Electrochemical Biosensor Arrays Utilising Bacteria and Aptamer Nano-bioreceptors for Toxic Chemicals Detection

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Electrochemical Biosensor Arrays Utilising Bacteria and Aptamer Nano-bioreceptors for Toxic Chemicals Detection

By

Hisham Faiadh Mohammad Abu-Ali

Thesis submitted in partial fulfillment of the requirements of Sheffield Hallam University for the degree of Doctor of Philosophy

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2019
DECLARATION

I hereby declare that this thesis submitted for the degree of PhD is the result of my own research and that this thesis has not been submitted for a higher degree to any other university or institution

Signed
DEDICATION

To my parents, specially my dearest father
To my mother soul your sacrifices shan't be forgotten
To my beloved and soulmate wife (Hala), and my angels (Um Al-Baneen and Mohammad)
Without your support and patience throughout my study,
This work would never has seen light

Hisham
ABSTRACT

This work was dedicated to development of novel biosensing technologies for detection of toxic chemicals, such as heavy metals, pesticides and petrochemicals, which possess a serious threat to humans and all living organisms in our planet nowadays. This was the main motivation for research in such important field. In the present work a novel approach in detection of heavy metal salts (HgCl$_2$, PbCl$_2$, ZnCl$_2$ and CdCl$_2$), pesticides (atrazine, simazine, DDVP), and petro-chemicals (hexane, octane, pentane, toluene, pyrene and ethanol) dissolved in water was proposed. It is based on a concept of inhibition sensor array utilising different whole bacteria cells. The main aim of this project is to develop novel, simple and cost-effective biosensing technologies for in-field detection of the above pollutants in water which effectively reduce the time and cost of analysis. Electrochemical detection appeared to be the most suitable for such task.

In this project, three types of bacteria, e.g. *Escherichia coli*, *Methylococcus capsulatus* (Bath) or *Methylosinus trichosporium* (OB3b) and *Shewanella oneidensis*, were selected because of their different inhibition patterns. The concentration of live bacteria (which is an indicator of the presence of pollutants) was first characterised by the optical analytical methods of optical density OD$_{600}$, fluorescence microscopy and flow cytometry. The main findings of this study were the facts that *E. coli* (K12 strain, gram-negative bacteria) are very sensitive to all above mentioned pollutants; methanotrophic bacteria (*Mc. capsulatus* Bath & *Ms. trichosporium* OB3b) appeared to be more resistant to petrochemicals; while *S. oneidensis* (MR-1 strain, gram negative bacteria) are more tolerant to heavy metals. A series of AC and DC electrochemical measurements were carried out on the same bacteria samples. As a first step, a correlation between optical and electrochemical characteristics of bacteria concentration in solution was established. The study of the effect of heavy metals, pesticides and petrochemicals on DC and electrical characteristics of bacteria in suspension revealed a similar inhibition pattern as was found in optical study. Then a similar study was carried out on samples of bacteria immobilized on the surface of screen-printed electrodes, which is more suitable for sensing applications. The results of DC (cyclic voltammograms) and AC (impedance spectroscopy) measurements were consistent with previous studies. A possibility of pattern recognition of pollutants by their inhibition effects on the selected bacteria was found. The classes of pollutants, e.g. heavy metals, pesticides, and petrochemicals, can be identified from pseudo-3D graphs of responses of the three sensing channels, e.g. electrodes with different immobilized bacteria. Much more accurate assessment of pollutants was achieved with Artificial Neural Network (ANN) software which was developed using MatLab. ANN programme was capable of both the identification of pollutants with 91% accuracy and rough estimation of their concentrations in five bands from 0.01 ng/ml to 1000 ng/ml (ppb). The developed bacteria sensor array could be suitable for simple, inexpensive, and quick preliminary in-field detection (screening) of water samples. The suspected highly contaminated samples could be easily identified and passed to specialized laboratories for further more detailed testing. In such way, the time and cost of analysis could be substantially reduced.

In addition to the inhibition sensor array utilising non-specific bio-receptors such as bacteria, the electrochemical detection of heavy metal ions (Hg$^{2+}$ and Pb$^{2+}$) was attempted using novel highly specific aptamer bio-receptors labelled with redox groups. Such experiments were successful; the above metal ions in very low concentrations down to 1 pg/ml (or 1 ppt) were detected using both cyclic voltammograms and impedance spectroscopy. The affinity of the aptamers used was found to be very high and similar to that of antibodies. Additional advantages of aptamers were their high stability and simple recovery by thermo-cycling. Considering fast evolvement of aptamer research, their advantages and low cost, the development of aptasensor arrays for accurate detection of large number of pollutants is possible in near future.
ACKNOWLEDGEMENTS

It is a difficult task to thank all the people who made this PhD thesis possible with so few words. However, I will try to do my best to extend my great appreciation to everyone who helped me scientifically and emotionally throughout this study. First of all, I would like to thank my god (Allah) for giving me the patience and stamina to overcome the difficulties that faced me during my PhD study. Secondly, I would like to acknowledge my county (Iraq) and the financial sponsor, Ministry of Higher Education and Scientific Research in Iraq and University of Basrah, Faculty of Science, Biology Department. Also I gratefully acknowledge the Iraqi Cultural Attaché in London for their support during my PhD research. I would like to express my sincere gratitude and thanks to my director of studies in Materials Engineering Research Institute (MERI), Professor Alexei V. Nabok for his excellent supervision, guidance and support during the study period, and for his discussion and interpretation of various aspects of science, experimental procedures and the results analysis. Without his support and encouragement, this work would not have been possible. My deepest appreciations and huge thanks go to my gorgeous supervisor, Professor Thomas J. Smith the head of research group of Molecular Microbiology in BioMolecular Research Centre (BMRC), for his advice, help, discussion and suggestion. My special thanks and love go to lovely wife and my children. My sincere thanks must go to Dr. Jim Yong (BMRC), for his help with fluorescence microscope measurements. Many thanks are also due to Dr. Sarah Small (BMRC) for her essential help and discussion about flow cytometer measurements. In addition, I would like to thanks Dr. Tim Nichol (BMRC), for his help and advice with molecular microbiology laboratory and bacteria samples preparation. Also big thanks go to Dr. Mohammad Akram Khan, for his essential help about the chemicals preparation. Many thanks must go to MERI reception staff for the help and assistance especially Corrie Houton, Amy McNally, Gail Hallewell and Rachael Toogood. I would also like to thank all technical staff of Sheffield Hallam University (MERI) and (BMRC) training team and workshop, especially Mr. Paul Allender for training and preparing the SEM and AFM imaging process measurements and Mr. Michael Cox for ICP-MS training and measurements. Finally I would like to thank all my brilliant Iraqi and English friends, PhD students at MERI and BMRC also my colleagues from University of Sheffield.

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<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td><em>E. coli</em></td>
<td><em>Escherichia coli</em> bacteria</td>
</tr>
<tr>
<td><em>S. oneidensis</em></td>
<td><em>Shewanella oneidensis</em> bacteria</td>
</tr>
<tr>
<td><em>Mc. Capsulatus</em></td>
<td><em>Methylococcus capsulatus</em> bacteria</td>
</tr>
<tr>
<td><em>Ms. trichosporium</em></td>
<td><em>Methylosinus trichosporium</em> bacteria</td>
</tr>
<tr>
<td>OD600</td>
<td>Optical Density at 600nm wavelength of light</td>
</tr>
<tr>
<td>UV/Vis</td>
<td>UltraViolet/Visible (spectroscopy)</td>
</tr>
<tr>
<td>MTB</td>
<td>Methanotrophic bacteria</td>
</tr>
<tr>
<td>K12</td>
<td>Strain of <em>E. coli</em> bacteria</td>
</tr>
<tr>
<td>MR-1</td>
<td>Strain of <em>S. oneidensis</em> bacteria</td>
</tr>
<tr>
<td>Bath</td>
<td>Strain of methanotrophic bacteria</td>
</tr>
<tr>
<td>OB3b</td>
<td>Strain of methanotrophic bacteria</td>
</tr>
<tr>
<td>DC</td>
<td>Direct Current</td>
</tr>
<tr>
<td>AC</td>
<td>Alternative Current</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleic Acid</td>
</tr>
<tr>
<td>BTEX</td>
<td>Benzene, Toluene, Ethylbenzene and Xylene</td>
</tr>
<tr>
<td>PAHs</td>
<td>Polycyclic Aromatic Hydrocarbons</td>
</tr>
<tr>
<td>PAH</td>
<td>Poly Allyamine Hydrochloride</td>
</tr>
<tr>
<td>pH</td>
<td>Measure of the acidity of an aqueous solution</td>
</tr>
<tr>
<td>DDVP</td>
<td>Dichlorvos (commercial pesticide)</td>
</tr>
<tr>
<td>ANN</td>
<td>Artificial Neural Network</td>
</tr>
<tr>
<td>PCA</td>
<td>Principal Component Analysis</td>
</tr>
<tr>
<td>LbL</td>
<td>layer-by-layer (deposition)</td>
</tr>
<tr>
<td>CV</td>
<td>Cyclic Voltammogram</td>
</tr>
<tr>
<td>EIS</td>
<td>Electrochemical Impedance Spectroscopy</td>
</tr>
<tr>
<td>GLC</td>
<td>Gas Liquid Chromatography</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>IDAM</td>
<td>Interdigitated Array Microelectrodes</td>
</tr>
<tr>
<td>NMS</td>
<td>Nitrate Mineral Salts</td>
</tr>
<tr>
<td>PEM</td>
<td>Poly Electrolyte Multilayers</td>
</tr>
<tr>
<td>ROS</td>
<td>Reactive Oxygen Species</td>
</tr>
<tr>
<td>LPO</td>
<td>Lipid Per Oxidation</td>
</tr>
<tr>
<td>H$_2$O$_2$</td>
<td>Hydrogen Peroxide</td>
</tr>
<tr>
<td>dsDNA</td>
<td>double strand DNA</td>
</tr>
<tr>
<td>ssDNA</td>
<td>single strand DNA</td>
</tr>
<tr>
<td>SPGEs</td>
<td>Screen printed gold electrodes</td>
</tr>
<tr>
<td>BESs</td>
<td>Bio-electrochemical systems</td>
</tr>
<tr>
<td>AAS</td>
<td>Atomic Absorption Spectroscopy</td>
</tr>
<tr>
<td>AES</td>
<td>Atomic Emission Spectroscopy</td>
</tr>
<tr>
<td>ICP-MS</td>
<td>Inductively Coupled Plasma Mass Spectroscopy</td>
</tr>
<tr>
<td>CVAFS</td>
<td>Cold Vapours Atomic Fluorescence Spectroscopy</td>
</tr>
<tr>
<td>HPLC</td>
<td>High-Performance Liquid Chromatography</td>
</tr>
<tr>
<td>GC-MS</td>
<td>Gas Chromatography–Mass Spectrometry</td>
</tr>
<tr>
<td>LC-MS</td>
<td>Liquid Chromatography–Mass Spectrometry</td>
</tr>
<tr>
<td>IFM</td>
<td>Infinite Focus Microscopy</td>
</tr>
<tr>
<td>AFM</td>
<td>Atomic Force Microscopy</td>
</tr>
<tr>
<td>SEM</td>
<td>Scanning Electron Microscopy</td>
</tr>
<tr>
<td>TEM</td>
<td>Transmission Electron Microscopy</td>
</tr>
<tr>
<td>US EPA</td>
<td>U.S. Environmental Protection Agency</td>
</tr>
<tr>
<td>IARC</td>
<td>International Agency for Research on Cancer</td>
</tr>
<tr>
<td>SELEX</td>
<td>Systematic Evolution of Ligands by Exponential Enrichment</td>
</tr>
<tr>
<td>QCM</td>
<td>Quartz Crystal Microbalance and</td>
</tr>
<tr>
<td>SAW</td>
<td>Surface Acoustic Wave</td>
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**Journals Publication**


**Conference Publications**

- **Hisham Abu-Ali**, A. Nabok, T. Smith and M. Al-Shanawa, Inhibition Biosensor Based on Bacteria for Environmental Pollution Detection Using Optical and Electrochemical Measurements (Poster), 18-20 May 2016, **MERI Student Symposium**, Sheffield Hallam University, Sheffield, **UK**.

- **Hisham Abu-Ali**, A. Nabok, T. Smith and M. Al-Shanawa Inhibition Biosensor Based on DC and AC Electrical Measurements of Bacteria Samples, **Biosensor 2016 Congress** (Poster), 25-27 May 2016, Gothenburg - **Sweden**.

- **Hisham Abu-Ali**, A. Nabok, T. Smith and M. Al-Shanawa, Using Optical and Electrochemical Measurements for Development Inhibition Biosensor Based on Bacteria for Environmental Pollution Detection (Poster), 16 December 2016, **MERI and BMRC Student Symposium**, Sheffield Hallam University, Sheffield, **UK**.

- **Hisham Abu-Ali**, A. Nabok, T. Smith and M. Al-Shanawa,(2017) Development of Novel Inhibition Biosensor Based on Bacteria for Environmental Pollutants Detection , **MERI Proceeding** , 16th -17th May (1), 6-8. Sheffield Hallam University, Sheffield, **UK**.

- **Hisham Abu-Ali**, A. Nabok, T. Smith Development of Novel Inhibition Biosensor Based on Bacteria for Environmental Pollution Detection (Oral), 23-24 May 2017, **MERI Student Symposium**, Sheffield Hallam University, Sheffield, **UK**.

- **Hisham Abu-Ali**, A. Nabok, T. Smith and M. Al-Shanawa, Development of a Novel Inhibition Biosensor Based on Immobilized Bacteria for Environmental Pollution Detection Using Optical and Electrochemical Measurements (Poster), **Biosensing Technology Congress**, 05-11 May 2017, Riva Del Garda - **Italy**.


- **Hisham Abu-Ali**, A. Nabok, T. Smith, The First Iraqi Student Conference in UK, Sheffield, University of Sheffield, 29 September 2017, Sheffield - **UK**.


Hisham Abu-Ali, A. Nabok, T. Smith Application of artificial neural network for preliminary detection of water pollution using inhibition electrochemical sensor array based on immobilized bacteria and modified screen-printed gold electrodes (Poster), 14 December 2018, MERI and BMRC Student Symposium, Sheffield Hallam University, Sheffield, UK.


PRIZES and AWARDS

1. The best presentation and paper placing in The 20th International Conference of Environmental Microbiology, (Zurich, 15th-16th January 2018, Switzerland).

2. The best poster and paper placing in The 21st International Conference of Lab on a chip device, system and technology (Barcelona, 23rd-24th May 2019, Spain).

3. Award from the Iraqi Cultural Attaché as one of excellence organizers in the First Iraqi Student Conference in UK, Sheffield, University of Sheffield (Sheffield, 29th September 2017, UK).

4. Award from the Iraqi Minister of Higher Education and Scientific Research as one of the best 20 student in the UK during the PhD study period. (London, 18th January 2018, UK).
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CHAPTER 1 INTRODUCTION

The need of biosensors and the use of sensing and biosensing technology approach are outlined in this chapter, also the invention history of electrochemical biosensor and classification of biosensors are covered in this chapter. The concept of the proposed inhibition biosensor based on bacteria, and inhibition biosensor array, has been outlined. At the end of this chapter, the aims and objectives of this research are given.

1.1 The needs of biosensors

The explosive development of industrial and agricultural activities contaminated the environment with large number of toxic chemicals, particularly, heavy metals, pesticides, petro-chemicals and BTEX compounds (which refer to benzene, toluene, ethylbenzene and xylene). High concentrations of these toxic chemical were observed at industrial and agricultural areas. The above chemicals considered among the most abundantly produced chemicals in the world spread in the atmosphere and aquatic environment, and have negative impacts on all the living organisms which led to call for fast and cost-effective analytical techniques to be used for extensive in field monitoring programs [1-5]. The detection of the above toxic chemicals in low concentrations is a quite difficult task, though not impossible and can be achieved with the existing advanced analytical methods such as atomic absorption or atomic emission spectrosopies (AAS, AES), inductively coupled plasma mass spectroscopy (ICP-MS), cold vapours atomic fluorescence spectroscopy (CVAFS) and high-performance liquid chromatography (HPLC). These methods are extremely sensitive but expensive, requiring specialized laboratory conditions and highly trained personnel [6-9]. As a result, both the time and cost of analysis become very high.

An alternative approach to those sophisticated methods is based on the use of biosensors, which could be much simpler, easy-to-use, and inexpensive. The main problem of biosensors, however, is the selection of bio-receptors which actually provide the function of recognition of target analyte molecules. Typical bio-receptors used in biosensors, i.e. enzymes, antibodies, aptamers, nucleic acids and peptides, can easily provide such functionality [10-14]. However the traditional biosensing approach may
struggle with a difficult task of detecting a large number of these toxic chemicals in a complex natural environment because every analyte may require a specific receptor. In this context, inhibition biosensors and biosensor array appear as suitable alternative to traditional biosensors.

### 1.2 Sensing and biosensing systems

Sensing and biosensing technologies have been expanded since 1962, when Clark and Lyons’ invented the first electrochemical biosensor for blood glucose detection [15], Chen and Chzo, were differentiated between sensor and biosensor, depending on their active recognizing material immobilized on the surface of electrode which was used. Selecting this material differs with respect to the nature of the target analyte and the kind of effective reaction between both of them [16]. A sensor is a device that responds to a physical stimulus (such as heat, light, sound, pressure, magnetism, or a particular motion) and transmits a resulting impulse (as for measurement or operating a control) [17]. While, biosensor is an analytical device which uses living bioreceptors or biological elements such as tissue, microorganisms, organelles, whole cell, enzymes, antibodies, nucleic acids to measure the presence of chemical substance by generating signals proportional to detect the concentration of an analyte in the specimens [18]. Figure (1-1) presents the scheme of biosensor application areas.
Heavy metals, pesticides, and petrochemicals possessing serious threat to humans and living organisms are of the main concern for the environmental security, nowadays. The most common sources of environmental pollution are manufacturing, automotive, agricultural, chemical, and medical industries. For instance, three of the most common heavy metals released from road travel are zinc, copper, and lead, accounting for at least 90% of the total metals in road runoff [19, 20].

These analytes do not remain where they originate. They can be transported to different locations in a number of different ways. Some of these toxic compounds (e.g. heavy metals, pesticides and hydrocarbons) can evaporate and drift away by winds before precipitating as rainfall. In addition, runoff from agricultural and urban areas into drainage pipes and sewers also contributes to significant pollution of surface and ground water with this kind of pollutants. The rain droplets included soluble heavy metal salts with the highest concentrations were zinc (Zn, 200 μg L\(^{-1}\)), iron (Fe, 88 μg L\(^{-1}\)), and
lead (Pb, 77 μg L\(^{-1}\)). TEM reveals that 76\% of cloud drops include metal particles that range from 50 nm to 1 μm diameter with a median diameter of 250 nm [21, 22]. A study from Switzerland revealed that much of the rain in Europe contains high levels of dissolved pesticides, actually 4 μg/l of 2,4-dinitrophenol, that it would be illegal to supply this water for drinking purposes [23]. A field conditions study in Hungary revealed the presence of 154 μg/l of atrazine, 89.1 μg/l of acetochlor, 47.4 μg/l of propisochlor, and 0.139 μg/l of chlorpyrifos in runoff water [24].

1.3 Biosensors and biosensor arrays

The emergent needs for a portable, rapid, sensitive, and specific screening instrument for multiple pollutants detection at the site of sample collection in the field could be solved via biosensors and biosensor arrays. Perumal and Hashim (2014), were defined biosensor array as a device composed of three elements (biological molecules, bioreceptors and transducers) which are intimately associated [25]. There are five of the most important features that must be available in the biosensor array which are (i), sensitivity; (ii), reproducibility; (iii), repeatability; (iv), stability and (v), simplicity in the procedure of surface modification and biological molecule immobilization [26]. The use of biosensor arrays have several very significant advantages over using a single biosensor for such applications lie in the following fact (1) Biosensor array adds new dimensions to the observation, helping to estimate more parameters and improve the estimation performance [27]. (2) The number of analyte which can be detected simultaneously can be expanded as need dictates and specific analyte become available (3) The biosensor arrays and tracer reagents are reusable if no target agent binds to the array surface. This feature significantly decreases the cost and operational burden for the user and simplifies automation for extended monitoring applications [28]. (4) The biosensor array is simple to use. It is easily portable for first responder applications. The insertion of the sensor array, tracer reagents and samples is very simple with no requirement for alignment operations by the user [29]. (5) The biosensor array is a low-cost system which can be made even more cost effective with mass production. (6) The biosensor array can be easily adapted for continuous monitoring operations by integration with a computer-controlled sampler to format automatic analytical system.
Because of these advantages, more and more biosensor arrays are applied in varied areas including environmental monitoring [30]. (7) The identification of many types of pollutants in the environment and the evaluation of their concentration is a much more difficult task which is impossible to solve using a single type of biosensor. However, the biosensor array approach utilising several types of biological materials being inhibited differently by different types of pollutants could solve the problem of environmental pollution detection.

1.4 The history of electrochemical biosensor

Electrochemical biosensors comprise potentiometric, amperometric, and impedimetric sensing techniques, with amperometric sensors which being the first type of biosensors described in 1953. Electrochemical biosensors have subsequently become the most developed group with greatest commercial success, largely due to amperometric glucose detection in diabetes monitoring. Their key advantages are low cost, point-of-care testing, and miniaturization capacity [31]. Electrochemical biosensor is a device composed of two intimately associated elements, a bio-receptor and transducer as shown in Figure (1-2) which presents this scheme. The electrochemical biosensor is the first form of all biosensor types and started to be common in labs of chemical industries, and many other searching fields, the history of electrochemical biosensor began when the pH thin glass electrode invented by Max Cremer, which was the oldest electrochemical sensor in 1906. After that, Søren Peder Lauritz Sørensen demonstrated the concept of pH (hydrogen ion concentration) in 1909, and the first electrode for pH measurements was brought to the world in 1922 by W.S. Hughes [32]. During the period from 1909 to 1922, Griffin and Nelson, demonstrated the first immobilisation of the enzyme invertase on aluminium hydroxide and charcoal [33], and then Leland Clark, was invented the first electrochemical biosensor for detection of blood oxygen in 1956. He is known as the ‘father of biosensors’, and his invention of the oxygen electrode bears his name: “Clark electrode” [34].

The feasibility and their recognition ability make electrochemical sensors a good candidate for application in many disciplines such as food, biomedical, environmental,
agricultural, and industrial fields. That is owing to their precise, cost effective, fast, and reliable detection of many organic and inorganic compounds. However, further development of these biosensors requires multidisciplinary research in the fields of material science, electronics, and computer science to meet the emergent needs in different fields [35].

**Figure 1-2:** Scheme of electrochemical biosensor: (A) bio-receptors, (B) modified SPGE, (C) DropSens μStat equipment (D) cyclic voltammogram.

The demonstration of an amperometric enzyme electrode for the detection of glucose by Leland Clark in 1962 was followed by the discovery of the first potentiometric biosensor to detect urea in 1969 by Guilbault and Montalvo Jr [36]. Eventually, in 1975 the first commercial electrochemical biosensor was developed by Yellow Spring Instruments (YSI). Table 1-1 shows the historical overview of biosensors in the period 1916 till 2018.
Table 1-1: Historical overview of biosensors development.

<table>
<thead>
<tr>
<th>Year</th>
<th>The invention and development event</th>
</tr>
</thead>
<tbody>
<tr>
<td>1916</td>
<td>First report on the immobilization of proteins [37].</td>
</tr>
<tr>
<td>1922</td>
<td>First glass pH electrode [38].</td>
</tr>
<tr>
<td>1956</td>
<td>Invention of the (Clark) oxygen electrode [39].</td>
</tr>
<tr>
<td>1962</td>
<td>First description of a biosensor: an amperometric enzyme electrode for glucose (Clark) [40].</td>
</tr>
<tr>
<td>1969</td>
<td>First potentiometric biosensor: urease immobilized on an ammonia electrode to detect urea [41].</td>
</tr>
<tr>
<td>1970</td>
<td>Discovery of ion-sensitive field-effect transistor (ISFET) by Bergveld [42].</td>
</tr>
<tr>
<td>1975</td>
<td>Fibre-optic biosensor being carbon dioxide and oxygen detection by Lubbers and Opitz [43].</td>
</tr>
<tr>
<td>1975</td>
<td>First commercial biosensor for glucose detection by YSI [44].</td>
</tr>
<tr>
<td>1975</td>
<td>First microbe-based immunosensor by Suzuki et al. [45].</td>
</tr>
<tr>
<td>1982</td>
<td>Fibre-optic biosensor for glucose detection by [46].</td>
</tr>
<tr>
<td>1983</td>
<td>Surface plasmon resonance (SPR) immunosensor by Liedberg et al. [47].</td>
</tr>
<tr>
<td>1984</td>
<td>First mediated amperometric biosensor: ferrocene used with glucose oxidase for glucose detection [48].</td>
</tr>
<tr>
<td>1990</td>
<td>SPR-based biosensor by Pharmacia Biacore [49].</td>
</tr>
<tr>
<td>1992</td>
<td>Handheld blood biosensor by i-STAT [50].</td>
</tr>
<tr>
<td>1996</td>
<td>Glucocard launched [51].</td>
</tr>
<tr>
<td>1996</td>
<td>Abbott acquires MediSense for $867 million [52].</td>
</tr>
<tr>
<td>1998</td>
<td>Launch of LifeScan FastTake blood glucose biosensor [53].</td>
</tr>
<tr>
<td>1998</td>
<td>Merger of Roche and Boehringer Mannheim to form Roche Diagnostics [54].</td>
</tr>
<tr>
<td>1999-2018</td>
<td>BioNMES, Quantum dots, Nanoparticles, Nanocantilever, Nanowire and Nanotube [55].</td>
</tr>
</tbody>
</table>

Ever since the development of the i-STAT sensor for glucose remarkable progress has been achieved in the field of biosensors. The field is now a multidisciplinary area of research that bridges the principles of basic sciences (physics, chemistry and biology) with fundamentals of micro/nanotechnology, electronics and applicatory medicine. The database ‘Web of Science’ has indexed over 84 000 reports on the topic of ‘biosensors’ in the last ten years. According to the categories typically used, the “enzyme electrode” introduced by Clark and Lyons in 1962 [56, 57] was an amperometric biosensor. This milestone in biosensor development was followed by other electrochemical biosensors, but it was not until ten years later that biosensors based on other transduction principles were published. Electrochemical biosensors, for instance, are meanwhile comparatively easy to miniaturize, which is one of the reasons for their widespread availability. In fact, detectors for biosensors used today depend mainly on electrochemical transduction, followed by optical and acoustic effects [58]. Thermal transduction is less frequently used, as are magnetic effects. However, the latter is increasingly employed as a separation tool in bioanalytical assays [59, 60].
1.5 Bio-electrochemical systems (BESs)

Electrochemical properties of biomaterials, such as whole cells and bacteria, were studied extensively in the past [61]. Recently, the subject of electrochemical characterisation of cells came back because of recent development in bio-cell sensors [62]. The electrochemical bio-cell sensor is a relatively young field, but is now achieving substantial success in science, engineering, and technology. There are several advantages of electrochemical techniques: (i), the measurements can be made quickly, which refer to the environment activities and can be easily transmitted, amplified and digitized, (ii), the measurements can be carried out in the laboratory, as a portable device (portable detector). Bio-electrochemical systems (BESs) take advantage of biological objects (enzymes, microbes, plants) for the catalysis of electrochemical reactions [63]. Some examples of BES are: Microbial Fuel Cells, Plant-Microbial Fuel Cells, Enzymatic Fuel Cells, Microbial Electrolysis Cells, Microbial Electro-synthesis Cells and Microbial Desalination.

BESs have recently emerged as a promising technology for energy recovery and for providing valuable products, such as hydrogen, ethanol and other organic molecules [64]. BES appears as a promising alternative for treating different types of wastewater. BESs use whole cell biocatalysts to drive oxidation and reduction reactions at solid-state electrodes. The most widespread application is presently the microbial fuel cell, which aims to generate power or at least decrease the usage of power associated with wastewater treatment. In the slipstream of microbial fuel cells, microbial electrolysis cells have emerged recently. The versatility of the latter has notably expanded the range of applications of BESs. Key applications are wastewater treatment, sediment-based electrical power generation, value added product generation, bioremediation and detecting of the biomass (biological elements concentration).

The electrical properties of biological material have been studied using suitable instrumentation. In addition, impedance techniques have been used to study and monitoring the growth rate of organs in-vivo, whole blood and erythrocytes, cultured cell suspensions and bacterial growth [65]. The integration of impedance with biological
recognition technology for detection of bacteria has led to the development of impedance biosensors that have come to be widely used in recent years. In addition, DC and AC properties of bacteria cell have been studied and monitored.

1.6 Inhibition electrochemical biosensor scenario

In the last 20 years many researchers described the realization of inhibition biosensors for environmental pollution detection. This study has focused on the development of inhibition bacteria based electrochemical biosensors for the determination of environmental pollutants. We have indeed designed, realized and applied inhibitor bacteria based bio-sensors.

Electrochemical devices are the most common in commercial sensor development because of their simplicity of construction and therefore low cost, reliability of performance, high sensitivity with minimal concentrations of analytes down to $10^{-8}$M, and wide dynamic range (up to 4 orders of magnitude in analyze concentration). The first biosensor, i.e. the glucose sensor based on Clark’s electrode, was actually electrochemical.

