of SfIP films are identical to a simple linear combination of the C_{60} and SubPc spectra, see figure 4.14. Thus neither a peak broadening nor a shift due to the intermixing process can be found.

In figure 4.14, three measured spectra (pure C_{60}, pure SubPc and two-component film) and one calculated linear-combination of two spectra are shown in the range close to \( E_F \). The linear-combination of both single component films was fitted to reproduce the spectra of the mixed layer. This simulated spectra (weighted sum of the single-component spectra)\(^9\) and the measured spectra of the mixed film match perfectly. The intermixed spectra can be

\(^9\)The threefold degenerated LUMO can maximally be filled with 6 electrons.  
\(^{10}\)The associated factors are not useful because the experimental settings are not absolute identical.
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interpreted as a simple superposition of features from both materials. No additional peaks and no broadening of the features close to $E_F$\(^{11}\) point towards a possible chemical reaction of both components. The system is *additive* in its electronic properties in regard to section 4.3. Therefore it is concluded that no strong hybridization of the electronic states takes place and no new chemical compound is created. A ground-state electron transfer between SubPc and C\(_{60}\) is not observed or too weak to be detected and therefore is not believed to be the main intermixing reason. Note that in this linear-combination, the C\(_{60}\) fraction has to match that in the intermixed phase because of the strong coverage dependence of the HOMO peaks.

\(^{11}\) The $\pi$-derived features are most sensitive for a chemical bonding.

![Graph 1](image1)

Figure 4.11.: He-I ($h\nu = 21.2$ eV) valence band spectra of various thickness of C\(_{60}\) layer on Ag(111). The characteristic HOMO and HOMO-1 peaks simultaneously shift towards smaller kinetic energies. The individual spectra are scaled by an arbitrary factor to stress the shifts.

![Graph 2](image2)

Figure 4.12.: Rigid band shift of the C\(_{60}\) HOMOs with increasing film thickness indicate a charge transfer and screening.
4.2. Discussion of Other Intermixing Models

(figure 4.12). Secondary, the whole valence spectrum of the SIMP film can be described as a linear-combination of its components (not shown here), but below 16eV K.E. the Ag features strongly dominate due to the low sub-monolayer coverage of molecules.

This additive behaviour is in contrast to other intermixed systems [89, 113] where the ground state electronic wave functions of two components actually do mix. For example Metal-Phthalocyanines (M-Pcs), combined with C₆₀ in a study of novel photoreceptor material [113]. For M=Ni, Cu, TiO or the metal-free H₂-Phthalocyanine (Pc), all Pc derived features in the UPS spectrum shift on the binding energy scale (0.2-0.7eV) towards the

![Graph](image1)

Figure 4.13.: UPS spectra close to $E_F$ of different films. In contrast to the SubPc layer, the C₆₀ and the two-component films show filled states at the Fermi edge.

![Graph](image2)

Figure 4.14.: Valence band spectra close to $E_F$ of pure C₆₀, pure SubPc and a two-component film on Ag(111) with approximately the same amount of the components, together with a calculated linear-combination of the pure films. All features of the SIMP spectra simply can be explained by the pure films, thus no strong interactions between the two components are believed to take place.
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Fermi energy upon contact with C_{60}. This observation of a change of the surface-band bending demonstrates that it is not sufficient to measure the spectral feature of the single components separately. Only a measurement of the two components in contact with each other gives the true values for the energy differences between the valence-band onset of C_{60} and the different M-Pcs. That system is strongly non-additive in its electronic properties, referring to section 4.3.

The SIMP spectra (see figure 4.14 and 4.11) show a metallic characteristics like the pure C_{60} film at low coverage \( \theta_{C_{60}} \). This is not mandatory linked to a SIMP property like “conducting nanowires” but could also derive from pure C_{60} phases, which form residual metallic \( 2\sqrt{3} \times 2\sqrt{3} \) islands.

**Limitations:** However, if there were new signals due to a chemical interaction, they are expected to be weak because of the relative small ordered SIMP domains within a large disordered gas-phase (ordered domains roughly estimated to 10%). The non-locality of the PES method leads to a fundamental restriction. In contrast to STM studies, large numbers of domains are statistically averaged. The handicap of small signals at thin films cannot be circumvented by using thick films, because the SIMP structures need close substrate contact and therefore are limited to 2D.\(^{12}\) A second problem as a matter of principle is the similar HOMO peak positions in the valence bands of both components. And also a satellite from the He-I_A line unfortunately falls into this range. Measurable effects upon catalytic promotion are hardly detectable even in atomic systems with strong interactions and, moreover, are known to be even weaker in molecular systems [89].

4.3. Thermodynamics of Mixtures

The two kinds of molecules in the mixture are called components. If one component strongly predominates, then one speaks about a solution, otherwise of a mixture. In these kind of mixtures, new phenomena can emerge that don’t exist in pure substances. Such two-component systems can contain aggregates of macromolecules, polymer-solutions, gels, soaps, liquid crystals, lipids, micelles, colloids and suspensions and are of major biological and technical importance [153]. The here observed phenomenon is known in 3D when two strongly unequal interactions between molecules leads to a demixing of a liquid phase [154]. Thereby, two or more liquid phases of different relative composition \( c \) and density \( \rho \) are formed and are in equilibrium with each other. This is known as partial miscibility of two liquids and as mixing gap, where the mixture is separated into two coexisting stable phases with different relative concentrations \( c \) (so-called demixing into conjugated solutions).

The properties of a multi-component system result from those of the components and of the effects which occur during the mixing process. The macroscopic behaviour of a multi-component system cannot be derived directly from the properties of the isolated components, but is determined also by the inter-molecular interactions. Generally, the physical interac-

\(^{12}\)And up to now, it was not possible to image a SubPc double layer at room-temperature, not even at very low tunneling currents.
tion that appear between the particles of a system stem from the charge of free ions, from free or missing electron pairs or from permanent or induced displacement of charge which leads to a dipole effect and is found in many molecules. These physical interactions which affect the free mobility by the attraction or repulsion between the molecules and by the own volume of the particles, lead to changes of the properties of the mixed particles compared to the isolated ones. As a consequence, the external observed macroscopic properties are always a sum of the molecular properties of the isolated particles and the part which comes from the interactions. The properties of a multi-component system consist of contributions from the single components proportionally to their quantity. Only if the pure materials are assembled as macroscopic sub-systems, the properties of the pure components are additive [153]. Otherwise, a spontaneous mixing process takes place and change the properties of the system. These changes are only due to the mixing process; the temperature and the pressure can remain constant. These changes in non-additive systems are caused by a dispersion of the molecules and by changes of the inter-molecular interactions. The former reason is independent of the material and the system and is called ideal part of the modification. This is the case if we theoretical keep the inter-molecular interactions constant during the mixing process. The result then is an ideal mixture. The changes in the intermolecular interplay, however, leads to the real effect or excess part. It contains the material or system specific properties. As every real system develops additional interactions that depend on the temperature, pressure, composition and the kind of partners, it always shows more or less variations from the ideal behavior. The total required space for example results from the particle volume and the interspaces, that depend on the interaction of the specific mixture. These effects of dispersion of the particles and the changes of interaction after mixing can be as tremendous as a changing in the state of aggregation. As is well known, a change of the melting point of a certain material often is applied by mixing with a suitable component, e.g. the defrosting of ice by NaCl. In the case of the SIMPs, real effects concern e.g. the arrangement, the symmetry, the apparent height of the fullerenes in STM images and the surface density, as previously discussed.

