Production and Characterisation of Self-Crosslinked Chitosan-Carrageenan Polyelectrolyte Complexes

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This dissertation is submitted for the degree of Doctor of Philosophy
Declaration

This dissertation is the original work of the author and does not include any research that is the outcome of work done in collaboration with others, except where specifically indicated in the text and acknowledgements.

The enclosed research was conducted in the Department of Materials Science and Metallurgy, University of Cambridge, under the supervision of Prof. Ruth Cameron and Prof. Serena Best between October 2013 and April 2017. This thesis has not been submitted, nor is in the process of being submitted, for any other qualifications at this or any other institution.

In accordance with the Degree Committee of the Faculty of Physics and Chemistry, this dissertation does not exceed 60,000 words.

Nawar Al-Zebari

April 2017
This thesis is dedicated to my loving parents, Mudhaffar Al-Zebari & Julyana Randquist.
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Lord, thank you for always being there for me.
“The journey of a thousand miles begins with one step.” - Lao Tzu
Abstract

Macromolecular biomaterials often require covalent crosslinking to achieve adequate stability and mechanical strength for their given application. However, the use of auxiliary chemicals may be associated with long-term toxicity in the body. Oppositely-charged polyelectrolytes (PEs) have the advantage that they can self-crosslink electrostatically and those derived from marine organisms are an inexpensive alternative to glycosaminoglycans present in the extracellular matrix of human tissues.

A range of different combinations of PEs and preparation conditions have been reported in the literature. However, although there has been some work on complex formation between chitosan (CS) and carrageenan (CRG), much of the work undertaken has ignored the effect of pH on the consequent physicochemical properties of self-crosslinked polyelectrolyte complex (PEC) gels, films and scaffolds. Chitosan is a positively-charged polysaccharide with NH$_3^+$ side groups derived from shrimp shells and, carrageenan is a negatively-charged polysaccharide with OSO$_3^-$ side groups derived from red seaweed. These abundant polysaccharides possess advantageous properties such as biodegradability and low toxicity. However, at present, there is no clear consensus on the cell binding properties of CS and CRG or CS-CRG PEC materials.

The aim of this study was to explore the properties of crosslinker-free PEC gels, solvent-cast PEC films and freeze-dried PEC scaffolds based on CS and CRG precursors for medical applications. The objective was to characterise the effect of pH of the production conditions on the physicochemical and biological properties of CS-CRG PECs. Experimental work focused on the interaction between PEs, the composition of PECs, the rheological properties of PEC gels and the mechanical properties of PEC films and scaffolds. In addition, cell and protein attachment to the PEC films was assessed to determine their interactions in a biological environment.

For biomedical applications, these materials should ideally be stable when produced such that they can be processed to form either a film or a scaffold and have mechanical properties comparable to those of collagenous soft tissues. FTIR was used to confirm PEC formation. Zeta potential measurements indicated that the PECs produced at pH 2-6 had a high strength of electrostatic interaction with the highest occurring at pH 4-5. This resulted in stronger intra-crosslinking in the PEC gels which led to the formation of higher yield, solid content, viscosity and fibre content in PEC gels. The weaker interaction at pH 7-12 resulted in higher levels of CS incorporated into the complex and the formation of inter-crosslinking through entanglements between PEC units. This resulted in the production of strong and stiff PEC films and scaffolds appropriate for soft tissue implants. The PECs prepared at pH 7.4 and 9 also exhibited low swelling and mass loss, which was thought to be due to
the high CS content and entanglements. From the range of samples tested, the PECs produced at pH 7.4 appeared to show the optimum combination of yield, stability and homogeneity for soft tissue implants.

Biological studies were performed on CS, CRG and PECs prepared at pH 3, 5, 7.4 and 9. All of the PE and PEC films were found to be non-cytotoxic. When the response of three different cell types and a high binding affinity protein (tropoelastin) was evaluated; it was found that the CS-CRG PEC films displayed anti-adhesive properties. Based on these experimental observations and previous studies, a mechanistic model of the anti-adhesive behaviour of PEC surfaces was proposed. It was therefore concluded that the CS-CRG PECs produced might be suitable for non-biofouling applications.
# Abbreviations

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<tr>
<td>% v/v</td>
<td>Percent volume/volume</td>
</tr>
<tr>
<td>% w/v</td>
<td>Percent weight/volume</td>
</tr>
<tr>
<td>°C</td>
<td>Degrees Celsius</td>
</tr>
<tr>
<td>µl</td>
<td>Microlitre</td>
</tr>
<tr>
<td>2-D</td>
<td>Two-dimensional</td>
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<tr>
<td>3-D</td>
<td>Three-dimensional</td>
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<tr>
<td>AFM</td>
<td>Atomic force microscopy</td>
</tr>
<tr>
<td>Alg</td>
<td>Alginate</td>
</tr>
<tr>
<td>ATCC</td>
<td>American Type Culture Collection</td>
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<tr>
<td>AU</td>
<td>Arbitrary units</td>
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<tr>
<td>BA-4</td>
<td>Mouse anti-elastin antibody</td>
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<tr>
<td>BSA</td>
<td>Bovine serum albumin</td>
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<td>C-4-S</td>
<td>Chondroitin-4-sulfate</td>
</tr>
<tr>
<td>CB</td>
<td>Carboxybetaine</td>
</tr>
<tr>
<td>CBR (3B-A)</td>
<td>Cibacron brilliant red 3B-A</td>
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<td>ChS</td>
<td>Chondroitin sulfate</td>
</tr>
<tr>
<td>Coll</td>
<td>Collagen</td>
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<tr>
<td>CRG</td>
<td>Carrageenan</td>
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<td>CS</td>
<td>Chitosan</td>
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<tr>
<td>DAPI</td>
<td>4′,6-diamidino-2-phenylindole</td>
</tr>
<tr>
<td>DDA</td>
<td>Degree of deacetylation</td>
</tr>
<tr>
<td>DMEM</td>
<td>Dulbecco’s Modified Eagle’s Medium</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
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<tr>
<td>DRG</td>
<td>Dorsal root ganglion</td>
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<td>DS</td>
<td>Dermatan sulfate</td>
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<td>DSC</td>
<td>Differential scanning calorimetry</td>
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<td>EB</td>
<td>Elongation to break</td>
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<tr>
<td>ECACC</td>
<td>European Collection of Animal Cell Cultures</td>
</tr>
<tr>
<td>ECM</td>
<td>Extracellular matrix</td>
</tr>
<tr>
<td>EDC</td>
<td>1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride</td>
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EDTA  Ethylenediaminetetraacetic acid
ELISA  Enzyme-linked immunosorbent assay
EtOH   Ethanol
FBS    Foetal bovine serum
FDA    Food and Drug Administration
FITC   Fluorescein isothiocyanate
FTIR   Fourier transform infrared spectroscopy
g     gram
G’     Storage modulus
G”    Loss modulus
GA     Glutaraldehyde
GAG    Glycosaminoglycan
GPC    Gel permeation chromatography
GRAS   Generally recognised as safe
h     Hour
hASCs Human adipose-derived stem cells
HMEC  Human microvascular endothelial cells
hMSCs Human mesenchymal stem cells
HPLC  High-performance liquid chromatography
HRP   Horseradish peroxidase
HT1080 Human fibrosarcoma cell line
HyA   Hyaluronic acid
Hz     Hertz
ICDD  International Centre for Diffraction Database
IgG-HRP Goat anti-mouse secondary antibody coupled with HRP
ITC   Isothermal titration calorimetry
l     Litre
LDPE  Low-density polyethylene
LVR   Linear viscoelastic region
m     Mass
m0    Initial mass
MB    Methylene blue
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<td>$m_d$</td>
<td>Dried mass</td>
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<tr>
<td><strong>Micro-CT</strong></td>
<td>X-ray micro-computed tomography</td>
</tr>
<tr>
<td>min</td>
<td>Minute</td>
</tr>
<tr>
<td>ml</td>
<td>Millilitre</td>
</tr>
<tr>
<td>$m_t$</td>
<td>Total mass</td>
</tr>
<tr>
<td>mV</td>
<td>Millivolt</td>
</tr>
<tr>
<td>MW</td>
<td>Molecular weight</td>
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<tr>
<td>$m_w$</td>
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</tr>
<tr>
<td>N</td>
<td>Newton</td>
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<td>NCS</td>
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<td>NHS</td>
<td>N-hydroxy-succinimide</td>
</tr>
<tr>
<td>nm</td>
<td>Nanometre</td>
</tr>
<tr>
<td>nX</td>
<td>Non-crosslinked</td>
</tr>
<tr>
<td>OM</td>
<td>Optical microscope</td>
</tr>
<tr>
<td>OPA</td>
<td>o-Phthalaldehyde</td>
</tr>
<tr>
<td>Pa</td>
<td>Pascal</td>
</tr>
<tr>
<td>PAA</td>
<td>Polymethylacrylate acid</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate buffer solution</td>
</tr>
<tr>
<td>PE</td>
<td>Polyelectrolyte</td>
</tr>
<tr>
<td>PEC</td>
<td>Polyelectrolyte complex</td>
</tr>
<tr>
<td>PEG</td>
<td>Polyethylene glycol</td>
</tr>
<tr>
<td>$pK_a$</td>
<td>Acid dissociation constant</td>
</tr>
<tr>
<td>PLL</td>
<td>Poly-L-lysine</td>
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<tr>
<td>pNPP</td>
<td>p-Nitrophenyl phosphate</td>
</tr>
<tr>
<td>PP</td>
<td>Polypropylene</td>
</tr>
<tr>
<td>PS</td>
<td>Polystyrene</td>
</tr>
<tr>
<td>PTFE</td>
<td>Polytetrafluoroethylene</td>
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<tr>
<td>PVA</td>
<td>Polyvinyl alcohol</td>
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<tr>
<td>Rad/s</td>
<td>Radiant/second</td>
</tr>
<tr>
<td>RGD</td>
<td>Arginyl-glycyl-aspartic acid</td>
</tr>
<tr>
<td>rpm</td>
<td>Rotation per minute</td>
</tr>
<tr>
<td>RT</td>
<td>Room temperature</td>
</tr>
<tr>
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<td>Definition</td>
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<td>--------------</td>
<td>------------</td>
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<tr>
<td>s</td>
<td>Second</td>
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<tr>
<td>SAXS</td>
<td>Small angle X-ray diffraction and spectroscopy</td>
</tr>
<tr>
<td>SB</td>
<td>Sulfobetaine</td>
</tr>
<tr>
<td>SE</td>
<td>Secondary electrons</td>
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<tr>
<td>SEC-MALLS</td>
<td>Size exclusion chromatography-multi-angle light scattering detection</td>
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<tr>
<td>SEI</td>
<td>Strength of electrostatic interaction</td>
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<td>SEM</td>
<td>Scanning electron microscope</td>
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<tr>
<td>SUP</td>
<td>Supernatant</td>
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<tr>
<td>SW1353</td>
<td>Chondrosarcoma cell line</td>
</tr>
<tr>
<td>TB</td>
<td>Toluidine blue</td>
</tr>
<tr>
<td>TCPP</td>
<td>Tissue culture plate polystyrene</td>
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<tr>
<td>TGA</td>
<td>Thermogravimetric analysis</td>
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<tr>
<td>TGF-β</td>
<td>Transforming growth factor beta</td>
</tr>
<tr>
<td>TMB</td>
<td>3,3',5,5'-tetramethylbenzidine</td>
</tr>
<tr>
<td>TS</td>
<td>Tensile strength</td>
</tr>
<tr>
<td>UMR</td>
<td>Unit molar ratio</td>
</tr>
<tr>
<td>UTS</td>
<td>Ultimate tensile strength</td>
</tr>
<tr>
<td>UV-Vis</td>
<td>Ultraviolet-Visible Spectroscopy</td>
</tr>
<tr>
<td>WAXS</td>
<td>Wide-angle X-ray scattering analysis</td>
</tr>
<tr>
<td>Wt. %</td>
<td>Weight percent</td>
</tr>
<tr>
<td>WVP</td>
<td>Water vapour permeability</td>
</tr>
<tr>
<td>XL</td>
<td>Crosslinked</td>
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<tr>
<td>XPS</td>
<td>X-ray photoelectron spectroscopy</td>
</tr>
<tr>
<td>ζ-potential</td>
<td>Zeta potential</td>
</tr>
<tr>
<td>ι-CRG</td>
<td>iota-Carrageenan</td>
</tr>
<tr>
<td>κ-CRG</td>
<td>kappa-Carrageenan</td>
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<tr>
<td>λ-CRG</td>
<td>lambda-Carrageenan</td>
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Chapter 1 - Overview and Motivation

