A systematic study of enzyme-nanomaterial interactions for application in active surface decontamination

Alan Steven Campbell
West Virginia University

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A SYSTEMATIC STUDY OF ENZYME-NANOMATERIAL INTERACTIONS FOR APPLICATION IN ACTIVE SURFACE DECONTAMINATION

Alan Steven Campbell

Thesis submitted to the Benjamin M. Statler College of Engineering and Mineral Resources at West Virginia University in partial fulfillment of the requirements for the degree of

Master of Science
in
Chemical Engineering

Cerasela Zoica Dinu, Ph.D., Chair
Nianqiang Wu, Ph.D.
Charter Stinespring, Ph.D.

Morgantown, West Virginia
2013

Keywords: Enzyme-Nanomaterial Interaction, Immobilization, Carbon-based Materials, Chloroperoxidase, Decontamination
ABSTRACT
ENZYME-NANOMATERIAL INTERACTIONS: PERSPECTIVES, APPLICATIONS AND OPPORTUNITIES

Alan Steven Campbell

Enzymes are a catalytic class of proteins that possess high specificity, selectivity and biocompatibility, which makes them ideal for multiple applications in industrial production and biotechnology. However, the use of enzymes in such applications is limited due to their low operational stability and increased cost attributed to difficulty of purification and reuse. Immobilization of enzymes onto nano-sized solid supports has emerged as a potential solution to these shortcomings with a trade-off of a percentage of activity loss upon immobilization. Herein, a comprehensive study of enzyme immobilization techniques with emphasis on active-surface decontamination applications is presented.

In Chapter 1, an overview of the potential uses of enzymes and enzyme immobilization techniques is given. Benefits of enzyme-nanosupport conjugates in industrial catalysis, energy production (i.e. biofuels and biofuel cells), biosensing and bioactive coatings are discussed with emphasis on enzyme-based decontamination coatings. It is emphasized that enzyme-based conjugates are capable of increasing enzyme stability at operational conditions used in industrial production of fine chemicals, pharmaceuticals and foods. Also, this Chapter emphasizes the benefits of enzyme immobilization in regard to the development of the next generation of biosensors with enhanced selectivity and specificity or biofuel cells that do not require a membrane, and thus allow miniaturization. Additionally, coatings capable of decontaminating pathogens such as bacteria and spores can be produced through the incorporation of enzyme-nanomaterial conjugates. Finally, the chapter provides new perspectives and future directions in enzyme-based biotechnology. A. Campbell, C. Dong, C. Xiang, N. Q. Wu and C. Z. Dinu, “Enzyme-Based Technologies: Perspectives and Opportunities” Accepted to Green Polymer Chemistry: Biocatalysis and Biomaterials, ACS Symposium Series 2012.

In Chapter 2, the impacts of the reactions that take place upon enzyme immobilization at the nanointerface are discussed and the effects of multiple variables present in the immobilization process on enzyme retained activity are identified. These variables include nanosupport characteristics (i.e. physical and chemical properties, rate of curvature), enzyme properties (i.e.
surface properties, molecular weight, isoelectric point) and immobilization technique (i.e. chemical or physical binding). Prior to immobilization of the selected enzymes (i.e. soybean peroxidase (SBP), chloroperoxidase (CPO) and glucose oxidase (GOX)) all nanosupports (i.e. single-walled carbon nanotubes (SWCNTs), multi-walled carbon nanotubes (MWCNTs) and graphene oxide nanosheets (GON)) are chemically functionalized under user-controlled conditions through strong acids treatment and characterized in terms of structure and morphology. A comparison of how the characteristics of both the nanosupports used as well as immobilization technique employed affect retained activity in an enzyme specific manner is also presented. A. Campbell, C. Dong, J. Hardinger, F. Meng, G. Perhinschi, N. Q. Wu and C. Z. Dinu, “Activity and Kinetics of Immobilized Enzyme Depend on the Enzyme-Interface Reaction” To be submitted to Langmuir.

In Chapter 3, the development of a self-decontaminating enzyme-nanosupport hybrid system is presented. This system is based on the generation of the strong decontaminant hypochlorous acid (HOCl) by CPO. Two strategies are investigated. First, the production of the required substrate (i.e. hydrogen peroxide (H2O2)) by photocatalytic titanium dioxide nanobelts to be used by immobilized CPO for in situ HOCl generation is tested. Secondly, the production of H2O2 by co-immobilized GOX onto MWCNTs in the presence of glucose to be further used by immobilized CPO for in situ HOCl generation is examined. Characterization of both conjugate systems as well as their capacity for HOCl generation is presented in detail. The decontaminant production capability of the CPO-MWCNTs-GOX system shows promise for the next generation of active surface decontamination coatings. A. Campbell, C. Dong, C. Xiang, N. Q. Wu, J. S. Dordick and C. Z. Dinu, “Bionano Engineering Hybrids for the Next Generation of Self-Sustainable Decontamination Coatings” Submitted to Process Biochemistry.

This thesis also contains Appendices in which supporting information in regard to the respective chapters is detailed. Also attached are other publications in which I have been a contributing author: (1) C. Dong, A. Campbell, R. Eldawud, G. Perhinschi, Y. Rojansakul and C. Z. Dinu, “Effects of Acid Treatment on Structure, Properties and Biocompatibility of Carbon Nanotubes” Applied Surface Science 2013, 264, 261-268. (2) C. Z. Dinu, I. Borkar, S. Bale, A. Campbell, R. Kane and J. S. Dordick “Perhydrolase-nanotube-paint sporicidal composites stabilized by intramolecular crosslinking” Journal of Molecular Catalysis B: Enzymatic 2012, 75,20-2.
DEDICATION

My parents Steve Campbell and Nancy Campbell
My brother Robert Campbell
ACKNOWLEDGEMENTS

The process of completing the research and coursework necessary to obtain a Master’s degree in Chemical Engineering is a very arduous one and would not be possible without constant support. I owe a debt of gratitude to all of those around me that have made it possible for me to reach this point.

I am indebted to my advisor Prof. Cerasela Dinu for her constant advice and assistance. I would not have even been involved in research if it was not for her guidance and the extra work she has put in on my account has not gone unnoticed. I must thank her for everything that has now become possible in my future career. I also want to thank Prof. Nick Wu whose collaborative efforts have made much of my research a possibility. Without these two esteemed scientists none of this work could have occurred.

I would also like to thank the other members of both Prof. Dinu’s and Prof. Wu’s lab groups with whom I have worked closely over the years as well as the other staff and faculty of the Chemical Engineering Department at WVU. Having been at WVU for so many years they have all had an impact on my career as a student.

Finally, I have to thank my friends and family who have always supported me in everything I have done.
# TABLE OF CONTENTS

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABSTRACT</td>
<td>ii</td>
</tr>
<tr>
<td>DEDICATION</td>
<td>iv</td>
</tr>
<tr>
<td>ACKNOWLEDGEMENTS</td>
<td>v</td>
</tr>
<tr>
<td>TABLE OF CONTENTS</td>
<td>vi</td>
</tr>
<tr>
<td>LIST OF FIGURES</td>
<td>vii</td>
</tr>
<tr>
<td>LIST OF TABLES</td>
<td>x</td>
</tr>
<tr>
<td>LIST OF SYMBOLS / NOMENCLATURE</td>
<td>xi</td>
</tr>
<tr>
<td>CHAPTER 1: ENZYME-BASED TECHNOLOGIES: PERSPECTIVES AND OPPORTUNITIES</td>
<td>1</td>
</tr>
<tr>
<td>CHAPTER 2: A SYSTEMATIC STUDY OF THE ENZYME-NANOSUPPORT INTERFACE</td>
<td>21</td>
</tr>
<tr>
<td>APPENDIX 1</td>
<td>46</td>
</tr>
<tr>
<td>CHAPTER 3: BIONANO ENGINEERED HYBRIDS FOR MICROBIAL DECONTAMINANT PRODUCTION</td>
<td>52</td>
</tr>
<tr>
<td>APPENDIX 2</td>
<td>79</td>
</tr>
</tbody>
</table>
LIST OF FIGURES

CHAPTER 1: ENZYME-BASED TECHNOLOGIES: PERSPECTIVES AND OPPORTUNITIES

Figure 1: Schematic diagrams for the applications of enzymes as biological catalysts currently used for industrial-based membrane separation (Figure 1a), biological fuel cell (Figure 1b), as core components in biosensors (Figure 1c), and as active constituents of surfaces with antifouling and decontamination properties (Figure 1d).................................3

Figure 2: a) Thermal stability of free S54V perhydrolase (AcT; filled diamond), AcT crosslinked with aldehyde dextran (filled squares) and AcT crosslinked with aldehyde dextran and immobilized onto SWCNTs (filled triangles) at 75°C. b) and c) Deactivation plots following second order deactivation model. d) Sporicidal activity of cross-linked AcT-nanotube based composites: control films (spores in buffer, filled diamond), films containing cross-linked AcT-nanotube (filled circles) and control spores in PGD and H$_2$O$_2$ reaction mixture (filled squares). (Reproduced with permission from reference 23. Copyright 2012 Elsevier).......9

Figure 3: Enzymes are immobilized onto nanosupports and incorporated in composites in a layered technology. When the activity of the enzyme on the outer layer of the coating has decreased below an acceptable level, that layer can be peeled away to expose the lower layer, thereby extending the functional lifetime of the coating.................................12
CHAPTER 2: A SYSTEMATIC STUDY OF THE ENZYME-NANOSUPPORT INTERFACE

Figure 1: FTIR spectra analysis of a) pristine and acid functionalized single-walled carbon nanotubes, b) pristine and acid functionalized multi-walled carbon nanotubes and c) pristine and acid functionalized graphene oxide nanosheets. FTIR spectra confirms the presence of carboxyl functionalizations in the acid functionalized samples………………………………………………………………………………………………………………………………30

Figure 2: Conceptual image depicting the reaction mechanic of each respective enzyme as well as the immobilization procedure. Surface curvature impacts enzyme structure upon immobilization………………………………………………………………………………………………………………………………32

Figure 3: Specific retained activity comparison of SBP, CPO, and GO\textsubscript{X} immobilized onto SWCNTs, MWCNTs and GON via a) physical adsorption, b) covalent binding and c) covalent binding via PEG linker. Enzyme molecular weight as well as nanosupport diameter increase from left to right………………………………………………………………………………………………………………………………34

Figure 4: Michaelis-Menten kinetics data of a) SBP immobilized onto SWCNTs (filled squares), MWCNTs (filled circles) and GON (filled triangles) via physical adsorption; b) CPO immobilized onto MWCNTs using physical adsorption (filled squares), covalent binding (filled circles) and covalent binding through a PEG linker (filled triangles); and c) GO\textsubscript{X} immobilized onto MWCNTs using physical adsorption (filled squares), covalent binding (filled circles) and covalent binding through a PEG linker (filled triangles)………………………………………………………………………………………………………………………………37
CHAPTER 3: BIONANO ENGINEERED HYBRIDS FOR MICROBIAL DECONTAMINANT PRODUCTION

Figure 1: a) Functionalization of photocatalyst pristine titanium dioxide results in the formation of carboxyl functionalized nanobelts or TiO$_2$-NBs. b) FTIR spectrum of TiO$_2$-NBs reveals the presence of the carboxyl peak at 1731 cm$^{-1}$, confirming -COOH functionalization. c) CPO enzyme immobilization onto TiO$_2$-NBs with and without the use of a PEG linker. The CPO-TiO$_2$-NBs-based conjugates HOCl generation strategy under UV irradiation

Figure 2: a) Acids treatment of MWCNTs leads to carboxyl functionalized MWCNTs (MWCNTs-COOH). Functionalization takes place at the defect sites in the MWCNTs structures; the resulting acid treated MWCNTs have hydrophilic residues (represented by the COOH groups) and hydrophobic walls. The MWCNTs-COOH are used as nanosupports for co-immobilization of CPO and GO$_X$. First, CPO is covalently attached to COOH-functionalized MWCNTs. Subsequently, the CPO-based conjugates are used for the physical attachment of GO$_X$; GO$_X$ will attach to the hydrophobic walls of the nanotubes to result in CPO-GO$_X$-MWCNTs conjugates. In the system containing the co-immobilized enzymes and through a chain reaction, GO$_X$ provides the H$_2$O$_2$ substrate needed by CPO for \textit{in situ} conversion of Cl$^-$ into HOCl. b) Raman spectra of pristine (black curve) and carboxyl functionalized (red curve) MWCNTs. The carboxyl functionalized MWCNTs have shifted peaks towards higher relative intensities confirming the COOH functionalization

Figure 3: a) Michaelis-Menten kinetics of enzyme-based MWCNTs conjugates. a) CPO-based conjugates (physically immobilized-open circles; covalently immobilized-filled triangles) kinetics relative to free CPO in solution (filled squares). b) GO$_X$-based conjugates (physically immobilized-open circles; covalently immobilized- filled triangles) kinetics relative to free GO$_X$ in solution (filled squares)
LIST OF TABLES

CHAPTER 3: BIONANO ENGINEERED HYBRIDS FOR MICROBIAL DECONTAMINANT PRODUCTION

Table 1: Characterization of CPO-TiO₂-NBs hybrid system for in situ generation of HOCl…61
Table 2: CPO-MWCNT-GOₓ conjugates generate HOCl in situ. The rate of HOCl generation is dependent on the enzyme immobilization conditions (i.e., through physical or covalent binding).…………………………………………………………………………………………………………………………..70
LIST OF SYMBOLS / NOMENCLATURE

1. SWCNTs: single-walled carbon nanotubes
2. MWNCTs: multi-walled carbon nanotubes
3. GON: graphene oxide nanosheets
4. SBP: soybean peroxidase
5. CPO: chloroperoxidase
6. GO$_X$: glucose oxidase
7. EDC: 1-ethyl-3-[3-dimethylaminopropyl] carbodiimide
8. NHS: N-hydroxysuccinimide
9. HOCl: hypochlorous acid
10. TiO$_2$: titanium dioxide
11. TiO$_2$-NBs: titanium dioxide nanobelts
12. PEG: Amino-dPEG$_8$-COOH linker
13. AcT: perhydrolase S54V
14. PGD: propylene glycol diacetate
15. FTIR: Fourier transform infrared spectroscopy
16. SEM: scanning electron microscopy
17. AFM: atomic force microscopy
18. EDX: energy-dispersive X-ray spectroscopy
19. ABTS: 2,2’-Azinobis[3-ethylbenzothiazoline-6-sulfonic acid]
20. MCD: monochlorodimedon
21. MRSA: methicillin-resistant *Staphylococcus aureus*
22. CMATs: carbon-based materials
23. CAB: citric acid buffer
24. PBS: phosphate buffered saline
25. MES: 2-(N-morpholino)ethanesulfonic acid sodium salt
26. BCA: bicinchonic acid assay
CHAPTER 1

ENZYME-BASED TECHNOLOGIES: PERSPECTIVES AND OPPORTUNITIES

ABSTRACT
Enzymes are biological catalysts that are currently used for biocatalysis, biofuel synthesis and biological fuel cell production, for biosensors, as well as active constituents of surfaces with antifouling and decontamination properties. This review is focused on recent literature covering enzyme-based technologies with emphasis on enzymes as preferred catalysts that provide environmentally friendly, inexpensive and easy to use alternatives to existing decontamination technologies against a wide variety of pathogens, from bacteria to spores.

INTRODUCTION
Enzymes are biological catalysts with high selectivity and specificity\(^1,2\) that are employed in a wide range of applications from industrial catalysis\(^3-6\), to biofuel\(^7-10\) and biofuel cell production\(^11-13\), from biosensing\(^14-16\), to pharmaceutical and agrochemical synthesis\(^17-19\), and in surface active materials with antifouling\(^20-22\) or decontamination\(^23,24\) capabilities. Their high specificity and selectivity have enabled enzyme-based industrial processes with high yields and fewer harmful byproducts than those resulting from traditional chemical processes\(^3,4,8\). Furthermore, enzymes operate at much milder conditions of temperature, pressure and pH than conventional catalysts\(^1,2\), thereby providing substantial energy and manufacturing costs savings\(^3,25\). However, there are a number of practical problems associated with the development of enzyme-based technologies \textit{in vitro}. For instance, enzyme isolation and purification is laborious and costly\(^18\) and most of the isolated enzymes have optimum activity in water-based environments. Further, in such applications\(^26\) their increased specificity and selectivity could lead to narrow-ranged and focused catalysis, thus enzyme-based systems with short operational lifetimes\(^1,2\).

Enzyme immobilization is used as a viable alternative to overcome the limitations of enzyme-based applications \textit{in vitro} and to ensure high enzyme activity retention and high operational stability\(^2,27\). The choice of immobilization technique is determined by considering both chemical and physical properties of the enzymes and of the support surfaces. As such,
immobilization has been achieved by entrapping enzymes into polymer matrices\textsuperscript{28, 29}, Langmuir-Blodgett films\textsuperscript{30, 31}, solid-\textsuperscript{32} or liquid-\textsuperscript{33} based membranes, or simply by attachment of enzymes onto solid supports (either by covalent or physical immobilization)\textsuperscript{16, 34, 35}. This review is focused on the current trends in enzyme-based technologies and our own research aimed at developing decontamination platforms based on enzymes and capable of neutralizing bacteria, viruses and spores\textsuperscript{23, 24, 36}. Various enzyme immobilization strategies are discussed and further insights into the next generation of surface decontamination technologies are provided, outlining the studies that are underway to enable these technologies to be self-sustainable (i.e. operate under ambient conditions without external addition of the enzyme substrate).

**INDUSTRIAL CATALYSIS**

Biocatalysis\textsuperscript{25} has gained widespread use across several industries including food processing, specialty and commodity chemicals, and in pharmaceuticals production\textsuperscript{5, 17, 18}. For example, in pharmaceutical and chemical industries, enzymes are used to circumvent the often complicated steps required by chemical synthesis and separation in order to generate compounds of high purity, typically chiral, while having a much lower environmental impact\textsuperscript{3, 17, 18}. A hypothetical process is shown in Figure 1a; the image shows a nanoparticle-enzyme-based packing technology developed for large-scale industrial reacting.

The industrial use of enzymes has been influenced by the emerging technologies that allowed recombinant technology or genetic engineering\textsuperscript{3, 17, 18} to be implemented for the generation of enzymes with improved catalytic properties and selectivity\textsuperscript{25, 37}, as well as by the development of immobilization and polymer-based crosslinking techniques that allow enhanced enzyme stability\textsuperscript{1, 2, 23}. Specifically, when an enzyme is immobilized onto the surface of a chosen support it can become partially denatured, i.e., the secondary and tertiary structural features of the enzyme can be altered, thus reducing its activity\textsuperscript{38}. Furthermore, enzyme-enzyme aggregation can occur at high surface loadings, which can further reduce enzyme activity\textsuperscript{39}. Immobilization and crosslinking of enzymes onto nanoscale supports, such as carbon nanotubes, are not only capable of increasing enzyme activity and stability in extreme conditions\textsuperscript{1, 23}, but could also allow for enzyme retention and thus reusability in several reaction processes. Activities of the enzymes immobilized at the nanoscale support have been found to be influenced by the properties of the support (i.e., surface curvature, surface chemistry, etc.) as well as by the
immobilization method being used (covalent versus physical)\textsuperscript{39}. For example, when Dinu et al. immobilized perhydrolase S54V (AcT) onto single-walled carbon nanotubes (SWCNTs), the immobilization process yielded \( \sim 20\% \) of the specific activity compared to the activity of free enzyme in solution. However, when the enzyme was crosslinked using aldehyde dextran prior to immobilization onto the SWCNTs, \( \sim 40\% \) specific activity was retained\textsuperscript{23}. These advantages of using enzyme immobilization or enzyme crosslinking might reduce the high cost associated with enzyme production and use\textsuperscript{18, 27}.

**Figure 1:** Schematic diagrams for the applications of enzymes as biological catalysts currently used for industrial-based membrane separation (Figure 1a), biological fuel cell (Figure 1b), as core components in biosensors (Figure 1c), and as active constituents of surfaces with antifouling and decontamination properties (Figure 1d).
Enzymes are at the forefront of several emerging energy technologies that will help to revolutionize energy production on both the macro- and micro-scales. Energy-based applications of enzymes include: biofuel synthesis and enzyme biofuel cell production.

With the costs of fossil fuels on the rise and a greater push for more environmentally friendly energy sources, biofuels represent a valuable alternative energy source, with enzymatic processing being a critical component of the process\textsuperscript{8,40}. Generally, biofuels are produced via the biochemical conversion (e.g., hydrolysis, esterification or transesterification) of renewable biomass, either chemically or enzymatically\textsuperscript{7,10}. Biofuels such as bioethanol and biodiesel are a classification of fuels derived from biomass conversion. In the United States, bioethanol manufactured from cornstarch was widely used in recent years\textsuperscript{41}. Biodiesel is produced from a variety of sources through the transesterification of alkyl esters from feedstock and not only is more environmentally-friendly but also can be used with a higher efficiency than traditional gasoline\textsuperscript{41}. The selectivity and biocompatibility of enzymes lead to a more efficient process with fewer unwanted byproducts than traditional chemical processing\textsuperscript{8}. The large loading requirements and inherent cost of enzymes have reduced the enthusiasm for industrial scale use of enzymes for biofuel production\textsuperscript{8}. However, the economic viability of enzymatic processes can be improved through enzyme immobilization onto solid supports to allow for large-scale production\textsuperscript{27} and reusability\textsuperscript{42}.

Biological fuel cells transform the chemical energy of organic compounds, such as glucose or ethanol, into electricity by using enzymes as the catalyst\textsuperscript{11,12,43}. Figure 1b shows a schematic diagram of an enzyme-based fuel cell. The biofuel reaction is catalyzed by two different enzymes; the oxidation of the enzyme at the anode interface transfers the electrons to the cathode and onto a second enzyme to lead to electric current production. Enzyme functionality and specificity allow the construction of the fuel cells without a membrane separating the anode and cathode\textsuperscript{12,43}. Due to this feature, enzyme-based fuel cells can be easily miniaturized to allow incorporation into implantable biomedical devices such as artificial organs, micro-pumps, micro-valves, pacemakers and sensors\textsuperscript{13,43} further decreasing the risk of cytotoxicity associated with the implants\textsuperscript{13}. 
ENZYMES AS BIOSENSORS

Enzyme-based biosensors can be used for recognition and quantification of various analytes from sugar\textsuperscript{44-46} to hydrogen peroxide\textsuperscript{47}, and from superoxide anions\textsuperscript{48}, to proteins\textsuperscript{49}. Enzyme-based biosensors are formed by immobilizing enzymes onto a wide range of transducers, including electrodes\textsuperscript{50}; the immobilized enzymes create an “open-gate-based electron communication window” with the electrode surface\textsuperscript{51, 52}. The general physical and chemical properties of the materials used in the construction of biosensors, as well as the working conditions being employed, play a significant role in the performance and the detection capability of the biosensor\textsuperscript{53}. For developing the next generation of viable biosensors with increased flexibility, accuracy, specificity and optimal performance, the proper support materials and enzyme immobilization conditions need to be carefully considered. The examples included below provide a comprehensive guide into current enzyme-based biosensors used in several laboratory and industrial settings.