If the molecules are distributed homogeneous over the whole system and the same interaction are dominating everywhere, a system is called homogeneous. On the other hand, several states of aggregation simultaneously can co-exist within a system. Such a system with a discontinuity in the molecular distribution and in its interactions are called heterogeneous. Individual homogeneous areas describe the phases, here in mixed systems mixing – phases, see figure 3.15. Different phases are separated by phase boundary lines, where the system properties change discretely. Thus phases are those parts of a heterogeneous system that have properties and composition different from other parts and are separated by interfaces. Furthermore, a phase is homogeneous, i.e. has identical physical and chemical properties in all locations. Under equilibrium condition, all phases in a phase-system are in equilibrium, also two that eventually are separated by a third phase. The equilibrium therefore is independent of the spatial arrangement of the phases [155]. A spatial separation of phases, however, may reduce the diffusion into the equilibrium. The phases in the schematic phase diagram (figure 3.15) agree with this thermodynamic definition, albeit they can be limited
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kinetically.

The miscibility generally is temperature-dependent. In certain temperature ranges, the components are not miscible in all concentrations. If two two-component phases co-exist, this state is the mixing gap. The mixing gap in liquid systems can have the forms shown in figure 4.15 (white areas). No states exist in the white areas, but two phases with the relative concentrations given by the boundary line (coexistence curve) coexist in equilibrium. In the dashed areas, a homogeneous phase of both components exists. The phase diagram of our SIMP system, so far only investigated at room-temperature, is believed to resemble case a) and c). 2D crystal structures are believed to exist only at certain temperatures: case c) for $T$ being the temperature during deposition, case a) for $T$ being the temperature of an already grown layer. Above the upper critical temperature, only a gas-phase containing both components (see section 3.4.2) is expected to exist. Cooling probably freezes the pattern and conserves it unchanged. A deposition below the lower critical temperature hinders the molecular diffusion and prevents a pattern formation.

Figure 4.15.: Isobar $T$, c-diagrams for the phase behavior of 3D binary systems with limited miscibility in the liquid phase. In the dashed areas, a homogeneous phase of both components exists. a) For instance the Phenol-Water-system: a Water-rich and a Phenol-rich phase can coexist up to an upper critical solubility-temperature $T_c$. Above the phase boundary line, a homogeneous mixture is found at any composition, at a certain higher temperature, a liquid-gas-transition takes place; b) System with lower critical solubility-temperature, e.g. Triethylamin-Water; c) System with lower and upper critical solubility-temperature, e.g. Nicotine-Water; The assignment of the SIMP system needs further investigations, most notably variable temperature STM experiments. Graphics from [153].

With help of the thermodynamic functions, it can be tried to predict if a system is homogeneous or heterogeneous. It is deciding if the decay of an original homogeneous system into various phases is associated with an increase in entropy or not. In the first case, a freshly built phase of microscopic range will gain size at the expense of the existing ones. The system will get heterogeneous or the original phase will even disappear completely. If the entropy decreases, the microscopic new phase disappears and the system remains homogeneous.

Yet in simpler systems like real gases in 3D, e.g. Ne-Kr, there exist some cases with a
limited miscibility that results from the Coulomb-interaction between the molecules [156]. If the potential energy $U_{AB}$ between various molecules $A$ and $B$ is larger than between similar components $U_{AA}$, resp. $U_{BB}$, then a mixing gap can appear, accompanied by a separation into two phases with different concentrations. But not only the orders of the potential energies are controlling the process. Furthermore, it is important how the entropy and the volume of the mixture depend on the concentration. Considering all this, the proper thermodynamic term of a mixture is its free enthalpy $G_m(P, T) = U_m - S_m T + V_m P$, $m$ being the index for mixing. The potential part of the internal energy $U_m$ from a mixture of two kind of molecules $A$ and $B$ with the concentration $c = \frac{N_A}{N_A + N_B}$ and the coordination number $z$ (number of nearest neighbors of a molecule) (if $N_A + N_B = N$) is [157]:

$$U_m(c) = \frac{N \cdot z}{2} [(1 - c)^2 U_{AA} + 2c(1 - c)U_{AB} + c^2 U_{BB}]$$  \hspace{1cm} (4.1)

This relation can be verified by counting the bonds of both kinds of molecules. If we assume a regular distribution of the molecules, the mean energy of a $A$-molecule in terms of its nearest neighbor is $U_A = (1 - c)U_{AA} + cU_{AB}$, the one of a $B$-molecule is $U_B = (1 - c)U_{AB} + cU_{BB}$. The total energy then is the sum of these average values over all molecules, i.e. $U_m = \frac{N}{2}[(1 - c)U_A + cU_B]$ which is equal to (4.1). $U_m(c)$ as a function of the relative concentration $c$ is drafted in figure 4.16 and supports the mixing.

In the case of the SIMP, $U_{AA} \approx 0.5$ eV, the hybridization strength of the C$_\infty$ with neighboring molecules [127], $U_{AB} \approx 0.42$ eV is the estimated\textsuperscript{13} strong C$_\infty$-SubPc intermixing term of Coulomb and Van der Waals interaction and $U_{BB} = 0.19$ eV the SubPc-SubPc interaction [75], which probably depends on the relative orientation and may alternate in sign with distance.

The entropy of a mixture is known from the thermodynamics. It is the part of the total entropy which depends only on the concentration $c$:

$$S_m(c) = -k \cdot N[(1 - c) \ln (1 - c) + c \cdot \ln (c)].$$  \hspace{1cm} (4.2)

This relation is obtained by counting the possibilities to distribute $N_B = cN$ molecules $B$ and $N_A = (1 - c)N$ molecules $A$ to $N = N_A + N_B$ places, namely $g(N, c) = \frac{N!}{(1 - c)N! \cdot cN!}$.

The mixing-entropy then is $S_m = k \cdot \ln g$. An approximation of the logarithm for $N \gg 1$ with help of the Stirling equation leads to equation (4.2). $S_m(c)$ as a function of the relative concentration $c$ is drafted in figure 4.16. The presence of two components in the mixed phase is preferred.