There is an increasing interest in the development of effective, sustainable and economical biomaterials that possess the appropriate physicochemical and biological cues for their intended applications. Biomacromolecules such as polysaccharides have wide potential as biomaterials 1,2, but often require chemical crosslinking to confer appropriate stability and mechanical strength 3. Some auxiliary crosslinker chemicals such as glutaraldehyde crosslinker are known to be toxic and mutagenic, and their fate in the body is not always well understood 4,5. “Self-crosslinking” via electrostatic interactions (polyelectrolyte complexing) can avoid the use of cytotoxic crosslinkers whilst maintaining stability at pH 7.4 and below 6. Unlike covalently crosslinked macromolecules, the advantage of polyelectrolyte complexes (PECs) is that no significant purification processes are required prior to administration 3. The PECs are dynamic materials owing to their reversible electrostatic bonds and charged groups which make them highly sensitive to their surrounding environment (especially to pH, ionic strength and PE concentration) 6–8. This dynamic structure can lead to controlled swelling and degradation, and may, therefore, be used for tissue engineering or drug-delivery applications in changing physiological environments.

This research focuses on the preparation, and physicochemical characterisation of PECs based on chitosan (CS) and κ-carrageenan (CRG) derived from shrimp shells and red seaweed, respectively. CS and CRG are less expensive than their glycosaminoglycan (GAG) counterparts found in the extracellular matrix of the body such as hyaluronic acid, chondroitin sulfate and heparin 9. The low costs are due to the vast abundance of waste shrimp shells 10 and farmed red seaweed 11 and their relatively simple extraction methods 9.

CS, a positively-charged polysaccharide is widely considered as the second most abundant organic material after cellulose 13. CS possesses appealing intrinsic properties such as non-toxicity 16 and biodegradability 17, as well as being bioadhesive 18 and bacteriostatic 19.

CRG, a negatively-charged polysaccharide, is an inexpensive alternative to heparin 20. It contains highly sulfated charged groups which result in immune-stimulating effects in organisms 21 or provide non-thrombogenic properties to coatings 20. CRG has also been shown previously to improve the formation of extracellular matrix components through bio-molecular crowding 22 and has been proposed for protein drug delivery 23 owing to its biocompatibility 24 and promotion of cell viability 25.
In the literature, there is a mixed view of the biological properties of CS and CRG materials. They have been reported to be suitable for both cell-adhesive and cell non-adhesive surfaces as shown in reviews by Liu et al. (2014) \(^{26}\) and Junter et al. (2016) \(^{27}\) and the references therein.

Although there have been wide interests in using CS and CRG as individual materials, there is currently little work published on CS-CRG PEC materials. PECs from CS and CRG have been prepared as freeze-dried 3-D scaffolds by Araujo et al. (2014) \(^{28}\). These scaffolds were found to be attractive for osteoblast cell attachment and showed potential for bone tissue engineering. However, CS-CRG PECs have also been widely reported to reduce cell adhesion such as in work carried out by Bratskaya et al. (2007) \(^{29}\) where CS-CRG PECs were used in a layer-by-layer film form. A reduction in the adhesion of two enterococcal strains was observed when compared with glass. Therefore, the biological properties of CS-CRG PECs merit further investigation in the form of films to exclude any potential complicating effects from the 3-D scaffold structure such as cell entrapment \(^{30}\) and 3-D architecture \(^{31}\).

The motivation for using CS-CRG PECs as opposed to CS and CRG alone is the potential for stable coatings, gels, films and scaffolds without further crosslinking, purification or modification. For example, once the PEC gel is produced, it can be solvent cast into non-porous films \(^{32}\) or freeze-dried into porous scaffolds \(^{33}\). CS alone can be too brittle for some applications due to its inherent chain rigidity \(^{34}\) and is difficult to control drug delivery when entangled alone \(^3\). CS is also highly soluble in acidic conditions due to its pK\(_a\) value of 6.4 \(^{35}\). CRG is soluble in water and therefore cannot be used in its pure form without covalent crosslinking \(^{36}\). CRG is also known to be more hydrophilic than CS \(^{37}\), and therefore the mechanical and hydration properties of CS-CRG PECs may be tailored by controlling the composition.

In this thesis, a rigorous approach to the synthesis and characterisation of PEC gels, films and scaffolds is presented. Previous research has shown that pH is a highly influential parameter on the charge density of PEs which could alter the physicochemical properties of PECs \(^{38}\). However, there are currently no detailed studies investigating the effect of pH on the physicochemical and biological properties of CS-CRG PECs. This study will improve the understanding of the effect of electrostatic complexing and self-crosslinking type upon the mechanical performance. It is hypothesised that varying the pH during PEC preparation will have a significant influence on the strength of the electrostatic interaction and ultimately on the stability of PECs produced.

The possibility of using polysaccharides for both tissue engineering and anti-fouling applications merits more detailed investigation as to whether the materials are adhesive or non-adhesive for cells. Therefore, the materials produced in this thesis will be tested with three different cell lines using cell attachment studies and a protein adhesion study. The origins of any effects observed will be discussed.
Chapter 2 – Literature Review

2.1 Polyelectrolytes

2.1.1 Introduction to Polyelectrolytes

Polyelectrolytes (PEs) are polymers with ionisable groups and exist as polycations (positively-charged polymers) and polyanions (negatively-charged polymers)\(^ {39}\). These materials become charged when their counterions are dissociated in an aqueous solution\(^ {40}\). They are similar to electrolytes, except that they consist of high molecular weight polymers and are therefore also known as polysalts because of their salt-like conductivity and polymer-like structure\(^ {41}\).

PEs can be divided into weak and strong types\(^ {42}\). The strong PEs usually consist of highly negatively-charged side groups such as sulfates (SO\(_3\)H) and phosphates (PO\(_3\)H\(_2\)) whereas weak PEs usually consist of carboxylic (COOH) side groups\(^ {42}\). Strong PEs dissociate at a wide pH range (e.g. pH 2-13), whereas weak PEs only dissociate at a more narrow pH range (e.g. pH 1-5)\(^ {44}\).

Generally, a non-charged polymer in a solution exists in a random conformation whereas a charged polymer will adopt a more expanded and rigid rod-like structure as previously measured with static and dynamic light scattering\(^ {45,46}\). The reason for the latter case is due to the high charge density of the side groups which causes the chains to repel each other\(^ {47,48}\).

The PE conformation is also largely influenced by the experimental conditions used\(^ {47,49,50}\). The charge density can be further altered with a change in counterion concentration (ionic strength)\(^ {42}\). The rigid chain structure may collapse back into a random neutral polymer like conformation when the ionic strength is increased\(^ {49}\). The random polymer conformation is due to the charge screening of the PEs by the counterions from the salt, and therefore reducing the overall charge density\(^ {51}\).

A similar random coil conformation of the PE may take place when the pH is near the pK\(_a\) value of the PE, inducing a neutral charge\(^ {38}\). Polymer conformation can affect the bulk properties i.e. viscosity and turbidity of solutions and can be studied using rheology, UV-Vis spectroscopy and static light scattering techniques\(^ {52,53}\).

A myriad of oppositely-charged PEs is available commercially for the production of self-crosslinked PECs. These PEs include polysaccharides\(^ {26}\), proteins\(^ {54}\), nucleic acids\(^ {55}\) and synthetic polymers\(^ {56,57}\). However, in this thesis, the emphasis will be placed on polysaccharides.
2.1.2 Polysaccharides as Polyelectrolytes

Polysaccharides can be divided into mammalian and non-mammalian types. CS is one of the few available positively-charged PEs. Therefore, the majority of PECs investigated are complexes between CS and other anionic PEs. Table 2.1 below shows a non-exhaustive list of the various types of polysaccharides and their associated charged side groups.

Table 2.1 – This table presents different types of existing PEs (mammalian and non-mammalian). The PEs available are mainly negatively-charged with the carboxylic side groups dominating the charges. The positively-charged CS may be used to complex with oppositely-charged polyanions.

<table>
<thead>
<tr>
<th>Polysaccharides</th>
<th>Charged Side group</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Non-mammalian</strong></td>
<td></td>
</tr>
<tr>
<td>Acacia (Aca)</td>
<td>COO&quot;</td>
</tr>
<tr>
<td>Alginate (Alg)</td>
<td>COO&quot;</td>
</tr>
<tr>
<td>Carboxymethyl cellulose (CMC)</td>
<td>COO&quot;</td>
</tr>
<tr>
<td>Gellan gum (GG)</td>
<td>COO&quot;</td>
</tr>
<tr>
<td>Gum kondagou (GK)</td>
<td>COO&quot;</td>
</tr>
<tr>
<td>Pectin (Pec)</td>
<td>COO&quot;</td>
</tr>
<tr>
<td>Xanthan gum (XG)</td>
<td>COO&quot;</td>
</tr>
<tr>
<td>Xylan (Xyl)</td>
<td>COO&quot;</td>
</tr>
<tr>
<td>Carrageenan (CRG)</td>
<td>OSO₃⁻</td>
</tr>
<tr>
<td>Fucoidan (FD)</td>
<td>OSO₃⁻</td>
</tr>
<tr>
<td>Porphyran (Por)</td>
<td>OSO₃⁻</td>
</tr>
<tr>
<td>Chitosan (CS)</td>
<td>NH₃⁺</td>
</tr>
<tr>
<td><strong>Mammalian</strong></td>
<td></td>
</tr>
<tr>
<td>Dextran sulfate (DexS)</td>
<td>OSO₃⁻</td>
</tr>
<tr>
<td>Heparan sulfate (HS)</td>
<td>OSO₃⁻</td>
</tr>
<tr>
<td>Keratan sulfate (KS)</td>
<td>OSO₃⁻, OSO₃⁻</td>
</tr>
<tr>
<td>Chondroitin sulfate (ChS)</td>
<td>OSO₃⁻, COO&quot;</td>
</tr>
<tr>
<td>Heparin (Hep)</td>
<td>OSO₃⁻, COO&quot;</td>
</tr>
<tr>
<td>Dermatan sulfate (DS)</td>
<td>OSO₃⁻, COO&quot;</td>
</tr>
<tr>
<td>Hyaluronic acid (HyA)</td>
<td>COO&quot;</td>
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</table>
Despite their disparate origins, non-mammalian and mammalian polysaccharides possess similar structures with recognised side groups (Fig. 2.1). Hence, non-mammalian polysaccharides have been used before as an alternative substitute to glycosaminoglycans (GAGs) for drug delivery and tissue engineering applications. It has been reported that non-mammalian polysaccharides like chitosan (CS) and κ-carrageenan (κ-CRG) have a lower immunogenic response and pathogen transmission risk than the bioactive GAGs. The next two sections will describe the production, structure and properties of CS and CRG in greater detail.

