Nano-layers against material deterioration in aqueous environment

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Nano-layers against material deterioration in aqueous environment

Ph.D Thesis

Talah Abohalkuma

This dissertation is presented as part of the requirements for the award of the Ph.D degree of Obuda University

Budapest
2018
Declaration

The research work reported in this thesis has been carried out under the supervision of Dr. Judit Telegdi member of the Obuda Doctoral School Comity. The subject matter of the thesis is original and has not previously been submitted in part or in full for any degree or diploma at this or any other tertiary educational institution.

Talah Abohalkuma
Dedication

TO MY BELOVED PARENTS
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<td>AFM</td>
<td>Atomic force microscopy</td>
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<tr>
<td>AC</td>
<td>Alternating current</td>
</tr>
<tr>
<td>AW</td>
<td>Atomic weight</td>
</tr>
<tr>
<td>CA</td>
<td>Contact angle</td>
</tr>
<tr>
<td>CE</td>
<td>Counter electrode</td>
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<tr>
<td>CS</td>
<td>Carbon steel</td>
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<tr>
<td>DC</td>
<td>Direct current</td>
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<td>EIS</td>
<td>Electrochemical impedance spectroscopy</td>
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<tr>
<td>IE</td>
<td>Inhibition efficiency</td>
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<tr>
<td>IR</td>
<td>Infrared</td>
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<tr>
<td>FTIR</td>
<td>Fourier-transform infrared spectroscopy</td>
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<tr>
<td>M</td>
<td>Metal</td>
</tr>
<tr>
<td>OCP</td>
<td>Open circuit potential</td>
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<tr>
<td>SAM</td>
<td>Self-assembly monolayers</td>
</tr>
<tr>
<td>SCE</td>
<td>Saturated calomel electrode</td>
</tr>
<tr>
<td>St. St</td>
<td>Stainless steel</td>
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<td>WE</td>
<td>Working electrode</td>
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List of Symbols

\( \rho \)  
Density of copper

\( F \)  
Faradays constant

\( \omega \)  
Frequency

\( \Omega \)  
Ohm

\( \eta \)  
Over-potential

\( \theta \)  
Phase shift

\( \alpha, \beta \)  
Transfer coefficients

\( \beta_a \)  
Anodic Tafel slope

\( \beta_c \)  
Cathodic Tafel slope

\( \text{Cl}^- \)  
Chloride ion

\( E \)  
Electrode potential

\( E_{\text{corr}} \)  
Corrosion potential

\( I \)  
Applied current

\( i \)  
Net current density

\( i_{\text{corr}} \)  
Corrosion current

\( R_p \)  
Polarization resistance

\( R_t \)  
Charge transfer resistance

\( t \)  
Time

\( Z \)  
Vertical distance factor

\( Z(j) \)  
Impedance

\( Z' \)  
Real part of the impedance

\( Z'' \)  
Imaginary part of the impedance
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Chapter One

1. Introduction

The metal corrosion is an important expense in all countries budgets every year. Its efficient control needs a lot of academic and industrial research, especially because the use of the most effective inhibitor – chromate – is banned since the last decade. Among the enormous number of chemicals used as inhibitors in case of different metals, the phosphonic acids - both in dissolved form as well as in molecular layers - can very effectively inhibit metal corrosion.

My work is focused on the research of self-assembled molecular layers (SAMs) of two special phosphonic acids. Both are amphiphiles; they consist of a –P (O)(OH)$_2$ head group (the geometry of the hydrophilic head group is nearly tetrahedral, and can act both as a hydrogen-bond donor via two P-OH groups and a hydrogen-bond acceptor through the P=O oxygen) and a hydrophobic side chain. These amphiphilic molecules differ in the hydrophobic molecular part. One has an alkenyl side chain with a double bond at the end position. I intended to polymerize this unsaturated bond to form a polymer net over the metal surface that could increase the compactness of the surface layer and parallel the anticorrosion activity. The other amphiphile is a semi-fluorinated alkyl phosphonic acid with several fluorine atoms along the carbon chain. It is well known that the fluorinated groups increase the hydrophobicity of the nanolayer-covered metal surface, and the barrier properties improve the anticorrosion protection in an aqueous solution.

The SAM layers are monolayers of organic molecules that form ordered molecular assemblies on the surface of an appropriate substrate and modify the metal surfaces by spontaneous adsorption of head groups that show affinity to the solid surface. This methodology is a versatile route for surface modification. The change in the tail group of the molecules in the SAM nanolayer easily results in different surface properties.

The complex investigation of self-assembled molecular layers, formed by the two amphiphiles mentioned above, makes up the main line of the dissertation. Some other amphiphilic molecules
were also investigated for comparison. The SAM nanolayers were deposited on iron alloys and aluminum surfaces from organic solvents of the amphiphiles. First, the effect of layer formation time on the quality of the SAM layers’, the amphiphile concentration as well as the post-treating of the undecenyl nanolayer was extensively investigated by specific techniques such as dynamic and static contact angle measurements in order to study the change in the wettability caused by the applied surface layers. The time-dependent layer formation was visualized by the maximum advancing and receding water contact angles. The SAM layer covered metal surfaces were visualized be atomic force microscopy (AFM). This technique, demonstrated in 3D and in surface section that allowed the observation of the change in the surface morphology caused by the presence of the nanolayers without or with post-treatment, on one hand, and, in the other hand, it monitored the influence of the electrolytes that could cause either general or pitting corrosion. The surface roughness parameters numerically characterized the change on the nanocoated surfaces subjected to corrosive environment. The anticorrosion behaviors of the SAM layers were investigated by electrochemical techniques like potentiodynamic polarization and electrochemical impedance spectroscopy (EIS). By these techniques not only the type of inhibition but also the effectiveness of the SAM layer was identified. All these complementary techniques demonstrated the usefulness of these two amphipiles as SAM layers in aggressive environment.

The extensive investigation of these SAM layers formed by the above-mentioned phosphonic acids could be in the future contribute to their use in different fields (e.g in electronic industry) as they can control the corrosion in molecular layer thickness (which eventuates decrease of materials applied against corrosion) in the presence of water (liquid or vapor).
Chapter Two

2. Literature survey

Metals such as iron and its alloys, copper as well as aluminum have found a wide use in technical and industrial applications and their corrosion has received a fundamental concern in the academic and industrial research (1). The metal surfaces are covered by inhomogeneous oxides, which are not stable against external influences.

2.1 Corrosion basics

Corrosion is known to be a chemical/electrochemical reaction between the metal or metal alloy and its environment, which leads to destruction. The energy of the chemical reaction that produces corrosion is equal to the energy amount needed to extract metals from their minerals. Consequently when a metal is extracted from its ore, it is transformed from a low energy state to a higher energy level due to the applied external energy. In return, metals do not stay in this higher energetic state but revert to a lower energy when different metal salts as corrosion products are formed.

In aqueous environment almost all corrosion processes of metals involve electron charge transfer. Accordingly, the electrochemical nature of a corrosion process is very important (2). Corrosion reaction depends on the type and rate of reaction, on environmental factors that are most important (3). It also depends on the microstructure, chemical nature, roughness, and heat treatment of the metal. The flow velocity of the electrolyte due to the mass transport distribution (4) influences the corrosion potential.

As soon as a metallic surface gets in contact with humidity, water or electrolyte, corrosion can take place by electrochemical reactions and two corrosion sites are formed on the metal surface; they are known as an anodic site and a cathodic sites. At the anodic site a charge transfer (electrons) occur which leads to a metal dissolution where metal atoms as metal ions dissolve
and form soluble ionic products or insoluble compounds of the metal that is generally an oxide. This electrochemical reaction is known as an oxidation reaction (5) and as electrons are produced, a less stable site is formed usually at surface areas that contain e.g., dislocation or imperfection. On the cathodic side the electrons are consumed and the reaction is reduction. The cathodic reaction depends on the pH of the media; either HO\(^{-}\) ions or hydrogen are evolved and on the metal surface oxide or hydroxide is formed (4, 6).

2.1.1 Electrochemical reaction

Corrosion reactions of metals can be symbolized with the following equations:

- Electrochemical reaction at the anodic site (metal dissolution) is:
  \[ M \rightarrow M^{2+} + 2e^{-} \] (2.1)

An example of the electrochemical anodic reactions is the dissolution of some metals such as:

- \[ \text{Fe} \rightarrow \text{Fe}^{2+} + 2e^{-} \] (2.2)
- \[ \text{Cu} \rightarrow \text{Cu}^{+} + e^{-} \] (2.3)
- \[ \text{Al} \rightarrow \text{Al}^{3+} + 3e^{-} \] (2.4)

- The pH-dependent electrochemical reactions at the cathodic site are:

1. In neutral or alkaline solutions:
   \[ O_2 + 2H_2O + 4e^{-} \rightarrow 4OH^- \ (\text{oxygen reduction}) \] (2.5)

2. In acidic solutions:
   i. in the absence of oxygen:
      \[ 2H^+ + 2e^{-} \rightarrow H_2 \ (\text{hydrogen evolution}) \] (2.6)
   ii. in the presence of oxygen:
      \[ O_2 + 4H^+ + 4e^{-} \rightarrow 2H_2O \ (\text{oxygen reduction}) \] (2.7)

2.1.2 Corrosion appearance

There are several corrosion forms such as uniform, crevice, galvanic, stress, intergranular, pitting, erosion corrosion, etc. My PhD work has focused on the investigation of uniform and
pitting corrosion that is the reason for me to give a short summary on these two forms of corrosion.

- **Uniform corrosion**: when the complete metal surface is submitted to the same corrosive environment, a large part of the metal surface is deteriorated (2, 7).

- **Pitting corrosion**: this is a localized form of corrosion that takes place at small areas on a metal surface and results in a rapid penetration. The size of orifice is much smaller than the depth of holes. Surface discontinuities can initiate pits (2). The presence of chloride ions increases the danger of pit formation where in the case of stainless steel in neutral and acidic solution the chloride ions increase the pitting corrosion. On iron and aluminum in alkaline chloride solution the mechanism of the pitting corrosion is the same (4).

2.1.3 Corrosion of metals

2.1.3.1. Corrosion of iron and its alloys

Products from the anodic and cathodic reactions can interact and form a solid corrosion product on the metal surface (7). An example of that is the interaction between the ferrous ions Fe$^{+2}$ (produced in anodic reaction) and the hydroxyl ions OH$^{-}$ (produced in cathodic reaction) as shown in equation (2.8).

$$\text{Fe}^{2+} + 2\text{OH}^- \rightarrow \text{Fe(OH)}_2$$  \hspace{1cm} (2.8)

The first formed ferrous hydroxide Fe(OH)$_2$ is transformed to ferric hydroxide, Fe(OH)$_3$ via oxidization by dissolved atmospheric oxygen:

$$4\text{Fe (OH)}_2 + \text{O}_2 + 2\text{H}_2\text{O} \rightarrow 4\text{Fe(OH)}_3$$  \hspace{1cm} (2.9)

Iron dissolution in neutral and alkaline media has similar mechanism (7, 8). These mechanisms are characterized by the formation of different oxide intermediates [Fe (OH)$_n$]$_{ads}$ depending on the pH and electrode potential.
Bessone’s proposal on the mechanism of iron dissolution in acidic media is that iron oxide is formed on the metal surface as the stability and protection of this film depends on several factors (9). In the pH range of 4 – 5.4 it is a time-dependent, porous oxide layer formation on the iron surface as reported Geana et al. (7,10) while \([\text{Fe(OH)}_2]_{\text{ads}}\) forms at a higher pH \(\geq 5.5\) when the iron dissolution is reduced. In alkaline solutions the first step is the formation of Fe (OH)\(_{\text{ads}}\) via the adsorption of OH\(^-\) ions on the iron surface (11, 12).

Guzman et. al. suggested structural rearrangements through a chemical reaction; the FeOOH and Fe(OH)\(_2\) transform into more stable compounds at higher potentials (13). Depending on the pH of the solution, the cathodic reaction can be either a diffusion controlled oxygen reduction or charge transfer controlled hydrogen evaluation, even though at all pH values the iron dissolution is the main reaction(14, 15). In aerated solutions at pH \(>4.2\) the dominant cathodic reaction is the oxygen reduction as reported by Lorbeer and Lorenz (16). Turgoose reported (17) that in unbuffered solution the pH increases up to 10 on the metal surface due to the oxygen reduction when the formation of a three dimensional oxide is favorable. A relationship between pH and passivation at a constant Fe\(^{2+}\) concentration and in alkaline state had been demonstrated by Nagayama and Cohen (18).

### 2.1.3.2. Corrosion of stainless steel

As a part of my research I worked with two stainless steels (304, 316), I summarize shortly the corrosion processes that would take place on their surfaces since they are the two most common stainless steel grades. The only difference is the presence of molybdenum in the composition of 316 stainless steel where as 304 stainless steel does not contain any.

- **Stainless steel 304** (which is mainly consists of 18% chromium and 8% nickel) is one of the most widely used stainless steel around the world, which is due to its excellent corrosion resistance. It can withstand corrosive attack of most oxidizing acids. Stainless steels do not corrode uniformly (i.e. they resist to general corrosion) as do the carbon steels do. The most common form of corrosion of stainless steels is the pitting corrosion when the passive layer on stainless steel is attacked by certain chemical species.
However, the stainless steel 304 does have one weakness: it is susceptible to corrosion in chloride solutions. The chloride ions (even in less than 25ppm concentration!) create localized areas of corrosion, called "pitting," which can spread beneath the protective chromium oxide barrier and interacts with the internal structures.

- **Stainless steel 316** is the second-most common form of steel used all over the world. It has almost the same physical and mechanical properties as stainless steel 304. The key difference is that stainless steel 316 contains about 2 to 3 percent molybdenum, which can drastically enhance the corrosion resistance, especially in environments that contain chloride ions.

2.1.3.3. Corrosion of aluminum

Aluminum and its alloys have good corrosion resistance because of the oxide layer formed as a protective film on the aluminum surface when it is exposed to oxygen. This coherent surface oxide hinders the further reactions with the environment and protects the aluminum surface from corrosion. In case of mechanical damage of the oxide layer, in the presence of oxygen it will be repaired immediately. The oxide layer, which provides protection against corrosion, is stable in the pH range 4-9. Out of this pH range, violent metal corrosion/dissolution will occur. Pitting corrosion is the most dangerous corrosion attack on aluminum; this could be very easily formed in the presence of chloride ions.

2.2 Corrosion protection

The corrosion processes i.e. the mechanism of metal dissolution and the cathodic reactions could be affected by the proper material selection, by dissolved corrosion inhibitors as well as by several methods such as cathodic protection, and surface treatments/coatings by paints, molecular layers and metallic layers (plating) (19, 20).

I will give a short summary on the corrosion inhibitors applied in dissolved form and then a more detailed discussion on organic nanolayers as this topic forms the main part of my research.
For a long time, corrosion inhibitors were used to protect metals against aggressive industrial environments (oil wells, refinery units, cooling systems, acid pickling processes etc.). At first, mainly inorganic chemicals (chromate, nitrite, nitrate, phosphates, etc.) were applied to protect metals against corrosion. Chromate is one of the most effective inhibitors, but it is already banned because of its toxicity (4). The need for developing chromate-, nitrite- and inorganic phosphorus free inhibitors drove the scientists to search for environmentally friendly organic inhibitors (21, 22).

Inhibitors are substances or mixtures that inhibit or minimize corrosion when added in small concentration to an aggressive environment (21). Corrosion inhibitors can slow down the rate of one or both of the anodic or cathodic reactions. It can happen through the formation of a barrier between the metal surface and the environment that would reduce the rate of the corrosion process. There is a wide range of inhibitor application almost in all industries and they can influence the corrosion kinetics. An efficient inhibitor should meet certain requirements such as formation of a barrier film, chemisorptions or precipitation on metal surfaces (23).

In general, the mechanism of inhibition can be one or more of the following:

i. Formation of an inhibitor thin film by interaction between inhibitor molecules and the metallic surface through chemical/physical adsorption.

ii. A film formation by oxide layer on the metal surface with the inhibitor.

iii. Production of a complex due to the reaction of the inhibitor with components present in the corrosive aqueous media (21).

Corrosion inhibitors can be classified according to their mechanism of action: cathodic, anodic or as a mix type of anodic- cathodic; adsorption; according to the chemical nature of the inhibitor as organic or inorganic, or oxidants or non oxidants (21). Inhibitors can also be classified concerning their retardation mechanism, whether they are interface or interphase inhibitors. As suggested by Lorenz and Mansfeld (24), interface inhibitors form a two dimensional (2D) films that reacts directly with the metal surface, assuming that the mechanism is a strong interaction.
between the metal surface and the inhibitor that forms a barrier due to the inhibitor adsorption as a two-dimensional layer. The 2D films react with a strong interaction directly with the metal surface and as the inhibitor adsorbs a barrier is formed.

The behavior of the inhibitor can be either by geometric blocking of metal surface and the solution; thus deactivation of the active sites by coverage, or by the inhibitor’s self-reaction that would take place instead of the metal layer. The other type of inhibition is the interphase inhibition when three-dimensional (3D) layers are formed by chemisorption or by reaction of the inhibitor with the film of the corrosion products; the inhibitors diffuse through the diffusion layer.

2.2.1 Organic inhibitors

Inhibitors used previously are of a wide range of chemicals and most are toxic in nature (25). Due to the environmental restrictions on heavy metal-based corrosion inhibitors, the researchers were motivated to study non-toxic and environmentally friendly corrosion inhibitors (26) such as organic compounds. Organic compounds are the majority of known inhibitors that contain hetero atoms (phosphorous, nitrogen, sulphur or oxygen), and a multiple bond for allowing adoption on the metal surface (25, 27). The efficiency of a corrosion inhibitor increases in the order of O < N < S < P (25). The adsorption depends on the electron density of the donor atom and of the functional group, which is influenced by the charge, type of electrolyte, the structure of the metal surface etc.

An organic inhibitor efficiency depends on the structure and the size of the inhibitor’s head group, hydrophobic part etc., the number and type of bonding groups (π or σ) or atoms in the molecule, ability to a complex formation with the atoms in the metal lattice, charges, and the nature of the metallic surface (substrate’s bonding strength). Existing data show that adsorption interaction between the inhibitor and the metal surface is how the most organic inhibitor behaves (28, 29). They form a hydrophobic film of adsorbed molecules, which acts as protective film on the metal surface. Adsorption of these organic inhibitors is dependent on the electron density at the donor atom on the functional group. It is
also influenced by the charge, type of electrolyte, and the structure of the metal surface (25). Organic phosphorus compounds as corrosion inhibitors are commonly applied on carbon steel, aluminum, and zinc due to their low toxicity (30, 31).

In my work amphiphilic phosphonic acids are in the focus. This is the reason that in the next part the phosphonic acid inhibitors are discussed.

2.2.1.1. Phosphonic acids as corrosion inhibitors

Organic phosphonic acids (which were developed instead of phosphates that increased the eutrophication of natural waters) are very effective in metal corrosion inhibition because of the stable P-C bond and the easy complexes formation of phosphonic groups with different metal/oxide/ions (6, 7, 10). They have been widely used in cooling water treatment because of their low toxicity, high stability and corrosion inhibition activity in aqueous media (32). They form strong bonds with several metal oxide substrates mostly through the formation of stable Me-O-P bonds (5, 33). The other factor that influences the effectiveness is the molecular structures of corrosion inhibitors that have important impact on the anticorrosion efficacy as pointed out by several authors (34-36).

Phosphonic acids as corrosion inhibitors employed often in case of different metals (iron, low-alloyed steel, stainless steels, zinc and aluminum) have been extensively studied because of their stable complexes with metal ions (24, 25, 28, 29, 37, 38).

i. Phosphonic acid corrosion inhibitors used in dissolved form

The use of dissolved inhibitors is one of the most practical methods for metal protection against corrosion. Inhibitors are those substances that inhibit, minimize corrosion rate when added in small concentration to an aggressive solution (19).

Researchers reported the protection of several metals against corrosion in aqueous solution by phosphonic acids (23, 39). They could be simple molecules with phosphono functional groups
like the hydroxyethan diphosphonic acid, nitrilotriphosphonic acid or the 1-phosphonobuthan tricarboxylic acid and there are several others where not only the phosphonic group, but also other molecular parts (e.g. amino, substituted amino etc) help to improve the anticorrosion properties. The anticorrosion efficiency of the phosphonic acids depends not only on the anchoring effect of the phosphonic group but on the hydrophobic molecular part, too. Shorter carbon chain results in less effective inhibition than a longer one although the activity depends on the water solubility. Also when the functional groups are not only in the α, but ω positions, they also can improve the anticorrosion efficacy (9, 11, 40).

As previously mentioned, the water-soluble phosphonic acids with shorter alkyl chain can effectively inhibit corrosion by adsorption to metal surfaces. In these cases the corrosion goes parallel with the deposition of the inhibitor molecules on the metal that could protect the surface from further corrosion, which means these are competitive reactions.

When metal surfaces - prior to corrosion attack - are coated with non water soluble phosphonic acids (with phosphono head group and bigger hydrophobic molecular part) the molecular film on the solid can control the metal dissolution in aggressive environment. This is the other possibility to control the undesired metal dissolution by coatings.

ii. Phosphonic acids in nanolayers

In contrast to inhibitors used in dissolved form, another possibility for corrosion protection is the application of coatings on metal surfaces. These layers could be macroscopic like in case of paints on metals, or a very thin molecular films and nanolayers that can also effectively hinder corrosion processes (6, 9, 41, 42). The selection of the protection techniques depends on the solids and on the corrosive environment. The organic nanolayers differ not only in the preparation method, but also in the thickness of the films formed on the metal surface.