The volume of a mixture is not easily calculated as a function of the concentration. Therefore, in place of using the free enthalpy $G_m$, we look at the free energy (Helmholtz-Potential, the energy available for doing work or the energy that gives the forces [139])

$$F_m = U_m - S_m T,$$  \hspace{1cm} (4.3)

\textsuperscript{13} Empirically varied to enable to two well-pronounced minima of the free energy $F_m$. 

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the convenient thermodynamic potential for constant $T$ and constant $V$. $F_m(c)$ as a function of the concentration $c$ is drafted in figure 4.16 for positive mixing energy. The competition between entropy and energy finally determines the situation of the phases. Only if $U_m$ is convex against the $c$-axis (positive mixing energy $U_{AB} > \frac{U_{AA} + U_{BB}}{2}$), $F_m = U_m - S_m T$ as a function of $c$ can have two minima, see figure 4.17 (when the temperature is low enough). This is equivalent to two stable phases, in which the system is demixing to (at two distinct concentrations). Only at the two minima, the free energy is minimized and therefore the mixture splits up into two phases. This can happen only below the critical mixing temperature $T_m$, which can easily be derived from the above given relation. Below this critical temperature $T_m$, we find $\frac{\partial F_m}{\partial c} = 0$. Another necessary condition to observe the intermixing effect is the value of the averaged fluctuations of the molecular kinetic energy $\langle E_{A,B}^{kin} \rangle < (U_{AB} - \frac{U_{AA} + U_{BB}}{2})$. In two-component gas systems, often high pressures are needed (Ne-Kr: $p>1240$ bar [156]). However, with complicated molecules, in particular such with a permanent electric dipole moment as used here, phase-separations in gas domains are common even at low pressures. But because of the complex potential, calculations are impracticable. This short calculation here proves the presence of a mixing gap in the system when $U_{AB}$ is chosen to 0.42 eV. However, the values for the potential energies still have to be determined more accurate.

![Figure 4.16: Relation between entropy and energy for $U_{AA}=0.5$ eV [127], $U_{BB}=0.19$ eV [75], $U_{AB}=0.42$ eV. At the relative concentration $c=0$, only $A$-molecules exist, $c=1$ stands for the pure $B$ phase. The competition between energy and entropy determines the emerging phases.](image)

With regard to figures 4.18 and 4.19 we recognize that the compact $(2\sqrt{3} \times 2\sqrt{3})R \pm 30^\circ$ structure of pure $C_{60}$ is closed by straight borderlines with $120^\circ$ angles. This can be interpreted as a minimization of the interface (phase boundary lines) and the phase boundary energy. In other words, the fullerenes tend to aggregate and maximize their number of

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14This is the change in energy that appears when a molecule is brought from the inside of the system to its surface.
4.3. Thermodynamics of Mixtures

neighbors. The result is the best approximation to a circle on a (111) lattice which is a hexagon itself.

Figure 4.17.: In a mixed system, the free energy can have two minima, leading to a demixing at distinct concentrations of the components.

Figure 4.18.: Rough Ag(111) surface with C\textsubscript{60} islands of hexagonal shape. The derivative of the piezo-movement is displayed to simultaneously show the details within the terrace and the huge step bunches. The C\textsubscript{60} molecules are individually resolved when scanned on a reduced area (not shown here). The fullerenes tend to aggregate and maximize their number of neighbors. This results in a hexagon with the preferred border/inside ratio. Scan range 170 nm x 170 nm. \( V_{bias} = 2 \text{V} \), \( I_t = 15 \text{pA} \).
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Figure 4.19.: Stripe domain containing a pure $C_{60}$ island. The interior angles of the island always are 120°. This conglomeration of the exceeding fullerenes shows the minimization of the phase boundary lines and the phase boundary energy. Scan Range 50nm x 30nm. $V_{bias} = 1.3\, V$, $I_t = 20\, pA$. 
4.4. Typically Segregating System: Perylen – \( C_{60} \) Codeposited on Ag(100)

As another system of coadsorbed molecules, \( C_{60} \) and Perylenes on Ag(100) was studied. Here, the molecules do not intermix, but segregate into single component phases. The (100) substrate orientation was selected because its symmetry agrees with the rectangular-shaped Perylenes, see figure 4.23. Details about the Ag(100) substrate and its preparation can be found in section 2.2.

4.4.1. \( C_{60} \) on Ag(100)

The single component \( C_{60} \)-phase on Ag(100) (see figure 4.20) shows that the \( C_{60} \) molecules are randomly ordered after the deposition process at room-temperature. There is no evidence of any preferential growth direction, and large holes are present inside the islands [117]. The (100) surface of this fcc crystal is rougher than the (111) surface [158] (see figure 2.3), and the molecules cannot cross the diffusion barrier to aggregate in compact islands. The lateral forces play a minor role. Only after an annealing cycle at 570 K for 10 minutes, they aggregate into dense islands, see figure 4.21. Remarkably, in these islands they can have two distinct heights, see figure 4.22. The islands are formed by bright and dark molecules which are organized along preferential directions. Looking in more detail, it is possible to distinguish two predominant arrangements of the bright \( C_{60} \) molecules (zigzag and square structure). The bright fullerenes align along preferential directions after annealing and form a rectangular c(6x4) superstructure. A third structure (linear) was seen by [117]. The number of bright \( C_{60} \) molecules is around 18% of the total molecules, and they do not change brightness in time, in contrast to other metal substrates. This can be attributed to a different chemical bonding to the substrate [112, 114]. On Ag(100) with a lattice unit of 0.409 nm, the intermolecular distance is 1.042 nm which is slightly larger than in bulk fcc \( C_{60} \) crystals (1.002 nm). The corrugation of the second layer (\(<0.1\) nm) is smaller than the first layer (\(<0.2\) nm) and without any difference in brightness, as is also seen on Ag(111) [142]. The reason could be a wobbling of the molecules during the scan, because it is known that the second layer is weaker bonded to the substrate than the first one and is not modified by direct substrate interaction [142]. This would enhance the mobility of the molecules in the 2nd layer and lead to a more smeared out appearance in STM images. Since there is no difference in brightness in the second layer, the adsorption site on the substrate is responsible for the observed film morphology [117].

The changeover from identical molecules in as-grown film to bright and dim molecules in annealed film is explained by the following model [117]: When a \( C_{60} \) molecule is deposited onto the Ag(100) surface at room-temperature, it is free to rotate at the bulk rate of 10^9 s^{-1} [112] like in the \( C_{60} \) crystal [159]. In these conditions, the molecule is not chemically bonded, and the charge transfer is inhibited or less effective. After annealing, the charge transfer sets in,\(^1\) promoting the bonding of \( C_{60} \) molecules and stopping the free rotational

\(^{15}\) 1.7 ± 0.08 electrons from the Ag(100) to the \( C_{60} \) molecule [160].
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motion or at least damping it down substantially [128]. At the end of this process, the molecules have assumed the equilibrium configuration, where the bright and dark ones differ in their orientation with respect to the metal surface and, consequently, with respect to the vacuum [112].