Electrochemical and electrical sensors are clearly distinguishable. Electrochemical sensors are based on detection of electrochemical reactions on electrodes in solutions, while electrical sensors detect variations in electrical parameters of materials or devices (current, voltage, conductance, resistance, impedance) during molecular adsorption, most-likely in a gaseous environment (gas sensors) as well as in liquids (ion sensors). These devices will be outlined in more detail in the sections below. However, before going into analysis of different electrical and electrochemical sensors, the physico-chemical processes in the electrolyte-solid state system have to be discussed first.

In this study the bio-cell sensor that included the microorganisms (bacteria) was employed for detection of heavy metals, pesticides and petrochemicals which is considered to be a cheap (cost effective), simple (easy to use), powerless (portable) and sensitive technique. Other and more sophisticated techniques, like HPLC, GC-MS and/or LC-MS should be used as confirmation techniques whenever a “positive” sample
is detected. The use of this and analogous inhibition biosensors in environmental field can represent an important tool for environmental monitoring activity. In particular inhibition based bio- sensors, for their simplicity of realization. In recent years the research in the field of the biosensors was directed to the development of highly selective and sensitive devices, but it is our personal opinion that in environmental analysis, as well as in food analysis, the use of biosensor is not going to be “exclusive” (like it may be for instance in the case of biocompatible implantable devices to be used in clinical chemistry). The role of biosensors in the determination of environmental pollutants is in our opinion not to completely replace the traditional, more sophisticated instrumental techniques, but to represent a valid complement to them, especially in all those situations where it is necessary to carry out measures “on the spot”, reducing the overall times of analysis and minimizing the sample pre-treatment process. In such situations a biosensor (or better an array of biosensors) would be the analytical tool supplying all the necessary information to monitor, in real time, the state of pollution of the matrix under investigation. In the case of a positive response of one or more biosensors “traditional” sampling procedure will be activated to carry out more specific assays aimed to confirm and quantify more precisely the extent of each case of environmental contamination. In this field the development of inhibition based biosensors would be a very powerful tool for the screening of huge populations of samples for different classes of pollutants, thus detecting any compound belonging to the same class on the basis of the common biological effects, and representing an effective and powerful aid for the early detection of environmental contamination.

1.7 Aims and objectives

The main aim of this project is the development of novel inhibition biosensing array for the detection of environmental pollution. The study utilizes three types of bacteria samples (Escherichia coli, Methylococcus capsulatus (Bath) or Methylosinus trichosporium (OB3b) and Shewanella oneidensis) for the study of their optical and electrochemical properties under effects of petrochemicals, pesticides and heavy metals. One of the main reasons of using bacteria is their versatility in detecting different pollutants. Another reason is the cost-effectiveness of bacterial inhibition sensors.
Potentially, this work may lead to the development of novel, inexpensive, low power, and portable sensor array for early detection (preliminary screening) of petrochemicals, pesticides and heavy metals.

These aims can be achieved through the following objectives:

1. To select the suitable bacteria species and use poly-L-lysine for their immobilization protocol.
2. To utilise several optical techniques, such as Optical Density (O.D600), Fluorescence microscopy, and flow cytometry for counting the bacteria number and percentage which is affected by the environmental pollutants such as petrochemicals, pesticides and heavy metals, also establishing a correlation between optical properties of bacteria (in suspension and immobilized forms).
3. To utilize AC and DC electrochemical measurements for detection of these pollutants and to establish a correlation between optical and electrochemical properties and bacteria concentration.
4. To study the effect of heavy metal salts (HgCl₂, PbCl₂, ZnCl₂ and CdCl₂), pesticides (atrazine, simazine, DDVP), and petro-chemicals (hexane, octane, pentane, toluene, pyrene and ethanol) on optical and electrochemical properties of bacteria suspension and immobilized bacteria.
5. To develop an inhibition bacterial biosensor array prototype suitable for identification and concentration evaluation of petrochemicals, pesticides and heavy metals in water using Artificial Neural Network (ANN) data processing. This study includes a possibility of using aptamers for electrochemical detection of (Hg²⁺ and Pb²⁺) ions analysis of real samples of water from different sources.
References


CHAPTER 2 LITERATURE REVIEW

This chapter covers the toxic chemicals which cause the environmental pollution, classification of environmental pollutants i.e. heavy metals, pesticides, petrochemicals and BTEX compounds) and their spread of these toxic chemicals and their impact on the environment as well and the methods for detection of these pollutants, and the effect of these pollutants on living organisms and bacteria mass in particular. Also the more detailed description of the sensing material (bacteria) is given at the end of this chapter.

1.2 Toxic chemicals and environmental pollution

Environmental security is one of the important requirements for protecting living organisms in our planet. However, it still remains a major global challenge [1, 2]. The environmental pollution with toxic chemicals is considered as a main and the vital issue of the global biosphere. The evolution in agriculture, pharmaceutical, and chemical manufacturing is essential to fulfill the requirements and demands of the growing human population. However, such developments often caused additional pollution of environment [3]. These pollutants are spread into the different parts of the environment and have negative impacts on the living organisms [4]. Environmental pollution has been described as any natural or industrial release of the chemical, thermal, biological or radioactive elements to some part of the environment which makes a threat to the health and wellbeing of the living species [5]. Today, environmental pollution has become a significant problem which threatening the wellbeing of the living organisms. There are so many other sources of hazardous and pollutants, such as chemical plants, roads which release heavy metals, agriculture which use pesticides and chemical industry using petrochemicals and other toxic components. These sites pose a serious threat to the human been, animals, plants and microorganisms [6].
Pollutants might cause either destruction with direct visible effects on the environment, or minor destruction in the system of living organism life cycle due to disturbance of a delicate balance of the biological nutrition which are noticeable after a certain time [7]. Water pollution is of particular concern nowadays, and the needs of the methods of detecting and eliminating pollutants is growing. There are many different pollutants with petrochemicals, heavy metals, and pesticides being the most common ones. Water pollution can be defined as the presence of any chemical or microbial objects in the fresh or seawater which reduce the water quality and affect the living bodies in it. [8].

A large number of unregulated chemicals are consistently introduced into watercourses. Furthermore, several of the mining processes cause discharges of chemical waste which damages the surrounding environment. Non-conventional waste product varies from naturally inactive substances, for instance, clay and ferrous remains [9].

2.2 Detection of toxic chemical pollutants

Removal of toxic pollutants from the environment, i.e. air, soil and water should start with the detection of these harmful elements. Ecological monitoring is all about sensing the pollution which currently occurs in the air, soil and in water. The treatment of environmental pollution with toxic chemicals problem needs sequence of processes, firstly, to have reliable methods of detection and identification of pollutants and secondly, adequate remediation procedures which can be applied.

The environmental pollutants detection and monitoring in air, water and soil is very important in the overall safety and security of humans, other animals and plants. Highly sensitive traditional analytical techniques such as chromatography and inductively coupled plasma mass spectroscopy are considered time consuming, expensive and require a lot of expertise. Many of analytical techniques for the detection of toxic pollutants and chemical pollutions have been established in the last 40-50 years, [10]. Powerful analytical tools such as gas liquid chromatography (GLC) and inductively coupled plasma mass spectrometry mass (ICP-MS) can identify any chemical contamination in very low concentrations down to the part per billion (ppb). However,
very often these methods are expensive, require the use of sophisticated equipment, specialized laboratories, and highly trained personnel, and thus cannot be used for in-field trials. At the present time, the needs for low-cost, portable, easy-to-use sensor devices, which can be used even by non-specialists in field conditions, is increasing. Therefore, there is need for simple, rapid, none expensive, highly sensitive, specific and portable device for analysing environmental pollutants.

There are many methods of toxic chemical environmental pollution detection, but this study focuses on the development of novel, portable and cost-effective inhibition biosensor array for preliminary analysis (screening) of the presence of petrochemicals, heavy metals, and pesticides in water and mostly on the development of simple and rapid method of their detection.

The main problem of sensor development, however, is the synthesis of specific receptors for a huge variety of pollutants. The use of natural bio-receptors such as enzymes and antibodies or their artificial analogs such as aptamers could be the way forward, though such bio receptors are quite expensive. The idea of using bio-objects such as enzymes, whole cell, and microorganisms) which can be inhibited by pollutants is a new trend in bio-sensing. The inhibition sensor arrays are capable of differentiation and quantification of pollutants [11].

The highly sensitive electrochemical sensors dominate biosensing market, and they are the most common biosensors for detection of different analytes [12]. In addition, the inexpensive, simple design and small size make them excellent candidates for the development of portable biosensors [13]. The new idea which recently appeared is the use of microorganisms which can resist the high concentrations of petrochemicals, pesticides and heavy metals. There are several reports about biosensing applications of microorganisms such as *Escherichia coli, Pseudomonas putida, Shewanella oneidensis* bacteria and *Anabaena algae* which have the ability to pollution resistivity [14]. Recently, the utilize of whole bacteria cell as a biosensor to detect the toxins in the environment was reported with *Shewanella oneidensis* MR-1 which is a
Gammaproteobacterium, that is mean it has the ability to detoxify many kinds of heavy metals and some organic solvent [15, 16].

In this project, it was proposed to use some of these microorganisms for detection of the presence of toxic chemicals (heavy metals, hydrocarbons, and pesticides) by monitoring live and dead bacteria count. The pollution detection processes of above pollutants were developing by discovering optical and electrochemical microbial biosensors [17, 18]. A number of experimental techniques will be used in this project, such as electrochemical methods such of cyclic voltammograms using three-electrode assemblies and DropSens potentiostat, also electrochemical impedance spectroscopy, optical methods such as fluorescence microscopy, optical density and flow cytometry. These measurements will be carried out before and after exposure of the bacteria in suspension as well as immobilized bacteria on the modified screen printed gold electrode to detect the above mentioned toxic compounds. The most favourable methods will be selected to provide the dependable response and high sensitivity. Other criteria for selection of experimental methods are the cost and the sensor development suitability. The new trend of the sensor array in this project will be explored about using different types of bacteria in the same sensor chip, for example, a screen printed gold electrode containing different immobilized bacteria colonies could be used. For this purpose, the bacterial responses such as anode or cathode current, AC impedance analysis and DC cyclic voltammograms could be utilized from each well. Different levels of different pollutants concentrations will be recorded and provide a database for pattern recognition. Such sensor array approach based on ANN (Artificial Neuro Network) will be applied for recognition of pollutants and evaluation of their concentration [19- 22]. For this work three strains of bacteria were selected, namely *Escherichia coli* (*E. coli*), Methanotrophic bacteria (*Methylococcus capsulatus* (Bath) and *Methylosinus trichosporium* (OB3b)) and *Shewanella oneidensis* MR-1 (*S.oneidensis*). *E.coli* and (*M. capsulatus* (Bath) or *M. trichosporium* (OB3b) being quite sensitive to heavy metal and pesticides, and could be suitable for monitoring the contamination at a low concentration. While *S. oneidensis* is know by it is resistance to these kinds of
pollutants [23, 24]. In the meantime, preliminary results were obtained by Al-Shanawa et al (2013, 2014) [25, 26], and this study is a further development of the concept of the inhibition bacterial sensor array, so these bacterial samples will be tested under treatment with heavy metals, pesticides and hydrocarbons. In order to acquire the fundamental knowledge and understanding of the mechanism of bacteria inhibition by different pollutants, the correlation between the bacterial concentration with the optical and electrochemical properties of liquid and immobilized bacteria samples have to be established. For this purpose, several optical methods, including fluorescent microscopy, optical density, and flow cytometry will be used in this study, along with electrochemical AC and DC measurements, in order to study the effect of heavy metal salts (HgCl\(_2\), PbCl\(_2\), ZnCl\(_2\) and CdCl\(_2\)), pesticides (atrazine, simazine, DDVP), and petro-chemicals (hexane, octane, pentane, toluene, pyrene and ethanol) on the above bacteria. Electrochemical measurements are considered promising for the development of simple inhibition bacteria-based biosensors for the detection of these pollutants. The use of three (or more) types of bacteria will be lead to the development of sensor arrays utilizing the principles of pattern recognition such as (ANN) for inhibition elements detection.

2.3 Classification of toxic chemical pollutants

2.3.1 Heavy metals

Heavy metal pollution is a serious global environmental problem and among the most abundant, toxic and persistent inorganic toxic chemicals which adversely affect living organisms and cause genetic mutation [27, 28]. Due to their high atomic weight and high density they are commonly referred to trace metals; many of these trace metals (e.g. Hg, Pb, Cd and Ni) are highly toxic to humans and other living organisms, and their presence in surface water at above background concentrations is undesirable [29]. The detection of toxic metal ions in aquatic environment is an important global issue because these contaminants may have severe effects on microorganisms, plants, animals
and human, and on ecosystem in general and these metals are able to accumulate in plant leaves [30]. Among the most toxic metallic water pollutants in aquatic environment are mercury and lead [31]. The extensive use of heavy metals in industries such as transport, medical and chemical manufacturing has led to their widespread presence in the world and caused increased concern about the potential impact of these pollutants on the life cycle of living organisms and biosphere. The toxicity of heavy metals depends on several factors which include the dose, route of exposure, and type of metal, as well as age, sex, genetics, and dietary conditions of the infected living species. These metallic elements are considered to be general contaminants which are known to cause many of genetic disorders, even at low doses of exposure. In addition some of heavy metals are classified as human carcinogens according to the U.S. Environmental Protection Agency (US EPA) and the International Agency for Research on Cancer (IARC). [32]. Heavy metal pollution is a problem associated with areas of intensive industry; roadways and automobiles are now considered as one of the largest sources of heavy metal pollution. Zinc, mercury, and lead are three of the most common heavy metals released from road travel, accounting for at least 90% of the total number of different type metals in road runoff. Lead concentrations, however, have been decreasing consistently since leaded gasoline was discontinued. Figure (2-1) shows the types of heavy metals source products.

![Figure 2-1: The sources and products containing heavy metals.](image)

Smaller amounts of other metals, such as nickel and cadmium, are also found in road runoff and car exhaust [33]. Heavy metals are considered as loosely defined subset of
elements that exhibit metallic properties. These mainly include transition metals, some
metalloids, lanthanides, and actinides. Several definitions have been proposed, some
based on density, some on atomic number or atomic weight, and some on chemical
properties or toxicity. Heavy metals occur naturally in ecosystems, with large variations
in concentration [34]. During precipitation on road surfaces, most heavy metals become
bound to the road dust or other particulates or become soluble. In either case, the metals
enter the soil or water resources. Whether in the soil or aquatic environment, metals can
be transported by several processes, which are governed by the chemical nature of
metals, soil and sediment particles, and the pH of the surrounding environment. Other
common sources of metal contaminants in the environment are mining and smelting
activities; other industrial emissions and effluents; urban development; vehicle
emissions; dumped waste materials; contaminated dust and rainfall [35]. In conclusion,
there are seven major categories of sources of metals contamination of the environment.

1. Natural sources, such as surface mineralization, volcanic gasses, spontaneous
   combustions or forest fires.
2. Metal containing agricultural sprays or soil amendments.
3. Emissions from large industrial sources, such as metal smelters and refineries,
   chemical and pharmaceutical industries.
4. The disposal of wastes from mines or mills.
5. Emissions from municipal utilities, such as coal or power stations or municipal
   incinerators.
6. Emissions from transport automobiles.
7. Other relatively minor sources of contamination, such as smaller scale industries
   that process metals.

**2.3.2 Petrochemicals and BTEX compounds**

The extensive use of petrochemicals and hydrocarbon products such as (hexane, octane,
pentane, pyrene and ethanol) and BTEX compounds (e.g. benzene, toluene,
ethylbenzene and xylene) leads to the contamination of almost all environmental resources. Particularly in the zones of petrochemical production, the surface soils and water environments are exposed to contamination by the industrial products. Nowadays, the growths of urban industry process lead to greater than before release of the petrochemical products into the eco-system. Petroleum production process is distribute many of environmental pollutants which enter the water environment due to releases of manufacturing products such as urban effluents, shipping activities, offshore oil production, oil spills, fossil fuel combustion, and natural seeps [36]. Petroleum production is costly and globally degrading; most of the petroleum oil sources are associated with visible large escapes, and many oil fields are found due to natural outflows. In addition to the main use of oil as a fuel, oil is extensively used in chemical industry for production of large varieties of chemicals. Figure (2-2) shows the number of petrochemical products. To make a full assessment of the petroleum extraction processes impact on the environment, the detailed characterization of the environment, such as water, air, vegetation, and soil, is required. For this reason, the detection of petrochemical pollution is vital to give information on environmental quality.

Figure 2-2: Common commercial petrochemicals.
It was described by some researchers that petroleum extraction processes commonly mount up heavy metal release from many sources, for instance rocks, sea salt intrusion, during migration to its present reservoir, refining, transportation, and handling [37]. Wastewater of some petrochemical plants, in addition to hydrocarbons, contains chlorinated chemicals. The biological treatment; particularly by the activated sludge process has been widely used for removal of organic compounds from petrochemical wastewater. The microbial composition of the activated sludge and its activity depends on the nature and availability of petroleum hydrocarbons, nutrient composition, and other environmental conditions (pH, temperature, dissolved oxygen, mixing system, plant configuration). The microbial degraders of organopollutants of contaminated areas are normally chemoorganotrophic species, which are able to use a huge number of natural and xenobiotic compounds as carbon sources and as electron donors for the generation of energy. Numerous microorganisms, including bacteria, fungi and yeasts, predominantly aerobics, are known for their ability to degrade these organopollutants [38, 39]. The typical aerobic degrading bacteria in organopolluted site belong to a spectrum of genera and species including, *Pseudomonas sp.*, *Acinetobacter sp.*, *Alcaligenes sp.*, *Flavobacterium cytophaga* group, *Xanthomonas sp.*, *Nocardia sp.*, *Mycobacterium sp.*, *Corynebacterium sp.*, *Arthrobacter sp.*, *Comamonas sp.*, and *Bacillus sp.* [40].

### 2.3.3 Pesticides and herbicides

The environmental impacts of pesticides and herbicides depend on their toxicity, solubility, distribution and the concentration in the environment. During the 1940s, the first synthetic pesticides come to be available, making huge benefits and increasing the food manufacture. Negative effect of pesticides on the eco-system and on the life cycle of living organisms started being noticed in the early 1960s [41]. Since then, the debate on the risks and benefits of pesticides has not ceased, and a large amount of research has been conducted into the impact of pesticides on the environment. 2.5 million tons of
pesticides have been estimated and applied to the crops agriculture worldwide yearly, the amount of pesticide coming in direct contact with or consumed by target pests is an extremely small percentage of the amount applied. In most studies the proportion of applied pesticides reaching the target pest has been found to be less than 0.3%, and 99.7% went 'somewhere else' in the environment [42]. The use of pesticides and herbicides in agriculture inevitably affects non-target organisms (including humans). Undesirable side-effects may occur in some species, communities or on ecosystems as a whole. The environmental effect of pesticide applications is increasing, and it has been taken into account by regulatory bodies, leading to increased restrictions on the use of pesticides or their complete ban. Although some of these pesticides have been eliminated the most harmful agents environmentally. The inputs of fertilisers and agrochemicals were reduced since the late 1970s and there has been considerable interest in 'integrated' arable farming systems, which attempt to reduce the use of pesticides [43]. Figure (2-3) shows the types of commercial pesticides and herbicides.

![Types of pesticides and herbicides used.](figure2-3.png)

In traditional cultivation farming systems, there are several reasons interfering with the farmer’s choice to make use of a specific pesticide such as the anticipated effectiveness against the pest, the risk of phytotoxicity to the crop and the cost of the application. In combined agricultural, the ecological influence of the pesticide has to be a most important condition to take into account. Several approaches were planned to assist
agriculturalists in estimating the environmental impact of pesticides in the last few years [44].

Manuel, et al (2008) [45] defined pesticides and herbicides as any substance or mixture of substances intended for preventing, destroying, repelling, or mitigating of any pest or weed. Frank, et al (1991) [46] reported that pesticides and herbicides are the most cost-effective means of pest and weed control known to contain numerous geno-toxic compounds. Pesticides can be classified according to their target, their mode or period of action, or their chemical structure. There are different types of pesticides which are herbicides, insecticides, fungicides and nematocides. More than 500 different pesticide formulations are being used in our environment, mostly in agriculture, although the control of biological public health hazards also continues to be an important field of application. In the last 50 years, the use of pesticides has greatly increased the quantity and improved the quality of food for the growing world population. Below, the three compounds used in this work are described in more details.

Atrazine (6-Chloro-N-ethyl-N-1-methylethyl-triazine-2,4,-diamine) is one of the herbicides discovered in 1958 and used first to control the agricultural weeds in 1960 and it is still in use, occupying 83% of U.S pesticides market. An atrazine monitoring program has been established in water systems due to atrazine being discovered and treated in polluted water. In addition, the program investigated the metabolites of atrazine derivatives which have the same poisonous effect as the parent compound [47]. Several studies in the U.S on the role of atrazine in causing cancer in human and animals have been performed but they concluded this program need to regulate the using of this pesticide rather than prevent it [48].

Simazine, (6-chloro-N2, N4-diethyl-1,3,5-triazine-2,4-diamine), was one of the first compound of atrazines [49], registered in 1957. From 1990 to 1993 it was among the most widely used herbicides in the US for pre and post-emergence weed control. It is a white crystalline solid with a melting point of 226 °C, slightly soluble in water (5 mg l\(^{-1}\)) and highly soluble in organic solvents. When applied to the soil it is absorbed
by leaves and roots, causing inhibition of photosynthesis in the entire plant [50] and [51]. It is biodegradable, metabolized in plants and soil through both chemical and microbial processes.

Simazine was used to prevent and control the growth of any unwanted plant and algae in water resources. Due to these negative impacts simazine use is banned in the countries of the European Union and currently simazine is considered dangerous for the environment.

Dichlorvos or 2, 2-dichlorovinyl dimethyl phosphate (commonly abbreviated as a DDVP) is an organophosphate widely used as an insecticide to control household pests, and protecting stored product from insects [52]. The compound has been commercially available since 1961 and has become controversial because of its prevalence in urban waterways and the fact that its toxicity extends well beyond insects. The insecticide has been banned in European Union since 1998 [53].

### 2.4 Effects of toxic chemical pollutants on living organisms

Heavy metals are one of the toxic and non-biodegradable pollutants released into the environment by industrial, mining and agricultural activities [54]. The density of heavy metals is 6.0 g/cm³ while the soils density is 2.65g/cm³ but concentrations are frequently elevated; because of the contamination of soil which occurs naturally. The most important heavy metals have potential hazards and occurrences in contaminated soils are cadmium (Cd), chromium (Cr), mercury (Hg), lead (Pb), nickel (Ni) and zinc (Zn).

Cadmium’s (Cd) toxicity has been linked to reproductive disorder by affecting the number, shape and activity of sperm, in addition reducing the weight of the neonate. Moreover it is causing possible carcinogenic anomalies and seems to be a causal factor in cardiovascular diseases and hypertension [55].
Chromium (Cr) in inorganic systems occurs in several chemical forms. Only Cr (III) and Cr (VI) are significant in biological systems. Trivalent chromium is an essential nutrient component [56]. Chromium is required for carbohydrate and lipid metabolism and the utilization of amino acids [57]. The biological function of Cr is also closely associated with that of insulin and most Cr-stimulated reactions depend on insulin [58]. However, excess of Cr (VI) in biological process has been implicated in specific forms of cancer.

Lead (Pb) is known to be toxic. It is a widespread contaminant in soils and water. Lead poisoning is one of the most prevalent public health problems in many parts of the world [59]. It was the first metal to be linked with failures in reproduction due to it is penetration the placenta easily, and it is also affects the brain, causing damaging brain development in infants [60].

Mercury (Hg) is toxic even at low concentrations to a wide range of organisms, including humans, the organic form of mercury can be particularly toxic [61].

Zinc (Zn) is essential micronutrient in plants, animals and human [62]. However, the excessive amount of zinc salts affects several organs simultaneously as exemplified by zinc phosphide. When this rodenticide is ingested, it reacts with water and stomach juice to release phosphine gas which can enter the blood stream and affect the lungs, liver, kidney, heart and central nervous system [63].

Nickel (Ni) occurs in the environment only at very low levels. Foodstuffs have a low natural content of nickel but high amounts can occur in food crops grown in polluted soils. Uptake of high quantities of nickel can cause cancer, respiratory failure, birth defects, allergies, and heart failure [64].

Recent research showed the ability of microorganism to survive in the presence of different types of heavy metals in a wide range of concentrations [65].
On the other hand, the possibility of using metal resistant bacteria as bio-indicators of polluted environment has been shown to be a sensitive and reliable tool in detecting the sub-lethal toxicity of these toxic compounds [66].

Saturated noncyclic hydrocarbons, cyclic hydrocarbons, olefinic hydrocarbons, aromatic hydrocarbons, sulphur compounds, nitrogen–oxygen compounds and heavy metals are considered the main contains of the petrochemical products. On the other hand, the chemical structures and the physical features were widely varies of any crude oil or refined product depending of its origin for production [67]. As a result of the petrochemical entrance to the marine environment, these pollutants may possibly suffer physical, chemical and biological changes due to the weathering processes, which could be considered as one of the main causes which inducing the toxicity and the potential eco-toxicological impacts of these ecological pollutants [68].

The most toxic components of petroleum products are polycyclic aromatic hydrocarbons (PAHs), alcohols, ketones, benzene derivatives or (BTEX), etc. Considering the adverse effects of the above pollutants on humans, animals, and wild life, the environmental agencies and World Health Organization set quite low limits (from 0.1 to 0.5 mg/l) for heavy metals, pesticides and some petrochemical (i.e. methyl alcohol and BTEX) pollutants in drinking water, food and feed. Aquatic organisms have the ability to take up these hydrophobic compounds due to their ability to connect with cellular molecules after binding to the lipophilic sites on organism cell wall. If the target substance is an important molecule of a cellular process, a toxic response may be induced, and, at the extreme, the integrity of the organism can be seriously compromised [69]. After being taken up by an organism, hydrocarbons and their metabolic products may enhance the production of reactive oxygen species (ROS) by several mechanisms that can lead to cellular damage through protein oxidation, lipid peroxidation (LPO) and DNA damage [70].
Reactive oxygen species (ROS) are generated in all living organisms mainly during mitochondrial metabolism [71]. ROS may include superoxide anion (O$_2^-$), hydroxyl radical (OH$^-$), hydrogen peroxide (H$_2$O$_2$) and nitric oxide (NO). Excessive amounts of ROS may overwhelm natural antioxidant defences promoting DNA, lipid and protein oxidative damage and oxidative stress, which may lead to cell apoptosis and death [72]. ROIs play a central role in the defence of plant and animal cell against pathogen attack [73].

Enhancement of DNA damage was due to oxidative stress, indicating that ROS accumulation in tissues caused subsequent DNA damage. A number of studies have shown that ROS are the major source of DNA damage by causing strand breaks, removal of nucleotides, and various modifications of the nucleotide bases [74]. There are many environmental factors which cause or induce the DNA damage and there are differences in DNA repair capacity [75].

Pesticides and herbicides usually cause unplanned ecological impacts, when they are not completely selective to the target organism. Living species possibly will uptake the pesticides during the digestion of contaminated supplements both food and water or by respiration of contaminated air and through direct skin contact. Agriculturalists and farmland workers suffer more than other people from higher dermal and respiratory uptake and genetic disorders due to direct contact with mixed and sprayed pesticides causing exposure [76]. Although the majority of pesticides were used to control pests in above-ground plant parts, they can reach the soil directly. The soil biological population consist of different types of micro and macro organisms such as bacteria, fungi, algae, earthworms and insects [77], which can be affected by pesticides.

Death of animals may be related to feeding on sources contaminated with pesticides and herbicides. Widespread mortality of wild animals in association with major pest control programmes was reported, when organochlorine pesticides were used in particular [78]. Peri and neonatal animals exposed to pesticides such as aldrin, atrazine, chlordane and
dieldrin, has shown these substances can elicit a variety of perturbations in the sexual differentiation. Birth defects in the male reproductive tract [79] and reductions in sperm count have been associated with the presence of endocrine disrupting chemicals in the environment [80].

In the assessment of pesticide toxicity to living species (humans and animals), the phenomena of carcinogenesis, immuno-disfunction, mutagenesis, neurotoxicity, endometriosis and teratogenesis should be considered along with toxicity in the restricted sense [81]. A recent report shows that toxic chemicals such as pesticides may damage the immune system [82], and can mimic hormones thus disrupting the endocrine system in both humans and animals, causing a variety of disorders [83]. Human health issues such as increased incidences of breast cancer, prostate cancer, testicular cancer, due to the highly toxicity of atrazine and simazine pesticides to human and other living organisms [84, 85].

2.5 Detection and remediation of toxic pollutants

There are many techniques used for detection of environmental pollutants as toxic chemicals, for example heavy metals, pesticides and hydrocarbons. A number of analytical methods, such as atomic absorption or atomic emission spectrosopies (AAS, AES), inductively coupled plasma mass spectroscopy (ICP-MS), cold vapour atomic fluorescence spectroscopy (CVAFS), and high pressure liquid chromatography (HPLC) are capable of detecting traces of toxic pollutants. However, these methods require sophisticated analytical equipment, specialised laboratories and highly qualified personnel, which make such analysis very expensive and time consuming. Conventional analytical techniques such as atomic absorption spectroscopy (AAS), inductively coupled plasma mass spectroscopy (ICP-MS) and chromatography are very sensitive and reliable [86]. However, they also suffer from the disadvantages of high cost, being time consuming, the need for highly trained technicians and the fact that they are mostly laboratory based [87]. Therefore, the development of alternative detection technologies,
for example, simple and inexpensive biosensor devices, capable of rapid detection of environmental pollutions, is urgently needed.