Glucose detection is of great importance in various fields such as the food industry, quality monitoring processes, and in clinical settings for diabetes diagnosis and therapeutic maintenance\textsuperscript{54}. Due to their high surface area-volume ratio, as well as their low toxicity and ease of fabrication, metal oxide-based and carbon-based nanomaterials are considered excellent candidates for immobilization of glucose oxidase to lead to the next generation of glucose-based biosensors (Figure 1c)\textsuperscript{55}. Zinc oxide nanotubes were recently used in biosensor fabrication that allowed linear detection of glucose in only 3 s, with a limit of detection between 50 µM to 12 mM\textsuperscript{56}; in this example the reaction is catalyzed by the glucose oxidase enzyme which transfers electrons to the support conductive material. Similarly, glucose oxidase-tetragonal pyramid-shaped zinc oxide nanostructure biosensors allowed detection in a range of 50 µM to 8.2 mM\textsuperscript{57}. In other settings, glucose oxidase was immobilized onto platinum multi-walled carbon nanotube-alumina-coated silica nanocomposites to form biosensors that displayed wide linear detection up to 10.5 mM and response time of less than 5 s\textsuperscript{58}. Lastly, bionanocomposites comprising glucose oxidase-platinum-functional graphene-chitosan complexes were used to achieve a detection limit of 0.6 µM\textsuperscript{59}. For clinical application, a multi-layer cadmium telluride quantum dot-glucose oxidase conjugate biosensor was developed to detect glucose concentrations in serum; such a biosensor allowed glucose detection with minimal pretreatment of the sample and with increased accuracy\textsuperscript{60}. 
Lactose is a metabolic byproduct regulated by the food industry\textsuperscript{61, 62}. Novel, rapid, simple and inexpensive biosensors that allow precise detection of lactose were constructed by integrating 3-mercapto propionic acid functionalized gold electrodes and beta-galactosidase-glucose oxidase-peroxidase-mediator tetrathiafulvalene combined membranes\textsuperscript{63}. Such biosensors exhibited a linear detection range of 1.5 µM to 120 µM, with a detection limit of 0.46 µM. Furthermore, such biosensors had a working lifetime of nearly 1 month.

Hydrogen peroxide is the byproduct of several biochemical oxidation processes, as well as an essential mediator in clinical, pharmaceutical and food industries as well as in the environment\textsuperscript{64}. Fast, accurate and reliable detection of hydrogen peroxide was achieved using horseradish or soybean peroxidase enzyme-based systems. For instance, horseradish peroxidase was immobilized onto gold functionalized titanium dioxide nanotubes\textsuperscript{65} or onto chitosan-based nanocomposites\textsuperscript{66} to allow the construction of biosensors with a hydrogen peroxide detection range of 5 µM to 400 µM (detection limit of 2 µM) and of 0.6 µM to 160 µM (detection limit of 0.15 µM), respectively. Similarly, soybean peroxidase-based biosensors were formed by immobilization of the enzyme onto single-walled carbon nanohorns and showed linear detection ranging from 20 µM to 1.2 mM (detection limit of 0.5 µM)\textsuperscript{67}.

Biological analytes ranging from superoxide anions to proteins have also been detected using enzyme-based biosensors. The superoxide anion is mostly regarded as toxic, leading to cellular death and mutagenesis\textsuperscript{68}. Recently, a novel, disposable superoxide anion biosensor based on the enzyme superoxide dismutase was fabricated\textsuperscript{48}. Such a biosensor was able to detect superoxide anions in a range from 0.08 µM to 0.64 µM; furthermore, this biosensor showed increased sensitivity, accuracy and long term stability. Also, a horseradish peroxidase-gold nanoparticles-carbon nanotube hybrid biosensor proved to have excellent ability to detect human IgG protein for advancing immuno-analysis assays\textsuperscript{69}.

Enzyme amperometric biosensors have also been developed and employed for the detection, monitoring and reporting of biochemical analytes related to a wide range of pathologies ranging from diabetes to trauma-associated hemorrhage\textsuperscript{53}. Implantable enzyme amperometric biosensors must recognize, transmute and generate physicochemical signals that are proportional to the chemical potential (concentration) of the analytes they are intended to be measured. Kotanen et al. have summarized the properties of such biosensors, as well as the conditions required to ensure enzyme biotransducer performance such as the stability, substrate interference, or
mediator selection. The failures associated with enzyme-based biosensors are mainly due to the degradation of the immobilized enzyme or its denaturation at the interface by unfolding which could lead to loss of biorecognition and thus loss of signal transduction\textsuperscript{51-53}.

**ENZYME-BASED BIOACTIVE COATINGS**

Enzymes can be used to provide biological function to non-biological materials, thus leading to a “bioactive” material or surface\textsuperscript{70}. In many such applications, enzymes are incorporated into paint or polymer-based coatings and subsequently applied to a desired surface\textsuperscript{22, 24, 71}. Two of the main areas in which this type of technology is being employed are in the development of antifouling surfaces\textsuperscript{20, 21} and surfaces with active decontamination capabilities\textsuperscript{23, 36}. Figure 1d illustrates the general principle of enzyme-based coatings; enzymes are immobilized onto nanosupports and upon entrapment in composite-based materials they can generate reactive species to prevent biofilm formation or to allow decontamination.

**ENZYME-BASED ANTIFOULING COATINGS**

The main aim of antifouling coatings is to prevent the attachment and growth of living organisms (referred to as a biofilm) onto a surface\textsuperscript{22}. This functionality is vital in many different applications including biomedical implants\textsuperscript{72}, biosensors\textsuperscript{73} and several types of equipment used in industrial and marine settings\textsuperscript{74, 75}. There are two major steps in biofilm formation: the initial adhesion of the fouling species, and the proliferation of that species\textsuperscript{22}. To combat adhesion or reduce adhesion strength\textsuperscript{76}, “non-sticky” coatings have been developed\textsuperscript{77}. To deter proliferation, enzyme-based coatings that generate reactive species to prevent biofilm formation have been developed\textsuperscript{22}. Such technologies offer viable alternatives to traditional antifouling coatings that rely on the use of broadly cytotoxic compounds\textsuperscript{78, 79}, and further provide safer and more environmentally friendly substitutes.

**ENZYME-BASED DECONTAMINATION COATINGS**

Enzyme-based decontamination platforms have been proposed as viable alternatives to currently available decontamination methods that use harsh chemicals and pose environmental and logistical burdens\textsuperscript{80-82}. Our groups have pioneered research into enzyme-nanomaterial-based coatings to be used as decontamination platforms that exhibit bactericidal, virucidal and
sporicidal activities23, 24, 36, 83. For instance, we have shown that upon enzyme immobilization onto carbon-based nanomaterials, including carbon nanotubes, enzyme S54V perhydrolase (AcT) stability is increased under adverse conditions such as high temperature (up to 75°C) as well as over long periods of time and room temperature storage conditions23, 38, 84 (Figure 2a,b,c). Also, the conjugates thus formed can further be incorporated into polymer or paint-based coatings without undesired leaching of the enzyme23, 71.

The decontamination capabilities of such coatings were tested against various pathogens. Peracetic acid generated by carbon nanotube-immobilized S54V perhydrolase in a latex-based coating was found to be able to decontaminate >99% of 10^6 CFU/mL B. cereus spores within 1 h (Figure 2d), 4x10^7 PFU/mL influenza virus in 15 min, and 10^6 CFU/mL E. coli in only 5 min, upon addition of the substrates propylene glycol diacetate and hydrogen peroxide23, 83. With a sustainable substrate source, such coatings can be used in the future as a passive decontamination measure to combat aerosolized anthrax. Additionally, Pangule et al. showed the antimicrobial capabilities of a lysostaphin-based coating. When such coatings were tested against 10^6 CFU/mL of methicillin-resistant Staphylococcus aureus (MRSA), >99% killing capability was achieved in only 2 h36. Borkar et al. tested the bactericidal and sporicidal capabilities of two other enzymes incorporated into paint-based coatings, namely laccase and chloroperoxidase. Hypochlorous acid produced by chloroperoxidase in the presence of hydrogen peroxide and Cl^− ions was found to be capable of killing >99% of 10^6 CFU/mL S. aureus and E. coli after 30 min. Immobilized laccase also showed bactericidal activity in the presence of several mediators with >99% killing achieved in 30 min for S. aureus and in 60 min for E. coli. The sporicidal capabilities of laccase were also demonstrated with >99% killing of 10^4 CFU/mL B. cereus and B. anthracis spores in 2 h24. All of these results show the enormous potential of enzyme-based systems for active surface decontamination in multiple situations including hospital and military scenarios23, 24, 36, 83.
Figure 2: a) Thermal stability of free S54 perhydrolase (AcT; filled diamond), AcT crosslinked with aldehyde dextran (filled squares) and AcT crosslinked with aldehyde dextran and immobilized onto SWCNTs (filled triangles) at 75°C. b) and c) Deactivation plots following second order deactivation model. d) Sporicidal activity of cross-linked AcT-nanotube based composites: control films (spores in buffer, filled diamond), films containing cross-linked AcT-nanotube (filled circles) and control spores in propylene glycol diacetate (PGD) and H₂O₂ reaction mixture (filled squares). (Reproduced with permission from reference 23. Copyright 2012 Elsevier).
CONCLUSIONS AND FUTURE DIRECTIONS

Recent advances in bioinformatics and molecular biology techniques have allowed production of enzymes with high activity, controlled specificity, and high catalytic power. Simultaneously, recent developments in immobilization of enzymes onto several nanoscale supports that have tailored properties controlled by the user, allowed the development of the next generation of enzyme-based applications as illustrated in this review. Growth in these areas will surely continue. For example, our groups continue to focus on enzyme-based decontamination strategies that will function without addition of external reagents, i.e., either the substrate or the enzyme mediator. Such enzyme-based decontamination strategies aim to be functional by simply relying on ambient conditions and will initiate *in situ* enzymatic generation of decontaminants; such systems are further defined as being self-sustainable. To achieve this goal, we are currently investigating a working strategy that allows immobilization of chloroperoxidase enzyme onto titanium dioxide nanosupports. Titanium dioxide is a widely studied photocatalyst that produces hydrogen peroxide from water when excited under UV-light. Hydrogen peroxide generated at the photocatalyst nanointerface could serve as the substrate for enzymatic *in situ* hypochlorous acid generation; hypochlorous acid is a much stronger decontaminant than H$_2$O$_2$ and thus has a broader activity range against both bacterial and sporicidal contaminants. Such strategy may be used in the development of the next generation of self-sustainable decontamination systems upon incorporation into a coating.

A major problem arising from the use of enzymes in a surface coating is enzyme deactivation over time. We envision the development of layered-based technologies that would allow user-controlled coating performance of such enzyme-based decontamination strategies (Figure 3). Specifically, in a layered system, when the activity of the enzyme on the outer layer of the coating has decreased below an acceptable level, that layer can be peeled away to expose the lower layer, thereby extending the functional lifetime of the coating. Ultimately, the potential for biotechnological application will be whether such systems can be durable and operate over a wide variety of conditions while having increased operational stability, shelf-life and being environmentally and user friendly.
Figure 3: Enzymes are immobilized onto nanosupports and incorporated in composites in a layered technology. When the activity of the enzyme on the outer layer of the coating has decreased below an acceptable level, that layer can be peeled away to expose the lower layer, thereby extending the functional lifetime of the coating.
REFERENCES


CHAPTER 2

A SYSTEMATIC STUDY OF THE ENZYME-NANOSUPPORT INTERFACE

ABSTRACT

Enzymes have potential applications in industrial catalysis, biosensing, drug delivery and decontamination, but have limited usage due to their low operational stability and yield loss in synthetic environments. Enzyme immobilization onto nano-sized solid supports has been proposed as an alternative to ensure enzyme stability retention and recovery. However, the nanosupport has often been shown to affect the enzyme activity; a deep understand of the enzyme-nanointerface reaction is thus needed if advances in the listed applications are to be made. Herein, we present a comprehensive and systematic study of the interface reactions between enzymes and nanosupports with different characteristics (i.e. surface area, charge and aspect-ratios). The interface reactions are discussed relative to the enzyme kinetics and retained activity at the nanosupports. Knowledge gained from this study can further be used to optimize enzyme-nanomaterial interactions in order to maintain optimal levels of enzyme activity while enhancing stability upon immobilization for use in a broad spectrum of applications.

INTRODUCTION

Enzymes are a naturally occurring class of proteins that possess excellent catalytic properties including activity, selectivity and specificity. They are environmentally friendly and generally produce fewer harsh byproducts than their chemical counterparts. Thus, they are very attractive to industries that generate products via chemical reaction such as in the fine chemical, pharmaceutical and agrochemical industries. In their review, Wandrey et al list the past and current states of industrial biocatalysis and elaborate on its promising future with new developments in enzyme technologies. However, before such applications can be implemented at an industrial scale there are several enzyme properties that must be improved, such as solubility, stability and inhibition. For instance, Aburto et al explain the usefulness of the enzyme chloroperoxidase in the petroleum refining industry, but also how its application in this process is limited due to low enzyme operational stability. Similarly, Garcia-Galan et al describe the potential uses of various enzymes in chemical industries ranging from
pharmaceuticals to energy or food production and focus on ways (methods) to improve their unsuitable characteristics in order to make their use a possibility.

A common approach used to enhance enzyme activity and stability for synthetic applications consists of immobilization of enzyme onto nano-sized solid supports. Immobilization was shown to enhance the stability of the enzymes while the high aspect ratio of the nanosupports allowed conjugate retention in solution and an ease of conjugate recovery via filtration. In addition to industrial advantages, interest in enzyme immobilization has also greatly increased recently due to its growing number of applications in biotechnology, such as biosensing, drug delivery, and decontamination. For example, Shi et al have shown how the immobilization of glucose oxidase improves its stability and specificity for use in glucose biosensors and Dinu et al have shown increased stability of the enzyme perhydrolase S54V at high temperatures upon immobilization for use in active surface decontamination.

However, enzyme immobilization can also have an adverse effect on its activity due to possible enzyme non-specific binding at the nanointerface and thus possible enzyme deformation of its active site upon attachment. It is thought that these effects are not only influenced by the properties of the enzyme but also by the characteristics of the nanosupports, as well as the immobilization method being used. However, to date, the preliminary studies have only looked at isolated enzymes and isolated nanosupports (e.g. Besteman et al report on the use of single-walled carbon nanotubes as a support for glucose oxidase immobilization for use in biosensing, but do not investigate any other supports or enzymes, and Luckarift et al examine the use of biomimetic silica supports to improve retained activity of butyrylcholinesterase but do not test any other enzymes or multiple immobilization methods), with no clear translation being suggested on how these studies could be transferred to other enzymes or how they could be tailored to the function of that enzyme implemented in alternative technologies. In particular, the activity of an enzyme immobilized at a nanosupport interface, and the correlation between the enzyme (i.e. surface properties, molecular weight, isoelectric point, structure) and the nanosupport characteristics (i.e. physical and chemical properties, the rate of curvature), or between the immobilization technique (i.e. physical or chemical binding) and the nanointerface reactions in water-based environments, is not well understood.

We propose to investigate the underlying complex mechanisms that influence enzyme behavior at nanointerfaces. We have focused our analyses on three model systems, i.e. enzymes
of different molecular weight and surface properties characteristics (soybean peroxidase (SBP, 40 kDa), glucose oxidase (GOX, 180 kDa), and chloroperoxidase (CPO, 42 kDa)) with unique potential for synthetic applications. The model systems have been immobilized at nanointerfaces with varying aspect ratios (i.e. SWCNTs, diameter 0.8-1.2 nm), multi-walled carbon nanotubes (MWCNTs, diameter 10-20 nm) and graphene oxide nanosheets (GON, dimensions 500-5,000 nm sheets)) using three separate methods (i.e. physical adsorption, covalent binding and covalent binding with an amino-dPEG₈-COOH linker (PEG)). The selected enzymes have interesting applications as well as similarities and differences in their properties to ensure a meaningful analysis at the nanointerface. SBP was chosen to be used as a model enzyme because of its relatively high inherent stability²⁸-³⁰. CPO was chosen due to its potential both as an oxidizer in industry and as the source of a strong decontaminant¹⁰, ²⁰. Grover et al showed the bactericidal capability of immobilized CPO against Escherichia coli when incorporated into a surface coating²⁰. GOX was chosen because of its widely studied use in biosensors to detect glucose concentrations¹⁶, ³¹-³³. Further, the effects of enzyme size on immobilization can be determined using these enzymes due to the much larger size of GOX compared to the similar smaller sizes of SBP and CPO. The characteristics of the enzyme and nanointerface were also investigated and the impact of the nanointerface on the enzyme kinetics is discussed. By determining how different factors at the enzyme-nanomaterial interface will affect the immobilized enzyme, a greater control over conjugate formation can be accomplished and thus further utilized in a wide array of applications. Further, the fundamental knowledge gained through such studies could lead to user-controlled tailoring of the biological material-nano interface for increased activity, stability and shelf life of enzyme-based synthetic applications (i.e. biosensors).

**MATERIALS AND METHODS**

**GRAPHENE OXIDE NANOSHEET SYNTHESIS**

Graphene oxide nanosheets (GON) were produced from regular graphite powder (Alfa Aesar, 99.8% purity). First, 10 g of the graphite powder and 5 g of NaNO₃ (Sigma Aldrich, 99.0%) were added to 230 mL of concentrated sulfuric acid (Fisher Scientific, 96.4%) in a 200 mL flask and placed in an ice bath with slow stirring. 30 mg of KMnO₄ (Sigma Aldrich, 99.0%) was then added slowly to ensure that the mixture temperature remained below 20°C. Next, the solution was heated to 35°C for 30 min, diluted in 460 mL of deionized (DI) water and quickly heated to
98°C for 15 min. The mixture was then rediluted in 710 mL of DI water preheated to 35°C and treated with 30 mL of 30% H₂O₂ (Sigma Aldrich). Finally, the solution was filtered and washed using DI water at 35°C until the effluent was mostly clear with a pH of 6. The resulting brown filter cake was then dried in a vacuum oven to obtain a distinctly brown powder, which was stored at room temperature until use.

**CARBON-BASED MATERIALS ACIDS FUNCTIONALIZATION**

Functionalized carbon-based materials (CMATs) (single wall carbon nanotubes - SWCNTs, 85% purity, Unidym Inc.; multi-walled carbon nanotubes- MWCNTs, 95% purity, NanoLab Inc.; or graphene oxide nanosheets- GON) were prepared via acids treatment as previously described. Briefly, 100 mg of pristine CMATs were added to a 60 mL mixture of 3:1 (V:V) sulfuric acid (Fisher Scientific, 96.4%) and nitric acid (Fisher Scientific, 69.6%). The mixture was subsequently ultrasonicated for 6 h (Branson 2510, Fisher Scientific) in ice at a constant temperature of approximately 25°C. Next, the solution was diluted with DI water and filtered through a GTTP 0.2 µm polycarbonate membrane (Fisher Scientific). Several cycles of redispersion and filtration in DI water were used to remove acidic residues or catalysts. The CMATs isolated on the filter were then dried in a vacuum desiccator and stored at room temperature until use.

**CARBON-BASED MATERIALS CHARACTERIZATION**

CMATs were characterized using scanning electron microscopy (SEM), energy-dispersive x-ray spectroscopy (EDX), and Fourier transform infrared spectroscopy (FTIR) for sample morphology and elemental composition, and to evaluate the chemical structure, respectively. For SEM and EDX characterizations, the samples (1 mg/mL in DI water) were deposited on silica wafers and dried under vacuum. Experiments were performed on a Hitachi S-4700 Field Emission Scanning Electron Microscope with a S-4700 detector combining secondary (SE) and backscattered (BSE) electron detection (in a single unit). FTIR data were collected in transmission mode using KBr pellet method on a Thermo Nicolet Instrument.

The length of pristine and functionalized SWCNTs and MWCNTs was quantified using atomic force microscopy (AFM) with a Si tip (Asylum Research, 50-90 KHz AC240TS) and tapping mode in air. Briefly, nanotube samples in DI water (0.1 mg/mL) were deposited onto
mica surfaces (9.5 mm diameter, 0.15-0.21 mm thickness, Electron Microscopy Sciences) and dried under vacuum overnight. Scan images of 10 µm x 10 µm and 1 µm x 1 µm areas were obtained.

To evaluate the degree of hydrophilicity/hydrophobicity of the CMATs, dispersity tests were performed in DI water (pH 6.25), phosphate buffered saline (PBS, 100 mM, pH 7, Sigma Aldrich) and citric acid buffer (CAB 50 mM, pH 4.8, Sigma Aldrich). Briefly, CMATs were first dispersed in each of the different solvents at a concentration of 3 mg/mL. The suspension was then centrifuged at 3000 rpm for 5 min and 0.8 mL of the resulting supernatant was removed and filtered through the GTTP 0.2 µm polycarbonate filter membrane. The filters were subsequently dried under vacuum and the amount of CMATs on the filter was weighed. Dispersity was calculated based on the volume suspended and the initial amount used in the dispersion test.

**ENZYME IMMOBILIZATION**

Soybean peroxidase (SBP, Bioresearch), glucose oxidase (GOX, Type VII, Sigma), and chloroperoxidase (CPO, Bioresearch) were immobilized onto CMATs using either physical or chemical binding. Briefly, for physical binding 2 mg of CMATs were first dispersed in 2 mL of enzyme solution (1 mg/ml in PBS for SBP, 0.5 mg/mL in PBS for GOX, or 0.5 mg/mL in CAB for CPO) via brief sonication. The solution was then incubated at room temperature for 2 h with shaking at 200 rpm. Next, the enzyme-CMAT conjugates were recovered by filtration using the GTTP 0.2 µm polycarbonate filter membrane. The supernatant was isolated and its volume recorded. The conjugates on the filter were then washed at least 6 times using the corresponding buffer (2 mL for each wash) to remove loosely bound enzyme, with the first two washes being isolated and their volumes recorded. Finally, the conjugates were redispersed in 2 mL of the corresponding buffer and stored at 4 °C.