Thin crystalline films grow near equilibrium by one of the following three mechanisms: the Frank–van–der–Merwe (FM), the Stranski-Krastanov (SK), and the Volmer-Weber (VW) modes [161]. These growth modes are determined based on relative magnitudes of the surface energies of the substrate, the film and their interfacial energy [115]. Figure 4.20 and other studies [112, 114] show that surface energies at the C$_{60}$-Ag(100) (and also at the C$_{60}$-Ag(111)) interface satisfy the conditions for FM growth. Because of charge transfer across the C$_{60}$-Ag interface, the forces binding the C$_{60}$ molecules to their substrate are stronger than van der Waals forces among these molecules, ensuring the first layer growth in the FM mode [115]. According to these STM studies the first-layer molecules form a full monolayer coverage, completing their growth before the nucleation in the second layer [112, 114]. These studies have also shown that the second-layer molecules form close-packed structures without altering orientations of the first-layer domains.

It is also possible that the surface structure of the substrate may undergo reconstruction, given the strength of interactions due to charge transfer at the interface. If these interactions displace the surface atoms from their bulk termination positions, the substrate surface may restructure to minimize its energy. However, this and other STM studies [112, 114] have found no evidence suggesting that C$_{60}$ molecules induce reconstruction on the Ag surface.

![Image](image_url)

Figure 4.20.: C$_{60}$ on Ag(100): After room-temperature deposition, the molecules are randomly distributed over several terraces. Only when the 1$^{st}$ layer is full, a 2$^{nd}$ layer can grow. Scan range 95 nm x 95 nm. $V_{\text{bias}} = 2.5$ V, $I_t = 30$ pA.
4.4. Typically Segregating System: Perylen – C\text{60} Codeposited on Ag(100)

Figure 4.21.: C\text{60} on Ag(100) after annealing to 570 K for 10 minutes. Densely-packed islands are formed. Due to different chemical bonding to the substrate, bright and dark C\text{60} molecules appear. The bright molecules form pseudo-periodic structures. Scan range 65 nm x 65 nm. $V_{\text{bias}} = 1.8$ V, $I_t = 30$ pA.

Figure 4.22.: C\text{60} on Ag(100) after annealing to 570 K for 10 minutes. The bright molecules are distributed in zigzag or square structures along a preferential orientation. Scan range 17 nm x 17 nm. $V_{\text{bias}} = 1.8$ V, $I_t = 30$ pA.
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4.4.2. Silated Perylenes on Ag(100)

In recent years, Perylene has become a model molecule for the investigation of the interaction of functionalized polycyclic aromatic molecules with solid surfaces [152, 162, 163]. Figures 4.23 and 4.27 show the \( N, N', N'', N''' \)-Tetra(trimethylsilyl)-4,9-diamino-perylene-chinon-3,10-diimin molecule (from now on called Perylene) in a space-filling model. In condensed 3D crystals, these molecules form a monoclinic structure (see figure 4.24) with space group \( p2_11/\alpha \), \( a=13.73(2) \, \text{Å}, b=10.587(10) \, \text{Å}, c=12.885(10) \, \text{Å}, \beta=111.32(7)^\circ, V=1744(3) \, \text{Å}^3, Z=2, \rho_{\text{calc}}=1.141 \, \text{mg/cm}^3 \). The melting point of \( C_{32}H_{40}N_4Si_4 \) is 485 K [164]. In 3D the molecules are connected to each other via the trimethyl-silyl ligands.

Figure 4.23.: Space-filling model of the \( N, N', N'', N''' \)-Tetra(trimethylsilyl)-4,9-diamino-perylene-chinon-3,10-diimin molecule. Colors specify different elements: white: H, black C, blue N, green Si. For the chemical structure see figure 4.27.

Figure 4.24.: 3D Crystal packing of \( N, N', N'', N''' \)-Tetra(trimethylsilyl)-4,9-diamino-perylene-chinon-3,10-diimin [164].

The monolayer structure is very different from the 3D crystal structure, see figure 4.24. When adsorbed onto Ag(100) (see section 2.2) and confined to 2D, they lay flat on the substrate because of \( \pi \)-metal interaction of the aromatic Perylene-core. The binding probably is not as strong as in the case of unsubstituted perylenes which bind with a quite strong
covalent character [165]. The trimethylsilyl ligands presumably elevate the perylen core and lead to a weaker physisorption of the molecules. In spite of the averaging over the lateral corrugation of the surface potential, each Perylen molecule adsorbs at a specific adsorption site: the molecules arrange in a so-called herringbone structure where each molecule is oriented perpendicular to its neighbor, see molecular-resolved islands in figures 4.25 and 4.26. The self-organization occurs at room-temperature without any additional annealing. The length of the overlayer unit-cell is 2.8 nm. The electron distribution in the molecules favors this orthogonal arrangement (see the electrostatic potential surfaces in figure 4.27). In this system, again the domain boundary energy plays an important role. The islands prefer to be closed by straight lines and angles of 90°, as can be seen in figure 4.26. Neighbored domains can differ in angle, preferred 54°, as can be seen in figure 4.28. Amusingly, this pattern is popular in macroscopic objects like floor panels, as can be seen in the photo, figure 4.29.

Figure 4.25.: Perylen arrangement in 2D on a Ag(100) surface: STM measurement reveal a molecular herringbone structure. Scan range 15 nm x 7 nm. $V_{\text{bias}} = 1.8$ V, $I_t = 60$ pA. The derivative of the piezo-movement is shown.

Figure 4.26.: Perylen molecules form a perfect herringbone structure on Ag(100). The island is preferential closed by straight lines. Scan range 33 nm x 33 nm. $V_{\text{bias}} = 1.8$ V, $I_t = 20$ pA.
4. Intermixing Mechanism

4.4.3. Mixture C<sub>60</sub> – Perylen

This system leads to segregation because the individual components strongly aggregate to pure phases and \( U_{AB} \), the \( C_{60} \)-Perylen-mixing energy, is not strong enough to overcome this process: \( U_{AB} < \frac{U_{AA} + U_{BB}}{2} \). Figure 4.30 shows a Perylen-island of herringbone pattern,

Figure 4.27.: The Perylen molecules arrange in a herringbone structure where every molecule is oriented perpendicular to its neighbors. The molecules are lying flat on the substrate for a strong \( \pi \)-orbital-substrate overlap. The electron distribution in the molecules, as is seen in the electrostatic potential surface, favors the orthogonal arrangement.

Figure 4.28.: STM measurement of Perylen herringbone domains with different angular orientation on a stepped Ag(100) surface. The molecules self-assemble without additional annealing. Scan range 71 nm x 71 nm. \( V_{bias} = 0.7 \text{ V} \), \( I_t = 20 \text{ pA} \). The derivative of the piezo-movement is shown.
surrounded by disordered $C_{60}$ molecules. No intermixing takes place. Potential mixed phases obviously have a higher free energy $F_m$ than the two segregating phases.