There are several possibilities for thin layer preparation (vapor deposition, layer-by-layer deposition, sol-gel technique, spin coating electrodeposition etc.).
Especially for organic molecular layer deposition generally there are two methods used. The techniques that are mainly applied for molecular film preparation are the Langmuir-Blodgett (LB) and the self-assembling molecular (SAM) layer formation.

In corrosive environments, stearic acid nanolayers could inhibit the corrosion on iron (23, 31). Other studies include palmitic acid on aluminum (25, 43) and 12-amino lauric acid (44), which also effectively mitigated the corrosion. Considering the type of corrosion there are differences in inhibiting effectiveness of nanolayers. For example, the alkyl hydroxamic acid nanolayers are more efficient than fatty carboxylic acids in the prevention of pitting corrosion of copper (23). Due to the ability of the alkyl phosphonic acids to form SAMs on a range of metal oxide layers, they have become one of the most important classes of self-assembling organic molecules in anticorrosive coatings. The surface modification by these organic materials with functional phosphono head groups can ascertain the modified solid surface characteristics by the self-assembling process, thus the structure and the chemical properties of the surface are controlled. A variety of phosphonic acids are commonly used to modify the surfaces of metals and their oxides for their corrosion protection.

It is important to understand the molecular interactions involved in the surface modification and the effects that the modification has on the electronic state of the surface (45). The functionalization of normal alkyl phosphonic acids is easy by the formation of thin films, not only on pure metals but on the metal alloys and metal oxide surfaces which is due to the strong interactions between the adsorbing molecules and the substrate surfaces (46-48).

Surface modification of stainless steel is also an important part of the research. The phosphonic acid-steel interaction is significant from industrial point of view. At room temperature SAM phosphonic acid monolayer are formed on stainless steels. The compact coverage of the metal surface was confirmed by contact angle measurement and atomic force microscopy (49). In the case of a shorter carbon chain, especially under strong basic condition, the stability decreases (50, 51). Long chain alkane phosphonic acids adsorb onto metal surfaces (9, 28, 37, 52, 53) and form dense layer. When copper corrosion was in the focus, the use of alkyl phosphonic acids turned to be effective. The phosphonic groups interact with the copper oxide layer via
condensation reaction between the phosphonic head groups and the surface-bound copper-hydroxyl species, copper-phosphonate and different by-products are formed. These nanolayers are useful in micro- and nano-electromechanical systems (54).

The barrier property of octadecyl phosphonic acid nanofilms on oxyhydroxide-covered aluminum surface is a result of a strong acid-base interaction of the phosphonate head group with the aluminum ions in the oxy-hydoxide film. The phosphonic amphiphile in SAM layer on aluminum strongly reduces the amount of adsorbed water (55). Aliphatic groups or fluorinated groups in phosphonic amphiphiles increase the hydrophobicity of the coated metal surface, and act as a barrier to the aqueous environment at the same time improve the anticorrosion activity (56).

Alkyl-, benzyl- and fluorinated alkyl phosphonic acids were studied at critical interfaces between transparent conductive oxides and organic active layers in photovoltaic devices (57). In some cases the efficiency of amphiphiles with the same chain length (C16) with and without fluorine substitution were compared and the influence of the higher hydrophobicity of fluorinated alkyl chain was demonstrated (58).

The application of molecular nanolayer coatings in the electronic industry up to now is not widespread (59). However, this could be an important application possibility because several metals are involved in these systems and the phosphonic acid nanolayers can control the corrosion processes of these components.

2.3. Self-Assembled Monolayers (SAM)

2.3.1 Self-Assembled Monolayers theory

In the previous part I gave examples on the nanolayer application against corrosion. Now I give a detailed description on the technique, which results in molecular films, i.e. on the self-assembling molecular layer preparation, characterization. The study of this layer formation technique started in the 1940s (60) and in the last 20 years organic self-assembled monolayers have attracted a significant interest among researchers in order to prepare a surface with tailored properties.
The self-assembly is a nanofabrication method that has a number of advantages: the self-assembly is inherently a parallel process; it creates a structure with sub-nanometer precision; this process at molecular level can generate three-dimensional structure; external forces and geometrical constraints can alter the self-assembling. The self-assembling film is a monolayer of the organic amphiphile that forms spontaneously an ordered structure by adsorption and organization on a solid surface.

The self-assembly is a key tool in supramolecular chemistry. As a system, it lies at the interface between molecular biology, chemistry, polymer science, materials sciences and engineering. The formation of the nanolayer is the consequence of multiple weak intermolecular forces that leads to formation of large, discrete, ordered structures from relatively simple units; it resembles on self-assembled natural phenomena (amphiphiles with bioactive moieties, self-assemblies of peptides etc.). There is a great potential for their use as smart materials and surfaces of non-fouling properties, of corrosion resistance, and of molecular electronics. They are important in a variety of fields (chemistry, physics, biology, materials science, nanoscience). The application of SAMs is very divergent: increase in the non-wetting surfaces properties combined together with higher lubrication and enhanced corrosion inhibition, higher biocompatibility, applicability in lithography, etc.

Self-assembled monolayer is a powerful, simple and highly flexible means for functionalizing solid surfaces. The self-assembly is a spontaneous process when an ordered pattern develops from a disordered state. In other words, during this process an assembly of molecules and organized structures are formed via intermolecular forces that include weak non-covalent interaction (hydrogen bonding, π-π stacking, electrostatic interaction, ion-dipole interaction etc.). Through the self-assembly a new class of materials at molecular level are formed. Mainly two kinds of self-assemblies are discussed. Static self-assembly is when, via ordered structure formation, the system reaches an energy minimum (and do not dissipate energy: nanorods, nanoparticles, structured block polymers etc.). In dynamic self-assembly the system dissipates energy via formation of patterned components (biological oscillation, electronic circuits). Another categorization of the molecular assembly is the electrostatic self-assembly (alternate adsorption of anionic and cationic electrolytes onto the proper structure, e.g. layer-by-layer
assembly) and the self-assembled monolayers when the basic building blocks evolve via weaker or stronger forces (adsorption, van der Waals bond, hydrogen and coordinate bonds, hydrophobic interaction etc.) and create a spontaneously formed, well-defined structure. There are several factors that influence the self-assembled molecular layer formation like roughness and charge of a surface, polarizability, as well as the molecular structure of the amphiphiles (dipole character, ionizable groups, and hydrophobic molecular part).

The self-assembly requires mobile molecules, the layer formation happens in fluid phase when a nanolayer is formed at the solid/liquid interface in a simple and inexpensive adsorption method. The formation of a nanofilm with well-ordered structure is spontaneous and happens upon immersion of a solid substrate into a dilute solution of amphiphilic molecules, which have ionic (or ionizable) head group and bulky hydrophobic part. The functional head groups of the amphiphiles interact with the solid surface by chemisorptions or physisorptions when the molecules are anchored to the solid substrate (16, 33); this is determined by the binding force intensity between the functional group and the solid surface. The chemisorption represents high adsorption energy and strong metal-amphiphile interaction. The layer is organized through van der Waals interactions among the hydrophobic molecular parts, mainly between the long aliphatic chains (33). Minimum 11 – 12 carbons in the backbone are required for formation of a closely packed monolayer. It is accepted that there is a subtle balance between substrate-head group interactions and chain length-dependent intermolecular interactions that determines the growth kinetics of a film.

The preparation of thin films by self-assembling method permits atomic/molecular level control over the structure and composition of the exposed interface. The coated metal surface properties are defined by the head and tail groups in the molecules involved in the SAM.

Self-assembled monolayer (SAM) preparation is a flexible and simple method to form thin and well-defined organic coatings. It is a considerably new potential alternative for the pre-treatment of metal surfaces by ultrathin organic films such as hydroxamic and phosphonic acids (1, 61, 62). It is applied on a variety of solid surfaces where the deposition process is spontaneous upon the immersion of a solid substrate into a dilute solution containing organic adsorbate molecules. A
relatively strong bond between atoms or moieties in the molecule and the substrate and additionally lateral interaction of molecules in the monolayer is required for SAM formation (63). The functional group is accountable for the strong metal-molecule interaction, which is commonly a chemisorption interaction. The hydrophobic tail groups form the outer surface of the film and changes the physical and chemical surface properties (40). The long hydrophobic chains interact with each other via different forces (e.g. hydrogen interaction, van der Waals interaction). The results of the formation of a highly ordered molecular assembly are summarized in some papers (40, 64, 65).

Apart from the SAM layer formation, there are a number of other methods like Langmuir-Blodgett (LB) techniques. The Langmuir-Blodgett film preparation starts with formation of a compact Langmuir monomolecular layer of well-ordered structure built from amphiphiles at the air-water interface. The head group of the organic compound faces the water while the tail groups, the hydrophobic parts, hang away from the water (66, 67).

The Langmuir films transformation onto a solid substrate results in the Langmuir-Blodgett (LB) films (68). By repeating the dipping process of the substrate into the solution containing the organic molecules, multilayers could be produced on the surface. The effectiveness of the LB layers deposited onto copper and iron surfaces was published in several papers (9, 69).

The SAM technique used in a wide range of functional groups (40) has advantages over the LB preparation such as it is a simpler and flexible method, also there are no specific experimental equipment requirements for the formation of SAM thin films, and there is a strong attachment between the substrate and the formed layers through electrostatic and/or chemisorptions interactions. The preparation of thin films by self-assembling method permits atomic level control over the structure and composition of the exposed interface. The metal surface properties are defined jointly by the head and tail groups in the molecules involved into the SAM (70).
2.3.2 Self-Assembled Monolayers applications

Large numbers of molecules were already used in nanolayers like alkyl amines and carboxylic as well as phosphonic and hydroxamic acids, though in the very first set of experiments thiol amphiphiles with various carbon chains formed nanolayers on copper, silver, and gold. The assortment of molecules is determined by the metal and the functional group in the amphiphile.

The use of organic coating is by far one of the main methods used in corrosion protection. They form barriers between the metal surface and the media. In modern areas of materials research such as microelectronic devices or micromechanics, the SAM films with thicknesses less than 10nm have become of a great interest.

The iron alloys like stainless steels have been extensively used in different industries (chemical plants, medical fields like in the manufacturing of vascular stents or orthopedic implants) due to its resistance against oxidation and corrosion, relative ease of fabrication, and good mechanical properties. Self-assembled monolayers of long-chain carboxylic acids with different terminal groups were formed on stainless steel 316L substrates using the solution-deposition technique. SAM layers of alkanoic acids e.g. on stainless steel 316L were formed in a one-step solution-deposition method forming a bidentate bond with the substrate (71). Amphiphilic phosphonic acid SAM layers on stainless steels resulted in high contact angle (108°) which is much higher than measured on the unmodified stainless steel. This proves the presence of an ordered film (72).

The SAM formation provides a simple strategy to preparation of ultrathin and thermodynamically stable organic films; for functionalization of a metal surface by phosphonic acids is an easy technique (73). The self-assembling of alkyl phosphonic acids monolayers on metals such as on steel, stainless steel, and aluminum, is an easy route to modify a metal surface (74, 75). The structure of the barrier layer formed under this condition increases the anticorrosion intensity as the metal dissolution is significantly depressed by the formation of a stable, densely packed hydrophobic film, which hinders the contact between the metal/metal oxide surface and the aggressive environment.
Some other literature examples are also presented when corrosion inhibiting nanolayers were applied in corrosive environments. When a phosphonic acid SAM monolayer is formed on stainless steel 316L, the amphiphilic molecule is covalently bound to the surface as a bidentate complex, which was determined by diffuse reflectance Fourier transform infrared spectroscopy and X-ray photoelectron spectroscopy. The compact coverage of the metal surface was confirmed by contact angle measurement and atomic force microscopy (49).

All alkyl phosphonic acids molecules with longer chain (C > 10) form ordered monolayers with hydrophobic properties (47, 69, 76, 77), with excellent stability, even until 30 days in acid, neutral and physiological solution and for up to 7 days under dry heating. The stability of a layer with a shorter alkyl chain decreases, especially under strong basic condition (46,40).

Fluorinated alkyl phosphonic acids were studied in photovoltaic devices (57). The importance of the fluorine in the alkyl chain was demonstrated by their increased effectiveness compared with the alkyl amphiphiles with the same chain length (56).

In the medical field one of the applications of these amphiphile molecular layers is the coverage of Co-Cr alloys with drug-eluting stents against inflammatory reactions. Other territory of surface modification with alkyl phosphonic acid SAM layers is the implant biomaterials (e.g. titanium alloys, stainless steel, alumina, calcium phosphates) (78).

**Summarizing** the information appeared in the literature, I can emphasize that phosphonic acid nanolayers were intensively studied on different metal surfaces (8, 9, 17-23). It is clear that the increased hydrophobic molecular character enhances the compactness of a SAM layer; disturbance in the compactness (e.g. substituent in α-position to the phosphonic group) decreases the efficacy of the nanolayer. A densely packed film structure results in a significantly improved anticorrosion efficiency, the stable hydrophobic film decreases the metal dissolution. Different conditions like temperature, type of solvent for dissolution of amphiphiles, concentration of functional molecules, adsorption time and metal surface smoothness/roughness all play important role in the adsorption of amphiphilic molecules on a metal surface. In the adhesion of phosphonic acid the presence of oxide layer on a metal surface is indispensable (unlike in case of alkyl thiols that can adsorb only on pure metal surface, without any oxide layer). Several factors
explain the success of phosphonic acid amphiphiles such as: they bind strongly to a relatively wide range of metals and inorganic surfaces; the densely packed ordered phosphonic acid SAM layers are stable under ambient conditions that facilitate their application and storage.
Chapter Three

3. Employed Experimental Techniques

The nanolayers have special morphology, electric properties, and spectroscopic character. For the characterization of a SAM layer, a number of methods are applied. The change in the wettability is represented by contact angle values. For visualization of the surface morphology the atomic force microscope and the scanning tunneling microscope are proper instruments; different spectroscopic techniques help in the determination of the coating compositions like surface enhanced Raman spectroscopy, Fourier transform infra-red spectroscopy, and X-ray photoelectron spectroscopy that demonstrates the bonding between coating and metal as well as characterize the bonding states of the metal surface.

In my work I used the contact angle values for characterization of the nanolayer-modified metal surfaces to demonstrate the change in the metal surface wettability caused by the amphiphilic nanolayer. The morphological change in 3D, the section analysis and the roughening of metal surface with and without SAM layers, before and after immersion into aqueous solution (in the presence of oxygen as well as chloride ions) were demonstrated by atomic force microscopy. To measure the anticorrosion efficiency of the SAM layers, I used electrochemical methods. They are sensitive to the electron transport through the nanolayer and gives information about the compactness of the molecular film that is responsible for the corrosion protection by the SAM layers and in other set of electrochemical measurements (polarization) reactions were demonstrated (anodic, cathodic or both) that are influenced by the nanolayers. In the next part, the techniques used in my research work will be introduced.

3.1. Contact angle

In surface science the understanding of the wetting phenomena is a critical subject (79) since the surface wettability plays a great role in several processes including lubrication, printing, and coating (important in corrosion). Studies of the wettability include the contact angle values as
basic data. Its definition is the angle formed at the interface of the solid-liquid and the vapor-liquid (80). These values indicate the hydrophobic and hydrophilic properties of a solid surface. The hydrophilicity or hydrophobicity of the surface is determined by the surface molecular groups, which would indicate the properties of the layer formed on the solid surface. The contact angle values predict the degree of surface protection. If the contact angle ($\Theta$) value in water is $< 90^\circ$, then the surface is hydrophilic, while at values of $\Theta > 90^\circ$ the wettability in water becomes very low and the surface is hydrophobic (Figure 3.1) (81).

The force of wetting ($f$) is given as following: (80)

$$f = \gamma_{lw} p \cos \Theta \quad (3.1)$$

*where:* $f =$ wetting force; $\gamma_{lw} =$ liquid surface tension; $p =$ perimeter of contact line (i.e., the same as the perimeter of solid sample’s cross-section); $\Theta =$ contact angle.

There are several techniques for measuring contact angles: telescope-goniometer, captive bubble, tilting plate and Wilhelmy balance method (80). The measurements could be static and dynamic.
3.1.1. Static contact angle

A surface wettability is best described by the angle between the tangent of the liquid/solid interface at the three phase contact line and the baseline of the sessile droplet of a defined liquid on the solid surface, which is known as the macroscopic contact angle.

The setup of the telescope-goniometer has measured the contact angle of various types of liquids on a polished surface and afterwards appeared the contact angle goniometer. The measurements are achieved by measuring the sessile drop’s tangent angle at the point of contact with the surface and reading the protractor along the eyepiece. Modification of the equipment was made over the years to improve the precision and accuracy. High magnification is used in order to enable a detailed testing of the intersection profile. A camera takes the drop profile’ photograph as the drop relaxes. Motor-driven syringe can be used in controlling the rate of liquid addition and removal in order to study the receding and advancing static contact angles (Figure 3.2) (80).

The resulting angle is due to the interfacial energies at the liquid/vapor $\gamma_{LV}$, solid/liquid $\gamma_{SL}$ and solid/vapor $\gamma_{SV}$. These boundaries are expressed by the Young’s equation for an ideal surface as:

$$\cos \theta = \frac{\gamma_{SV} - \gamma_{SL}}{\gamma_{LV}} \quad (3.2)$$
In case of a real surface, several factors like roughness and chemical heterogeneity are not taken into account by the Young’s expression (79).

![Schematic presentation of static contact angle.](image)

Figure 3.3: Schematic presentation of static contact angle.

The advantages of this method is its simplicity and that it uses few microliters of liquid and a few square millimeters of the substrate’s surface; although the risk not to realize the impurities is higher because of the small size of the investigated substrate surface (80). Also a value less than 20° is not measurable precisely because it is hard to assign the tangent line. Figure 3.3 demonstrates schematically of the instrument.

### 3.1.2. Dynamic contact angle values measured by Wilhelmy balance

The surface tension of a liquid by a Wilhelmy plate apparatus is represented by the force of the liquid pulling down on a plate and measures the contact angle between the plate and the liquid. When a vertically suspended plate touches a liquid surface, then a force (F) acts on this plate, that correlates with the surface tension and with the contact angle (θ) according to the following equation: \( \sigma = \frac{F}{L} \cdot \cos \theta \), where \( \sigma \) is the liquid surface tension, \( L \) is the perimeter of the probe. Two different angles can be seen: as the plate enters the liquid, there will be an advancing angle, and as the plate exits, this is represented by a receding angle.
This technique is widely used on solid samples, which measures the contact angle indirectly. This method is based on the surface tension measurements whereas the flat plate is immersed in a liquid having a horizontal upper surface; the total force needed in keeping the plate fixed is related to the weight of the plate resistance forces and the surface tension $\sigma$. The advancing contact angle is measured because of the liquid advances on the immersed plate into the liquid. As a result of the withdrawing of the plate from the liquid, the liquid would recede so the receding dynamic contact angle is measured (Figure 3.4). The force change detected on the balance is the buoyancy and the wetting force where the force of gravity remains the same (80,81). The high difference in the advancing and receding contact angle values is either due to the inhomogeneous or the roughness of the surface.

![Figure 3.4: Schematic representation of the dynamic contact angle measurement.](image)

### 3.2 Atomic force microscopy (AFM)

The history of the scanning probe microscopy (where to the AFM belongs) started in the eighties of the last century. First the scanning tunneling microscope (STM) was developed (the inventors deserved the Nobel Prize for it) where the tunneling current is measured along the scanning. As the STM could visualize only the conducting materials, the researcher worked further and they invented the AFM which can visualize all surfaces irrespectively they are conductive or not. In the case of AFM (which is the most popular type of the scanning probe microscopy) the force
between a very sharp tip and the solid surface under investigation is recorded. This revolutionized the nanoscale characterization. The important parts of the AFM are as follows: a spring (cantilever) equipped with a tip, a piezo tube which can move the sample in the nanometer (nm) and some hundred micrometer range, a laser beam that illuminates the cantilever from where the laser light will be reflected (through a mirror) into a position sensitive photodetector. The heart of the AFM is the cantilever-tip assembly that interacts with the sample in a raster scanning mode. The AFM can characterize a wide variety of material properties achieving resolution down to the nanometer scale and beyond. The AFM can operation on air, in vacuum, under liquid, that is why it is applicable in a wide range of sciences (materials science, chemistry, physics, and biology) (82). The AFM can operate in non-contact mode (attractive force range), in contact mode (repulsive force range) and in tapping mode when the tip constantly changes its position between the two modes. This technique allows the surface visualization in 2D and 3D, as well as the section analysis gives numerical information about the surface irregularity and by the roughness parameters the solid surface could also be characterized. The operating principle of atomic force microscopy is illustrated in Figure 3.5.

Figure 3.5: Schematic representation of the AFM set-up (83)
3.3 Infrared (IR) spectroscopy

IR spectroscopy is one of the common spectroscopic techniques used by organic and inorganic chemists. By interaction of a molecule with the IR photons the molecular vibrations and rotations of the molecule are excited. A vibration (or rotation) is IR active only, when a change in the dipole moment of the molecule (or bond) occurs as a result of the molecular vibration (rotation) excitation. For example, symmetric molecules (or bonds) have no IR spectrum. Common bands of molecular bond and functional groups absorb at characteristic frequencies, called group frequencies, which enables identification and/or structural analysis of samples by means of IR spectroscopy.