For chemical binding, 2 mg of CMATs were first activated using 1-ethyl-3-[3-dimethylaminopropyl] carbodiimide (EDC, Acros Organics) and N-hydroxysuccinimide (NHS, Pierce). Specifically, the CMATs were dispersed in 160 mM EDC and 80 NHS in 2-(N-morpholino)ethanesulfonic acid sodium salt buffer (MES, 50 mM, pH 4.7) with a final volume of 2 mL via sonication, and incubated at room temperature for 15 min with shaking at 200 rpm. Subsequently, the mixture was filtered through the GTTP 0.2 µm polycarbonate filter membrane and washed thoroughly with MES to remove any ester residues. Next, the activated CMATs
were immediately dispersed in 2 mL of the selected enzyme solution (consistent with physical binding) and incubated at room temperature for 3 h with shaking at 200 rpm. Enzyme-CMAT conjugates were then recovered and washed, with the supernatant and the two washes recovered, as previously described (see physical binding). Finally, the conjugates were redispersed in 2 mL of the corresponding buffer and stored at 4 °C.

For chemical binding through a spacer, 2 mg of the selected CMATs were first activated with EDC and NHS chemistry as previously described (see covalent binding), and subsequently dispersed in 5 mL of 1 mg/mL Amino-dPEG₈-COOH (PEG, 32.2 Å, Quanta Biodesign) in the necessary buffer and incubated at room temperature for 3 h with shaking at 200 rpm. The resulting conjugates were then filtered and washed with the necessary buffer. Finally, the selected enzyme was attached to the PEG linker using the covalent attachment protocol as previously described. Conjugates were then redispersed in 2 mL of the required buffer and stored at 4 °C.

**ENZYME LOADING**

The amount of enzyme attached to the CMATs (defined as enzyme loading) was determined using the standard BCA Assay (Pierce) and by subtracting the amount of enzyme washed off in the supernatant and the first two washes collected after the immobilization from the initial amount added to the CMATs in the immobilization process. Briefly, 1 mL working reagent containing 50 parts reagent A with 1 part reagent B (reagents were provided stock with the BCA Assay kit) was mixed with 50 µL of enzyme solution (either from the supernatant or the washes) and incubated at 37°C for 30 min. Absorbance at 562 nm was recorded for each sample using a UV-Vis spectrophotometer (Thermo Scientific EVO300) and compared to a calibration curve of known concentrations of the respective enzyme (free in solution) in the working reagent. Loadings were estimated as the difference between the amount of enzyme washed out and the known initial amount of enzyme added during the incubation.

**ENZYME ACTIVITY ASSAY**

Enzyme retained activity was determined using colorimetric reactions monitored in a UV-Vis spectrophotometer (Thermo Scientific EVO300). Enzyme retained activity was calculated by
comparing the activity of immobilized enzyme to the activity of the free enzyme in solution at the same concentration.

Specifically, the activity of SBP was determined by monitoring the oxidation of (2,2’-Azinobis[3-ethylbenzothiazoline-6-sulfonic acid]) (ABTS, Sigma Aldrich) by SBP in the presence of hydrogen peroxide (H$_2$O$_2$, Sigma Aldrich) at 412 nm. Briefly, 20 µL of the SBP solution to be tested (free or immobilized) was added to 650 µL of 0.25 mg/mL ABTS and mixed in a plastic cuvette. Next, 20 µL of 6.5 mM H$_2$O$_2$ was added to initiate the reaction and the cuvette was immediately placed in the spectrophotometer and rate of absorbance change monitored for 2 min. The initial reaction rate was calculated from the slope of the linear time-course and reported in µM µg$^{-1}$ s$^{-1}$.

The reaction used to determine the activity of SBP was also used to determine the activity of GO$_X$. However, in this assay, the GO$_X$ produces the required H$_2$O$_2$. Briefly, 400 µL of PBS, 250 µL of 0.25 mM glucose (ACROS), 250 µL of 0.25 mg/mL ABTS, and 50 µL of 0.5 mg/mL SBP were first mixed in a plastic cuvette. Then, 50 µL of the GO$_X$ sample to be tested was added to initiate the reaction and the cuvette was immediately placed in the spectrophotometer and rate of absorbance change monitored for 2 min. The initial reaction rate was calculated from the slope of the linear time-course and reported in µM µg$^{-1}$ s$^{-1}$.

The activity of CPO was determined by monitoring the conversion of 2-chloro-5,5-dimethyl-1,3-cyclohexanedione (monochlorodimedon, Alfa Aesar) to dichlorodimedon by CPO in the presence of Cl$^-$ and H$_2$O$_2$ at 278 nm. Briefly, 500 µL of CAB, 440 µL of 227.27 mM NaCl (ACROS), 20 µL of 5 mM monochlorodimedon, and 20 µL of the CPO sample to be tested were first mixed in a quartz cuvette. Then, 20 µL of 50 mM H$_2$O$_2$ was added to initiate the reaction and the cuvette was immediately placed in the spectrophotometer and rate of absorbance change monitored for 2 min. The initial reaction rate was calculated from the slope of the linear time-course and reported in µM µg$^{-1}$ s$^{-1}$.

**ENZYME KINETIC PARAMETERS DETERMINATION**

The kinetic parameters, $K_m$ and $V_{max}$ values, of the free and immobilized enzyme (onto all CMATs) were determined by measuring the initial rates of reaction in the respective activity assays (previously described), with varying substrate concentrations. Specifically, for SBP the concentration of H$_2$O$_2$ was varied from 0 to 0.04 mM final concentration, for GO$_X$ the
concentration of glucose was varied from 0 to 100 mM final concentration and for CPO the concentration of H$_2$O$_2$ was varied from 0 to 4 mM final concentration. The K$_m$ and V$_{max}$ values for the free and immobilized enzyme were calculated using Line weaver–Burk plots using the initial rate of the enzymatic reaction$^{36,37}$:

$$\frac{1}{v} = \frac{k_m}{v_m [S]} + \frac{1}{v_m},$$

where [S] is the concentration of the substrate in mM, V and V$_{max}$ represent the initial and maximum rates of reaction, respectively reported in µM µg$^{-1}$ s$^{-1}$, and K$_m$ is the Michaelis-Menten constant reported in mM.

**STATISTICAL ANALYSIS**

All results are presented as mean ± standard deviation with at least six trials for each statistic.

**RESULTS AND DISCUSSION**

Three different carbon-based materials with different dimensions have been used in this study. These nanosupports vary in rate of surface curvature, as well as surface hydrophilicity/hydrophobicity. Both of these characteristics have been shown to affect the retained activity of immobilized enzymes$^{11,17,38}$. It was hypothesized that supports with a lower rate of curvature would have a more detrimental effect on the enzyme activity and kinetics as more of the enzyme will come into contact with the surface of the nanosupport and thus increase loss of surface area available for substrate interaction as well as enzyme denaturation. These nanosupports were chosen not only because of their varying characteristics but also for their proven benefits. MWCNTs$^{12,32,33,39,40}$, SWCNTs$^{9,11,13,39}$ and GON$^{41-43}$ have all been widely studied as supports for enzyme immobilization with promising results in multiple applications. The level of carboxyl functionality and thus hydrophilicity of these nanosupports was enhanced via acid functionalization prior to enzyme immobilization$^{34}$.

**MORPHOLOGY AND STRUCTURE CHARACTERIZATION OF CARBON-BASED MATERIALS (CMATS)**

In particular, pristine SWCNTS (diameter = 0.8-1.2 nm, length = 760 ± 276 nm), MWCNTs (diameter = 10-20 nm, length = 6,049 ± 2,954 nm) and GON (dimensions = 500 – 5,000 nm) were acid functionalized as previously described$^{12,34}$. Briefly, the CMATs were incubated in a
strong acids mixture, containing nitric and sulfuric acid for 6 h. To evaluate the effect of acids functionalization on the structure and morphology of the nanomaterials, the CMATs were examined using scanning electron microscopy (SEM), atomic force microscopy (AFM), energy dispersive X-ray analysis (EDX) and Fourier transform infrared spectroscopy (FTIR) before and after acids functionalization.

The typical morphologies of the pristine and functionalized SWCNTs and MWCNTs as investigated by Scanning Electron Microscopy (SEM) are shown in Figure S1 (Appendix 1). The SEM investigations of CMATs showed that acid treatment did not affect the overall morphology of the samples (Figure S1, Appendix 1); however, atomic force microscopy (AFM) analysis in tapping mode showed that acid treatment reduced the length of both SW- and MWCNTs. Briefly, upon acid functionalization SWCNTs were shortened from 760 ± 276 nm to 516 ± 277 nm and MWCNTs were shortened from 6,049 ± 2,954 nm to 452 ± 213 nm. Diameters of the SWCNTs and MWCNTs were unaffected by acid treatment and were 0.8-1.2 nm and 10-20 nm, respectively. The dimensions of the GON were 500-5,000 nm (sheets). EDX analysis (Figure S1, Appendix 1) showed that the oxygen content increased when compared to carbon and other elements for all the acids treated samples. Similar to previous research, this increase was due to acids treatment suggesting carboxyl functionalization onto the CMATs surface.

FTIR data further confirmed grafting of carboxyl functionalities upon acid treatment onto CMATs surface (Figure 1). In particular, the FTIR spectra analysis of acid treated SWCNTs and MWCNTs revealed the presence of intensive bands at wavenumbers around 3450, 2900, 1750-1550, and 1300-950 cm⁻¹ (Figure 1 a, b). The ~2900 and 3450 cm⁻¹ peaks were generally associated with the presence of hydroxyl moieties and/or hydroxyl moieties in carboxyl groups. The presence of the spectrum band in between 1750 to 1550 cm⁻¹ was associated with the formation of carbonyl and carbon-carbon bonds, respectively. The intensity of the bands in the range of 1300-950 cm⁻¹ are characteristic of carbon-oxygen bond formation, confirming the presence of large amounts of hydrated surface oxides and thus carboxyl functionalization. Further, the FTIR spectra of the GON is presented in Figure 1 c. Prominent peaks can be observed at wavenumbers 3400-3200 cm⁻¹, ~1740 cm⁻¹, ~1620 cm⁻¹ and 1400-1060 cm⁻¹. The large peak in the range 3400-3200 cm⁻¹ was indicative of –OH groups present at the surface of the nanosupport in the form of carboxylic acids or intercalated water. The peak at ~1740 cm⁻¹ resulted from the C=O bond in carboxylic acids groups as well as carbonyl moieties. The ~1620...
cm\(^{-1}\) peak showed the presence of C=C bonds resulting from un-oxidized regions of the graphene. Finally, the large band at 1400-1060 cm\(^{-1}\) was a product of carboxyl groups and epoxy or alkoxy groups at the surface of the material\(^{47}\). Our results were in agreement with previous studies, which showed that that liquid phase oxidation with strong acids mixtures introduces chemical property changes and adds free carboxylic acid groups to CMATs\(^{46, 47}\).
Figure 1: FTIR spectra analysis of a) pristine and acid functionalized single-walled carbon nanotubes, b) pristine and acid functionalized multi-walled carbon nanotubes and c) pristine and acid functionalized graphene oxide nanosheets. FTIR spectra confirms the presence of carboxyl functionalizations in the acid functionalized samples.

Carboxyl functionalization upon acid treatment also improved the CMATs dispersions in several water-based environments\textsuperscript{34}. Specifically, Table S1 (Appendix 1) shows that in all solutions used, i.e. DI water (pH 6.25), phosphate saline buffer (PBS, pH 7, 100 mM) and citric acid buffer (CAB, pH 4.9, 50 mM), the solubility of CMATs was improved upon the acid treatment up to a 13.5 fold increase observed in CAB for GON. For example, in DI water, the solubility of SWCNTs, MWCNTs and GON improved 9.3 fold, 6.8 fold and 6.5 fold, respectively.

**INFLUENCE OF THE IMMOBILIZATION CONDITIONS AND THE CMATS PROPERTIES ON THE ENZYME IMMOBILIZATION**

The acid functionalized CMATs were further used for enzyme immobilization. Three different enzymes, namely, soybean peroxidase (SBP), chloroperoxidase (CPO) and glucose oxidase (GOX), were immobilized onto SWCNTs and MWCNTs using three immobilization conditions: physical adsorption, covalent binding, and covalent binding through a PEG linker. SBP was further immobilized onto GON using each immobilization technique in order to further
test the impact of nanosupport characteristics. This investigation is conceptually represented in Figure 2.

**Figure 2**: Conceptual image depicting the reaction mechanic of each respective enzyme as well as the immobilization procedure. Surface curvature impacts enzyme structure upon immobilization.

Direct physical adsorption is a simple method of protein immobilization with generally high loadings, but can result in unwanted protein-protein interactions (i.e. two or more enzymes in contact resulting in decreased available surface area) and enzyme leaching over time. Covalent binding was accomplished through the use of the zero length cross-linker 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide (EDC). This slightly more complex method allows for stronger binding that will also decrease protein-protein interactions due to more directed binding. Directed covalent binding through a PEG linker serves to bring each immobilized enzyme away from other enzymes as well as the surface of the nanosupport to prevent protein deformation and increase available surface area.

After immobilization, the amount of enzyme attached to the surface, denoted as enzyme loading, and the activity of the enzyme compared to free enzyme at the same amount were determined. Loading amounts were estimated by subtracting the amount of enzyme washed out after the incubation process from the initial amount of enzyme added, and are reported as mg of enzyme per mg of nanosupport. All loading and specific retained activity data for each trial
involving physical adsorption, covalent binding and covalent binding through a PEG linker are respectively shown in supporting information Tables S2, S3 and S4 (Appendix 1).

A comparison of the specific retained activities of each enzyme immobilized onto the three different nanosupports using physical adsorption, covalent binding and covalent binding through a PEG linker are shown in Figure 3 a, b and c, respectively. Each graph shows the retained activity of SBP, CPO and GOX arranged in order of enzyme increasing molecular weight as immobilized onto SWNCTs, MWCNTs and GON. The CMATs supports are also arranged in order of their increasing diameter (decreasing surface curvature) from left to right for each respective immobilization technique. As can be seen in all three graphs, the specific retained activity varies significantly with the nanosupports being tested. Previous reports have shown that enzymes immobilized onto nanosupports with smaller diameters and thus higher rates of curvature tend to retain higher levels of activity\textsuperscript{38, 54}. Reasoning for this finding states that the higher rate of curvature of the support results in an increased center-to-center distance between two adjacent immobilized enzymes, which would reduce or totally prevent the unwanted interactions between these neighboring proteins\textsuperscript{38}. Additionally, when an enzyme is attached to the surface of a nanomaterial, and because of the increased enzyme deformation at the nanointerface, the enzyme surface area that interacts with the nanosupport increases along with increasing nanosupport diameter. This could potentially increase enzyme denaturation as well as decrease substrate interaction (Figure 2)\textsuperscript{1, 9}. These effects are also compounded by the previously mentioned protein-protein interactions\textsuperscript{38}. Specifically, Asuri et al found that not only did the suggested increased protein-protein interactions caused by a less curved surface cause lower initial activity loss upon immobilization but also caused a more dramatic activity loss over time in harsh environment\textsuperscript{54}. 
Figure 3: Specific retained activity comparison of SBP, CPO, and GO\textsubscript{X} immobilized onto SWCNTs, MWCNTs and GON via a) physical adsorption, b) covalent binding and c) covalent binding via PEG linker. Enzyme molecular weight as well as nanosupport diameter increase from left to right.

Our study found that enzymes bound to SWCNTs (0.8-1.2 nm diameter) retained lower percentages of activity compared to enzymes bound to MWCNTs (10-20 nm diameter) (Figure 3 a, b and c). For example, CPO physically bound to SWCNTs retained only around 2% of native activity whereas CPO physically bound to MWCNTs retained around 41% of native activity. This could be caused by an actual increase in protein-protein interaction between adjacent enzymes immobilized onto the same support\textsuperscript{54}. The interactions caused by an abundance of enzyme immobilized onto the same support, potentially even forming a monolayer, could outweigh the benefits of enzyme denaturation and active site loss by using a support with a smaller diameter\textsuperscript{55}. The much more dramatic decrease in rate of surface curvature observed with SBP immobilization onto GON (500-5,000 nm sheets) results in a specific retained activity also much lower than when MWCNTs were used, showing that the effect of surface curvature on enzyme denaturation and enzyme surface area lost to substrate interaction once again control the level of retained activity\textsuperscript{38}.

Our studies, which could further be used to design an optimum enzyme immobilization strategy to maximize retained activity while enhancing stability, yield an optimum nanosupport diameter (MWCNTs, 10-20 nm). For example, when examining the physical immobilization of
SBP onto the various nanosupports, MWCNTs yielded a specific retained activity of about 25%, whereas SWCNTs and GON yielded activities of about 15 and 2%, respectively. For covalent binding, this difference increased with MWCNTs maintaining about 28% and SWCNTs and GON only about 4% each.

The impact of various immobilization methods on specific retained activity was tested by attaching SBP, CPO and GOX to the CMATs using physical adsorption, covalent binding and covalent binding through a PEG linker (Figure 3 a, b and c, respectively). As can be seen in Figure 3, the most dramatic effect of immobilization method on retained activity occurs in the trials of CPO and GOX bound to MWCNTs. Specifically, CPO bound to MWCNTs physically, covalently and covalently with a PEG linker yielded retained activities of around 41, 53 and 30%, respectively. GOX bound to MWCNTs physically, covalently and covalently with a PEG linker resulted in retained activities of around 10, 38 and 63%, respectively. Comparing physical to covalent immobilization, the retained activities of the CPO samples increased 12% whereas the GOX samples increased 28%. This is attributed to non-specific binding of the enzyme at the hydrophobic walls of the MWCNTs. Both enzymes have an isoelectric point of around 456, 57, but the working pH of the GOX (PBS pH = 7) trials is higher than that of CPO (CAB pH = 4.8) meaning that the GOX is more negatively charged and thus has a higher affinity for hydrophobic surfaces. Therefore, more non-directed binding occurs in the physical immobilization trials of GOX, which leads to extensive protein-protein and protein-nanosupport interactions9, 38.

The covalent binding procedure helps to direct binding to the carboxyl functionalizations present on the surface of the CMATs50-52, 55. As the CPO has a lower affinity for hydrophobic surfaces compared to that of GOX, the transition to covalent binding from physical binding has less of an improvement on retained activity, although there is an improvement due to the more ordered binding mechanic. Further, covalent binding with the PEG linker is meant to bring the enzyme away from the surface of the nanosupport and thus more extensively reduce non-specific enzyme interaction9, 53. This effect is seen in the 25% increase in activity seen between GOX covalently bound and covalently bound with the linker. This significant increase further confirms the hindering of the enzyme at the nanosupports interface due to protein interaction as mentioned when comparing nanosupports physical and covalent binding. The increased size of GOX relative to SBP and CPO could also add to the extent of this effect. Alternately, CPO exhibited a 13% decrease in retained activity when covalent binding is compared to covalent binding with the
PEG linker. It is possible that the size and structure of the CPO allows for interaction of the PEG spacer arm with the active site of the enzyme, thus hindering activity, as well as the increased mobility provided by the spacer arm causes agglomeration away from the nanointerface. These results show that variables such as immobilization technique affect enzyme-nanomaterial conjugates in an enzyme specific manner, which is in agreement with previous studies\textsuperscript{8,11}.

**KINETICS OF THE ENZYME-CMATS CONJUGATES**

Kinetic parameters were evaluated for all enzyme-nanomaterial conjugate systems. Namely, $V_{\text{max}}$ and $K_m$ values were calculated using Lineweaver-Burk plots\textsuperscript{37}, which allowed for the determination of $K_{\text{cat}}$ values. All kinetic parameters for the enzymatic systems, along with the kinetic parameters of free enzyme for comparison, are included in supporting information Tables S5, S6 and S7, respectively (Appendix 1). These kinetics data support the retained activity data previously discussed. For example, in the trials with SBP physically bound to various nanosupports, the trends follow those of the specific retained activity. Specifically, SBP physically adsorbed onto SWCNTs, MWCNTs and GON yielded $V_{\text{max}}$ values of 0.005, 0.017 and 0.003, respectively (Figure 4a). Similar to the immobilization onto various nanosupport comparisons, immobilization onto MWCNTs yielded a decrease of about 87% of native SBP whereas samples immobilized onto SWCNTs and GON resulted in $V_{\text{max}}$ decreases of 96 and 98% when compared to the native enzyme, respectively. However, the $K_m$ values were on the same order of magnitude for all samples, indicating that there was no significant conformational change upon immobilization.
Figure 4: Michaelis-Menten kinetics data of a) SBP immobilized onto SWCNTs (filled square), MWCNTs (filled circle) and GON (filled triangle) via physical adsorption; b) CPO immobilized onto MWCNTs using physical adsorption (filled square), covalent binding (filled circle) and covalent binding through a PEG linker (filled triangle; and c) GOX immobilized onto MWCNTs using physical adsorption (filled square), covalent binding (filled circle) and covalent binding through a PEG linker (filled triangle).

Additionally, when comparing various immobilization methods, the kinetics reinforce specific retained activity observations. CPO bound to MWCNTs physically, covalently and covalently through a PEG linker yielded $V_{\text{max}}$ values of around 12, 13 and 8, respectively (Figure 4b). This represents decreases of about 55, 53 and 71% from native enzyme, respectively, which show the effect of more directed binding discussed previously.
The more dramatic effect of binding technique used was observed with the kinetics of the GO\textsubscript{X} samples. Specifically, GO\textsubscript{X} bound to MWCNTs physically, covalently and covalently through a PEG linker yielded $V_{\text{max}}$ values of around 0.029, 0.196 and 0.234, respectively (Figure 4c). These values correspond to decreases of around 94, 60 and 52% from native enzyme, respectively. Km values for both CPO and GO\textsubscript{X} in these trials were on the same order of magnitude, respectively, confirming that no significant conformational change occurred upon immobilization.

**CONCLUSIONS**

We have studied the behavior of multiple enzymes immobilized onto nanosupports of varying characteristics using three immobilization techniques. The results show that the retained activity and kinetics of the immobilized enzyme are influenced by the properties of the support and the conditions of the immobilization being used, in an enzyme specific manner. Such an understanding of how to optimize the amount of enzyme activity and kinetic behavior retained upon immobilization is crucial for enzyme based applications such as biosensors and biofuel cells.
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**Figure S1:** EDX data and SEM image of a) pristine SWCNTs, b) acid functionalized SWCNTs, c) pristine MWCNTs, d) acid functionalized MWCNTs; and EDX data of e) pristine GON and f) acid functionalized GON. SEM images of GON could not be obtained due to sample charging.