2.5.1 Atomic absorption spectroscopy (AAS)

Atomic absorption spectrometry has been described as a spectral analytical procedure for the quantitative determination of chemical elements employing the absorption of light by free atoms in a gaseous state. AAS is one of the most important instrumental techniques for both quantitative and qualitative analysis of metallic and non-metallic elements in organic or inorganic chemicals [88]. The atomic absorption phenomenon has been noticed firstly in 1802 with Wollaston’s observation when he detected the dark bands in the emission spectrum of sunlight [89]. In 1859, Kirchoff and Bunsen correctly explained Wollaston’s observation by showing that the dark bands were due to the absorption of solar radiation by ground-state gas-phase atoms [90]. However, the process of absorption by atomic vapours would not be used as a quantitative analytical tool for nearly a century until Alan Walsh fabricated the first analytical atomic absorption spectrophotometer in 1953 [91]. Since its invention, atomic absorption spectroscopy (AAS) has gained acceptance as a standard method for the analysis of both metallic and non-metallic elements. AAS is widely accepted because of its high sensitivity at the parts-per-million level and below. Examples of application of AAS as a routine method of chemical analysis include various forms of industrial manufacturing, geology, medicine, and agriculture. The AAS technique has been divided into two stages: the conversion of an analyte molecule into its constituent atoms (atomization), and the subsequent absorption of radiation by these free atoms. AAS quantization is accomplished by measuring the amount of absorbing species at a given wavelength. AAS quantitation principles are based on the Beer-Lambert law, which establishes a linear relationship between the absorbance and the concentration of gas-phase atoms. The AAS instrumentation is similar to other high-resolution spectroscopic techniques, with the main difference in the radiation sources and the need of heating to vapourise materials. The atomic spectrometer schematically shown in Figure (2-4)
which operates as the follows radiation source. In short, electrons of atoms in the atomizer can be promoted to higher orbitals (excited state) for a short period of time (typically in nanoseconds) by absorbing a defined quantity of energy (radiation of a given wavelength). This amount of energy, i.e., wavelength, is specific to a particular electron transition in a particular element. In general, each wavelength corresponds to only one element, and the width of an absorption line is only of the order of a few picometers (pm), which gives the technique its elemental selectivity. The radiation flux without a sample and with a sample in the atomizer is measured using a detector, and the ratio between the two values (the absorbance) is converted to analyte concentration or mass using the Beer-Lambert Law [92]. Usually, this requirement is met by using a spectral source, such as the hollow cathode lamp [93]. AAS can be used to determine over 70 different elements in solution or directly in solid samples employed in pharmacology, biophysics and toxicology research. The technique makes use of absorption spectrometry to assess the concentration of an analyte in a sample. It requires standards with known analyte content to establish the relation between the measured absorbance and the analyte concentration [94].

![Figure 2-4: Schematic diagram of atomic absorption spectroscopy (AAS) [92].](image-url)
2.5.2 Inductively coupled plasma mass spectrometry (ICP-MS)

The quantitative analysis of priority hazardous substances in the environment becomes more and more challenging because new legislation often requires more sensitive methods, or even completely new approaches, for the determination at very low concentrations (pg/L levels) of already defined priority compounds or newly emerging contaminants that show up in the environment as substitutes for already banned substances or as a result of changing industrial processes [95]. Since its introduction in the 1980s, inductively coupled plasma mass spectrometry (ICP-MS) has evolved to become arguably the most versatile, element-specific detection technique [96].

In parallel, because of the fast developments in the field of elemental speciation, the utilization concept of ICP-MS has undergone a significant change. It is a type of mass spectrometry which is capable of detecting metals and several non-metals at concentrations as low as one part in $10^{12}$ ppt (parts per trillion). Inductively coupled plasma-mass spectrometry (ICP-MS) is a very sensitive analytical technique with a high linear dynamic range (ultra-trace to main components). It is capable of analysing all elements from Li to U and can be applied to solutions, solids and gasses [97]. ICP-MS sampled material is transferred by an argon flow as shown in Figure 2-5 into inductively coupled plasma in which an effective temperature of 7000°K results in atomisation and ionisation of the material. Subsequently, the ions are extracted into a mass spectrometer, with which the elemental composition of the material is determined [98]. This is achieved by ionizing the sample with inductively coupled plasma and then using a mass spectrometer to separate and quantify those ions. Compared to atomic absorption techniques, ICP-MS has greater speed, precision, and sensitivity [99].
Figure 2-5: Schematic diagram of inductively coupled plasma mass spectrometry (ICP-MS) [100].

2.5.3 Chromatography

Chromatography is one of the most important analytical techniques used in pollutants residue analysis. It has two advantages which are, firstly, the sensitivity and specificity of the detection systems, secondly, the ability to separate the mixture of analytes in the column. Until recently gas/liquid chromatography (GLC) of pesticides was conducted using packed columns containing a variety of liquid phases and supports [101]. A wide range of volatilities and specific responses of pesticides necessitated numerous analytical conditions in order to chromatograph several classes of pesticides in a single sample. Many pesticides are too polar or do not respond on a packed column while others are thermally unstable and degrade during the chromatographic analysis. Electron capture is equipped to determine low amounts of residues from small samples of various substrates. Although total reliance should not be placed on the analytical data obtained from GLC for the identification of a pesticide residue, there is a need to compare it with other methods like GC-MS and LC-MS also using alternate column packings.

GLC deploys a physical method of separation of two phases, one stationary phase and the other mobile phase moving in a particular direction. Chromatography is the collective term for a set of laboratory techniques for the separation of mixtures. The
mixture is dissolved in a fluid, the mobile phase (as it shown in Figure 2-6), which carries it through a structure holding another material called as the stationary phase. The different components of the mixture travel at different speeds, causing them to separate. The separation is based on differential partitioning between the mobile and stationary phases. A subtle difference in a compound's adsorption coefficient results in differential retention in the stationary phase and thus in the separation of compounds. Chromatography may be preparative or analytical.

The purpose of preparative chromatography is to separate the components of a mixture for more advanced use (this is a form of purification) [102]. Analytical chromatography is done normally using small amounts of material and it measures the relative proportions of the analytes in a mixture. The two are not mutually exclusive. The above mentioned techniques are very expensive in both the actual equipment cost and exploitation cost which requires special laboratories and highly qualified personnel.

![Figure 2-6: Schematic diagram of the chromatographic measuring process [103].](image)

**2.6 Biosensors and sensor arrays**

Much more affordable option of pollution detection is the use of chemical sensors or biosensors which involves integration of recognition elements with transducing
materials for identifying the analyte activity or concentration which present in the sample. Since the mid-1980s there has been a continuous growth in the use of biosensors for environmental analysis owing to their advantages, such as screening of various contaminants in environmental matrices, minimizing sample pre-treatment, reducing the cost and time of analysis, and displaying sufficient sensitivity and selectivity [104]. Biosensors are formed by a combining molecular recognition, (such as an enzyme, an antibody, a microorganism, whole plant cell and tissue) and physicochemical transducer which convert chemical reaction into physical measurable parameter such as the signal (for example: an electrochemical, optical, piezoelectric and so on). A particular subclass of biosensors is represented by the electrochemical biosensors or bio-electrodes, in which combined enzymes and electrochemical transducer.

The general mechanism on which all bio-electrodes are based depends on the interaction between the analyte, present in the sample, and the biochemical component (typically enzyme) immobilized on the surface of the electrode: the consequent formation of one or more electro active species generates an electrical signal or a variation of the existing electrical signal, such signal can be easily recorded using suitable electrochemical apparatus and therefore it is proportional to the concentration or the activity of the chemical species to be determined.

Inhibition biosensors have been used for indirect monitoring of organic (e.g., pesticides) or inorganic substances (e.g., heavy metals) which inhibit its bio-catalytic properties of enzyme. The problem with these types of biosensors based in enzymatic inhibition is that only a few enzymes are sensitive to heavy metals [105] and majority of enzymes are extremely sensitive to environmental conditions such as temperature and pH and thus can not function for long.

The development and research of (bio) sensors is becoming one of the most popular scientific areas at the intersection of the biological and the engineering sciences. Semiconducting technology has developed so much that we see now a rapid infiltration
of new nanotechnology-based approaches in the field of sensors. The resulting new discipline of nanobiosensing is a good example of how engineering sciences, biosensor-related research has experienced an explosive growth over the last two decades. A biosensor is generally defined as an analytical device which converts a biological response into a quantifiable and processable signal [106]. Figure (2-7) shows schematically the following parts comprising a typical biosensor: (a), bioreceptor that specifically binds the analyte; (b), an interface architecture where a specific biological event takes place and gives rise to a signal picked up by (c), the transducer element. The transducer signal which could be anything from the in-coupling angle of a laser beam to the current produced at an electrode, is converted to an electronic signal and amplified by a detector circuit using the appropriate reference and sent for processing to (d), computer software which converts a physical parameter describing the process to electrical signals. Finally, the resulting signals are presented through (e), an interface to the human operator.

Biosensors can be applied to a large variety of samples including body fluids, food samples, cell cultures and be used to analyse environmental samples.

Figure 2-7: Typical biosensor elements [107].
In order to construct a reliable biosensor suitable for use by the general public, a number of conditions must be met:

1. The biocatalyst (i.e., enzyme, antibody, and whole cell) must be highly specific for particular analyte of interest, be stable under normal storage conditions and show a low variation of performance between batches.

2. The reaction should be independent from variations of basic such physical parameters of the environment as temperature, pH, and highly controlled stirring. This will allow the analysis of samples with minimal pre-treatment. If the reaction involves co-factors or co-enzymes, these should also be immobilized on the surface of the transducer.

3. The response should be accurate, precise, reproducible, and linear over the concentration range of interest, without dilution or concentration of samples. It should also be free from electrical or other transducer-induced noise.

4. If the biosensor is to be used for invasive monitoring in the clinical environment, the probe must be tiny and biocompatible, not causing toxic or antigenic effects. Furthermore, the biosensor should not be prone to inactivation or proteolysis.

5. For rapid measurements of analytes from human samples, it is desirable that the biosensor can provide real-time analysis.

6. The complete biosensor should be cheap, small, portable, and capable of being used by semi-skilled operators.

2.6.1 Classification of biosensors
Biosensors can be broadly classified on the basis of bioreceptors types and transducing elements used:
I) **On the basis of biological elements**

a) Enzyme biosensors: enzymes having high selectivity and activity towards specific substrata are the best candidates to be used as bio-receptors. Most of the enzymes used in biosensor are oxidizes but there are certain limitations as their activity is susceptible to pH, temperature, ionic strength etc.

b) Microbial biosensors: they use either cell or microorganisms as biological element. The metabolism of cell or microorganism is used as a basis of their activity. They are cheaper and more versatile as compared to other sensor elements.

c) Immune biosensors: they use antibodies as a bioreceptors which are immobilized on the surfaces of transducer

d) Chemical biosensors: they use aptamers or DNA oligomers or synthetic binding proteins named affimers as a bioreceptors which are immobilized on the surfaces of transducer.

II) **On the basis of transducing elements**

a) Electrochemical transducers: the three common types of electrochemical transducers are impedimetric, amperometric and potentiometric transducers. In amperometric transducers the potential between the two electrodes is set, and the current produced by oxidation / reduction of the electro-active species is measured and correlated with the concentration of the analyte of interest. Potentiometric sensors measure the potential of electrochemical cell with zero current. Impedimetric sensors are based on AC measurements of impedance.

b) Optical transducers: they are used for determining the concentration of analyte on the basis of change in optical density or other optical parameters at appropriate wavelength of light, for example absorption, reflection, polarization, interference Photodetector as a function of incident angle. The examples are optical biosensors based on SPR surface plasmon resonance, wave guide interferometers and optical fibres

c) Calorimetric transducers: they used for calculating the heat of biochemical reactions by measuring the temperature difference between the reaction vessel and isothermal heat sink surrounding it [108].
d) Gravimetric transducers: they are based on measurements of resonance frequency of piezoelectric materials which depends on the mass of adsorbed analyte. Typical examples are QCM quartz crystal microbalance and SAW surface acoustic wave sensors [109].

2.6.2 Whole-cell bacterial sensor array

Whole-cell bacterial biosensors have recently been used in environmental studies to quantify heavy metals and xenobiotic and antibiotic compounds [110]. Bacteria display many surface epitopes that can lead to nonspecific interactions with the sensor surface [111].

In the last few years, there have been dramatic advances in new analytical formats such as microarrays which have revolutionized our ability to characterize and quantify biologically relevant molecules [112]. The principle in all cases is the same. A large family of well-defined reactive molecules is fixed onto a mapped solid surface grid and exposed to a multi-component analyte mixture. Sites on the chip in which a recognition event has occurred (e.g. by a complementary nucleic acid sequence) are identified by one of several possible detection techniques, for example fluorescence. The characteristics of the sample can then be recognized from the nature of the bioreceptor molecules which binding to these sites. Using this principle, an increasingly large number of applications are being developed in medicine, biology, toxicology, drug screening and more. The idea of whole-cell arrays has been advanced by Van Dyk and co-workers [113].

A whole-cell-based biosensor is an analytical device that assimilates a whole cell as the biological component of the sensor. It gives high selectivity and has a physical transducer to generate an assessable signal relative to the concentration of the analyte [11]. Whole-cell biosensors are very specific in nature and the cost of such devices is generally low. They have many advantages such as the ease of use, transferability, and
the capacity to provide constant real-time signals [114]. The concept of synthetic biology offers a platform for redesigning of novel gene composition of a cell. This can be done by pairing a reporter gene that generates a signal with a contaminant-sensing component that responds to chemical or physical change to exposure to a specific analyte. A schematic of whole-cell biosensor is given in Figure (2-8)

![Figure 2-8: Schematic of whole-cell-based biosensor [115].](image)

### 2.6.3 DNA Biosensors and microarrays

In recent years, there has been an enormous increase in the utilization of nucleic acids to recognize toxic compounds, specifically metal ions, because many toxic chemicals such as pollutants show a high affinity toward DNA/RNA. Metal ions exhibit a very high affinity toward DNA, as the interaction between DNA and metal ions either gives favourable or adverse effects such as DNA damage and gene mutation. For this reason, nucleic acid-based biosensors have been developed and used. DNA biosensors exploit the preferential binding of complementary single-stranded nucleic acid sequences. They usually rely on the immobilization of a single-stranded DNA probe onto a surface able to recognize its complementary DNA target sequence by hybridization [116].

Recently, an electrochemical DNA biosensor was developed to study DNA damage caused by several pesticides, such as atrazine, 2,4-D, glufosinate ammonium, carbofuran, paraoxonethyl and difluorobenzuron [117]. A biotinylated DNA probe was immobilized on a streptavidin-modified electrode surface. This DNA probe was hybridized with biotinylated complementary DNA target analyte. Streptavidin labelled
with ferrocene was further attached to the hybridized biotinylated DNA. The close proximity of ferrocene to the electrode surface induced a current signal. The presence of pesticides caused an un-winding of the DNA and thus a decrease of the ferrocene oxidation current which was observed (see Figure 2-9) in voltammetry experiments. Paraoxonethyl and atrazine caused the fastest and most severe damage to DNA.

![Schematic of electrochemical DNA biosensor](image)

**Figure 2-9**: Schematic of electrochemical DNA biosensor [118].

### 2.6.4 Apta-sensors

In 1990, Ellington’s group [119] reported the development of an *in vitro* selection technique which allowed the discovery of specific nucleic acid sequences that bind non-nucleic acid targets with high affinity and specificity. The technique was called SELEX (systematic evolution of ligands by exponential enrichment) and the resulting DNA or RNA oligonucleotides are referred to as aptamers [120].

Aptamers show high affinity towards a wide range of target analytes, including proteins, metal ions, and various organic and pathogenic microorganisms. Aptamers possess several competitive advantages over antibodies, first of all, their accurate and reproducible chemical production [121]. The selected nucleic acids bind their targets with affinity and specificity comparable to those of anti-bodies. Aptamers are more stable than antibodies. They can be selected in extreme conditions whereas antibodies are only stable in physiological conditions. Aptamers can undergo reversible denaturation and they can be easily modified with new functional groups without affecting their activity. Also aptamers are able to change their secondary structure by
engulfing the target. Such transformations can be detected using either ferrocene or electrochemical labels as shown in Figure (2-10). Because of these advantages, numerous aptamer-based biosensors have been developed for the detection of a wide range of targets [122]. There is little information for the detection of pesticides using aptamers. However, a single strand DNA aptamer with specific binding to acetamiprid was described [123]. The potential of aptamers for the pesticide and petrochemical detection has not been exploited yet but aptamer-based biosensors could be an alternative to the conventional methods of pollutants analysis.

![Figure 2-10: Schematic of aptamer-based biosensors, (A) hairpin and quadruplex shapes; (B) linear labelled and label free aptmer](image)

2.6.4.1 Aptamer selection process (SELEX)

Much of the success of nucleic acid aptamers is due to SELEX (systematic evolution of ligands by exponential enrichment; an elegant process by which aptamers can be generated for a given target (e.g protein) [125]. In SELEX, a single-stranded DNA library is first generated and exposed to the target. Any oligonucleotides within the library that bind to the target are retained, while the non-binding sequences are washed away [126]. Typically, the sequences in the library that bind to the target will be a small fraction of the total library so, after this first step, most of the library will have been washed away, and the overall concentration of DNA will been significantly reduced. The remaining (bound) sequences are then eluted and retained. In the next step, the
remaining (binding) sequences are amplified by polymerase chain reaction (PCR). This increases the concentration of DNA in preparation for the next round of SELEX. The library is now biased, or enriched, towards sequences that bind to the target [127].

The process is repeated many times. Often, the binding conditions are made more stringent as SELEX proceeds through each round, increasing the selection pressure. Successive selection and amplification result in a library that, at the end of the SELEX process (typically 8-15 rounds), contains only sequences that bind strongly to the target. After SELEX is complete, the binding sequences need to be identified, which is done by sequencing [128]. Figure (2-11), shows the SELEX process.

**Figure 2-11:** SELEX process [129].
2.6.4.2 Structure and types of aptamers

Aptamers have different types of secondary structure which depends on the nucleotide sequence used to accommodate particular targets. Typical 3D secondary structures are shown below and illustrated in Figure (2-12)

1. Hairpin (or stem-loop) occurs when two regions of the same strand are complementary to one another and can form Watson-Crick base pairs.
2. Quadruplex occurs in guanine-rich sequences, when four guanine bases can associate through hydrogen bonding.
3. Kissing complex formed when the unpaired nucleotides in one hairpin loop base pair with the unpaired nucleotides in another hairpin loop. Usually occurs in RNA.

![Aptamer secondary structures](image)

**Figure 2-12:** Aptamer secondary structures [130].

2.6.5 Immobilization of bioreceptors

The basic requirement of a biosensor is that the biological material should bring the physicochemical changes in close proximity to the transducer. In this sense, the immobilization technology has played a major role to get the solution [131]. Immobilization not only helps in forming the required close proximity between the biomaterial and the transducer, but also helps in stabilising it for reuse. The biological
material has been immobilised directly on the transducer or in most cases, in membranes, which can subsequently be mounted on the transducer. Biomaterials can be immobilised either through adsorption, entrapment, covalent or electrostatic binding, cross-linking or a combination of these techniques.

Several immobilization techniques such as covalent binding, physical entrapment, adsorption, cross-linking, and encapsulation have been reported in literature but not a single technique can be considered as a universal method of immobilization to achieve the better biosensor response. The most important factor to be considered while designing a biosensor is to attain higher sensitivity, and functional stability. For immobilization of whole bacteria as bio-sensitive cells on the sensor surface, a bioreceiving membrane was coated using an electrostatic layer-by-layer (LbL) deposition method. In recent years indeed, polyelectrolyte multilayers (PEM) represented a new attractive way for creating bio-functionalized surface coatings. The LbL assembly technique consists in the alternate deposition of polyanions and polycations from aqueous solutions to build ultrathin multi-layered films on flat substrates [132]. It was first introduced to immobilize biomacromolecules such as doxorubicin [133].

2.6.6 Methods of immobilization

Immobilization of bioreceptors in biological processes can occur either as a natural phenomenon or through artificial process [134]. Different immobilization types have been defined: covalent coupling/cross linking, capture behind semi-permeable membrane or encapsulation, entrapment and adsorption [135]. The types of immobilization can be grouped as “passive” (using the natural tendency of molecules to attach to surfaces-natural or synthetic, and grow on them) and “active” (floculants agents, chemical attachment and gel encapsulation) [136].

2.6.6.1 Covalent binding/cross linking processes

The mechanism involved in this method is based on covalent bonds formation between activated inorganic support and the cell in the presence of a binding (crosslinking)
agent. For covalent linking, chemical modification of the surface is necessary. Covalent attachment and cross-linking are effective and durable to enzymes and antibodies, but it is rarely applied for immobilization of cells. It is caused mainly by the fact that agents used for covalent bonds formation are usually cytotoxic and it is difficult to find conditions when cells can be immobilized without any damage [134]. There are few reports of successful covalent binding of cells and most of them are related to yeast. Navarro and Durand (1977) [137], published an article describing a successful way of covalent binding of *Saccharomyces carlsbergensis* on porous silica beads. Two years later, there was another publication concerning yeast (*Saccharomyces cerevisiae, Saccharomyces amuneca*) immobilization with this method on borosilicate glass and zirconia ceramics [138].

2.6.6.2 Entrapment process

Entrapment is an irreversible method, where immobilized cells are entrapped in a support matrix or inside fibres. This technique creates a protective barrier around the immobilized microbes, ensuring their prolonged viability during processing and storage in polymers [139]. Entrapment is the most extensively studied method in cell immobilization. The matrices used are agar, alginate, carrageenan, cellulose and its derivatives, collagen, gelatine, epoxy resin, photo cross-linkable resins, polyacrylamide, polyester, polystyrene and polyurethane [134]. Entrapment of the microorganisms in porous polymer carrier was often used to capture the microorganisms from suspended solution and then obtain the immobilized microorganisms. As a rule, the entrapment methods are based on the inclusion of cells within a rigid network to prevent the cells from diffusing into surrounding medium while still allowing penetration of the analyte. Entrapment of cells in alginate gel is popular because of the requirement for mild conditions and the simplicity of the used procedure. Entrapment allows high mechanical strength, but has some disadvantages, such as, cell leakage, costs of immobilization, diffusion limitations, and deactivation during immobilization and abrasion of support material during usage. Another disadvantage is low loading capacity as biocatalysts have to be incorporated into the support matrix [140].

50
2.6.6.3 Encapsulation process
Encapsulation is another irreversible immobilization method, similar to entrapment. In this process, bioreceptors are restricted by the membrane walls (usually in a form of a capsule), but free-floating within the core space [139]. The membrane itself is semi-permeable, allowing for free flow of substrates and nutrients (when cells are used as a biocatalyst), yet keeping the biocatalyst inside. The factor determining this phenomenon is the proper pore size of the membrane, attuned to the size of core material. This limited access to the microcapsule interior is one of the main advantages of microencapsulation, for it protects the biocatalyst from the harsh environmental conditions. As most immobilization method, it prevents biocatalyst leakage, increasing the process efficiency as a result [141]. However, even though in encapsulation, high cell loading can be achieved, but the capsules are still very weak [142]. The diffusion limitation is one of the inevitable drawbacks associated with encapsulation method [143].

2.6.6.4 Adsorption process
Cell immobilization is probably the simplest method of reversible immobilization [144]. This technique is based on the physical interaction between the microorganism and the carrier surfaces, simple, cheap and effective. The immobilization of microorganisms on properly chosen adsorbents stimulates microbial metabolism, protects cells from unfavorable agents, and preserves their physiological activity [145]. This cell immobilization technique involves the transport of the cells from the bulk phase to the surface of support (porous and inert support materials), followed by the adhesion of cells, and subsequent colonization of the support surface [146]. Adsorption is based on weak forces. However, it still enables efficient binding process. Usually in bonds formation, several forces are involved: van der Waals forces, ionic and hydrophobic interactions and hydrogen bonds. Both electrostatic and hydrophobic interactions govern the cell-support adhesion, which is the key step in controlling the cell immobilization on the support [139].
Due to their high specific surface area, void volume, mechanical and permeability, low pressure drop, diffusion problems and toxicity, maximum loading, biodegradability and durability and low cost and high availability, they are widely applicable [147].

2.6.6.5 Metal-link/chelation process
A partial explanation of this immobilization method seems to be the formation of covalent bonds between the gelatinous hydrous metal oxides used as support materials and suitable ligands contributed by proteins and carbohydrates present on the cell surface [148]. Hydrous titanium (IV) and zirconium (IV) oxides seem to have minimal effects on the activity of enzymes immobilized to these supports, and yeast cells immobilized in this way have remained viable, according to measurements of their external invertase activity in the periplastic space [149]. The cells also appeared to be quite firmly attached to the surface of the metal oxide. Cells of *Arthrobacter globiformis* have also been covalently bound to silica gel after activation with chromium (III) chloride and such immobilized cells were capable of steroid transformation. Cells immobilized on transition-metal-activated inorganic supports appeared to be operationally stable [148].

2.6.7 Inhibition bacterial sensor array
The main problem of using chemical and biosensors for detection of any toxic chemicals which considered as environmental pollutants is the requirements of specific receptors for every analyte of interest. Advances in synthetic chemistry and biochemistry allow these requirements, to be fulfilled though it is not easy and appeared to be the most expensive part of sensor development. An alternative approach lies in the development of inhibition sensors which utilize enzymes, cells, microorganisms as bioreceptors whose function can be inhibited by pollutants. Such reactions are not specific, e.g. bioreceptors could be inhibited simultaneously by different analytes, so that a single sensor can not differentiate the analytes. This can be done using a sensor array approach in which each sensor is affected differently by different analytes. Such a
sensor array equipped with the appropriate data processing system such as an artificial neural network (ANN) is capable of differentiating responses to several analytes. Our recent research, proved the concept of such inhibition sensor array using bacteria as sensing elements. Series of optical and electrochemical measurements of two types of bacteria (*E.coli* and *Deinococcus radiodurance*) allowed discrimination of two types of pollution, i.e. heavy metals (Ni, Cd) and radionuclides which emit $\gamma$-radiation [150]. There are many previous studies focused on using bacteria as sensing elements for different analytes detection and different sample sources based on optical and electrochemical transducers which are summarized in Table (2-1) as bacterial biosensors for detection of different toxic chemicals and pollutants.

**Table 2-1.** Bacterial biosensors for different pollutants detection.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Bacterial type Biosensor</th>
<th>Transducer type</th>
<th>Sample</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zinc, copper, cadmium, and nickel</td>
<td><em>Pseudomonas fluorescens</em> 10586s pUCD607 with the lux insertion on a Plasmid</td>
<td>Optical</td>
<td>Soil</td>
<td>[151]</td>
</tr>
<tr>
<td>Zinc, copper, cadmium, nickel, lead, iron, and aluminium</td>
<td><em>Chlorella vulgaris</em> strain CCAP 211/12</td>
<td>Electrochemical</td>
<td>Waste</td>
<td>[152]</td>
</tr>
<tr>
<td>Zinc, cobalt, and Copper</td>
<td><em>Pseudomonas</em> sp. B4251, <em>Bacillus cereus</em> B4368, and <em>Escherichia coli</em> 1257</td>
<td>Electrochemical</td>
<td>Water</td>
<td>[153]</td>
</tr>
<tr>
<td>Organophosphates, urea, and ethanol</td>
<td><em>Flavobacteium</em> sp., <em>Bacillus</em> sp., <em>Saccharomyces Ellipsoideus</em></td>
<td>Potentiometric</td>
<td>Laboratory Isolate</td>
<td>[154]</td>
</tr>
<tr>
<td>Mercury, lead, zinc, cadmium, atrazine, simazine, DDVP, hexane, octane pentane, pyrene, toluene and ethanol.</td>
<td><em>E.coli</em> k12 <em>Shewanella oneidensis</em> <em>Methylococcus capsulatus</em> (<em>Bath</em>) and <em>Methylosinus trichosporium</em> (OB3b)</td>
<td>Electrochemical</td>
<td>Laboratory Isolate</td>
<td>This work</td>
</tr>
</tbody>
</table>
2.6.8 Electrochemical sensors based on immobilized bacteria cell

The examples which are given in the previous table are important as a further step towards the development of bacteria-based inhibition sensor array, but it is still far away from real sensor development. Dealing with liquid bacteria samples is not the way forward because of natural variations in bacterial concentration even in laboratory samples not to mention “real” samples taken for analysis. The problem of having a reliable reference for such measurements is a very difficult one. It would be much more useful for real sensor development to use bacteria immobilized on the electrode surface. Changes in biological parameters, including the cells number or cells size, can be studied and monitored in the electrical properties of the bio-cells, i.e. with a bio-cell sensor. A typical biosensor based whole cell consists of an anode and cathode, separated with an electrolyte solution, where the cathode electrode is coated with a cell-culture [155]. The current that easily passed through the aqueous electrolyte solutions can be affected by less conductive bacteria or cells which contain the lipid bilayer of the bacteria cells' membrane acting as an insulator. These measurements can be performed on two or three electrode systems, with working electrodes coated with cell cultures. The applied external electrical field catalyses the live bacteria (that had a lipid layer) accumulating on the working electrode, creating an isolation layer coating the electrode, the conductivity decreasing as a result of this effect. Different artefacts, such as some toxic chemicals and radiation, may affect the functioning of cells (increasing or reducing the number of viable bacteria) that appear as conductivity decreases or increases [156]. Therefore, comparing the results of electrochemical tests performed on a fresh cell culture and after exposure to the artefact can provide information on the concentration of pollutants or radiation dose. In the current project, this idea was explored using bacteria. The three electrodes electrochemical set up is more stable than the two electrodes. Firstly, it contains an additional reference electrode having a stable electrochemical potential invariant of applied voltage and chemical composition of the
solution; the potentials of other two electrodes are measured in respect to a reference electrode [157].