2.1.3 Chitin/Chitosan

Chitin is found in the shells of crustaceans (shrimp shell, crab shell), the exoskeleton of insects and cell walls of fungi. Every year 6-8 million tonnes of crustacean waste is produced which makes chitin highly abundant as well as economically and environmentally sustainable. Since chitin does not dissolve in acids or alkalis, it is usually deacetylated with strong alkalis such as NaOH (replacing the acetyl groups with amine groups) to form chitosan. The extraction of chitosan from raw crustacean shells can be described in four main stages: 1) cleaning and grinding of shells, 2) deproteination of the shell to chitin/calcium carbonate, 3) demineralisation of chitin/calcium carbonate to chitin, 4) deacetylation of chitin to chitosan. A more detailed description of the production process of CS is described below:
1) Shrimp shells are used because the walls are thin and therefore the isolation of chitin becomes easier. The selected shells are then cleaned, dried, and ground into small shell pieces.

2) The chemical bonds between the chitin and the proteins are disrupted when using NaOH at a concentration ranging from 0.125 to 0.5 M, at varying temperatures (up to 160 °C) and treatment duration (from few minutes up to few days). Invariably, the chemical NaOH used can also result in partial deacetylation or hydrolysis of CS. The protein is removed since this is the primary cause for an allergic reaction in humans.

3) Dilute HCl (up to 10 % w/v) at room temperature, and reaction time ranging from 15 min to 48 h, are used for the decomposition of calcium carbonate present in the chitin matrix. Longer demineralisation time results in lower ash (salt) contents but also causes polymer degradation.

4) Alkali deacetylation is used more frequently than acid deacetylation because the glycosidic bonds are more susceptible to acid than to alkali which results in lower MW. The N-deacetylation of chitin is carried out by a hot concentrated solution of NaOH for a few hours to obtain a degree of deacetylation (DDA) of 85-99 %. Alternatively, chitin is placed in concentrated NaOH (30 g NaOH/45 g H2O/3 g of chitin) at 25 °C for 3 hours or more, followed by dissolution in crushed ice around 0 °C. This method produced a soluble CS with a DDA of 48-55 %.

As a linear polysaccharide, CS is composed of β-1,4-linked glucosamine (deacetylated) and N-acetyl-D-glucosamine (acetylated units) (Fig. 2.2) 14. Commercially, the polymer can be obtained with a deacetylation between 50 to 95 % with a molecular weight between 10 to 1000 kDa 59,65,66. The polymer can be classified as low molecular weight (< 150 kDa) and high molecular weight (> 700 kDa) 59. When the MW reaches < 10 kDa, the material is known as an oligochitosan 14.
CS has been widely used in biomedical applications and is considered safe for human use in topical, subcutaneous and ocular applications. The glycosidic bonds present in CS can be broken by enzymes (i.e. lysozyme) present in the body into oligosaccharides such as glucosamine (present in the body as a constituent for joint lubrication). This allows films or scaffolds to be broken down in the long-term into biocompatible components and could ultimately reduce the chronic inflammatory response and potentially help to reduce fibrotic encapsulation. CS has also found applications in weight loss treatments, although the outcomes have not been very effective. The positive charges present in CS molecules make it potentially bactericidal. The antibacterial effect is thought to be due to the interaction between the positively-charged groups in CS and the negatively-charged anions located in bacterial cell walls. This interaction disrupts the cell walls and releases their inner contents. CS has been found to have mucoadhesive properties which increases retention at the site of implantation. The effect of pH on the solubility of CS in water and the low viscosity at high concentration makes CS suitable as a cell penetration enhancer for genes or as a drug delivery system. CS is also a highly effective haemostatic agent and can aggregate platelets which result in rapid blood clotting in major liver injuries. For this reason, CS is widely used as a form of coating or replacement of cellulose fibre as a form of haemostatic gauzes to stop critical bleeding injuries encountered in the battlefield. Other researchers have also reported that CS is non-adhesive to cells or proteins. Therefore, CS has found uses in both medical implants for tissue engineering or anti-fouling applications. Finally, CS can be complexed with negatively-charged molecules including growth factors, nucleic acids, cytokines, anionic glycosaminoglycans (GAGs) and proteoglycans, which are essential components of the extracellular matrix (ECM) found throughout the body.

Figure 2.2 - Molecular structures of chitin, chitosan, and protonated chitosan polymer.
2.1.4 Carrageenan

CRG is a natural linear sulfated polysaccharide and is obtained from edible red algae (Rhodophyceae) \(^{83}\). It is also known as Irish moss because it historically originates from the county of Carraghen situated on the south coast of Ireland where it has reportedly been used for the past 600 years \(^{83}\). Essentially, dried red algae are washed and soaked in water for 24 h \(^{84}\). The pulp (cut and pressed algae) is then mixed with water at a concentration of 1:80 (v/v). The mixture is adjusted to pH 9 with Ca(OH)\(_2\) solution followed by warming at 90 °C for 2 h by stirring. The extracted viscous filtrate is separated from the cellulose (solid material) of the algae. The filtrate is neutralised to pH 7 with 1 % HCl solution and reheated to 60 °C for 30 min. The filtrate is then coagulated using KCl solution at a concentration range of 1.5-3.5 % for 15 min, and the CRG gel is subsequently filtered to remove the water. Finally, the CRG gel is soaked in 96 % ethanol, dried at 70 °C in air for 24 h and milled.

CRG consists of a linear backbone built up by β-D-galactose and 3,6-anhydro-α-D-galactose linked with α-1,3 and β-1,4-glycosidic bonds \(^{85}\). When the polymer is dissolved in water, the sulfate groups ionise to give a negatively-charged molecule. There are three main commercial types of CRGs: kappa (κ) - (one sulfate group per disaccharide), iota (ι) - (two sulfate groups per disaccharide) and lambda (λ) - (three sulfate groups per disaccharide) CRG (Fig. 2.3) \(^{86}\). κ-, ι-CRG can change from a random coil into a double helix conformation and could, therefore, gel when the critical concentration is achieved at a critical temperature \(^{87}\). On the other hand, λ-CRG does not form a gel but only increases in viscosity with an increase in concentration. The US Food and Drug Administration (FDA) has considered CRG “generally recognised as safe” (GRAS) for consumption and topical applications \(^{83}\). CRG is widely used as a thickener in ice creams, stabiliser in toothpaste and dispersant for barium sulfate suspension used in medical imaging \(^{83}\). It has also been proposed for tissue engineering and drug delivery systems, due to its biocompatibility \(^{24,88}\) and its ability to enhance cell viability \(^{23,25}\). On the other hand, CRG has also been shown to reduce cell adhesion of which some of the factors were thought to be due to the highly hydrophilic nature and highly repulsive sulfate groups preventing negatively-charged cells to adhere \(^{29,89}\).

Figure 2.3 – The molecular structures of commercially available CRG types: κ-CRG, ι-CRG and λ-CRG with one, two and three sulfate groups per monomer unit, respectively \(^{83}\).
2.2 Polyelectrolyte Complexes

2.2.1 Introduction to Polyelectrolyte Complexes

Polyelectrolyte complexing is similar to ionic crosslinking (complexing). The difference is that in ionic complexing, a small and well-defined MW is used compared to the polyelectrolytes containing a broad MW distribution with charged side groups 69.

The formation of PECs can be an attractive alternative to covalent crosslinking since PECs have a dynamic structure due to their reversible bonds 5. These reversible bonds are more sensitive to any external factors such as ionic strength and pH compared to covalent bonds. As a consequence, the swelling characteristics can be changed with the change in charge density. Unlike covalently crosslinked macromolecules, dissolution can occur at certain pH ranges in PECs 6. Hence, smart polymeric systems may be created that can deliver drug controllably at changes in physiological conditions. In addition, since no covalent crosslinkers are required for stability, these PEC materials may be combined with biologically stimulating ingredients such as growth factors, drugs, antibiotics, bioactive materials, cells, cytokines, proteins, enzymes and nucleic acids for specific medical applications 34,90,91.

The advantage of PECs is that no significant purification methods are required after preparation since no crosslinkers are needed to be removed. Also, the strengths and stiffness of PEC gels are usually lower than those in covalently crosslinked macromolecules which may be used for softer tissues 92. The interactions between PEs can be strong and long-term under physiological conditions which make PEC materials well tolerated and potentially stable as implants 69. These PECs are relatively versatile as they can be shaped into many different assemblies (e.g. coatings, films, gels, nanoparticles, colloids, fibres, and scaffolds) 93.

However, the main drawback of the PEC system is that preparation can be quite laborious and the production yield in large scale may be difficult 94. The final properties of the PECs can also change significantly depending on pH, ionic strength, temperature, time of mixing, the degree of ionisation of each of the oppositely-charged PEs, the density of the charges on the PEs, charge distribution over the polymeric chains, concentrations of the PEs and polymer chain flexibility 95. Reproducibility can be difficult since small changes in many parameters can affect the physicochemical properties of PECs during preparation.
2.2.2 General Concept of Polyelectrolyte Complex Formation

The theory of formation and organisation of PECs was pioneered in the early 1980s by Japanese scientists E. Tsuchida, Y. Osada and H. Ohno \(^96\) and Russian scientists V. A. Kabanov and A. B. Zezin \(^97\). The equation below signifies a polyelectrolyte complexation reaction:

\[
(A^-c^+)_n + (C^+a^-)_m \rightleftharpoons (A^-C^+)_{nx} + (A^-c^+)_{n-x} + (C^+a^-)_{m-x} + xc^+ + xa^-
\]  \(2.1\)

where \(A^-\) and \(C^+\) are PEs, \(a^-\) and \(c^+\) are the corresponding counterions, \(n\) and \(m\) are the numbers of ionic groups, and \(x\) is the molar mixing ratio i.e. \(n/m = x\) or \(m/n = x\) \(^59\). Complex formation is entropically driven since the counterions are released when the macromolecules are reacted \(^43,54\). The entropic increase of the liberation of counterions is higher than the entropic decrease caused by the condensation of the two PEs \(^98\). For complexing to occur, the oppositely-charged PEs need to be in an aqueous environment at a pH interval between the \(pK_a\) values of the two polymers, CS (\(pK_a = 6.5\)) \(^99\) and CRG (\(pK_a = 2\)) \(^100\) e.g. pH 4.25.

Secondary interactions such as hydrogen bonding, ion-dipole and dipole-dipole attractions may also take place, but the primary form of binding is due to the electrostatic attraction (ionic bonds) \(^101\). The ionic bonds are associated with high water content and electrical charge density \(^5\). New ionic bonds formed between oppositely-charged PEs may be indicated using FTIR spectroscopy where a new and/or shifted wavenumber is detected \(^102–104\). The polyelectrolyte complexation process can also be followed with microscopy \(^105\), viscosity of supernatant \(^48,106\), conductometric and potentiometric titration of mixture \(^34,107,108\) and turbidity of mixture or supernatant solutions \(^34,106,109–111\).