3.3.1 FT-IR Spectrometer

The development of the IR spectrometers where the data is collected converted to a spectrum from an interface pattern is known as the Fourier Transform Infrared Spectroscopy (FTIR) term.

3.3.1.1 The Michelson Interferometer

The Michelson Interferometer Experimental Setup is the base of the FTIR spectrometers where the interferometer consists of a fixed mirror, a moving mirror (back and forth), a beam splitter of a special material. The radiation from the source strikes the beam splitter where it is separated into two beams, one of is transmitted to the fixed mirror and other is reflected to the moving mirror. Afterwards, the radiation is reflected back to the beam splitter by the fixed and the moving mirrors. Half of the reflected radiation is transmitted and the other half is reflected to the beam splitter which would result in a one beam that passes to the detector the other goes back to the source (84).

3.3.2.1 Microscopy and imaging

The IR microscope permits the observation and spectra measurements of regions as small as just 5 microns across. Images are generated by the combination of a microscope with a 2D array
detector with a resolution of 5 microns and thousands of pixels. The resulting images indicate spectrum for each of the pixel and is viewed as maps at any wavelength. This would specify the different chemical species of the sample (84).

The Michelson Interferometer Experimental Setup is the base of the FTIR where the interferometer consists of a fixed mirror, a moving mirror (back and forth), a beam splitter of a special material. The radiation from the source strikes the beam splitter where it is separated into two beams, one of is transmitted to the fixed mirror and other is reflected to the moving mirror. Afterwards, the radiation is reflected back to the beam splitter by the fixed and the moving mirrors. Half of the reflected radiation is transmitted and the other half is reflected to the beam splitter which would result in a one beam that passes to the detector the other goes back to the source (84) as shown in Figure 3.6.

![Figure 3.6: Schematic of Michelson Interferometer (85).](image)

3.4 Electrochemical Methods

Corrosion phenomena in aqueous environments being of an electrochemical nature and controlled by the corrosion potential of the metal, which makes the electrochemical methods
have an important impact on corrosion testing and research (86). As the corrosion is an electrochemical process, it is logical to use electrochemical techniques to evaluate the properties of nanocoatings.

The polarization is defined according to the ASTM as “the change of the open-circuit electrode potential as a result of the passage of current”. The change of the potential of the working electrode and detecting the produced current, which is a function of time, is the work of the polarization methods. During the last 75 years, the researchers had applied electrochemical techniques to the corrosion processes. In the 1930’s C. Wagner and W. Traud were the first developers publishers of the mixed-potential theory, that helped in the explanation of experimental results (86). The determination of the corrosion rate at which a metal dissolves under certain conditions and environment is of a great importance in the field and laboratory corrosion studies; several other techniques have been developed throughout the years in determining the corrosion, such as linear polarization resistance (LPR), potentiodynamic polarization, weight loss measurements, electrochemical impedance spectroscopy (EIS), Tafel extrapolation method, and electrochemical noise analysis, which is a more recent technique (86, 87). These methods are based on the detection of corrosion by the observation of the charge-transfer process response to a controlled electrochemical disturbance.

3.4.1 DC Methods

Techniques such as the LPR, potentiodynamic polarization are so-called direct current (DC) methods where the current is measured by applying a DC potential excitation.

3.4.1.1. Potentiodynamic technique

Potentiodynamic polarization is an electrochemical technique that presents a significant amount of information such as corrosion kinetics, corrosion rate of the working electrode, pitting, and passivity. It also determines the type of the electrochemical reaction, whether it is a cathodic (reduction) or an anodic (oxidation) reaction.
The electrode is submitted to a potential, and the outcome current is measured. The plotting of the potential (V) vs. the current density (A/cm²) results in the Tafel plots. The current density values are plotted in logarithmic scale.

The corrosion kinetic parameters such as corrosion potential (E$_{corr}$) and the current density (i$_{corr}$) are determined by intersecting both straight lines in the linear regions on the anodic and the cathodic curves. The slopes of the linear regions are represented by Tafel constants: $\beta_a$ for the anodic and $\beta_c$ for the cathodic reactions. All information is shown in Figure 3.7.

![Figure 3.7: Tafel extrapolation showing the anodic ($\beta_a$) and cathodic ($\beta_c$) Tafel constants.](image)

The corrosion current ($i_{corr}$) can be determined as:

\[
i_{corr} = \frac{\beta_a \beta_c}{2.303(\beta_a + \beta_c)R_p}
\]

where:
\begin{align*}
\beta_a &= \text{anodic Tafel constant expressed in Volts per decade current} \\
\beta_c &= \text{cathodic Tafel constant expressed in Volts per decade of current} \\
i_{corr} &= \text{corrosion current density expressed in current units per unit area}
\end{align*}
At the measurement setup, the scan rate has to be carefully chosen in order to obtain full information taking place on the electrode surface; thus relatively slow potential of e.g. 0.1mV/sec is preferable where at high potential less information can be obtained on the rate establishing step of the working electrode process and information on the fast reaction step.

This technique is widely used in corrosion studies as it can evaluate the efficiency of corrosion inhibition (9). It has also been used for the evaluation of self-assembled monolayers (SAM) formed e.g. by hydroxamic acids on carbon steel (9). In my study on carbon steel surfaces the polarization method had been used to measure the effect of layer formation time and the pH effect of the electrolyte.

3.4.2 AC method

In this method, an AC potential excitation is used instead of the DC potential excitation.

3.4.2.1 Electrochemical impedance spectroscopy (EIS)

The use of electrochemical impedance spectroscopy in evaluation of electrochemical and corrosion mechanism is due to its ability of probing electrochemical systems even at very low frequencies. This technique was developed after the elaboration of the potentiostat in the 1940s followed by the frequency response analyzer in the 1970s (13). The EIS has become a powerful tool in the study of metal electrode/electrolyte interface, coatings, porous electrodes, electrode/layer/ electrolyte interface (55). Additionally, electrochemical reaction mechanisms, kinetic, passive surfaces, materials of dielectric and transport properties of an electrode, protection properties of corrosion inhibitors, e.g: corrosion inhibition via phosphonic acids formed by absorption on metal surfaces (7, 88), metal coatings and thin film had been studied (89). It is a non-destructive method (advantages over the DC methods) since it uses very small excitation potential amplitudes, and has a minimal perturbation to the working system.

EIS as an electrochemical measuring method quantifies the protective behavior of coatings. EIS results can indicate the state of coating degradation, the efficacy of corrosion protection and can even identify protective mechanisms. EIS measurements on organic coatings apply alternating
voltage over a counter electrode and the coated metal substrate, and the response of the system is registered. Impedance \((Z)\) is a measure of a circuit’s tendency to resist (impede) the flow of an alternating electrical current and is the AC equivalent of electrical resistance \((R)\) in DC applications.

A sine wave voltage applied to an electrochemical cell gives a current response, which is shifted in time due to the influence of the system. This time shift is expressed as the phase angle. Since the impedance is frequency dependent, the measurements go over a wide range of frequencies. By measuring the high frequency impedance \((10^5 - 10^3 \text{ Hz})\) and using equivalent circuit modeling the time of measurement can be kept to a minimum.

Data achieved by impedance spectroscopy are fitted to an equivalent circuit when enough parameters (elements in the equivalent circuit) are used. It is preferred to fit the data to the most probable equivalent circuits. The fitting parameters are as follows: electrolyte resistance \((R_s)\): the solution resistance between the working electrode and the reference electrode, constant phase elements \((CPE)\): used instead of a capacitor to represent the non-homogeneity such as the roughness or the energetic inhomogeneity of the system, coating resistance \((R_c)\): electrochemical (corrosion) behavior of the system, metal double layer capacitance \((C_{dl})\): capacitance between the working electrode and the electrolyte; controlled by many variables such as temperature, electrode potential, types of ions, impurity adsorption, ionic concentrations, oxide layers, electrode roughness, etc), charge transfer resistance \((R_{ct})\). Two parameters can characterize the CPE. The diffusion on the metal surface is represented by the Warburg element \((W)\) that would appear at low frequency of the Nyquist plot (90, 91).

The EIS data can be displayed by two common methods, which are Nyquist and Bode plots. Nyquist plots are usually used for mechanistic analysis due to mechanistic implications (e.g. planar diffusion vs. pore diffusion) of number of relaxations (88). In the Nyquist plots the real part \((Z')\) is plotted on the x axis while the imaginary part \((Z'')\) on the y axis. The low frequencies resemble to slower processes arising on the metal interface while the higher frequencies resemble reactions that are kinetically controlled as well as to the charge transfer processes that is usually related to the coating interface.
By the application of sinusoidal AC potential centered at the open circuit potential ($E_{OCP}$) of the sample that causes it to cycle between the anodic and cathodic polarization as a result of a small AC current. Amplitude signal $\leq 10\text{mV}$ is applied (81) to the sample with the use of a potentiostat; the resulting current is analyzed to extract the phase and the amplitude relationship between the current and the voltage signals. Usually the impedance is measured as a function of frequency at a number of decades from 100 kHz to 1 mHz as an example (92). The low frequency choice is usually compromise between the resolution and the acquisition time, a common compromise between the time needed for data collection and accuracy is the choice of 5-10 points per decade and a low frequency limit of 10-1 mHz. In Bode plots, the frequency is an independent variable where an accurate compression can be applied between the experimental and the calculated impedance spectra. In Bode graphs with the same data the logarithm of the modulus of the impedance where $\log |Z|$ vs. $\log \omega$ is the Bode module plot and the phase angle ($\Theta$) vs. $\log \omega$ is the Bode phase plot (86) (Figure 3.8).

![Nyquist plot](image)

![Bode plots](image)

**Figure 3.8:** EIS data interpretations.
3.5 Adsorption on metal surface

Adsorption is known to be the adhesion of ions, atoms, molecules of dissolved solids, liquid or gas that are called adsorbate to create a film on a surface, which is the adsorbent (93). This happens when a self-assembled monolayer of molecules is formed on a solid surface (94). The adsorption process on a metal surface is usually visualized by plots that show the concentration dependent change of the surface coverage on a solid surface. These graphs known as adsorption isotherms have several types of isotherms such as Freundlich, Langmuir and the Stephen Brunauer, Paul Emmet and Edward Teller which is called the BET isotherm. There are several factors that can affect the process of adsorption such as (93):

- Surface area.
- Condition of experiments, eg, pressure temperature, pH, etc.
- Nature of adsorbent and adsorbate.
- Activation of adsorbent.

3.5.1 Types of adsorption:

There are two types of adsorption depending on the nature of the forces between the adsorbent and the adsorbate molecules. In the case of the physical adsorption (physisorption) the attraction forces are weak (like van der Waal’s and hydrogen forces). At these week forces the adsorption can be reversed according to the circumstances (pressure, concentration, and change in heating). The second type of the adsorption is the chemical adsorption (chemisorption) where the forces between the adsorbate and the adsorbent almost equal to the strength of the chemical bonds thus resulting in a very strong attraction; therefore the process cannot be reversed.

3.5.2 Adsorption Isotherms

The relationship between the equilibrium concentration of a substrate that is in contact with an adsorbent and the adsorbed adsorbate on a surface at a constant temperature is known as the adsorption isotherm (95).
vi. Langmuir adsorption isotherm

This type of adsorption describes quantitatively the formation of a monolayer on the solid surface and after that no further adsorption takes place. The Langmuir adsorption represents the equilibrium between the solid and liquid phase, which contains a finite number of identical sites. This type of adsorption isotherm that can predict chemisorption as well as physisorption of a monolayer were driven by Langmuir in 1916. The Langmuir derive was a relation between adsorbed material and its equilibrium concentration as Figure 3.9 (b) shows. Some main assumptions were proposed by Langmuir such as (93):

- There are many identical active sites on the surface known as adsorbing.
- Only one molecule can be adsorbed by each active site.
- No multilayer can be adsorbed on the surface but a monolayer.
- Constant heat of adsorption is on all sites of the surface.
- Interaction of the adsorbed molecules does not take place on the surface.
- The surface of the active sites is seamlessly as a flat plane containing no corrugations.

vii. BET adsorption isotherm

In this type of isotherm the adsorption on a single site does not affect the other sites the same as in the Langmuir theory. Adsorbed molecules can form multiple layers as well as the assumption is that the surface possess is uniform as shown in Figure 3.9(c).

viii. Freundlich adsorption isotherm

This type of adsorption describes the characteristics for a heterogeneous surface. There is an empirical relation between the quantities of the adsorbed adsorbate and the adsorbate concentration at equilibrium (Figure 3.9(d)) (93).
Figure 3.9: Different types of the isotherms; (a): linear (when the surface coverage linearly depends on the adsorbate concentration); (b) Langmuir type; (c) BET type; (d) Freundlich type (96).

There are some other types of adsorptions such as:

**ix. Temkin isotherm**

This isotherm contains a factor that explicitly takes into account the interaction between the adsorbate and the adsorbed molecules. This model assumes that the heat of adsorption of all molecules in the lazer would decrease linearly rather that logarithmic with the coverage.

**x. Dubinin-Radushkevich model**

The Dubinin-Radushkevich isotherm is generally applied to express the adsorption mechanisms with a Gaussian energy distribution onto a heterogeneous surface.
Chapter Four

4. Experimental Work

4.1 Materials and methods

4.1.1 Chemicals used for SAM layer formation

- **Fluorophosphonic acid**: $C_6F_{13}-C_2H_4-PO(OH)_2$ is a fluoroalkyl chemical with phosphonic acid functional group, Specific Polymers, Castries, France: SP-01-003; MW: 428.08; solubility: ethanol, methanol; pKa: first acidity: 2.08, second acidity: 7.63

- **Undecenyl phosphonic acid**: $CH_2=CH-[CH_2]_9-PO(OH)_2$, Product of: Specific Polymers, Castries, France: SP-61-003, MW: 234.27; solubility in: acetone, ethanol, THF; pKa: first acidity 2.42, second acidity: 8.08
• **Styren-co-styphos acid**

  ![Styren-co-styphos acid structure]

  $x=0.5; y=0.5; \text{MW: } 12250; \text{solubility: DMSO}$

  ![1H NMR (DMSO)]  
  ![31P NMR (DMSO)]

• **Dodecyl and hexadecyl phosphonic acid**: prepared and purified in the laboratory.

• **Sodium perchlorate** (NaClO₄)H₂O; M=140.46g/mol. Merck, D-6100 Darmstadt, F.R. Germany

• **Sodium chloride** (NaCl) M: 58.441g/mol, Reanal, Budapest, Hungary
- **Sodium hydroxide** (NaOH) Reanal, Budapest, Hungary.
- **Sulfuric acid** (H\textsubscript{2}SO\textsubscript{4}) 96%, Farmitalia Carloerba S.P.A, Milan
- **Methanol, ethanol, chloroform acetone**, Reanal, Budapest, Hungary
- **Water**: in all experiments ultra-clear water (Milli-Q water) was used (conductivity: 18MΩ cm\textsuperscript{-1} (5.5x10\textsuperscript{-6} S/m at 25 °C).

- All chemicals were used without further purification.

### 4.1.2 Metals of study

The chemical compositions of metals under investigation are the follows:

i. **Carbon steel composition**: C:2.29%; Fe:97.71%

ii. **Stainless steel 316**: C:0.08%; Mn:2.0%; Si:0.75%; P:0.045%; S::0.03%; Cr:18%; Mo: 3.00%; Ni: 14%; N: 0.1%

In the chromium-nickel stainless steel the molybdenum content increases the corrosion resistance, improves resistance to pitting in chloride ion solution.

iii. **Stainless steel 304**: C:0.08%; Mn:2.0%; Si:0.75%; P:0.045%; S::0.03%; Cr:20%; Ni: 10.5%; N: 0.1%

The significant difference between the two stainless steels is that the 304 type does not contain molybdenum.

iv. **Aluminum 5052**: Mg: 0.8%, Al:99.8%

### 4.1.3 Metal sample preparation

#### 4.1.3.1 Samples for contact angle measurements and for atomic force microscopy

Iron alloy samples in dimension of 10x11x2 mm were grinded using silicon carbide emery paper (grit sizes: 220, 400, 600, 800, 1200, 2200 and 4000). Afterwards, the samples were polished with diamond pastes of different grain sizes (15-9-6-3µ), washed with distilled water, degreased with acetone and dried at atmospheric conditions. In the case of aluminum to the last surface finishing Al\textsubscript{2}O\textsubscript{3} powder (grain size:3µ) was used.
4.1.3.2 Samples for electrochemical measurements

The metal samples of 14 mm in diameter were connected to a copper wire by welding, embedded into epoxy resin (Araldite AY 103 Resin and Hardener HY 956 Adhesive, Huntsman), then were grinded with silicon carbide emery paper of different grits (200, 400, 600, 800, to 1200 grit), washed with distilled water, acetone and dried in air.

4.1.3.3 Additional oxide layer formation

On aluminum alloy: after surface finishing by Al$_2$O$_3$ powder (grain size: 9 μm), the coupons were kept in boiling water for one hour, dried on air at room temperature and kept under normal atmosphere before SAM layer preparation.

4.1.4 Self-assembled molecular layer preparation

The solid phosphonic acid powders were dissolved in organic solvents (methanol, chloroform) in the concentration range of $5 \times 10^{-2}$ and $5 \times 10^{-3}$ M. Metal samples were immersed and left for predefined times ranging from 30 min, 2, 3, 4, 24, 48, 72 h in the phosphonic acid solutions. After the required contact time, the superfluous of the amphiphile solution was removed by letting to drop the solution and the last small remedy from the metal surface was removed by dipping the solids into the pure organic solvent. Finally the SAM-coated metal samples with self-assembled molecular layers were left to dry in air.

4.1.4.1 Modification of undecenyl phosphonic SAM

I have investigated the influence of irradiation and illumination on the CH$_2$=CH- groups in the undecenyl phosphonic acid SAM layers.

i. Irradiation by $^{60}$Co gamma source

Irradiation of undecenyl phosphonic acid SAM layers deposited on carbon steel was performed at room temperature in an SLL-01 type pilot scale facility equipped with $^{60}$Co gamma source
applying a dose rate of 11.5 kGy/h (equal to 11.5 kJ/kg,h) (Institute of Isotopes Co. Ltd., Budapest, Hungary). In one case 2kGy, in the other 20kGy was the absorbed dose.

ii. Illumination by UV light

The SAM layer built from undecenyl phosphonic acid, deposited onto carbon steel was illuminated by UV light (λ=254nm) for 30 and 60 min. In case of undecenyl phosphonic acid SAM layer developed on aluminum alloy surface longer (3 h) illumination was also used.

4.2 Methods of SAM layer characterization

4.2.1 Surface characterization by contact angle values

4.2.1.1 Static contact angle

Values were determined by a 2μL sessile drop measurement on a home-built device equipped with Hamilton micropipette and CCD camera. The contact angle was determined by geometrical evaluation of the spherical drop (97).

4.2.1.2 Dynamic contact angle measurement

A digital surface tensiometer (NIMA Ltd, Model DST 9005, UK) as shown in Figure 4.1, allowed the determination of dynamic contact angles based on the Wilhelmy theory. The metal samples with and without nanolayers were automatically dipped into the pure water (advancing contact angle) and pulled out from it (receding contact angle). The depth of immersion was 8 mm, the rate of the sample movement was 5 mm/min, and the apparent change in weight of the sample was registered (98). This was automatically converted to the advancing and receding contact angle values. In the static and dynamic contact angle measurements Milli Q water was used.
4.2.2 Atomic force microscopy

The surface visualization was performed by atomic force microscope (Digital Instruments, NanoScope III). The tip was Si$_3$N$_4$ (Veeco, Santa Barbara, CA, USA). The change in the morphology of the metal surface with and without nanolayers was imaged in contact mode, on air, before and after the corrosive attack. The section profile and the roughness of the surfaces were numerically characterized. Figure 4.2 shows the device used for these measurements.

4.2.3 Fourier transform infrared (FTIR) spectroscopy

Fourier transform infrared (FTIR) spectroscopy was used to identify structural changes of UPA starting material caused by UV treatment or irradiation. IR spectra were collected by the means
of a Varian 2000 (Scimitar Series) FTIR spectrometer equipped with an MCT (Mercury-Cadmium-Telluride) detector and fitted with a single reflection diamond ATR accessory (Specac Ltd, UK). No sample preparation was needed before spectrum collection. For each spectrum 64 individual spectra were averaged with a spectral resolution of 4 cm\(^{-1}\).

**4.2.4 Electrochemical techniques**

The experiments were carried out in a 0.1M NaClO\(_4\) electrolyte (saturated with air). The pH of the solutions was controlled by diluted NaOH solution to reach the alkalinity level and by H\(_2\)SO\(_4\) when the measurement goes at pH= 3.

**4.2.4.1 Electrochemical cell**

A three electrode cell (500ml) was used that consisted of a platinum sheet (surface area: 7.5cm\(^{2}\)) as an auxiliary or a counter electrode (CCE), a saturated calomel electrode (SCE) as a reference electrode (RE) and coated/non-coated metal surfaces as the working electrodes (WE) with a surface area: 1.54cm\(^{2}\).

**4.2.4.2 Open circuit potential (OCP) values vs time**

The potential change as a function of time was monitored during a time-period of 20 minutes using the Autolab PGSTAT 30 Autolab Instrument (Metrohm). Potentiostat / Galvanostat PGP201 VoltaLab (Figure 4.3).

![Potentiostat used in the experiments](image)
4.2.4.3 Potentiodynamic polarization measurement

The experiments were carried out in 0.1M NaClO₄ electrolyte (saturated with air). For controlling the pH values of the electrolytes, NaOH (pH=9 and pH=11) and H₂SO₄ (to pH=3) solutions were used. The measurement started after the settling of the open circuit potential (E\text{ocp}, 20min.). The measurements were performed with a sweep rate of 0.166mV/s (Autolab PGSTAT 30 Autolab Instrument (Metrohm)).