Two parallel plate electrodes are indicated as the working and counter electrodes, the third reference electrode was placed close to the working electrode. The electrical potentials of both working and counter electrodes are measured in respect to the reference electrode having a constant potential in electrolytes solutions. Typically Ag/AgCl reference electrodes are used in such measurements. Working and counter electrodes are typically made of metal or carbon. The working electrode is coated with the investigated substance, in this project cell-culture. Gold-coated glass slides seem to be the most common for working electrodes, since the chemistry of modification of gold is well established for coating gold with different biomaterials. Counter electrodes should be chemically inert; such as platinum or gold which is commonly used for this purpose. However, even chemically inert metals, such as Au and Pt, show instability of surface potential during electrochemical reactions, when a current flowing between electrodes is accompanied with chemical ion exchange electrode reactions. In this case, the role of the reference electrode is crucial for performing accurate electrochemical measurements that are typically carried out using potentiostat. In some cases, however, when electrochemical reactions are not essential, simple measurements can be performed in a two-electrode system without using reference electrode and potentiostat. These simple electrochemical measurements are used in order to establish the correlation between electrical properties (conductivity, capacitance, current or resistance) of liquid bacteria samples and live bacterial counts, after that studying the effect of γ-radiation and heavy metal ions on bacteria. The use bacteria in suspension were essential for preliminary in-vitro study of the effect of pollutants. However, it is of limited use in biosensors. Immobilized bacteria are much more promising for such applications. The measurements have to be taken twice, first on freshly immobilized bacteria, then on on bacteria exposed to pollutants. Electrochemical measurements were successfully used for studying electrical properties of cells deposited on metal
electrodes and showed great prospects in using such cell-based sensors for detection of various analytic [25, 26]. In the previous study, the principles of cell-sensors were extended further to more complex organisms, such as *E. coli*, and another type of bacteria, *D. radiodurans*, which known for its high resistance to γ-radiation [150].

### 2.7 Bacteria as sensing elements and pollutants re-mediators

Different types of microorganisms (bacteria) were utilised as a sensor elements in this study for detection of a large number of toxic chemicals, depending on if they were highly resistant or sensitive to these pollutants. The types of bacteria which can resist or tolerant some of environmental pollutants which are in general classified as gram negative, may be due to the presence of an extra membrane layer. Atypical example of such species is *S. oneidensis* known as a highly resistance and tolerant bacteria to different pollutants. On other hand, highly sensitive Gram negative bacteria such as *E. coli* which is considered to be sensitive to the pollutants while *Methylococcus capsulatus* (Bath) and *Methylosinus trichosporium* (OB3b) may be highly sensitive to some pollutants, for instance heavy metals and pesticides and highly resistant to others such as petrochemical. In this study HgCl₂, PbCl₂, ZnCl₂ and CdCl₂ were used to study the effect of heavy metals ions on *E. coli, Mc. capsulatus* (Bath) & *Ms. trichosporium* and *S. oneidensis* samples. In the current project this approach will be developed further using simple electrochemical measurements on electrodes with immobilized bacteria, for extended range of analytes including heavy metals, pesticides and petrochemicals. The selection of bacteria can serve the purpose since *E. coli* is predominantly affected by heavy metals, while *S. oneidensis* requires them as nutrient and Methanotrophic bacteria thrive in the presence of some hydrocarbons. The desirable outcome of this PhD project will be the development a sensor array prototype capable of differentiating the above pollutants. Such an inhibition sensors array not be sensitive enough for all environmental pollutants, however, they can be used for quick preliminary detection
(screening) of water samples utilizing ANN subsequently these suspected samples could be transferred to conventional laboratory for further analysis.

2.7.1 General description of bacteria

Bacteria are typically between 0.5 µm and 5 µm in size, displaying different shapes, including spherical cocci, rod-shaped bacilli, and spiral-shaped spirilla or spirochetes, among others. Unlike eukaryotic cells, most bacteria are encapsulated by a cell wall which is present on the outside of the cytoplasmic membrane. The cell wall comprises mainly peptidoglycan, a negatively charged polymer matrix comprising of cross-linked chains of amino sugars, namely, N-acetylglucosamine and N-acetylmuramic acid. Bacteria can be classified as either Gram positive or Gram negative depending upon the architecture and thickness of the cell wall. Gram-positive bacteria retain the violet Gram stain due to their thick peptidoglycan layer on the outside of the cell membrane. In contrast, Gram-negative bacteria do not take up the stain, as their thinner peptidoglycan layer is sandwiched between two cell membranes. The outer lipid membrane of Gram-negative bacteria also contains lipopolysaccharides (LPS), which act as endotoxins and elicit a strong immune response in humans, as well as various proteins, including porins. The thick peptidoglycan wall surrounding Gram-positive bacteria contains extra components such as lipids, surface proteins, and glycoproteins. Pathogenic Gram-negative bacteria include *Escherichia coli*, *Salmonella*, *Shigella*, *Legionella*, *Haemophilis influenzae*, *Neisseria gonorrhoeae*, and *Neisseria meningitides*. Examples of pathogenic Gram-positive bacteria include *Streptococcus*, *Staphylococcus*, *Bacillus*, and *Clostridium* [158].

Bacteria comprise a large domain of prokaryotic microorganisms. Typically, a few micrometres in length, bacteria have a wide range of shapes, ranging from spheres to rods and spirals [159]. They were among the first life forms to appear on Earth, and are present in most habitats on the planet, including soil, acidic hot springs, radioactive waste, water, and deep in the Earth's shell, as well as in organic matter and the live
bodies of plants and animals, providing outstanding examples of mutualism in the digestive tracts of humans [160]. There are typically 40 million bacterial cells in a gram of soil and a million bacterial cells in a millilitre of fresh water. In all, there are approximately $5 \times 10^{30}$ bacteria living on our planet [161, 162], forming a biomass that exceeds that of all plants and animals. Bacteria are vital in recycling nutrients, with many steps in nutrient cycles depending on these organisms, such as the fixation of nitrogen from the atmosphere and putrefaction [163]. In biological communities surrounding hydrothermal vents and cold seeps, bacteria provides the nutrients needed to sustain life by converting dissolved compounds such as hydrogen sulphide and methane. Most bacteria have not been characterised, and only about half of the phyla of bacteria have species that can be grown in the laboratory. Bacterial cells are about one tenth the sizes of eukaryotic cells and are typically 0.5-5.0 µm in length. However, a few species are up to half a millimetre long and are visible by the naked eye [164], for example, *Epulopiscium fishelsoni* reaches 0.7 mm [165]. Among the smallest bacteria are members of the genus *Mycoplasma*, which are parasitic organism in animal cell and measure only 0.3 µm, as small as the largest viruses. Some bacteria may be even smaller [166].

Most bacterial species are either spherical, called cocci or rod-shaped, called bacilli. The bacterial cell are surrounded by a lipid membrane, or cell membrane, which encloses the contents of the cell and acts as a barrier to hold nutrients, proteins and other essential components of the cytoplasm within the cell. As they are prokaryotes, bacteria do not tend to have membrane-bound organelles in their cytoplasm and thus contains few large intracellular structures. They consequently lack a true nucleus, mitochondria, chloroplasts and other organelles present in eukaryotic cells, such as the Golgi apparatus and endoplasmic reticulum [167]. Bacteria were once seen as simple bags of cytoplasm, but elements such as prokaryotic cytoskeleton [168] and the localization of proteins to specific locations within the cytoplasm have been found to show levels of complexity. These sub-cellular compartments have been called "bacterial hyper structures". Micro-
compartments such as carboxysomes supply a further level of organization, which are compartments within bacteria that are surrounded by polyhedral protein shells, rather than by lipid membranes. These "polyhedral organelles" restrict and compartmentalize bacterial metabolism, a function performed by the membrane-bound organelles in eukaryotes.

Many important biochemical reactions, such as energy generation, occur by concentration gradients across membranes. The general lack of internal membranes in bacteria means reactions such as electron transport occur across the cell membrane between the cytoplasm and the periplasmic space. However, in many photosynthetic bacteria the plasma membrane is highly folded and fills most of the cell with layers of light-gathering membrane. These light-gathering complexes may even form lipid-enclosed structures called chlorosomes in green sulphur bacteria [169]. Other proteins import nutrients across the cell membrane, or expel undesired molecules from the cytoplasm.

Bacteria do not have a membrane-bound nucleus, and their genetic material is typically a single circular chromosome located in the cytoplasm in an irregularly-shaped body, called the nucleoid. The nucleoid contains the chromosome, with associated proteins and RNA. The order Planctomycetes are an exception to the general absence of internal membranes in bacteria because they have a double membrane around their nucleoids and contain other membrane-bound cellular structures. Like all living organisms, bacteria contain ribosomes for the production of proteins, but the structure of the bacterial ribosome is different from those of eukaryotes and Archaea. Some bacteria produce intracellular nutrient storage granules, such as glycogen, polyphosphate, and sulfur or poly hydroxyal kanoates. These granules enable bacteria to store compounds for later use [170]. One of bacterial phyla, named photosynthetic cyanobacteria, are able to produce internal gas vesicles, which they use to regulate their buoyancy, allowing them to move up or down into water layers with different light intensities and nutrient levels and split H₂O and make O₂.
2.7.2 Bacterial cell wall structure

In most bacteria a rough external layer, the cell wall, protects the delicate protoplast from mechanical damage and osmotic lysis; it also determines a cell’s shape. Additionally, the cell wall acts as a molecular filter, a permeability barrier that excludes various molecules. It also plays an active role in regulating the transport of ions and molecules. The cell walls of different species may differ greatly in thickness, structure and composition. There are broadly speaking two different types of cell wall in bacteria, whether a given cell has one or the other type of wall can generally be determined by the cells reaction to certain dyes, these two types called Gram-positive and Gram-negative. The names originate from the reaction of cells to the Gram stain (red stain), a test long-employed for the classification of bacterial species. Gram-positive bacteria (are a class of bacteria that take up the crystal violet stain used in the gram staining method of bacterial differentiation), which possess a thick cell wall (about 30-100 nm) and it generally has a simple, uniform appearance under the electron microscope. Some 40-80% of the wall is made of a tough, complex polymer, peptidoglycan. Essentially, peptidoglycan consists of linear heteropolysaccharide chains and teichoic acids [171]. In contrast, Gram-negative bacteria (are a class of bacteria that do not retain the crystal violet stain (stained red) used in the Gram staining method of bacterial differentiation), which it have a relatively thin cell wall (20-30 nm) with a distinctly layered appearance under the electron microscope. The inner layer nearest the cytoplasmic membrane is widely believed to consist of a few layers of peptidoglycan (15 nm thick) surrounded by a second lipid membrane, containing lipopolysaccharides and lipoproteins. Most bacteria have the Gram-negative cell wall, and only the Firmicutes and Actinobacteria (which are known as the low G+C and high G+C Gram-positive bacteria, respectively) have the alternative Gram-positive arrangement [172]. These differences in structure can produce differences in antibiotic susceptibility. Therefore, the lipid membrane reaction for gram stain helped the researcher to distinguish between the bacteria. The simple structure of a bacteria cell is shown in Figure (2-13).
Figure 2-13: The schematic structural image of a typical Gram-positive and Gram-negative bacteria cell [173].

The lipid membrane is a thin polar membrane consisting of two layers of lipid molecules. These membranes are flat sheets that form a continuous barrier around cells. The cell membrane of almost all living organisms and many viruses are covered by a lipid, as are the membranes surrounding the cell nucleus and other sub-cellular structures. The lipid layer is the barrier that keeps ions, proteins and other molecules where they are needed and prevents them from diffusing into areas where they should not be. Lipid layers are ideally suited to this role because, even though they are only a few nanometres in width, they are impermeable to water-soluble (hydrophilic) molecules and are particularly impermeable to ions, which allow cells to regulate salt concentrations and pH by pumping ions across their membranes using proteins called ion pumps. In many bacteria there are fine, hair like proteinaceous filaments extending from the cell surface; these filaments can be divided into three main types: flagella, fimbriae and pili. Sex pili (singular: pilus) are elongated or hair-like proteinaceous structures which project from a cell’s surface; they are found specifically on those
Gram-negative cells which have the ability to transfer DNA to other cells by conjugation, a process in which the pili themselves play an essential role. The various types of pili differ in size and shape: for example, some are long, thin and flexible, while others are short, rigid and nail-like, the type of pilus correlates with the physical condition under which conjugation can take place.

2.7.3 Growth of bacteria

Bacteria cell growth involves a coordinated increase in the mass of its constituent parts; it is not simply an increase in total mass, since this could be due, for example, to the accumulation of a storage compound within the cell. Usually, growth leads to the division of a cell into two similar or identical cells. Thus, growth and reproduction are closely linked in bacteria, and the term growth is generally used to cover both processes. Bacteria grow only if their environment is suitable; if it’s not optimal, growth may occur at a lower rate or not at all or the bacteria may die, depending on species and condition. Essential requirements for growth include, (i) a supply of suitable nutrients; (ii) a source of energy; (ii) water; (iv) an appropriate temperature; (v) an appropriate pH; (vi) appropriate levels (or the absence) of oxygen. Consider the growth of bacteria on a solid medium of one common type of solid medium widely used in bacteriological laboratories, which is a jelly like substance (an agar gel) containing nutrients and other ingredients. Suppose that a single bacterial cell is placed on the surface of such a medium and given everything necessary for growth and division. The cell grows, division continues, the progeny of the original cell eventually reach such immense numbers that they form a compact heap of the cells that is usually visible to the naked eye; this mass of cells is called a colony. In addition, either bacterium can move freely through a liquid medium by diffusion or, in motile species, by active movement; thus, as cells grow and divide, the progeny are commonly dispersed throughout the medium. Usually, as the concentration of cells increases, the medium becomes increasingly turbid (cloudy), so that each bacterial species need specific medium and particular environmental conditions for pure growth [174]. This study utilized three kinds of bacteria to scan a wide range of environmental pollution.
2.8 Types of Bacteria used as sensing elements in this work

2.8.1 Escherichia coli K-12 strain

*E. coli* K-12 is a Gram-negative, rod-shaped bacterium commonly found in the intestine of warm-blooded organisms (endotherms). Most of *E. coli* strains are non-pathogenic, but some of them can cause serious food poisoning in human e.g. *E. coli* O157 and several kinds of disease in animals, which are occasionally responsible for product recalls due to food contamination [175]. The non-pathogenic strains are part of the normal flora of the gut, and can benefit their hosts by producing vitamin K2, and by preventing the establishment of pathogenic bacteria within the intestine. This work aims at the development of novel sensing technologies based on different types of bacteria as inhibition sensing array for detection of petrochemicals, pesticides and heavy metals. *E. coli* belongs to the Gram-negative bacteria, and was selected for this task. Also, it is facultative anaerobic, rod shape. As mentioned in previous sections, there are hundreds of different strains of *E. coli*: some are harmless; others cause serious illness. A non-pathogenic strain of *E. coli* (K12) normally present in the intestinal tract in humans and animals was used in this study. Figure (2-14) shows the *E. coli* bacteria cells and their colonies.

![Figure 2-14](image)

**Figure 2-14:** (A) SEM image of *E. coli* (K12 strain) Gram-Negative bacteria cells (B); their culture colonies.
2.8.2 *Shewanella oneidensis* MR-1 strain

*Shewanella oneidensis* MR-1, a facultative anaerobe classified as a gamma proteobacterium, can utilize numerous inorganic compounds as electron acceptors (e.g., oxygen, nitrate, and metals). *Shewanella oneidensis* MR-1 (formerly *Alteromonas putrefaciens* and *Shewanella putrefaciens*) is best known for its respiratory versatility, including the ability to reduce metal oxides and radionuclides. Like many other members of the *Shewanella* genus, it can also grow aerobically or use any of a broad variety of organic (fumarate, trimethyl-amine oxide, dimethyl sulfoxide, and glycine) and inorganic (nitrate, elemental sulphur, thiosulfate, and sulphite) compounds as terminal electron acceptors in the absence of O$_2$. This strain was originally isolated from anaerobic sediments of Oneida Lake in New York through the selective enrichment of bacteria that could respire Mn (IV) [176]. Members of the *Shewanella* genus are frequently isolated from redox-stratified freshwater and marine environments, where they are proposed to play an important role in the geochemical cycling of nitrogen [177], metals [178] and sulphur [179]. Figure (2-15). Shows the SEM image of *S.oneidensis* (MR-1 strain) Gram-Negative bacteria cell (A), and their colonies (B)

**Figure 2-15:** (A) SEM image of *S.oneidensis* (MR-1 strain) Gram-Negative bacteria cell (B); their culture colonies.
2.8.3 Methanotrophic bacteria

Methane-utilizing bacteria (methanotrophs) are a diverse group of gram-negative bacteria that are related to other members of the Proteobacteria. These bacteria are classified into three groups based on the pathways used for assimilation of formaldehyde, the major source of cell carbon, and other physiological and morphological features. The type I and type X methanotrophs are found within the gamma subdivision of the Proteobacteria and employ the ribulose monophosphate pathway for formaldehyde assimilation, whereas type II methanotrophs, which employ the serine pathway for formaldehyde assimilation, form a coherent cluster within the beta subdivision of the Proteobacteria. Methanotrophic bacteria are ubiquitous. The growth of type II bacteria appears to be favored in environments that contain relatively high levels of methane, low levels of dissolved oxygen, and limiting concentrations of combined nitrogen and/or copper. Type I methanotrophs appear to be dominant in environments in which methane is limiting and combined nitrogen and copper levels are relatively high. These bacteria serve as biofilters for the oxidation of methane produced in anaerobic environments, and when oxygen is present in soil, atmospheric methane is oxidized. A limited number of methanotrophs have the genetic capacity to synthesize a soluble methane monooxygenase which catalyzes the rapid oxidation of environmental pollutants including trichloroethylene. [180]. (See Figure 2-16)

Figure 2-16: (A) SEM image of *Mc. capsulatus* (Bath strain) Gram-Negative bacteria cell (B); their culture colonies.
References


CHAPTER 3  EXPERIMENTAL METHODOLOGY

This chapter describes with more details a number of experimental techniques, such as fluorescence microscopy, flow cytometry, and optical density (OD\textsubscript{600}) which were utilized in this study for the purpose of counting the percentage of live and dead bacteria and then studying the effect of environmental pollution on bacterial concentration. The development of sensor array devices for detection of heavy metals, pesticides and petrochemical pollutions is the desirable outcome of this work. In addition, to the above mentioned optical techniques, scanning electron microscopy (SEM) and atom forces microscopy (AFM) will be described as well. Also, the use of simple DC and AC electrochemical measurements e.g. cyclic voltammogram (CV) and electrochemical impedance spectroscopy (EIS) for studying the effect of these pollutants on bacterial suspension and immobilized bacteria will be described.

3.1 Experimental equipment used

3.1.1 Optical methods

The effect of the above pollutants on the bacterial cultures was examined using three different optical experimental techniques: fluorescence microscopy, UV–visible spectrophotometry, and flow cytometry. A Becton-Dickinson FACSCalibur flow cytometer was used to characterise bacteria after staining them with the BacLight live/dead bacterial viability kit (Molecular Probes). Fluorescence microscopy of bacterial suspension and bacteria immobilized on the screen printed gold electrodes was performed with an Olympus-BX60 instrument using the BacLight kit for staining bacteria. Optical density of bacterial cultures was measured at 600 nm with a 6715 UV/Vis spectrophotometer (Jenway).
3.1.1.1 Fluorescence microscopy using Olympus-BX60 instrument

The most common imaging technique used to study the morphological structures of different materials including organic films and biological objects is optical microscopy. The limitations in resolution of optical microscopy due to the diffraction limit and aberrations in optical elements do not allow observing objects in sub-micron dimensions. Alternative modern microscopic techniques, such as electron microscopy and scanning probe microscopy, offer much better resolution down to 10 nanometers (nm). However, they can be invasive due to the use of high vacuum or direct contact of nano-probes and thus not suitable for biological objects. In this respect, non-invasive optical microscopy is advantageous for studying biological objects. Recent advances of optical microscopy, such as near field optical microscopy, managed to overcome the diffraction limit. The use of modern image-processing software can also enhance the performance of optical instrumentation, such as interfacial force microscope (IFM). Another problem in optical microscopy is the lack of contrast of some objects, such as organic and biological thin films. This problem can be solved using polarised light in dark-field microscopy as shown in Figure (3-1A,B), or staining the material with light-absorbing or fluorescent dyes [1]. The biological cells are typical example of non-contrast objects, and the most common way to increase it is to stain the cell culture with selective dyes for example (BacLight™ Bacterial Viability Kit, for microscopy and flow cytometry) [2]. Therefore, the fluorescence dyes that used in this study is BacLight bacterial viability kit contain two dyes the first one is SYTO 9 green fluorescence DNA dye, and the other is red fluorescence DNA (propidium iodide) dye. Those dyes have different physical and biological effect in their spectrum and permeability into the living bacteria cell membrane. In short, bacteria with intact cell membranes stain fluorescent green (live cell), whereas bacteria with damaged membranes stain fluorescent red (dead cell) depends on the charge [3]. Typical dark-field images of bacteria stained with BacLight are given in Chapter 4. The background remains virtually non-fluorescent. The ratio of (excitation/emission) wavelength about (480/500 nm) for SYTO9 stain
which show green spot, and (490/635 nm) for propidium iodide enabling to see the red spots which refers to the dead cells.

**Figure 3-1:** (A) Fluorescence microscope Olympus-BX60, (B); its schematic diagram.

### 3.1.1.2 Optical density measurements using 6715 UV/Vis spectrophotometer

One of the most common methods of detecting optical density of cell culture at 600 nm is (OD<sub>600</sub>) which used to determinate the cells density [4]. A linear relationship exists between the number or density of cells) and the absorption rate at 600 nm as in Figure (3-2A,B).

**Figure 3-2:** (A) 6715 UV/Vis spectrophotometer (B); Schematic of a typical optical density device imaging process.
The calibrate OD\textsubscript{600} measurements in concentration units (cell/ml) the calibration has to be performed using an independent technique such as optical microscopy [5]. The feature single wavelength allows simple absorbance (Abs) or transmission (T %) measurements at a single wavelength defined by user. However, when light hits bacteria, the light may be scattered and/or absorbed [6]. Because bacteria are usually transparent the (OD\textsubscript{600}) of a bacteria liquid culture is mostly due to light scattering. The (OD\textsubscript{600}) outcome represents the amount of light passed through a sample relative to a reference. In this study, the optical density (OD\textsubscript{600}) was studies and plotted as a function of time exposed to pollutants. The concept of OD\textsubscript{600} device is based on the principle of light scattering. When measuring light scattering, it is important to consider the wavelength of light used, so that the light absorption is minimal for most bacterial cultures. A wavelength of around 600 nm is a good choice, and it is the middle of visible range. So the measured absorbance in such samples is due to light scattering, and not the result of molecular absorption and OD\textsubscript{600} gives true account of viable bacteria assuming that dead bacteria are not motile and sediment [7].

### 3.1.1.3 Flow cytometry using Becton-Dickinson FACSCalibur

Quantification of the total counting of bacterial numbers is a basic and essential task in several areas of microbiology. During the last two decades, the direct counting methods that use fluorochrome stains and fluorescence microscopy have become increasingly popular because most naturally occurring bacteria cannot be enumerated accurately as a colony-forming unit (CFU) by culturing on agar media. Advances in fluorescent dye technology and flow cytometry now allow the application of this rapid, automated technique to such studies.

Flow cytometry has become increasingly popular because it offers the advantage over microscopy in rapid, easy, and accurate bacteria enumeration [8]. This increasing popularity is also due to the recent development of low-cost compact flow cytometers. In this work flow cytometry was used to count bacterial cells of different types: i.e.
E. coli, S. oneidensis and Mc capsulatus (Bath). To verify the accuracy and the precision of this technique, total bacteria counts made by flow cytometry were compared with those obtained using fluorescence microscopy. The results of this study showed that flow cytometry was a reliable technique for counting a mixture of bacteria in samples from aquatic ecosystems.

Flow cytometry has been described as automated microscopy that has the advantages of automation, objectivity and speed (flow cytometers can analyse thousands of cells per second). To achieve this flow cytometers quantitatively measure the optical characteristics of cells (or other particles) as they are presented in single file in front of a focused light beam. In order to present cells in single file they are introduced, in the flow cell, into a fast flowing fluid stream termed the sheath flow.

The basis of the flow is a jet of isotonic sheath fluid (approximately 100 µm in diameter) into which samples are injected at a controlled rate, typically between 10 and 60 ml/min. The light source used to illuminate samples is either a high-pressure mercury vapour lamp or an assortment of lasers. As particles pass through the light beam the following three parameters are measured using photomultiplier tubes as shown in Figure (3-3A,B); these are forward light scattering (also called FALS or FSC), side scattering (also called 908 LS SALS or SSC) and fluorescence (FL).

The amount of light scattered forward and at right angles by any particle tends to increase with the bacterial cell size or numbers. In addition, the cell refraction ability is related to surface properties and internal structure, and these also affect FSC and SSC. Natural fluorescence (auto-fluorescence) is emitted by cellular components such as flavin nucleotides, pyridine and photosynthetic pigments. A typical flow cytometer measures fluorescence in three wavelength ranges. Light of defined wavelengths is channelled to particular detectors, for example detector FL1 will typically measure green fluorescence, FL2 orange fluorescence and FL3 red fluorescence. All these information referred to Shapiro (1995) [9], and Robinson (1997) [10].
Figure 3-3: (A) Flow cytometry instrument (B); Schematic diagram of flow cytometry measurements.

3.1.1.4 Scanning Electron Microscopy using FEI-Nova SEM

Scanning electron microscopy is one of the traditional and well-established methods in surface science. The SEM operates in a vacuum, with a high energy electron beam (typically from 5 to 20 keV) focused into a spot of several tens of nanometers in diameter (or fractions of nanometers in modern high-resolution instruments) [11]. Because of a very short wavelength of the electrons used (for example, electrons with an energy of 100 eV have a wavelength $\lambda_e = \text{equal to} \ 0.12 \ \text{nm}$), the resolution of SEM is limited by the beam diameter and image distortions, (astigmatism) introduced by the focusing system. A complex electromagnetic system provides both the formation of the electron beam and its scanning over the sample (see Figure 3-4). The electrons that are reflected (backscattered) from the surface, or the emitted secondary electrons, are collected with a sensitive detector to provide imaging of the studied surfaces. SEM can be used to study samples of different nature and morphologies, practically without any limitations. Ideally, the samples for SEM study should be conductive to avoid electrical charging of the sample with an electron beam which causes image distortion. Therefore, the samples of insulating materials for SEM study are recommended to coat (decorate)
with thin of carbon or gold film which reproduces the surface profile. Another possible drawback of SEM lies in the film damage caused by high energy electrons, which is particularly important for organic film study. The interpretation of SEM images of composite samples may not always be straightforward. This is especially true for secondary electron images, which represent a combination of both the surface and the work function profiles [11]. In scanning electron microscopy (SEM), beams of electrons are focused by the electromagnetic lenses on the sample with energy up to 40 KeV in high vacuum chamber and scanned with parallel lines. Consequently, the reflected signals backscattered or secondary electrons are collected to estimate the sample morphology and provide an image. [12]. Up to date, SEM is widely used in several fields, for instance in semiconductor manufacturing, medical and material science research. Figure (3-4 A,B) shows the SEM scheme and operation with the instrument image. An SEM instrument contains electron source which is the electron gun, two magnetic or condenser lenses, scanning coils which facilitates the deviation of electron beam in x and y dimensions, objective lens and detectors for backscattered and secondary electrons[13]. The scanning electron microscope FEI-Nova SEM has been used in this research.

Figure 3-4: (A) Scanning Electron Microscope FEI-Nova (B); Schematic diagram of a typical SEM.
3.1.1.5 Atomic Force Microscopy using Nanoscope IIIa Brucker AFM

Atomic force microscopy (AFM) is a scanning probe technique widely used in research and industrial applications for visualizing surfaces at the nanoscale as well as measuring surface physical properties. Even though electron microscopy techniques like transmission electron microscopy (TEM) or scanning electron microscopy (SEM) can be utilized for similar surface investigations, the high vacuum environment and specific sample preparation required, prevents the real time imaging and in-situ monitoring of biological systems.

AFM, with its ability to obtain high resolution images under ambient (air or fluid) conditions, therefore, has been used to follow kinetic processes in many systems ranging from polymeric and crystalline materials [14], to cellular membranes and biomolecules [15]. Atomic force microscopy (AFM) is a high-resolution scanning microscopy with a verified nanometer resolution. Gerd Binnig and Heinrich Rohrer were developed both SEM and AFM in the early 1980s at IBM Research - Zurich, an achievement that earned them the Nobel Prize in Physics in 1986. The first atomic force microscope was available in the market in 1989.

AFM includes a cantilever with a sharp tip which is used to scan the film surface. This probe follows the surface profile and the information recorded using a laser as shown in Figure (3-5A), and Figure (3-5B) shows the AFM principles operation [16].