Different annealing cycles up to 560 K for 15 minutes were performed to overcome potential mixing barrier. Even with help of this activation energy\textsuperscript{16}, the molecules do not mix. No changes can be observed in the pattern, except a slight reorganization of the fullerenes into a more regular arrangement.

\textsuperscript{16}so as applied to pure $C_{60}$

Figure 4.29.: Photo of a floor panel with herring-bone structure. Image size 1 m x 0.66 m.

Figure 4.30.: This image shows the segregating system Perylen – $C_{60}$ on Ag(100). The Perylenes form a perfect herringbone structure, while the fullerenes arrange around the Perylen-island. Imaged after annealing to 560 K for 15 minutes. Longer annealing induces pseudo-periodic structures in the $C_{60}$ islands but no intermixing. Scan range 52 nm x 52 nm, $V_{bias} = 1.8$ V, $I_t = 24$ pA.
5. Conclusion and Outlook

The production of such self-assembled intermixed molecular monolayers is an important step on the route to the creation of self-assembled functional devices such as molecular data storage or arrays of nano-wires. Thus the here presented technique has the potential for a novel concept for the growth of organic anisotropic thin films and nano-structures at surfaces. The observed intermixed pattern asks for further use in experiments and later maybe even in applications. Some ideas are discussed in this chapter. The potential of this new method, which currently is available for licensing [166], is shown. Finally, some concluding remarks close this chapter.

**Bistability:** A missing fullerene in a stripe leads to dynamics of the adjacent $C_{60}$. Figure 5.1 shows reversible changes of the molecular position as probably induced by the scanning tip at room-temperature. The $C_{60}$ close to the vacancy, as shown in the top of this figure, has been shown to hop between two different positions at a frequency (at least) in the order of the scanning frequency ($\sim$ Hz). This experiment clearly demonstrates the reproducibility of the positional switching.

![Image](image_url)

Figure 5.1.: Within a stripe, a single $C_{60}$ molecule reversibly jumps between two metastable positions which could be assigned to bit-states. In this illustration, the fast scan direction is vertical.
**Self-repairing:** All patterns of the schematic room-temperature phase diagram (figure 3.15) can be found directly after the room-temperature deposition, i.e. without any additional annealing. But for an energetic consideration we have to take into account the kinetic energy which is provided during the deposition. This could in principle supply the activation energy needed for the pattern formation.

Figure 5.2a shows the known *stripe* pattern. In this experiment, a deliberate destruction of the pattern with the STM tip through lowering the tunneling resistance was carried out. The pattern then is distorted (figure 5.2b). Later, the pattern is slowly reshaping towards the original arrangement (figure 5.2c and d). This proofs the ample mixing energy at room-temperature. The rebuilding process was monitored over time and is shown in figure 5.3.

![Self-healing of the stripe pattern](image)

Figure 5.2.: Self-healing of the *stripe* pattern: a) Perfect *stripe* domain, recorded at $V_{bias} = 1.6 \text{ V}$, $I_t = 20 \text{ pA}$, tunneling resistance $R_t = 80 \text{ G}\Omega$, scan range $22 \text{ nm} \times 22 \text{ nm}$, time 0 s. Notice that the SubPc molecules are hardly visible due to the rather low bias voltage $V_{bias}$ of 0.7 V (section 4.2.3). b) Same area after a tip-induced distortion at the center of the image (16 mV bias voltage $V_{bias}$ during $\approx 50 \text{ ms}$ for a lower tunneling resistance of $R_t = 800 \text{ M}\Omega$). Four stripes are perturbed. Image recorded with the same tunneling and scan parameters as in a). Time 170 s. c) Left row is recovered. Three rows still are broken. Time 227 s. d) Only two rows remain broken. Time 332 s.

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1The $C_{60}$ molecules are deposited at $\approx 570 \text{ K}$ which gives them a kinetic energy of $kT = 1.38 \times 10^{-23} \text{ JK}^{-1}$. $570 \text{ K} = 7.9 \times 10^{-21} \text{ J} \approx 50 \text{ meV}$. 

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Improvements of production: It was shown that islands of this new material that are \( \approx 100 \text{ nm} \) in extent can be created. Analysis of many such images taken from several samples indicates that these ordered regions represent about 10\% of the surface. Preliminary experiments indicate that it should be possible to grow these islands much larger, perhaps to entirely cover a metal support. Parameters that could produce changes in island size may be (1) the deposition rate, (2) the relative rates of arrival of the components, (3) the substrate temperature during deposition, (4) the post-deposition annealing, (5) using derivatives of the chemical components, and (6) using single crystals as substrates in place of multidomain films. Modifications of these layers in terms of both electron transport properties and the addition of substituents are possible to engineer. I believe this is only the first in a broad new class of materials based both upon weak intermolecular interactions and image charges associated with adsorption on metal surfaces. What remains to be seen is whether these structures can be extended upward by sequential deposition to produce new three-dimensional materials and whether the individual components can be manipulated reproducible.

Nanowires: For future application, it would be interesting to have conducting nanowires. It is not clear a priori, if the chains in the stripe pattern are conducting, i.e. if there is enough overlap between individual C\(_{60}\) molecules. It is however possible to polymerize molecules of a SAM pattern to make them electrical conducting by electron radiation of ca. 1000 \( \mu \text{C/cm}^2 \) [167]. The thiol's themselves are better decoupled from the substrate because they are attached by \( \sigma \)-bonds. \( \pi \)-adsorption (unfortunately) leads to a better substrate coupling. The polymerization and functionalization is an active research field [168, 169]. For an application, of course, the wires have to be electronically decoupled from their support. The wires therefore have to be grown on an insulator which is not at all clear to work at the moment. Experiments have to be done with scanning probe techniques to check pattern formation on insulators.

![Graph](image)

Figure 5.3.: After a tip-crash, the stripe pattern restores naturally. 500 sec after the destruction, 40\% of the initially replaced molecules are “naturally” reassembled in the original pattern.
Storage device: In a visionary context and on the basis of the described inventions, it seems plausible that there may be technological implementations of single molecule storage on the SIMP materials. The structures have a potential to serve as data storage devices [170]. The large spacing between the molecules enables a variety of different assignments between molecular arrangement and “information bit” patterns: The pseudo linear arrays of C∞ molecules can be taken as “hard” formatted track marks to identify the positions of each SubPc molecule within one line. After re-positioning of individual SubPc molecules [171-178], the modified pattern can be associated with a bit pattern. A global “reset” could be applied by a controlled annealing cycle as discussed in the previous section. A treatment at elevated temperature could be used to assemble the molecules in their original position. This may be useful to erase written information in a self-assembled data storage layer. Therefore the next experimental step would be the use a variable temperature STM and additionally the capability of manipulate single molecules.