**Table S1:** Nanosupport Solubility

<table>
<thead>
<tr>
<th>Nanosupport</th>
<th>Solubility (mg/mL)</th>
<th>CAB (50 mM, pH 4.8)</th>
<th>DI Water (pH 6.25)</th>
<th>PBS (100 mM, pH 7)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pristine SWCNTs</td>
<td>0.06</td>
<td>0.06</td>
<td>0.13</td>
<td></td>
</tr>
<tr>
<td>Acid Treated SWCNTs</td>
<td>0.50</td>
<td>0.56</td>
<td>0.63</td>
<td></td>
</tr>
<tr>
<td>Pristine MWCNTs</td>
<td>0.06</td>
<td>0.13</td>
<td>0.13</td>
<td></td>
</tr>
<tr>
<td>Acid Treated MWCNTs</td>
<td>0.56</td>
<td>0.88</td>
<td>0.50</td>
<td></td>
</tr>
<tr>
<td>Pristine GON</td>
<td>0.13</td>
<td>0.31</td>
<td>0.81</td>
<td></td>
</tr>
<tr>
<td>Acid Treated GON</td>
<td>1.75</td>
<td>2.01</td>
<td>1.13</td>
<td></td>
</tr>
</tbody>
</table>

**Table S2:** Physical Adsorption Loading and Activity Data

<table>
<thead>
<tr>
<th>Enzyme-Nanosupport</th>
<th>Loading (mg enzyme / mg nanosupport)</th>
<th>Specific Retained Activity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SBP-SWCNTs</td>
<td>0.19 ± 0.03</td>
<td>14.81 ± 6.77</td>
</tr>
<tr>
<td>CPO-SWCNTs</td>
<td>0.09 ± 0.02</td>
<td>1.49 ± 0.16</td>
</tr>
<tr>
<td>GOX-SWCNTs</td>
<td>0.17 ± 0.03</td>
<td>1.99 ± 0.45</td>
</tr>
<tr>
<td>SBP-MWCNTs</td>
<td>0.15 ± 0.05</td>
<td>25.28 ± 4.04</td>
</tr>
<tr>
<td>CPO-MWCNTs</td>
<td>0.10 ± 0.02</td>
<td>41.36 ± 6.65</td>
</tr>
<tr>
<td>GOX-MWCNTs</td>
<td>0.24 ± 0.02</td>
<td>9.89 ± 3.30</td>
</tr>
<tr>
<td>SBP-GON</td>
<td>0.27 ± 0.06</td>
<td>1.71 ± 0.77</td>
</tr>
</tbody>
</table>
**Table S3:** Covalent Binding Loading and Activity Data

<table>
<thead>
<tr>
<th>Enzyme-Nanosupport</th>
<th>Loading (mg enzyme / mg nanosupport)</th>
<th>Specific Retained Activity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SBP-SWCNTs</td>
<td>0.08 ± 0.02</td>
<td>4.38 ± 1.49</td>
</tr>
<tr>
<td>CPO-SWCNTs</td>
<td>0.06 ± 0.01</td>
<td>2.06 ± 0.35</td>
</tr>
<tr>
<td>GO(_X)-SWCNTs</td>
<td>0.25 ± 0.03</td>
<td>14.04 ± 1.69</td>
</tr>
<tr>
<td>SBP-MWCNTs</td>
<td>0.24 ± 0.10</td>
<td>28.01 ± 5.01</td>
</tr>
<tr>
<td>CPO-MWCNTs</td>
<td>0.07 ± 0.02</td>
<td>52.70 ± 6.26</td>
</tr>
<tr>
<td>GO(_X)-MWCNTs</td>
<td>0.25 ± 0.02</td>
<td>37.67 ± 4.70</td>
</tr>
<tr>
<td>SBP-GON</td>
<td>0.16 ± 0.04</td>
<td>4.03 ± 1.98</td>
</tr>
</tbody>
</table>

**Table S4:** Covalent Binding through PEG Linker Loading and Activity Data

<table>
<thead>
<tr>
<th>Enzyme-Nanosupport</th>
<th>Loading (mg enzyme / mg nanosupport)</th>
<th>Specific Retained Activity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SBP-SWCNTs</td>
<td>0.23 ± 0.05</td>
<td>7.99 ± 3.73</td>
</tr>
<tr>
<td>CPO-SWCNTs</td>
<td>0.04 ± 0.02</td>
<td>3.46 ± 1.81</td>
</tr>
<tr>
<td>GO(_X)-SWCNTs</td>
<td>0.06 ± 0.02</td>
<td>20.32 ± 8.08</td>
</tr>
<tr>
<td>SBP-MWCNTs</td>
<td>0.10 ± 0.03</td>
<td>19.70 ± 3.82</td>
</tr>
<tr>
<td>CPO-MWCNTs</td>
<td>0.18 ± 0.05</td>
<td>30.26 ± 6.59</td>
</tr>
<tr>
<td>GO(_X)-MWCNTs</td>
<td>0.04 ± 0.01</td>
<td>63.06 ± 3.37</td>
</tr>
<tr>
<td>SBP-GON</td>
<td>0.16 ± 0.02</td>
<td>1.59 ± 0.60</td>
</tr>
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</table>
**Table S5:** Soybean Peroxidase Michaelis-Menten Kinetics

<table>
<thead>
<tr>
<th>Nanosupport</th>
<th>Maximum Production Rate (µM µg(^{-1}) s(^{-1}))</th>
<th>(K_m) (µM H(_2)O(_2))</th>
<th>(K_{cat}) (s(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Free SBP</td>
<td>0.128 ± 0.042</td>
<td>1.9 ± 0.8</td>
<td>3.53 ± 1.64</td>
</tr>
<tr>
<td>MWCNTs (Physical)</td>
<td>0.017 ± 0.007</td>
<td>7.2 ± 2.3</td>
<td>0.47 ± 0.27</td>
</tr>
<tr>
<td>MWCNTs (Covalent)</td>
<td>0.011 ± 0.004</td>
<td>1.6 ± 0.4</td>
<td>0.30 ± 0.16</td>
</tr>
<tr>
<td>MWCNTs (Covalent with PEG)</td>
<td>0.008 ± 0.003</td>
<td>2.9 ± 0.4</td>
<td>0.22 ± 0.12</td>
</tr>
<tr>
<td>SWCNT (Physical)</td>
<td>0.005 ± 0.001</td>
<td>3.7 ± 1.0</td>
<td>0.14 ± 0.04</td>
</tr>
<tr>
<td>SWCNTs (Covalent)</td>
<td>0.012 ± 0.003</td>
<td>3.7 ± 1.1</td>
<td>0.33 ± 0.12</td>
</tr>
<tr>
<td>SWCNTs (Covalent with PEG)</td>
<td>0.022 ± 0.011</td>
<td>3.4 ± 1.7</td>
<td>0.61 ± 0.43</td>
</tr>
<tr>
<td>GON (Physical)</td>
<td>0.003 ± 0.001</td>
<td>1.4 ± 1.0</td>
<td>0.08 ± 0.04</td>
</tr>
<tr>
<td>GON (Covalent)</td>
<td>0.005 ± 0.001</td>
<td>2.6 ± 0.3</td>
<td>0.14 ± 0.04</td>
</tr>
<tr>
<td>GON (Covalent with PEG)</td>
<td>0.002 ± 0.001</td>
<td>2.8 ± 2.7</td>
<td>0.03 ± 0.04</td>
</tr>
</tbody>
</table>

**Table S6:** Chloroperoxidase Michaelis-Menten Kinetics

<table>
<thead>
<tr>
<th>Nanosupport</th>
<th>Maximum Production Rate (µM µg(^{-1}) s(^{-1}))</th>
<th>(K_m) (µM H(_2)O(_2))</th>
<th>(K_{cat}) (s(^{-1}))</th>
</tr>
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<tbody>
<tr>
<td>Free CPO</td>
<td>27.14 ± 2.60</td>
<td>480 ± 70</td>
<td>1,139.88 ± 109.20</td>
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<tr>
<td>MWCNTs (Physical)</td>
<td>12.28 ± 2.56</td>
<td>340 ± 70</td>
<td>515.76 ± 107.52</td>
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<tr>
<td>MWCNTs (Covalent)</td>
<td>12.79 ± 1.97</td>
<td>530 ± 60</td>
<td>537.18 ± 82.74</td>
</tr>
<tr>
<td>MWCNTs (Covalent with PEG)</td>
<td>7.90 ± 3.16</td>
<td>550 ± 130</td>
<td>331.8 ± 187.69</td>
</tr>
<tr>
<td>SWCNT (Physical)</td>
<td>0.42 ± 0.12</td>
<td>240 ± 40</td>
<td>17.64 ± 7.13</td>
</tr>
<tr>
<td>SWCNTs (Covalent)</td>
<td>0.56 ± 0.10</td>
<td>140 ± 30</td>
<td>23.52 ± 5.94</td>
</tr>
<tr>
<td>SWCNTs (Covalent with PEG)</td>
<td>0.94 ± 0.49</td>
<td>130 ± 20</td>
<td>39.48 ± 19.10</td>
</tr>
<tr>
<td>Nanosupport</td>
<td>Maximum Production Rate (µM µg⁻¹ s⁻¹)</td>
<td>Kₘ (µM glucose)</td>
<td>Kₜₐₜ (s⁻¹)</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>---------------------------------------</td>
<td>-----------------</td>
<td>------------</td>
</tr>
<tr>
<td>Free GOx</td>
<td>0.488 ± 0.040</td>
<td>2,600 ± 500</td>
<td>78.08 ± 6.40</td>
</tr>
<tr>
<td>MWCNTs (Physical)</td>
<td>0.029 ± 0.003</td>
<td>1,800 ± 800</td>
<td>4.64 ± 0.48</td>
</tr>
<tr>
<td>MWCNTs (Covalent)</td>
<td>0.196 ± 0.032</td>
<td>3,200 ± 700</td>
<td>31.36 ± 5.12</td>
</tr>
<tr>
<td>MWCNTs (Covalent with PEG)</td>
<td>0.234 ± 0.032</td>
<td>2,600 ± 700</td>
<td>42.12 ± 8.15</td>
</tr>
<tr>
<td>SWCNT (Physical)</td>
<td>0.043 ± 0.010</td>
<td>2,300 ± 600</td>
<td>7.74 ± 2.55</td>
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<tr>
<td>SWCNTs (Covalent)</td>
<td>0.067 ± 0.005</td>
<td>1,000 ± 100</td>
<td>12.06 ± 1.27</td>
</tr>
<tr>
<td>SWCNTs (Covalent with PEG)</td>
<td>0.099 ± 0.039</td>
<td>2,300 ± 500</td>
<td>17.82 ± 9.93</td>
</tr>
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</table>
CHAPTER 3

BIONANO ENGINEERED HYBRIDS FOR MICROBIAL DECONTAMINANT PRODUCTION

ABSTRACT

Current methods for microbial decontamination are corrosive and/or toxic and can cause collateral damage to goods and people. Herein a non-toxic, non-corrosive and easily deployable enzyme-nanosupport hybrid system was developed for in situ generation of hypochlorous acid (HOCl), a strong decontaminant. Chloroperoxidase working enzyme was immobilized onto two different nanosupports (i.e., a photocatalyst and a carbon-based nanomaterial) and the enzyme loading and activity at the nanosupport interfaces, as well as its potential to generate HOCl was evaluated. The enzyme-photocatalyst hybrid system showed negligible capability of HOCl generation under UV light irradiation. In contrast, the enzyme-carbon-based hybrid system exhibited strong capability of HOCl generation, which further has the potential to be employed for decontamination of bacteria and spores.

INTRODUCTION

With the high demand for disinfection of contaminated surfaces in hospital settings\(^1\text{--}^6\) and for removal of chemical or biological warfare agents in military scenarios\(^7\text{--}^{10}\), it is of great importance to develop coatings that are fast in decontaminating, readily deployable, and user-and environmentally benign. Conventionally used microbial decontamination agents such as glutaraldehyde, alcohols, ammonium compounds or halides (i.e. chlorine or iodine) are required in high concentrations and can be harmful to both the environment and humans\(^11\text{--}^{14}\). Further, they cannot be incorporated into coatings and are usually deployed after contamination has occurred\(^7\text{,}^8\). Thus, there is an increased interest in developing microbial decontamination coatings that can decontaminate during contamination or immediately upon contamination, while protecting personnel and infrastructure and allowing for self-cleaning\(^15\text{,}^16\). Ideally, such coatings should also allow decontamination of large areas as well as decontamination of sensitive substrates\(^17\).

Current methods to produce coatings with decontamination capabilities rely mainly on encapsulating oxide-based photocatalysts such as titanium dioxide\(^18\text{--}^{20}\), zinc oxide\(^21\text{,}^22\), or silver
nanoparticles\textsuperscript{23, 24} into composites to produce reactive oxygen species (i.e. hydrogen peroxide (H\textsubscript{2}O\textsubscript{2}), superoxide radical, etc.). However, such coatings have shown limitations with regard to the photocatalyst leaching out\textsuperscript{25}, or the H\textsubscript{2}O\textsubscript{2} decontaminant not being strong enough to allow decontamination of more resilient pathogens such as spores\textsuperscript{23, 26-28}. Recently, our group has shown that enzyme-carbon nanomaterial-based conjugates entrapped into polymer or paint-based coatings can generate potent decontamination agents such as hypochlorous or peracetic acid\textsuperscript{29, 30}. Using such coatings, decontamination was achieved upon addition of specific enzyme substrates, e.g. for perhydrolase S54V (AcT)-carbon nanotubes conjugates encapsulated into latex-based paints, peracetic acid was produced in the presence of propylene glycol diacetate and H\textsubscript{2}O\textsubscript{2} as substrates, yielding the efficient decontamination of both \textit{E. coli} and \textit{B. cereus}\textsuperscript{31}. Similarly, laccase- or chloroperoxidase-carbon nanotubes conjugates encapsulated into paints allowed decontamination of \textit{E. coli} and \textit{S. aureus}, with laccase showing further decontamination capability against \textit{B. cereus} and \textit{B. anthracis}, after the addition of the respective substrates for each enzyme (i.e., H\textsubscript{2}O\textsubscript{2} for chloroperoxidase and I\textsuperscript{-} anions for laccase)\textsuperscript{30}. In these studies the carbon nanotubes enhanced enzyme stability, while their high surface area to volume aspect ratios allowed for high enzyme loadings and thus retention of the enzyme-nanosupport conjugates into the coatings\textsuperscript{31-35}.

We now aim to advance towards creating a self-decontaminating enzyme-nanosupport hybrid system to be incorporated into coatings and allow \textit{in situ} generation of hypochlorous acid (HOCl). HOCl has a wide spectrum of activity against a variety of pathogens such as spores and bacteria\textsuperscript{30, 31, 36}. Specifically, we are employing two different strategies to allow the working enzyme chloroperoxidase (CPO) to use H\textsubscript{2}O\textsubscript{2} produced at nanosupport interface to convert (Cl\textsuperscript{-}) into HOCl. The H\textsubscript{2}O\textsubscript{2} is produced either by a titanium dioxide (TiO\textsubscript{2}-NBs) or by a multi-walled carbon nanotube (MWCNT-COOH) modified nanosupport through a photocatalytic or an enzymatic chain reaction, respectively. We characterized the biocatalysts at the nanosupport interface and compared the two strategies in terms of HOCl generation rate, as well as the retained CPO activity upon immobilization. Such strategies could provide viable means for the next generation of self-sustainable coatings with enhanced stability and activity to be used for surface-decontamination of model bacteria or spores.
EXPERIMENTAL
SYNTHESIS OF TITANIUM DIOXIDE NANOBELTS (TiO$_2$-NBS)

Pristine anatase titanium dioxide nanosupports were synthesized by hydrothermal processing. Specifically, 1.2 g of anatase titanium dioxide nanoparticles (Alfa Aesar, USA) were added to 85 mL of 10 M sodium hydroxide (Sigma, USA) aqueous solution. The mixture was vigorously stirred for 1 h at room temperature and transferred to a 100 mL Teflon-lined stainless steel autoclave and heated at 190°C for 24 h. The resulting white, fluffy powder was collected and washed with copious amounts of DI water and 0.1 M hydrochloric acid (Fisher Scientific, USA) until the pH of the washing solution was less than 7. The powder was further dried overnight at 80°C and heated in a quartz tube furnace at 700°C for 30 min with a ramp rate of 1°C/min to obtain pristine single-crystalline anatase titanium dioxide nanosupports (TiO$_2$). The pristine nanosupports were subsequently functionalized with 3-triethoxysilylpropyl succinic anhydride (TESPSA, Fisher Scientific, USA) to lead to titanium dioxide nanosupports with carboxyl group functionalities (TiO$_2$-NBs). For this, TESPSA was added to a toluene solution (Fisher Scientific, USA) of titanium dioxide nanosupports and stirred at room temperature for 24 h. Then, the mixture was centrifuged and washed with methanol (Fisher Scientific, USA) to remove excess TESPSA. The TiO$_2$-NBs were dried overnight at 80°C, and stored at room temperature until use.

MULTI-WALLED CARBON NANOTUBE (MWCNTS) FUNCTIONALIZATION

Commercial multi-walled carbon nanotubes (MWCNTs, 95% purity, 10-20 nm in diameter, Nanolab Inc., PD15L5-20, USA) were acid functionalized to obtain carboxyl group functionalities$^{37,38}$. Briefly, 100 mg of MWCNTs were added to a 60 mL mixture of 3:1 (V/V) sulfuric (Fisher Scientific, 96.4%, USA) and nitric (Fisher Scientific, 69.6%, USA) acids and sonicated in an ice bath at a constant temperature of approximately 23°C for 6 h. The MWCNTs-acids mixture was then diluted, filtered, and washed extensively with DI water to remove any catalysts or residues. Thus, carboxyl functionalized MWCNTs (MWCNT-COOH) were dried under vacuum and stored at room temperature until use$^{29,37,38}$.

NANOSUPPORT CHARACTERIZATION

Samples (TiO$_2$ and TiO$_2$-NBs or MWCNTs and MWCNT-COOH) were characterized using Scanning Electron Microscopy (SEM) and Atomic Force Microscopy (AFM). Briefly, samples
(1mg/mL in DI water) were deposited on silica wafers, dried under vacuum, and then examined using a Hitachi S-4700 Field Emission Scanning Electron Microscope with a S-4700 detector combining secondary (SE) and backscattered (BSE) electron detection (in a single unit). For AFM, a Si tip (Asylum Research, 50-90 KHz AC240TS, USA) was employed to investigate the length of pristine and carboxyl functionalized MWCNTs. The samples (0.1mg/mL in DI water) were mounted on a mica substrate (9.5 mm diameter, 0.15-0.21 mm thickness, Electron Microscopy Sciences, USA) and dried under vacuum. Scans were performed in air and images of 10 µm x 10 µm were obtained. At least 30 individual nanotubes from different images were measured to acquire the average length distribution.

TiO₂-NBs were further characterized using Fourier Transform Infrared Spectroscopy (FTIR) in order to confirm carboxyl functionalization. FTIR data were collected in transmission mode using KBr pellet method on a Thermo Nicolet Instrument (USA).

Raman spectroscopy (Renishaw InVia Raman Spectrometer, CL532-100, 100 mW, USA) was used to investigate the chemical structure and properties of MWCNTs (both pristine and carboxyl functionalized MWCNTs). Samples (1mg/mL) were deposited on glass slides (Fisher, USA) and irradiated through a 20x microscope objective using an argon ion (Ar⁺) laser beam performing at 514.5 nm and having a spot size of < 0.01 mm². Low laser energy (i.e., < 0.5 mV) and short exposure time (~10 sec) were set to prevent unexpected heating effects of the sample. Successive scans ranging from 100 to 3200 cm⁻¹ were acquired.

**ENZYME IMMOBILIZATION**

Three different strategies were employed for enzyme immobilization onto the nanosupports (either TiO₂-NBs or MWCNTs-COOH). First, covalent attachment of the enzyme was carried out as previously demonstrated²⁹. Generally, 2 mg of nanosupport was dispersed in 2 mL of 2-(N-morpholino)ethanesulfonic acid sodium salt (MES) buffer (50 mM, pH 4.7, Fisher, USA) containing 160 mM 1-ethyl-3-[3-dimethylaminopropyl]carbodiimide hydrochloride (EDC, Acros Organics, USA) and 80 mM N-hydroxysuccinimide (NHS, Pierce, USA) by brief sonication. After 15 min incubation with shaking at 200 rpm and at room temperature, the nanosupports were filtered and washed thoroughly with MES buffer. Subsequently, the nanosupports were redispersed in a 0.5 mg/mL solution of either chloroperoxidase (CPO, Bioresearch, USA) or glucose oxidase (GOx, Sigma, USA). After 3 h incubation at room temperature with shaking at
200 rpm, the nanosupport-enzyme conjugates were filtered and washed thoroughly with citric acid buffer (CAB, 50 mM, pH 4.8, Sigma, USA) for CPO or sodium phosphate buffer (PBS, 100 mM, pH 7, Sigma, USA) for GOX.

Secondly, covalent binding through a PEG linker was carried out in an effort to bring the enzyme away from the nanosupport\textsuperscript{29}. Briefly, the nanosupports were first activated using EDC/NHS chemistry as previously described, and then incubated for 3 h in 5 mL of 1 mg/ml amino-dPEG\textsubscript{8}-acid in PBS (3.2 nm length, Quanta Biodesign, USA). After 3 h, the PEG-nanosupports were extensively washed with PBS and redispersed in the EDC/NHS=160 mM/80 mM solution for 15 min at 200 rpm and room temperature. Upon activation the nanosupports were filtered, washed thoroughly with MES and redispersed in 0.5mg/mL solution of enzyme (either CPO or GOX) for covalent binding as previously described \textsuperscript{29}.

Thirdly, for the CPO and GOX immobilization onto the same nanosupport, a combined binding strategy was used. Specifically, one enzyme was first covalently attached to MWCNTs-COOH as previously described\textsuperscript{29}. The conjugates were then extensively washed with the necessary buffer and subsequently used for the physical attachment of the second enzyme (initial concentration of 0.5 mg/mL of either CPO or GOX were used during both immobilizations). Physical adsorption was performed for 2 h at room temperature, with shaking at 200 rpm.

After the incubation, the conjugates were extensively washed with the designated buffer. When immobilization was completed, the supernatants and first two washes of each immobilization strategy were isolated to determine enzyme loading.