Many of the above-mentioned factors are mainly determined by the charge density of the polyions which is influenced by the pH used \(^112\). When the pH is adjusted to reduce the charge density of a single PE, the PEC becomes softened and turns into a complex coacervate like structure, whereas at optimal interaction pH, the charge density for both PEs is high and therefore a precipitate is likely to occur due to stronger interaction \(^43\). Complex coacervation is also used as a term for polyelectrolyte complexing where oppositely-charged colloids complex together. Bungenberg de Jong and Kruyt \(^113\) coined the term complex coacervation to distinguish it from the mere coacervation of a single polymer as cited by Kizilay et al. (2011) \(^114\). However, the term complex coacervate is also widely used for weakly bound oppositely-charged PEs exhibiting a gel-like or fluid-like consistency distinguished from the solid consistency found in complex precipitates \(^43\). Precipitation can be avoided when the charge interaction is weakened by adding adequate amounts of NaCl or adjusting the pH close to the \(pK_a\) of the weak PE \(^7,115\). However, at high ionic strength, the PEC can be suppressed, and therefore no phase separation such as complex coacervation is likely to occur. Depending on the strength of interaction...
between PEs the time-scale of diffusion measured with a stopped-flow instrument can range from milliseconds to days and even longer in precipitates since it is harder to rearrange strongly attracted PEs. The structure is, therefore, dynamic and reversible and can vary with the strength of interaction which ultimately depends on the PEs and the conditions of the solvents used.

To improve the stability of PECs, further ionic crosslinking can be induced by the addition of ions such as Ca$^{2+}$ in alginate and K$^+$ in CRG. Ionic crosslinking is governed by the strength of electrostatic interaction between the polymers, and this depends on the global charge densities which control the relative amount of individual PE in a PEC. The PE with a lower charge density is usually present in higher amounts within the PEC as more polymer is required to cancel out the opposite charge of the other PE. The stability of PEC solids can also be enhanced by annealing the complex at high temperatures which may result in covalent bond formation under dry conditions.

PECs can exist in various forms, and their structures can generally be divided into two types i.e. the ladder-like and the scrambled egg-like model (Fig. 2.4). The ladder-like model was one of the earliest models used to depict PEC structures, where the PEs were proposed to be only extended and zipped together. The scrambled egg model represents more flexible PEs ionically linked into a random coil-like conformation. PECs represent a combination of both models, however the latter being more representative of most PEC structures. The irregular scrambled egg structure makes some of the free ionic charges not always accessible due to steric effects. The structures of PECs have been determined by using a combination of characterisation techniques including x-ray analysis, polarised light and electron microscopy, potentiometric titration, dynamic light scattering, viscosity, turbidity, circular dichroism and calorimetry.

Figure 2.4 – Schematic depiction of two possible PEC structures (ladder-like and scrambled egg-like model).
In order to understand the structure of PECs and its influence on the stability and mechanical properties of PEC gels and solids, it is important to recognise how PECs are formed. E. Tsuchida was the first to present the structures of PECs as it is known today. PECs are generally formed in the following three processes described below and shown schematically in Figure 2.5:

1. Primary complex formation (Coulomb forces): mainly a rapid electrostatic attraction between the charged molecules results in the formation of an amorphous aggregate which is also known as the random primary complex. The structure formed is referred to as the scrambled egg-like structure due to the random arrangement of the macromolecules.

2. Ordered secondary complex: formation of new bonds and/or the correction of distorted bonds of PE chains within intracomplexes, mainly through rearrangement and hydrogen bonding between the macromolecules. The structure formed is known as the ladder-like structure due to the ordered rearrangement of the macromolecules.

3. Intercomplex aggregation processes: complex aggregates are mainly formed through hydrophobic interactions and subsequent entanglements to form intercomplexed fibrils that can further develop into networks. Note, that the random primary complex can also turn into a stable intercomplex aggregate through entanglements without the need to reform into an intracomplex first.

PEC formation is mainly driven by the ion-exchange reaction that occurs at a high rate even when dilute solutions are used. When weak PEs are utilised then the effect of pH can dramatically influence their charge density (supported by zeta potential measurements), and therefore the extent of reaction can be varied.
2.2.3 Effect of Experimental Conditions on Polyelectrolyte Complex Formation

PEC preparation is a versatile process, and the final properties are highly dependent on the physicochemical properties of the polymers and the reaction conditions used \(^{126,127}\). Understanding the effect of reaction conditions could potentially control the strength of electrostatic interaction and ultimately the mechanics and stability of films and scaffolds in the search for self-crosslinked biomaterials.

The strength of electrostatic interaction between oppositely-charged PEs can be altered depending on several factors such as the degree of ionisation of each PE, degree of deacetylation of CS, the charge distribution over the polymeric chains (charges can be evenly distributed or concentrated in one end of chain), the nature of the ionic groups on the polymeric chains (carboxylic groups, sulfate groups and amine groups), the molecular weight of the PEs (i.e. higher molecular weight may result in greater entanglement) and the polymer chain flexibility \(^{95,128}\). Other parameters can be altered externally such as the concentration of the PEs, their mixing molar ratio, the mixing order of PEs, speed of mixing (dropwise addition or one-shot addition), the duration of mixing, as well as the temperature (some polymers need heat to dissolve), ionic strength, and pH used to fully dissolve the material (the right pH needs to be used in the medium to ionise the PE fully) \(^{58}\). PEC formation can be either stoichiometric or non-stoichiometric depending on the molar mixing ratio, acidity/basicity, chain length and chain flexibility of PEs in PECs \(^{129}\). In theory, when equal amounts of oppositely-charged groups are combined in a stoichiometric manner, the overall net charge becomes zero, and therefore the complex formed is an insoluble precipitate \(^{130}\).

Other indirect factors that affect the stability and formation of PECs are the diffusion coefficient, chain conformation (straight chains, helical chains or random folded chains), viscosity (viscous solutions reduce complex formation), miscibility (the more miscible the more reactive) between PE solutions, centrifugation step, stirring rate, acid type and washing steps \(^{131}\). In the next few paragraphs, the most important parameters on PEC formation are discussed in more detail.

The pH has been found to be an important parameter affecting PEC formation due to its influence on the degree of ionisation on polyions \(^{38,44}\). When both polyions are weak PEs, their ionisation degree is highly dependent on the pH. It was found that the strength of electrostatic interaction as measured with zeta potential was highest, and drug release was lowest when the PEs have complexed at an optimal pH range i.e. pH in the range between the pK\(_a\) of both PEs \(^{33,132}\). For the case of CS complexed with a strong PE, a decrease in pH results in higher ionisation degree of CS and the polyanion which results in strong interaction \(^{38}\). However, at extreme acidic and alkaline conditions in the presence of
weak PEs, one of the reacting PEs would either be fully ionised or non-ionised and therefore electrostatic interaction would not likely occur. In a different study, large insoluble complexes were formed when β-lactoglobulin was complexed with alginate at pH 3 and 4 due to the strong electrostatic interactions. At pH 5, soluble complexes were formed whereas, at pH 6 and 7, no complexes were formed due to the charge reversal of the polyampholyte β-lactoglobulin which resulted in electrostatic repulsion with negatively-charged alginate. The pH was also found to strongly affect the zeta potential of the individual chitosan and gum arabic PEs, with the highest PEC yield and viscoelastic properties at pH 4.5. Similarly, the interaction between whey protein isolate and κ-carrageenan displayed higher viscosity at pH 4.5 that could be associated with a more compact microstructure and robust PECs prepared at pH 4. These results show that only a small change in pH can largely influence the electrostatic interaction and ultimately the physical properties of PECs.

The ionic strength is another parameter that can significantly affect the complex formation. The addition of large amounts of inorganic salts results in the screening of the charged groups present in the PEs and can, therefore, reduce the overall charge density. This leads to the weakening of the electrostatic interaction of polyions. Thus, complex coacervation of PECs may occur. Coacervates can break when excess salt is added but may not break complex precipitates since these structures are usually irreversible in optimal pH conditions.

The initial concentration of the PEs is of great importance for polyelectrolyte complexation. Work carried out by Chieng and Chen have shown that at low concentrations of poly(diallyldimethylammonium chloride) (PDADMAC) result in vesicle complex formation between PDADMAC and phospholipids. At very high concentration and chain length of PDADMAC, the vesicles become interconnected, forming a supramolecular network. This factor influences the composition, size, surface charge, and stability of PECs. Ultimately, it is of great interests to control the self-crosslinking of PECs by varying the production conditions to control film and scaffold mechanics and stability for specific applications.

The degree of deacetylation (DDA) of CS is directly related to the number of amino groups present per disaccharide (monomer unit), and therefore the interaction with other polyanions is strongly dependent on this parameter. It was confirmed by Gåserød et al. (1998) that an increased number of acetyl groups resulted in weaker interaction between CS and alginate due to the decrease in overall charge density (charges per unit length) of CS as supported by zeta potential.

Molecular weight is an important parameter since it directly relates to the chain length of the PE and can be a limiting factor to the binding to other polyanions. It was previously found by others that a higher MW resulted in less intensive and surface limited binding than with low MW CS as measured.
using a fluorescent microscope to observe the fluorescently stained CS. This was due to the ability of small chain CS molecules to diffuse and penetrate more easily into the core of the PEC. On the other hand, the drug release was shown to be prolonged with higher MW CS compared with low MW CS. This could be ascribed to the thicker, more viscous and less permeable outer wall produced on the PEC when using higher MW CS. Higher MW can also lead to a higher drug encapsulation efficiency. However, the high viscosity can hinder diffusion and binding of CS molecules to other polyanions. Therefore, complexation between CS and polyanions may be completely or partially halted.

Several facts can be drawn from the complexation studies presented in the literature. It seems that when:

- One of the polions is a strong PE then a precipitate is more likely to occur rather than a complex coacervate gel. Precipitates are more likely to happen in polysaccharides with strongly charged side groups containing sulfates or phosphates than they are in polysaccharides with weakly charged side groups containing carboxyls.
- The mixing ratio is equal, the resultant complex is neutral.
- An excess PE is present, the overall complex is only to some extent charged with the charge of the PE in excess.
- The pH of the solution is adjusted close to the pK\textsubscript{a} of the PEs, the complexation may be suppressed.
- Salt is added, complexation may be suppressed.

Additional information on complexation can be found in the extensive review on PECs written by Thünemann et al. (2004).  

### 2.2.4 Chitosan-Carrageenan Polyelectrolyte Complexes

This section of the review spans work carried out on PECs based on CS-CRG from two decades i.e. from the years 1997 to 2017. The review includes information on hydrogels, films and scaffolds and their associated properties obtained such as stability, mechanics and degradability. Table 2.2, summarises the parameters controlled, methods and results of the typical PECs produced based on CS and CRG.

With respect to the CS-CRG PEC gels, the main parameters studied previously include the effects of CRG concentration, ionic strength (NaCl concentration), CRG type (\(\kappa\), \(\iota\), \(\lambda\)\)), different CS and CRG concentration, and varying degree of deacetylation of CS with various CRG types (\(\kappa\), \(\iota\), \(\lambda\))\). Homogeneous complex coacervate gels have been produced when reacting CS and CRG with 5.7 % NaCl and extended mixing times (5-7 days).
There have been several studies reported in the literature on the production and characterisation of CS-CRG films. Carneiro et al. (2013)\(^{147}\) have solvent cast CS-CRG PEC gels into films to produce non-thrombogenic materials for blood contacting applications. These were prepared by using four different volumetric proportions of CS and CRG. The swelling of the PECs was found to decrease when the amount of CS was increased. The PECs were found to be non-toxic to fibroblasts making them suitable for cell therapies. In another study, CS-CRG PEC films were also made by solvent casting the mixtures using different acids (acetic, ascorbic, lactic, malic and citric acid) for potential wound healing applications\(^{148}\). Further information about the results can be found in the summary written in Table 2.2.