The anticorrosion efficiency of the surface layers was calculated by the following equation:

\[ \eta = (1 - \frac{j}{j_{\text{corr}}}) \times 100 \quad (4.1) \]

where:
- \( \eta \) = efficacy (%)
- \( j_{\text{corr}} \) = current density measured on the uncoated surface (mA/cm\(^2\))
- \( j \) = current density measured on the SAM-coated surface (mA/cm\(^2\))

The surface coverage (\( \Theta \)) is calculated from the efficiency data (99):

\[ \Theta = \frac{\eta}{100} \quad (4.2) \]

4.2.4.4 Electrochemical Impedance Spectroscopy (EIS)

The metal samples with and without SAM layer were placed in a teflon holder with an 1cm\(^2\) exposed metal surface. Impedance spectra were measured at the open circuit potential \([E_{\text{ocp}}]\) with 10 mHz signal amplitude in the frequency range of 10 kHz and 0.1 mHz with a 5 points per decade. The experiments, which began after the settling of the \( E_{\text{ocp}} \) of the working electrode, were carried out using a Solartron 1286 electrochemical interface and 1250 frequency response analyzer. The measurements were applied using the ZPlot software and the data fitting with the ZView software.

Figure 4.4 illustrates the electrochemical cell and the equivalent circuit for both of the electrochemical measurements applied in the study.
- All measurements were carried out at atmospheric pressure and room temperature.

Figure 4.4: Schematic of electrochemical cell (a) and the equivalent electric circuit of the working system (b).
Chapter Five

5. Results and discussion

Carbon steel, stainless steel types 304 and 316, and aluminum as bare metal samples were SAM-layer coated with different amphiphiles such as fluorophosphonic acid, undecenyl phosphonic acid, styrene-co-styphos acid, dodecyl- and hexadecyl phosphonic acids. Results and characterizations of the influences of the metal types, type of amphiphilic molecules, layer formation time, surface roughness, and irradiation/illumination on the SAM layer are illustrated and discussed in this chapter.

5.1 SAM layer formation

The self-assembly of surfactant-type molecules/amphiphiles in solution has been widely investigated both experimentally and theoretically, because of the numerous practical application possibilities of molecular layers.

In a self-assembling system, individual components (molecules and solids) interact in a predefined way and result in a spontaneous self-organization of the molecules into higher ordered structure. It is easy to obtain different surface properties when the tail groups of the amphiphilic molecules in the SAM layer are changed. For the SAM layer formation on metal surfaces covered by native metallic oxide layer (which is more-or-less hydrated) by alkyl phosphonic acids, an acid-base condensation mechanism in three steps is proposed (54). The first step is a hydrogen bond formation between the polar head group of phosphonic acid and the surface hydroxyl groups; this is followed by a reaction between the P-OH groups of phosphonic acid containing two hydroxyl groups by the surface via acid-base condensation when monodentate and bidentate bonds are formed. This reaction produces phosphonate salts and water. The formation of tridentate bond (which enables the phosphonic acids to form self-assembled monolayers on metal surfaces) is the consequence of hydrogen bond formation between the phosphoryl oxygen and the surface hydroxyl groups (P=O…H-O) (100-103). The hydrophobic tail of the phosphonic amphiphiles helps in formation of supramolecular SAM
structures (104, 105). The hydrogen bonding, the acid-base interaction as well as covalent bonding between the -P(O)(OH)$_2$ groups and the metal oxide layer can attribute to the formation of a densely packed, and well organized organic SAM film on the metal surface (106).

It is essential to characterize the SAM-layer coated metal surfaces. The compactness of the SAM layer can determine its anticorrosion efficiency. My work focused on those SAM layers that were supposed to significantly decrease the corrosion. The SAM layers of phosphonic acids were developed on the native metal oxide surface using a simple room-temperature solution deposition method. As the head group faces the metal surface, the hydrophobic molecular part is perpendicular to it. Later, when already a total surface is covered by the amphiphiles, generally the molecules are not in perpendicular position, but tilted to the surface. A dense hydrophobic layer can well control the access of water (with or without dissolved gases and ions) to the metal surface, i.e. the possibility of corrosion. The measurement of the water wettability is by the surface contact angle. I have prepared SAM layers that could significantly decrease the surface wettability i.e. increase the hydrophobic character of the surface coating and, as a consequence, the anticorrosion activity. The results of the changes in the parameters of the SAM formation (amphiphile concentration, layer formation time, metals, surface roughness, pH values) are demonstrated in the next part of the thesis by different characterization techniques of SAM-layer coated metal surfaces.

5.2. SAM layer characterization

5.2.1 Characterization by contact angle measurements

One of the most obvious surface characteristics that can be affected by the environment is the wetting properties. The wettability of solid surfaces by liquid materials is an important aspect of materials science and surface chemistry that plays an important role in the corrosion coatings. In wetting studies, usually the contact angle is one of the most important data that show the wetting values when a liquid and a solid are in contact. The wettability is summarized in the next Table 5.1.
Table 5.1: Wettability in the mirror of the contact angles (79).

<table>
<thead>
<tr>
<th>Contact angle</th>
<th>Degree of wetting</th>
<th>Strength of solid/liquid interaction</th>
<th>Strength of liquid/liquid interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>(S) ( \theta = 0 )</td>
<td>perfect wetting</td>
<td>strong</td>
<td>weak</td>
</tr>
<tr>
<td>(C) ( 0 &lt; \theta &lt; 90^\circ )</td>
<td>high wettability</td>
<td>strong</td>
<td>strong</td>
</tr>
<tr>
<td>(B) ( 90^\circ \leq \theta &lt; 180^\circ )</td>
<td>low wettability</td>
<td>weak</td>
<td>strong</td>
</tr>
<tr>
<td>(A) ( \theta = 180^\circ )</td>
<td>perfect non-wetting</td>
<td>weak</td>
<td>strong</td>
</tr>
</tbody>
</table>

In my experiments, the liquid was water as the corrosion starts always in the presence of water. The metal surfaces with or without SAM layers were analyzed and characterized by dynamic and static contact angle values.

### 5.2.1.2 Surface characterization by dynamic contact angle values

The dynamic contact angle measurement allows measuring of two different angles: the advancing contact angle (when the solid sample enters the water) and the receding contact angle (when the solid sample exits the water). Almost in all cases, there are differences between the two values. The magnitude of the difference indicates that the compactness of the layer is not dense enough to produce a superhydrophobic nanofilm; in other words, the nanolayer does not cover the solid surface perfectly. A water contact angle higher than 100° indicates the formation of an ordered film, which is more hydrophobic than the control sample. According to the literature (49) the phosphonic groups are bond to the metal surface, covered by oxide layer, in a bi- and tridentate mode, determined by diffuse reflectance FTIR spectroscopy.

Lower average advancing contact angle represents SAM layers when the amphiphile has a shorter carbon chain / smaller hydrophobic part or when in the amphiphilic carbon chain there are substituent or double bonds. Not only the inhomogeneity of the molecular layer, but also the surface roughness can influence the change in the advancing and receding contact angle values.
5.2.1.2.1 Wettability change caused by SAM layers on different metal surfaces

i. Influence of SAM layers on carbon steel surfaces

a. Effect of the amphiphiles and their concentration

In the first set of contact angle experiments the question to answer was how do the phosphonic acid amphiphiles with different hydrophobic parts influence the surface wettability, i.e. under identical condition, the layers formed on the metal surface (covered by native oxide layer) can increase the surface hydrophobicity with the same rate or not. Dynamic contact angle values measured on carbon steel covered by different phosphonic acid SAM layers, formed in 24 h are shown in Figure 5.1. First the behavior of the fluorinated alkyl phosphonic acid SAM layer, developed on carbon steel is discussed. Fluorophosphonic acid layer achieved the highest contact angle values: 129° / 80° (advancing and receding angles, respectively). These values prove the formation of a compact layer on the metal surface and the influence of the superhydrophobic fluorine atoms; both factors result in reduced water wettability. Surfactants, like amphiphiles, are used for lowering the surface energy, and the decrease in the surface energy results in formation of superhydrophobic surfaces. Fluorine atoms in the alkyl chain have a great impact on reducing the surface energy, the presence of CF$_3$ and CF$_2$ groups assign high hydrophobicity to the fluorophosphonic acid SAM layer.

Figure 5.1: Influence of hydrophobic molecular part in SAM layers on the wettability of coated carbon steel (layer formation: 24 h; U.P:undecenyl phosphonic acid, Sty: styren-co-styphos acid; F.P: fluorophosphonic acid).
The hydrophobicity of the SAM layers formed by the second amphiphilic molecules, the **undecenyl phosphonic acid**, could not reach the level of the fluorophosphonic amphiphile. It is also evident that the highest difference between the advancing and receding contact angles is in the case of the undecenyl phosphonic acid SAM layer. The explanation is clear: in this case the nanolayer is less compact than at the other two amphiphiles.

The third chemical is the **styrene-co-styphos acid**. As its structure suggests, the anchoring phosphonic groups fix the molecules to the metal surface and, additionally, a horizontal hydrophobic net is formed from the polymeric chain. This is reflected in the higher advancing as well as in receding contact angle values.

The influence of the concentration of the amphiphile solutions is demonstrated in Figure 5.2. It is evident that in the case of the fluorophosphonic acid already a less concentrated amphiphile solution is enough to reach low water wettability, which is almost equivalent with that value measured in case at its more concentrated solution (Figure 5.2).

![Figure 5.2: Influence of concentration on the wettability of carbon steel with fluorophosphonic acid and undecenyl phosphonic acid SAM layers formed at 24 h (5E-3=5x10^{-3} M; 5E-2=5x10^{-2} M).](image)
On the other hand, the lower concentration of the undecenyl phosphonic acid does not allow the formation of a compact layer. It is reflected in the change in the receding contact angle values: at lower concentration this is much lower than measured in the case of the SAM layer formed from the more concentrated solution. Though the increase in the amphiphile concentration resulted in a better surface coverage, i.e. SAM layer formed in more concentrated solution shows higher receding contact angle value. The consequence is that SAM layers formed from more concentrated solution ($5 \times 10^{-2}$ M) resulted in better surface coverage (Figure 5.2).

b. Effect of the SAM layer formation time

In the case of the fluorophosphonic acid the layer formation time has less important impact on the increase in the contact angle than in case of the undecenyl phosphonic acid; i.e. in a short time the layer formed from the fluoro amphiphilic molecule shows a real well ordered structure, compactness.

Evaluating the influence of the film formation time in case of undecenyl phosphonic acid SAM layer, there was an increase in the advancing contact angle values: after 4 h the contact angle was $83^\circ$ that increased to $107^\circ$ when more time (24 h) was left for the self-assembling process. It shows that in a shorter time less amphiphilic molecules can absorb onto the carbon steel, and domains of disordered molecules are present. With the increase in the layer formation time, more phosphonic groups attach to the solid, thus less free space is for the hydrophobic molecular parts, which results in an increased interaction among the hydrophobic chains. At the end of long SAM formation time a more hydrophobic surface is formed on the carbon steel surface. Even in this case, the high advancing contact angle ($107^\circ$) was followed by low receding angle ($15^\circ$). This is the consequence of the double bond at the end of the alkenyl chain, which does not allow the formation of a densely packed surface film. It is clear that the phosphonic head groups interact with the metal surface through oxygen bridges anchoring the molecules to the solid surface. On the other hand, the alkyl part of the alkenyl chain that is nearer to the solid, can keep together this hydrophobic part of the chain through different forces (hydrogen bond, van der Waals force), but the double bonds at the end of the carbon chain does not allow a firm association. As a result, the water molecules can approach the metal surface easier.
Table 5.2 illustrates the influence of the immersion cycles during the dynamic contact angle measurement. When a self-assembled molecular layer of undecenyl phosphonic acid covers the carbon steel surface (formed in 24h), at the first dip high contact angle is observed (advancing: 107°). However, after the second, third and fourth dip this value decreases drastically (advancing: 60°), which immediately indicates that the layer is not compact, the alkenyl chain cannot form a densely packed surface layer (due to the double bond). The low receding contact angle value (15°) shows the same. When the amphiphile is the fluorophosphonic acid (SAM layer formed in 24h), the formation of a molecular layer with well-defined, compact structure is mirrored in the contact angle values; the consequence is that the contact angle measured at the first dip (advancing: 129°) have only slight decrease after the second, third and fourth immersion cycle (advancing: 125°). It proves unequivocally that the fluorophosphonic acid nanolayer has a close-packed structure. The same is reflected in the high receding contact angle value: 80°. This indicates that the monolayer is stable in neutral water. Generally, the magnitude of the contact angle hysteresis is correlated with the monolayer packing densities. The high contact angle values suggest that the hydrophobic barrier effect originates from the well-structured hydrophobic molecular part.

Table 5.2: Contact angle values measured on carbon steel surfaces covered by SAM layers formed by different chemicals; influence of the layer formation time, concentrations and dipping number.

<table>
<thead>
<tr>
<th>Metal</th>
<th>Chemical</th>
<th>Time</th>
<th>C. A (Adv, Rec)</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>carbon steel</td>
<td>Undecyl phosphonic acid(5x10⁻³ M)</td>
<td>4 h</td>
<td>83, 45</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>24 h</td>
<td>107, 15 (1st dip) 60, 15 (4 other dips)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Undecyl phosphonic acid(5x10⁻³ M)</td>
<td>24 h</td>
<td>108, 52(1st dip) 105, 45 (4 other dips)</td>
<td>Exposed to UV for 30min each side</td>
</tr>
<tr>
<td></td>
<td>Undecyl phosphonic acid(5x10⁻² M)</td>
<td>24 h</td>
<td>110,45</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Styrene co Styphos acid(5x10⁻² M)</td>
<td>24 h</td>
<td>115, 70</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Fluorophosphonic acid(5x10⁻³ M)</td>
<td>24 h</td>
<td>129, 80(1st dip) 125, 80 (4 other dips)</td>
<td></td>
</tr>
</tbody>
</table>
ii. **Influence of SAM coatings on stainless steel surfaces**

The stainless steels are important alloys, their resistance against corrosion determines their usage. Under special conditions (e.g. in body fluid) the release of the iron, chromium, nickel and any other alloying metals can generate allergic and carcinogenic effect. Therefore, the surface properties of stainless steels determine their application conditions. This explains why it is necessary to treat properly their surfaces that lead to enhanced applicability without losing the physical and mechanical properties.

The SAM layers of the phosphonic acid amphiphiles used in my experiments were deposited onto stainless steel surfaces by room-temperature solution deposition.

*a. Stainless steel 304*

Results achieved on SAM coated **304 stainless steel** (Figure 5.3) show that the highest contact angle value was achieved with the fluorophosphonic acid SAM layer formed at 24h (advancing angle: 127°). The high contact angle value can be related to the dense oxide layer that covered the stainless steel surface. It is interesting that there was almost no change in the contact angle values when the concentration increased from $5 \times 10^{-3}$ M to $5 \times 10^{-2}$ M. This shows that a well-defined, densely packed SAM layer is formed already in the less concentrated amphiphile solution. For comparison, alkyl phosphonic acids with shorter (dodecyl: C12) and longer (hexadecyl: C16) carbon chains were investigated. In the case of these two chemicals, I also could follow the influence of the layer formation time on the change in the surface hydrophobicity. When the SAM layer was developed from dodecyl phosphonic acid solution, the layer formation time affected the formation of a dense layer. The contact angle values increased from 95° achieved at 15min to 118° measured at 24 h (Table 5.3), which indicates that longer time was needed for the formation of a more condensed, compact layer in the case of the amphiphile with shorter carbon chain. It is interesting that the longer alkyl chain does not create a more hydrophobic surface: the layer wettabiliy is almost the same as measured on C12 layer. Nevertheless, the influence of the longer carbon chain is reflected in the higher receding contact angel (proving a more compact state of the nanolayer). In both cases (C12 and C16) increasing
the layer formation time as well as the concentration of the amphiphiles more hydrophobic nanofilms were produced on the metal surface; I noticed similar effect in the case of the carbon steel covered by the same SAM films.

It is interesting that the dynamic contact angle values measured on stainless steel 304 with SAM layers of styrene-co-styphos acid, fluorophosphonic acid and undecenyl phosphonic acid are similar to those measured on carbon steel. The highest hydrophobicity – superhydrophobicity - was observed on the fluorophosphonic SAM layer; the increase in the concentration during the film formation resulted in a slightly enhanced contact angle. On the other hand, the higher concentration eventuates in smaller hysteresis, i.e. the receding contact angle value is higher; the layer is more compact (Figure 5.4). In case of the undecenyl phosphonic acid SAM layer the nanofilm shows denser structure as the receding contact angle is much higher than was on carbon steel.

![Figure 5.3: Influence of amphiphile types of SAMs on stainless steel 304 (layer formation time: 24h; The surfaces are covered by SAM of: C12P: dodecyl phosphonic acid; C16P: hexadecyl phosphonic acid; U.P:undecenyl phosphonic acid, Sty: styren-co-styphos acid; F.P: fluorophosphonic acid ).](image-url)
Table 5.3: Contact angle values measured on stainless steel 304 surfaces covered by SAM layers; influence of the chemicals, layer formation time, concentrations and dipping time.

<table>
<thead>
<tr>
<th>Metal</th>
<th>Chemical</th>
<th>Time</th>
<th>C. A. (Adv, Rec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>stainless steel 304</td>
<td>dodecyl phosphonic acid</td>
<td>15 min</td>
<td>95, 60</td>
</tr>
<tr>
<td></td>
<td></td>
<td>24 h</td>
<td>118, 65(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>112, 65(4 other dips)</td>
</tr>
<tr>
<td></td>
<td>hexadecyl phosphonic acid(5x10^-3 M)</td>
<td>15 min</td>
<td>103, 55</td>
</tr>
<tr>
<td></td>
<td></td>
<td>97, 60</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>4 h</td>
<td>100, 72</td>
</tr>
<tr>
<td></td>
<td></td>
<td>24 h</td>
<td>107, 75</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>100, 55</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5 days</td>
<td>120, 80</td>
</tr>
<tr>
<td></td>
<td></td>
<td>7 days</td>
<td>95, 45</td>
</tr>
<tr>
<td></td>
<td>hexadecyl phosphonic acid(8x10^-3 M)</td>
<td>4 h</td>
<td>103, 46 (1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>95, 46 (4 other dips)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>24 h</td>
<td>103, 45 (1st dip)</td>
</tr>
<tr>
<td></td>
<td>Undecyl phosphonic acid (5x10^-3 M)</td>
<td>4 h</td>
<td>115, 45(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>85, 45 (others 4 dips)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>24 h</td>
<td>115, 80</td>
</tr>
<tr>
<td></td>
<td>Styrene-co-styphos acid (5x10^-2 M)</td>
<td>24 h</td>
<td>110, 60</td>
</tr>
<tr>
<td></td>
<td>Fluorophosphonic acid (5x10^-2 M)</td>
<td>24 h</td>
<td>127, 70(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>123,70 (others 4 dips)</td>
</tr>
<tr>
<td></td>
<td>Fluorophosphonic acid (5x10^-3 M)</td>
<td>24 h</td>
<td>128, 60(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>120, 58(2nd dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>117, 57 (others 3 dips)</td>
</tr>
</tbody>
</table>
Figure 5.4: Influence of the concentration on the wettability of stainless steel 304 with fluorophosphonic acid (F.P) and undecenyl phosphonic acid (U.P) SAM layers formed at 24h (5E-3=5x10^{-3} M; 5E-2=5x10^{-2} M).

### b. Stainless steel 316

The effect of layer formation time in cases of undeceyl phosphonic acid, styrene-co-styphos acid and fluorophosphonic acid on **stainless steel 316** samples is illustrated in Figure 5.5 and data are depicted in Table 5.4. One can notice that the highest advancing contact angle of 118° was again achieved with the fluorophosphonic acid. The undecenyl phosphonic acid and the styrene-co-styphos acid solution after 24 h immersion gave almost the same contact angles. It is important to mention that on the stainless steel 316 the receding values are lower; the differences between the advancing and receding contact angle values are higher than in the case of the stainless steel 304. This could be explained by the differences in the surface oxide layer. In the case of stainless steel 316 the interaction between the molybdenum oxide that partly covers the metal surface and the phosphonic groups is less powerful than on the chromium oxide and iron oxide surface (e.g. in case of 304 oxide layer).
iii Influence of SAM coatings on aluminum surfaces

In this part I present the contact angle values measurements on aluminum surfaces covered by phosphonic acid SAM layers and the numerical data are discussed. It is well known that the aluminum alloys are often used in different industries like aircraft, automobiles, construction because of their strength to weight ratio, ductility and low cost. The wide application is due to the corrosion resistance caused by the easily formed passive oxide layer on the aluminum.
The surface hydrophobicity increases when the SAM films of alkenyl and fluorophosphonic acids cover the metal as these molecular layers act as barrier to the aqueous solutions and the anticorrosion activity improves. The phosphonic amphiphiles form the SAMs by condensation reactions of the acid functional group (that show strong affinity to the aluminum) with the basic surface bound aluminohydroxil species via Al-O-P bond:

\[
R\cdot\text{PO(OH)}_2 + \text{Al-OH} \rightarrow R\cdot\text{OH} \cdot \text{OP-O-Al} + \text{H}_2\text{O} \quad (5.1)
\]

To understand the anticorrosion protection of the SAM layers on the aluminum surface it is important to characterize the layers. The next Table 5.5 summarizes the results of the dynamic contact angle measurements. The high contact angle values show that the phosphonic groups adsorb very well on the metal surface, and reduce the metal solubility in the aqueous media with decreasing the active spots on the surface.