**ENZYME LOADING**

The concentration of enzyme in the washing solutions or in the supernatants was determined using standard bicinchonic acid (BCA) assay\textsuperscript{29, 31, 39, 40}. Specifically, a working reagent was first prepared by mixing 50 parts of reagent A with 1 part of reagent B (reagents were provided with the BCA Assay kit, Pierce, USA). 50 µL of each washing or supernatant solution was then added to 1 mL of the working reagent and incubated in a water bath at 37°C for 30 min. The absorbance of each sample was monitored at 562 nm using an UV-Vis spectrophotometer (Thermo Scientific EVO300, USA). Standard calibration curves were prepared using the corresponding native enzyme and serial dilutions (from 1 mg/mL to 0.03 mg/mL). The enzyme loadings onto the nanosupports were estimated as the difference between the initial amount of
enzyme added in the immobilization process, and the total amount of enzyme washed out in the isolated supernatants and two washes. Results are reported as mean ± standard deviation with at least six trials for each statistic.

ENZYME ACTIVITY

The enzyme activity upon immobilization was spectrophotometrically assessed using an UV-Vis spectrophotometer. The immobilized enzyme activity is reported as the specific retained activity relative to the same amount of free enzyme in solution. For measuring CPO activity, the conversion of monochlorodimedon (MCD) to dichlorodimedon in the presence of H₂O₂ and sodium chloride (NaCl) was monitored at 278 nm for 2 min. The reaction mixture contained 50 mM CAB, 100 mM NaCl, 0.1 mM MCD, 1 mM H₂O₂, and 20 μL of CPO sample (all reagents were purchased from Sigma, USA unless otherwise specified). For GOX, the production of H₂O₂ was monitored by the oxidation of 0.25 mg/mL ABTS by soybean peroxidase (SBP, Bioresource, USA) in excess at 412 nm for 2 min. The reaction mixture contained 50 mM PBS, 62.5 mM glucose (Acros Scientific, USA), 0.05 mg/ml SBP, 12.5 μg/mL ABTS, and 50 μL of GOX sample. For the multiple enzyme system, the CPO activity assay used glucose (62.5 mM final concentration) instead of free H₂O₂. The rate of HOCl generation was calculated using the extinction coefficient of CPO at 278 nm (ε = 12.2 mM⁻¹ cm⁻¹). The rate of HOCl generation is proportional to the rate of MCD conversion which is calculated knowing the amount of enzyme in the sample being tested, the enzyme extinction coefficient and the beam path length in the spectrophotometer (1 cm in this setup). Results are reported as mean ± standard deviation with at least six trials for each statistic.

DETERMINATION OF TiO₂-NB PHOTOCATALYTIC CAPABILITIES

Hydrogen peroxide (H₂O₂) generated by the TiO₂-NBs was quantified upon incubation of the nanosupport in DI water, under UV light for various time periods, and using two independent assays, i.e. the chloroperoxidase activity assay (CPO, Bioresource, USA), and the Amplex® Red Hydrogen Peroxide/Peroxidase Assay Kit (Invitrogen, USA). For the CPO assay, 5 mL of 1 mg/mL TiO₂-NBs was incubated under UV-A (λ = 316-400 nm) or UV-C (λ = 235-280 nm) irradiation, in ice for 10, 30 or 60 min. The solution was immediately filtered, and the filtrate containing H₂O₂ generated was used as the H₂O₂ source for the previously described CPO
activity assay. Specifically, 20 μL of the filtrate was added to a solution containing 50 mM CAB, 100 mM NaCl, 0.1 mM MCD, and 0.007 mg/mL of free CPO (average concentration in CPO activity assay used for immobilized enzyme) (final volume 1 mL) in a quartz cuvette, and the absorbance was monitored at 278 nm for 5 min on a UV-vis spectrophotometer. For the Amplex® Red Hydrogen Peroxide Assay 500 μL of filtrate was added to 500 μL of Amplex® Red working solution containing 100 μM Amplex® Red reagent and 0.2 U/mL horseradish peroxidase. All reagents are included in assay kit. After 30 min incubation at room temperature, the absorbance of the solution was measured at 560 nm and compared to a calibration curve of known H₂O₂ concentrations.

**CPO-BASED-TiO₂-NB CONJUGATES FOR HOCL GENERATION**

The HOCl generation rate of CPO-TiO₂-NB conjugates was determined based on the CPO activity assay previously described with modifications. Specifically, a 2 mL sample of 1 mg/mL CPO-TiO₂-NB conjugates was incubated under UV-A (λ = 316-400 nm) or UV-C (λ = 235-280 nm) in ice for 10, 30 or 60 min. Immediately following incubation, 40 μL of the solution was added to a reaction mixture consisting of 50 mM CAB, 100 mM NaCl and 0.1 mM MCD (final volume 1 mL) in a quartz cuvette, and the absorbance was monitored at 278 nm for 5 min on a UV-vis spectrophotometer. The rate of HOCl generation was calculated using the extinction coefficient of CPO at 278 nm (ε = 12.2 mM⁻¹ cm⁻¹) as previously described.

**ENZYME KINETIC PARAMETERS DETERMINATION**

The kinetic parameters, Kₘ and Vₘₐₓ values, of the free and immobilized CPO and GOₓ (onto both nanosupports, i.e. TiO₂-NB and MWCNT-COOH) were determined by measuring the initial rates of reaction in the CPO or GOₓ activity assays previously described, with H₂O₂ (0-4 mM final concentration) or glucose (0-100 mM final concentration), respectively, as the substrate. The Kₘ and Vₘₐₓ values for the free and immobilized enzymes were calculated using Lineweaver–Burk plots by using the initial rate of the enzymatic reaction data:

\[
\frac{1}{v} = \frac{k_m}{V_{\text{max}}[S]} + \frac{1}{V_{\text{max}}}, \text{ where } [S] \text{ is the concentration of the substrate, } V \text{ and } V_{\text{max}} \text{ represent the initial and maximum rates of reaction, respectively, and } K_m \text{ is the Michaelis-Menten constant.}
\]
STORAGE STABILITY OF ENZYME-BASED CONJUGATES

The storage stability of the free and CPO-carbon nanomaterial-based conjugates was investigated by measuring the activities of the conjugates stored at 4 or 23 °C for a 4 week period. The activities were measured after 1, 2 and 4 weeks using the previously reported MCD assay.

RESULTS AND DISCUSSION

We are focusing our efforts on creating an active enzyme-nanosupport hybrid system to be incorporated into coatings and allow in situ generation of hypochlorous acid (HOCl). Our hypothesis is that the working enzyme, chloroperoxidase (CPO), immobilized onto nanosupports can use H$_2$O$_2$ produced at the nanosupport interface to convert (Cl$^-$) into HOCl. HOCl has a wide spectrum of activity against a variety of pathogens from spores to bacteria$^{30, 31, 36}$. Two different strategies were proposed and evaluated to determine the efficiency and yield of HOCl generation. Further, the enzyme kinetics and stability were assessed.

STRATEGY 1: ENZYME-PHOTOCATALYST HYBRID SYSTEMS FOR IN SITU GENERATION OF HOCl

In the first strategy, we hypothesized that titanium dioxide can produce H$_2$O$_2$ as the substrate to be used by CPO to for the conversion of (Cl$^-$) into HOCl. Titanium dioxide is a widely studied photocatalyst that produces reactive oxygen species from water when excited under UV light$^{42-44}$. Even though the photocatalyst has been extensively studied for the decontamination of a wide variety of water contaminants$^{43, 45}$ as well as contaminants in air$^{46}$, to our knowledge there are no previous studies that attempted using the titanium dioxide as a scaffold nanosupport to generate H$_2$O$_2$ for further enzyme kinetics.

The scaffold nanosupport was produced from pristine anatase titanium dioxide via hydrothermal processing$^{47}$. Specifically, pristine anatase titanium dioxide nanosupports were carboxyl functionalized using 3-triethoxysilylpropyl succinic anhydride to allow the formation of titanium dioxide nanobelts or TiO$_2$-NBs (Figure 1a). Carboxyl functionalities were confirmed using Fourier Transform Infrared Spectroscopy (FTIR; Figure 1b). A large peak was identified at 1731 cm$^{-1}$ confirming the presence of the C=O bond onto the TiO$_2$-NBs$^{48}$.
Figure 1: a) Functionalization of photocatalyst pristine titanium dioxide results in the formation of carboxyl functionalized nanobelts or TiO₂-NBs. b) FTIR spectrum of TiO₂-NBs reveals the presence of the carboxyl peak at 1731 cm⁻¹, confirming -COOH functionalization. c) CPO enzyme immobilization onto TiO₂-NBs with and without the use of a PEG linker. The CPO-TiO₂-NBs-based conjugates generate HOCl under UV irradiation.
To determine whether the carboxyl functionalization changed the nanosupport morphology, pristine anatase titanium dioxide and TiO$_2$-NBs nanosupports were investigated using Scanning Electron Microscopy (SEM). SEM images of the nanosupports before and after carboxyl functionalization are shown in Figure S1 (Appendix 2). The nanosupports were found to be 60-300 nm wide and several micrometers in length. No significant changes were identified in the TiO$_2$-NBs morphology or length distribution upon carboxyl functionalization.

In order to generate enzyme-hybrid systems, we used the TiO$_2$-NBs as scaffolds for the covalent immobilization of CPO enzyme through EDC/NHS chemistry$^{49, 50}$ or EDC/NHS chemistry with a PEG spacer$^{29}$ (Figure 1c). The PEG spacer was highly hydrophilic, with a length of 3.2 nm; previous studies have shown that such a spacer has little or no chemical effect on enzyme immobilization and allows for improved solubility of the nanosupport$^{51}$. The CPO-TiO$_2$-NBs and CPO-PEG-TiO$_2$-NBs conjugates showed enzyme loadings of 0.10 ± 0.03 and 0.04 ± 0.02 mg enzyme/mg nanosupport, respectively, which represented 20 % and 8 % of the amount of protein that was offered, respectively (Table 1).

**Table 1: Characterization of CPO-TiO$_2$ hybrid system for *in situ* generation of HOCl.**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>CPO-TiO$_2$</th>
<th>CPO-PEG-TiO$_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Loading (mg protein/mg nanosupport)</td>
<td>0.10 ± 0.03</td>
<td>0.04 ± 0.02</td>
</tr>
<tr>
<td>Loading function of enzyme offered (%)</td>
<td>20</td>
<td>8</td>
</tr>
<tr>
<td>Specific retained activity relative to free enzyme (%)</td>
<td>0.30 ± 0.13</td>
<td>12.00 ± 1.63</td>
</tr>
<tr>
<td>$V_{\text{max}}$ ($\mu$M $\mu$g$^{-1}$ s$^{-1}$)</td>
<td>0.54 ± 0.11</td>
<td>4.64 ± 0.87</td>
</tr>
<tr>
<td>$K_m$ ($\mu$M H$_2$O$_2$)</td>
<td>270 ± 80</td>
<td>490 ± 70</td>
</tr>
<tr>
<td>$K_{\text{cat}}$ ($s^{-1}$)</td>
<td>22.68 ± 4.62</td>
<td>194.88 ± 36.54</td>
</tr>
<tr>
<td>η</td>
<td>0.02</td>
<td>0.17</td>
</tr>
</tbody>
</table>
The activity of the CPO immobilized at the TiO$_2$-NBs interface was evaluated using the colorimetric reaction showing the conversion of monochlorodimedon to dichlorodimedon in the presence of H$_2$O$_2$. The CPO-TiO$_2$-NBs conjugates retained about 0.3%, while CPO-PEG-TiO$_2$-NBs conjugates retained about 12% activity when compared to the activity of the same amount of free CPO in solution (Table 1). The low activity observed for CPO-TiO$_2$-NBs is attributed to the interface reactions that take place at the TiO$_2$-NBs nanosupport surface. Specifically, previous studies have shown that nanosupports with lower curvature favor enzyme-enzyme interactions and non-specific attachment of enzymes that could lead to enzyme denaturation.$^{8,29,30}$ Further, at the working pH of 4.8 that is slightly above the isoelectric point of CPO (pI = 4)$^{52}$ the CPO enzyme has a net negative charge and the TiO$_2$-NBs (isoelectric point pI = 6.5)$^{53}$ has a net positive charge, thus favoring the adsorption or non-specific binding of the CPO to the nanosupport and accounting for the reduced enzyme activity. Meanwhile, the higher activity observed for the CPO-PEG-TiO$_2$-NBs was presumably due to the PEG spacer bringing the enzyme away from the nanosupport thus reducing non-specific enzyme interaction with the nanosupport and thus enzyme denaturation at the nanointerface.$^{31}$

Kinetic constants were evaluated for the free CPO and compared to the kinetics of the CPO-based conjugates, i.e., CPO-TiO$_2$-NBs and CPO-PEG-TiO$_2$-NBs, by using Lineweaver–Burk plots$^{54}$. Namely, the $K_m$ (substrate concentration at which the initial reaction rate is half maximal) and $V_{max}$ (maximum initial rate of an enzyme catalyzed reaction) values of the CPO-TiO$_2$-NBs and CPO-PEG-TiO$_2$-NBs conjugates are shown in Table 1 and compared to the free CPO in solution. The $K_m$ values were on the same order of magnitude for all analyzed samples; specifically found to be 480, 270 and 490 μM, respectively, for the free, CPO-TiO$_2$-NBs and CPO-PEG-TiO$_2$-NBs conjugates indicating that there was no significant conformational change of the enzyme active site upon immobilization. The apparent $K_m$ for the directly covalently conjugated enzyme was decreased by about 40% when compared to the free enzyme or the PEG covalently conjugated enzyme that showed no significant change. $V_{max}$ values were on the same order of magnitude (i.e., 27, 0.5 and 4.6 μM mg$^{-1}$ s$^{-1}$, respectively for the free, CPO-TiO$_2$-NBs and CPO-PEG-TiO$_2$-NBs conjugates); the $V_{max}$ for the covalently immobilized enzyme decreased about 98% and about 81% for the PEG immobilized samples when compared to the free enzyme in solution. The smaller $V_{max}$ obtained when the enzyme was immobilized directly via covalent binding was presumably due to the enzyme coming into direct contact with the
nanosupport, decreasing the chance of a reaction to occur and consequently slowing the reaction rate. Our reported \( K_m \) and \( V_{\text{max}} \) results are on the same order of magnitude with previous studies reporting on the CPO immobilization at the carbon nanomaterial interface\(^{30}\), polymer coated magnetic nanoparticles interface\(^{55}\) or onto mesoporous silicate material\(^{56}\). Any deviations observed may result from the environmental differences in which the experiments were performed.

The efficiency factor \( \eta \) was calculated from the maximum reaction rates of the immobilized CPO (both CPO-TiO\(_2\)-NBs and CPO-PEG-TiO\(_2\)-NBs) relative to the rate of the free enzyme in solution using

\[
\eta = \frac{v_{\text{immobilized}}}{v_{\text{free}}},
\]

where \( v_{\text{immobilized}} \) is the reaction rate of the immobilized enzyme (directly through covalent binding or through PEG and covalent binding) and \( v_{\text{free}} \) is the reaction rate of the free enzyme (Table 1). The efficiency factor for the CPO-TiO\(_2\)-NBs was 0.02 while the \( \eta \) for the CPO-PEG-TiO\(_2\)-NBs was 0.17. The reduction in the efficiency factor observed for the immobilized enzyme further confirmed the non-specific interactions of the enzymes at the nanosupport interface and thus enzyme denaturation\(^{8, 29, 30}\).

To test whether the enzyme-nanosupport hybrid systems can generate \textit{in situ} H\(_2\)O\(_2\), we irradiated the systems under UV (i.e. UV-A (\( \lambda = 316-400 \) nm) or UV-C (\( \lambda = 235-280 \) nm), Figure 1c). Our data showed that there was negligible capability to allow the conversion of monochlorodimedon to dichlorodimedon in the presence of Cl\(^-\). Further, the enzyme-nanosupport hybrid systems showed negligible capability for HOCl generation even after 10, 30 or 60 min of continuous UV irradiation (both UV-A and UV-C). The negligible capabilities of the enzyme-nanosupport hybrid systems to ensure the conversion of monochlorodimedon to dichlorodimedon, or to generate HOCl upon UV irradiation, was presumably due to the limited H\(_2\)O\(_2\) substrate concentration being available at the nanosupport interface. Specifically, with the substrate being exhausted in the CPO conversions (either of the monochlorodimedon to dichlorodimedon, or of the Cl\(^-\)), the initially zero order reaction becomes a “leading” reaction, which does not allow the formation of high enough concentrations of enzyme-substrate complexes to become detectable. This was confirmed in our control experiments performed with free CPO (the same amount of free enzyme in solution as the amount of enzyme immobilized
onto nanosupport and determined by the loading assay) that showed that for the reaction to be detectable, at least 1 µM of H₂O₂ substrate needs to be present in solution (Figure S2, Appendix 2). Complementary, the suggested “leading reaction” is also supported by previous reports that have shown that the rate of H₂O₂ decomposition at the nanosupport interface is sharply increased at pH values above 4⁵⁷ (our working pH is 4.8). Further, this rate of H₂O₂ decomposition could be accentuated as result of an initial adsorption step of H₂O₂ onto the nanosupport followed by a second process that consists of the cleavage of H₂O₂ at the oxide-based nanosupport interface⁵⁸,⁵⁹. Additionally, previous studies have shown the possibility of enzyme deactivation in enzyme-TiO₂ systems in the presence of UV light, which would add to loss of activity³⁹,⁴⁰.

While this strategy did not demonstrate the capability to detect HOCl in situ produced by CPO immobilized onto TiO₂-NBs nanosupports, they demonstrated the feasibility of covalently immobilizing CPO at oxide-based interfaces to lead to active enzyme-based conjugates, with higher activities and kinetics being observed for the enzyme immobilized using a spacer. Such enzyme-nanosupport hybrid systems can further be exploited for the next generation of biosensors with electrochemical performances to be studied by cyclic voltammetry (CV) and amperometric methods⁶⁰-⁶².

STRATEGY 2: CO-IMMOLIZED ENZYME-CARBON-BASED MATERIAL HYBRID SYSTEMS FOR IN SITU GENERATION OF HOCL

In our second strategy, we hypothesized that carboxyl functionalized multi-walled carbon nanotubes (MWCNTs-COOH) can be used as an alternative to TiO₂-NBs nanosupports to allow generation of H₂O₂ substrate necessary for the conversion of (Cl⁻) into HOCl in the presence of CPO. Our hypothesis was that H₂O₂ substrate can be produced through an enzymatic chain reaction at the MWCNT interface. Specifically, glucose oxidase (GOₓ) can be used for H₂O₂ generation⁶³ that would further serve as substrate for the co-immobilized CPO to convert (Cl⁻) into HOCl (Figure 2). MWCNTs-COOH were chosen as nanosupports because they have a significantly smaller diameter (MWCNTs diameter: 10-20 nm) and thus a higher surface curvature when compared to TiO₂-NBs (TiO₂-NBs diameter: 60-300 nm). The higher surface curvature will presumably reduce the non-specific interactions of the CPO with the nanosupport interface⁵⁹,⁶¹,⁶⁴, thus reducing enzyme denaturation and activity loss observed in Strategy 1. Further, the choice of the MWCNTs-COOH will avoid any H₂O₂ decomposition at the
nanosupport interface thus making the hypothesis for the HOCl generation viable. Moreover, the high aspect to surface area ratio of the MWCNTs-COOH will allow high CPO loadings and ease of isolation of the enzyme-based hybrid systems through filtration\textsuperscript{29, 31}.

**Figure 2:** a) Acids treatment of MWCNTs leads to carboxyl functionalized MWCNTs (MWCNTs-COOH). Functionalization takes place at the defect sites in the MWCNTs structures; the resulting acid treated MWCNTs have hydrophilic residues (represented by the COOH groups) and hydrophobic walls. The MWCNTs-COOH are used as nanosupports for co-immobilization of CPO and GO\textsubscript{X}. First, CPO is covalently attached to COOH-functionalized MWCNTs. Subsequently, the CPO-based conjugates are used for the physical attachment of GO\textsubscript{X}; GO\textsubscript{X} will attach to the hydrophobic walls of the nanotubes to result in CPO-GO\textsubscript{X}-MWCNTs conjugates. In the system containing the co-immobilized enzymes and through a chain reaction, GO\textsubscript{X} provides the H\textsubscript{2}O\textsubscript{2} substrate needed by CPO for in situ conversion of Cl\textsuperscript{-} into HOCl. b) Raman spectra of pristine (black curve) and carboxyl functionalized (red curve) MWCNTs. The carboxyl functionalized MWCNTs have shifted peaks towards higher relative intensities confirming the COOH functionalization.
MWCNTs were first carboxyl functionalized through acid treatment as previously described\textsuperscript{29} (Figure 2a). To evaluate whether there were any morphological and structural changes upon acid treatment, the resulting carboxyl functionalized MWCNTs (MWCNT-COOH) were characterized using Scanning Electron Microscopy (SEM), Atomic Force Microscopy (AFM) and Raman spectroscopy. No morphological changes were observed for the acid treated samples (Figure S3, Appendix 3) relative to pristine MWCNTs. However, the carboxyl functionalized MWCNTs were significantly shorter than pristine MWCNTs, with average lengths of $443 \pm 238$ when compared to the pristine original lengths of $5126 \pm 2283$ nm. The carboxyl functionalization was confirmed using Raman spectroscopy (Figure 2b). Specifically, the Raman spectrum showed the presence of 4 bands for both pristine and carboxyl functionalized MWCNTs samples, i.e. the so-called D band at around $1340$ cm\textsuperscript{-1}, G band at around $1580$ cm\textsuperscript{-1}, G’ band at around $2670$ cm\textsuperscript{-1} and another band at around $2920$ cm\textsuperscript{-1}. The D band is associated with non-crystalline carbon species, such as defect sites on the MWCNTs wall surface\textsuperscript{65}. Compared to the pristine MWCNTs samples (black curve), the spectrum of the carboxyl functionalized samples (red curve) showed a wider D band shifted towards a higher frequency, indicating the addition of carboxyl groups on the nanotubes sidewall. The G band observed around $1585$ cm\textsuperscript{-1} showed the high degree of ordering of the MWCNTs\textsuperscript{66}. Shifting the G band in carboxyl functionalized MWCNTs was associated with either the removal of metal catalyst particles, the increase of electron-accepting functional groups, or generation of amorphous carbon species\textsuperscript{37}. Further, the $I_D/I_G$ ratio (ratio of the relative intensity of the D band relative to the G band indicates the level of functionalization) of MWCNTs increased from 0.457 to 0.817, further confirming the carboxyl group functionalities (Table S1, Appendix 2).