Three-dimensional structures made from CS-CRG PECs have only been reported by Araujo et al. (2014)\(^{6}\). Araujo and co-workers have shown that stable PEC scaffolds can be produced at pH 7.4 and below at different CS-CRG molar ratios (1:1, 2:1 and 3:1). However, they did not investigate the effect of varying pH on the PEC scaffold formation. Instead, they investigated the stability of the PECs under different pH conditions (4.5, 7.4, 9 and 11). They found that the PECs at very high pH (9 and 11) can induce solubilisation of the PEC whereas, at pH 7.4, the complexes were stable.

To the best of the author’s knowledge, there appear to be no previous studies looking at the effect of pH on the preparation of CS-CRG PEC gels. In addition, the PEC gels in the literature have not been characterised systematically or in great detail. Some characterisation techniques for CS-CRG PEC gels such as ζ-potential and microscopy remain unexplored. In addition, the effects of the strength of electrostatic interaction and the type of crosslinking (electrostatic vs. secondary interactions) on the mechanical performance in terms of gels, films and scaffolds are limited. Finally, a systematic study of the biological response using a variety of cells and proteins to PECs with controlled composition is still missing in the literature.
Table 2.2 – A table summarising the parameters controlled during CS-CRG PEC gels, films and scaffold preparation.

<table>
<thead>
<tr>
<th>PEC Type &amp; Reference</th>
<th>Parameters Controlled During PEC Preparation</th>
<th>Summary of PEC Preparation Method</th>
<th>Summary of Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gel 7</td>
<td>• Ionic Strength</td>
<td>CS was dissolved in dilute acetic acid at 80 °C. CRG and NaCMC were mixed with NaCl. The solutions were then mixed and kept at 5 °C for two days.</td>
<td>Aggregates were formed due to the strong electrostatic interaction between CS and highly charged CRG. As NaCMC content was increased, more homogeneous PECs were formed, but at very high NaCMC, no PEC was formed due to the weaker (−ve) carboxylic groups. The addition of salts suppressed the high charge density and reduced the swelling.</td>
</tr>
<tr>
<td>Gel 21</td>
<td>• CRG Type (κ, ι, λ)</td>
<td>CS was dissolved in dilute sodium acetate buffer (0.02 M). CRG was dissolved in water at 50 °C. The PEs were mixed for 5-7 days.</td>
<td>At low concentrations of CS and λ-CRG, insoluble precipitates were formed. At higher concentrations, complex coacervate gels were formed. Concentrations higher than 1 % (w/v) of either PE becomes too viscous, and no complexation would occur. The mechanical strength was highest at κ-CRG and lowest at λ-CRG. This effect was explained by the formation of additional crosslinks by double helixes in κ-CRG.</td>
</tr>
<tr>
<td>Gel 145</td>
<td>• Ionic Strength</td>
<td>CS and κ-CRG were dissolved in 1 % acetic acid and in water at 70-80 °C, respectively. The PEs were mixed at varying NaCl content. The suspension was then centrifuged and washed.</td>
<td>Considerable amounts of NaCl (4 % or 6 % w/w) inhibits PEC formation due to the screening the PE charges. Swelling of PEC gel at pH 10-12 was 10.2 x higher than the rest of the pHs. The maximum swelling occurred at pH 10.5. At pH &lt; 9 and pH &gt; 13, no swelling was observed.</td>
</tr>
<tr>
<td>Gel 146</td>
<td>• CS MW</td>
<td>CRG and CS solutions were mixed in PBS. CS and CRG were determined spectrophotometrically using oppositely-charged dyes.</td>
<td>At high CRG concentration, soluble complexes were formed. Low MW CS bonds more efficiently to CRG than high MW CS due to the lower steric hindrance in the shorter chain length. High MW CS is in a random coil conformation and can, therefore, screen some of the binding sites.</td>
</tr>
<tr>
<td>Gel 143</td>
<td>• CRG Type (κ-, ι-, λ-)</td>
<td>CRG was dissolved in water at 50 °C for 30 min. PECs were formed by adding CS and CRG at different concentrations. The PECs were centrifuged and the amount of non-reacted PEs was detected</td>
<td>At increased ionic strength, the coil-helix transition is inhibited and therefore complexes with charge ratios slightly less than unity was formed. Hydrophobic forces were thought to dominate at high salt concentrations.</td>
</tr>
<tr>
<td>Film 147</td>
<td>Molar Ratio of CS and CRG</td>
<td>CS was dissolved in acetic acid solution and cast into films followed by neutralisation in 1 M NaOH solution for 24 h. κ-CRG was dissolved in water under stirring at 80 °C for 1 h and cast and immersed in 2 M KCl solution for 24 h and washed with water and dried for 2 days. CS dissolved in acetic acid was mixed with CRG dissolved in water at different proportions. The films were neutralised in NaOH for 24 h and dried.</td>
<td>The swelling of PECs decreased when the amount of CS increased. Bovine serum albumin (BSA) adsorption was decreased on PECs but fibrinogen adsorption increased when compared to the individual CS and CRG films. Although all films showed similar cell expansion and viability, the PEC 50:50 vol. % CS/κ-CRG has been proposed as an acceptable scaffold material for new cell therapies, due to their positive effect on cell survival.</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Film 148</td>
<td>Acid Type</td>
<td>Either acetic acid, ascorbic acid, lactic acid, malic acid or citric acid were used with ascorbic acid to dissolve CS. The PECs were prepared by mixing κ-CRG and CS with glycerine and PEG as plasticisers. The films were dried in an oven at 70 °C for 12 hours.</td>
<td>Malic acid exhibited overall highest tensile strength (TS), elongation (EB) and low water vapour permeability (WVP). Citric acid exhibited the lowest TS, Young’s modulus (E) and WVP properties. Acetic acid showed a relatively high TS but a low EB and WVP. Lactic acid showed average TS and E but with highest WVP of all acid types used.</td>
</tr>
<tr>
<td>Scaffold 6</td>
<td>Molar Ratio of CS and CRG</td>
<td>CS solution was dissolved in dilute HCl. CRG was dissolved in water at 80 °C. The CS solution was added to the CRG solution and mixed vigorously. The PEC was centrifuged and the amount of CRG in the supernatant was detected spectrophotometrically. The PEC was washed and freeze-dried to obtain a scaffold.</td>
<td>The PECs were stable at pH ≤ 7.4 but dissolved as the pH increased to non-physiological values of 9 and 11. The PECs precipitated at pH 4.5 with some hysteresis. This showed that the PEC formation was reversible during preparation at different conditions. The use of higher CRG concentration resulted in an increase in hydrophilicity and water absorption in scaffolds. An increase in CS content resulted in an increase in mechanical properties. Overall, it was shown that the PEC scaffolds were very stable under physiological conditions.</td>
</tr>
</tbody>
</table>
2.3 Conclusions and Unanswered Questions

Previous studies have shown that pH is highly influential on the PEC formation in different PEC systems. However, very little work has been conducted on the effect of pH on the physicochemical properties of CS-CRG PECs. This thesis considers the effect of pH on the PEC interaction between weak CS and strong CRG. The resulting properties of PEC gels, films and scaffolds are then characterised and discussed.

2.4 Project Aims

The ability of self-crosslinking materials without having to use toxic chemicals is highly desirable for both tissue engineering or anti-fouling applications. This work aims to provide a comprehensive analysis of the effect of a change in pH on the production and characterisation of PECs, as gels, films and scaffolds. This study will aid the understanding of the binding between PEs under wet and dry conditions. The use of films offers the potential to understand the properties of a surrogate for a scaffold strut. Once there is sufficient understanding of the key parameters and their effects on the physicochemical properties of PECs, then the biological response to the films can be measured. The effects observed in scaffolds may be both chemistry-related, and due to the physical effects of the scaffold architecture.

This thesis will explore the production, and physicochemical characterisation of PECs and compositions optimised for stability will subsequently be evaluated using biological testing. The aims of this thesis are as follows:

- Establish a suitable methodology to quantitatively measure the concentration of PEs present in the PEC mixture.
- Production and characterisation of CS-CRG PEC gels, films and scaffolds.
- Establish and understand the effect of pH on the interaction and composition of prepared PECs using physicochemical analytical techniques.
- Establish the biological properties of the PECs using three different general cell lines and a protein adhesion study to find out whether the materials are cell adhesive or anti-cell adhesive in nature.
2.5 Thesis Structure

The structure of this thesis is as follows:

- Chapter 3 explores the potential of a developmental study of a spectrophotometric method to potentially determine the level unreacted PEs during PEC formation and ascertain the composition of the PECs.

- Chapter 4 describes the preparation of self-crosslinked CS-CRG PEC gels. These PEC gels were characterised in their native state to understand the interaction between CS and CRG macromolecules. The strength of electrostatic interaction and composition of PEC gels were controlled by preparing the PECs at various pH.

- Chapter 5 describes the preparation of self-crosslinked CS-CRG PEC films. These films were developed as free-standing materials through solvent casting the PEC gel suspensions to test the effect of self-crosslinking on mechanical properties.

- Chapter 6 concerns the development of self-crosslinked CS-CRG PEC scaffolds. These scaffolds were formed by freeze-drying the PEC suspension to form a porous three-dimensional structure. The effect of self-crosslinking was tested on the mechanical properties of scaffolds.

- Chapter 7 considers the biological properties of the self-crosslinked CS-CRG films. Three different cell lines and a protein were used to determine the applicability of CS-CRG PEC materials for tissue engineering or non-biofouling applications.

- Chapter 8 reports the overall conclusions and outlines suggestions for future work.
Chapter 3 – Spectrophotometric Study

3.1 General Introduction

In this thesis, a method is required to quantitatively measure polyelectrolyte (PE) concentrations in dilute mixed polyelectrolyte systems. The dilute mixed PE systems studied consisted of two oppositely-charged PEs namely, chitosan (CS) and carrageenan (CRG).

Methods previously employed in the literature to confirm PEC formation have included visual observation, fourier transform infrared spectroscopy (FTIR)\(^{34,46,149–155}\), thermogravimetric analysis (TGA)\(^{149,150,155}\), differential scanning calorimetry (DSC)\(^{151,34}\), X-ray diffraction spectroscopy (XRD)\(^{151,153–155}\), the viscosity of supernatant\(^{156}\), the viscosity of PEC\(^{34}\), potentiometric titration\(^{149}\), conductometric titration\(^{149}\), turbidimetric titration\(^{149,157}\), and pH\(^{34}\). However, most of these techniques can only provide a global evaluation of complex reaction and do not explicitly provide quantitative information on complex interaction and composition\(^{46}\). Quantitative methods that were previously used to determine the composition of the complex include elemental analysis techniques\(^{46}\) such as energy-dispersive X-ray spectroscopy (EDX)\(^{158}\), X-ray photoelectron spectroscopy (XPS)\(^{46,147}\), nitrogen, carbon and sulfur (NCS) elemental analysis\(^{112,137,159,160}\), solid-state nuclear magnetic resonance spectroscopy (NMR)\(^{159}\) or high-performance liquid chromatography (HPLC)\(^{161}\).