The fluorophosphonic acid resulted in the highest contact angle values (168° and 95°) (Figure 5.5) among all three chemicals used. Although after 48 h immersion time the results of styrene-co-styphos acid and fluorophosphonic acid were almost the same, in the case of the styrene-co-styphos acid there was a decrease in the contact angle values as the number of dips increased. This indicates that fast surface adsorption of amphiphiles occurs within a short time; however, well-oriented high-quality SAMs are formed after longer period (some hours). The nanolayer formed in short period could have some inhomogeneity and in this case, the water molecules could approach the metal surface. There was almost no change in the contact angle values with time-increase for both the undecenyl phosphonic acid and fluorophosphonic acid whereas for styrene-co-styphos acid the value increased from 150° at 2 h to 165° at 48 h. Nevertheless, one has to keep in mind that the fluorophosphono amphiphile is the only one where the repeated dipping into the water did not cause significant decrease in the contact angle; in other words, this is the most compact layer comparing with the other two SAMs. It is important to mention that the increase in the layer formation time and in the amphiphile concentration influenced the contact angle values in the same direction as in the case of the other alloys i.e. the wettability in water decreased. In order to see the influence of the pure solvents used for developing the SAM layers, the metals were immersed into methanol and chloroform, and the contact angles were followed. As contact angle values proved, the organic solvents did not cause change in the wettability of the bare metal; the advancing angle was 63°.
Table 5.5: Contact angle values measured on SAMs covered aluminum surfaces; influence of the amphiphiles, layer formation time, and concentrations

<table>
<thead>
<tr>
<th>Metal</th>
<th>Chemical</th>
<th>Time</th>
<th>C. A (Adv, Rec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aluminum</td>
<td>Undecyl phosphonic acid(5x10^-3 M)</td>
<td>24 h</td>
<td>138, 20(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>45, 20(2nd dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>40, 15(other 3 dips)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>48 h</td>
<td>145, 45(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>85, 45(2nd dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>70, 45(3rd dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>63, 45(4th dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>50, 45(5th dip)</td>
</tr>
<tr>
<td></td>
<td>Styrene-co-styphos acid(5x10^-3 M)</td>
<td>2 h</td>
<td>150, 38(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>55, 38(other dips)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4 h</td>
<td>170, 45(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>60, 45(other 4 dips)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>24 h</td>
<td>170, 45(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>60, 45(other dips)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>48 h</td>
<td>165, 50(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>100, 50(2nd dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>95-80,50(other 3 dips)</td>
</tr>
<tr>
<td></td>
<td>Styrene-co-styphos acid (5x10^-2 M)</td>
<td>24 h</td>
<td>160, 40(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>60, 40(2nd dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>48, 40(other 3 dips)</td>
</tr>
<tr>
<td></td>
<td>Fluorophosphonic acid(5x10^-3 M)</td>
<td>2 h</td>
<td>162, 80(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>140, 60(2nd dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>132, 55(3rd dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>125, 50(4th dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>120, 50(5th dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4 h</td>
<td>175, 65(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>130, 57(2nd dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>118, 53(3rd dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>110, 50(4th dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>103, 50(5th dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>48 h</td>
<td>168, 95</td>
</tr>
<tr>
<td></td>
<td>Fluorophosphonic acid(5x10^-2 M)</td>
<td>48 h</td>
<td>175, 135(1st dip)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>170, 130(other 4dips)</td>
</tr>
</tbody>
</table>
It is important to mention that on aluminum the increase in the layer formation time and in the amphiphile concentration influenced the contact angle in the same way as in the case of the other iron alloys. For the sake of the comparison, I show the wettability changes on all SAM-covered metals that were investigated (carbon steel, stainless steels 304 and 316, aluminum). The most hydrophobic, so-called superhydrophobic surfaces were developed on aluminum surface by these phosphonic acid amphiphiles (Figure 5.7 and 5.8). These figures confirm the statement by showing the contact angle values of nanolayers formed from fluorophosphonic and undecenyl phosphonic acids. On all metals both amphiphiles form very hydrophobic layers and it is important to mention that on the stainless steel 304 developed compact, dense layer, which is reflected in high receding values. The most hydrophobic – superhydrophobic - SAM layer is formed from the fluorophosphonic acid on aluminum.
Figure 5.7: Comparizon of wettabilities measured on different metal surfaces covered by fluorophosphonic acid SAM layer formed in 24 h (amphiphile concentration: $5 \times 10^{-3}$ M; St.St. = stainless steel).

Figure 5.8: Wettability measured on different metal surfaces covered by undecenyl phosphonic acid SAM layer formed in 24 h (amphiphile concentration: $5 \times 10^{-3}$ M; St.St. = stainless steel).
As for aluminum samples, the contact angle value was higher than those on carbon steel due to the hard aluminum oxide that formed on the surface where the chemical interacted with the oxide layer on the metal surface to form a protective layer.

From the achieved results, it can be observed that the fluorophosphonic acid forms a more hydrophobic coating than the undecenyl phosphonic acid, which is due to the structure of the molecules involved into the self-assembling process.

5.2.1.1.2. Influence of the surface roughness on the SAM layer formation

In the literature, there are information about the surface roughness and the bias toward to corrosion. The roughness of a surface can determine the characteristics of SAM nanolayers. With increasing roughness the surface area increases, more amphiphilic molecules will be absorbed through adhesion of head groups.

As previously I had mentioned, the surface free energy (influenced by the surface chemistry), surface roughness and its structure/homogeneity can control the surface wettability as they can trap air on the surface and stabilize the Cassie-Baxter state. When the metal surface is flat, and, additionally, covered by nanolayers, the surface energy is reduced (107, 108). In my experiments, I have investigated the influence of the surface finishing (roughness) on the change in the dynamic water contact angle. Figure 5.9 summarizes what happens when the metal surface, before the SAM layer deposition, was grinded with a series of emery paper (started from 400 up to 4000-grain size). Then on the bare metal of different roughness SAM layers of the two amphiphiles (fluorophosphonic and undecenyl phosphonic acid) were deposited. The increase in the contact angle values with increasing emery paper grain sizes is well demonstrated in Figure 5.9. It is interesting that the fluorophosphonic amphiphile (which generally produces more hydrophobic surface than the undecenyl one) cannot deposit a more hydrophobic SAM layer of high contact angle on the roughest surface, contrary to the undecenyl phosphonic acid.

As the bare metal surface is more and more smother, the tendency changes and, at last, on the smoothest surface the SAM nanolayer of the fluorophosphonic amphiphile is more hydrophobic.
than the undecenyl one. The structure of the undecenyl phosphonic acid assists to this behavior, as the double bond at the end of the carbon chain helps to disguise air bubbles. The hydrophobic character of the bare metal disappears after the second dip, in the case of the undecenyl phosphono amphiphile layer it deceases, the wetting ability of the fluorophosphonic layer does not change; it stays very hydrophobic as the results of dynamic contact angles demonstrate.

![Figure 5.9: Influence of the aluminum surface finishing on the changes in the contact angle values (concentration: 5x10^3 M; Und phosphonic acid= undecenyl phosphonic acid).](image)

5.2.1.3. Influence of the SAM layer curing by UV light on the wettability

The undecenyl phosphonic acid with a double bound in the alkenyl chain is capable for polymerization. Its SAM layer showed less ordered structure then the other amphiphiles due to the destruction of the ordering by the CH\(_2\)=CH- group at the end of the alkenyl chain. The idea to polymerize these molecules to form a “hydrophobic net” over the metal surface (to increase the anticorrosion activity) was achieved by two techniques.
After exposing the carbon steel metal sample coated by an undecenyl phosphonic acid layer to a UV light ($\lambda = 254$ nm) for 30 minutes, the advancing angle value was 108° for the first dip and 105° for the other dips. This indicates that the layer became more compact due to the influence of the UV light that led to the polymerization of the double bounds (Table 5.2).

UV exposition of undecenyl phosphonic acid layers on aluminum showed no change for samples that were immersed for 2hrs into the amphiphile solution and exposed to UV light for 30 minutes. While those samples when the exposure time was longer (3hrs) on samples when the SAM layer formation time was 2 and 48hrs, the contact angle values decrease which indicate that the layer became looser and the water molecules could penetrate to the metal surface.

It is clear when the contact angle value without illumination was already high; the layer was already compact enough (due to the dense oxide surface layer which allows adsorption of much more phosphonic acid molecules notwithstanding the presence of the interfering double bonds. In the next experiments the change in the wettability will be analyzed by measuring the static contact angles.

### 5.2.1.3 Surface characterization by static contact angle values

A liquid drop can spread across a surface because of the adhesive forces between the liquid and the solid. When the cohesive force is high in the liquid, the drop avoids the contact with the surface. When a contact angle is less than 90° (case:C), the surface wetting is favorable; the water will spread over the surface area. At above 90° (case:A), the surface wetting is unfavorable, a compact liquid drop will be formed (Figure 5.10).

![Figure 5.10:Wetting properties by different fluids](image)

$\theta = 0; C: 0 < \theta < 90^\circ; B: 90^\circ \leq \theta < 180^\circ; A: \theta = 180^\circ$
5.2.1.2.1. Influence of the SAM layer curing by UV light

The wetting of the carbon steel surface (coated and uncoated with fluorophosphonic or undecenyl phosphonic acid layers) was investigated in water. Static contact angle results are illustrated in Figures 5.11 and in Table 5.6. In case of measuring the static contact angle, small drop of water is placed on the solid surface (80).

In the first set of experiments carbon steel surface was investigated in uncoated and coated form with undecenyl phosphonic acid SAM layer. For the sake of comparison result got by dynamic and static contact angle measurements, the fluorophosphonic acid SAM layer was also analyzed. The static contact angle values are summarized in Table 5.6. As long as the fluoro amphiphile in SAM nanolayer gave a high value (124°) the undecenyl phosphonic acid in nanofilm produced a less value of 88°.

In order to polymerize the double bond in the SAM layer formed by undecenyl phosphonic acid, two methods were chosen to carry out these experiments and the influence was measured with the use of the static contact angle since only one side of the surface could be treated while for the dynamic contact angle measurements both sides needed treatment.

In order to investigate the effect of curing of the double bond in the undecenyl phosphonic acid SAM layer on the carbon steel sample formed in 24hrs, it was undertaken to UV light (λ = 254 nm) for 30 and 60 minutes. The influence of the shorter and longer illumination time, which was followed by contact angle measurement showed that the shorter time has a much less important effect on the layer hydrophobicity (contact angle of SAM without illumination: 88°; after illumination for 30 min: 81°). With other words, this time was enough for polymerization of the minority of the double bounds. On the other hand, 60 min UV light illumination resulted in an increased contact angle value: from 88° to 102°. This indicates that the layer became more compact due to the influence of the UV light that led to the polymerization of the majority of the double bounds (Table 5.6).
5.2.1.2.2. Influence of irradiation by $^{60}$Co gamma source

The other possibility for the polymerization of a double bond in an alkenyl chain is the application of gamma irradiation. In one case 2kGy and in the other 20kGy was absorbed by the undecenyl phosphonic acid SAM layer deposited onto the carbon steel surface. The surface film handled with 2kGy did not really change the contact angle: on the uncured SAM it was $88\pm4^\circ$ that of the cured sample with 2kGy was $85\pm2^\circ$. However, the absorption of 20kGy increased the contact angle up to $118^\circ$. This indicates that the irradiation by 2kGy did not affected the great part of the double bonds in the SAM layer, but the 20kGy absorption interacted with almost all double bonds in the nanofilm, which rearranged the SAM layer surface that became more compact and formed a real barrier between the metal surface and the aqueous environment (Figure 5.10, Table 5.6).

Table 5.6: Static contact angle values measured on carbon steel samples with or without coatings, before and after curing.

<table>
<thead>
<tr>
<th>Chemical</th>
<th>Treatment</th>
<th>Contact angle [$^\circ$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bare metal</td>
<td>-</td>
<td>68</td>
</tr>
<tr>
<td>SAM of fluorophosphonic acid (24 h)</td>
<td>-</td>
<td>124</td>
</tr>
<tr>
<td>SAM of undecenyl phosphonic acid (4 h)</td>
<td>-</td>
<td>86</td>
</tr>
<tr>
<td>SAM of undecenyl phosphonic acid (24 h)</td>
<td>-</td>
<td>88</td>
</tr>
<tr>
<td>SAM of undecenyl phosphonic acid (48 h)</td>
<td>-</td>
<td>85</td>
</tr>
<tr>
<td>SAM of undecenyl phosphonic acid (24 h)</td>
<td>2kGy</td>
<td>85</td>
</tr>
<tr>
<td>SAM of undecenyl phosphonic acid (24 h)</td>
<td>20kGy</td>
<td>118</td>
</tr>
<tr>
<td>SAM of undecenyl phosphonic acid (24 h)</td>
<td>30min. UV</td>
<td>81</td>
</tr>
<tr>
<td>SAM of undecenyl phosphonic acid (24 h)</td>
<td>60min UV</td>
<td>102</td>
</tr>
</tbody>
</table>
Fig. 5.11: Static contact angle values measured on cured and not cured SAM layers deposited onto carbon steel samples.

The comparison of the dynamic and static contact angle values measured on similar surfaces in case of fluorophosphonic acid SAM layer gave almost the same value (129° and 124° respectively), but the undecenyl phosphonic acid nanofilm behaved differently which could be due to the slightly different roughness of the metal surface.

5.3 Surface visualization by atomic force microscopy (AFM)

The SAM coated surfaces were characterized by AFM, too. This technique allows the surface visualization in 3D as well as the numerical characterization of the surface (measure of roughness, section analysis).
5.3.1 Influence of short SAM layer formation time on the surface morphology and on the anticorrosion activity

Carbon steel samples, with and without coatings, were visualized by AFM. The images on the surface morphology showed that both chemicals (fluorophosphonic and undecenyl phosphonic acids) formed compact layers on the metal surface. By this technique not only the SAM layer covered metal surfaces are characterized, but the anticorrosion activity of these layers, too.

In the first set of AFM experiments, I was interested not only in the surface morphology of the SAM layer covered metal surfaces but in the influence of the layer formation time on the anticorrosion activity. In Figures 5.12 and 5.13 the AFM images visualize the carbon steel surface without and with SAM layers built in 30 minutes, handled with different (NaCl, NaClO₄) electrolytes. The carbon steel shows a smooth, partly irregular metal surface, which, after nanolayer deposition has already more regular patterns on the surface. After immersion of SAM coated metal into perchlorate, small depositions appear on the surface showing the formation of corrosion products in the aerated electrolyte. The sodium chloride deteriorated the surface; the formation of deep pits is visible, which is clearly demonstrated on the 3D images. The comparison of the surface roughness of the bare metal and of the nanocoated surface after immersion into NaCl assures that the layer is much less rough when the nanocoating is present than without it.

![Carbon steel on air](image1.png) ![Carbon steel immersed in NaCl for 1 h](image2.png)

Figure 5.12: Carbon steel surface before and after immersion into sodium chloride.
Figure 5.13: Influence electrolytes on SAM layers on carbon steel (layer formation time: 30 min).

The last two images in Figure 5.13 show that even in the case of short coating time the fluorophosphonic acid SAM layer can inhibit the pitting corrosion. There is not any pit on the metal surface after 1 h long immersion into sodium chloride.
5.3.2 Influence of longer SAM layer formation time and irradiation on the surface morphology of different metals

The results presented here show the influence of the longer nanolayer formation time (24 h) on the surface morphology, without and with curing the undecenyl phosphonic acid SAM layer, and the influence of the two electrolytes. On Figure 5.14 the change caused by longer nanolayer deposition is visualized in 3D as well as in section images. The surface of the fluoro amphiphile-covered metal surface is similar either 30 minutes or 24 h was left for layer deposition. It was interesting that the undecenyl phosphonic acid after adsorption onto the metal surface made a special surface appearance, which differs from that one shown after 30 min deposition. Previous XPS results confirmed the possibility of multilayer appearance on the surface.

Figure 5.14: SAM layers of fluorophosphonic acid (a) and undecenyl phosphonic acid (b) and its section (c) on carbon steel surfaces (layer formation time: 24 h).
Images in Figure 5.15 and 5.16 show the influence of the irradiation on the undecenyl SAM layers.

Figure 5.15: Undecenyl phosphonic acid SAM layer (layer formation time: 24 h) after irradiation: absorption of 2 kGy; carbon steel.

Figure 5.16: Undecenyl phosphonic acid SAM layer (layer formation time: 24 hrs) after irradiation: absorption 20kGy; carbon steel.
Figure 5.17: Oxide layer covered aluminum surface (a, b) and fluorophosphonic acid SAM layer covered aluminum surface (c, d) visualized in 3D and by section analysis.

The 3D images and the section analysis proved that the presence of the molecular layers changes the surface morphology. It is visible that these amphiphiles develop special coverage on the metals. On the other hand, the irradiation left “special signs” on the SAM layers. Even in the case of 2 kGy absorption, the surface morphology did not significantly changed, however, the 20 kGy altered the surface. This drastic change appeared already in the contact angle values.

The same SAM-covered samples were further investigated to study the influence of the SAM layers on the general and pitting corrosion.
5.3.3 Influence of SAM layers on pitting corrosion

The influence of sodium chloride solution on carbon steel surfaces covered by nanolayers was investigated. After one hour of immersion in sodium chloride, the carbon steel surface coated with fluorophosphonic acid SAM layer –formed in 2 hours - did not suffer from pit formation, not any sign of pits was observed. It means that the layer worked as a good, dense barrier between the metal and the solution. Similar observation was done when the layer deposition time was short (Figure 5.13). The undecenyl phosphonic acid SAM layer could only protect partly the metal surface from pitting; the surface roughening was much less, than without SAM layer (Figure 5.18), though this nanolayer is less effective than the fluoro amphiphile in SAM layer. It is due to the nanolayer structure, which is negatively influenced by the double bond in the molecules and this structure allows the Cl⁻ ion penetration onto the metal surface.

Figure 5.18: Influence of sodium chloride solution on carbon steel: (a): bare metal surface before and (c): after immersion in chloride solution for 1 h; (b): undecenyl phosphonic acid SAM nanolayer covered carbon steel surface and (d) after immersion into chloride solution for 1 h.
Figure 5.19: Influence of sodium chloride solution on the aluminum surface (a, b): metal surface visualized in 3D and its section; (c, d): aluminum surface after immersion into chloride solution for 1 h.
Figure 5.20: Influence of chloride solution on aluminum surface covered by fluorophosphonic acid SAM layer after 1 h (a, b): SAM surface morphology and its section before corrosion test (c, d): the SAM layer after immersion into the chloride solution for 1h.

Similar experiments were done with aluminum. SAM layers were formed from both amphiphiles. As Figure 5.19 and 5.20 show, when the metal without nanolayer, was immersed into sodium chloride, the metal surface became very rough. However, the presence of the fluoro amphihile in SAM film could control the pitting corrosion: there was not any sign of pits on the surface. This is very important in the case of aluminum as this metal is very sensitive to the pitting corrosion.
5.3.4 Influence of SAM layers on general corrosion

The influence of fluorophosphonic and undecenyl phosphonic acid SAM layers on general corrosion was investigated in aerated sodium perchlorate solution. AFM images taken after one hour immersion in the electrolyte solution are shown in Figures 5.21 and 5.22.

Figure 5.21: Aluminum alloy surface without nanolayer in sodium perchlorate for 1 h, visualized by section analysis and in 3D.

Figure 5.22: Aluminum alloy surface covered by fluorophosphonic acid SAM layer after immersion in sodium perchlorate for 1 h, visualized by section analysis and in 3D. The first set of images demonstrates the significant roughening of the aluminum surface in perchlorate solution (see the section lines). The fluorophosphonic acid SAM nanolayer could inhibit the surface deterioration, the corrosion, which is reflected in the smooth section line.
The AFM images showed in 3D as well as the section lines prove that the nanolayers developed on different metals inhibit both the general and the pitting corrosion. The inhibition efficiency depends on the amphiphile structure.

5.3.5 Influence of the SAM layer post-treatment on the anticorrosion efficacy

![AFM images showing the influence of the SAM layer post-treatment on the anticorrosion efficacy](image)

Figure 5.23: Influence of the SAM modification by irradiation on the anticorrosion activity (UP: undecenyl phosphonic acid; SAM layer formation time: 24 h; irradiation by 2 kGy and 20 kGy; immersion into NaCl solution).
I previously mentioned the change in the wettability caused by irradiated undecenyl phosphonic acid SAM surface. This modification of the SAM layer caused alteration in the anticorrosion activity, too. Figure 5.23 demonstrates the increased corrosion inhibition caused by the irradiated nanolayer coating on the carbon steel surfaces. It is clear that even the 2 kGy absorption made the nanolayer more resistant against corrosion, i.e. the layer compactness increased.

**5.3.6 Surface characterization by roughness parameters**

In order to characterize the influence of the corrosive environment numerically, I used the roughness parameters offered by the AFM software.

With increasing surface roughness the possibility of pitting corrosion increases, with other words a smoother surface is less corroded than a rougher one. On a smoother surface the defect places are reduced. Two different processes contribute to the pitting corrosion on smooth and rougher surface:

1. The pit formation ability is contributed to the presence of surface defects, which are more numerous on rough surface. On smooth surface the number of the defect places is reduced because of passivation (109, 110).