The topographic image of the surface can be acquired by recording the deflection of the cantilever, which results from the difference of interaction force between cantilever tip and the sample surface [17]. AFM can operate in three different modes: contact mode, noncontact mode, and tapping mode [11]. Figure (3-5B) shows the dependence of the interaction force on the distance between the tip and the sample surface. At larger distances from the surface, the attraction force is dominating, while the repulsion force starts to dominate at smaller distances (in the range of a few angstroms). The basic idea of AFM is slightly different from SEM, Although the main unit is the same [X,Y,Z
piezo-ceramic positioner], the principle of registration of vertical movement of the tip, which is usually made of Si₃N₄ by CVD, on a flexible cantilever, is optical.

The laser beam reflected from the cantilever is detected with the position sensitive photodetector. This construction is universal, and is adopted by the majority of scanning nanoprobe instruments [11]. In the contact mode, the tip is brought into close contact with the sample surface, so that the force between the tip and the sample becomes repulsive. The deflection of the cantilever caused by this force is registered with a photodetector. The signal is compared to the predefined value of deflection (force), and the DC feedback system generates a certain voltage applied to the Z-part of the piezo-ceramic to keep the value of deflection (force) constant. This DC voltage therefore measures the surface roughness. The vertical resolution of the AFM contact mode is in the range of $10^{-2}$ nm, while the lateral resolution can reach the values less than 1 nm, depending mostly on the tip radius [11]. The disadvantage of contact mode is that it can damage both the sample and the tip. Comparison of AFM tapping and contact modes shows clearly the advantage of the former for this particular application.

**Figure 3-5:** (A) Atomic force microscope Nanoscope IIIa (Brucker) (B); Schematic diagram of a typical atomic force microscope.
3.2 Electrochemical DC and AC methods for biosensing approaches

Electrochemical sensors can be classified by their operational principles, e.g. amperometric, conductometric (impedimetric), and potentiometric, which are based, respectively, on measurements of electric current, conductivity in DC measurements or general impedance in AC measurements, and voltage (potential). Different biological reactions and processes can be monitored using their electrochemical characteristics; this approach is not invasive and is relatively cheap. The parameters and metabolic activity of cells and microorganisms can be studied and monitored using their electrochemical properties. Cell growth, cell activity, changes in cell composition, numbers, shapes or cell locations are only some examples of characteristics can be detected by microelectrode cell sensors. The electrochemical properties of a biological sample reflect actual physical properties of the cell membrane.

3.2.1 DC electrochemical measurements

DC measurement is a very reliable technique for investigating the electrical properties of a liquid bacteria suspension. If a DC potential is applied across the electrodes, a current may flow under certain conditions. Thus, it is important to consider the addition of a resistive path in parallel with the capacitive in the electrical model of this surface. This resistor can be non-linear with the applied voltage. The flow of a current through this metal-electrolyte interface requires the net movement of a charge in response to an electric field due to the applied voltage.

The conductivity of metals and semiconductors is due to electrons (holes in semiconductors are missing electrons) while electrolytes conduct the current due to ions. Therefore the processes of discharging ions must take place at the interface between metal as semiconductor and electrolyte, for example, in the case of a simple system of two metal electrodes dipped into NaCl aqueous solution and connected to a battery (see Figure 3-6) the following electrochemical reactions are taking place:
\[ 2H^+ + 2e^- \rightarrow H_2 \uparrow \] and \[ Na^+ + e^- \rightarrow Na \uparrow \] on cathode, \[ O^{-2} \rightarrow 2e^- + O_2 \uparrow \] and \[ 2Cl^- \rightarrow 2e^- + Cl_2 \uparrow \] on anode.

**Figure 3-6:** Charges exchange on metal electrodes in contact with electrolyte

In reality, the reactions are more complex: highly reactive Na will most likely react with water to form NaOH near cathode, while \( O_2 \) and \( Cl_2 \) gas may cause the corrosion of anode by forming metal oxides and metal chlorides, respectively; alternatively \( Cl_2 \) may react with water to form HCl near the anode.

As a result of such charge exchange processes electrical double layers will form on the surface of both electrodes. For example on the cathode, the negative electron charge on the surface of metal will be counterbalanced by positive ions in solution, as shown in Figure 3-7. Because the concentration of ions in solution is much lower than concentration of electrons in metals, the space charge of ions will be extended (by few microns) into solution.
Figure 3-7: Electrical double layer (EDL) at metal-electrolyte interface and the resulting distribution of surface potential in EDL.

The diagram of an EDL in Figure 3-7 is schematic and in reality both cations and anions in the electrolyte are solvated by polar water molecules. The main feature of an EDL is however represented correctly, with an actual double-layer called the Helmholtz layer at the very surface and an extended diffusion layer called as Gouy-Chapman layer. According to the Sterns model [18] the resulted surface potential distribution is described by a linear potential drop within the Helmholtz layer between 0 and outer Helmholtz plane \((x_{OHP})\) also called as Stern plane

\[
\Psi = \Psi_0 \left(1 - \frac{x}{x_{OHP}}\right) \quad (3.1)
\]

and an exponential dependence within the Gouy-Chapman layer beyond the Stern plane

\[
\Psi = \Psi_{St} \exp\left(-\frac{x - x_{OHP}}{\delta}\right). \quad (3.2)
\]

The potential \(\zeta\) at shear plane (boundary of diffuse layer) is known as \(\zeta\)-potential or electrokinetic potential. \((x > x_{OHP})\). Because Stern and shear planes are very close \(\zeta \approx \Psi_{St}\). Another characteristic parameter, e.g. Debye length \((\delta)\), is introduced at the point of \(\Psi_{St} / e\). The Debye length depends on the ion concentration \(C\) in solution as \(\delta \approx 1/\sqrt{C}\), and typically is in the nanometres range. \(\zeta\)-potential and thus the voltage
drop on a particular metal-electrolyte contact depend on many parameters. There include the surface potential of the metal, electrolyte contents, concentrations of chemicals present, pH, and temperature. Passing current through the contact may alter the chemical composition and concentrations in the vicinity of the electrode and therefore affect the potential drop at the contact. Such instability of the contacts’ potentials should be taken into account during electrical measurements in electrolytic cells. The simplest electrochemical cell shown in Figure (3-8a) consists of two electrodes traditionally called as working electrode (WE) and counter electrode (CE). The working electrode is usually the electrode being studied, i.e. the electrochemical processes on this electrode which could be functionalised with biological receptors, such as enzymes, are of interest. The distribution of the electric potential in this system in Figure (3-8b) exhibits potential drops at both electrodes as well as a potential drop in the electrolyte in the middle. Since both contact potentials are not stable the electrical measurements in two-electrode system have some uncertainty. In order to eliminate the instability of one of the electrode (typically WE), its potential is be recorded against the third reference electrode which a stable potential.

**Figure 3-8:** Two-electrode system (a) and potential distribution in it (b); three-electrode (c) and four-electrode systems (d).
as shown in Figure (3-8c). The construction of the RE is such that potential is invariant of the electrolyte content. One of the most popular RE designs is Ag/AgCl electrode which consists of an Ag wire in contact with the saturated AgCl solution incapsulated into a glass probe connected to the cell via porous glass tip. Such three-electrode systems are very common and widely accepted in electrochemistry. If, however, the stability of potentials of both WE and CE are of concern, a second reference electrode (RE1) coupled to CE can be introduced in four-electrode cell as shown in Figure (3-8d). One more condition should be obeyed when using reference electrodes in three- or four-electrode systems, the electric current should not pass between WE and RE (or CE and RE1). The circuit design for basic DC electrical measurements of current voltage characteristics in three-electrode cell, called voltammetry, is shown in Figure (3-9a). The current is passed between WE and CE and recorded with the amperometer while the voltage at WE is measured against the RE. Usually such measurements are carried out using a potentiostat (see Figure 3-9b) which automatically provide such functionality. The potentiostat allows operating in different modes, for example, cyclic voltammetry during which the current is recorded during scanning over the predefined voltage range.

![Circuit for voltammetry in three-electrode cell (a), the same circuit with potentiostat (b).](image)

**Figure 3-9:** Circuit for voltammetry in three-electrode cell (a), the same circuit with potentiostat (b).
Let’s discuss the processes of electric charge exchange on electrodes in more detail. The electroactive chemicals which take part in the electric charge exchange could exist in either reduced (Red) or oxidized (Ox) forms, thus the charge exchange reaction on electrodes can be described as: \( Ox + z e^- \leftrightarrow Red \), where \( z \) is the number of electrons discharged at the electrode. According to Faraday’s law the electric current between two electrodes immersed in an electrolyte consists of two components, i.e. cathode (or reduction) current \( I_c \) and anode (or oxidation) current \( I_a \) which is given by Faraday law:

\[
I = I_c + I_a = zFk_c C_{ox} - zFk_a C_{red},
\]

where \( F \) is Faraday constant, \( C_{ox} \) and \( C_{red} \) are concentrations of Ox and Red components, respectively, while \( k_c \) and \( k_a \) are, respectively, cathode and anode reactions rates which in turn depend on respective electrodes’ potentials:

\[
k_c = k_0 \exp \left( -\alpha Z F \frac{(E - E_0)}{RT} \right) \quad \text{and} \quad k_a = k_0 \exp \left( (1 - \alpha) Z F \frac{(E - E_0)}{RT} \right);
\]

where \( k_0 \) and \( E_0 \) are, respectively, standard rate constant and standard redox potential, \( R \) is gas (or Redberger) constant and \( T \) is temperature. Assuming that \( I_c = I_a \) in equilibrium, the ratio of concentrations of Ox and Red components is given by Nernst’s equation:

\[
\frac{C_{Ox}}{C_{Red}} = \exp \left[ \frac{Z F (E - E_0)}{RT} \right]
\]

Current-voltage characteristics of electrolytic cell are usually recorded by scanning the applied voltage from negative (cathode) to positive (anode) potentials and back; such characteristics are called as cyclic voltammograms (CV) in electrochemistry. Typical CV characteristics of redox reaction are shown schematically in Figure (3-10). The peak current corresponding to the oxidation process is observed on the forward pass, while almost symmetrical negative current peak associated with the reduction process appears on the reverse pass.
As has been mentioned previously, the most common and therefore accurate measurements are taken in three-electrode systems using a potentiostat. Typically, the voltage on the plot corresponds to the voltage of working electrode measured against the reference electrode. The current values from peaks may deviate from zero value due to so-called non-Faradaic processes not related to charge exchange, i.e. polarization of electrodes, adsorption and desorption of molecules on electrodes and corresponding redistribution of electrical charges in the system.

The most common approach is based on measurements of cyclic voltammograms (CV) in three-electrode electrochemical cell as described in previous (3.3.1) section. The first electrochemical sensor, known as Clarks’ oxygen electrode, was developed by L.C. Clark in 1956 [19]. As shown schematically in Figure (3-11), it is constructed on a bases of Ag/AgCl electrode with inserted Pt wire acting as a catalytic working electrode; the electrode is separated from the investigated volume by an oxygen-permeable membrane. An amperometer connecting Ag/AgCl and Pt electrodes registers a current caused by oxygen reduction on the Pt electrode acting as a cathode:

\[ O_2 + 4e^- + 2H_2O \rightarrow 4OH^- . \]
Further modification of Clark electrode by coating it with the enzyme glucose-oxidase (GOD) lead to development of glucose sensor. Enzymes are folded proteins which can capture selectively some small molecules (known as substrata) and catalyze their decomposition. In this case, glucose-oxidase catalyses decomposition of glucose with one of the side products being hydrogen peroxide;

Figure 3-11: Clark oxygen electrode (a), Glucose sensor (b).
Glucose + $O_2 \rightarrow H_2O_2 +$ Gluconolactone.

Hydrogen peroxide is further reduced to oxygen ($2H_2O_2 \rightarrow O_2 + 2H_2O$) which can be detected with the Clark electrode.

Also $H_2O_2$ can be directly detected with Clark electrode, but this time the Pt electrode has to be set at anode potential of 0.68V, and the current was due to the following electrochemical anode reaction: $H_2O_2 \rightarrow O_2 + 4e^- + 2H^+$. 

Later, the glucose sensor as well as other enzyme sensors, have been developed further in using direct amperometry on electrodes with entrapped enzymes [20]. Different enzyme reactions can be utilised for sensor development; in addition to decomposition of glucose by glucose-oxidase, other typical enzyme/substratum pairs are: cholinesterase/choline, urease/urea, lactase/lactose, alcohol dehydrogenase converting alcohols to aldehydes or ketones, etc. It is interesting that glucose sensors are the most common on the biosensing market because of the unique stability of glucose-oxidase. But due to diabetes being an increasingly common and controllable condition, so long as blood glucose is known.

These days the most common approach in enzyme amperometric sensing is based on the use of mediators as alternative oxydising agents [21]. Mediators are typically redox pairs, i.e. chemicals having distinctive oxidized and reduced states with corresponding peaks on CV. Such mediated enzyme electrochemical reaction is shown schematically in Figure (3-12):

![Figure 3-12: The scheme of mediated enzyme reaction.](image)

100
Typical examples of such mediated enzyme sensors are: sensor for glucose utilising enzyme glucose oxidase and ferrocene (as mediator) [22], sensors for fructose using the enzyme fructose dehydrogenase and mediator ferricyanide [23], sensor for morphine based on the enzyme morphine dehydrogenase and mediator phenazine methasulfate [24].

These days, the design of amperometric enzyme sensors is mostly based on standard screen-printed three-electrode assemblies produced commercially, for example by DropSens. Typical electrode materials are gold, platinum, and carbon and the assembly typically has a solid Ag/AgCl electrode on board. Portable plug-and-play DropSens potentiostats allows various types of measurements including CV. Immobilization of active enzymes on the surface of electrodes can be achieved by different means. In addition to traditional methods of enzyme encapsulation into a porous matrix such as glutaraldehyde/BSA mixture [25] or hydrogels [26] the method of electrostatic self-assembly proved to be successful [27]. Deposition of alternating layers of enzyme urease or cholinesterase and the polycation PAH proved to be successful and provided good stability of enzymes samples could be kept in the fridge for few weeks as well as good permeation of substrata into the matrix [11].

3.2.1.1 Cyclic voltammetry using μSTAT 4000 (DropSens)

Cyclic voltammetry is the most common technique used for acquiring versatile information on the redox processes and the kinetics of heterogeneous electron-transfer reactions, and adsorption processes. It is often the first experiment performed in an electrochemical analytical study and usually used to learn the electrochemical properties of an analyte in solution [28].

μSTAT 4000 is a portable multi potentiostat/galvanostat from DropSens for use with electrochemical sensors or electrochemical cells. The instrument contains a microprocessor which controls up to four independent electrochemical nodes; each one is able to apply potential or current to the electrodes and measure the current or
potential response. Each node is used with electrochemical sensors or electrochemical cells with three electrodes: working electrode, reference electrode and auxiliary electrode. With μStat 4000 you can perform up to four different electrochemical techniques at the same time, or carry out the study using the same electrochemical technique in several nodes but selecting different parameters to record. Also, μSTAT 4000 can be used in multichannel mode, with up to four working electrodes sharing one auxiliary electrode and one reference electrode. μSTAT 4000 can be connected to a PC via Bluetooth or by means of a USB cable [29].

Cyclic voltammetry has proven to be a convenient and relatively fast method of screening electro-catalysts, while in situ cyclic voltammetry has been widely used for evaluating electrode preparation procedures and characterizing electrode changes during fuel cell durability testing. However, the limitation of cyclic voltammetry is that it mainly provides qualitative and quantitative information, so the thorough investigation of electrochemical reactions is often always associated with other electro analytical methods. Cyclic voltammetry is an important and very frequently utilized electrochemical technique because it offers a wealth of experimental information and insights into the kinetic and thermodynamic details of many chemical systems. It was first reported in 1938, and then described theoretically by Randles in 1948 [28]. Cyclic voltammetry measurement is accomplished with a two- or three-electrode arrangement, whereby the potential relative to some reference electrode is scanned at a working electrode while the resulting current flowing through a counter electrode is monitored as it shown in Figure (3-13A,B). It is rarely used for quantitative determination, but it is ideally suited for a quick search of redox couples, for understanding reaction intermediates, and for obtaining stability in reaction products [30].
Figure 3-13: (A) Four-channel cyclic voltammetry experimental set-up based on µSTAT 4000 potentiostat (B); Typical voltammogram on PC screen, portable fixed-voltage potentiostat and screen-printed three electrode assemblies.

3.2.1.2 Cyclic voltammetric analysis

In cyclic voltammetry, the electrode potential changes linearly versus time. This ramping is known as the experiment's test rate (V/s). The voltage is applied between the electrodes and the current is measured between the working electrode and the counter-electrode. The data are most often plotted as current (I) against voltage (V). Cyclic voltammetry (CV) has become a broadly used electro analytical technique in many areas of chemistry. It is widely used to study a diversity of redox processes, to achieve stability of reaction products, the presence of intermediates in oxidation-reduction reactions, reaction and electron transfer kinetics [28], and the reversibility of a reaction [30].

The curve resulting from a particular set of cyclic voltammetry measurements is known as a cyclic voltammogram. A typical cyclic voltammogram recorded for a reversible redox system is shown in Figure (3-14A). The characteristics of the peaks in a cyclic voltammogram can be used to acquire qualitative information about the relative rates of reaction and reactant diffusion in a given electrochemical system. In Figure (3-14B), it can be observed that when the potential of the working electrode is more positive than
that of a redox couple, the corresponding reactants may be oxidized and produce an anodic current. The peak current occurs when the potential reaches a value at which all the electrochemically active reactants at the electrode surface are completely consumed. When the potential is controlled at this value, the mass transport rate from the bulk to the electrode surface reaches a maximum, driven by the largest concentration gradient between them. After this peak, the current will decline because the double-layer thickness increases, resulting in a less steep concentration gradient for the active reactant. On the reverse voltage scan, as the working electrode potential becomes more negative than the reduction potential of a redox couple, reduction may occur and cause a cathodic current. A cyclic voltammogram can have several cathodic and anodic peaks due to intrinsic reaction mechanisms.

The important parameters in a cyclic voltammogram are the peak potential and peak currents. If the electron-transfer process is fast as compared to other processes (such as diffusion), the reaction is regarded as electrochemically reversible, and the peak separation due to the potential scan rate.

Therefore, the reversibility of an electrochemical reaction is always a relative term, related to the potential scan rate. A reaction that is reversible at low-scan rates may become quasi reversible or even irreversible at high-scan rates. In these cases, the anodic peak potential becomes more positive and the cathodic peak potential becomes more negative. This occurs because the current takes more time to respond to the applied voltage than in the reversible case. The separation of the two peaks also becomes larger than in the reversible case, as shown in Figure (3-14B). Therefore, cyclic voltammetry can be used to elucidate the kinetics of electrochemical reactions taking place at electrode surfaces. From the sweep-rate-dependent peak amplitudes, and the widths and potentials of the peaks in the voltammogram, information can be obtained about adsorption, desorption, diffusion, and coupled homogeneous electrochemical reaction mechanisms. If the rate of the electron-transfer is slow in
comparison to other processes (such as diffusion) the electrochemical reactions will be quasi-reversible or irreversible rather than completely reversible [29].

![Figure 3-14](image)

**Figure 3-14:** (A) Typical cyclic voltammogram; (B) The effects of scan rate (a) electron transfer rate (b) on cyclic voltammogram.

### 3.2.2 AC electrochemical measurements

Today, AC electrochemical measurement might be one of the most powerful and sensitive technologies, for examining the surface electrical properties of an electrode and its coatings which used for determining surface area of the electrode and analyte reaction. These techniques fall into two general categories. The first type uses a surface-limited chemical reaction to quantify the surface area of the electrode. In contrast, the second type measures a physical characteristic that is proportional to the surface area.

#### 3.2.2.1 Electrochemical Impedance Spectroscopy using PARSTAT 4000A

The PARSTAT 4000 teamed with the VersaStudio software package, comprises a simple-to-use, flexible, and extremely powerful system for performing a wide range of electrochemical testing. The PARSTAT 4000 is a potentiostat/galvanostat with frequency response analyzer (FRA) combined in a single unit. It is controlled from any
suitably equipped PC using the VersaStudio electrochemistry software package. The usefulness of impedance spectroscopy lies in the ability to distinguish the dielectric and electrical properties of individual contributions of components under investigation. Electrochemical impedance involves the analysis of resistive and capacitive (or inductive) elements responses to the small amplitude sinusoidal excitation signal response [31].

Impedance spectroscopy has been widely used for detection of a variety of analytes from small molecules, such as toxins, to large protein molecules and even larger objects such as viruses and bacteria [32].

The detection principle of impedance spectroscopy (IS) lies in recording the changes of impedance caused by different phenomena: (i), binding of target molecules to receptors antibodies, DNA, proteins, whole cells, bacteria and other bio-recognition elements immobilized on the surface of the electrodes [33] (ii), changes in the conductivity of the medium caused by the growth of bacteria [34] (iii), due to suspension of target molecules in the aqueous medium [35], or (iv), capturing bacterial cells on the surface of electrodes using dielectrophoresis (DEP) [36], or (v), changes in the ionic concentration of the medium caused by the activity of enzyme used as labels for the signal amplification [37].

Generally impedance measurements are divided into two categories: Faradaic and non-Faradaic [38]. Faradaic measurements require the presence of a redox probe providing electron-ion exchange at the electrodes, while non-Faradsic measurements can be performed in the absence of a redox probe. Traditionally, macro-sized metal rods or wires were used as electrodes immersed in the medium to measure the impedance [39], shown in Figure (3-15 A,B). In attempt to miniaturize the IS sensors and improve their sensitivity, micro-electrodes have been considered as potential candidates to combine with traditional detection systems. Microelectrodes favor a greater rate of reactant supply while macro electrodes cause greater depletion of reactants and require lower
concentrations of electro-active ions to form double layer as compared to macro electrodes [40].

As a result, microelectrodes can perform impedance measurement even in solutions of low conductivity, where macro electrodes may not be sensitive. Among microelectrodes, interdigitated array microelectrodes (IDAM) present promising advantages in terms of low ohmic drop, fast establishment of steady-state, rapid reaction kinetics, and increased signal-to-noise ratio [41]. IDAM consist of a series of parallel microband electrodes in which alternating microbands are connected together, forming a set of interdigitating electrode fingers. Due to proximity of cathodic and anodic electrodes, minute amounts of ionic species can be efficiently cycled between the electrodes resulting in very large (>0.98) collection efficiencies, giving the IDAM an advantage in detecting small amounts of generated electrode products [42]. IS using IDAM is illustrated in Figure (3-15 A, B).

Additionally, IDAM eliminates the need for a reference electrode because of small amplitudes of AC signal and provides simple means for obtaining a steady-state current response, which is comparatively simpler to detect in comparison to three or four electrode systems [43]. Impedance spectroscopy is one of the principal electrical/electrochemical transductions methods providing the means of label-free, real-time, and non-invasive detection of different analytes for a wide range of applications in biological and biomedical areas.

The following examples clearly show the great potential of impedance spectroscopy which still is not fully explored [44]. Electrochemical impedance spectroscopy is being applied for detection of the concentration of cells adhered on surfaces and present in solution. This method has been extensively used in microbiology for detection, quantification, and identification of bacteria, including micro-machined devices for cell counting or cell differentiation. It has also been used for selective capturing of bacterial cells, antibodies, aptamers, or bacteriophages [45]. Another common application of
impedance spectroscopy is the monitoring of bacterial growth; this technique allows distinguishing between viable and dead bacteria cells. A novel impedance biosensor for bacterial cell detection is constructed by immobilizing antibodies that are specific to the target bacterial cells on an electrode surface. The presence of intact cell membranes on the electrodes determines the current and thus the sensor signal. In this method, the impedance is recorded as a function of the interrogating frequency, and thus having the name of electrical/electrochemical impedance spectroscopy (EIS) [46].

![Figure 3-15: (A) PARSTAT 4000A impedance analyzer instrument; (B) Schematic diagram of impedance measurements, and DropSens interdigitated electrode used.](image)

### 3.2.2.2 Electrochemical impedance analysis

The conductometric electrochemical sensors are based on measurements of conductivity or resistivity of electrolyte solutions containing some active organic or bio-molecules acting as receptors. Since electrochemical DC measurements involve ion-exchange processes on electrodes described in the previous section (3.2.2.1). Conductometry is usually associated with AC measurements of electrical impedance; this method is often referred to impedimetry. These measurements are usually performed in a two-electrode configuration using very small amplitudes of AC voltage in order to avoid any electrochemical processes on the electrodes. The electrodes are functionalised with
molecular receptors capable of binding analyte molecules; these could be antibody-antigen pairs. Alternatively, the active chemicals may exist in solution. The design of electrodes and cells can vary widely. However, these days commercially produced interdigitated metal (gold or platinum) electrodes are the most common. An interdigitated electrode design is schematically shown in Figure (3-16).

![Diagram showing interdigitated electrodes and AC measurements](image)

**Figure 3-16:** Design of interdigitated electrodes and the scheme of AC measurements.

Depending on the manufacturer, the number of fringes \( N \) is varied from tens to hundreds, the distance between fringes (\( \delta \)) is from single to tens of microns, and the overlap (\( l \)) is typically of few millimetres. Such design allows increasing significantly the effective area between electrodes (\( A \)): \( A = l \times \delta \times (2N - 1) \) For example, DropSens produce platinum or gold interdigitated electrode assemblies having 50 or 100 fringes with the spacing of 5\( \mu m \) or 10 \( \mu m \). The AC impedance measurements are usually carried out using impedance analysers operating over a wide frequency range from fractions of Hz up to hundreds of MHz. The dependences of impedance (\( Z \)) versus frequency (\( f \)) called as impedance spectrum is recorded at small amplitudes of AC voltage, typically 100mV, with or without DC off-set. The impedance spectrum obtained can be presented in different ways. For example as spectra of real (\( Z' \)) and imaginary (\( Z'' \)) parts of impedance known as a Nyquist plot. Alternatively, the spectrum
can be presented as spectrum of parallel conductance (G_p) vs capacitance (C_p), or spectrum of the magnitude of Z and phase angle (tanθ). The results obtained are analysed using dedicated software, which allows fitting the data to the equivalent circuit model and evaluate its parameters.

Typical equivalent circuits explaining the majority of experimental features of impedance measurements in metal-electrolyte–metal systems shown in Figure (3-11) splits in two parts: (i), the electrolyte solution which is described by the resistance of solution (R_S), and (ii), a double layer near the metal electrodes, which represents polarisation of the capacitance of a double layer C_DL in parallel with resistor (R_DL). An addition, a constant phase element Zw is associated with the diffusion of chemicals in the vicinity of the double layer; at frequencies higher than 10Hz this element can be neglected (see Fig. 3-17 a,b).

**Figure 3-17:** Equivalent circuit for impedance measurement in metal-electrolyte-metal system (a), simplified equivalent circuit (b).

It was assumed that two electrodes are identical, thus elements related to double layers on two electrodes can be combined be in one parallel circuit.

The impedance of the simplified circuit can be calculated as
\[ Z = Z' - jZ''; \quad Z' = \frac{R_{DL}}{1 + \omega^2 R_{DL}^2 C_{DL}^2} + R_S; \quad Z'' = \frac{\omega R_{DL}^2 C_{DL}}{1 + \omega^2 R_{DL}^2 C_{DL}^2}. \]

At low frequencies \((\omega \to 0)\) \(Z' = R_{DL} + R_S\) and \(Z'' = 0\), while at high frequencies \((\omega \to \infty)\) \(Z' = R_S\) and \(Z'' = 0\).

The reciprocal of impedance is admittance (i.e., admittance is the current-to-voltage ratio). Impedance is represented as a complex quantity \(Z\) and the term ‘complex impedance’ and real part is the resistance \(R\), and the imaginary part is the reactance \(X\). The reactance and impedance of a capacitor are respectively, where \(\omega\) is the angular frequency of the sinusoidal signal and \(j\) phase indicates. (see Appendix-B).

The results of impedance spectra measurements are often presented as the dependence of \(Z''\) vs \(Z'\) known as a Nyquist plot (see schematic graph in Figure 3-18 a,b).

**Figure 3-18:** Nyquist plot \(Z''\) vs \(Z'\); Dotted line indicate the outcome of binding analyte to receptor (a) schematic graph of the affinity (b).

Typically, the affinity of reactions, i.e. binding the analyte to receptor immobilized on the surface of interdigitated electrodes, causes changes in the Nyquist plot as shown
schematically in Figure 3-18 (a,b). The conductometric sensor response can be easily quantified by the parameter $R_{DL}$ which can be easily extracted from data using appropriate data fitting software.

Alternatively, $Z$ can be presented by parallel conductivity (or resistivity) and $G_p$ ($R_p$) and capacitance $C_p$.

$$Z = \frac{R_p}{1 + \omega^2 R_p^2 C_p^2} - j \frac{\omega R_p^2 C_p}{1 + \omega^2 R_p^2 C_p^2},$$

(3.7)

Both representations are very close. For example, at low frequencies ($\omega \to 0$)

$$G_p = \frac{1}{R_{DL} + R_S}, \quad C_p = C_{DL} \frac{R_{DL}^2}{(R_{DL} + R_S)^2},$$

while at high frequencies ($\omega \to \infty$) $G_p \to \infty$ and $C_p = C_{DL}$.

One more possibility is to present the results as spectra of $|Z|$ and $\tan \theta$ which can be easily worked out from eq (3.7). The critical cases are: $|Z| = R_{DL} + R_S$ and $\tan \theta = 0$ when $\omega \to 0$, $|Z| = 0$ and $\tan \theta = 0$ when $\omega \to \infty$. The choice of representation of the experimental results and their analysis depends on the particular application.