In this chapter, the potential of a spectrophotometric method to measure the concentration of CS and CRG in dilute mixed PE solutions will be investigated. The spectrophotometric method measures the change in dye absorbance with the change in polyelectrolyte (PE) concentration. It has been previously shown that small quantities between 0 to 80 µg/ml\(^{162,163}\) of chitosan could be measured in this way. However, it is yet unknown whether the CS or CRG molecules can be accurately determined in the presence of the other interfering PE within the same solution. CS and CRG concentration will be measured using oppositely-charged dyes such as cibacron brilliant red 3B-A (CBR) and methylene blue (MB), respectively. The use of dyes is necessary since the individual PEs are not detectable under UV or visible light. This change in colour then will be detected using sensitive colorimetric instruments such as a UV-Vis spectrometer or a general visible light spectrum plate reader. The accuracy and reliability of both techniques are compared and outlined in this chapter.

The primary objective of this study was to identify whether the spectrophotometric method can be accurately and reliably used to determine the concentration of unreacted CS and CRG that remained in the supernatants after the PEC reaction\(^{46,6}\). By knowing the amounts of PEs present in the supernatant, the composition of the remaining non-dilute PEC system can then be derived.
Previously, Araujo et al. (2014) investigated the composition of CS-CRG PECs using the MB dye only. The composition was found by adding the dye to the supernatants to observe the presence of any remaining unreacted CRG. However, it was assumed that no chitosan remained in the supernatant and any effect from interference between PEs was not considered. In the present study, the effect of interference between the competing PE was investigated by using two dyes as mentioned above.

The objectives of this chapter were to investigate:

1. The quantification of chitosan and carrageenan separately using oppositely-charged dyes.
2. The difference between UV-Vis spectrophotometer and plate reader to measure the absorbance values. The speed, accuracy and reliability were compared.
3. The stability of the prepared dye solutions.
4. The optimised experiment presented in the literature.
5. The polyelectrolyte-dye interactions and the quantification of PEs in mixed PE systems.

### 3.2 Quantification of Chitosan and Carrageenan

#### 3.2.1 Introduction

Solubilised CS has been quantitatively determined using Orange II, Alizarin S, Alizarin GG, Congo Red, Reactive Red 123, Reactive Yellow 14S and Bromocresol purple dyes. Some of these listed dyes posed several disadvantages. For example, Orange II dye required long mixing times with CS and the CS-dye complex needed to be centrifuged for extended periods before the measurements could take place. The use of picric acid and ninhydrin dyes required pre-hydrolysis of the CS, and the preparation was therefore too lengthy. Also, ninhydrin is known to be a harmful and irritant material. More sensitive methods have been developed to detect CS in solution. These methods include fluorescent biomarkers such as o-phthalaldehyde that can be conjugated to primary amino groups present on the CS molecules. This method gives the advantage of detecting CS in the presence of other primary amino groups, i.e. in the presence of proteins and other polyelectrolytes. However, the method requires extra preparation steps. Other fluorescent methods employed involve hydrolysis of CS with sodium nitrite, and the amine group is replaced with thiobarbituric acid dye, fluorescein isothiocyanate (FITC) conjugated and ortho-phthalaldehyde (OPA) conjugated CS nanoparticles. In this study, CBR was chosen because the dye dissolves readily in water, and the sodium ions dissociate to form four negatively-charged sulfate groups per molecule. The sulfate groups are situated on opposite sides and therefore promote the interaction with polycations. CBR is a red anionic dye that turns magenta when the sulfonic acid groups in the dye bind to the
cationic amino groups in CS. The standard curves of CBR were previously shown to be linear which makes the measurements of unknown concentrations more convenient \(^{162,163}\). Finally, CBR was also recommended specifically for the detection of CS in the presence of PEC systems \(^{162}\) but has not yet been used before in that setting.

![Figure 3.1](image1.png)

**Figure 3.1** – Molecular structure of cibacron brilliant red 3B-A (CBR). Figure obtained from Sigma-Aldrich.

CRG is usually detected with cationic dyes. The most common cationic dyes are toluidine blue (TB) \(^{46,166}\), alcian blue (AB) \(^{173}\) and methylene blue (MB) \(^{174}\). MB was chosen as the preferred option as this dye has been previously utilised for the detection of CRG in the supernatant of CS-CRG complexes \(^{6}\). MB is soluble in water and therefore does not require buffer preparations and centrifugation steps. It also does not require tedious heating-cooling cycles as previously reported \(^{174}\). The molecular structure of MB is shown below in Figure 3.2, showing the positively-charged nitrogen atom. It was previously found that the interaction between the dyes and the PEs are electrostatic \(^{175}\).

![Figure 3.2](image2.png)

**Figure 3.2** – Molecular structure of methylene blue (MB). Figure obtained from Sigma-Aldrich.
3.2.2 Materials and Methods

3.2.2.1 Quantification of CS

An original method for CS quantitation using CBR published by Muzzarelli \( ^{162} \) was slightly modified to suit the CS-CRG PEC system. HCl was used instead of lactic acid because the HCl was utilised in the formation of PECs as later shown (Chapter 4). Furthermore, the use of salt in the buffer was eliminated to avoid electrostatic interactions and charge screening of charged groups \( ^{176} \). In summary, 1 litre of the glycine-HCl buffer at 0.1 M (pH 1.5) and 100 ml of CBR (0.15 % w/v) dye solutions were prepared. Then 50 ml of CBR (0.15 % w/v) was added to 950 ml glycine-HCl buffer (0.1 M) to obtain a concentration of 0.0075 % w/v of CBR in glycine-HCl buffer solution (CBR dye buffer). CS solution of 0.05 % w/v was made using 0.05 g of CS in 99.5 ml ultrapure type 1 water with 0.5 ml of 12.2 M HCl. The standards were prepared by pipetting 0, 15, 30, 45, 60, 80, 100, 150, 200, and 250 µl of CS solution into 7 ml-bijou bottles and made up to 300 µl with the glycine-HCl buffer (0.1 M). Then, 3 ml of the dye-buffer solution was added to each concentration standard. The standards were mixed in a vortex mixer (SCILOGEX MX-S Vortex Mixer, Germany) at medium speed (approximately 1300 rpm) for 3 s at RT. A summary of the samples used for the CS standard calibration is shown in Table 3.1. The absorbances were measured at a wavelength of 570 nm with a UV-Vis spectrometer (Lambda 25; Perkin Elmer Inc., USA) set at a resolution of 1 nm and speed of 960 nm/min. All measurements were carried out in triplicate.

Table 3.1 – Production method of calibration standard solution at different CS concentrations using CBR dye. The solutions are added in sequential order.

<table>
<thead>
<tr>
<th>Sample No.:</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st Add CS 0.05 % w/v, µl</td>
<td>0</td>
<td>15</td>
<td>30</td>
<td>45</td>
<td>60</td>
<td>80</td>
<td>100</td>
<td>150</td>
<td>200</td>
<td>250</td>
</tr>
<tr>
<td>2nd Add HCl-Glycine buffer, µl</td>
<td>300</td>
<td>285</td>
<td>270</td>
<td>255</td>
<td>240</td>
<td>220</td>
<td>200</td>
<td>150</td>
<td>100</td>
<td>50</td>
</tr>
<tr>
<td>3rd Add CBR dye buffer, ml</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Final CS concn., µg/ml</td>
<td>0</td>
<td>2.3</td>
<td>4.5</td>
<td>6.8</td>
<td>9.1</td>
<td>12.1</td>
<td>15.2</td>
<td>22.7</td>
<td>30.3</td>
<td>37.9</td>
</tr>
</tbody>
</table>

3.2.2.2 Quantification of CRG

The method utilised in this study is based on a technique first reported by Michon \( ^{174} \). In summary, 1 l of 0.0015 % w/v MB solution was diluted from 0.05 % w/v stock solution. A concentration of 0.1 % w/v CRG in 100 ml water was prepared as the stock solution. Then 50 ml of 0.1 % w/v CRG stock solution was added to 50 ml deionised water to produce 0.05 % w/v CRG solution (adjusted to pH 7). The sample was then diluted further, by using 50 ml of 0.05 % w/v CRG solution to 50 ml water to produce a CRG concentration of 0.025 % w/v. The solution was then diluted six more times (Table 3.2). Then 1 ml of each CRG concentration solution was added to 9 ml of MB in a 30-ml beaker. The
standards were mixed in a vortex mixer (SCILOGEX MX-S Vortex Mixer, Germany) at medium speed for 3 s at 25 °C. Subsequently, the optical densities of the standards were measured at 554 nm, 615 nm, and 663 nm using a UV-Vis spectrometer (Lambda 25; Perkin Elmer Inc., USA) set at a resolution of 1 nm and speed of 960 nm/min. All measurements were carried out in triplicate.

Table 3.2 – Production method of calibration standard solution at different CRG concentrations using MB dye. The solutions are added in sequential order.

<table>
<thead>
<tr>
<th>Sample No.</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st Add MB solution, ml</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td>2nd Add 1 ml of CRG, (X) % w/v</td>
<td>0</td>
<td>7.81 x 10^-5</td>
<td>1.56 x 10^-4</td>
<td>3.13 x 10^-4</td>
<td>6.25 x 10^-4</td>
<td>1.25 x 10^-3</td>
<td>18.75 x 10^-3</td>
<td>2.5 x 10^-3</td>
<td>3.75 x 10^-3</td>
<td>5 x 10^-3</td>
<td>1 x 10^-2</td>
</tr>
<tr>
<td>Final CRG concn. µg/ml</td>
<td>0</td>
<td>0.78</td>
<td>1.56</td>
<td>3.13</td>
<td>6.25</td>
<td>12.5</td>
<td>18.75</td>
<td>25</td>
<td>37.5</td>
<td>50</td>
<td>100</td>
</tr>
</tbody>
</table>

3.2.3 Results

3.2.3.1 Quantification of CS

Figure 3.3 shows the full visible light spectrum (400-700 nm) of CS at different concentrations. A standard curve was obtained from the absorbance values recorded at 515 nm and 570 nm (Fig. 3.4). The absorbance values of the main peak at 570 nm and shoulder peak at 515 nm were shown to increase and decrease respectively in a linear fashion with CS concentration. CBR changed colour from red to magenta during the interaction between the dye and the polycation.

Figure 3.3 – Full visible-light spectrum of different CS concentration obtained by UV-Vis spectroscopy over a wavelength range of 400-700 nm. The legend on the right shows the concentration of CS (µg/ml) in ascending order.
Figure 3.4 (a) – CS concentration standard curve at $A_{515}$. Mean ± SD, $n = 3$.

Figure 3.4 (b) – CS concentration standard curve at $A_{570}$. Mean ± SD, $n = 3$. 
3.2.3.2 Quantification of CRG

Figure 3.5 shows the spectrum obtained from 400-700 nm at different concentrations of CRG. A high absorbance value at the wavelength of 663 nm corresponds to a low concentration of CRG in solution. In Figure 3.6, three CRG standard curves are presented based on three different wavelengths. All wavelengths can be used for the detection of CRG concentration. The MB changed colour from blue to purple immediately after the addition of CRG. The CRG concentration could only be measured until 60 µg/ml before the absorbance values plateaued.