To test the feasibility of the proposed approach, we first assessed whether MWCNTs would serve as viable nanosupports for CPO and GOx immobilization through either physical or covalent binding, and whether they ensure high activity and loading for each one of the enzymes being immobilized. The CPO-MWCNTs physically and covalently bound conjugates showed loadings of $0.10 \pm 0.02$ and $0.07 \pm 0.02$ mg enzyme/mg nanosupport, respectively, which represented $20\%$ and $14\%$ of the amount of protein that was offered, respectively (Table S2). The specific activity of the physically bound conjugates was about $41\%$ while the activity of the covalently bound conjugates was about $52\%$ of the activity of the free enzyme in solution. The GOx-MWCNTs physically and covalently bound conjugates showed loadings of $0.24 \pm 0.02$ and
0.25 ± 0.02 mg enzyme/mg nanosupport, which represented 48% and 50% of the amount of protein that was offered, respectively (Table S3, Appendix 2). The specific activity of the physically bound conjugates was about 10%, while the activity of the covalently bound conjugates was about 38% of the activity of the free enzyme in solution. The higher activities observed for both covalently bound CPO- and GOX-MWCNTs conjugates was a result of the reduced non-specific binding of the enzyme onto the nanosupport and reduced enzyme-enzyme interactions\(^{29, 31, 64}\). Specifically, the smaller curvature of the cylindrical nanotube will result in an increase in the center-to-center distance between adjacent enzymes when compared to the enzyme immobilized onto TiO\(_2\)-NBs nanosupport thus reducing unfavorable lateral interactions that could lead to enzyme deactivation\(^{67}\).

Kinetic constants were also evaluated and compared to the free enzyme in solution. The \(K_m\) and \(V_{max}\) values of the free and CPO-based conjugates are shown in Figure 3a and Table S2 (Appendix 2). The \(K_m\) values were on the same order of magnitude for all analyzed samples indicating that, similarly to the CPO-TiO\(_2\)-based conjugates, the CPO-MWCNTs-based conjugates (either physically or covalently immobilized) did not show significant conformational change of the enzyme active site upon immobilization. Specifically, the values found were 480, 340 and 530 \(\mu\)M, for the free, CPO physically adsorbed and CPO covalently bound, respectively. The apparent \(K_m\) for the directly covalently conjugated enzyme was increased by about 10% when compared to the free enzyme. The \(V_{max}\) values were also in the same order of magnitude and found to be 27, 12 and 13 \(\mu\)M \(\text{mg}^{-1} \text{s}^{-1}\), respectively for the free CPO, CPO physically adsorbed and CPO covalently immobilized onto MWCNTs. The \(V_{max}\) values of both the covalently and physically immobilized enzyme decreased by only about 50% when compared to the free enzyme in solution. This represents an increase of about 61% when compared to the \(V_{max}\) of the CPO-TiO\(_2\)-based conjugates further supporting that MWCNTs nanosupports provide a viable alternative in terms of reducing the enzyme non-specific interaction to the interface that could have slowed down the reaction rate.
Figure 3: a) Michaelis-Menten kinetics of enzyme-based MWCNTs conjugates. a) CPO-based conjugates (physically immobilized-open circles; covalently immobilized- filled triangles) kinetics relative to free CPO in solution (filled squares). b) GOx-based conjugates (physically immobilized-open circles; covalently immobilized- filled triangles) kinetics relative to free GOx in solution (filled squares).
The $K_m$ and $V_{max}$ values of the free and GOX-based conjugates (both physically and covalently immobilized) are shown in Figure 3b and Table S3 (Appendix 2). Specifically, the $K_m$ values were 2,600, 1,800, and 3,200 μM for free GOX, GOX physically and GOX covalently immobilized, respectively. The apparent $K_m$ for the covalently bound enzyme decreased about 23% whereas the value for the physically bound enzyme increased about 30%. The $V_{max}$ values were found to be 0.49, 0.03 and 0.20 μM mg$^{-1}$ s$^{-1}$, respectively for free GOX, GOX physically and GOX covalently bound. The 94% and 60% decreases in $V_{max}$ for covalently bound and physically adsorbed GOX, were presumably due to the non-specific attachment of the GOX or enzyme-enzyme interaction at the nanointerface that could have slowed the reaction rate.

The efficiency factor $\eta$ was also calculated from the maximum reaction rates of the immobilized CPO or GOX (both through physical and covalent binding) and relative to the rate of the free enzyme in solution (see Strategy 1 and Tables S2 and S3, respectively, Appendix 2). The efficiency factor for the CPO-physically adsorbed conjugates was 0.45 while the $\eta$ for the CPO immobilized via covalent binding was 0.47. The efficiency factor for the GOX-physically adsorbed conjugates was 0.06 while the $\eta$ for the GOX immobilized via covalent binding was 0.40. The catalytic efficiency was increased when compared to the covalently bound CPO-TiO$_2$-based conjugates further supporting the conjugates obtained at the MWCNTs interface were highly active and viable platform for the enzyme co-immobilization strategy.

Once it was confirmed that individual CPO and GOX resulted in highly active conjugates at the MWCNTs interface, MWCNTs-COOH were used for the co-immobilization of CPO and GOX to lead to the next generation of in situ microbial decontamination hybrid systems (Figure 2a). For such a system to be viable, both enzymes must be successfully immobilized onto the surface of the same nanosupport, remain active upon immobilization and allow the promotion of a chain reaction through their co-immobilization conditions (i.e. the product of one enzyme to serve as the substrate of the co-immobilized enzyme in order to generate the HOCl decontaminant). We rationalized that using a -COOH functionalized nanosupport is a viable platform to ensure a successful co-immobilization strategy since the -COOH groups could serve as active groups for covalent immobilization of one enzyme, while the hydrophobic walls of the nanotube will provide additional space for physical adsorption of a second enzyme$^{68}$. Briefly, either CPO or GOX was covalently immobilized onto MWCNTs-COOH using EDC/NHS
chemistry followed by the physical adsorption of the second enzyme. Both combinations were investigated (i.e. CPO covalent and GOX physical; and GOX covalent and CPO physical). The resulting loadings of the two co-immobilized enzymes onto the same nanosupport are shown in Table 2. CPO-GOX-based MWCNTs-COOH hybrid systems were further used to test whether they allow the generation of HOCl through a chain reaction at the nanotube interface (Figure 2a; i.e. to test whether GOX will generate enough H2O2 for the conversion of (Cl−) in the presence of CPO into HOCl). Our data has shown that the hypothesized co-immobilization strategy was highly efficient in that both enzymes maintained their individual activities and kinetics characteristics and were able to generate HOCl as the final product from their chain reaction at the nanosupport interface. The rate of in situ HOCl generation correlated with the loadings, activities and kinetics of the two enzymes. Our data showed that the chain reaction at the nanotube interface allowed the highest rate of HOCl generation, i.e. 0.11 ± 0.03 µM µg−1 s−1, to be achieved by the conjugates that contained covalently immobilized CPO and physically adsorbed GOX. The conjugates that contained covalently immobilized GOX and physically adsorbed CPO allowed 0.04 ± 0.01 µM µg−1 s−1 rate of HOCl generation. Such rate was previously shown to adequately induce microbial and bacterial decontamination. Further, our strategy demonstrated that these conjugates retained about 50% of their activities even after 4 weeks storage at 4°C and about 40% at 23°C (Table S4, Appendix 2).

**Table 2:** CPO-MWCNT-GOX conjugates generate HOCl in situ. The rate of HOCl generation is dependent on the enzyme immobilization conditions (i.e., through physical or covalent binding).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>CPO-MWCNT-GOX (CPO covalent GOX physical)</th>
<th>CPO-MWCNT-GOX (GOX covalent CPO physical)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Loading (mg protein/mg nanosupport)</td>
<td>CPO: 0.07 ± 0.02</td>
<td>CPO: 0.09 ± 0.02</td>
</tr>
<tr>
<td></td>
<td>GOX: 0.19 ± 0.04</td>
<td>GOX: 0.17 ± 0.02</td>
</tr>
<tr>
<td>Loading function of enzyme offered (%)</td>
<td>CPO: 14</td>
<td>CPO: 18</td>
</tr>
<tr>
<td></td>
<td>GOX: 38</td>
<td>GOX: 34</td>
</tr>
<tr>
<td>Rate of HOCl (µM µg−1 s−1)</td>
<td>0.11 ± 0.03</td>
<td>0.04 ± 0.01</td>
</tr>
</tbody>
</table>
Our 2\textsuperscript{nd} Strategy showed the feasibility of \textit{in situ} HOCl generation at a nanosupport interface through an enzymatic chain reaction. This strategy can overcome the disadvantages of immobilizing enzymes onto different nanosupports and mixing them upon use, such as the reductions in reactivity and stability that are usually correlated with differences in optimal pH’s\textsuperscript{70}. Further, such enzyme-based conjugates can be incorporated into coatings to create self-sustainable surfaces with enhanced microbial decontamination capabilities\textsuperscript{8, 29-31}. For instance, one could envision the encapsulation of dextrin or dextrin-derivates into paint or polymer-based coatings\textsuperscript{29, 31, 71, 72} together with these prepared enzyme-carbon-based material hybrid systems; further exposure of such coatings to ambient (Cl\textsuperscript{–})\textsuperscript{73} will lead to \textit{in situ} generation of HOCl and thus provide a self-cleaning and self-sustainable microbial decontamination coating.

**CONCLUSIONS**

Two different strategies were explored for \textit{in situ} generation of HOCl through an enzymatic reaction. In our strategies CPO working enzyme was immobilized either at a photocatalyst- (TiO\textsubscript{2}-NB\textsubscript{s}) or at a carbon-based nanosupport (MWCNTs-COOH) interface. CPO immobilized onto MWCNTs showed 52\% increase in the specific retained activity when compared to the CPO immobilized onto TiO\textsubscript{2}-NB\textsubscript{s}. CPO-MWCNT-based hybrid systems were capable of generating HOCl at a high rate known to be feasible for microbial decontamination. This research can be viewed as an important first step toward creating self-sustainable microbial decontamination coatings to be used against various pathogens such as bacteria and spores. Further, these co-immobilized-enzyme-MWCNTs-based conjugates are interesting as active biohybrid nanomaterials; for instance, one can arrange given enzymes onto the same nanosupport according to a specific function thus making sequential enzymatic reactions at nanobiointerfaces become feasible for biosensor applications.
REFERENCES


APPENDIX 2

Figure S1: SEM images of a) pristine titanium dioxide and b) TiO$_2$-NBs. No changes in the morphology of the samples have been identified after -COOH functionalization.

Figure S2: Colorimetric reaction showing the conversion of MCD at constant CPO concentration and various H$_2$O$_2$ concentrations. The minimum H$_2$O$_2$ for which conversion was observed was 1 µM.
Figure S3: SEM images of a) Pristine MWCNTs and b) Carboxyl functionalized MWCNTs. No morphological differences were recorded between the pristine and the acids treated samples.

Table S1: MWCNTs Raman analysis data.

<table>
<thead>
<tr>
<th>Sample</th>
<th>D band position (cm$^{-1}$)</th>
<th>G band position (cm$^{-1}$)</th>
<th>$I_D/I_G$ Intensity ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pristine MWCNTs</td>
<td>1341</td>
<td>1571</td>
<td>0.457</td>
</tr>
<tr>
<td>6h cut MWCNTs</td>
<td>1346</td>
<td>1580</td>
<td>0.817</td>
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</table>
Table S2: CPO-MWCNT conjugates immobilization data and kinetics.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>CPO-MWCNT (covalent)</th>
<th>CPO-MWCNT (physical)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Loading (mg protein/mg nanosupport)</td>
<td>0.07 ± 0.02</td>
<td>0.10 ± 0.02</td>
</tr>
<tr>
<td>Percentage of enzyme offered (%)</td>
<td>14</td>
<td>20</td>
</tr>
<tr>
<td>Specific Retained Activity (%)</td>
<td>52.70 ± 6.26</td>
<td>41.36 ± 6.65</td>
</tr>
<tr>
<td>$V_{\text{max}}$ (µM µg$^{-1}$ s$^{-1}$)</td>
<td>12.79 ± 1.97</td>
<td>12.28 ± 2.56</td>
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<tr>
<td>$K_{\text{m}}$ (µM H$_2$O$_2$)</td>
<td>530 ± 60</td>
<td>340 ± 70</td>
</tr>
<tr>
<td>$K_{\text{cat}}$ (s$^{-1}$)</td>
<td>537.18 ± 82.74</td>
<td>515.76 ± 107.52</td>
</tr>
<tr>
<td>$\eta$</td>
<td>0.47</td>
<td>0.45</td>
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**Table S3:** GOx-MWCNT conjugates immobilization data and kinetics.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>GOx-MWCNT (covalent)</th>
<th>GOx-MWCNT (physical)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Loading (mg protein/mg nanosupport)</td>
<td>0.25 ± 0.02</td>
<td>0.24 ± 0.02</td>
</tr>
<tr>
<td>Percentage of enzyme offered (%)</td>
<td>50</td>
<td>48</td>
</tr>
<tr>
<td>Specific Retained Activity (%)</td>
<td>37.67 ± 4.70</td>
<td>9.89 ± 3.30</td>
</tr>
<tr>
<td>$V_{\text{max}}$ ($\mu$M $\mu$g$^{-1}$ s$^{-1}$)</td>
<td>0.196 ± 0.032</td>
<td>0.029 ± 0.003</td>
</tr>
<tr>
<td>$K_m$ ($\mu$M glucose)</td>
<td>3,200 ± 700</td>
<td>1,800 ± 800</td>
</tr>
<tr>
<td>$K_{\text{cat}}$ (s$^{-1}$)</td>
<td>31.36 ± 5.12</td>
<td>4.64 ± 0.48</td>
</tr>
<tr>
<td>$\eta$</td>
<td>0.40</td>
<td>0.06</td>
</tr>
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</table>

**Table S4:** CPO-MWNT-GOx conjugates stability data

<table>
<thead>
<tr>
<th>Storage Temperature (°C)</th>
<th>Immobilization Method</th>
<th>Rate after 1 Week (% of original)</th>
<th>Rate after 2 Weeks (% of original)</th>
<th>Rate after 4 Weeks (% of original)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>CPO covalent GOx physical</td>
<td>80.50 ± 2.77</td>
<td>58.41 ± 9.73</td>
<td>51.22 ± 7.32</td>
</tr>
<tr>
<td>4</td>
<td>GOx covalent CPO physical</td>
<td>98.86 ± 3.54</td>
<td>96.29 ± 3.71</td>
<td>71.02 ± 9.50</td>
</tr>
<tr>
<td>23</td>
<td>CPO covalent GOx physical</td>
<td>52.12 ± 12.11</td>
<td>26.04 ± 10.66</td>
<td>39.23 ± 17.18</td>
</tr>
<tr>
<td>23</td>
<td>GOx covalent CPO physical</td>
<td>68.94 ± 16.82</td>
<td>54.08 ± 16.31</td>
<td>39.45 ± 11.53</td>
</tr>
</tbody>
</table>
Effects of acid treatment on structure, properties and biocompatibility of carbon nanotubes

Chenbo Dong, Alan S. Campell, Reem Eldawud, Gabriela Perhinschi, Yon Rojanasakul, Cerasela Zoica Dinu

A R T I C L E   I N F O

Article history:
Received 28 July 2012
Received in revised form 25 September 2012
Accepted 28 September 2012
Available online 23 October 2012

Keywords:
Nanotubes
Acid treatment
Structure modification
Cytotoxicity
Biocompatibility

A B S T R A C T

Carbon nanotubes (CNTs) are promising to be the next generation of viable tools for bioapplications. Further advances in such bioapplications may depend on improved understanding of CNTs physical and chemical properties as well as control over their biocompatibility. Herein we performed a systematic study to show how acid oxidation treatment changes CNTs physical and chemical properties and leads to improved CNTs biocompatibility. Specifically, by incubating CNTs in a strong acid mixture we created a user-defined library of CNTs samples with different characteristics as recorded using Raman energy dispersive X-ray spectroscopy, atomic force microscopy, or solubility tests. Systematically characterized CNTs were subsequently tested for their biocompatibility in relation to human epithelial cells or enzymes. Such selected examples are building pertinent relationships between CNTs biocompatibility and their intrinsic properties by showing that acid oxidation treatment lowers CNTs toxicity providing feasible platforms to be used for biomedical applications or the next generation of biosensors.

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1. Introduction

Carbon nanotubes (CNTs) are nanoscale diameter materials of tubular shape and micrometer length with many interesting properties that make them viable candidates for a wide range of applications including electrical circuits [1], hydrogen storage [2], fiber optics [3], and conductive plastics [4]. In recent years, CNTs functionalization with biomolecules such as proteins [5], enzymes [6,7] or nucleic acids [8] opened up exciting bioapplications in biolabeling [9], biosensing [10], drug delivery [11], bioseparation [12] and tissue engineering [13]. However, further development of such bioapplications is hindered by: (1) CNTs limited available surface area for biomolecule functionalization [14], (2) lack of understanding of CNTs growth mechanisms in uncontaminated forms [15], (3) CNTs structural instability since larger nanotubes are prone to kinking and collapsing [16,17], and (4) CNTs cytotoxicity and associated health risks posed during their manufacturing and processing [18]. These challenges are mainly associated with the fact that as-produced CNTs form large aggregates in liquid environments since their hydrophobic walls are prone to van der Waals interactions [19]. Thus, in order to increase CNTs bioapplications [20] and reduce their aggregation [21] and cytotoxicity [22], it is critical to overcome their intrinsic hydrophobicity and tendency to form conglomerates in solution. Numerous attempts have been made to overcome CNTs hydrophobicity and increase their hydrophilicity; these include gas- [23] and liquid-phase activation [24], and oxidation with strong oxidants including hydrogen peroxide [25], potassium permanganate [26], potassium hydroxide [27], and nitric and/or sulfuric acid [6,7,28]. Among these attempts, nitric and sulfuric acid oxidation is regarded as the most prevalent treatment since it is easy to implement in both laboratory and industrial settings [20]. When CNTs are oxidized with such aggressive acids, their hydrophilicity is increased by the introduction of oxygen-containing functional groups, i.e., carboxyl [29], carbonyl [26,29], and phenol groups [30]. Moreover, during such oxidation treatments amorphous carbon [31] and residual metal catalyst particles are removed, possibly resulting in reduced intrinsic toxicity of CNTs [22]. Despite the fact that wide evaluations of the effects of acid oxidation on CNTs have been carried out, systematic investigations of changes in physical and chemical properties and how such changes can be further employed for increasing CNTs biocompatibility and thus bioapplications are still lacking.

Herein we performed a systematic study of the changes in physical and chemical properties of pristine CNTs upon user-controlled
treatment with nitric and sulfuric acids. Further, we assessed how these changes affect CNTs biocompatibility in relation to cellular and enzymatic systems [6,7,10]. Our hypothesis was that selected biological examples will help build pertinent relationships between CNTs biocompatibility and their intrinsic properties and demonstrate how interface reactions between a biological molecule and the nanomaterial can be further used to provide systems with lower toxicity to be used for selected bioapplications as well as feasible platforms for the next generation of biosensors.

2. Materials and methods

2.1. Acid oxidation of CNTs

Acid oxidation treatment of single- and multi-walled carbon nanotubes (SW- and MWCNTs, respectively) was employed to generate a library of samples with different physical and chemical properties. Specifically, commercial SWCNTs (85% purity, Unidym Inc.) and MWCNTs (85% purity, Nanolab Inc. (PD15L5-20)) were incubated in a concentrated sulfuric (96.4%, Fisher, USA) and nitric acid (69.5%, Fisher, USA) mixture in a ratio of 3:1 (V/V). The CNTs/acid mixture (where the CNTs can refer to either SW- or MWCNTs) was subsequently sonicated in an iced bath (Branson 2510, Fisher, USA) for 1, 3, or 6 h, at a constant temperature of 23 °C. When the required time elapsed, CNTs/acid mixture was diluted with deionized (di) water and filtered through a 0.2 μm polycarbonate filter membrane (Fisher, USA). Several cycles of resuspension in di water were employed to remove acidic residues or catalysts. The CNTs were isolated on the filter, subsequently dried in a vacuum desiccator and stored at room temperature for further use.

2.2. Energy dispersive X-ray analysis (EDX) of CNTs

Energy dispersive X-ray analysis (EDX) was used for quantitative elemental analysis of pristine and acid oxidized CNTs. Samples (1 mg/ml in di water) were deposited on silica wafers and dried under vacuum. The experiments were performed on a Hitachi S-4700 Field Emission Scanning Electron Microscope (USA) with a S-4700 detector combining secondary (SE) and backscattered (BSE) electron detection (all in a single unit), operating at 20 kV. Results are presented as a percent of elements relative to the most dominant element.

2.3. Scanning Electron Microscopy (SEM) of CNTs

Samples (1 mg/ml in di water of both pristine and acid treated CNTs) were dried on silica wafers under vacuum and imaged using a Hitachi S-4700 Field Emission Scanning Electron Microscope (USA) with a field emission at 10 kV.

2.4. Raman spectroscopy of CNTs

Raman spectroscopy (performed on a Renishaw InVia Raman Spectrometer, CLS52-100, 100 mW, USA) allowed determination of the chemical structure and any modifications resulted from the acids oxidation of both pristine and acid treated CNTs. Briefly, CNTs deposited on glass slides (Fisher, USA) were excited through a 20× microscope objective using an Argon ion (Ar+) laser beam with a spot size of <0.01 mm² operating at 514.5 nm. Detailed scans were taken in the 100–3200 cm⁻¹ range; low laser energy (i.e., <0.5 mV) and exposure time of 10 s were used to prevent unexpected heating effects.

2.5. CNTs solubility measurement

The solubility of CNTs (pristine and acids oxidized) was evaluated in di water (pH 6.25) and Phosphate Saline Buffer (PBS, pH 7, 100 mM ionic strength). Briefly, CNTs were diluted in the solvent of interest to yield to a 3 mg/ml solution. The suspension was then centrifuged at 3000 rpm for 5 min; subsequently, part of the supernatant (0.8 ml) was removed and filtered through a 0.2 μm GF filter membrane. The filter membrane was then dried under vacuum and the amount of CNTs was weighted. The solubility of the CNTs was calculated based on the volume used for suspension and the initial starting amount.

2.6. CNTs length measurement

An atomic force microscope (AFM, Asylum Research, USA) was used to evaluate the length of pristine and acids treated CNTs. A Si tip (Asylum Research, 50–90 kHz AC240TS, USA) helped perform tapping mode in air. CNTs samples (i.e., pristine, 1, 3 or 6 h acids oxidized SW and MWCNTs) were dispersed in di water (to yield solutions of 0.1 mg/ml concentration), deposited on mica surfaces (9.5 mm diameter, 0.15–0.21 mm thickness, Electron Microscopy Sciences, USA) and allowed to dry overnight under vacuum. Scan images of 10, 5 or 1 (μm × μm) areas were acquired. For each sample, at least 30 individual CNTs were counted and measured to obtain average length distribution.