Therefore, electrochemical methods (both DC and AC) are very common in bio-sensing due to several advantages mainly high sensitivity, low-cost, and simplicity of use. Amperometric sensors based on DC measurements, e.g. CV, are typically associated with enzyme sensors, while conductometric sensors based on AC measurements can be utilised for detection of affinity reactions such as immune reactions.
References

CHAPTER 4 Optical and electrochemical detection of toxic pollutants: data obtained on bacterial suspensions

This chapter covers in more detail the experimental procedure of bacterial growth in culture media and focused on the optical and electrochemical data of effect three types of environmental pollutants i.e. heavy metals ($\text{Hg}^{2+}$, $\text{Pb}^{2+}$, $\text{Zn}^{2+}$ and $\text{Cd}^{2+}$), pesticides (atrazine, simazine and DDVP) and petrochemicals (hexane, pentane, pyrene, toluene, octane and ethanol), on three types of bacterial isolates which are $\text{E. coli}$ K12, $\text{S. oneidensis}$ MR-1, and the methanotrophic bacteria ($\text{Mc. capsulatus}$ Bath or $\text{Ms. trichosporium}$ OB3b). The effects of these toxic chemicals on the optical and electrochemical properties of bacteria in suspensions are measured.

4.1 Bacterial culture conditions

Three diverse bacterial strains were selected for this work: (i), the model Gram negative $\text{E. coli}$ K12, known to be sensitive to various types of pollutants including heavy metals, pesticides, and hydrocarbons [1] (ii), $\text{S. oneidensis}$ MR-1, a Gram-negative bacterium known to tolerate and interact with heavy metals [2], and (iii), methanotrophic bacteria ($\text{Mc. capsulatus}$ Bath and $\text{Ms. trichosporium}$ OB3b) which are Gram-negative bacteria that grow on methane and are also able to co-oxidise a range of other hydrocarbons and hydrophobic organic molecules [3,4]. $\text{E. coli}$ cultures were grown at 37 °C for 16 h in LB broth and LB agar [5], $\text{S. oniedensis}$ cultures were grown at 30 °C for 24 h in the same medium [6]., while methanotrophs bacteria ($\text{Mc. capsulatus}$ Bath or $\text{Ms. trichosporium}$ OB3b) were grown at 30 °C for 2 weeks in flask cultures in nitrate mineral salts (NMS) medium and on NMS agar plates, using methane as the carbon source, as described previously [7].

4.2 Preparation of analyte samples

The samples of $\text{Hg}^{2+}$, $\text{Pb}^{2+}$, $\text{Zn}^{2+}$, $\text{Cd}^{2+}$, atrazine, simazine, DDVP, hexane, pentane, pyrene, toluene, octane and ethanol (all from Sigma-Aldrich) were prepared at concentrations of 0.1, 1, 10, 100 mM by consecutive dilution of 1 M stock solutions in
deionised water. The stock solutions of hydrocarbons were prepared in a 40% (v:v) ethanol: water mixture. Liquid bacterial culture samples were mixed with these solutions in 1:1 ratio and incubated for 2 hr at (22-25) °C.

4.3 Optical Measurements of bacteria suspension samples

The numbers of live and dead bacteria were determined with fluorescence microscopy and OD$_{600}$ similarly to that described in [8] and earlier in Chapter 3. Live and dead bacteria appeared under microscope as green and red cells, respectively. Typical fluorescence microscopy images are shown in Figure (4-1) for E. coli, Figure (4-2) for Ms. trichosporium OB3b and Figure (4-3) for S. oneidensis, respectively.

Figure 4-1: Fluorescence microscopy images of E. coli before (A) and after (B) treatment with HgCl$_2$ (1 M) for 2 hr.

Figure 4-2: Fluorescence microscopy images of Ms. trichosporium (OB3b) before (A) and after (B) treatment with HgCl$_2$ (1 M) for 2 hr.
It is clear that the exposure to HgCl$_2$, (simazine, and octane which will be shown in appendix-A1,2) reduces the number of live (green) bacteria and increases the dead ones (red), though $S.$ $oneidensis$ are much less affected than $E.$ $coli$ and $Ms.$ $trichosporium$ (OB3b). A similar though even more pronounced pattern was observed in flow cytometry experiments where bacteria were stained with the same L7012 Live/Dead Bacterial Viability Kit. The results in Figure 4-4, show live and dead bacteria as blue (live) and orange (dead) dots respectively. Increase in the dead $E.$ $coli$ bacteria count after exposure to HgCl$_2$ salt, (simazine, and octane which will be shown in appendix-A1,2) is visually apparent. Analysis of Figure 4-4A(b) yields the percentage of live $E.$ $coli$ cells as 23.88% and 76.12% for dead cells. In addition, dead $E.$ $coli$ bacteria appear mostly in the bottom-left quadrant of the graph in Figure 4-4A(b) indicating that increase in the bacterial size is most-likely due to the damage of cell membranes. In contrast, $S.$ $oneidensis$ bacteria were affected much less, the percentages of live and dead bacteria after exposure to 1M HgCl$_2$ solution were 83.36% and 16.64%, respectively. Again, dead bacteria appeared slightly enlarged since they were shifted to the bottom-left in Figure 4-4 B (b). Figure 4-4 C (a , b) shows flow cytometry results for $Ms.$ $trichosporium$ (OB3b), where the percentage was 43.57% for live bacteria and 56.43% for dead bacteria.
Figure 4-4: Flow cytometry results for *E. coli* Bacteria (A), *S. oneidensis* (B) and *Ms. trichosporium* (OB3b) (C); graphs (a) and (b) were obtained, respectively, before and after treatment with HgCl$_2$ (1M) for 2 hr.

The result of optical density measurements of all bacteria species and the effect of their treatment with HgCl$_2$, simazine, and octane prepared at different concentrations are shown in Table 4-1. The bacterial density was assessed and presented as absorbance by losses of light intensity in the middle of visible range (600nm) as a result of light scattering on the bacteria. The reduction in optical density upon increasing the HgCl$_2$ concentration was much more pronounced for *E. coli* and *Ms. trichosporium* (OB3b) than that for *S. oneidensis*. 
Table 4-1: The results of OD600 for all bacteria samples after exposure to HgCl$_2$ for 2 hr.

<table>
<thead>
<tr>
<th>Bacteria</th>
<th>HgCl$_2$ concentration</th>
<th>STD(n=3)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Before treatment</td>
<td>0.1mM</td>
</tr>
<tr>
<td>E. coli</td>
<td>0.813</td>
<td>0.773</td>
</tr>
<tr>
<td>S. oneidensis</td>
<td>0.827</td>
<td>0.869</td>
</tr>
<tr>
<td>Ms. trichosporium OB3b</td>
<td>0.754</td>
<td>0.728</td>
</tr>
</tbody>
</table>

Among the three optical methods used, flow cytometry appeared to be the most reliable and not affected by different motility of *E. coli*, *Ms. trichosporium* (OB3b) and *S. oneidensis*. It is known that dead *E. coli* and *Ms. trichosporium* (OB3b) bacteria are not motile and tend to sediment which may affect the results of static fluorescent microscopy and optical density measurements. Nevertheless, the results of optical testing of bacterial samples provided a background for further study using much simpler electrochemical methods.

**4.4 Electrochemical measurements on bacterial suspension samples**

**4.4.1 CV measurements**

All CV electrochemical measurements were carried out on a DropSTAT4000P potentiostat instrument (from DropSens) controlled by Autolab software using DropSens screen printed gold electrodes (SPGEs). These electrodes have a conventional three electrode configuration with gold working electrode (4-mm diameter disk) and counter electrode (16 mm×1.5 mm curved line), and the potentials were ± 0.2 and ±0.5, at a scan rate of 100mV/s recorded against Ag/AgCl (16 mm×1.5 mm straight line) pseudo-reference electrode. Typical cyclic voltammograms for *E. coli*,
*Ms. trichosporium* (OB3b) and *S. oneidensis* of different cell densities (i.e. dilutions with nutrient broth) are shown in Figure (4-5 A, B ,C).
Figure 4-5: Cyclic voltamgram recorded on *E. coli* (A), *Ms. trichosporium* (OB3b) (B) and *S. oneidensis* (C) of different dilutions (1:1, 1:5, 1:10 and stock solutions); CV curves for clear broth are shown on all the graphs.

Generally, the CV graphs in Figure (4-5) are almost featureless for all bacterial species in the selected voltage range from -0.5V to +0.5V which was chosen deliberately in order to avoid electrochemical reactions on the electrodes. A slight increase in the anode current at -0.2V indicates the beginning of hydrogen reduction. The values of cathode current at -0.5V appear to decrease with the increase in bacteria concentration (or dilution ratio 1:10, 1:5, 1:2, 1:1) with the largest current shown for clear LB broth and the lowest for undiluted bacterial stock suspension. This indicates that the bacteria adsorbed onto the surface of gold electrodes and act as an insulating layer, reducing the current. These data are very important since they establish a correlation between the values of cathode current and bacterial cell density in the LB broth.
The next step was to study the effect of Hg$^{2+}$ ions, simazine, and octane on CV characteristics of *S. oneidensis*, *Ms. trichosporium* (OB3b) and *E. coli* bacteria. A series of CV measurements were carried out on liquid samples of bacterial suspensions in LB broth which was mixed in 1:1 ratio with different concentrations of HgCl$_2$, simazine, and octane and kept for 2 hours in (22-25)$^\circ$C prior to CV measurements. The results of these measurements in Figure (4-6) show substantial increase in cathode current (Ic) for *E. coli* samples (Figure 4-6A) upon increasing the concentration of HgCl$_2$, simazine, and octane. This effect was much less pronounced for *S. oneidensis* samples (Figure 4-6B). In addition, oxidation and reduction peaks appeared on CV curves for *S. oneidensis* and *Ms. trichosporium* (OB3b) samples in (Figure 4-6A,C) which are definitely related to electrochemical reactions associated with the presence of HgCl$_2$ and octane. The anodic peak increases with the increase in HgCl$_2$ concentration. However, such electrochemical reactions do not appear with *E. coli* samples in (Figure 4-6B). It is known that HgCl$_2$ and octane act as cofactors for *S. oneidensis* and *Ms. trichosporium* (OB3b) bacterial cell growth [9].
Figure 4-6: Cyclic voltammogram of E.coli (A) S. oneidensis (B) and Ms. trichosporium (OB3b) (C) bacteria solutions in nutrient broth which treated with different concentration of HgCl$_2$. 
When analysing the effect of heavy metal salts on CV characteristics of liquid bacterial culture samples the effect of extra Hg$^{2+}$ and Cl$^{-}$ ions on conductivity of liquid medium has to be taken into account. In order to find out the true effect of heavy metal ions on the bacteria, the values of anode current ($I_A$) of bacterial samples has to be normalized by the reference current $I_{ref}$ of the LB broth diluted 1:1 with particular concentration of HgCl$_2$. The values of relative changes of anode current $\Delta I_A/I_{A0} = (I_A - I_{A0})/I_{A0}$ at -0.5V of *S. oneidensis*, *Ms. trichosporium* OB3b and *E. coli* bacteria are plotted in Figure 4-7 against the concentration of HgCl$_2$.

![Figure 4-7: The dependence of relative changes in cathode current at –0.5 V for S. oneidensis, M. trichosporium OB3b and E. coli on the concentration of HgCl$_2$.](image)

The effects of HgCl$_2$ on *S. oneidensis*, *trichosporium* OB3b and *E. coli* are completely different: $\Delta I_A/I_{ref}$ goes up with the increase in HgCl$_2$ concentration for *E. coli* and *M. trichosporium* OB3b which means that *E. coli* and *M. trichosporium* OB3b bacteria are
inhibited by Hg$^{2+}$ ions becoming less electrically resisting, while $\Delta I_{N}/I_{ref}$ is almost flat at low concentrations of HgCl$_2$ and slightly increases at high concentration of 1M. This means that *S. oneidensis* are practically not affected by HgCl$_2$ at low concentrations but inhibited at high concentrations. This is a very promising result showing a possibility of pattern recognition of heavy metals using the two bacteria.

### 4.4.2 Electrochemical Impedance Spectroscopy measurement

Impedance spectra were measured using an impedance analyzer (2000A) and gold interdigitated electrodes (from DropSens), 5 µm dimension for bands/gaps are available with reference number (G-IDEAU5) and the overlapping length is 6.76 mm. The AC voltage amplitude was 5 mV with the frequency varied from 100 mHz to 100 kHz; no DC bias was applied. Similarly to the CVs, the impedance spectra measurements were carried out on electrodes coated with bacterial suspension which were treated with solutions containing different concentrations of pollutants. Typical results are shown as Nyquist plots in Figure (4-8).

**Figure 4-8:** The Nyquist plots (-Zim vs Zre) for interdigitated electrodes with *E. coli* bacteria suspension treated with Hg$^{2+}$ ions of different concentrations (A); equivalent circuit (B).
The analysis of impedance spectra using an equivalent circuit model (shown as inset in Figure 4-8 was carried out according to the simplified circuit model impedance when the diffusion impedance \( Z_{\text{diff}} \) is neglected [10]. Study of the inhibition effect of toxic pollutants on optical and electrochemical properties of bacteria in suspension from proved the principle of pollution detection, from a practical point of view, the use of bacteria immobilized on the surface electrodes is much more promising for such sensor development.

A decrease in impedance was observed with increasing \( \text{Hg}^{2+} \) concentration, as evidenced by the decreasing height and diameter of the semi-circular Nyquist traces, which correspond to decreased capacitance and resistance of the sensor surface respectively. \( R_s \) (solution resistance), \( R_{\text{dl}} \) (double layer resistance) are presented in Table 4-2. Good stability of the sensor was observed as the solution resistance was nearly constant at all stages of biosensor construction and testing; while \( R_{\text{dl}} \) decrease due to treat the bacteria with \( \text{Hg}^{2+} \) ions.

**Table 4-2:** Values for the EIS parameters obtained from fitting the Nyquist plots shown in Figure 4-8 to the equivalent circuit model.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>( R_s ) (Ω)</th>
<th>( R_{\text{dl}} ) (Ω)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacteria suspension</td>
<td>54.903</td>
<td>172.63</td>
</tr>
<tr>
<td>0.1 ( \text{µM} ) \text{Hg}^{2+}</td>
<td>50.529</td>
<td>143.75</td>
</tr>
<tr>
<td>1 ( \text{µM} ) \text{Hg}^{2+}</td>
<td>48.726</td>
<td>125.82</td>
</tr>
<tr>
<td>10 ( \text{µM} ) \text{Hg}^{2+}</td>
<td>48.038</td>
<td>105.14</td>
</tr>
<tr>
<td>100 ( \text{µM} ) \text{Hg}^{2+}</td>
<td>53.414</td>
<td>93.666</td>
</tr>
<tr>
<td>1000 ( \text{µM} ) \text{Hg}^{2+}</td>
<td>51.053</td>
<td>81.606</td>
</tr>
</tbody>
</table>
References
CHAPTER 5 Optical and electrochemical detection of toxic pollutants: data obtained on immobilized bacteria

The results given in the previous chapter are important as a preliminary step towards the development of bacteria-based inhibition sensor array, but it is still far away from real sensor development. Dealing with liquid bacterial samples is not the way forward because of natural variations of bacteria concentration even in laboratory samples not to mention “real” samples taken for analysis. This chapter will focus on the problem of having a reliable reference for such measurements. It would be much more useful for sensor array development to use bacteria immobilized on the electrode surface.

5.1 Bacterial immobilization process

Three selected types of bacteria e.g. *E. coli*, *Mc. capsulatus* Bath and *S. oneidensis* were immobilized on the surface of the screen-printed gold electrodes modified with poly L-lysine (PLl) [1, 2]. This was achieved by incubating samples in 1:1000 mixture on Dropsens electrodes (Figure 5-1 a,b) of PL1 (0.1 mg/ml) with deionized water for 1 h at 37 °C. Then bacteria were immobilized by dropping stock suspensions of *E. coli*, *Mc. capsulatus* Bath and *S. oneidensis* on the modified electrodes, keeping it there for 1 h, then washing out non-bound bacteria with PBS as illustrated in Figure (5-1c). The electrodes with immobilized bacteria could be kept at 4°C for 24 hours without compromising bacterial activity.

**Figure 5-1**: DropSens three-electrode assembly (a), DropSens interdigitated electrodes (b), Schematic diagram of bacteria immobilization procedure (c).
5.2 Preparation of analyte solutions

The inhibition effects on the above mentioned bacteria was studied by exposing them to the following chemicals (pollutants): HgCl$_2$, PbCl$_2$, ZnCl$_2$ and CdCl$_2$, atrazine, simazine, DDVP, hexane, pentane, octane, ethanol, toluene and pyrene. Solutions of different concentrations (0.1, 1, 10, 100, and 1000 µM) were prepared by multiple dilution of 1 mM stock solution of each analyte dissolved in deionised water. 40% ethanol solution in water was used for dissolving the hydrocarbons, including toluene and pyrene. The samples of immobilized bacteria were treated by immersing them into the required solutions of the above chemicals for 2 hours.

5.3 Optical and SEM characterization of immobilized bacteria

5.3.1 Fluorescent microscopy study of immobilized bacteria

In this section we deployed fluorescent microscopy for characterization of bacteria immobilized on the surface of screen printed gold electrode. Fluorescence microscopy images in Figures (5-2 and 5-3) show the effect of Pb$^{2+}$ and Zn$^{2+}$ ions on *S. oneidensis* bacteria immobilized on modified screen printed gold electrodes where live and dead bacteria appear as green and red spots, respectively. It is clear that the exposure to 1M solution of PbCl$_2$, ZnCl$_2$ and CdCl$_2$ for 2 hours reduced the number of live bacteria (green spots) and increases the dead ones (red spots). Such experiments were carried out for all three types of bacteria and all analyte types used. The results of this study are presented in Figure (5-2) and Figure (5-3) and Table 5-1 as the numbers of live (green) and dead (red) bacteria on recorded images of identical dimensions.

5.3.2 SEM study of immobilized bacteria

For scanning electron microscopy (SEM) immobilized bacteria were fixed on double-sides carbon tape mounted on a sample holder and coated with a few-nanometer-thick
layer of carbon using a carbon evaporator (Edwards E306A; Edwards, United Kingdom). The samples were examined with a LEO1550VP field emission scanning electron microscope equipped with an Oxford INCA energy-dispersive X-ray spectrophotometer. The system was operated at 1 to 15 kV for high-resolution secondary electron imaging and elemental analysis (see Figure 5-4).

5.3.3 Fixation of immobilized bacteria samples for SEM measurements
Bacteria amended culture (1.5 mL) were pelleted by centrifugation (11000 × g; 10 min; room temperature), and washed with 0.1 M sodium phosphate buffer (pH 7.4). The specimens were then fixed in 3% (v/v) glutaraldehyde in the same buffer overnight at room temperature and washed again in the same buffer. Secondary fixation was carried out in 1% (w/v) aqueous osmium tetroxide for 1 hour at room temperature followed by the same wash step. Fixed cells were dehydrated through a graded series of ethanol dehydration steps 75%, 95% and 100% (v/v), and then placed in a 50/50 (v/v) mixture of 100% ethanol and 100% hexamethyldisilazane for 30 min followed by 30 min in 100% hexamethyldisilazane [3].

5.3.4 AFM imaging of immobilized live bacteria
A drop of the bacterial suspension was deposited on the surface of the screen printed gold electrode and characterized by atomic force microscopy. The immobilized bacterial cells were imaged in 5500 AFM microscope outfitted with a 90 μm scanner in combination with an inverted Olympus IX81 light microscope. Silicon nitride cantilevers (0.1 N/m) were used to scan the sample in non-contact mode at a rate of 6–13 μm/s at 512 or 256 points per line resolution. Images were processed on first-order flattening (Figure 5-5 A,B) [4].
Figure 5-2: Fluorescence microscopy images of immobilized *S. oneidensis* bacteria before (A) and after (B) treatment with PbCl$_2$ (1 M) for 2 hours.

Figure 5-3: Fluorescence microscopy images of immobilized *S. oneidensis* before (A) and after (B) treatment with ZnCl$_2$ (1M) for 2 hours.
Table 5-1: The numbers of live and dead bacteria immobilized on microscopic images of modified screen printed gold electrodes obtained from fluorescence microscopy images for all three bacteria before and after treatment with 1M solutions of the three pollutants for 2 hours.

<table>
<thead>
<tr>
<th>Types of Bacteria used</th>
<th>Types of Pollutants</th>
<th>Bacteria count Before exposure</th>
<th>After exposure</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Live</td>
<td>Dead</td>
<td>Live</td>
</tr>
<tr>
<td>Escherichia coli</td>
<td>93</td>
<td>20</td>
<td>21</td>
</tr>
<tr>
<td>Shewanella oneidensis</td>
<td>149</td>
<td>22</td>
<td>72</td>
</tr>
<tr>
<td>Mc. capsulatus Bath</td>
<td>43</td>
<td>13</td>
<td>16</td>
</tr>
<tr>
<td>Escherichia coli</td>
<td>81</td>
<td>25</td>
<td>18</td>
</tr>
<tr>
<td>Shewanella oneidensis</td>
<td>79</td>
<td>18</td>
<td>15</td>
</tr>
<tr>
<td>Mc. capsulatus Bath</td>
<td>62</td>
<td>17</td>
<td>19</td>
</tr>
<tr>
<td>Escherichia coli</td>
<td>69</td>
<td>21</td>
<td>20</td>
</tr>
<tr>
<td>Shewanella oneidensis</td>
<td>57</td>
<td>11</td>
<td>28</td>
</tr>
<tr>
<td>Mc. capsulatus Bath</td>
<td>75</td>
<td>19</td>
<td>71</td>
</tr>
</tbody>
</table>

Analysis of fluorescence microscopy data in Table. (5-1) revealed that *E. coli* and *Mc. capsulatus* Bath were badly affected by large concentrations of Hg\(^{2+}\) ions, while *S. oneidensis* are less affected. The negative effect of atrazine is dramatic and more or less similar for all three bacteria. Toluene, however, did not affect *Mc. capsulatus* Bath; though it inhibited both *E. coli* and *S. oneidensis*. Such behaviour of immobilized bacteria is similar to those bacteria in solution.
The direct evidence of cell enlargement was obtained from SEM study. SEM images in Figure 5-4 show the enlargement of *E. coli* bacteria (Figure 5-4B) and rupture of *Mc. capsulatus Bath* bacteria cells (Figure 5-4D) caused by exposure to high concentration (1M) of ZnCl$_2$. These observations are similar to previously reported SEM studies of bacteria [5,6].
Figure 5-5: AFM images of modified SPGE before (A) and after (B) immobilized *E.coli* bacteria with poly-L-lysine.

In contrast, *S. oneidensis* bacteria were affected much less by ZnCl₂ than the other bacterial strains and appeared slightly elongated. Similar elongation has been observed in *S. aureus* due to exposure to high salt concentration as a specific response to other stress conditions [7]. Also, a significant increase in bacteria length was found in *S. oneidensis* exposed to UV radiation [8].

### 5.4 Electrochemical measurements of immobilized bacteria samples

#### 5.4.1 Cyclic voltammograms (CVs) measurement

The effect of HgCl₂, atrazine, and toluene on all three bacteria, in immobilized bacteria using cyclic voltammograms (CVs) was studied. Typical series of CVs recorded on *E. coli*, *S. oneidensis*, and *Mc. capsulatus* Bath samples are shown in Figure (5-6). The graphs of CV in Figure (5-6) show a characteristic peak of anodic current at about 0.3V and peak of cathodic current between -0.1V and -0.2V which are most likely correspond to oxidation and reduction reaction of PBS buffer. The current peak values which are maximal on bare electrodes in PBS reduced substantially on the electrodes with immobilized bacteria which act as an insulating layer reducing both anodic and cathodic currents. Exposure of bacteria to toxic chemicals probably causes damage to bacteria.
cell wall thus affecting their insulating properties which is why the peak currents are rising with the increase in pollutants concentrations. (see Appendix - A3)
Figure 5-6: Cyclic voltammograms in the selected voltage range from (-0.5 V to +0.5 V) and scan rate at 100mV/s, for: immobilized *E.coli* treated with toluene from 0.1 µM-1mM concentration (A), immobilized *Mc. capsulatus* Bath treated with atrazine from 0.1 µM-1mM concentration (B), and immobilized *S. oneidensis* treated with HgCl₂ from 0.1 µM-1mM concentration (C).

In Figure (5-6A,B,C) the CV cycles appear to shift upwards upon increasing the pollutants concentration from 0 (untreated bacteria) to 0.1 mM, 1mM, 10 mM, 100 mM, and 1M. The characteristic parameter in this study, e.g. the value of anodic current at +0.5 V increases with the increase in pollutant concentration for all three bacteria in both liquid and immobilized forms. This means that the electrical conductivity is controlled by bacteria adsorbed on the surface of the surface of screen printed gold electrodes and acting as insulating layer which reducing the current. The correlation between bacterial concentration and the electric current (or conductivity) values is very important for further study of the effect of pollutants, and such measurements were
always carried out first [9]. The presence of pollutants (Hg^{2+} ions, atrazine, and toluene in our case) causes damage to the bacterial cells, and therefore bacteria became less insulating, in-turn leading to an increase in the anodic current, which is observed in Figure 5-6 (A,B,C).

Similar analysis could have been carried out using cathodic peaks, however the use of anodic peaks is sufficient for this study. To analyze the effect of pollutants on electrical properties of immobilized bacteria, the values of anodic current ($I_A$) at +0.5V from CV measurements were normalized to the currents values of uncoated electrodes in PBS with the addition of a particular pollution of particular concentrations ($I_{A0}$) to construct the values of relative changes of anodic current. $\Delta I_A/I_{A0} = (I_A - I_{A0})/I_{A0}$ For example, for S. oneidensis bacteria treated with 1mM solution of PbCl$_2$ (Figure 5-6 C), the reference was recorded on uncoated electrodes in PBS containing 1mM PbCl$_2$.

The relative changes in anodic current are presented in Figure (5-7) for all three bacteria studied as concentration dependences of the three pollutants. As one can see the effects of HgCl$_2$, atrazine, and toluene on S. oneidensis, Mc. capsulatus Bath and E. coli are completely different. E. coli appeared to be affected by HgCl$_2$, atrazine, and toluene even at low concentrations since the $\Delta I_A/I_{A0}$ values increase monotonically in Figures 5-7 (A), 5-7 (B), and 5-7 (C), respectively. This means that E. coli is equally inhibited by all three pollutants and becoming less electrically resisting. In contrast, S. oneidensis is almost unaffected by HgCl$_2$ at low concentrations of all pollutants up to 10mM, and then $\Delta I_A/I_{A0}$ started to increase at high concentrations of 100mM and 1M.
Relative changes in $I_A$ for different concentrations of HgCl$_2$ (A) and Atrazine (B) for E.coli, M.capsulatus, and S.oneidensis.
Such behaviour of immobilized *E. coli* and *S. oneidensis* bacteria is similar to those free in liquid as reported in our previous study. *M. capsulatus* (Bath) respond to HgCl₂ (Figure 5-7A) and atrazine (Figure 5-7B) similarly to the other two bacteria studied though the changes in $\Delta I_A/I_{A0}$ are more pronounced at high concentrations, particularly for atrazine. However, *M. capsulatus* (Bath) bacteria are not affected by toluene (see Figure 5-7C) even at high concentration; moreover an overall trend to small decrease in $\Delta I_A/I_{A0}$ is observed. Such behavior was expected since *M. capsulatus* (Bath) consume some hydrocarbons [10]. (see Appendix - A3 for more results of author immobilized bacteria)
5.5 Sensor array data analysis

5.5.1 Identification of water pollutants using pseudo 3D plots of sensor responses

All the data obtained from the sensor array containing three electrodes functionalized by three different types of bacteria, namely *E. coli*, *S. oneidensis*, and *M. capsulatus*, treated by 12 different pollutants, e.g. heavy metal ions (Hg$^{2+}$, Pb$^{2+}$, and Cd$^{2+}$), pesticides (atrazine, simazine, and DDVP) and petrochemicals (hexane, octane, pentane, ethanol, pyrene and toluene) at concentrations of 0.1, 1, 10, 100, 1000 µM were analysed using pseudo-3D plots of sensor responses in Figure 5-8 (A, B, C). In Figure 5-8A the experimental points for toluene, simazine and HgCl$_2$ are well separated up to their concentrations of 100 µM. Similarly, in Figure 5-8B, the data for pyrene, DDVP and PbCl$_2$ are well separated up to their concentrations of 100 µM. Also octane, atrazine and CdCl$_2$ show similar pattern in Figure 5-8C. An attempt of pattern recognition has been done by presenting the relative responses of the three channels, e.g. three bacteria (*E. coli*, *M. capsulatus* (Bath) and *Shewanella oneidensis*) immobilized on three screen-printed electrodes, to the three pollutants (HgCl$_2$, atrazine, and toluene) in a pseudo-3D plot in Figure 5-8(A, B, C). It can be also noticed that heavy metal ions (Hg$^{2+}$, Pb$^{2+}$, and Cd$^{2+}$) give responses mostly in the "north-west" section of respective 3D graphs, while pesticides (atrazine, simazine, and DDVP) appeared in the "south", "south-west" sections, and petrochemicals (hexane, octane, pentane, ethanol, pyrene and toluene) lie mostly in the "north-east" sections. This is a clear indication that pattern recognition principles can be applied for identification of toxic pollutants using different types of bacteria. The concentration of these pollutants could be evaluated too using the appropriate calibration and data extrapolation.
Figure 5-8: 3D plot of relative changes $I_a/I_{ref}$ in anodic current for *E.coli*, *M. capsulatus*, and *S. oneidensis* caused by different pollutants. Arrows show the direction of the pollutants’ concentration increase from 0.1 mM to 100 mM.