Figure 3.5 – Full visible-light spectrum of different CRG concentration obtained by UV-Vis spectroscopy over a wavelength range of 400-700 nm. The legend on the right shows the concentration of CRG (µg/ml) at descending order.
Figure 3.6 – CRG concentration standard curves at wavelengths of (a) 663 nm (b) 615 nm and (c) 554 nm. Mean ± SD, n = 3. It is assumed that the best fit is an exponential curve as previously denoted by Michon et al. (2002)\textsuperscript{174} and Araujo (2013)\textsuperscript{28}.
3.2.4 Discussion

3.2.4.1 Quantification of CS

The formation of CS-CBR complexes resulted in an increase in $A_{570}$ and a decrease in $A_{615}$, known as a bathochromic shift or red shift. The bathochromic shift is the change in colour from a short wavelength to a longer wavelength. After complex formation between CBR and polycation, the $A_{570}$ increased and the $A_{615}$ decreased obeying the Lambert-Beer law (Fig. 3.4). The Lambert-Beer law is the linear relationship between the absorbance and concentration of the absorbing species. The standard curves in this study are comparable to the ones previously reported by Muzzarelli (1998) and Wischke & Borchert (2006).

3.2.4.2 Quantification of CRG

For CRG, a decrease in $A_{663}$ & $A_{615}$ and an increase in $A_{554}$ relates to the formation of new CRG-MB complexes (metachromatic complexes). As the concentration of CRG increases, the absorbance reaches a plateau due to the depletion of the dye. Once the concentration limit was reached for CRG at 100 μg/ml, it can be assumed that all MB molecules were saturated by sulfate groups present in CRG. In the case of CRG-MB complexes, metachromatic complexes between MB molecules stabilised by water through long-range forces result in the absorbance at 554 nm. However, the absorbance at 663 nm ($\alpha$-band) is more widely used because of its greater separation points in absorbance values at different CRG concentrations. Free MB molecules result in the absorbance at 663 nm. The long-range forces can only occur when the MB molecules were close enough to each other i.e. between 0.35 and 0.7 nm. The spectrophotometric method has also been previously used to characterise the coil to helix transition by measuring the absorbance of the metachromatic complex at different temperatures. At 60 °C, the CRG chains are in a coil conformation which results in a greater separation distance (1 nm) between the sulfate groups of $\iota$-CRG molecules, and in theory, no long-range forces can form to have an absorbance at 554 nm. However, at a lower temperature (between 50 and 30 °C), the CRG molecules turn into a helix and become stiffer. This helical conformation results in the shortening of the distance between the sulfate groups in CRG (0.66 nm) and hence the formation of metachromatic complexes.
3.2.5 Conclusion

A quantitative colorimetric method for both CS and CRG were presented. The CS-CBR showed two peaks at 515 and 570 nm, whereas the peaks for CRG-MB were at 554, 615 and 663 nm. The calibration curves obtained for CS were linear, while those for CRG plateaued at high CRG content. The wavelengths of 554 nm and 663 nm were chosen for the measurement of CS and CRG, respectively. These wavelengths were selected because the absorbances were found to be very sensitive to changes in PE concentrations. The CS standard curves were linear over the concentration range (0-37 µg/ml) tested, whereas CRG could be measured from 0-60 µg/ml before absorbance value plateaued.

3.3 Comparative Study of UV-Vis Spectrometer and Plate Reader

3.3.1 Introduction

A UV-Vis spectrometer is more widely used than a plate reader for the detection of polyelectrolyte concentrations. In this study, the accuracy and reliability of the plate reader were evaluated because of its advantages over UV-Vis such as rapid analysis and small volume requirements. The aim of this sub-study was to assess the speed and accuracy of both instruments and to identify the best apparatus for spectrophotometric studies in this thesis.

3.3.2 Method

The standard curves for CS and CRG were compared using a UV-Vis spectrometer (Lambda 25, USA) and a plate reader (Spectro Star Nano, Germany). The absorbance peaks for CS and CRG were measured at 570 nm and 663 nm, respectively.

3.3.3 Results

The plate reader was a faster method than the UV-Vis spectrometer. However, the absorbance values obtained from the UV-Vis spectrometer were more than double the absorbance values obtained with the plate reader (Fig. 3.7 and 3.8). The UV-Vis spectrometer also produced more reproducible results when the measurements were taken on different days. In other words, the UV-Vis spectrometer was more reliable than the plate reader. Therefore there is a trade-off between speed and accuracy when comparing the two instruments.
Chapter 3 – Spectrophotometric Study

Figure 3.7 – CS standard curves produced using UV-Vis spectrometer and plate reader.

\[ y = 0.02x + 0.55 \quad R^2 = 0.98 \]

\[ y = 0.01x + 0.22 \quad R^2 = 0.95 \]

Figure 3.8 – CRG standard curves produced using UV-Vis spectrometer and plate reader.

\[ y = 2.01 \times e^{\frac{x}{25.3}} + 0.64 \quad R^2 = 0.97 \]

\[ y = 0.81 \times e^{\frac{x}{74.2}} + 0.18 \quad R^2 = 0.97 \]
3.3.4 Discussion

The UV-Vis spectrometer was more sensitive to measurements than the plate reader. This can be observed by the greater absorbance range found in the UV-Vis spectrometer. The higher absorbance values obtained in UV-Vis spectrometer may be due to the longer path length of the cuvette sample found in UV-Vis spectrometer. It is known from the Lambert-Beer Law that an increased light path through the cuvette increases the absorbance values \(^{178}\). The standard curves produced seem to be relatively reproducible with small error bars. This signifies that the colorimetric assay is a reliable and sensitive technique.

3.3.5 Conclusion

Since the concentrations of PEs were minuscule, the method with the highest sensitivity and accuracy was required. Therefore, it is concluded that the UV-Vis spectrometer is the preferred choice for the detection of polyelectrolytes in this thesis. Where specified, the plate reader will be used for its rapidness and where the measurement of PEs is not precisely required such as in time measurement studies presented in the next sub-study presented in Section 3.4.

3.4 Time Measurements to Study Dye Decay

3.4.1 Introduction

This study was conducted to measure the decay of dyes with time. The aim of this study was to find out whether the dye solutions needed to be prepared freshly every single time or whether the same batch of solution can be used over some days without the need to create a new standard curve.

3.4.2 Method

The absorbance of dye-polyelectrolyte complexes was measured on three different days using the same dye stock solution produced from the same batch. The time points used for CS were 1, 5 and 15 days and for CRG were 1, 7 and 17 days. Between measurements, the dye solutions were kept at 4 °C and covered with aluminium foil to prevent decay. Absorbance values were determined using a plate reader.

3.4.3 Results

The absorbance values are shown for CS (Fig. 3.9) and CRG (Fig. 3.10) as a function of PE concentration value. Both dyes decayed with time, illustrated by the shift of the standard curves to lower absorbance values with time. However, the gradients of each graph were unchanged with time.
Figure 3.9 – Original standard calibration curves at different time points to observe decay of CBR 3B-A dye.

Figure 3.10 – Original standard calibration curves at different time points to observe decay of MB dye.
3.4.4 Discussion

Figures 3.9 and 3.10 indicate that a new standard calibration curve is required for every single time point which supports the data reported by Soedjak (1994) \(^{175}\). Therefore, the best option is to obtain the supernatant samples and conduct the measurements together using the plate reader to avoid time-colour change discrepancy. It was previously reported that MB decomposes very slowly in the aqueous solution \(^{179}\). The decay of the dye with time may be due to the conversion of MB to methylene violet by hydrolysis of dimethylamino groups \(^{179,180}\). Although the dye solutions were covered with aluminium foil, some illumination may have resulted in photo-reduction of MB dye \(^{179}\). Photo-reduction may occur by receiving one or more electrons by the photo-excited species \(^{181}\). It may be that a similar photo-reduction mechanism was occurring for CBR. Overall, both dyes showed only little decay with time, which may be due to the low temperature and light conditions used during storage.

3.4.5 Conclusion

The data showed that the calibration curves changed with time due to the ageing of the dye solutions, even under appropriate storage conditions. Therefore, new standard curves should always be produced when measuring absorbance values at different time points.

3.5 Confirmation of Wischke and Borchert’s Study

3.5.1 Introduction

It was previously claimed by Wischke and Borchert (2006) that the sensitivity and the detection limit of the technique carried out with CBR to detect CS can be increased by reducing the volume of CBR-buffer solution used \(^{163}\). The aim of this sub-study was to confirm this claim and whether the result could be reproduced.

3.5.2 Method

Various volumes of 0.05 % w/v CS solutions were made up to 300 µl using the buffer solution as outlined before in Section 3.2.2.1. This mixture was then mixed with CBR-buffer solutions as used in the earlier section or reduced level of 1 ml and 2 ml. The absorbance was measured using a plate reader.

3.5.3 Results

Figure 3.11 shows the effect of CBR volume on the formation of standard curves. The sensitivity
increases with decreasing CBR-buffer volumes used. However, the reduction in dye volume also resulted in a lower detection limit of the CS concentration due to the loss of linearity in absorbance measurements beyond 12 µg/ml.

Figure 3.11 – Effect of CBR volume on standard curves produced. Using different dye volumes of CBR-buffer at 0.075 % w/v.

3.5.4 Discussion

Figure 3.11 shows that the 3 ml CBR-buffer solution used by Muzzarelli (1998) provided a linear gradient across the wider concentration range. This result is contrary to that reported by Wischke and Borchert (2006) where the data indicated that the concentration limit could be increased to 100 µg/ml by using lower dye volumes. While slightly more sensitive at lower dye volumes, in these results the signal plateaued at a lower CS concentration range. Unfortunately, the method reported by Wischke and Borchert (2006) could not be reproduced in this study.

A possible explanation for the plateau at low dye volumes is that the number of dye molecules present was not sufficient to interact with a large number of CS molecules, and therefore, the dye molecules of CBR were saturated early with CS. In other words, smaller dye volumes increased the sensitivity, but the concentration detection limit was reduced.

3.5.5 Conclusion

In conclusion, the method proposed by Wischke and Borchert (2006) to increase the sensitivity and CS detection range could not be reproduced. At higher CBR dye volumes, the sensitivity reduces and the concentration detection range increases. For this thesis, 3 ml dye volume was chosen as the best method for a relatively sensitive measurement combined with a high concentration detection range.
3.6 Polyelectrolyte-Dye Interaction

3.6.1 Introduction

In earlier reports, it was found that the addition of interfering compounds such as acids, sugars, salts, milk, proteins, dyes, emulsifiers and neutral hydrocolloids may be diluted out to prevent the interference with the absorbance measurements of CRG-MB complexes. This suggests that the MB dye can be used in mixed PE systems and may therefore be used to measure the concentrations of PEs in supernatants in the presence of oppositely-charged PEs. This study aims to test the claim above and to deduce the competitive interactions between CS and CRG with either CBR or MB dyes. The competitive binding of MB with other cationic PEs towards CRG has been studied before using MB reacted with iota-CRG/poly-L-lysine or iota-CRG/gelatin or pectate/poly-L-lysine. In all the studies, it was shown that the addition of the opposing PE to the MB-PE complex increases the absorbance at 665 nm and decreases the absorbance at 570 nm. On the other hand, the use of CBR with other PEs is relatively new in PECs and therefore no competitive binding assays have been carried out before. Nevertheless, the interaction between CS and CRG have never been tested before by either MB or CBR. Therefore, the objective of this study was to observe whether concentration measurements were affected by interference from complex formation between the dyes (CBR and MB) and the oppositely-charged PEs (CS and CRG). Ultimately, this study may provide information as to whether the use of a particular dye can determine the concentration of a specific PE in the presence of an oppositely-charged PE in the supernatant.