Using these patterns, a limit in the order of Tbyte/Inch² could be approached, which is 2-3 magnitudes higher than that of the currently established “edge of technology” magnetic or optical storage techniques. The exact 2D registry of the SIMP and the recognition of molecules and their position allows for numerous variations in data bit to 2D pattern assignments: Redundant storage, parity bits and the assignment of “words” as multiple bytes are straightforward generalizations of the here proposed basic scheme.

One of the obvious limitations of these SPM based data storage techniques is the speed. The “Millipede”[179] approach serves as a parallel interface to circumvent the limitations of the inherently slow read/write cycles in polymer indentations. Our approach has an exceptionally high signal to noise ratio in STM: In Figure 5.4 it is shown, that a data trace (cross section of STM image over the stripe with a C∞ vacancy in figure 3.10) taken at 1 kHz bandwidth leads to a clear identification of vacancy and occupancy site. This high signal to noise ratio in the order of 20 suggests that significant speed can still be gained in the individual read/write cycles.

![Image](image_url)

Figure 5.4.: The line-scan over the stripe with two missing C∞ molecules from figure 3.10 shows the high signal/noise ratio of the potential storage device.

Concluding remarks: We conclude that the observed mode of self-assembly and intermixing emerges from a delicate balance of electrostatic and Van der Waals interactions between
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static and fluctuating charge distributions of predominantly polar and polarizable molecules which are bound non covalently onto a polarizable metallic substrate. Each SIMP has its specific structure, composition, symmetry and surface density. The type of pattern can be controlled by the ratio of the two molecular components during the deposition. The exact arrangement of the molecules within the 2D layers described here, offers possibilities beyond the properties which have made SAMs so useful. For the first time, control over the ordering of intermixed arrays of functionalizable molecules created by self-assembly has been demonstrated over hundreds of nanometers. The relatively large spacing between the molecules leaves room for local rearrangements which are currently under study. This motivates the exploration of nanoscale data storage concepts, and the study of electronic [180] and mechanical [140] processes at a single-molecule level, which has often been hampered by the difficulty to identify individual incorporated molecules when using other techniques of self-assembly [181].
A. Porphyrins: A Model Molecule to Study Conformational Flexure and Switching

In this chapter, molecular conformation of a surface adsorbate, Porphyrin, will be discussed, as well as its recognition by Scanning Tunneling Microscope. When a large molecule is adsorbed on a metallic surface, its internal conformation is generally modified leading to a change in its mechanics, electronic transparency and chemical reactivity. It is therefore of fundamental importance to be able to determine the internal configuration of a molecule when adsorbed on a metallic surface. In this respect, STM is a unique experimental method, which contemporarily allows to image the molecule on a surface [90, 182, 183] and to manipulate it [171-176].

The special quality of a molecule is that in its entity, it can be readily modified by synthetic chemists such that its interaction with both the tip and surface are specially designed and controlled. Such “synthetic” molecules can also be used as “building blocks” for artificial supra-molecular structures [68]. By their larger sub-units, molecular orientation can be recognized in STM data which is not possible for single atoms. The free conformation of a molecule in vacuum can be modified when adsorbed an a surface and also the substrate can undergo adsorbate-induced reconstructions [184]. In conjunction with chapter 3, it is interesting to note that the conformation of an adsorbed molecule can be changed by a coadsorbate. Benzene for instance, undergoes a triangular distortion towards Kekule-benzene on Ru(001) when NO [93] or CO [88] is codeposited on Pt(111).

The molecular structure and vacuum conformation of the investigated Porphyrin molecule are described in section A.1. Section A.2 gives an introduction into the subject of molecular conformation. The molecule, adsorbed on Ag(100), with all its features like step-edge decoration, island formation, asymmetric conformation, and switching is discussed in section A.3. Section A.4 finally concludes the observations.

A.1. Structure of a Free Cu-TBPP Molecule

Cu-tetra-[3,5 di-ter-butyl-phenyl] porphyrin (Cu-TBPP) belongs to a class of porphyrins with four phenyl-based substituents of symmetrically bonded, interconnecting carbon atoms on pyrrole rings [185], see figure A.3. Figures A.1 and A.2 show a space-filling model of the molecule. Cu-TBPP is an organic molecule where well-defined structural changes can be found: the substituents can be rotated around the $\sigma$-single bonds. Two pairs of sterically interacting H electron shells exist for the planar orientation of the substituent at the top right of figure A.2. The overall structure of TPP derivatives can be ascribed to a steric repulsion between the ortho-substituents on the phenyl ring and the outermost substituents on the pyrrole ring. Together with the competing effect of the degree of overlap with the porphyrin

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ring, this repulsion determines the rotation of the phenyl groups. Thus, the rotation angle
around each of the four phenyl-porphyrin $\sigma$-single bonds is the predominant factor that
determines the shape of Cu-TBPP. X-ray diffraction has shown that tetraphenyl porphyrins
TPP derivatives exhibit dihedral angles of 60-90° between the porphyrin ring and the phenyl
substituents in bulk crystal samples, depending on the nature of the substitution and their
local environment. In vacuum, the $d_{i}t$-butylphenyl substituents are oriented out of plane
with the porphyrin in the minimum energy state, see figure A.1.

![Figure A.1. Relaxed geometry (vacuum conformation) of a single free Cu-TBPP molecule. Colors specify different elements: C, dark gray; H, white; N, blue; Cu, red (see figure A.3). The molecule has a diameter of about 2nm and is about 0.9nm high.](image)

![Figure A.2. Series of space-filling models, showing the rotation of the upper right TBP-group around a $\sigma$-single bond.](image)

**A.2. Molecular Conformation**

Properties of molecular material generally derive from the chemical and physical properties
of each molecular sub-unit and from cooperative phenomena like conductivity. Next to
the chemical structure of a molecule which is fixed during chemical synthesis, the specific
shape of the molecule is defined by bond rotation, bending and flexure. This shape which is
generally referred to as a molecule’s conformation is a predominant factor for “functional”
properties of the molecular material in all molecular sciences. Both physical and chemical
properties of functional molecules as well as cooperative bulk behavior can be modified
by changing conformation. The particular important role of molecular conformation in life-science systems was recognized when the 3D electron density maps of bio-molecules were resolved in X-ray crystallography [186]. Later, this method was complemented with NMR techniques [187, 188] and molecular modeling (see section 2.6) to understand and visualize complex proteins. Up to now, one central challenge in life sciences remains the understanding of “Structure Function Relationship” towards the prediction of molecular function and properties given the synthesized structure of a molecule.