2.7. Cell culture and treatment with CNTs

Non-tumorigenic human bronchial epithelial cells (BEAS-2B) were purchased from American Type Culture Collection (ATCC, USA). The cells were cultured in DMEM medium supplemented with 5% fetal bovine serum (FBS), 2 mM l-glutamine and 100 units/ml penicillin/streptomycin (all reagents were purchased from Invitrogen, USA). Cells were passaged weekly using 0.05% trypsin (Invitrogen, USA) and kept in 5% CO₂ at 37 °C. Pristine and acids oxidized SWCNTs were dispersed in di water by sonication, filtered through the 0.2 μm GF filter membrane, resuspended in cellular media and sonicated at room temperature to form stable dispersions. For treatment, BEAS-2B cells were seeded overnight in a 12 well plates (Fisher, USA) at a density of 3.5E5 cells/well, and allowed to reach confluence. Subsequently, the cells were exposed to 100 μg/ml SWCNTs; 24 h post exposure, the cells were incubated with 6.5 μg/ml Hoechst 33342 dye (Molecular Probes, USA) for 30 min at 37 °C and analyzed by microscopy by scoring the percentage of cells with intensely condensed chromatin and/or fragmented nuclei using fluorescence microscopy (Leica Microsystems, USA). Approximately 1000 cell nuclei from ten random fields were analyzed for each sample. The apoptotic index was calculated as the percentage of cells with apoptotic nuclei relative to the total number of cells. At least 3 independent trials were performed for each sample.

2.8. Functionalization of CNTs with enzyme

Soybean peroxidase (SBP, Biossearch, USA) was covalently attached to 1, 3 or 6 h acid treated MWCNTs using 1-ethyl-3-[3-dimethylaminopropyl] carbodiimide hydrochloride (EDC; Acros Organics, USA) and N-hydroxysuccinimide (NHS, Pierce, USA) [32]. Briefly, 2 mg CNTs (MWCNTs) were dispersed in 160 mM EDC and 80 mM NHS (total volume of 2 ml in MES (2-(N-morpholino)ethanesulfonic acid sodium salt, 50 mM, pH 4.7, Sigma, USA) for 15 min at room temperature with shaking at 200 rpm. The activated MWCNTs were next filtered through the 0.2 μm GF filter membrane, washed thoroughly with MES buffer to remove any ester residues, immediately dispersed in 2 ml of 1 mg/ml SBP
solution in PBS (100 mM, pH 7.0) and incubated for 3 h at room temperature at room temperature with shaking at 200 rpm. The resulting SBP–MWCNT conjugates were filtered and washed extensively with PBS to remove any unbound enzyme [32]. The supernatants and washes were collected to quantify enzyme loading.

2.9. Enzyme loading

The amount of SBP attached to MWCNTs (i.e., SBP loading) was determined using standard BCA assay kit (Pierce, USA) and subtracting the amount of enzyme washed out in the supernatant and washes from the amount of SBP initially added to the MWCNTs. Briefly, the working reagent (1000 μl) was prepared by mixing 50 parts of reagent A with 1 part of reagent B (the reagents are provided with the kit). The mixture of reagents A and B was further added to 50 μl solutions of SBP-containing samples (i.e., the samples isolated in the form of the supernatant and washes). The resulting solutions were incubated at 37 °C for 30 min. Absorbance at 562 nm was determined on a spectrophotometer (Fisher, USA). Control calibration curves were prepared by serial dilutions of SBP (free in solution) into the working reagent.

2.10. Enzyme activity assay

The activity of SBP was measured by monitoring the oxidation reaction of (2,2’-Azinobis [3-ethylbenzothiazoline-6-sulfonic acid]) (ABTS, Sigma, USA) in the presence of hydrogen peroxide (H₂O₂, Sigma, USA). 20 μl of the SBP–MWCNTs conjugates were added to 0.65 ml ABTS solution (0.5 mM final concentration, Pierce, USA) and mixed; subsequently, 20 μl H₂O₂ solution (0.2 mM final concentration) was added to the sample in order to initiate the reaction. The change in absorbance was monitored spectrophotometrically at 412 nm immediately upon addition of H₂O₂. The initial reaction rate was calculated from the slope of the linear time-course. The extinction coefficient of the oxidized ABTS product is 32,400 M⁻¹ cm⁻¹ at 412 nm [33]. The activity of the immobilized enzyme is reported as specific activity relative to free enzyme activity. The activity of the free enzyme was determined using an equivalent amount of free enzyme (based on loading data) and the protocol provided above.

2.11. Statistical analysis

All results are presented as mean ± standard deviation.

3. Results and discussion

We prepared a library of single- and multi-walled carbon nanotubes (SW- and MWCNTs) using liquid phase oxidation with a strong nitric and sulfuric acids mixture [6,7]. The approach is shown in Scheme 1; sonication in the acids mixture attacks the graphene sheets on the C–C bands [34], introduces defects and oxidizes the CNTs at the defect sites leading to shorter nanotubes. To reduce the reaction rate of acids attack, the water bath sonicator was maintained at room temperature. The carboxylic acidic groups introduced in SW- and MWCNTs were determined previously using acid–base titrations [35,36] or the formation of a dodecyamine zwitterions [37].

We further investigated the chemical composition of pristine and acids oxidized CNTs using energy dispersive X-ray analysis (EDX) [20,38]. EDX spectra of pristine SW- and MWCNTs are shown in Fig. 1a and b, respectively, as a plot of X-ray counts vs. energy (in keV). The analysis revealed the presence of high contents of carbon (C) and oxygen (O), with iron (Fe) as metal catalyst in both pristine SW- and MWCNTs samples. The energy peaks correspond to the various elements in the sample, with Fe yielding two peaks at 7.07 keV and 6.40 keV [39]. Other elements (e.g., Al, Si, Cl, S, etc.) were also present but in very low amount. The Fe peak was larger for the SWCNTs sample when compared to the MWCNTs one. The difference was reflective of their pristine characteristics since SWCNTs purity was 85% while the purity of pristine MWCNTs was 95%, per manufacturer information (see Section 2). The insets in Fig. 1 show the changes in the O and Fe contents with the acids oxidation treatment time for both SW- and MWCNTs samples. As shown, Fe content decreased with the treatment time for both SW- and MWCNTs samples indicating removal of the metal catalyst. The decrease in the Fe content was more pronounced for the SWCNTs when compared to MWCNTs samples. This is a reflection of the different purities of the two samples chosen in these experiments. For the O content, the change was also dependent on the sample characteristics. The relative low purity SWCNTs samples contain more amorphous carbon [40] than the higher purity MWCNTs [41]. Thus, the acids treatment led to a significant increase of the O content with the acids treatment time for the SWCNTs (Fig. 1a, inset) while compared to a smaller increase for the MWCNTs samples.

Fig. 2 shows the SEM images of the pristine and acids treated samples (both SW- and MWCNTs). As shown, user-controlled acids treatment did not lead to significant morphological changes either for SW- (Fig. 2a) or SWCNTs while Fig. 2c shows 6 h acids treated SWCNTs or MWCNTs (Fig. 2c) shows pristine MWCNTs while Fig. 2d shows 6 h treated MWCNTs samples.

The structural changes upon acids treatment of the CNTs samples were investigated using Raman resonance spectroscopy [42–44]. Fig. 3 shows the Raman spectra of pristine and acids treated SW- and MWCNTs. The Raman analysis of the SWCNTs reveals the presence of 4 bands (Fig. 3a), the so-called D (disorder mode) band around 1340 cm⁻¹, G⁻ and G' bands at around 1545 cm⁻¹ and 1590 cm⁻¹ respectively, and G' band at 2650 cm⁻¹ [22,45]. The Raman analysis of the MWCNTs also shows the presence of 4 bands (Fig. 3b), with the D band around 1340 cm⁻¹, G band at 1585 cm⁻¹, G' band at 2650 cm⁻¹, and another band at 2920 cm⁻¹ [46,47]. The D band around 1340 cm⁻¹ is related to the non-crystalline C species, i.e., defects in the CNTs [48], while the G band observed around 1585 cm⁻¹ is indicative of a high degree of ordering and well-structured C-based structures [42]. The size of the D band relative to the G band can be used as a qualitative measurement for the formation of undesired forms of C [49]. Both pristine and acids treated CNTs (SW- and MWCNTs) have a relatively small D band at around 1350 cm⁻¹, with the D band being wider and shifted toward higher frequency in the acids treated CNTs.

Scheme 1. Time-dependent incubation of pristine CNTs (SW- and MWCNTs) with a mixture of sulfuric and nitric acids leads to acids oxidized CNTs.
samples when compared with the pristine ones. The ratio of intensity of D peak relative to the G peak represents the degree of CNTs functionalization [49]. Higher \( I_D/I_G \) ratio suggests higher level of functionalization (\( I \) represents the peak’s relative intensity).

D band, G band and \( I_D/I_G \) ratio of the various CNTs samples (both SW- and MWCNTs) are shown in Table 1. The ratio of \( I_D/I_G \) for SWCNTs changed minimally from 0.237 for pristine to 0.263 after 6 h acid treatment. For 1 and 3 h acid oxidized SWCNTs, the \( I_D/I_G \) ratio seemed to have decreased. Previous reports have shown that for relatively low purity CNTs (in this particular example the SWCNT’s purity is 85%; see Section 2) the \( I_D/I_G \) does not provide precise overall information on the sample structure [50], and the \( I_D/I_G \) ratio might be both a reflection of washing away amorphous carbon while simultaneously inducing carboxylic acid groups [20].

For instance, in the initial 1 h SWCNTs acids oxidation, the effect of washing away amorphous C (which is known to lead to decreased \( I_D/I_G \) [51]) suppressed the effect of adding carboxylic acid groups (which is known to lead to increased \( I_D/I_G \) [52]). However, after 6 h, most of the amorphous C was removed and the \( I_D/I_G \) became indicative only of the degree of functionalization with carboxylic groups.

\( I_D/I_G \) for MWCNTs increased from 0.457 for pristine to 0.788 for 3 h, and 0.796 after 6 h acids oxidation. This increase in the level of functionalization has a similar trend to the increase in the O or decrease in the Fe catalyst content as observed through the EDX analyses (Fig. 1). Specifically, for the high purity MWCNTs most of the Fe catalysts are removed during the 3 h treatment time (see inset Fig. 1b) this leading to removal of the defects in the MWCNTs structure. Since defects are where the promotion of the carboxylic groups formation takes place [53], and since for the MWCNTs there was a small decrease in the Fe and a small increase in the O content (Fig. 1b inset) from the 3 h to 6 h treatment time, the \( I_D/I_G \) for MWCNTs will be minimally changed between these time points as indicated in Table 1. Such analyses confirm that the acids oxidation introduced CNTs chemical property changes i.e., added functional free carboxylic acid groups, to both SW- and MWCNTs sample.

We further investigated how the degree of CNTs dispersion in water-based environments is influenced by the acids oxidation time. We used two solvents with different pHs and ionic strengths, i.e., di water (pH 6.25) and Phosphate Saline Buffer (PBS, pH 7, 100 mM). The results (Fig. 4) indicated that the solubility of CNTs

![Fig. 1. EDX elemental analysis of pristine SWCNTs (a) and MWCNTs (b). The insets show the changes in the O and Fe contents with the acids treatment time employed under user-control.](image-url)
in both di water and PBS was improved upon the acids oxidation, with increased acids oxidation times leading to increased solubility. Generally, pristine and acid oxidized SWCNTs (either 1, 3 or 6 h cut) were more dispersed in PBS when compared to di water (Fig. 4a). MWCNTs did not show a similar trend; specifically, pristine and 1 h cut MWCNTs were more soluble in PBS, however, after longer acids oxidation times (i.e., 3 and 6 h) the solubility was higher in water when compared to PBS (Fig. 4b). The changes in the solubility observed for the MWCNTs samples after longer acids oxidation times are correlated with the changes in the functionality of these samples and number of carboxylic acidic groups being generated. Specifically, longer acids oxidation times will lead to higher number of carboxyl groups being generated (see Figs. 1 and 3).

When the MWCNTs acids treated samples are placed in water-based environments, carboxylate anions groups are generated by the deprotonation of carboxylic acid groups [54]. At high ionic strength, the probability for these anions to form aggregates [55] increases thus leading to the lower solubility observed for the 3 and 6 h acids oxidized MWCNTs placed in PBS when compared to solubility of these samples placed in water.

Atomic force microscopy (AFM) and tapping mode [56] was used to analyze the morphology and quantify the length of the CNTs samples. Specifically, cross sectional areas from $(10 \times 10)$ to $(1 \times 1) \mu m \times \mu m$ were scanned to derive the length of at least 30 CNTs/sample (both SW- and MWCNTs; pristine, 1, 3 and 6 h cut). Pristine and acids oxidized CNTs length distributions are shown in

![Fig. 2. SEM image of (a) pristine SWCNTs, (b) pristine MWCNTs and (c) 6 h acids treated SWCNTs (d) 6 h acids treated MWCNTs; the scale bar is 1 μm.](image)

![Fig. 3. Raman spectra of pristine, 1, 3 and 6 h acids oxidized SWCNTs (a) and MWCNTs (b).](image)
Having established that the acids oxidation influences the chemical and physical properties of pristine CNTs (both SW- and MWCNTs), we proceeded to examine whether user-controlled acids oxidation would also affect CNTs biocompatibility. First, we performed a systematic study on the cellular toxicity resulted from the incubation of immortalized human bronchial epithelial cells with acids oxidized SWCNTs. Previous in vivo studies have shown that cellular exposure to SWCNTs results in macrophages without nuclei [57,58], with SWCNTs inducing chromosome aberration [18]. However, to our knowledge, no studies that looked at the influence of the different acids oxidation times to BEAS-2B immortalized human bronchial epithelial cells have been performed. Moreover, to our knowledge, there is no correlation in the literature on how cellular toxicity depends on the SWCNTs physical and chemical properties as impaired by the acids oxidation time and how such toxicity can be controlled. In our experiments, BEAS-2B cells were exposed to SWCNTs for 24–72 h at Permissible Exposure Limit for particulates not otherwise regulated (i.e., 100 µg/ml of SWCNTs, based on previous laboratory exposure levels [58,59]). Fig. 6 shows the percentage of apoptotic BEAS-2B cells upon exposure to SWCNTs; our data shows that the cytotoxicity of the 6 h acids treated SWCNTs is lower than that of pristine SWCNTs. Specifically, the percentage of apoptotic cells for pristine SWCNTs is about 19% while the percentage of apoptotic cells for 6 h acids treated SWCNTs is about 15% upon 72 h incubation. These results are comparable to control cells (cells that have not been exposed to SWCNTs) and they emphasize that user-controlled acids oxidation time can be employed to create a library of sample of SWCNTs that have high biocompatibility with cellular system. We hypothesized that the observed trend is due to the changes in the chemical and physical structure of the SWCNTs upon acid functionalization. Specifically, shorter and more hydrophilic SWCNTs (see our previous EDX and AFM results) would be predominantly taken up by the cells through endocytosis [60], while for the longer SWCNTs the uptake mechanism is predominantly through piercing [61]. Furthermore, the longer SWCNTs once taken up by the cells can localize at the cell nucleus and interfere with the normal progression of cells to

**Fig. 4.** Solubility of pristine and acids oxidized SWCNTs (a) and MWCNTs (b) in deionized (di) water and phosphate buffer saline (PBS).

**Fig. 5.** The average length distribution and the standard deviation of SWCNTs (a) and MWCNTs (b) with the acids treatment time.

**Fig. 6.** Cytotoxicity of pristine and 6 h acids treated SWCNTs to BEAS-2B human epithelial cells after 24, 48 and 72 h, respectively.
Table 2
Loading and retained specific activity of immobilized SBP onto acids treated MWCNTs.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Loading (mg SBP/mg MWCNTs)</th>
<th>Retained specific activity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 h cut covalent</td>
<td>0.254 ± 0.05</td>
<td>9.40 ± 1.68</td>
</tr>
<tr>
<td>3 h cut covalent</td>
<td>0.282 ± 0.06</td>
<td>28.18 ± 6.52</td>
</tr>
<tr>
<td>6 h cut covalent</td>
<td>0.265 ± 0.15</td>
<td>33.97 ± 9.82</td>
</tr>
</tbody>
</table>

4. Conclusion

Our results have shown that user-controlled acid oxidation of CNTs led to the formation of a library of samples with different physical and chemical properties. Specifically, we have shown that CNTs oxidation with a nitric and sulfuric acids mixture results in removal of metal catalyst, an increase in the number of functional groups having electron accepting ability, and generation of shorter CNTs with higher solubility in aqueous environments. Our results were confirmed by Raman spectroscopy, SEM, AFM, EDX and solubility tests. Further, we have shown that CNTs acids oxidation improves nanotube biocompatibility as tested by direct incubation with human epithelial cells or with test enzymes. User-controlled design of CNTs biocompatibility can lead to new types of analytical tools for life science and biotechnology [75–77].

Acknowledgements

This work is support by the NSF/CBET 1033266 and NSF-EPS-1003907. The authors acknowledge Adrienne McGraw, Chemical Engineering/WVU for her help with EDX/SEM analysis and Dr. Weiqiang Ding/WVNano for his help with Raman analysis. Authors acknowledge use of the WVU Shared Research Facilities.

References

H.

268

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walled

Defects

multi-walled

nanotubes,


Perhydrolase-nanotube-paint sporidical composites stabilized by intramolecular crosslinking

Cerasela Zoica Dinu a,*,1, Indrakant V. Borkar b, c, 1, Shyam Sundhar Bale b, c, Alan S. Campbell a, Ravi S. Kane b, c, Jonathan S. Dordick b, c, **

a Department of Chemical Engineering, College of Engineering and Mineral Resources, West Virginia University, Morgantown, WV 26506, USA
b Department of Chemical and Biological Engineering, Rensselaer Polytechnic Institute, Troy, NY 12180, USA
c Center for Biotechnology & Interdisciplinary Studies, Rensselaer Polytechnic Institute, Troy, NY 12180, USA

ABBREVIATIONS
AcT, isolated from Mycobacterium smegmatis, is a homo-octamer of 184 kDa with 72 × 72 × 60 Å dimensions [1] that effectively catalyzes the perhydrolysis of propylene glycol diacetate (PGD) to generate peracetic acid (PAA) [Scheme 1], a potent decontaminant effective against bacteria, yeasts, fungi, and spores [2–4]. AcT has a perhydrolysis to hydrolysis ratio greater than 1 and an activity 50-fold higher than that of the best lipase tested [5,6]. This makes AcT a potentially valuable biocatalyst for decontaminating various surfaces if highly active and stable enzyme-based surface formulations could be developed. Nonetheless, identifying methods to improve enzyme activity and stability, particularly upon extended exposure to the PAA product remains a formidable challenge. Moreover, the large size of AcT coupled to its relatively high degree of surface hydrophobicity may also limit its long-term use [1,6].

We have developed a strategy to preserve the activity and operational stability of a large multi-subunit enzyme immobilized onto carbon nanotubes and incorporated into latex paint. Our strategy involved the intramolecular crosslinking of perhydrolase SS4V (AcT, a homo-octamer) and the subsequent immobilization of the crosslinked AcT onto single-walled carbon nanotubes (SWNTs). We employed aldehyde dextran – a bulky polymeric aldehyde obtained by oxidation of dextran with sodium metaperiodate – as a crosslinking reagent. The activity of AcT crosslinked with aldehyde dextran and covalently attached to SWNTs (AcT-dex-SWNTs) was ~40% of that of native AcT and more than two-fold higher than that of enzyme immobilized directly, i.e., without crosslinking. This relatively high retention of AcT activity was consistent with the nearly complete retention of the enzyme’s secondary structure upon attachment to the nanoscale support. Further incorporation of the AcT-dex-SWNTs conjugates into a latex-based paint led to active composites that were used to decontaminate Bacillus spores.

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1. Introduction

AcT, isolated from Mycobacterium smegmatis, is a homo-octamer of 184 kDa with 72 × 72 × 60 Å dimensions [1] that effectively catalyzes the perhydrolysis of propylene glycol diacetate (PGD) to generate peracetic acid (PAA) [Scheme 1], a potent decontaminant effective against bacteria, yeasts, fungi, and spores [2–4]. AcT has a perhydrolysis to hydrolysis ratio greater than 1 and an activity 50-fold higher than that of the best lipase tested [5,6]. This makes AcT a potentially valuable biocatalyst for decontaminating various surfaces if highly active and stable enzyme-based surface formulations could be developed. Nonetheless, identifying methods to improve enzyme activity and stability, particularly upon extended exposure to the PAA product remains a formidable challenge. Moreover, the large size of AcT coupled to its relatively high degree of surface hydrophobicity may also limit its long-term use [1,6].

We have focused on stabilizing enzymes by attachment, both covalently and non-covalently, onto carbon nanotubes. Indeed, in the case of AcT, we have begun to test the biological decontamination of Bacillus cereus, a simulant of Bacillus anthracis, by AcT immobilized onto multi-walled carbon nanotube (MWNTs) and incorporated into polymeric films and paint composites [1]. Carbon nanotubes have excellent support characteristics due to their high surface area to volume ratios that allow relatively high enzyme loadings [7–11], ease of recovery by filtration for the enzyme-nanotube conjugate [1], and high aspect ratios that result in entrapment of the support in coatings, films and paints, thereby preventing leaching of any attached biocatalyst from the surface [1]. Critically, we hypothesize that the operational and thermal stability of a large multi-subunit enzyme such as AcT may be increased on a surface if greater rigidity of the enzyme were induced, for example, via crosslinking prior to attachment onto the nanoscale support. Along these lines, homo-bi- or poly-functional aldehydes, such as glutaraldehyde or aldehyde dextran, respectively, have been used to crosslink multimeric enzymes [12], thus increasing the rigidity of the enzyme and avoiding the formation of non-specific protein–protein associations.

* Corresponding author at: Department of Chemical Engineering, West Virginia University, College of Engineering and Mineral Resources, PO Box 6102, ESF 445, Morgantown, WV 26506, USA. Tel.: +1 304 293 9338; fax: +1 304 293 4139.
** Corresponding author at: Department of Chemical and Biological Engineering, Department of Biology, 2213 Center for Biotechnology Interdisciplinary Studies, Rensselaer Polytechnic Institute, 110 8th Street, Troy, NY 12180, USA. Tel.: +1 518 276 2899; fax: +1 518 276 2207.
E-mail addresses: cerasela-zoica.dinu@mail.wvu.edu (C.Z. Dinu), dordick@rpi.edu (J.S. Dordick).