2. The diffusion of corrosion causing species (e.g. chloride ions) is affected by the surface roughness.

In case of the aggressive species are in contact with the surface and the diffusion of the corrosion products are limited, the repassivation of the metal is reduced and that allows the growing of a pit (109, 111, 112). The deep grooves trap the aggressive species, and the corrosion product cannot diffuse out of the groove; this increases the growth of the pit. On the other hand, when corrosion products can diffuse out quickly, no accumulation of the aggressive species can happen, the metal surface could be repassivated. In case of repassivation or stable oxide layer formation pit nucleation and growth is reduced (113).

The visualization of a surface by atomic force microscope permits the calculation of the surface roughness at nanoscale. It far exceeds the resolution achieved by other (e.g. optical) methods and is becoming increasingly important.
The measured surface roughness depends on the spatial and vertical resolution of an instrument. A real surface exhibits roughness on many length scales. Two important factors affect the resolution of roughness values measured at the metal surface: the vertical resolution, which is limited by the noise, and the spatial resolution limited by the tip radius.

The roughness can be characterized by several parameters.

i. *Maximum height of the profile* (R_{\text{max}}): the most significant parameter: the vertical distance between the deepest valley and highest peak around the surface profile.

ii. *Roughness average* (Ra): this is the most widely used parameter as it is easy to measure. The average roughness is the mean absolute profile; it does not make any distinction between valleys and peaks, the measure depends on the average heights profile.

iii. *Root means square roughness* (R_{q}): this is a statistical measure, which is similar to the roughness average; the only difference is the mean squared absolute value of surface roughness.

I used atomic force microscopy to determine the roughness parameters, which are summarized in Table 5.7 and 5.8 (the solid surface was carbon steel) and in Table 5.9 (the SAM layers were deposited onto aluminum surface). Both the undecenyl and fluorophosphonic acid SAM layers were investigated in aggressive solutions.

The average roughness profile (Ra) values measured on the bare *carbon steel* metal and on the nanolayer did not show significant difference, but after irradiation the Ra values increase a little. The root means square roughness (R_{q}) values, which are more sensitive to the valleys and peaks than the average roughness, show similar trend than the Ra values. Additionally, the R_{\text{max}} parameters followed the same tendency.
Table 5.7: Roughness parameters of carbon steel surfaces covered by undecenyl phosphonic acid SAM layer formed at 24h with and without radiation treatment.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Treatment</th>
<th>Rq [nm]</th>
<th>Ra [nm]</th>
<th>Rmax [nm]</th>
</tr>
</thead>
<tbody>
<tr>
<td>CS bare</td>
<td>-</td>
<td>5.3</td>
<td>4.2</td>
<td>53.1</td>
</tr>
<tr>
<td>CS U. P</td>
<td>-</td>
<td>4.9</td>
<td>3.7</td>
<td>40.4</td>
</tr>
<tr>
<td>CS U. P</td>
<td>2kGy</td>
<td>6.6</td>
<td>5.0</td>
<td>73.0</td>
</tr>
<tr>
<td>CS U. P</td>
<td>20kGy</td>
<td>8.8</td>
<td>7.2</td>
<td>67.6</td>
</tr>
</tbody>
</table>

On the other hand, when the metal sample without coating was dipped into chloride solution, all three roughness values increased drastically; the Rq and Ra values about twenty times, the Rmax around twelve times are higher (Table 5.8). The situation is different in the presence of the SAM layer. After immersion of SAM covered metal into sodium chloride solution, the Ra and Rq roughness parameters increased less than ten times, the Rmax value is around ten times higher. The influence of the irradiation on the layer structure makes an appearance conspicuously; in the case of 20kGy absorption, all three roughness parameters are much less than without the chloride solution. Even the 2kGy irradiation could decrease the roughness values after the corrosion test, and, along with, the anticorrosion efficiency.

Table 5.8: Effect of NaCl on roughness parameters of carbon steel surfaces covered with undecenyl phosphonic acid SAM layer (formed at 24h) with and without irradiation.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Treatment</th>
<th>Rq (nm)</th>
<th>Ra (nm)</th>
<th>Rmax (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CS bare</td>
<td>1h NaCl</td>
<td>103</td>
<td>84.1</td>
<td>644</td>
</tr>
<tr>
<td>CS U. P</td>
<td>1h NaCl</td>
<td>38.5</td>
<td>29.5</td>
<td>472</td>
</tr>
<tr>
<td>CS U. P</td>
<td>2kGy+1h NaCl</td>
<td>14.6</td>
<td>11.4</td>
<td>144</td>
</tr>
<tr>
<td>CS U. P</td>
<td>20kGy+1h NaCl</td>
<td>6.7</td>
<td>4.9</td>
<td>94</td>
</tr>
</tbody>
</table>

Similar experiments were carried out on aluminum. Salient features are the increase in Rq and Rmax parameters when the metal was immersed into sodium chloride and perchlorate solutions. In both cases, the electrolytes caused remarkable roughening. The presence of the fluorophosphonic SAM layer could control the surface properties: neither the chloride, nor the perchlorate electrolytes affected the metal surface. The surfaces are much less rough than in case the aluminum is dipped into the electrolytes without coating (Table 5.9). The roughness
parameters observed on the AFM images show the importance of these phosphonic acid amphiphiles applied in SAM layers and prove that they can effectively decrease both the general and the pitting corrosion though the fluoro amphipile is more effective. SAM layers on the carbon steel and aluminum drastically decreased the Rmax values that represent – as it was mentioned - the vertical distance between the deepest valley and highest peak around the surface profile.

Table 5.9: Effect of NaCl and NaClO₄ electrolytes on the roughness parameters and depth analysis data of aluminum surfaces covered with fluorophosphonic acid SAM layer formed at 4h.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Treatment</th>
<th>Rq (nm)</th>
<th>Rmax (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>-</td>
<td>9.4</td>
<td>122</td>
</tr>
<tr>
<td>Al 1h NaCl</td>
<td>86.2</td>
<td>413</td>
<td></td>
</tr>
<tr>
<td>Al 1h NaClO₄</td>
<td>139.0</td>
<td>422</td>
<td></td>
</tr>
<tr>
<td>Al F. P SAM</td>
<td>-</td>
<td>11.5</td>
<td>88</td>
</tr>
<tr>
<td>Al F. P SAM 1h NaCl</td>
<td>14.5</td>
<td>145</td>
<td></td>
</tr>
<tr>
<td>Al F. P SAM 1h NaClO₄</td>
<td>26.2</td>
<td>217</td>
<td></td>
</tr>
</tbody>
</table>

5.5 Characterization by IR spectroscopy

5.4.1. The effect of UV and irradiation on solid UPA revealed by IR spectroscopy

On the right spectra of Figure 5.24, IR1 the C-H stretching region of UPA samples is shown. The spectra are dominated by the antisymmetric ($v_{as}$) and symmetric ($v_s$) methylene stretching modes at 2918 and 2851 cm⁻¹, respectively. The position and shape of these two bands reflect an ordered conformation of the aliphatic chains (114). No changes were observed on the spectra of irradiated samples suggesting that the alkyl chains of the molecules remained intact. Small shift towards lower wavenumber were witnessed for sample UV treated for 180min. The band shift from higher to lower wavenumbers means that the number of gauche conformers decreases and the number of highly ordered all-trans conformers of alkyl chain increases. It seems plausible that the C=C double bonds of UPA are saturated upon UV treatment leading to the formation of a more ordered structure. The presence of C=C double bond produce only a very weak C-H stretching band at 3065 cm⁻¹ ($v$=C-H) not suitable for direct monitoring of structural changes.
Figure 5.24: IR1 C-H vibration regions of UPA samples’ spectra: right: C-H stretching region; left: C-H scissoring region.

Additional information about alkyl chain packing can be obtained also from the position and shape of methylene scissoring mode (δ\text{CH}_2). This spectral feature is very sensitive to side-by-side chain interactions as well as the packing organization of the methylene chain. IR1 two medium intense bands are observed for UPA samples. The high frequency band around 1471 cm\(^{-1}\) indicates partially ordered chains, while the low frequency band is characteristic for reduced side-by-side interactions and an increase in chain motion (115). It is worth to note that upon irradiation and 180min UV treatment the relative intensity of the higher frequency band of CH\(_2\) scissoring mode is increasing suggesting a more pronounced side-by-side interaction of alkyl chains parallel with reducing gauche defects caused by C=C moieties.
The relative intensities and shape of stretching vibrations of phosphonic group are presented in Figure 5.25. IR2. The most prominent is the shift of the P=O stretching band from 1178 towards 1184 cm\(^{-1}\) for UV treated samples. It seems plausible that the UV treatment disrupt some of the P=O bonds. An increase in band intensity can be observed also for C=C-H deformation modes at
993 and 907 cm$^{-1}$, respectively, suggesting that 180min UV treatment and irradiation may affect the C=C bond of starting UPA.

Table 5.10: IR band positions with possible assignments and explanations

<table>
<thead>
<tr>
<th>UPA</th>
<th>UPA_UV 120min</th>
<th>UPA_UV 180min</th>
<th>UPA 2kGy</th>
<th>UPA 20 kGy</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>3078 w</td>
<td>3078 w</td>
<td>3078 w</td>
<td>3078 w</td>
<td>3078 vw</td>
<td>-OH stretching</td>
</tr>
<tr>
<td>3065 w</td>
<td>3065 w</td>
<td>3065 w</td>
<td>3065 w</td>
<td>3065 vw</td>
<td>=C-H stretching</td>
</tr>
<tr>
<td>2999 w</td>
<td>2999 w</td>
<td>2996 w</td>
<td>2999 w</td>
<td>2999 vw</td>
<td>CH2 stretching</td>
</tr>
<tr>
<td>2978 w</td>
<td>2978 w</td>
<td>2976 w</td>
<td>2978 w</td>
<td>2978 vw</td>
<td>CH2 stretching</td>
</tr>
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<td>2960 w</td>
<td>2960 w</td>
<td>2959 w</td>
<td>2960 w</td>
<td>2959 vw</td>
<td>CH2 stretching</td>
</tr>
<tr>
<td>2918 vs</td>
<td>2918 vs</td>
<td>2916 vs</td>
<td>2918 vs</td>
<td>2919 vs</td>
<td>CH2 antisymmetric stretching of ordered chain</td>
</tr>
<tr>
<td>2851 vs</td>
<td>2851 vs</td>
<td>2849 vs</td>
<td>2851 vs</td>
<td>2851 vs</td>
<td>CH2 symmetric stretching of ordered chain</td>
</tr>
<tr>
<td>1643 m</td>
<td>1643 m</td>
<td>1642 m</td>
<td>1644 m</td>
<td>1644 m</td>
<td>C=C stretching</td>
</tr>
<tr>
<td>1471 m</td>
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<td>1469 m</td>
<td>1469 m</td>
<td>1469 m</td>
<td>CH2 scissoring</td>
</tr>
<tr>
<td>1462 m</td>
<td>1462 m</td>
<td>1462 m</td>
<td>1462 m</td>
<td>1462 m</td>
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<td>1402 w</td>
<td>1402 w</td>
<td>CH2 wagging</td>
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<tr>
<td>1287 w</td>
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<td>1288 w</td>
<td>1288 w</td>
<td>1288 w</td>
<td>CH2 twisting</td>
</tr>
<tr>
<td>-</td>
<td>1253 w</td>
<td>1253 w</td>
<td>-</td>
<td>-</td>
<td>P-OH deformation</td>
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<td>1204 w</td>
<td>1204 w</td>
<td>1204 w</td>
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<td>1179 m</td>
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<td>1183 m</td>
<td>1178 m</td>
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<td>P=O stretching</td>
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<td>1115 m</td>
<td>1116 m</td>
<td>1115 m</td>
<td>1115 m</td>
<td>PO$_3^{2-}$</td>
</tr>
<tr>
<td>1045 w</td>
<td>1044</td>
<td>1045 w</td>
<td>1043 w</td>
<td>1043 w</td>
<td>PO$_4^{3-}$</td>
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<td>993 s</td>
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<td>C=C-H deformation</td>
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<td>907 s</td>
<td>907 s</td>
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<td>907 s</td>
<td>907 s</td>
<td>C=C-H deformation</td>
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<td>779 m</td>
<td>779 m</td>
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<tr>
<td>718 m</td>
<td>718 m</td>
<td>718 m</td>
<td>718 m</td>
<td>717 m</td>
<td>CH2 rocking</td>
</tr>
</tbody>
</table>
5.5 Electrochemical measurements

During the corrosion of metals electrochemical processes take place in electrolytes that could be followed by different electrochemical techniques that all can contribute to the understanding of processes happen on the anode and cathode and help in elucidation of the mechanisms of the corrosion. The corrosion resistance of the SAM coated metal surfaces was evaluated either by the change of the open circuit potential in time, by potentiodynamic polarization, or by electrochemical impedance spectroscopy.

5.5.1 Open circuit potential ($E_{ocp}$) vs. time

The value of the open circuit potential ($E_{ocp}$) is a parameter determined by the rates of cathodic and anodic reactions on the corroding surface; then the rate of the anodic and cathodic reactions are equal and the net current is zero. It depends on the corrosion media, and chemical composition of the metal. The change in the open circuit potential as a function of time ($E_{ocp}$ vs time), under static conditions on bare and fluorophosphonic as well as undecenyl phosphonic acid SAM layer-coated carbon steel surfaces are presented in Figures 5.26 and 5.27 and in Tables 5.11 and 5.12. The time-dependent change of the open circuit potentials goes parallel with the self-assembled layer formation time: with increasing molecular layer formation time, the values of the $E_{ocp}$ change in the more positive direction, which shows that the surface coatings formed at longer time, altered significantly the metal surface. Comparing the shift of the $E_{ocp}$ of the coated samples with the uncoated, the highest shift was measured for SAM layer of undecenyl phosphonic acid formed at 24h, when the potential shifted from -502 mV (bare metal) to -209 mV. In the case of the more hydrophobic fluorophosphonic acid, already at shorter film formation time (0.5 h), there was a significant positive shift in the $E_{ocp}$ value: from -502 mV to -313 mV. Longer film formation time with fluoro amphiphile did not resulted in significant increase in the $E_{ocp}$. The 0.5 h gave an $E_{ocp}$ value of -313mV while that of 24h formed layer was -320mV. This indicates that the metal surface - already after half an hour - was mostly covered by fluoro amphiphile molecules. On the other hand, when the SAM layer was formed from the less hydrophobic undecenyl phosphonic acid molecules (which, additionally, contain double bond) to reach the same, significant shift in the $E_{ocp}$ needed more time; after 2 h film formation the $E_{ocp}$
shifted to -243mV. It means that the molecules needed more time to arrange themselves in a compact layer that can cover the more active sites on the metal surface.

Figure 5.26: Open circuit potential results for layers formed by fluorophosphonic acid.

Figure 5.27: Open circuit potential results of layers formed by undecenyl phosphonic acid.
Table 5.11: Steady-state potential values of carbon steel covered by SAM nanolayer of fluorophosphonic acid (formed in 20min).

<table>
<thead>
<tr>
<th>Layer formation time (h)</th>
<th>E (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bare metal</td>
<td>-502</td>
</tr>
<tr>
<td>0.5</td>
<td>-313</td>
</tr>
<tr>
<td>1</td>
<td>-298</td>
</tr>
<tr>
<td>2</td>
<td>-319</td>
</tr>
<tr>
<td>24</td>
<td>-320</td>
</tr>
<tr>
<td>48</td>
<td>-305</td>
</tr>
</tbody>
</table>

Table 5.12: Steady-state potential values of carbon steel covered by SAM nanolayer of undecenyl phosphonic acid.

<table>
<thead>
<tr>
<th>Layer formation time (h)</th>
<th>E (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ø</td>
<td>-502</td>
</tr>
<tr>
<td>0.5</td>
<td>-413</td>
</tr>
<tr>
<td>1</td>
<td>-336</td>
</tr>
<tr>
<td>2</td>
<td>-243</td>
</tr>
<tr>
<td>24</td>
<td>-209</td>
</tr>
</tbody>
</table>

5.5.2 Effect of layer formation time on the electrochemical processes

5.5.2.1 Potentiodynamic polarization results

The inhibitor efficiencies of the SAM layers formed on carbon steel surfaces were analyzed by electrochemical potentiodynamic measurements. This technique helps in the determination of the type and the function of the nanolayers. The results can answer whether the layers controlled the anodic reaction (metal dissolution) or the cathodic process. The carbon steel electrodes with or
without nanolayers were investigated in perchlorate solution. The results are plotted as potential (mV) versus logarithm of the current density (mA/cm²).

In cases of the fluorophosphonic and undecenyl phosphonic acid SAM layers, with increasing layer formation time there is a shift in the $E_{corr}$ potential in the more positive direction (Figures 5.28 and 5.29). It indicates that the anodic process (i.e. the metal dissolution) is inhibited. It is clear that the anticorrosion efficiency of the films increases with increasing layer formation time by both amphiphile nanolayers. The curves also show that in short nanolayer formation time inhomogeneous, not very compact layers are formed.

Effective SAM layers were formed by undecenyl phosphonic acid at 24hrs where the $i_{corr}$ was shifted to a very low value, similarly to the fluoro amphiphilic SAM layers. This demonstrates that with increasing layer formation time there is an improvement in the quality of the film structure, which is due to the formation of well-organized molecular structure within the films. In longer time, after the amphiphiles anchor to the metal surface through the phosphonic head groups, the molecules cover the whole surface and the hydrophobic molecular part vertically cover the area, which gives a hydrophobic character to the surface. Even in a short layer formation time (0.5 h), there was a decrease in the $i_{corr}$ (0.7 x10⁻⁶ A/cm² for fluorophosphonic acid SAM layer and 3 x10⁻⁶ A/cm² for the undecenyl phosphonic acid). We should keep in mind that the same influence of the layer formation time was observed on the corrosion potential ($E_{corr}$); it decreased as the time increased. Corrosion kinetic parameters $i_{corr}$, $E_{corr}$, efficiency: $\eta$ [%], are shown in Tables 5.13 and 5.14.
Figure 5.28: Effect of SAM layer formation time of fluorophosphonic acid on the corrosion reactions.

Figure 5.29: Effect of SAM layer formation time on the corrosion reactions in the case of undecenyl phosphonic acid.
Table 5.13: Corrosion parameters measured on carbon steel coated by fluorophosphonic acid SAM layers

<table>
<thead>
<tr>
<th>Chemical</th>
<th>Layer formation time (h)</th>
<th>(i_{\text{corr}}) (A/cm(^2)) ((10^{-6}))</th>
<th>(E_{\text{corr}}) (mV)</th>
<th>Efficiency ((\eta\ %))</th>
<th>Surface coverage((\Theta))</th>
</tr>
</thead>
<tbody>
<tr>
<td>bare</td>
<td>-</td>
<td>9</td>
<td>-608</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>F.P</td>
<td>0.5</td>
<td>0.7</td>
<td>-230</td>
<td>92</td>
<td>0.92</td>
</tr>
<tr>
<td>F.P</td>
<td>1</td>
<td>0.4</td>
<td>-170</td>
<td>96</td>
<td>0.96</td>
</tr>
<tr>
<td>F.P</td>
<td>2</td>
<td>0.3</td>
<td>-178</td>
<td>97</td>
<td>0.97</td>
</tr>
<tr>
<td>F.P</td>
<td>24</td>
<td>0.06</td>
<td>-175</td>
<td>99</td>
<td>0.99</td>
</tr>
<tr>
<td>F.P</td>
<td>48</td>
<td>1</td>
<td>-382</td>
<td>89</td>
<td>0.89</td>
</tr>
</tbody>
</table>

Table 5.14: Corrosion parameters measured on carbon steel coated by undecenyl phosphonic acid SAM layers

<table>
<thead>
<tr>
<th>Chemical</th>
<th>Layer formation time (h)</th>
<th>(i_{\text{corr}}) (A/cm(^2)) (10^{-6})</th>
<th>(E_{\text{corr}}) (mV)</th>
<th>Efficiency ((\eta\ %))</th>
<th>Surface coverage((\Theta))</th>
</tr>
</thead>
<tbody>
<tr>
<td>bare</td>
<td>-</td>
<td>9</td>
<td>-608</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>U. P</td>
<td>0.5</td>
<td>3</td>
<td>-285</td>
<td>67</td>
<td>0.67</td>
</tr>
<tr>
<td>U. P</td>
<td>1</td>
<td>2</td>
<td>-245</td>
<td>78</td>
<td>0.78</td>
</tr>
<tr>
<td>U. P</td>
<td>2</td>
<td>0.9</td>
<td>-221</td>
<td>90</td>
<td>0.90</td>
</tr>
<tr>
<td>U. P</td>
<td>24</td>
<td>0.7</td>
<td>-218</td>
<td>92</td>
<td>0.92</td>
</tr>
<tr>
<td>U. P</td>
<td>48</td>
<td>0.8</td>
<td>-223</td>
<td>91</td>
<td>0.91</td>
</tr>
</tbody>
</table>

If I want to summarize the results of the potentiodynamic polarization experiments, I can mention that within the first short period a fast surface adsorption occurs, when the low coverage is correlated with the molecules “laying along the surface”. In this period there is a difference between the corrosion current dependence. High quality SAM layer with a high coverage (the molecules are “standing” on the surface) is reached after 24 h in case of fluorophosphonic acid.
It is interesting that after 48 h molecular layer formation time, a small decrease in the efficiency of fluoro amphiphile was observed. It could be explained with adsorption of further molecules onto the first layer forming a second layer, which disturbs the orientation of the first line of molecules. In this case a small part of the head groups will face the aqueous environment that decreases the water contact angle value, i.e. increases the hydrophilicity. There is similar observation given in the literature. In case of the undecenyl phosphonic acid, a dense nanolayer forms in 24 h. It shows that the undecenyl amphiphile needs more time for formation of a well-structured nanolayer. The film formation happens through continuous molecular adsorption and desorption as well as by molecular reorientation. The well-oriented dense structure is responsible for the high anticorrosion efficiency.