In this chapter, the goal is to shed some light at the concept of conformation for molecules when adsorbed at surfaces and interfaces. First it is important to note that none of the above-mentioned experimental techniques has given clues about the exact structure of molecules when in contact with a solid or liquid surface. Single parametric techniques like UV, vis. or X-ray Spectroscopy give only scarce insight into the exact structure of molecules. Diffusion measurements and surface X-ray Diffraction have revealed important statistical information about molecular motion and packing. For single molecular objects and small numbers of molecular assemblies, however, all these techniques fail due to their statistical nature. Small amounts of molecular material at surfaces, do not provide enough signal for analysis and furthermore, there is little clue about the exact atomic positions. Only recently, Jung et al. [140] have proven the conformational identification of individual molecules through high quality Scanning Tunneling Microscopy images with submolecular resolution. Donhauser [181] observed conformational switching of single chain-like molecules by STM and reported that the local environment is playing a critical role in the conformational relaxation. This process was further identified as an effect of the electric field rather than of the tunneling current.

In the history of biomolecular conformation, porphyrins have played a core role, as they are the group involved in oxygen binding to the heme and in photosynthesis. This role was identified from the early synthesis of the so called “picket fence” porphyrins where the porphyrin is substituted with asymmetric substituents. The asymmetry of these substituents allowed for the identification of different conformational isomers (atropisomers) which originate from the sterically hindered bond rotation of the phenyl-porphyrin bond. This was first studied by Gottwald and Ulman in 1969 [189] for tetra-arylporphyrins and compared to the simpler case of the bi-phenyl molecule. In this early experiment, column chromatography was used to identify the different isomers after isomerization which occurs at room temperature for the studied system. Hindered ligand rotation and ligand exchange are of prime interest, as they allow for the construction of bi- and multilabile mechanisms using molecular architecture.

Porphyrin overlayers were found to order orientational by annealing (periodic conformation orientation). The H$_2$-TBPP on Au(111), showed a conformational change [190] where the ground-state conformation undergoes a bending and leads to a saddle shaped structure: two opposite legs are lower. The “bending axes” of the molecules the as grown films are random, but after annealing, an ordered superstructure can be found over large areas: neighboring molecules then show perpendicular “bending axes”.

People from the Rieder group at the Freie Universität Berlin showed a reversible modification of a porphyrin molecule on Cu(211) with an STM tip at low temperatures [176, 191].

With an Atomic Force Microscope (AFM) the behavior of the Cu-TBPP under an ex-
A. Porphyrins: A Model Molecule to Study Conformational Flexure and Switching

external load was investigated at room-temperature [192, 193]. Frequency-distance curves were made on the Porphyrin and indicated a conformational change of the molecule. The measured force needed to bend down a leg was approximate 0.4nN.

The here used molecule was found to adapt itself to the substrate geometry when deposited to various substrates [140, 194]: due to its flexibility and internal degrees of freedom, the molecule undergoes substrate-induced conformational changes, as was observed with STM on Cu(100), Cu(111), Cu(211) [194], Au(110), and Ag(110) [140]. Thereby, the substituents undergo a rotation close to 90°.

Figure A.3.: Chemical structure of Cu-tetra-[3,5 di-ter-butyl-phenyl] porphyrin (Cu-TBPP).

A.3. Cu-TBPP on Ag(100)

The porphyrins were deposited by evaporation at approximate 570 K from a tantalum crucible onto atomically clean Ag(100) substrate which was kept at room-temperature, see section 2.2 for the substrate preparation and and section 2.3 for details of the molecular deposition. In figure A.4 with low molecular coverage ($\approx 0.05$ ML), one can recognize single molecules as square shaped objects. In an overview STM image, figure A.5, one can see that the molecules are nicely aligned along the substrate step-edges. The molecules decorate the steps because they are attracted by electrostatic forces occurring at this substrate discontinuity. At the steps, the electron density does not change abrupt, but is smeared out and leads to a surface dipole, the upper level being slightly positive, the lower negative [195].

At a higher coverage, condensed islands are created. Because of the high packing density, diffusion of the molecules is further hindered and sub-molecular features are visible in high resolution STM images. Individual Cu-TBPP molecules are roughly identified as four protrusions of square shape, see for example figure A.6. Both their dimensions and theoretical STM electron-scattering quantum chemistry calculations (ESQC) confirm that the lobes correspond to the di-ter-butylphenyl (DBP) ligands [194]. The porphyrin core does not contribute significantly to the image and appears dark because it is supported approximately 0.7nm above the surface and consequently is electronically decoupled from the substrate, in agreement with previous investigations [174].

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1 Theoretical calculations demonstrate [194]
that the tunneling current mainly flows through the legs, because the associated orbitals are coupled to the metal surface.

\(^1\)Owing to the weak overlap between the \(\pi\)-states of the porphyrin core and the phenyl-ligands at this bond angle, the porphyrin-phenyl bond of the TBPP is predominantly a single \(\sigma\)-bond that does not participate in the conjugated electron system.

Figure A.4.: At low molecular coverage, the Cu-TBPP molecules decorate the steps of Ag(100). The molecules can be identified as single square-like objects. Scan range 40 nm x 40 nm. \(V_{bias} = 1 \text{ V}, I_t = 150 \text{ pA}\).

Figure A.5.: Step decoration of Cu-TBPP on Ag(100) at low molecular coverage. Scan range 112 nm x 112 nm. \(V_{bias} = 1 \text{ V}, I_t = 150 \text{ pA}\). A slight median filter and superimposition of the derivative was applied. Artificially colorized.
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When the molecules are adsorbed onto Ag(100) at this higher coverage, they can appear in various forms, see figure A.6: There exists three different brightness’ of the four legs and most of the molecules have “mickey-mouse” shape (one bright leg that is neighbored by two dimmed legs and the opposite leg is dark, see figure A.7). The asymmetric STM image

![STM Image]

Figure A.6.: At higher coverage, the Cu-TBPP molecules form condensed islands on Ag(100). The individual legs appear with different heights. A height-profile on the marked line can be found in figure A.7. Scan range 11 nm x 8.5 nm. $V_{bias} = 1$ V, $I_t = 40$ pA.

![Height-profile]

Figure A.7.: Height-profile on the line drawn in figure A.6. Different heights are assigned to different leg-orientations.
of Cu-TBPP, as is also seen on Cu(211) [194], can be explained by a conformally flexed molecule. The determination of single-molecule conformation relies on the shape of the four lobes at each molecule. The relative position of the uppermost t-butyl group corresponding to each lobe of the molecule is determined by the rotation angle for the phenyl-porphyrin bond of each DBP substituent. The turned leg leads to a different electron distribution in the molecule and therefore to a different tunneling resistance. Conformational changes affect the symmetry of the lobes and the transport probability through molecular subgroups. Unflexed Cu-TBPP molecules with four identical legs can be identified as nearest neighbors. There exist three levels of leg height, however only faint contrast variations are observed for one class of legs. Figure A.8, the histogram of illustration A.6, shows the distribution of the recorded heights: The histogram is fitted by a sum of four gaussian curves representing the Ag substrate, the low, medium and high legs, respectively. From the fit, the relative apparent heights of the three cases can be derived, as well as their percentage. The emerging heights behave like low : medium : high = 1 : 1.34 : 2.08. Thereby, the measured height of the high leg is approximate 0.6nm. The areas scale like low : medium : high = 1 : 1.96 : 0.4, which means that twice as much legs are in medium position than in low position, and only half as much are in the high conformation. This is clear when we consider that the flexed molecules have one low, one high and two medium legs, but some molecules are not deformed and have four low legs.