1 These authors contributed equally to this work.

1381-1177/5 – see front matter © 2011 Elsevier B.V. All rights reserved. doi:10.1016/j.molcatb.2011.11.003
In the current work we sought to improve AcT activity and stability when attached to single-walled carbon nanotubes (SWNTs). SWNTs were chosen as supports, as their higher surface curvature when compared with MWNTs, is expected to reduce lateral interactions between adjacent protein molecules adsorbed, leading to greater retention of native protein structure and activity. Specifically, we show that by crosslinking AcT with the polyfunctional aldehyde dextran followed by covalent attachment onto SWNTs, we can dramatically improve enzyme operational stability, thermostability, and activity of the resulting conjugates. Further incorporation of the conjugates into paint led to composites that showed complete decontamination of $10^6$ Colony Forming Units (CFU)/mL of B. cereus spores in 60 min.

2. Results and discussion

Biological decontamination is aimed at eliminating biological hazards associated with pathogens infecting solid surfaces in laboratories [13], pilot plants [14,15], hospitals [16] or battlefield settings [17]. Decontamination involves rapid removal or neutralization of the pathogen using physical and/or chemical methods. An ideal decontaminant must be non-corrosive, non-toxic, and environmentally friendly. Enzymatic decontamination methods possess such ideal properties, since they are biodegradable, safe, easy to use, environmentally benign, and effective in low-volume doses [1,17,18]. AcT-catalyzed synthesis of PAA (Scheme 1) represents an excellent example of one such biologically driven decontamination, and thus, serves as a practical model system in this study.

Following our previous methodology [1], AcT was initially covalently attached to acid–oxidized SWNTs, which possessed carboxylic acid “handles” ideal for EDC/NHS coupling chemistry (Scheme 2a). The hydrophilic carboxyl moieties also increased nanotube dispersion and solubility [1]. Covalent attachment of AcT onto SWNTs led to enzyme loadings of $0.15 \pm 0.01$ mg AcT per mg SWNTs. However, AcT-SWNT conjugates retained <20% of the native solution specific activity of AcT (Fig. 1a). In contrast, when MWNTs were used as supports for AcT covalent attachment, only 8% specific activity was retained at a similar loading as that for SWNTs (see Supporting materials). This is in agreement with previous reports that show that enzyme structure and function when bound to SWNTs or silica nanoparticles are more native-like than when bound to flatter surfaces [1,19]. The hydrophobic nature of AcT (aliphatic index of 95.66, grand average hydrophaticity (GRAVY) of 0.117 based on computational analysis [1]), may also lead to non-specific (and potentially unfavorable) hydrophobic interactions between the enzyme and non-functionalized hydrophobic regions of the SWNTs and MWNTs. Such interactions could be strong enough to alter enzyme structure and reduce catalytic activity. Encouraged by the higher activity retained by AcT immobilized on the SWNTs, we proceeded to use these supports in this work.

Chemical modification of proteins with crosslinkers is known to reduce non-specific interactions [12,20–22]; however, to our knowledge no work has been performed on crosslinking enzymes followed by attachment onto nanoscale supports. To that end, we performed light crosslinking of AcT with 0.25 and 0.50% (w/w) glutaraldehyde. In both cases, the relatively light crosslinking did not result in inter-enzyme molecular linkages, as SDS-PAGE gels showed a band at roughly the same molecular weight as free AcT (Scheme 2b). No higher molecular weight bands were observed. The activity of the crosslinked AcT (AcT-glu) was ~34% of the free
enzyme activity (Fig. 1a). We rationalized that the loss in activity may have been due to reaction of glutaraldehyde with key amino acid residues near the active site of the enzyme and/or because the small size of glutaraldehyde blocks the active site channel of AcT [12, 22]. To overcome either of these deleterious outcomes, we performed light crosslinking with the polymeric aldehyde dextran prepared by oxidizing dextran (20kDa) with sodium periodate. As with glutaraldehyde, light crosslinking with 0.25 and 0.50% (w/w) dextran aldehyde resulted in exclusively intramolecular AcT crosslinks (Scheme 2b). The somewhat diffuse band with increased molecular weight was due to the polydispersity of the dextran aldehyde. In no case did we observe molecular weights of 2× or 3× of native AcT, indicating that only intramolecular crosslinking had occurred. The activity of the crosslinked Act-dex was ~52% of that of free AcT (Fig. 1a). The relatively high retention of catalytic activity of the AcT-dex would suggest that the enzyme’s secondary structure remained intact. Indeed, comparison of the circular dichroism (CD) spectrum of AcT-dex to that of native AcT (Fig. 1b) revealed that the enzyme retained the majority of its secondary structure, specifically ~93% of the native AcT α-helix content. Act-glu, however, was
structurally perturbed (Fig. 1b) with only 67% secondary structure retention even at 0.25% glutaraldehyde. This result is consistent with greater loss of activity of AcT-glu vs. AcT-dex.

To assess whether the pre-immobilization crosslinking-induced AcT stabilization by aldehyde dextran could carry over to AcT-based conjugates, we used EDC/NHS coupling to attach both glutaraldehyde- (as a comparison) and aldehyde dextran-crosslinked AcT to SWNTs in a methodology similar to that shown in Scheme 2a. AcT-glu-SWNT conjugates retained ~19% of native AcT activity, while AcT-dex-SWNT conjugates retained ~40% of native AcT activity (Fig. 1a) and nearly 80% of the activity of the AcT-dex pre-immobilized crosslinked enzyme form. In another strategy, attachment of crosslinked AcT to SWNTs was performed using an amphiphilic poly(ethylene glycol) (PEG) linker [1]. Theoretically, this type of linker could improve enzyme activity by reducing non-specific binding between the enzyme and the nanosupport, as well as it can provide more favorable orientation of the protein at the nanoscale surface [1,17]. However, no improvement in enzyme activity was observed for AcT attached to the SWNTs via the PEG linker (see Supporting material). Thus, crosslinking with the polymeric aldehyde dextran stabilizes AcT against nanotube facilitated protein deactivation.

The kinetics of AcT-dex and AcT-dex-SWNT (2 μg free or equivalent of immobilized enzyme) was studied by measuring the initial reaction rates at different substrate concentrations. The concentration of hydrogen peroxide was varied from 0.1 mM to 428 mM while the concentration of PGD was kept constant at 200 mM. AcT-dex and AcT-dex-SWNT both followed Michaelis–Menten kinetics as a function of H₂O₂ concentration (Fig. 2) with fairly similar (Vₘₐₓ/Kₘ) values (Table 1) and ca. 25–33% of that of the native enzyme, as a result of roughly equal contributions of lower Vₘₐₓ and higher Kₘ. Importantly, the relatively minor changes in both Vₘₐₓ and Kₘ indicate that the enzyme retained its intrinsic function following both crosslinking with aldehyde dextran and attachment to SWNTs.

The combination of retained secondary structure and high activity as a result of AcT-dex crosslinking indicates that the polymeric aldehyde is restricted from gaining access to the active site and is likely causing rigidification of the multi-subunit enzyme thus leading to maintaining activity on the heterogeneous SWNT. Such rigidification would be expected to stabilize the enzyme under harsh conditions, for example, high temperature. To test this stabilization, we incubated AcT, AcT-dex, and AcT-dex-SWNTs at 75 °C for up to 2 h. While the free enzyme lost nearly 80% of its activity at 75 °C after 2 h, AcT-dex and AcT-dex-SWNTs retained more than 65% activity under these conditions (Fig. 3a). Moreover, free AcT and AcT-dex followed second order thermal deactivation (Fig. 3b and c). A typical second order thermal deactivation model is reflected

![Figure 2](image_url)

**Fig. 2.** Kinetics of parameters of free Act (filled diamond), Act crosslinked with aldehyde dextran (filled squares), and Act crosslinked with aldehyde dextran and immobilized onto SWNTs (filled triangle). The concentration of hydrogen peroxide was varied from 0.1 mM to 428 mM while the concentration of PGD was kept constant at 200 mM. At least 5 replicates were performed.

<table>
<thead>
<tr>
<th></th>
<th>Vₘₐₓ (mM/min)</th>
<th>Kₘ (mM)</th>
<th>Vₘₐₓ/Kₘ (min⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Free Act</td>
<td>2.08</td>
<td>55</td>
<td>0.038</td>
</tr>
<tr>
<td>AcT-Dex</td>
<td>1.12</td>
<td>88</td>
<td>0.0125</td>
</tr>
<tr>
<td>AcT-Dex-SWNTs</td>
<td>1.02</td>
<td>110</td>
<td>0.0095</td>
</tr>
</tbody>
</table>

* The values represent an average of measurements with standard error <7%.
in Eq. (1), where A is the residual AcT activity and \( k_d \) is the second order deactivation rate constant.

\[
\frac{dA}{dt} = -k_d A^2 \tag{1}
\]

Integration of Eq. (1) within limits leads to Eq. (2) where \( A_0 \) is the initial activity of AcT and \( X \) is the fraction of enzyme deactivated.

\[
\frac{X}{1-X} = A_0 k_d t \tag{2}
\]

Plotting \( X/(1-X) \) versus time confirmed second-order deactivation kinetics (Fig. 3c) and allowed determination of \( A_0 k_d \) values. Half-life times (\( t_{1/2} \)) at 75 °C were calculated using Eq. (3), which gave values of 33 and 313 min for free AcT and AcT-dex, respectively.

\[
t_{1/2} = \frac{1}{A_0 k_d} \tag{3}
\]

These results are consistent with the primary deactivation of native AcT at elevated temperature being due to aggregation. AcT-dex-SWNTs do not appear to follow second order thermal deactivation (Fig. 3b), consistent with a restricted rotational degree of freedom resulting from its multi-point attachment to the nanoscale support. Further stabilization from aggregation may be due to decreased protein–protein interactions on the highly curved surface of the SWNTs [23]. In addition to thermal stability, we also studied the operational stability of AcT-dex-SWNT incubated at 4 °C for up to 180 days. No loss of activity was observed under these typical storage conditions, further confirming the high stability afforded by crosslinking of the enzyme with aldehyde dextran and attachment to the SWNT.

Motivated by the increased stability of the crosslinked enzyme bound to SWNTs, we proceeded to incorporate the crosslinked-Conjugates into a latex paint (Scheme 2c) to form nanocomposites that can decontaminate B. cereus spores [24–29]. The high aspect ratio of the SWNTs allowed retention of the enzyme within the composite with no enzyme leaching being observed after 15 days incubation in buffer [17]. As a control, when free AcT or AcT-dex was directly added to the latex solution and dried (i.e., no nanotubes), nearly 50% of the enzyme leached out from that paint within the first 30 min. When challenged with 10^6 CFU/mL, AcT-dex-SWNT paints (containing 0.04%, w/w, AcT) incubated in a solution containing 100 mM each of PGD and H_2O_2 generated ca. 10 mM PAA and killed >99% of the spores within 1 h (Fig. 4). Specifically, in only 15 min we achieved approximately 60% spore killing, which is in agreement with previous literature reports for direct PAA addition [30–32]. Moreover, when the composites were challenged with 10^6 CFU/mL of a simple non-spore forming bacterium, E. coli, the killing time was reduced to 5 min.

3. Conclusions

We have shown that the use of aldehyde dextran as a crosslinking agent stabilizes the native structure of AcT. The enzyme retains high activity (>40% of native aqueous solution activity) and can be incorporated into paint-based composites that have decontamination properties against B. cereus (a simulant of B. anthracis) and E. coli. Further work is underway to assess composite stability and activity in a wide variety of conditions (i.e., temperature, humidity, various paint compositions and polymers, etc.). The capability of generating sufficiently high concentrations of PAA makes these composites particularly useful as surface coatings for the disinfection of a wide range of pathogenic agents including bacteria and spores.

4. Experimental

4.1. AcT crosslinking with glutaraldehyde

Perhydrolase S54 (AcT, 1 mg/mL, gift from Genencor International, Inc. Palo Alto, CA, USA) was incubated at 4 °C with 0.25 and 0.50% of glutaraldehyde (Sigma, USA), respectively, for 24 h. Thus crosslinked AcT was treated with 10% (v/v) Tris–HCl buffer (1 M, pH 8, Sigma, USA). Excess reactants were removed by extensive dialysis against water at 4 °C for 24 h [33,34].

4.2. AcT crosslinking with aldehyde dextran

Aldehyde dextran was obtained by fully oxidizing 50 mL of dextran (20 kDa, 33.3 mg/mL, Sigma, USA) with sodium periodate (4.36 g, Sigma, USA) in distilled water (21, 37). After 2 h incubation in dark, the oxidized dextran was extensively dialyzed against distilled water at 4 °C for 24 h. The purified aldehyde dextran was then lyophilized for long-term storage. AcT (5 mL) in 50 mM sodium phosphate buffer (100 mM, pH 7) was added to 150 mg of aldehyde dextran (final AcT concentration was 1 mg/mL) in the presence of 150 mM trimethylammonioborate (Sigma, USA) for 24 h at 25 °C. Schiff bases, formed between the primary amino groups of the enzyme and the aldehyde groups of the polymer were reduced by the addition of 3 mg/mL sodium borohydride (Sigma, USA) at pH 10 [12]. After 30 min, the pH was decreased to 7 by the addition of HCl. Aldehyde dextran was further treated with 10% Tris–HCl (1 M, pH 8) in order to avoid non-specific crosslinking.

4.3. Crosslinking of the AcT confirmed by gel electrophoresis

Gel electrophoresis was used to confirm enzyme crosslinking. Specifically, NuPAGE® Novex® Tris–Acetate Gels (Native-PAGE) (Invitrogen, USA) of 4–12% gradient, 10-well was loaded with 25 µl sample of 0.01 mg/mL AcT or equivalent of AcT-derivates (i.e., AcT-dex, AcT-glu, etc.) prepared in Novex® Tris–Glycine Native Sample Buffer (Invitrogen, USA). Appropriate unstained molecular weight marker NativeMark™ (Invitrogen, USA) was also used. The gel run at 80 V for 120 min in Novex® Tris–Glycine Native Running Buffer and was stained with polyacrylamide pre-cast gel SimplyBlue™ Coomassie protein stain (Invitrogen, USA), 1 x pre-mixed solution.

4.4. Acid oxidation of carbon nanotubes

SWNTs were purchased from Unidym, Inc. (USA) and oxidized as previously described [1]. Briefly, 100 mg SWNTs were suspended
in 60 mL of 3:1 (v/v) sulfuric acid to nitric acid (H₂SO₄: HNO₃) mixture (Fisher Scientific, USA) and sonicated at room temperature for 6 h. The acid oxidized SWNT suspension was diluted in distilled water and the mixture was filtered through 0.2 μm polycarbonate filter membrane (Millipore, USA). The SWNT “cake” that formed on the filter was resuspended in distilled water by sonication and the filtration step was repeated until water-soluble SWNTs were obtained and any insoluble impurities were removed. The SWNTs were dried under vacuum and stored at room temperature.

4.5. Functionalization of acid oxidized SWNTs with AcT and crosslinked AcT

Free and crosslinked AcT was covalently attached to 6 h oxidized SWNTs using 1-ethyl-3-[3-dimethylaminopropyl] carbodiimide hydrochloride (EDC; Acros Organics, USA) and N-hydroxysuccinimide (NHS, Pierce, USA), respectively (1). Briefly, 2 mg of SWNTs, 6 h oxidized nanotubes) were dispersed in 160 μM EDC and 80 mM NHS (total volume of 2 mL in MES (2-(N-morpholino)ethanesulfonic acid sodium salt, 50 mM, pH 4.7, Sigma) for 15 min at room temperature and 200 rpm. The activated SWNTs were next filtered through the 0.2 μm filter, washed thoroughly with MES buffer to remove any ester residues, and immediately dispersed in 1 mg/mL AcT (free, crosslinked with glutaraldehyde, or crosslinked with aldehyde dextran respectively) solution in phosphate buffer (50 mM, pH 7.0) and incubated for 3 h at room temperature with shaking at 200 rpm. The resulting AcT-based-SWNT conjugates were filtered and washed extensively with buffer to remove any unbound enzymes (1) while the supernatants and washes were collected to quantify enzyme loading.

4.6. Enzyme loading

The amount of AcT attached to SWNTs (i.e., AcT loading) was determined using standard BCA assay (bicinchoninic acid, Pierce, USA) and subtracting the amount of enzyme washed out in the filtrates from the amount of AcT initially added to the SWNTs. Briefly, the working reagent was prepared by mixing 50 parts of reagent A (BCA Protein Assay Reagent A Formulation: Bicinchoninic acid and tartrate in an alkaline carbonate buffer, http://www.piercenet.com/browse.cfm?flID=02020101) with 1 part of reagent B (BCA Protein Assay Reagent B Formulation: 4% copper sulfate pentahydrate solution, http://www.piercenet.com/browse.cfm?flID=02020101); subsequently 200 μL of the working reagent was incubated with 25 μL AcT-based solution. The resulting solution was incubated in 96-well plate with a clear flat bottom (Thermo Scientific, USA) at 37 °C for 30 min. Absorbance at 562 nm was determined on a Microplate reader (SpectraMax M5, Molecular Devices, Sunnyvale, CA, USA). Control calibration curves were prepared using serial dilutions of AcT (free in solution) into the working reagent.

4.7. Activity assay

AcT activity was determined by measuring the peracetic acid (PAA) generated by the free or immobilized enzyme (1). In a typical reaction, 10.6 μL hydrogen peroxide (H₂O₂, 30%, v/v, from Sigma, USA) stock solution was added to a mixture of 0.8 mL propylene glycol diacetate (PGD, final concentration 100 mM in potassium phosphate buffer, 50 mM, pH 7.1, Sigma, USA) and 0.2 mL AcT solution (2 μg/mL final concentration for free AcT or equivalent concentration for AcT for AcT-based conjugates). The mixture was shaken at room temperature and 200 rpm for 20 min. The PAA assay was conducted by diluting 25 μL of reaction solution 100-fold in deionized water and subsequently mixing 25 μL of the diluted solution with 75 μL deionized water and 0.9 mL assay reagent (the assay reagent was prepared by mixing 5 mL potassium citrate buffer, 125 mM, pH 5.0 with 50 μL ABTS water solution, 100 mM, and 10 μL KI water solution, 25 mM; all the reagents were purchased from Sigma, USA). The mixture was then incubated at room temperature for 3 min and the absorbance at 420 nm was measured on a UV-vis spectrophotometer. PAA concentration was calculated as [PAA] (mM) = A₄₂₀nm × 0.242 × 400 (400 is the dilution factor). The specific activity of AcT-based conjugates was calculated as the ratio of the normalized activity of the conjugates to that of the native AcT.

4.8. Circular dichroism of AcT, crosslinked AcT and AcT-based conjugates

Circular dichroism (CD) analysis was performed using a Jasco 815 Circular dichroism Spectrometer (Jasco Analytical Instrument, Inc., Easton, MD, USA). Free AcT, crosslinked AcT, and AcT-based conjugate samples were dispersed in phosphate buffer to a final concentration of ca. 10 μg/mL AcT. The CD data were collected in the range of 205–260 nm and the molar ellipticity, θ, was calculated using equation (4) where molecular weight of AcT is 184 kDa, the number of amino acids is 216 (PDB-2Q0S) and the cuvette path length is 1 cm.

[θ] = \frac{\text{Molecular weight (kDa)}}{\text{No. amino acids}} \times \frac{10 \times \text{path length (cm)}}{\text{[conc](mg/mL)}} \times \text{[conc](mg/mL)}}

Subsequently, the residual structure of the protein in each of the cases was calculated using Eq. (5) (38).

% α-helix = \frac{-[θ]_{222 \text{ nm}} + 3000}{39, 000}

4.9. Kinetics of AcT, crosslinked AcT and AcT-based conjugates

Kinetics of free AcT, crosslinked AcT and AcT–nanotube conjugates was studied by measuring the initial reaction rates of the samples at different substrate concentrations. Specifically, the concentration of H₂O₂ was varied from 0.1 mM to 428 mM while the PGD concentration was maintained at 200 mM.

4.10. Thermal stability of AcT, crosslinked AcT and AcT-based conjugates

Thermal stabilities of free, crosslinked AcT and AcT-based conjugates were investigated by incubating enzyme-containing solutions in a water bath at 75 °C. Samples were collected periodically, diluted, and the activity was evaluated as previously described. The activities of different enzyme compositions were compared to the free enzyme activity.

4.11. Preparation of spores

B. cereus 4342 was purchased from ATCC (USA) and cultured in nutrient broth (3 g/L beef extract, 5 g/L peptone, Difco, USA) prepared in distilled water for 48 h. The samples were next centrifuged at 3000 rpm for 3 min and sporulation was induced by resuspending the cells in Difco Sporulation Media (DSM) at 37 °C and 200 rpm for 72 h. All reagents were purchased from Sigma, unless otherwise specified. To terminate sporulation, the solution was centrifuged at 3000 rpm for 3 min and the sediment was resuspended in distilled water; the procedure was repeated 5 times. Spore purity was determined by DIC confocal microscopy at 100× magnification (Nikon, NY). The spores were visibly free of germinating cells and spore concentration was estimated using standard plate count technique.
4.12. Sporicidal efficiency of biocatalytic composites

Enzyme-nanotube based composites were prepared as described previously [1,17]. Briefly, water-soluble crosslinked AcT-nanotube conjugates were mixed with eco-friendly paint (Freshaire Choice™, with no volatile organic compounds, from ICI paints, Strongsville, OH, USA) in a glass vial (2.5 cm diameter, VWR, USA). The mixture was air-dried for 2 days; the resulting composite had a thickness of ~450 µm as measured by surface profilometry (Dektak 8 Surface Profiler, Veeco Instruments Inc., Plainview, NY, USA). Decontamination of spores was evaluated by incubating 10^6 CFU/mL spores with the Act-based composite in a reaction mixture containing 100 mM PGD and 100 mM H_2O_2 in 1 mL phosphate buffer (50 mM, pH 7), after shaking for 1 h at room temperature and 200 rpm. Aliquots from this reaction mixture were withdrawn periodically, diluted in phosphate buffer, spread onto a nutrient agar, and incubated at 37°C for 12 h. Sporidal efficiency was determined by counting colonies grown on the agar surface and by comparing corresponding colony counts with those obtained from different controls (paint and paint with reaction mixture without enzyme, respectively).

Acknowledgements

This work was supported by DTRA (HDTRA1-08-1-0022). We thank Gregg Whitey and Karl Sanford (Genencor) for the gift of AcT.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.molcatb.2011.11.003.

References