![Figure 5.30: Time dependent effectiveness values of nanolayers formed on carbon steel surfaces.](image)

Correlation was found between the layer formation time and the efficiencies (Figure 5.30) and also between the calculated surface coverage from the efficiencies (Tables 5.13, 5.14). In the case of these amphiphiles the curves followed Langmuir kinetics; i.e. the number of the active locations is infrequent, there is a specific bound and the result is a monolayer. To prove the Langmuir type correlation, the time (Time (h)) and the time/coverage (Time (h) / Θ) were plotted (Figure 5.31). As for the undecenyl amphiphile the correlation between time and coverage was a little similar to a Freundlich isotherm, which is also a classical isotherm, like the Langmuir one. The difference between them is that during the layer formation, the binding force decreases and...
there is not a well measurable saturation. The efficiency curves, which go parallel with the surface coverage curves, show the differences in the adsorption in the first 2 hours: the undecenyl amphiphile molecules can adhere to the metal surface but form a well-structured layer less easily than the fluorophosphonic one. Both amphiphiles follow the Langmuir adsorption kinetics as the straight lines in Figure 5.31.

Figure 5.31: Correlation between the SAM layer formation time (h) of the fluorophosphonic acid as well as of the undecenyl phosphonic acid and the formation time divided by the surface coverage (time (h) / \( \Theta \)); a: change in 48 hours; b: change in the first 2 hours.
In both cases, the correlation gave straight lines not only in the early period of the adsorption but at the longer layer formation time, too. It proves that the process of the layer formation follows Langmuir adsorption as shown in Figure 5.30. This is an important correlation as originally the Langmuir adsorption is correlated with the concentration of the molecular concentration but I could replace this value with the SAM layer formation time.

5.5.2.2 Electrochemical Impedance Spectroscopy (EIS) results

Organic SAM nanofilms were developed that could successfully resist to the aggressive environments metals are exposed to. In order to assist the development and characterization of these films, in the evaluation of their performance the electrochemical impedance spectroscopy is of great importance.

The EIS is a non-destructive measurement, provides information on surfaces in corrosive medium. A small amplitude sinusoidal alternating signal is applied on the SAM-modified metal surface. The analysis of the impedance spectrum results in an equivalent circuit model, which allows the evaluation of the electrochemical information measured on the nanocoated surface.

The equivalent circuit, which interprets data and gives physical meanings of the studied system, is used in electrochemical systems. In my case, the equivalent circuit is composed of $R_s$ bulk solution resistance, of the constant phase element (CPE: it replaces the double layer capacitance models the behavior of an imperfect capacitor), and parallel with it, the $R_{ct}$ charge transfer resistance that describes the electrochemical reactions under control. This technique is very useful to investigate organic films on metallic surfaces used as corrosion inhibitors. Using the $R_{ct}$ values, we can evaluate the corrosion behavior of the nanolayers as the charge transfer resistance is inversely related to corrosion rate.

EIS technique was utilized to probe the quality and the behavior of nanolayers on electrodes under the effects of layer formation time, pH of the electrolyte, and the time of immersion in the electrolyte. The results are shown in forms of: 1. Nyquist plot where the real vs. imaginary
amplitudes are plotted; a Nyquist plot is a parametric plot of a frequency response used in automatic control and signal processing as well as is applied for assessing the stability of a system; 2. Bode plots are very useful to represent the gain and phase of a system as a function of frequency. This is referred to as the frequency domain behavior of a system. In Bode graphs with the same data the logarithm of the modulus of the impedance $\log |Z|$ vs. $\log \omega$ is the Bode module plot and the phase angle $(\Theta)$ vs. $\log \omega$ is the Bode phase plot.

The EIS results were evaluated by the Zview software in order to fit the measured data to the best electric circuit model in which the $R_s$, $R_{ct}$ and the CPE values are calculated. The CPE is used to replace a capacitor that represents the double layer or non-homogeneity of the surface. Two components are represented by the CPE, the $(Q)$ reflects the capacitance of the CPE impedance and the $(n)$ reflects the phase shift degree of the non-ideality in the capacitive behavior and the uniformity of the surface.

All electrochemical results achieved by the EIS technique were represented by the equivalent circuit as shown in Figure 5.32.

![Figure 5.32: Equivalent electric circuit for the EIS results.](image)

As the Nyquist and Bode plots show (Figures 5.33 and 5.34), the nanolayers of both amphiphiles on carbon steel surfaces affected the surface properties; this resulted in higher protection as the layer formation time increased. In the case of the fluorophosphonic acid nanolayer the charge transfer resistance (which informs about the surface compactness) results in the highest value ($R_p$: 54550 $\Omega$.cm$^2$) in case when the layer formed in 24 h. At shorter nanofilm formation (0.5 h) this value is much less: 10718 $\Omega$.cm$^2$, which is due to the submolecular island nucleation; they will gradually grow and coalesce in longer time. With other words: in shorter time the molecules cannot occupy all the active places on the metal surface, active sites are left on the metal surface.
More time is needed for seize the full surface. Yet even at a short time (0.5 h) the layer formed by fluorophosphonic acid gave 8-times better result when compared with the $R_p$ of bare metal (Table 5.15).

Figure 5.33: Fluorophosphonic acid-SAM layer formed on carbon steel; time dependent Nyquist and Bode plots.
Figure 5.34: Undecenyl phosphonic acid-SAM layer formed on carbon steel; time dependent Nyquist and Bode plots.

Results of the undecenyl phosphonic acid (Table 5.16) showed the same trend in the function of layer formation time as the fluorophosphonic acid. For the layer formed at 0.5 h, the $R_p$ value
recorded was 8671 Ω.cm²; when the time increased up to 1 h and 2 h, the $R_p$ increased to 19975 Ω.cm² and 29638 Ω.cm², respectively, resulting in an efficiency increase from 91% to 95%. An $R_p$ value of 44942 Ω.cm² was recorded for the layer formed at 24 h, which is almost five fold of that achieved by the 0.5 h layer, which is due to the formation of a more compact, protective thin film that inhibited the corrosion sites on the metal surface. Evaluation of these results, the electrochemical process taking place on the metal surface is charge transfer and the diffusion does not really influence; the layers managed to a certain point to prevent the oxygen diffusion to the base metal. It can also be noticed that as the $R_p$ increases, the Q decreases; the n does not change significantly. It means that the surface uniformity is almost constant. These results indicate that the charge transfer from the metal decreased due to the blocking of some corrosion sites on the metal surface.

This reveals itself in the change of the $R_p$ as well as in the Q values. The higher $R_p$ values represent the increase in the charge transfer resistance; the decrease in the Q values the positive change in the capacitance, the almost constant n values inform about the surface homogeneity.

Even though the results of the undecenyl phosphonic acid nanolayer were good in protecting the metal surface, yet again the results of fluorophosphonic acid nanolayer were better due to the more compact and more hydrophobic fluorophosphonic acid SAM layer. The contact angle values prognosticated these results: the more hydrophobic nanofilms can stand against better to the attack of a corrosive environment.

<table>
<thead>
<tr>
<th>Chemical</th>
<th>Layer formation time (h)</th>
<th>Q (μF/cm²)</th>
<th>n</th>
<th>$R_p$ (Ω. cm²)</th>
<th>η (%)</th>
<th>Surface coverage(θ)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bare</td>
<td>-</td>
<td>78</td>
<td>0.68</td>
<td>1246</td>
<td>-</td>
<td>0.76</td>
</tr>
<tr>
<td>FP</td>
<td>0.5</td>
<td>21</td>
<td>0.79</td>
<td>10718</td>
<td>76</td>
<td>0.76</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>9</td>
<td>0.76</td>
<td>24254</td>
<td>95</td>
<td>0.95</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>8</td>
<td>0.75</td>
<td>28923</td>
<td>96</td>
<td>0.96</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>4</td>
<td>0.73</td>
<td>54550</td>
<td>98</td>
<td>0.98</td>
</tr>
<tr>
<td></td>
<td>48</td>
<td>4</td>
<td>0.76</td>
<td>47703</td>
<td>97</td>
<td>0.97</td>
</tr>
</tbody>
</table>

Table 5.15: EIS parameters for SAM layers formed by fluorophosphonic acid on carbon steel.
Table 5.16: EIS parameters for SAM layers formed by undecenyl phosphonic acid on carbon steel

<table>
<thead>
<tr>
<th>Chemical</th>
<th>Layer formation time (h)</th>
<th>Q (μF/cm²)</th>
<th>n</th>
<th>R_p (Ω·cm²)</th>
<th>η (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bare</td>
<td>-</td>
<td>78</td>
<td>0.68</td>
<td>1246</td>
<td>-</td>
</tr>
<tr>
<td>UP</td>
<td>0.5</td>
<td>23</td>
<td>0.81</td>
<td>8671</td>
<td>59</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>9</td>
<td>0.75</td>
<td>19975</td>
<td>91</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>8</td>
<td>0.80</td>
<td>29638</td>
<td>95</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>9</td>
<td>0.77</td>
<td>44942</td>
<td>97</td>
</tr>
<tr>
<td></td>
<td>48</td>
<td>20</td>
<td>0.73</td>
<td>31243</td>
<td>97</td>
</tr>
</tbody>
</table>

Figure 5.35: Correlation between the polarization resistance and the layer formation time measured on fluorophosphonic acid (FP) and undecenyl phosphonic acid (UP) SAM coated carbon steel.

These curves depicted in Figure 5.35 support the observation given in case of correlation between the layer formation time and the surface coverage. The fluorophosphonic acid follows saturation already in the first period of the layer formation (a rapid increase in the R_p values reaching saturation) but when the undecenyl phosphonic acid builds the nanolayer, the saturations is reached at a much longer time.
5.5.3 Effect of electrolyte pH on the anticorrosion efficiency

5.5.3.1 Influence of the pH measured by potentiodynamic polarization

It was interesting how these SAM layers behave in electrolytes with different pH values. Results are summarized in Figures 5.36-5.38 and Table 5.17-5.19. At pH=3 both chemicals behaved in the same trend as the bare metal in spite of a slight shift in the $E_{corr}$ to the positive range. There was a rapid increase in the $i_{corr}$ values above the $E_{corr}$ that might be related to the distorted SAM layer structure, which could be caused by the $\text{H}^+$ ions’ effect on the phosphono head group structure. We should keep in mind that the $pK_1$ value of these amphiphiles are between 2 and 3 (fluorophosphonic acid: 2.08; undecenyl phosphonic acid: 2.42), the acidic environment changes the place the head group occupy and the ratio among the mono- bi- or tridentate- adsorption types could also be changed that results in a less compact SAM structure. This leads to an enhanced metal dissolution.

Figure 5.36: Behavior of the bare metal, fluorophosphonic acid and undecenyl phosphonic acid SAM layers covered carbon steel in electrolyte at pH = 3.
Table 5.17: Effect of pH on the anticorrosion behavior of layers formed by fluorophosphonic acid and undecenyl phosphonic acid on carbon steel in electrolyte with pH = 3

<table>
<thead>
<tr>
<th>Chemical</th>
<th>pH</th>
<th>$i_{corr}$ (A/cm$^2$) ($10^{-6}$)</th>
<th>$E_{corr}$ (mV)</th>
<th>Efficiency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bare metal</td>
<td>3</td>
<td>8</td>
<td>-475</td>
<td>-</td>
</tr>
<tr>
<td>F. P</td>
<td>3</td>
<td>7</td>
<td>-370</td>
<td>13</td>
</tr>
<tr>
<td>U. P</td>
<td>3</td>
<td>4</td>
<td>-370</td>
<td>50</td>
</tr>
</tbody>
</table>

As the pH value increased to pH 7, the carbon steel electrode covered by fluorophosphonic acid SAM layer showed a rapid decrease in the corrosion current density ($i_{corr}$) up to $0.02 \times 10^{-6}$ A/cm$^2$ value (at pH 3: $7 \times 10^{-6}$ A/cm$^2$) and the $E_{corr}$ also shifted to the anodic direction. These parameters indicate that the layer could control much better the corrosion processes in neutral solution. The same was observed on the undecenyl phosphonic acid SAM at pH 7 value: decrease in the $i_{corr}$ and positive shift in the $E_{corr}$. Accordingly, the rate of the corrosive metal dissolution decreased significantly. At pH 9 and 11 values there were changes in the $E_{corr}$ and $i_{corr}$ values compared to that of the pH 7 measured on the fluorophosphonic and undecenyl phosphonic acid nanolayers. There is a small increase in the metal dissolution. These changes could be due to the start in the amphiphile dissolution and to change in the structure of the head group within the SAM layers; both effects are influenced by the second pK values (undecenyl: 8.08; fluorophosphonic: 7.63) and the result is a little decrease in the anticorrosion activity of both SAM layers at higher than neutral pH values (Figures 5.37 and 5.38, Tables 5.18 and 5.19).
Figure 5.37: Influence of the electrolyte pH values on the anticorrosion activity of the SAM layers formed by fluorophosphonic acid on carbon steel (NaClO$_4$; 24 h SAM layer formation time).

Figure 5.38: Influence of the electrolyte pH values on the anticorrosion activity of the SAM layers formed by undecenyl phosphonic acid on carbon steel (NaClO$_4$; 24 h SAM layer formation time).
Table 5.18: Corrosion parameters measured at different pH values in the presence of fluorophosphonic acid SAM layers on carbon steel (NaClO₄; 24 h layer formation time).

<table>
<thead>
<tr>
<th>Chemical</th>
<th>pH</th>
<th>(i_{corr}) (A/cm²) ((10^{-6}))</th>
<th>(E_{corr}) (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F. P</td>
<td>3</td>
<td>7</td>
<td>-370</td>
</tr>
<tr>
<td>F. P</td>
<td>7</td>
<td>0.02</td>
<td>-253</td>
</tr>
<tr>
<td>F. P</td>
<td>9</td>
<td>0.06</td>
<td>-321</td>
</tr>
<tr>
<td>F. P</td>
<td>11</td>
<td>0.05</td>
<td>-208</td>
</tr>
</tbody>
</table>

Table 5.19: Corrosion parameters measured at different pH values in the presence of undecenyl phosphonic acid SAM layers on carbon steel (NaClO₄; 24 h layer formation time).

<table>
<thead>
<tr>
<th>Chemical</th>
<th>pH</th>
<th>(i_{corr}) (A/cm²) ((10^{-6}))</th>
<th>(E_{corr}) (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>U. P</td>
<td>3</td>
<td>4</td>
<td>-370</td>
</tr>
<tr>
<td>U. P</td>
<td>7</td>
<td>0.06</td>
<td>-342</td>
</tr>
<tr>
<td>U. P</td>
<td>9</td>
<td>0.06</td>
<td>-398</td>
</tr>
<tr>
<td>U. P</td>
<td>11</td>
<td>0.04</td>
<td>-462</td>
</tr>
</tbody>
</table>

5.5.3.2 Influence of the pH measured by electrochemical impedance spectroscopy

The pH effect on the fluorophosphonic acid and undecenyl phosphonic acid nanolayers along with the bare metal in electrolyte at pH=3 are shown in Figure 5.39. The effect is the same that was presented with the other electrochemical technique and the explanation is the same: there is a change in the head group structures initialized by the acidic environment. The \(R_p\) values were quite the same for all three surfaces (fluorophosphonic acid layer, undecenyl phosphonic acid nanolayer and bare metal) as shown in Table 5.20.

Table 5.20: EIS parameters measured on carbon steel as bare metal, SAM layers formed by fluorophosphonic acid and undecenyl phosphonic acid on carbon steel in electrolyte at pH=3.

<table>
<thead>
<tr>
<th>Chemical</th>
<th>pH</th>
<th>Q (µF/cm²)</th>
<th>n</th>
<th>(R_p) (Ω. cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bare metal</td>
<td>3</td>
<td>38</td>
<td>0.78</td>
<td>1650</td>
</tr>
<tr>
<td>F. P</td>
<td>3</td>
<td>38</td>
<td>0.72</td>
<td>1790</td>
</tr>
<tr>
<td>U. P</td>
<td>3</td>
<td>40</td>
<td>0.70</td>
<td>1691</td>
</tr>
</tbody>
</table>
Figure 5.39: Comparison of EIS results got in corrosion experiment by bare metal, layers formed by fluorophosphonic acid and undecenyl phosphonic acid on carbon steel in electrolyte at pH=3.
Figure 5.40: Nyquest and Bode plots for layers formed by fluorophosphonic acid on carbon steel in electrolyte at different pH values.
Figure 5.41: Nyquist and Bode plots measured on carbon steel surface covered by undecenyl phosphonic acid SAM layer in electrolyte at different pH values.
The SAM layer performed best at pH=7, the efficiency was the highest at neutral pH. As the pH increased above this value (pH=9 and pH=11) the Rp decreased, the anticorrosion efficacy decreased. The Rp of fluorophosphonic acid layer in a solution with a pH of 9 resulted in an Rp value of just 6546 $\Omega \cdot \text{cm}^2$ while for pH=11 it was 3371 $\Omega \cdot \text{cm}^2$. This also might be related to the change in the head group structure within the SAM layers because of the second pK values. Similar behavior was noticed at the undecenyl phosphonic acid SAM layers as Figures 5.40 and 5.41 and Tables 5.21 and 5.22 illustrate.

Table 5.21: EIS parameters for SAM layers formed by fluorophosphonic acid on carbon steel in electrolyte at different pH values

<table>
<thead>
<tr>
<th>Chemical</th>
<th>pH</th>
<th>Q ($\mu\text{F/cm}^2$)</th>
<th>n</th>
<th>Rp ($\Omega \cdot \text{cm}^2$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FP</td>
<td>3</td>
<td>38</td>
<td>0.72</td>
<td>1790</td>
</tr>
<tr>
<td>FP</td>
<td>7</td>
<td>5</td>
<td>0.78</td>
<td>40366</td>
</tr>
<tr>
<td>FP</td>
<td>9</td>
<td>7</td>
<td>0.75</td>
<td>6546</td>
</tr>
<tr>
<td>FP</td>
<td>11</td>
<td>3</td>
<td>0.75</td>
<td>3371</td>
</tr>
</tbody>
</table>

Table 5.22: EIS parameters measured on SAM layers formed by undecenyl phosphonic acid on carbon steel in electrolyte at different pH values

<table>
<thead>
<tr>
<th>Chemical</th>
<th>pH</th>
<th>Q ($\mu\text{F/cm}^2$)</th>
<th>n</th>
<th>Rp ($\Omega \cdot \text{cm}^2$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>UP</td>
<td>3</td>
<td>40</td>
<td>0.7</td>
<td>1691</td>
</tr>
<tr>
<td>UP</td>
<td>7</td>
<td>9</td>
<td>0.77</td>
<td>38065</td>
</tr>
<tr>
<td>UP</td>
<td>9</td>
<td>16</td>
<td>0.76</td>
<td>29061</td>
</tr>
<tr>
<td>UP</td>
<td>11</td>
<td>4</td>
<td>0.9</td>
<td>35454</td>
</tr>
</tbody>
</table>

5.5.4 Time dependence of protectively effect on of SAM layer covered metal

The effect of the immersion of metals (with and without SAM coating, formed at 24 h) in electrolyte had been studied for both of the SAM layers formed from fluorophosphonic acid and undecenyl phosphonic acid.

As shown in Figures 5.42 and 5.43 the Rp values decrease as the immersion time in electrolyte increases, which indicates that the nanolayer starts to change the structure, becomes less
compact, more permeable, and thus less protective. The $R_p$ value of the fluorophosphonic acid SAM layer formed at 24h and immersed in the electrolyte for 20 min was 54550 $\Omega$. cm$^2$ (Table 5.23) and after 6h of immersion dropped to 27390 $\Omega$. cm$^2$, even though this value is about 22-times higher than that of the bare metal (1246 $\Omega$. cm$^2$) handled under the same conditions. Increasing the immersion to 24 h, there was a decrease to $R_p$ (21438 $\Omega$. cm$^2$). This means that in 24 h the layer effectiveness decreased to almost the half, but this value is still 17-time higher than that of the bare metal. The reason for the decrease in the anticorrosion activity could be that the longer SAM layer formation time allows the adhesion of a secondary molecular layer, which results in a more hydrophilic surface.

The behavior of the undecenyl-phosphonic acid SAM layer is similar to the fluoro amphiphiles; there is a decrease in the $R_p$ values after dipping the coated samples into the electrolyte. The starting $R_p$ value measured at 20 min was 44942 $\Omega$. cm$^2$, which sharply decreased to 1799 $\Omega$. cm$^2$ at 24 h of immersion time (Table 5.24). This value is close to the bare metal’s, which shows that the surface metal was corroded, even though there were no signs of corrosive attacks by visual inspection, the nanolayer turned to be permeable for the electrolyte, the metal corrosion was initiated.