5.5.2 Electrochemical Impedance Spectroscopy (EIS) measurements

Impedance spectra were measured using an impedance analyzer (4000A) and gold interdigitated electrodes (from DropSens), 5 μm dimension for bands/gaps are available with reference number (G-IDEAU5) and the overlapping length is 6.76 mm. The AC voltage amplitude was 5 mV with the frequency varied from 100 mHz to100 kHz; no DC bias was applied. Similarly to CVs, the impedance spectra measurements were carried out on electrodes modified with immobilized bacteria which treated with solutions containing different concentrations of pollutants. A typical example of data is shown as Nyquist plots in Figure (5-9) for immobilized *E. coli* treated with different concentrations of pyrene. (see Appendix-A4 figures 1,2).
Figure 5-9: The Nyquist plots (-Zim vs Zre) for interdigitated electrodes modified with immobilized *E.coli* bacteria treated with pyrene of different concentrations (A); equivalent circuit (B).

The EIS data are presented as Nyquist plots in Figure 5-9. The analysis of impedance spectra using an equivalent circuit model shown as inset in Figure (5-9 B) was carried out according to the simplified circuit model impedance (when the diffusion impedance $Z_{\text{diff}}$ is neglected) [10]. A decrease in impedance was observed with increasing $\text{Hg}^{2+}$ concentration, as evidenced by the decreasing height and diameter of the semi-circular Nyquist traces, which correspond to decreased capacitance and resistance of the sensor surface respectively. Rs (solution resistance), $R_{\text{dl}}$ (double layer resistance) are presented in Table 5-1. Good stability of the sensor was observed as the solution resistance was nearly constant at all stages of biosensor construction and testing. Whilst the $R_{\text{dl}}$ values decreased upon addition of $\text{Hg}^{2+}$, showing a clear trend. The effect of analyte to a
biosensor surface causes an increase in impedance, and this has been widely reported [11].

Table 5-1: Values for the EIS parameters obtained from fitting the Nyquist plots shown in Figure 5-9 to the equivalent circuit model.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Rs (Ω)</th>
<th>Rd (Ω)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Immobilized bacteria</td>
<td>50.986</td>
<td>758.47</td>
</tr>
<tr>
<td>0.1 µM Hg²⁺</td>
<td>49.366</td>
<td>433.25</td>
</tr>
<tr>
<td>1 µM Hg²⁺</td>
<td>48.205</td>
<td>297.08</td>
</tr>
<tr>
<td>10 µM Hg²⁺</td>
<td>45.510</td>
<td>239.33</td>
</tr>
<tr>
<td>100 µM Hg²⁺</td>
<td>43.311</td>
<td>112.17</td>
</tr>
<tr>
<td>1000 µM Hg²⁺</td>
<td>40.412</td>
<td>74.777</td>
</tr>
</tbody>
</table>

5.6 Discussion of the optical and electrochemical measurement results

The observed effects of the above pollutants on the three selected bacteria are somehow expected. In general terms, different chemicals of both organic and inorganic origin may affect microorganisms in two possible ways, e.g. acting as either catalyzers enhancing bacterial metabolism or as inhibitors having an opposite effect of reducing bacteria metabolism and even damaging bacteria membranes and causing their death. In our case, *E. coli* is obviously inhibited by the pollutants used. This results in the reduction of live bacteria count which was confirmed by the optical study. Consequently, the increased number of damaged or dead bacteria reduces their insulating properties, thus causing an increase in both anodic and cathodic currents as well as the reduction in resistance (Rs) is very small and (Rd) is much effected in impedance spectroscopy. *Shewanella oneidensis* bacteria are known to be tolerant to heavy metals at low concentration, which may have even be growth stimulating, and which can be used in water treatment [12]. High concentrations of heavy metals are damaging. This explains the observed tolerance of *S. oneidensis* to heavy metals at low concentrations, while
other pollutants are still acting as inhibitors. *M. capsulatus* (Bath), in contrast, are known by their ability to use some organic chemicals (hydrocarbons, alcohols) as a food source [13], and therefore are used in sewage treatment [14]. In other words, *M. capsulatus* bacteria are stimulated by some petrochemicals, while heavy metals and pesticides still act as inhibitors. Optical and electrochemical study of both *S. oneidensis* and *M. capsulatus* (Bath) showed the characteristics changes, respectively, in the live bacterial concentration and anodic current in line with their expected catalytic-inhibition patterns. Combining the above three types of bacteria in a sensor array was logical and therefore enabled the array to identifying the type of pollutants. This could be achieved using optical methods with flow cytometry being perhaps the most suitable method for this task.

However, very simple electrochemical measurements of anodic current could do a similar a job at substantially reduced cost. Modified screen-printed electrodes with immobilized bacteria can be prepared in advance and kept active for few weeks when stored at 4 °C. Such electrical tests can be used for quick preliminary analysis of water samples. The samples indicating a presence of certain pollutants can be passed to specialized laboratories for further more detailed and accurate testing. The overall cost and time of analysis would be substantially reduced as a result. The sensor stability depends on the activity of the immobilized bacteria. We found that bacteria were still alive and active after 24 hours storing in the refrigerator at 4 °C. After 48 hours the live bacterial concentration reduced slightly (10-15%), and after 72 hours reduced further to over 50%. Therefore, we can conclude, that currently the sensors stability is limited to 24 hours. Ideally, electrodes with freshly immobilized bacteria have to be used for sensing.
References

CHAPTER 6  Analysis of Environmental Pollutants using Artificial Neural (ANN) Network Algorithm

6.1 Statistical analysis of sensor array data
The experimental data obtained by CV measurements in Chapter 5, shows distinct patterns of sensor responses to certain pollutants in water. According to the literature, such data patterns can be treated by various pattern recognition algorithms including principal component analysis [1], partial least squares [2], multiple correspondence analysis [3], multiple linear regression [4], and artificial neural networks [5]. Shaffer et al. (1999), demonstrated that among the above methods, the neural network based algorithms have the highest accuracy in classifying sets of data obtained from chemical and biological sensor arrays [6].
Artificial neural networks (ANNs) also performed very well in solving overlapping signal distributions or difficult non-linear quantifications [7]. A variety of applications that deployed ANNs as a tool for multi-component analysis have been reported, particularly in chemical sensing and biosensing. Examples of such applications are: the determination of pesticides using enzyme sensors and immunosensors [8, 9], the classification of neurotoxins [10], the analysis of ethanol-glucose mixtures [11] and the quantification of metals and inorganic pollutants in groundwater [12]. These complex analytical systems are sometimes defined as an ‘electronic tongue’ due to their capability to recognise both the quantitative and qualitative composition of solutions by mimicking the concept of human sensing [13]. Previous studies [17, 18, 19, 20, 21, 22 and 23] have been established that ANN based algorithm has a high accuracy in classifying experimental data obtained from enzyme sensor arrays.

6.2 The concept of artificial neural network
Much more accurate recognition of pollutants was achieved with the use of an artificial neural network (ANN) programme written using Neural Network Toolbox, version 4.0
within MATLAB 6.1 (Mathworks, Natick, MA). The ANN model shown in Figure 6.1 is a simple general example of ANN which consists of three layers: (i), the input layer of the responses of three sensing channels containing different bacteria; (ii), the hidden layer containing a number of neurons corresponding to the number of pollutants studied (in this example it is 7); and (iii), the output layer representing a binary code (in this case a 3-digit code) which identifies the type of pollutants. The concept of artificial neural network has been inspired from examining the central nervous system of humans. The artificial neural network consists of simple computational units called neurons, which represent the processing elements in the network. Neurons are connected together using weighted links which pass signals between them. The weights of these links express the importance of its input, i.e. it is the long-term memory that the net uses to learn. Learning or training a neural network means finding the right values of weights. Each neuron accepts multi-inputs, computes a new activation level then produces a single output which could be transmitted to other neurons. The input signal to a neuron can be simple facts or outputs of other neurons while the output signal can be either an input to other neurons or a final solution to the problem.

Figure 6-1: Schematic diagram of ANN.
Commonly, ANN consists of simple processing elements or ‘neurons’ grouped together to form three types of layers known as input, hidden, and output layers (see the example in Figure 6-1). The neurons are linked with each other in a particular configuration, so that the output from neurons of one layer becomes the input to the neurons in the next layer. In each neuron, the input signal \( (x_i) \) is multiplied by a weight factor \( (w_i) \). The weighted input signals are added together and transferred to an activation function \( (F) \) that generates an output signal \( (y_{new}) \) of a neuron. The activation function can have any form, from a pure linear dependence to an elaborate exponential function, like the hyperbolic tangent (tan-sigmoid) function which was used in our case:

\[
F_{\text{tan-sigmoid}}(S) = \frac{\exp(S) - \exp(-S)}{\exp(S) + \exp(-S)},
\]

(6.1)

where \( S \) is the sum of \( p \) weighted input signals

\[
S = \sum_{i=1}^{p} w_i x_i
\]

(6.2)

ANN typically requires a large set of data and long training times. Generally, in trained networks particular input leads to a specific target output. During training, the weights of the network are iteratively adjusted to minimize the average squared error between the network outputs \( (a) \) and the target outputs \( (t) \), normally known as mean square error (MSE):

\[
\text{MSE} = \frac{1}{n} \sum_{i=1}^{n} (e_i)^2 = \frac{1}{n} \sum_{i=1}^{n} (t_i - a_i)^2,
\]

(6.3)

where \( n \) is the total number of the network inputs.

The weights of the network are optimised by several different training algorithms, such as Levenberg-Marquardt [6], resilient backpropagation [8,11], scaled conjugate gradient [9] and Bayesian regularization [7] algorithms.
6.3 The design strategy of ANN for data analysis of bacteria sensor array

The ANN used in this work for data analysis of the developed inhibition bacteria biosensor array was built using MATLAB software (version 6.1, Mathworks, Natick, MA) using the supplied functions and algorithms in MATLAB Neural Network Toolbox (version 4.0, Mathworks, Natick, MA). The strategy of the ANN design is outlined at the diagram in Figure 6-2

Figure 6-2: The strategy of ANN development for water pollutants identification
It starts with collecting the set of experimental data which contains the values of responses of all channels to all pollutants studied in all concentrations used. The second stage is the development of the ANN itself by selecting the required numbers of neurons in the input, hidden, and output layers. Then the designed ANN has to be “trained”. The training process required feeding the training data set, which is a table of all sensor responses to all pollutants in known concentrations, and the desired output, i.e. the binary code corresponding to particular pollutant and its concentration. The data are fed many times (up to $10^5$-$10^6$ times) and results in tuning link weights between the neurons. The MSE value is calculated after each iteration; and when the MSE reaches the target value the training is terminated. After that, the ANN is ready for testing.

The testing is carried out by a single feeding the testing data set, e.g. the responses of all sensors to particular pollutants in known concentrations (though different from those used in training). The results of the test show whether the ANN system works properly; if both the type of pollutant and its concentration are identified correctly, then ANN is working properly, if not, some changes have to be made in the program.

In the final step of ANN testing the pollutants of unknown concentrations have to be used either separately or in their mixture. A properly designed and trained ANN should be able to identify pollutants and to evaluate their concentrations.

6.4 The ANN design

The ANN design used in this project is shown in Figure 6-3. It consists of three neuron layers: (i), the input layer containing three neurons corresponding to the relative sensor responses of each bacteria, e.g. *E. coli*, *M. capsulatus*, and *S. oneidensis* (see Chapter 5); (ii), the hidden layer consisting of 12 neurons corresponding to 3 heavy metals (Hg$^{2+}$, Pb$^{2+}$, Cd$^{2+}$), 3 pesticides (atrazine, simazine, DDVP) and 6 petrochemicals (hexane, octane, pentane, toluene, pyrene, and ethanol); and (iii), the output layer consisting of 6 neurons representing a 6-digit binary code which combines the type of pollutant and its
The concentration is split into 5 bands, e.g. 0.1 μM, 1 μM, 10 μM, 100 μM, and 1 mM; and the ANN rounds the concentration to the nearest lower band. A hyperbolic tangent was used as the activation function for the hidden neurons. The actual ANN MatLab code is given in Appendix C.

6.5 ANN training

The training data shown in Table 6-1 were presented as Excel files consisting of three significant parts: the concentration of analytes (C) in μM, the relative responses of the three bacteria, and the 6-digit binary code corresponding to the type of pollutant and its concentration. For example, the code (000001) corresponds to Hg^{2+} in concentration of 0.1 μM, while (001101) corresponds to Cd^{2+} in concentration of 10 μM, etc.
Table 6-1: Data set for ANN training.

<table>
<thead>
<tr>
<th>C μM</th>
<th></th>
<th></th>
<th></th>
<th>E. coli</th>
<th>M. capsulatus</th>
<th>S. oneidensis</th>
<th></th>
<th>Binary code</th>
<th>target analyte</th>
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154
The training was carried out using Multilayer Perceptron Back Propagation Algorithm (MLP BPA) [15]. The neuron link weights are consequently tuned utilizing the purelin function for expanding the precision of the output. The error is ascertained as the contrast between the objective output and the system output. During the calculation the weights and biases of the system are modified to minimize the LMSE (Least Mean Square Error).

The ANN training procedure exploited the Levenberg-Marquardt algorithm to optimize the weights of neurons in a hidden layer. This algorithm appeared to be the fastest method for the network training using the limited experimental data of this study. A hyperbolic tangent was used as the activation function for the hidden neurons and a log-sigmoid function was used for the output neurons. The training was performed for 250000 epochs (e.g. 250,000 repetitions of data feeding) with the mean square error (MSE) goal set to $10^{-10}$. Figure 6-4 shows the saturation of MSE at about 250,000 epochs, which indicates the completion of the ANN training.
Figure 6-4: ANN training: Reduction of MSE during the 250,000 epochs of data feeding.

6.6 ANN testing (simulation)

After the training, the ANN programme was tested by feeding the data obtained from the bacterial sensor array for PBS solutions spiked with a particular concentration of pollutants randomly selected within the 1 μM - 100 μM concentration ranges. Both the input testing data and the outcomes of pollutants classification are shown in Table 6-2.

Table 6-2: The results of ANN identification of pollutants and estimation their concentration.

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The ANN outcome is a 6-digit code representing the type of pollutant and its concentration rounded to the nearest quantized concentration value. Despite the limited amount of data for ANN training, the programme managed to identify the pollutants correctly. The comparison of values in the last two columns representing, respectively, the obtained and actual concentrations of pollutants showed that the concentration was estimated correctly. For example, the sample spiked with 1.45 \( \mu M \) of atrazine was identified by a binary code 010001 which corresponds to atrazine in concentration of 1 \( \mu M \); the sample spiked with 0.66 \( \mu M \) of HgCl\(_2\) was identified by a code 000001 corresponding to Hg\(^{2+}\) in concentration of 1 \( \mu M \); the sample spiked with 83 \( \mu M \) of pyrene was identified by a code 110110 as pyrene in concentration of 100 \( \mu M \).
The precision of the network model was evaluated using 36 randomly selected experimental data points. This test shows that 91% of the test data was correctly classified, using the equation of True negatives / True negatives + False positives to include the calculation of that 91% precision [16]. Classified pollutants were then quantified using the optimized individual network algorithm. The results were reasonable since the network models were trained using a limited amount of data. A very good correlation between the target and the evaluated concentration of pollutants was observed with an acceptable average error of 10^{-10}. The successful classification and quantification of randomly selected data sets proves that the neural network models were able to process the information given and could be used to analyse unknown compounds or their mixtures, assuming that they belong to the group of 12 analytes used to build the models.

6.7 Summary

The use of ANN software for data processing allowed the more accurate identification of water pollutants, e.g. heavy metals, pesticides, and hydrocarbons as well as the estimation of their concentration in the range from 0.1 μM to 1 mM.

The designed ANN programme is capable of identification of pollutants as well as the rough estimation of their concentration by rounding the output to the nearest quantized concentration value, e.g. 0.1μM, 1μM, 10μM, 100μM, and 1 mM. The ANN was trained by multiple feeding the experimental results, e.g. the responses of all three sensing channels to all 12 pollutants (Hg^{2+}, Pb^{2+}, Cd^{2+}, atrazine, simazine, DDVP, hexane, pentane, pyrene, toluene, octane and ethanol) in five concentrations (0.1μM, 1μM, 10μM, 100μM, and 1 mM). Despite a rather small amount of experimental data, the trained ANN was able to classify and quantify the pollutants with acceptable errors of less than 10%.
The results obtained are very promising since simple electrochemical measurements of anodic current at +0.5V combined with ANN-based data processing allow both the identification of pollutants studied as well as rough estimation of their concentrations in the range from 0.1 μM to 1000 μM. This has emphasised the implementation of the artificial neural network algorithm as a tool for the analysis of the distinct pattern of sensor responses.

It has to be said that ANN approach in sensor array data processing is not ideal though better than the other statistical methods mentioned earlier in section 6-1. The main problem is that the ANN will only identify the analytes (pollutants in our case) which the ANN was trained on; the unknown chemicals could be either misdiagnosed, e.g. mixed up with some of the known analytes and the concentration could be wrong. That is why the analysis of “real samples” was always problematic because of a background of a number of chemicals which the ANN was not trained on.

Nevertheless, the ANN system combined with the bacteria sensor array could be successfully used for preliminary detection (screening) of water samples, in which the suspected samples containing particular pollutants in large concentrations are identified and passed to specialized laboratories for further testing.
References


CHAPTER 7 Detection of heavy metals using aptamer-based assays

As an alternative to the use of non-specific bio-receptors such as bacteria in inhibition sensor array format, highly specific receptors such as aptamers were used in current study, so this chapter describes the use of specific aptamers against Hg$^{2+}$ and Pb$^{2+}$ ions having electrochemically active labels of ferrocene and methylene blue, attached respectively, at 5' termini. The sensing mechanism of these aptamer probes is based on the changes in the DNA strand's conformation from the linear to a folded structure upon binding the metal ions, which in turn affects the electron transfer between the redox label and the metal electrode.

7.1 Experimental Methodology

7.1.1 Aptamers and other chemicals used

The following modified deoxyribonucleotides (P1 and P2) selected as specific aptamers for Hg$^{2+}$ and Pb$^{2+}$ ions, respectively, were purchased from Sangon-Biotech (China):
P1: Ferrocene-5’-TTC TTT CTT CCC CTT GTT TGT T-3’-SH [1].

P2: Methylene blue-5’-CAA CGG TTG GTG TGG TTGG-3’-SH [2].

The thiol groups (SH) at the C3 termini or prime were attached to provide strong and oriented binding of the aptamers to screen printed gold electrodes. The redox functional groups, e.g. ferrocene or methylene blue, were attached to C5 termini or prime in order to provide distinctive electrochemical properties, such as current peaks on CV characteristics associated with oxidation and reduction reactions.

The other chemicals used (all from Sigma Aldrich) were Hepes and phosphate binding buffers (HBB and PBB), also 1,4-dithiothreitol (DTT). Hepes binding buffer (HBB) was prepared by dissolving 50 mM Hepes sodium salt, 3 mM MgCl$_2$, 120 mM NaCl, and 5 mM KCl in deionized Milli-Q water. The pH of the buffer was adjusted to 7.4. Similarly, phosphate binding buffer (PBB) was prepared by dissolving 10 mM Na$_2$HPO$_4$, 1.76 mM KH$_2$PO$_4$, 3 mM MgCl$_2$, 2.7 mM KCl, and 137 mM NaCl. The pH
of the buffer was adjusted to 7.4. The addition of MgCl₂ to the buffers was essential to preserve the aptamer single DNA strand from self-coiling. For long-term storage, the 100 µM solutions of received aptamer was prepared in sterilized deionized water and stored at -20 °C in small aliquots in order to avoid repeated thaw-freeze cycles.

7.1.2 Immobilization of aptamers
The aptamers were immobilized on gold surface via thiol groups on the 3’- termini in the following procedure. Stock solution of the required aptamer was diluted to 1 µM with HBB or PBB supplemented with 1 mM of 1,4-dithiothretiol (DTT) and 3 mM of MgCl₂. DTT causes the removal of the protecting group from the SH moiety and released the aptamers with free SH end groups that could then bind to the surface of screen printed gold electrode in the presence of Mg²⁺ aptamers are initially stretched. Before immobilization, the aptamers samples were activated by rapid (1 min) heating up to 95 °C followed by 1 min cooling at (4 °C) using a conventional thermocycler polymerase chain reaction unit (TECHNE PCR version TC-3000) as appeared in Figure 7-1. Immobilization was carried out by casting aptamers solution onto the screen printed gold electrode surface; the samples were then incubated for 4 h at room temperature in a humidity chamber. The unreacted aptamers were removed from the electrode surface by several rinses with non-folding buffer (HBB), then the screen printed gold electrode with immobilized aptamers were kept in HBB to prevent aptamers from coiling.

7.2 ICP-MS measurements
A PerkinElmer NexION™ 350X ICP mass spectrometer was used as a test method for the determination of Hg²⁺ and Pb²⁺ in the real water samples. The ICP-MS instrument was equipped with a PE-AS91 auto-sampler. Samples were introduced via a cross-flow nebulizer with a Scott-type spray chamber. The NexION™ 350X and auto-sampler and peristaltic pump are controlled by the NexION™ 350X Windows NT software and are fully automated.
7.3 Electrochemical measurements

Electrochemical measurements provide the means for rapid and portable detection of heavy metal ions [3]. All CV electrochemical measurements were carried out using a DropSTAT4000P potentiostat instrument (from DropSens) controlled by Autolab software and using DropSens screen printed gold electrodes (SPGEs). These electrodes have a conventional three electrode configuration with gold working electrode (4-mm diameter disk) and counter electrode (16 mm×1.5 mm curved line), and Ag/AgCl (16 mm×1.5 mm straight line) pseudo-reference electrode and scan rate 100 mV/s [4]. CV measurements were carried out on electrodes with the immobilized aptamers, first, in pure buffer solution (HBB or PBB), then in buffer solutions with the addition of either HgCl$_2$ or PbCl$_2$ salts at different concentrations starting from 1 ng/l up to 10 µg/l. Because the addition of heavy metal salts increases the conductivity of buffer solutions, control measurements were carried out in on electrodes without aptamers at each concentration of metal salts. These measurements provide a calibration which was later used for analysis of real samples of water taken from different natural resources in the area were tested using the same screen-printed electrodes with immobilized aptamers. The reference in this case was bottled drinking water. The aptamer/metal ion binding kinetics was studied by recording current in SPGEs at fixed potentials: (-0.2 V) for HgCl$_2$ of different concentrations and (-0.4V) for PbCl$_2$ of different concentrations.

Impedance spectra were measured using an impedance spectroscope (4000A) and gold interdigitated electrodes (from Metrohm DropSens) containing 250 fringes on each side spaced by 5 µm; the overlapping length 6.76 mm. The AC voltage amplitude was 5 mV with the frequency varied from 100 mHz to100 kHz; no DC bias was applied. Similarly to CVs, the impedance spectra measurements were carried out on electrodes both coated and non-coated with aptamers, in buffer solutions containing different concentrations of metal ions.
7.4 Results and discussion

7.4.1 Design strategy of the apta-sensor

The strategy in this study (illustrated in Figure 7-2) was based on the principle that the aptamers could act as chelating factor for the analytes and undergo conformational changes that lead to changes in the electrochemical properties of aptamers containing the redox label. According to this model, in the absence of target analyte, the aptamer would remain unfolded with an extended conformation (the presence of MgCl$_2$ in the buffer is vital for that). However, upon addition of target analyte, the conformation of aptamer changes to a coiled oligonucleotide chain wrapping around the target, that would bring the probe closer to the electrode surface, resulting in increase in both cathodic and anodic currents due to the enhancement of the electron transfer between the electrode and the redox label. The increase in concentration of target analyte would increase the concentration of coiled aptamers on the surface and subsequently increase the electrochemical current. The screen printed gold electrodes activated with the anti-Hg$^{2+}$ or anti-Pb$^{2+}$ aptamers were prepared as described in the 7.1.2 section. These electrodes functionalized with Hg$^{2+}$ and Pb$^{2+}$-specific aptamers after reactivated using PCR as appeared below in Figure 7-1, were used directly as apta-sensors or stored submerged in HBB at 4 °C or room temperature for several days without any decrease in sensitivity.

![Figure 7-1: PCR activation process](image)

(A) SPGEs functionalized with Hg$^{2+}$ and Pb$^{2+}$ specific aptamers, (B) heating up to 95 °C, and (C) cooling aptamers at (4 °C).
Figure 7-2: Schematic diagram of electrochemical detection of heavy metal ions Hg$^{2+}$ (A) and Pb$^{2+}$ (B) using redox-labelled aptamers.

7.4.2 Cyclic Electrochemical measurements

Typical cyclic voltammograms (CVs) recorded on electrodes with immobilized aptamers in PBB solution with added heavy metal salts in different concentrations are shown Figure 7-2(A,B). The anti-Hg$^{2+}$ aptamer with ferrocene redox group showed well-resolved anodic and cathodic current peaks at about +0.2V and -0.2V, respectively, corresponding to oxidation and reduction of ferrocene (see Figure 7-2A). There is a clear correlation between the amplitudes of those two peaks and the concentrations of HgCl$_2$ in the buffer solution; the current goes up with the increase in Hg$^{2+}$ ions contents as was explained in the schematic diagram in Figure (7-2). Similar results were observed for anti-Pb$^{2+}$ aptamer with methylene blue redox label shown in Figure (7-2B), though the current peaks were not well-pronounce but rather appeared as shoulders at
potentials of about ± 0.2 V. Again, a correlation between the values of current and PbCl₂ salt concentration is apparent and proves the apta-sensing concept in Figure (7-2). In order to eliminate the effect of HgCl₂ and PbCl₂ salts on the conductivity of HBB solutions the control measurements were carried out on three-electrode assemblies without immobilized aptamers. These results are shown in Figure 7-4(A,B) and demonstrated no current peaks associated with redox groups, but just monotonous increase in anodic and cathodic currents above the respective potentials of about ±0.4 V upon increasing the heavy metal salts concentrations. The main point is that the current increase at the voltages of ±0.2 V was very small as compared to those on Figure (7-3).
Figure 7-3: Cyclic voltammograms of screen-printed electrodes with immobilized anti Hg$^{2+}$ (A) and anti-Pb$^{2+}$ (B) aptamers in HBB solutions with different concentrations of HgCl$_2$ and PbCl$_2$ salts. The reference curves were recorded in pure HBB without heavy metal salts added.
Figure 7-4: Cyclic voltammograms of screen-printed electrodes without immobilized aptamers in HBB solutions with different concentrations of HgCl$_2$ and PbCl$_2$ salts.

The dependence of the anodic current changes (at +0.2V) against the concentrations of heavy metals ions are shown in Figure 7-5(A,B). The background currents caused by the conductivity of HBB with added heavy metal salts was subtracted.
Figure 7-5: The concentration dependences of changes in anodic current at 0.2V upon binding of Hg$^{2+}$ (A) and Pb$^{2+}$ (B) ions to respective aptamers. Insets show the zoomed-in calibration curves with the data of five real water samples (marked as star points).
The cross-sensitivity tests were carried out by measuring CV curves on samples with immobilized anti-Hg and anti-Pb aptamers to other metal (e.g. Zn, Cu, Cd, and Pb or Hg) chloride salts. The results of such tests presented in Figure 7-6 showed no responses for non-complementary metals, which confirm high specificity of the aptamers used.

![Figure 7-6](image.png)

**Figure 7-6:** The cross-sensitivity tests: The CV responses of anti-Hg\(^{2+}\) aptamer (blue) and anti-Pb\(^{2+}\) aptamer (red) to Zn\(^{2+}\), Cu\(^{2+}\), Cd\(^{2+}\), Hg\(^{2+}\), and Pb\(^{2+}\) ions in 100 ng/ml concentrations.

Real samples of water were taken from different natural resources in the area were tested by CV measurements on screen-printed electrodes with immobilized anti-Hg\(^{2+}\) and anti-Pb aptamers. The results are lined up on the respective calibration graphs in Figure 7-5. For the sake of evaluation of Hg\(^{2+}\) and Pb\(^{2+}\) content in these samples the concentration dependences in Figure (7-6) were extrapolated to lower concentrations of Hg\(^{2+}\) and Pb\(^{2+}\) ions below the 0.01 ng/ml level. The estimated values for Hg\(^{2+}\) and Pb\(^{2+}\) contents are shown near respective data points. The obtained data for real samples of water are compared in Table 7-1 with the results of ICP-MS testing of the same water.
samples. The correlation between the two sets of data is present in terms of order of the concentration increase, however the ICP-MS values are typically from 3 to 10 times larger than those obtained with CVs. The semi-logarithmic calibration graphs of both sensors in Figure 7-6 (A,B), showed linear dependences on concentrations of respective ions. The studied concentration range was from 0.1 to 1000 ng/ml. The cross-sensitivity tests, e.g. exposures of anti-Hg$^{2+}$ aptamer to Pb$^{2+}$ and anti-Pb$^{2+}$ aptamer to Hg$^{2+}$, showed no responses which confirmed high specificity of aptamers. The data for real samples of water taken from different natural resources in the area are lined up on the above calibration graphs. For the sake of evaluation of these samples the concentration dependences were extrapolated to lower concentrations of Hg$^{2+}$ and Pb$^{2+}$ ions. The estimated values for Hg$^{2+}$ and Pb$^{2+}$ contents are shown near respective data points. The obtained data are compared in Table 7-1 with the results of ICP-MS testing of the same samples. Reasonable correlation was achieved for mercury ions, with CV data being in the same increasing order from samples 1 to 5 as ICP-MS data. Unfortunately, there is no correlation for lead ions; CV measurements showed much high values as compared to ICP-MS.