![Histogram of STM image](image)

Figure A.8.: Histogram of the STM image in figure A.6: The four fitted gaussian curves represent the Ag substrate, low, medium and high legs, respectively. Most of the legs show the medium conformation and are approximately 35% smaller than the bright legs.

Because topographic STM images represent a convolution of the electronic and topographic structure of the surface (see equation (2.4)), a change of apparent height observed in STM images can thus be due to a change in the physical height of the molecule, a change in the conductivity of the molecule or both. F. Moresco [191] has shown by theoretical calculation (ESQC) that the tunneling current depends strongly on the leg orientation (several orders of magnitude). The core of the porphyrin is decoupled more in the deformed configuration than in unflexed molecules probably because the molecule is tilted and the center porphyrin plane is raised.
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It is assumed that the legs are turned out of the perpendicular equilibrium state because of the interaction with the Ag(100) substrate (substrate induced conformational change) that leads to a strained molecular film. The distribution of the dim and bright legs is not purely random: The bright legs are distributed in a pseudo periodic super-structure. It looks like the bright legs of different molecules repel each other, maybe due to steric interaction. Since the porphyrin powder is of high purity, it can be ruled out that the observed brightness variation is due to the presence of other molecules or defect porphyrins.

This conformational adaption within the molecules occurs in addition to the general tendency for an adsorbat molecular lattice to rotate with respect to the substrate in order to minimize the inequivalent number of adsorption sites. The different apparent heights in the STM images could in principle also be explained by different adsorption sites or by an electronic effect as in the case of the C_{60} (section 4.4.1). In connection with the discussed observation of the conformational changes with AFM and STM and the observed flexibility on different substrates, however, this conformational interpretation is justified.

The substrate cannot easily be imaged simultaneously with the molecules. But by lowering the tunnelling resistance, the Ag lattice gets visible. Figure A.9 shows both, the substrate and the adsorbates, and can help to derive the exact adsorption registry.

![3D illustration of an STM image with a Cu-TBPP island and atomic resolved Ag(100) substrate. The tunneling parameters were changed close to the upper end during the scan from molecule-sensitive to atom-sensitive. V_{bias} = 1 V, I_t = 40 pA for molecules (lower part); V_{bias} = 0.1 V, I_t = 150 pA, 10x faster scan-speed (0.015 s/line) for atoms (upper part). The substrate lattice (with a lattice constant of 0.289 nm) and the Porphyrin superstructure (with an intermolecular-distance of 1.4 nm) are rotated approximately 6° against each other. Scan range 15 nm x 15 nm.](image-url)


**Spontaneous conformational change** We also found that some legs change brightness, i.e., from dark to bright and vise versa during sequential imaging of the same surface area at the same scanning conditions, see figures A.10, A.11, and A.12. This effect is assumed to derive from a conformational change of the molecules. A reason for this spontaneous (or probably slightly tip-induced) intramolecular dynamic could be the overall strain in the monolayer, which is a consequence of the intermolecular and strong molecule-substrate interaction. This is also believed to lead to the asymmetric conformation because the substituents can more closely pack in this conformation. This view is supported by the observation, that neighboring molecules show correlated changes in shape: The “high” substituent avoid to exist in the nearest neighbor positions. It looks like the turned legs repel each other and a conformational change of one molecule thereby can induce a change in a second molecule, see figure A.10 a-c). The pseudo-periodic superstructure then is better conserved. The “blinking” molecules seem to be concentrated to certain regions, maybe caused by underlying substrate defects.

![Figure A.10](image)

*Figure A.10:* Three successive STM images a), b) and c) of the same Cu-TBPP island on Ag(100), recorded with identical scan and tunneling parameters. The time between the images is 103 s. The two marked molecules change the appearance during the scans: the molecule labeled by a circle changes from *south* to *west* conformation, whereas the molecule labeled by a square changes from *west* to *north* conformation. All other molecules are unaffected. Scan range 9 nm x 6.5 nm. $V_{bias} = 1$ V, $I_t = 40$ pA.

In some images, see figure A.13, there exist regions without any bright molecules. In the next scan, however, the bright features can be back at the original positions. This could be due to a tip change that turns the tip insensible for the identification of different legs and that this is an imaging effect. Thus, not every tip is sensitive enough to identify single molecules conformation.

### A.4. Conclusion

The changes in the conformation and its detection render this molecule a model to study single molecular behavior. With porphyrins a single molecular storage device could be im-
A. Porphyrins: A Model Molecule to Study Conformational Flexure and Switching

implemented [197]: Apparent heights could represent the ON and OFF states of a switch. The molecules could probably be switched by an SPM tip. Or the molecules could be attached between two crossed electrodes of a nanowire array. A voltage pulse could possibly switch the molecule between two discrete states. Such devices already are under investigation, but at the moment still need thousands of molecules between the electrodes for reproducible switching [198, 199].

The idea of molecular switches is quite clear [200–202], but we have to take into account that the properties of such a switch will change dramatically when connected to wiring interfaces [203]. The molecular geometry will adapt to the environment which in turn affects the electronic structure of the first. A further drawback of the proposed storage idea is the coupling between the individual porphyrins.

In a more general sense, the observation of conformational effects for single molecules is an astounding achievement in experimental molecular science which may give us even deeper insight into important processes which occur at surfaces, like catalysis, molecular electronics and opto-electronics and tribology.

![Figure A.11.: Cu-TBPP on Ag(100). The scans were completed 103 s after each other. The conformation of the two imaged molecules has repeatedly changed between the scans. This image is rendered by the POV-Ray™ Raytrace program packet [196]. \( V_{\text{bias}} = 1 \text{V}, I_t = 40 \text{pA}. \)](image1)

![Figure A.12.: This color image is a composition of two successive STM image (from figure A.10), each with its own color (bright blue and yellow). The differences between the two images appear in the original color, whereas the unchanged areas result in purple. Two special molecules stand out due to an alteration of their conformation.](image2)
Figure A.13.: Three successive STM images of the same condensed Cu-TBPP island on Ag(100). The time between the images is 50 s. Only in some time slices during the scan, the tip had changed to become sensitive enough to distinguish between the intramolecular features of molecules with various conformation. Scan range 90 nm x 90 nm. $V_{bias} = 2.2 V$, $I_i = 20 \mu A$. 
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  *Multiphase adsorption of molecules in a two dimensional bottle: Two dimensional molecular motion and dynamic equilibrium.*

  *Conformational Flexure and Switching of a Single Porphyrin*

- Frühjahrstagung der Deutschen Physikalischen Gesellschaft, 11.-15.3.2002, Regensburg (Germany):
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Publications

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