This big difference in the $R_p$ values observed between those achieved by fluorophosphonic acid SAM layers and those of undecenyl phosphonic acid SAM nanocoating (layer formation time in both cases was 24h) is related to the differences in the structure of the two amphiphiles: the undecenyl has a double bond that does not allow the formation of a compact layer, but the fluorophosphonic amphiphile has a hydrophobic alkyl chain, which supports the formation of a densely packed, well-ordered nanofilm. All these observations are reflected in the Nyquist and Bode plots.
Figure 5.42: Effect of immersion time into electrolyte on fluorophosphonic acid SAM layer covered carbon steel surface.
Figure 5.43: Effect of immersion time into electrolyte on undecenyl phosphonic acid SAM layer covered carbon steel surface.
Table 5.23: Effect of immersion time into electrolyte on fluorophosphonic acid SAM layer covered carbon steel surface.

<table>
<thead>
<tr>
<th>Chemical</th>
<th>Immersion time (h)</th>
<th>Q (μF/cm²)</th>
<th>n</th>
<th>R_p (Ω. cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F. P</td>
<td>20min</td>
<td>2</td>
<td>0.87074</td>
<td>54550</td>
</tr>
<tr>
<td></td>
<td>1 h</td>
<td>3</td>
<td>0.85764</td>
<td>42770</td>
</tr>
<tr>
<td></td>
<td>2 h</td>
<td>3</td>
<td>0.8555</td>
<td>42030</td>
</tr>
<tr>
<td></td>
<td>3 h</td>
<td>3</td>
<td>0.85064</td>
<td>35690</td>
</tr>
<tr>
<td></td>
<td>4 h</td>
<td>4</td>
<td>0.8451</td>
<td>29350</td>
</tr>
<tr>
<td></td>
<td>5 h</td>
<td>11</td>
<td>0.8195</td>
<td>27390</td>
</tr>
<tr>
<td></td>
<td>6 h</td>
<td>13</td>
<td>0.8109</td>
<td>17270</td>
</tr>
<tr>
<td></td>
<td>24 h</td>
<td>6</td>
<td>0.7834</td>
<td>21438</td>
</tr>
</tbody>
</table>

Table 5.24: Effect of immersion time into electrolyte on undecenyl phosphonic acid SAM layer covered carbon steel surface.

<table>
<thead>
<tr>
<th>Chemical</th>
<th>Immersion time (h)</th>
<th>Q (μF/cm²)</th>
<th>n</th>
<th>R_p (Ω. cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>U. P</td>
<td>20min</td>
<td>4</td>
<td>0.84307</td>
<td>44942</td>
</tr>
<tr>
<td></td>
<td>1hr</td>
<td>10</td>
<td>0.8351</td>
<td>15115</td>
</tr>
<tr>
<td></td>
<td>2hrs</td>
<td>10</td>
<td>0.83617</td>
<td>17232</td>
</tr>
<tr>
<td></td>
<td>3hrs</td>
<td>16</td>
<td>0.78094</td>
<td>5936</td>
</tr>
<tr>
<td></td>
<td>4hrs</td>
<td>28</td>
<td>0.72632</td>
<td>3904</td>
</tr>
<tr>
<td></td>
<td>5hrs</td>
<td>44</td>
<td>0.70875</td>
<td>3816</td>
</tr>
<tr>
<td></td>
<td>6hrs</td>
<td>58</td>
<td>0.71363</td>
<td>4029</td>
</tr>
<tr>
<td></td>
<td>24hrs</td>
<td>61</td>
<td>0.71554</td>
<td>1799</td>
</tr>
</tbody>
</table>
Chapter Six

6. Summary

6.1 Preparation of SAM layers; amphiphiles and metals

My PhD work was dedicated to the investigation of special self-assembled molecular layers formed by two phosphonic acid amphiphiles:

- **Undecenyl phosphonic acid**: \(\text{CH}_2=\text{CH-(CH}_2)_9\text{-PO(OH)}_2\), that contains a double bond at the end of the carbon chain;

- **Fluorophosphonic acid**: \(\text{C}_6\text{F}_{13}-\text{C}_2\text{H}_4\text{-PO(OH)}_2\), where the majority of the hydrogen atoms in the carbon chain is replaced by fluorine atoms.

Parameters influenced the preparation of compact self-assembled molecular layers:

- the layer formation time
- the amphiphile concentration

In order to compare the influence of the molecular structure on the properties of the deposited layers, some other amphiphiles were also used in SAM film such as:

- **styren-co-styphos** acid that had a special structure; when this molecules (like the other phosphonic amphiphiles) are anchored to the metal surfaces through the phosphonic head groups, the remarkable polymeric aralkyl chain could form a hydrophobic net over the metal surface.

- **dodecyl and hexadecyl phosphonic acids**: The difference in the carbon chain length increases the hydrophobicity that influences the compactness of a SAM layer.

Metals under investigation

- **Iron alloys**: carbon steel, stainless steels 304 and 316 as these are the mostly used iron alloys.

- **Aluminum** alloy with a small Mg content is also important industrial metal.
6.2 Characterization of the SAM layers

The techniques listed below were used for the characterization of the SAM nanolayers.

6.2.1 Change in the wettability followed by dynamic and static contact angle values

At first the SAM layers formed on metal surfaces were characterized by the changes in the contact angle/wettability caused by the presence of the nanolayers, due to the different chemicals, deposition time and the increase in the amphiphile concentration.

The conclusions drawn from the contact angle data are as following:

i. **Influence of the layer deposition time:** With increasing layer deposition time the wettability measured in water decreases. On all types of metals, the formation of a very compact layer needs longer time (around 24 h). But a very long time (around 48 h) does not always increase the surface hydrophobicity, which could be due to deposition of a second layer (in this case the outer groups are the hydrophilic phosphonic ones).

ii. **Influence of the amphiphile type:** Considering the chemicals that formed the SAM layers, the fluorophosphonic acid produced the most hydrophobic nanolayers on all metals. This is understandable as the presence of fluorine atoms in the alkyl chain increases the water repulsive effect significantly. It is clearly demonstrated when I compared the contact angles of the SAM layers formed by the dodecyl and hexadecyl phosphonic acid with that one of fluorophosphonic acid. The presence of even longer alkyl chains in C12P (the number of carbon atoms is twelve) and C16P (the number of carbon atoms is sixteen) cannot overcome the hydrophobicity of fluorophosphonic acid (where the alkyl chain is shorter, the number of carbon atoms in the backbone is eight) SAM layers. The wettability value of the styren-co-styphos acid SAM nanofilms was between those values produced by the undecenyl and fluorophosphonic acid nanolayers. The undecenyl phosphonic acid amphiphile does not produce as hydrophobic surface as the fluorophosphonic acid. The contact angle data clearly proved that the undecenyl amphiphile cannot form very densely packed nanolayer because of the double bond at the end of
the alkenyl chain. It is not striking as a CH$_2$=CH- groups hinder the formation of a well-ordered molecular structure. The SAM never reaches a very well ordered state, even at longer assembly time and at higher concentration.

### iii. Influence of the amphiphile concentration:
Other information derived from the contact angle measurements is that the increase in the concentration of the amphiphiles (from where the deposition of nanolayer happens) results in lower water wettability, which means that the metal surface turned to be more hydrophobic. It is also important to mention that the increasing immersion time has more significant impact on the contact angle values than the increase in the concentration. In a shorter time the amphiphilic molecules start to cover the metal surface and the layer does not cover fully the solid, though the surface is more hydrophobic than the bare metal. At higher concentration (and at longer deposition time) the SAM layer islands grow together and cover homogeneously the metal surface.

### iv. Analysis of the advancing and receding contact angle values after multiple dipping:
The layer compactness was characterized by the change in the advancing and receding contact angle values after several dipping of the coated metal samples into water. In case of amphiphiles like the undecenyl phosphonic acid the compactness of the SAM layer never reaches a very well ordered state, even at longer assembly time and at higher amphiphile concentration. It is reflected in the differences between the advancing and receding contact angle values. After the second, third etc. immersion, both the advancing and the receding contact angle values decreased, which is the consequence of the non-ideal ordering of the molecules in the nanolayer.

### v. Influence of the post-treatment of the undecenyl phosphonic acid SAM layer:
When the SAM nanolayers of the undecenyl phosphonic acid were modified via external actions, i.e. by illumination with UV light and irradiation by $^{60}$CO–gamma ray, according to the wettability of the coated surfaces, the original undecenyl SAM films were converted into a more compact ones as not only the advancing but the receding contact angle values increased. The two different influences were supposed to interact with the double bonds at the SAM surface. These modifications were supposed to change the hydrophobicity i.e. the compactness of the nanofilms. The illumination by UV light increased the contact angle (i.e. the SAM layer turned to be more
compact) especially when it was used for longer time. The irradiation with 2kGy could less interfere with the nanolayers, but the 10-times stronger irradiation (20kGy) modified already the coating surface wettability significantly causing a film with much higher hydrophobicity. The altered structures were investigated by infrared spectroscopy. The results confirmed the change in the structure of the post-treated surface in both cases. The increase in the compactness of the nanofilms is important from the point of view of application possibilities of this type of coatings (e.g. as anticorrosion layers). The influence of the post-treatment was evidenced by other techniques (AFM, IR), too.

i. **Influence of metals:** There were significant differences in the surface hydrophobicity measured on certain metals covered by the same amphiphilic SAM. The highest contact angle values, so-called superhydrophobic surfaces were produced on aluminum by SAM layers built of the undecenyl- and fluorophosphonic acids as well as of the styren-co-styphos acid. These layers were not “destroyed” by further dipping into water, which is due to the very compact oxide layer on the aluminum surface whereto the phosphonic groups could anchor; and the result was a high density of the amphiphile on the metal surface. The second highest hydrophobic value was measured on the stainless steel 304. The difference in the wettability of SAM layer-covered stainless steel 304 and 316 is due to the presence of the molybdenum, which is present (though in small concentration) in the stainless steel 316; it disturbs the homogeneity of the surface oxide layer whereto the phosphonic groups are attached.

ii. **Influence of the surface roughness on the SAM layer formation:** The impact of the roughness factor showed that the SAMs on rougher surface behave interestingly. The wetting behavior of undecenyl nanolayers differed from the characteristic of the nanolayer formed from fluorophosphonic acid amphiphile: generally, on smooth metal surfaces always the fluorophosphonic molecules produce more hydrophobic coating (under identical conditions). On rougher surface the undecenyl amphiphile produced more hydrophobic layer, which is due to the structure of this molecule (double bond at the end of the alkenyl chain). This molecular part could support the retention of air bubbles near to the solid surface. This is clearly shown by the behavior of the bare metal. The increasing smoothness on the surface allowed the formation of
more compact film of the fluoro amphiphilic molecule; the undecenyl one on smoother surface produced less hydrophobic surface due to the molecular structure.

iii. **Influence of the post-treatment on the undecenyl amphiphile SAM layer:** The molecular structure of the undecenyl phosphonic acid SAM layers was altered via illumination with UV light and by irradiation by $^{60}$Co-gamma ray. The two different influences were supposed to interact with the double bonds at the SAM surface producing a polymer network which should result in a more compact nanofilm. The illumination by UV light increased the contact angle (i.e. the SAM layer turned to be more hydrophobic) especially when it was used for longer time. The irradiation with 2kGy could less interfere with the nanolayer, but the 10-times stronger irradiation (20kGy) modified already the coating surface significantly causing a film with much higher hydrophobicity. The altered structures were investigated by XPS technique and infrared spectroscopy and in both cases the preliminary results confirmed the change in the structure of the post-treated surface. This was important from the point of view of anticorrosion activity (as the AFM measurements proved).

6.2.2 **Visualization of surface morphology of SAM nanolayer coated metal surfaces on air and in corrosive environment**

The contact angle values characterized the change in the wettability of SAM layers of different amphiphiles deposited onto several metal surfaces, under different deposition conditions. Subsequent methods were used to demonstrate the preparation-dependent change in the surface morphology and, additionally, their influence on the anticorrosion activity of the SAM layers. The AFM technique allowed the presentation of the change in the surface morphology of different metals caused by the coatings as well as by the illumination/irradiation. All results were demonstrated by 3D imaging, by section analysis and by roughness parameters, too.

The SAM nanofilm behaviors were characterized in the case of two types of corrosion: the general corrosion measurements were carried out in aerated sodium perchlorate and the pitting corrosion in sodium chloride solution.
i. **Influence of the SAM layers on the general corrosion**

The immersion of uncoated sample in perchlorate solution resulted in rough surface. This was especially visible on aluminum surface. But both amphiphile under investigation could effectively control the general corrosion. The fluorophosphonic SAM nanolayer was more effective, it did not allow approaching the aerated electrolyte to the metal surface and so, the corrosion was efficiently inhibited. The diminution of general corrosion was demonstrated by 3D images and characterized numerically by section analysis.

ii. **Influence of the SAM layers on the pitting corrosion**

The sodium chloride electrolyte causes pitting corrosion. The question arose: can these SAM layers control only the general corrosion or they are active against the pit formation. The pitting inhibition activities of both SAM amphiphiles were checked on different metals. It was clearly demonstrated that the metal surfaces without coatings suffered pitting corrosion. The fluoro amphiphile SAM-covered solids were almost intact on carbon steel and on aluminum. Important to mention, that this layer could control the pit formation, even in case of short layer formation time. The undecenyl nanolayer was less active but it also decreased the pitting corrosion. It was proved not only by the 3D images but by the section analysis: the deepness of the pits informs us about the seriousness of corrosive effect.

iii. **Anticorrosion effectiveness of the post-treated undecenyl amphiphile in SAM nanolayer**

The same technique, i.e. the AFM proved the usefulness of the irradiation: the SAM covered metal surface built from undecenyl phosphonic acid either after UV treatment or after irradiation can much better control the pit formation even at lower irradiation (2kGy); but the efficiency of the post-treated nanofilms are much better after 20 kGy irradiation. The chloride ions cannot destroy the surfaces, it stayed very smooth.

iv. **Change in roughness parameters measured on surfaces without and with coatings, before and after corrosion tests**

The AFM software furnishes three different roughness parameters ($R_q$: rout mean square roughness, $R_a$: roughness average, $R_{\text{max}}$: maximum height of the profile) on the visualized
surfaces. All these numbers unequivocally showed that the nanocoatings decreased the surface roughness. Both electrolytes caused significant increase in the roughness parameters on uncoated metals. The presence of SAM nanolayers decreased the roughness parameters not only before, but after immersion into the both electrolytes. The highest decrease was achieved when the undecenyl SAM layer was irradiated by 20kGy and then immersed into the chloride solution.

6.2.3 Demonstration of the anticorrosion activity of SAM-coated metal surfaces by electrochemical techniques

Potentiodynamic measurement and electrochemical impedance spectroscopy helped in the characterization of the anticorrosion activity of the SAM layers. The first type of measurement informed me about the type of corrosion (anodic or cathodic inhibition). The EIS could answer the question how the layers control the corrosive deterioration (inhibition of charge transfer). The influence of the layer formation time on the corrosion inhibitive efficacy was demonstrated by both electrochemical measurements and proved that with increasing layer formation time the anticorrosion efficiency increased. In the case of fluorophosphonic amphiphile coating the anticorrosion efficiency reaches already very high value when the layer formation time is shorter (2 h). At the undecenyl phosphonic acid the time-dependent increase in the corrosion inhibiting efficacy is slower; to get a more effective anticorrosion nanocoating needs longer time.

According to the Tafel curves, at both SAM layers the $E_{corr}$ values shifted into the positive direction and the anodic current decreased indicating that these nanofilms control the metal dissolution, they behave like anodic inhibitors. According to the EIS results the increased anticorrosion effectiveness of the layers are due to increase in the polarization resistance of the nanolayers.

I have found an interesting relationship between the layer formation time and the efficiency: both amphiphiles follow Langmuir-type correlation (generally not the layer formation time, but the concentration is correlated with the efficiency in Langmuir isotherm). The undecenyl phosphonic acid adsorption takes more time (as it was already shown by other techniques). The Langmuir-type correlation was confirmed by the graph when the layer formation time/surface
coverage was depicted as a function of the layer formation time. At both SAM layers it gave straight lines proving the validity of the Langmuir correlation.

When the pH of electrolyte was changed, both electrochemical techniques proved that the efficiency of either SAM layers is pH-dependent. At low pH values these SAM film cannot control the corrosion, the metal dissolution increases. The explanation could be that the head group structure of both amphiphiles in SAM films changes in acidic solution and occupies smaller place because of their pK values, and the so-called tridentate bonding (which is effectual for the SAM layer formation) is altered to di- and mono-dentate bonding. So, more free locations are available for the deteriorating aqueous solvent components and, as a consequence, for the metal dissolution. Around neutral pH values and above them the corrosion current decreases drastically, i.e. the anticorrosion effect increases. Both the electrochemical impedance spectroscopy and the potentiodynamic technique gave similar results. When the layer formation time increased, the anticorrosion efficacy increased. A saturation type correlation was found between the layer formation time and the polarization resistance values.
References

83. Schematic representation of the AFM set-up (http://web.mit.edu/cortiz/www)
85. Schematic of Michelson Interferometer https://www.researchgate.net/figure/Schematic-figure-of-Michelson-interferometer_fig1_306050242
96. IS.Alexandar,M.Pharm, Adsorption pharmaceutical analysis Pin:636008 https://www.slideshare.net/alexmpharm/adsorption-pharmaceutical-analysis.
- **Thesis points**

1. I demonstrated that the time of layer formation had a major influence on the performance of the coated surface and it has a more decisive impact on the nanolayer structure than the increase in the amphiphile concentration. (T. Abohalkuma, F. Shawish, J. Telegdi; “Phosphonic acid derivatives in self assembled layers against metal corrosion” *International Journal of Corrosion and Scale Inhibition* 3(3) (2014) 151-159).

2. I have proven by contact angle values as well as by the decreased anticorrosion activity that the double bond in the undecenyl phosphonic acid nanolayer disturbs the SAM layer structure. These amphiphilic molecules in SAM layers were less active than that of the fluorophosphonic amphiphile. This is due to the less ordered structure in the case of the undecenyl amphiphile. At the same time, the influence of hydrophobic fluorine atoms in the amphiphile on the increased hydrophobicity was demonstrated, which resulted in a very condensed molecular layer that was correlated with the very effective corrosion inhibition. (Talah Abohalkuma, Abdul Shaban, Judit Telegdi “Corrosion Processes Controlled by Phosphonic Acid Nano-layers” *Periodica Polytechnica Chemical Engineering* 60(3), 165-168, (2016) DOI: 10.3311/PPch.8260).

3. In case of the undecenyl phosphonic acid SAM layer, (which has a less ordered structure and, as a consequence, it can control the corrosion processes with less activity,) I was who first modified the SAM layer by post-treatments (by illumination and irradiation); both types of handleings changed not only the surface wettability (because of the polymerization of the double bonds in the nanolayer), but the anticorrosion efficacy was also significantly increased. In the case of the UV light a longer-time treatment (1 h) could successfully modify the layer structure. As to the irradiation, already 2 kGy of γ-ray altered the structure of the nanofilm that resulted in an increased prevention of the metal surface against the pitting corrosion; a much higher dose (20 kGy) produced already a compact nanolayer that was effective against the general and the pitting corrosion. . (J. Telegdi and T. Abohalkuma, “Influence of the nanolayer’ post-treatment on the
4. The importance of the oxide layer was demonstrated in my work. It was proven that an increased oxide layer thickness on aluminum adsorbs the phosphonic acid amphiphiles more effectively than a thinner one like in the case of the carbon steel oxide layer. On the other hand, I have shown the importance of the alloying elements on the surface characteristics: the different wettability of the two stainless steels (304 and 3016) was confirmed with the differences in the surface oxide layer composition. (Judit Telegdi, Giorgio Luciano, Soumitro Mahantry, Talah Abohalkuma, “Inhibition of aluminum alloy corrosion in electrolytes by self assembled fluorophosphonic acid molecular layer” Materials and Corrosion, 67(10) 1027-1033 (2016) DOI: 10.1002/maco.201508304).

5. I proved that the SAM layers formed by fluorophosphonic acid show anticorrosion performance towards both general and pitting corrosion while those formed by undecenyl phosphonic acid are active mainly against the general corrosion. (Talah Abohalkuma, Judit Telegdi “Corrosion protection of carbon steel by special phosphonic acid nano-layers” Materials and Corrosion 66(12) 1382-1390 (2015) DOI: 10.1002/maco.201508304).

6. On the basis of electrochemical results, I have found special correlation between the layer formation time and the anticorrosion activity, which follows Langmuir-type adsorption; in my case, instead of the amphiphile concentration, the layer formation time and the anticorrosion activity is correlated. (J. Telegdi and T. Abohalkuma, “Influence of the nanolayer’ post-treatment on the anticorrosion activity”, International Journal of Corrosion and Scale Inhibition, 7 (3) 352–365 (2018) DOI: 10.17675/2305-6894-2018-7-3-6).

7. The importance of the environmental pH was demonstrated and explained: the SAM layer formed at 24h by fluorophosphonic acid and by undecenyl phosphonic acid could not keep their anticorrosion activity at pH=3, but these layers are very active at neutral and
- Publications

i. Journal papers:


- Cited by


IF: 1.37
• **Cited by**


IF: 0.13
4. Judit Telegdi, Giorgio Luciano, Soumitro Mahantry, 
Talah Abohalkuma, “Inhibition of 
aluminum alloy corrosion in electrolytes by self assembled fluorophosphonic acid 
IF: 1.4

- Cited by


ii. Conference publication:

- Oral presentations


8. Six presentations at the end of each of the first six semesters.

- Poster presentations:


4. T. Abohalkuma and J. Telegdi “Special phosphonic acids used in nanolayers against corrosion” at the EUROCORR 2013 September 1 - 5, 2013, Estoril, Portugal.