Table 7-1: ICP-MS testing results of real samples.

<table>
<thead>
<tr>
<th>Sample number</th>
<th>Hg$^{2+}$ ions (ng/ml)</th>
<th>Pb$^{2+}$ ions (ng/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CV results</td>
<td>ICP-MS result</td>
</tr>
<tr>
<td>Sample 1</td>
<td>0.00035</td>
<td>0.00170</td>
</tr>
<tr>
<td>Sample 2</td>
<td>0.00133</td>
<td>0.01062</td>
</tr>
<tr>
<td>Sample 3</td>
<td>0.01611</td>
<td>0.07831</td>
</tr>
<tr>
<td>Sample 4</td>
<td>0.08133</td>
<td>0.70311</td>
</tr>
<tr>
<td>Sample 5</td>
<td>0.71010</td>
<td>0.90013</td>
</tr>
</tbody>
</table>
7.4.3 Impedance spectroscopy measurements

The impedance spectroscopy scans were carried out on interdigitated electrodes with immobilized aptamers (as well as on bare electrodes) in HBB solutions containing heavy metal salts. Typical results are shown as Nyquist plots in Figure 7-7(A,B). As one can see, there is a major difference between these two graphs; in the presence of aptamers the Nyquist plots shift to the left (lower resistance) and reduces in radius with the increase in Hg$^{2+}$ concentration, while in the absence of aptamers the plots remain almost unchanged with a small decrease in the -$Z_{im}$ values.
Figure 7-7: The Nyquist plots (Zim vs Zre) for interdigitated electrodes with immobilized anti-Hg$^{2+}$ aptamers binding Hg$^{2+}$ ions of different concentrations (A); the Nyquist plot for bare interdigitated electrodes in solutions with different concentrations of HgCl$_2$ (B).

Similar results were obtained for PbCl$_2$ (added in the appendix-A4 Fig.3). The analysis of impedance spectra using an equivalent circuit model (shown as inset in Figure 7-7A) was carried out. According to the simplified circuit model impedance (when the diffusion impedance $Z_{\text{diff}}$ is neglected) [5], the real part of the impedance at critical points of the Nyquist plot is given as:

$$Z_{\text{re}} = R_b + R_{ct} \text{ at } \omega = 0 \text{ and } Z_{\text{re}} = R_b \text{ at } \omega = \infty,$$

where $R_b$ is the bulk resistance of the electrolyte solution, $R_{ct}$ and $C_{dl}$ connected in parallel are, respectively, the charge transfer resistance and capacitance associated with an electrical charge double layer on the surface of gold electrodes.
A decrease in impedance was observed with increasing Hg$^{2+}$ concentration, as evidenced by the decreasing height and diameter of the semi-circular Nyquist traces, which correspond to decreased capacitance and resistance of the sensor surface respectively. $R_s$ (surface resistance), $R_{dl}$ (double layer resistance) are presented in Table 7-2. Good stability of the sensor was observed as the solution resistance was partially constant at all stages of biosensor construction and testing. Whilst the $R_{dl}$ values decreased upon addition of Hg$^{2+}$, showing a clear trend. The binding of aptamers to the analyte as a biosensor surface causes a decrease in impedance.

**Table 7-2:** Values for the EIS parameters obtained from fitting the Nyquist plots shown in Figure 7-7A to the equivalent circuit model.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>$R_s$ (Ω)</th>
<th>$R_{dl}$ (Ω)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.1 ng/ml Hg$^{2+}$</td>
<td>38.026</td>
<td>932.5964</td>
</tr>
<tr>
<td>1 ng/ml Hg$^{2+}$</td>
<td>40.412</td>
<td>409.7153</td>
</tr>
<tr>
<td>10 ng/ml Hg$^{2+}$</td>
<td>42.970</td>
<td>174.7767</td>
</tr>
<tr>
<td>100 ng/ml Hg$^{2+}$</td>
<td>45.160</td>
<td>78.67283</td>
</tr>
<tr>
<td>1000 ng/ml Hg$^{2+}$</td>
<td>51.043</td>
<td>56.78813</td>
</tr>
</tbody>
</table>
Figure 7-8: Typical kinetics for anti-Hg$^{2+}$ aptamers binding Hg$^{2+}$ ions of different concentrations. Inset shows the dependence of $1/\tau$ against the concentration of Hg$^{2+}$ ions.

As one can see from Figure (7-7), the value of $R_b$ is very small (typically in single Ohms), while $R_{ct}$ is much larger (above 900 Ω at low concentrations of heavy metals) and decreases upon increasing concentration of Hg$^{2+}$ ions most-likely due to the enhancement of the electron transfer between the ferrocene label and the electrode, very much in line with the scheme shown in Figure (7-2). At the same time, the increase of Hg$^{2+}$ concentration without aptamers immobilized on the surface does not show any significant effect on the $R_{ct}$ values which corresponds well to data in Figure (7-4). Similar results were observed for interdigitated electrodes with immobilized anti-Pb$^{2+}$ aptamers in solutions containing PbCl$_2$. (see Appendix-A4 Fig.3 for Pb$^{2+}$)
The kinetics of aptamers (Hg$^{2+}$ and Pb$^{2+}$) binding

The kinetics of Hg$^{2+}$ and Pb$^{2+}$ ions binding to specific aptamers were studied by recording the time dependencies of cathodic current (at -0.2V) of three-electrode assemblies with immobilized aptamers for different concentrations of both metals ions. Cathodic current was chosen because of smaller shift of the reduction peak upon binding Hg$^{2+}$ ions. Typical time dependences for anti-Hg$^{2+}$ aptamers are shown in Figure (7-8), for different concentrations of HgCl$_2$ salt varied from 0.01 ng/ml to 1 µg/ml. Then, the characteristic time constants ($\tau$) were evaluated by fitting kinetics curves to rising exponential function. Following the Langmuir adsorption law, the reciprocal time constant (1/$\tau$) depends on the analyte adsorption and desorption rates ($k_a$ and $k_d$) and the concentration ($C$) of analytes (in our case Hg$^{2+}$ and Pb$^{2+}$) as:

$$\frac{1}{\tau} = k_a C + k_d.$$

The values of $k_a$ and $k_d$ can be found, respectively, as the gradient and intercept from the linear dependence 1/$\tau$ against C. Then both the association constant ($K_A$) and affinity constant ($K_D$) can be found as $K_A = k_a/k_d$ and $K_D = 1/K_A$ [6-8]. 1/$\tau$ vs C dependence for Hg$^{2+}$ ions was shown in Figure (7-8) as an inset. The values of $k_a$ and $k_d$ are calculated as: $k_a = 0.1471$ (sec$^{-1}$ ng$^{-1}$ ml$^{-1}$) $\times$ 271.52 (g mol$^{-1}$) $\times$ 10$^3$ (ml ng$^{-1}$) $\approx$ 3.94.10$^4$ (sec$^{-1}$ mol$^{-1}$) and $k_d = 0.0363$ (sec$^{-1}$), where 271.52 (g mol$^{-1}$) is molecular weight of HgCl$_2$, and 10$^3$ factor was used as conversion from ng/ml to µg/ml. Therefore $K_A = k_a/k_d = 1.1 \times 10^{-6}$ (mol$^{-1}$) and $K_D = 9.08 \times 10^{-7}$ (mol) for anti-Hg$^{2+}$ aptamer. Similar values, e.g. $K_A = 1.2 \times 10^{-6}$ (mol$^{-1}$) and $K_D = 8.55 \times 10^{-7}$ (mol) were found for anti-Pb$^{2+}$ aptamer. The obtained values correspond well to the aptamers affinity evaluated in the process of their synthesis [9, 10], and they are typical for highly specific binding of analytes to aptamers or antibodies. Similar analysis was carried out for binding kinetics of anti-Pb$^{2+}$ aptamer, and quite similar values of $K_A = 1.2 \times 10^{-6}$ (mol$^{-1}$) and $K_D = 8.55 \times 10^{-7}$ (mol) were found. The obtained $K_A$ and $K_D$ values for both anti-Hg$^{2+}$ and anti-Pb$^{2+}$ aptamers correspond well to the aptamers affinity evaluated in the process of their synthesis [11, 12] and they are typical for highly specific binding reactions of analytes to aptamers or antibodies.
**7.5 Discussion**

The concept of electrochemical apta-sensor for heavy metal ions was proved, and the results obtained were encouraging. The selectivity and sensitivity of this apta-sensor to the heavy metals ions, e.g. Hg$^{2+}$ and Pb$^{2+}$, is promising for development of novel, simple, and cost-effective electrochemical apta-sensors for rapid detection of heavy metals in water. A series of cyclic voltammogram and impedance spectroscopy measurements allowed the investigation of the mechanism of aptamer/heavy metal binding. The proposed model electrochemical apta-sensing based on changing the conformation of aptamer oligonucleotide chain from linear to the folded one, thus bringing the redox label closer to metal surface and increasing the electron charge transfer was proved. A simple detection of anodic (or cathodic) current at fixed voltage corresponding to oxidation (or reduction) peak potential is sufficient for detection of Hg$^{2+}$ and Pb$^{2+}$ in a wide range of concentrations down to 0.1 ng/ml (or 0.1 ppb). The detection of heavy metals in real water samples was attempted and was partially successful, however further work is required for developing methodology of real samples testing. The study of aptamer/target binding kinetics yielded the values for the association constant $K_A = 1.1 \times 10^{-6}$ (mol$^{-1}$) and the affinity constant $K_D = 9.08 \times 10^{-7}$ (mol) for aptamer/Hg$^{2+}$ binding; similar values of the association constant $K_A = 1.2 \times 10^{-6}$ (mol$^{-1}$) and the affinity constant $K_D = 8.55 \times 10^{-7}$ (mol) were found for aptamer/Pb$^{2+}$ binding. This study proved highly specific interaction between heavy metal ions and their specific aptamers. Further work could focus on development of the apta-sensor array for detection of other heavy metals (chromium, cadmium, arsenic, nickel, copper, silver, zinc, etc.) using simple DC electrochemical transducers. The different redox-labels can be used in future for simultaneous detection of different heavy metals and apply the possibility of using multiplexing aptamer arrays.
References


2. S. E. Wang, S. Si, Aptamer biosensing platform based on carbon nanotube long-range energy transfer for sensitive, selective and multicolor fluorescent heavy metal ion analysis. Analytical Methods, 5(12), (2013) 2947-2953.


CHAPTER 8 Conclusion and future work

8.1 Thesis conclusion

The optical and electrochemical characteristics of both bacteria i.e. (bacteria in solution and immobilized bacteria) and the effect of toxic chemical pollutants i.e. heavy metal salts (HgCl₂, PbCl₂, ZnCl₂ and CdCl₂), pesticides (atrazine, simazine, DDVP), and petro-chemicals (hexane, octane, pentane, toluene, pyrene and ethanol) in water on all three bacteria types (Escherichia coli, Methylococcus capsulatus (Bath) or Methylosinus trichosporium (OB3b) and Shewanella oneidensis) and a possibility of exploiting the principles of pattern recognition for identification of these mentioned pollutants is the main goal of this thesis. This work concentrated on studying the effects of the above mentioned contaminants on living bacteria. The results were shown that many types of bacteria isolates have the ability to survive at high levels of environmental contamination. The types of bacteria strain that can resist above mentioned contaminants are in general classified as gram negative, a typical example of which is S. oneidensis, known for having the highest resistance to heavy metals and pesticides pollution. On the other hand, the highly sensitive types of bacteria, such as E. coli, are sensitive to the all of these pollutants. Also in this work Methanotrophic bacteria were used to study the effects of heavy metals, pesticides and petrochemical toxicity as bacteria in solution samples and immobilized bacteria. It was shown highly resistant to hydrocarbons and extremely sensitive to heavy metals and pesticides.

Characterisation of samples were carried out using a variety of experimental techniques, i.e. optical methods including optical density measurements, fluorescent microscopy and flow cytometry for studying light scattering in bacteria samples, also complementary methods such as SEM, AFM and electrochemical methods both DC and AC.

To study the effect of these pollutants, fluorescence microscopy measurements were carried out on samples of (E. coli, Methylococcus capsulatus (Bath) or Methylosinus
trichosporium (OB3b) and S. oneidensis) bacteria stained with two organic dyes: green and red, associated respectively with living and dead bacteria. The results showed a significant difference in images of bacteria before and after exposure to the pollutants. The numbers of live E. coli bacteria decreased exponentially with the increase in each pollutant concentration. On other hand, S. oneidensis which is a Gram-negative bacteria resisted the heavy metals at low concentration, and the concentration of live bacteria increase slightly. However, they were damaged at high concentration and the bacteria concentration gradually decreased. The method of fluorescence was illustrative, allowing direct observation of live and dead bacteria. However, this method did not provide correct bacteria counts because of the limited resolution of the analysis of microscopy images. An improved analysis was based on the calculation of total intensities of green and red fluorescence.

Optical density (OD\textsubscript{600}) techniques have been used to estimate the bacteria cell’s density as a function of exposure time and concentration of pollutants. The same optical technique, namely fluorescence microscopy, was employed to study the effect of heavy metals, pesticides and petrochemical on immobilized bacteria. The effect of heavy metal salts (HgCl\textsubscript{2}, PbCl\textsubscript{2}, ZnCl\textsubscript{2}, CdCl\textsubscript{2}), pesticides (atrazine, simazine, DDVP), and petrochemicals (hexane, octane, pentane, toluene, pyrene and ethanol) appeared to be quite similar on bacteria in solution. Fluorescence microscopy seems to give the most reliable count of live bacteria concentrations. Therefore, the comparison of the inhibition effects of heavy metals, pesticides and petrochemical on (E. coli, Methylococcus capsulatus (Bath) or Methylosinus trichosporium (OB3b) and S. oneidensis) bacteria were carried out using true live and dead count of fluorescence microscopy. It showed clearly the possibility of pattern recognition of the three inhibition factors, e.g. heavy metals, pesticides and petrochemicals.

A simpler way of detecting pollutants was developed using the electrochemical properties of bacteria. The effect of heavy metals, pesticides and petrochemicals on electrochemical characteristics of microorganisms was studied.
The obtained data for DC and AC electrical study of three types of bacteria (E. coli, Methylococcus capsulatus (Bath) or Methylosinus trichosporium (OB3b) and S. oneidensis) correlated, as which did with the data of the optical study. The measurements of DC anodic and cathodic current were used for quantification of live bacteria concentrations, and thus the effect of heavy metals, pesticides and petrochemicals on bacteria. In addition, the AC characteristics, that included conductance and capacitance, were depicted as a function of pollutants exposure concentration. The AC capacitance increases when the bacteria concentration increases; in contrast AC conductance decreases. The capacitance and conductance were scanned for a wide range of frequencies; the big difference in the results of the three types of bacteria was very clearly related to the electrical properties change, which is related to the change in bacteria density or their concentration. The results at high frequency were very interesting, which encouraged us to utilize it to evaluate pollution levels.

The electrochemical technique was engaged to study the effect of heavy metals heavy metal salts (HgCl2, PbCl2, ZnCl2 and CdCl2), pesticides (atrazine, simazine, DDVP), and petro-chemicals (hexane, octane, pentane, toluene, pyrene and ethanol) on bacteria in solution and immobilized bacteria. The effect of above mentioned pollutants appeared to be comparable on (E. coli, Methylococcus capsulatus (Bath) or Methylosinus trichosporium (OB3b) and S. oneidensis) bacteria. AC and DC properties of electrochemical solutions that contained (E. coli, Methylococcus capsulatus (Bath) or Methylosinus trichosporium (OB3b) and S. oneidensis) bacteria were studied and the results were compared to immobilized bacteria and normalized to the results of samples not mixed with pollutants. Comparative results can be used to estimate pollutant concentration and the effect of each toxicant on bacteria.

Moreover, the difference in the responses of (E. coli, Methylococcus capsulatus (Bath) or Methylosinus trichosporium (OB3b) and S. oneidensis) bacteria to heavy metal salts (HgCl2, PbCl2, ZnCl2 and CdCl2), pesticides (atrazine, simazine, DDVP), and petro-chemicals
(hexane, octane, pentane, toluene, pyrene and ethanol) allows the application of the principle of pattern recognition for identification and quantification of pollutants. This work has proved the concept of a simple and cost effective electrical bacteria-based sensor and sensor array for preliminary assessment of the presence of toxins in water. This part of work has been achieved, through calculated and plot the pattern recognition of (live and dead) count for (E. coli, Methylococcus capsulatus (Bath) or Methylosinus trichosporium (OB3b) and S. oneidensis) bacteria, in addition the anodic and cathodic current subtract and AC conductance subtract, all of these calculations were done for heavy metal salts (HgCl\textsubscript{2}, PbCl\textsubscript{2}, ZnCl\textsubscript{2} and CdCl\textsubscript{2}), pesticides (atrazine, simazine, DDVP), and petro-chemicals (hexane, octane, pentane, toluene, pyrene and ethanol) effect. Meanwhile, the electrical equivalent circuit of the bacteria cell sensor was estimated. The simplest idea for this circuit consists of surface resistance in parallel with surface capacitance, both in series with block resistance. The capacitance and conductance of equivalent circuits were calculated at low frequency (\(~0\)) and at high frequency (\(\sim\infty\)). For some tests, the theoretical results showed a clear identification with practical results. This identification in results confirms the validity of results obtained, whether practical or theoretical.

Also, this work involved development of an electrochemical biosensor for highly selective, and rapid detection of Hg\textsuperscript{2+} and Pb\textsuperscript{2+} ions using DNA-based specific aptamer probes labeled with ferrocene (or methylene blue) and thiol groups at their 5’ and 3’ termini, respectively. The concept of electrochemical apta-sensor for heavy metal ions was proved, and the results obtained were encouraging. The selectivity and sensitivity of this apta-sensor to heavy metals ions, e.g. Hg\textsuperscript{2+} and Pb\textsuperscript{2+}, is high and thus promising for development of novel, simple, and cost- effective electrochemical apta-sensors for rapid detection of heavy metals in water.

Aptamers were immobilized onto the surface of screen-printed gold electrodes via the SH groups, and then cyclic voltammetry and impedance spectra measurements were performed in buffer solutions with the addition of the HgCl\textsubscript{2} and PbCl\textsubscript{2} salts at different
concentrations. Changes in 3D conformation of aptamers caused by binding their respective targets, e.g. Hg$^{2+}$ and Pb$^{2+}$ ions, are accompanied by the increase in the electron-transfer between the redox label and the electrode, and thus the presence of the above ions can be detected electrochemically. The detection of Hg$^{2+}$ and Pb$^{2+}$ ions in a wide range of concentrations down to 0.1 ng/ml (or 0.1 ppb) was achieved. The study of the kinetics of aptamer/heavy metal ions binding gave the values of the affinity constants of around $9 \times 10^{-7}$ mol, which proved high specificity of the aptamers used.

Finally, the analysis of a large amount of experimental data was carried out using artificial neural network (ANN) programme for more accurate identification of pollutants as well as the estimation of their concentration. The results are encouraging for the development of a simple and cost-effective bio-sensing technology for preliminary in-field analysis (screening) of water samples for the presence of environmental pollutants. The use of ANN software for data processing allowed the more accurate identification of water pollutants, e.g. heavy metals, pesticides, and hydrocarbons as well as the estimation of their concentration in the range from 0.1$\mu$M to 1mM.

### 8.2 Suggestion for future work

Since the current project was focused mostly on fundamental research aspects of environmental pollution detection, particularly on establishing novel detection principles using different types of bio-receptors, e.g. microorganisms (bacteria) and aptamers, further R&D work is required for development of sensor devices.

In terms of fundamental research, it will be of great interest to study in more detail the effect pollutants on bacteria cell membrane and cellular components using different methods, for instance confocal microscopy and AFM.

Further work is required in the development of bacteria sensor array devices which includes both the expansion of the array by adding new types of bacteria and widening
the range of analytes, particularly in pesticides and petro-chemicals. Detection of BETX and polycyclic aromatic hydrocarbons are of particular interest these days. Radioactive pollutants, such as depleted uranium, can also be included in the list of analytes; in that case *Deinococcus radiodurans* bacteria could be added to the sensor array.

The improvement of the ANN data processing is also required mainly for more precise evaluation of pollutants' concentration. This could be achieved using two-stage ANN processing: identification of pollutants with the general ANN, and the evaluation of pollutants' concentration using several dedicated ANN for each pollutant. This work is currently underway.

So far, the *in-vitro* detection of pollutants was carried out in this project. In future the focus should be on pollutants detection in real samples of water using the developed bacteria sensor array. Complementary analytical techniques, such as Chromatography or Inductively Coupled Plasma Mass Spectrometry (ICP-MS) can be used to verify the results obtained with bacteria-sensor array. This work is also underway.

The use of aptamers for detection of environmental pollutants has great potential. An aptamer sensor array containing several channels which specifically detect particular analytes could be a very interesting future project. An alternative (to sensor array) approach could involve a range of aptamers for different pollutants having different labels. In that way, the sensor responses could be multiplexed in one sensor chip.

Another important problem to address is the remediation of environmental pollutants, which could be done with the use of suitable genetically modified bacteria. In future, the detection and remediation of pollutants could be combined together in one piece of equipment.
Appendix-A1: Fig. 1. Fluorescence microscopy images of different bacterial samples before (a) and after (b) treated with simazine in 1 M for 2 hours.

- *E.coli* with simazine in 1M
  - a = before treatment, b = after treatment

- *M.capsulatus* with simazine in 1M
  - a = before treatment, b = after treatment

- *S.oneidensis* with simazine in 1M
  - a = before treatment, b = after treatment

Appendix-A2: Fig. 2. Fluorescence microscopy images of different bacterial samples before (a) and after (b) treated with octane in 1M for 2 hours.

- *E.coli* with octane in 1M
  - a = before treatment, b = after treatment

- *M. trichosporium* with octane in 1M
  - a = before treatment, b = after treatment

- *S.oneidensis* with octane in 1M
  - A = before treatment, B = after treatment
Appendix-A3: Figures (1-12) showed the cyclic voltammograms of different bacterial samples (bacteria suspension and immobilized bacteria after treated with different types of pollutants.

Fig.1: *E.coli* suspension with simazine in different concentrations.

Fig.2: *M. capsulatus* suspension with octane in different concentrations.
Fig. 3: *S. oneidensis* suspension with toluene in different concentration.

Fig. 4: *M. trichosporum* suspension with HgCl$_2$ in different concentration.
Fig. 5: *E. coli* suspension with hexane in different concentrations.

Fig. 6: DDVP with immobilized *E. coli* in different concentrations.
Fig. 7: immobilized *M. trichomonos* with hexane in different concentrations.

Fig. 8: immobilized *E. coli* with Cd^{2+} in different concentrations.
Fig. 9: immobilized *S. oneidensis* with Cd$^{2+}$ in different concentrations.

Fig. 10: immobilized *M. capsulatus* with Cd$^{2+}$ in different concentrations.
Fig. 11: immobilized *S. oneidensis* with Zn$^{+2}$ in different concentrations.

Fig. 12: immobilized *S. oneidensis* with Pb$^{+2}$ in different concentrations.
Appendix -A4: Figures (1,2) showed the EIS of different of immobilized bacteria after treated with different types of pollutants. While Figure 3 for aptamer biniding Pb$^{+2}$. 

Fig. 1. The Nyquist plots (-Zim vs Zre) for interdigitated electrodes with immobilized *M. capsulatus* with DDVP in different concentrations.
Fig. 2. The Nyquist plots (Zim vs Zre) for interdigitated electrodes with immobilized *S. oneidensis* with Pb^{2+} in different concentrations.

Fig. 3. The Nyquist plots (Zim vs Zre) for interdigitated electrodes with immobilized anti-Pb^{2+} aptamers binding Pb^{2+} ions of different concentrations.
Appendix - B  Analysis of impedance spectroscopy measurements of bacteria samples

The data of impedance spectroscopy can be fitted into the following equivalent circuit:

$$\frac{1}{Z_s} = \frac{1}{R_s} + \frac{1}{-j\frac{1}{\omega C_s}} = \frac{1}{R_s} - j\frac{1}{\omega C_s} = \frac{1 + j\omega R_s C_s}{R_s};$$

$$Z_s = \frac{R_s}{1 + j\omega R_s C_s} = \frac{R_s - j\omega R_s C_s}{1 + \omega^2 R_s^2 C_s^2};$$

$$Z = Z_s + R_B = \frac{R_s - j\omega R_s C_s}{1 + \omega^2 R_s^2 C_s^2} + R_B$$

after separating real and imaginary parts $Z = Z' + jZ''$

$$Z' = \frac{R_s}{1 + \omega^2 R_s^2 C_s^2} + R_B = \frac{R_s + R_B + \omega^2 R_s^2 C_s^2}{1 + \omega^2 R_s^2 C_s^2};$$

$$Z'' = -\frac{\omega R_s C_s}{1 + \omega^2 R_s^2 C_s^2}.$$

if $\omega = 0$, $Z' = R_s + R_B; Z'' = 0$.

if $\omega = \infty$, $Z' = R_B; Z'' = \frac{\omega R_s C_s}{\omega^2 R_s^2 C_s^2} = \frac{1}{\omega} = 0$.

at $Z'$ is maximal when $\frac{dZ'}{d\omega} = 0$

This gives $\omega^2 R_s^2 C_s^2 = 1$, therefore $C_s = \frac{1}{\omega_{\text{max}} R_s} = \frac{1}{2\pi f_{\text{max}} R_s}$;
Another representation of $Z$ is $Z = \sqrt{(Z')^2 + (Z'')^2}$; \[ \tan \theta = \frac{Z''}{Z'} \].

\[ Z = \frac{\sqrt{(R_S + R_B + \omega^2 R_B R_S^2 C_S^2)^2 + \omega^4 R_S^2 C_S^2}}{1 + \omega^2 R_S^2 C_S^2} \] \[ \tan \theta = -\frac{\omega R_S C_S}{R_S + R_B (1 + \omega^2 R_S^2 C_S^2)} \].

if $\omega = 0$, $Z = R_S + R_B$; \[ \tan \theta = 0 \).

if $\omega = \infty$, $Z = \sqrt{R_B + 1} \approx R_B$; \[ \tan \theta = \frac{\omega R_S C_S}{\omega^2 R_B R_S^2 C_S^2} = \frac{1}{\omega R_B} = 0. \]

$Z$ can be also presented as a series connection of $R$ an $C$: $Z = R - \frac{j}{\omega C}$.

where \[ R = \frac{R_S}{1 + \omega^2 R_S^2 C_S^2} + R_B = \frac{R_S + R_B + \omega^2 R_B R_S^2 C_S^2}{1 + \omega^2 R_S^2 C_S^2} \],

and \[ \frac{1}{\omega C} = \frac{\omega R_S C_S}{1 + \omega^2 R_S^2 C_S^2} \] \[ \omega C = \frac{1 + \omega^2 R_S^2 C_S^2}{\omega R_S C_S} \]; so that \[ C = \frac{1 + \omega^2 R_S^2 C_S^2}{\omega^2 R_S C_S} \].

if $\omega = 0$, $R = R_S + R_B$; \[ C = \infty. \]

if $\omega = \infty$, $R = R_B$; \[ C = R_S C_S. \]
Appendix C

Matlab code for ANN analysis of data of bacteria sensor array

The following code below has written and executed by using MATLAB.

```matlab
targets=[0 0 0 0 0 1
         0 0 0 0 1 0
         0 0 0 1 0 0
         0 0 0 1 0 1
         0 0 0 1 1 0
         0 0 0 1 1 1
         0 0 1 0 0 0
         0 0 1 0 0 1
         0 0 1 0 1 0
         0 0 1 0 1 1
         0 0 1 1 0 0
         0 0 1 1 0 1
         0 0 1 1 1 0
         0 0 1 1 1 1
         0 1 0 0 0 0
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         1 1 1 1 1 1
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         2 2 2 0 0 1
         2 2 2 0 1 0
         2 2 2 0 1 1
         2 2 2 1 0 0
         2 2 2 1 0 1
         2 2 2 1 1 0
         2 2 2 1 1 1
         ];

% create a feedforward network
net = feedforwardnet();

net.layers{1}.size = 60; % number of nodes in hidden layer
net.layers{2}.size = 6; % number of nodes in output layer

% initialise the network weights and biases
net = init(net);
```
net.trainFcn = 'traingdm';
% net.trainFcn = 'traingd';

% There are seven training parameters associated with traingd: epochs, show, 
goal, time, min_grad, max_fail, and lr.

net.trainParam.show = 10;       % show after specified iteration
net.trainParam.lr = 0.02;       % learning rate
% net.trainParam.mc = 0.05;       % momentum
net.trainParam.epochs = 250000; % training iteration
% net.trainParam.goal = 1e-10;    % stopping goal

net.divideParam.trainRatio = 1; % training set [%]
net.divideParam.valRatio = 0;   % validation set [%]
net.divideParam.testRatio = 0;  % test set [%]

net.layers{1}.transferFcn = 'tansig';
net.layers{2}.transferFcn = 'tansig';

% Train
net = train(net,d,targets);
view(net);

a = sim(net,d);
a=a';
a=(a)