3.6.2 Addition of CRG to CS-CBR

3.6.2.1 Method

Competitively charged dyes were used to examine the interaction between CS and CRG at different PE concentrations. PE interactions were carried out at two different concentrations. The method is described briefly below. All measurements were performed in triplicate.

The samples were prepared as described in Section 3.2.2, with the exception that 50 µl of 0.05 % w/v or 0.5 % w/v CRG were added to have a CRG concentration of 7.58 µg/ml or 75.76 µg/ml, respectively. The method was carried out in the following order: different volumes of CS 0.05 % w/v were pipetted and made up to 250 µl with HCl-glycine buffer (CBR-dye buffer). Next, 3 ml CBR dye-buffer solution was added to each sample, and finally, 50 µl of either 0.05 % w/v or 0.5 % w/v of CRG were added to the mixture. Thus, a chance was given for competitive interaction to occur between CRG and CBR with CS. In other words, the dye and the oppositely-charged PE were added together first. Then the
competitive PE was added. In practice, the situation will be the other way in that the two PEs will be present before the dye is added (Fig. 3.12). Therefore, the competitive binding may be even greater in the supernatant system. If in the first experiment, the CRG displaces the CBR dye, then it is very likely that the CBR will not displace the CRG molecules found in the supernatant. Table 3.3 shows a detailed outline of the amounts and addition order used for each standard solution produced.

Table 3.3 - The content used to produce the standard solutions CS-CBR with CRG as a competing PE at different concentrations. For 7.58 or 75.76 µg/ml CRG (competition method). The solutions are added in sequential order.

<table>
<thead>
<tr>
<th>1st Add</th>
<th>CS 0.05 % w/v, µl</th>
<th>0</th>
<th>15</th>
<th>30</th>
<th>45</th>
<th>60</th>
<th>80</th>
<th>100</th>
<th>150</th>
<th>200</th>
<th>250</th>
</tr>
</thead>
<tbody>
<tr>
<td>2nd Add</td>
<td>HCl-Glycine buffer, µl</td>
<td>250</td>
<td>235</td>
<td>220</td>
<td>205</td>
<td>190</td>
<td>170</td>
<td>150</td>
<td>100</td>
<td>50</td>
<td>0</td>
</tr>
<tr>
<td>3rd Add</td>
<td>CBR dye buffer, ml</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>4th Add</td>
<td>CRG 0.05 % w/v or 0.5 % w/v, µl</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Final CS concn., µg/ml</td>
<td>0</td>
<td>2.3</td>
<td>4.5</td>
<td>6.8</td>
<td>9.1</td>
<td>12.1</td>
<td>15.2</td>
<td>22.7</td>
<td>30.3</td>
<td>37.9</td>
<td></td>
</tr>
</tbody>
</table>

3.6.2.2 Results

Figure 3.13 shows the standard curves of the interaction between CS-CBR-CRG at two different CRG concentrations. As can be observed from $A_{570}$ (Fig. 3.14), the addition of 7.58 µg/ml of CRG to the CS standard curve did not have a significant effect on the interaction between CS and CBR. However, the addition of 75.76 µg/ml lowered the absorbance after reaching a CS concentration of 12 µg/ml. As the amounts of CS and CRG increase, the interference between CRG and CS-CBR complex becomes greater.
Figure 3.13 (a) - Full visible-light spectrum of CS-CBR absorbance with 0 % w/v CRG. The legend on the right shows the concentration of CS (µg/ml).

Figure 3.13 (b) - Full visible-light spectrum of CS-CBR absorbance with 0.05 % w/v CRG. The legend on the right shows the concentration of CS (µg/ml).

Figure 3.13 (c) - Full visible-light spectrum of CS-CBR absorbance with 0.5 % w/v CRG. The legend on the right shows the concentration of CS (µg/ml).
3.6.2.3 Discussion

The change in absorbance during the addition of the CRG to the CS-CBR complex may have been due to the dissociation of the CS-CBR (bathochromic complexes) and the formation of CS-CRG complexes. The dissociation results in the release of free CBR dyes into the medium. Therefore the affinity of CS-CRG binding is stronger than that found in CS-CBR complexes (Fig. 3.15).

On the other hand, the results showed that the addition of small concentrations of CRG to the CS-CBR complex did not change the absorption values significantly. Therefore, it is likely that concentrations of CS in the supernatant can be adequately quantified when a small concentration of interfering CRG is present. When necessary, the supernatant may be diluted to a low concentration range to access the non-interfering linear region.

Figure 3.15 - A representation of the competition between negatively-charged carrageenan (CRG) and negatively-charged cibacron brilliant red (CBR) for the positively-charged chitosan (CS). At very high CRG concentration, a small number of dye molecules are displaced, producing a shift in absorbance.
3.6.3 Addition of CS to CRG-MB

3.6.3.1 Method

The samples were prepared as outlined in Section 3.2.2 with the difference that 0.1 ml of 0.05 % w/v or 0.5 % w/v CS was added to 8.9 ml of MB solution. Then, 1 ml of CRG at different concentrations was added to the previous mixture. Therefore, the amount of CS present in 10 ml CRG-MB solution was either 5 µg/ml or 50 µg/ml. If in the first experiment, the CS displaces the MB dye, then it is very likely that the MB will not displace the CRG molecules found in the supernatant (Fig. 3.16). Table 3.4 shows a detailed outline of the amounts and addition order used for each standard solution produced.

Figure 3.16 – The methods above are showing the steps involved in adding the reagents followed by absorbance measurements using the UV-Vis spectrometer. In the competitive binding experiment here (shown left), CRG is added to MB first before adding the CS molecule as the competing PE. In the PEC manufacture experiment (shown right), CBR is added to the CS-CRG found in the supernatant. If the CS displaces the MB dye in the first experiment, then it is very likely that the MB will not displace the CS molecules found in the supernatant.

Table 3.4 – The content used to produce the standard solutions of CRG-MB with CS as a competing PE at different concentrations. For 5 or 50 µg/ml CS (competition method). The solutions are added in sequential order.

<table>
<thead>
<tr>
<th>1st Add</th>
<th>MB solution, ml</th>
<th>8.9</th>
<th>8.9</th>
<th>8.9</th>
<th>8.9</th>
<th>8.9</th>
<th>8.9</th>
<th>8.9</th>
<th>8.9</th>
<th>8.9</th>
</tr>
</thead>
<tbody>
<tr>
<td>2nd Add</td>
<td>1 ml of CRG, (X) % w/v</td>
<td>7.81 x 10^-5</td>
<td>1.56 x 10^-4</td>
<td>3.13 x 10^-4</td>
<td>6.25 x 10^-4</td>
<td>1.25 x 10^-3</td>
<td>18.75 x 10^-3</td>
<td>25 x 10^-3</td>
<td>37.5 x 10^-3</td>
<td>5 x 10^-3</td>
</tr>
<tr>
<td>3rd Add</td>
<td>CS 0.05 % w/v or 0.5 % w/v, ml</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
</tr>
<tr>
<td>Final CRG concn., µg/ml</td>
<td>0.78</td>
<td>1.56</td>
<td>3.13</td>
<td>6.25</td>
<td>12.5</td>
<td>18.75</td>
<td>25</td>
<td>37.5</td>
<td>50</td>
<td>100</td>
</tr>
</tbody>
</table>

3.6.3.2 Results

The spectra of CRG standard curves with CS are shown below in Figure 3.17. The absorbance values of all the conditions measured at 663 nm are presented in Figure 3.18. When a low concentration of CS (5 µg/ml) was added to the CRG-MB complex, the absorbance at 663 nm remained at the same level.
of the MB-dye solution. When a higher concentration of CS (50 µg/ml) was added, the absorbance was slightly changed due to the interference from the competing PE.

![Figure 3.17](a) - Full visible-light spectrum of CRG at a CS concentration of 0 % w/v. The absorbance of various concentrations of CRG was measured at 663 nm. The legend on the right shows the concentration of CRG in µg/ml.

![Figure 3.17](b) - Full visible-light spectrum of CRG at a CS concentration of 0.05 % w/v. The absorbance of various concentrations of CRG was measured at 663 nm. The legend on the right shows the concentration of CRG in µg/ml.

![Figure 3.17](c) - Full visible-light spectrum of CRG at a CS concentration of 0.5 % w/v. The absorbance of various concentrations of CRG was measured at 663 nm. The legend on the right shows the concentration of CRG in µg/ml.
3.6.3.3 Discussion

The addition of CS to CRG-MB complexes increased $A_{554}$ and decreased $A_{663}$ corresponding to a decline in metachromatic complexes and release of free MB molecules in the medium (Fig. 3.17). The results showed that the addition of small amounts of CS (5 µg/ml) reduced the interaction between CRG-MB. This was despite the fact that the CRG and MB molecules were added together before the CS was added. This showed that the interaction between CS-CRG was stronger and displaced the interaction between CRG-MB (Fig. 3.19). A possible explanation for this may be due to the positioning of the positively-charged groups present on CS and MB. The amino group in CS is positioned outwards and therefore increasing the potential for interaction. The sulfur group in MB is slightly positive which reduces the degree of interactions. Finally, the flexibility of the PE chains are higher than in the dye molecules and therefore the interaction between the PE-PE molecules is higher than in the PE-dye molecules.
As the amount of competing PE was increased, the interference resulted in the dissociation of the PE-dye complex. The effect was greater in CRG-MB assays, where the addition of CS at 50 µg/ml completely dissociated the interaction between CRG and MB. Similarly, but less affected, the CS-CBR bond separated in favour of adding CRG at 75.76 µg/ml to form a CS-CRG bond. Therefore, the interaction between PEs was significantly stronger than the interaction between the PE and the dye. As a result, it can be predicted that the unknown CRG concentration may not be identified accurately within the supernatant in the presence of a low concentration of oppositely-charged CS. It is projected that the spectrophotometric method will underestimate the amount of PE present in the supernatant. Hence, for this thesis, it is important to have another assay that can determine the concentration of CRG.
3.7 Overall Conclusions

It has been of interest to develop a protocol to measure the composition and interaction of PECs. The spectrophotometric method was effective in detecting small concentrations of PEs. CS and CRG can be measured separately with high sensitivity and accuracy. The UV-Vis spectrometer was more reliable, sensitive and accurate than the plate reader. However, the plate reader can measure small sample volumes and provide rapid measurements. The dye solutions were found to be affected by some decay with time. Therefore, sample measurements need to be conducted together to provide a more reliable assay. The reportedly improved method devised by Wischke and Borchert (2006) for the detection of CS could not be reproduced, and a higher CS concentration detection limit was obtained when greater volumes of CBR-dye buffers were used. Finally, it was found that the addition of interfering PEs at high and in some cases at low concentrations resulted in the interference between the PE-dye complex. It would be likely that the exact presence of PEs in the supernatant can be underestimated. Therefore, other techniques used for the determination of PEC composition should be considered.
Chapter 4 – Polyelectrolyte Complex Gels

4.1 Introduction

The definition of a gel is still a matter of debate, but according to Peppas, gels are defined as macromolecular networks swollen in water or biological fluids. PECs can be categorised into two types of gels; a complex coacervate gel (liquid with a dynamic structure like a particle dispersion) and a fibrous precipitate gel (concentrated viscoelastic polymer solution with a pseudo-plastic behaviour). When the interaction between PEs is weak or strong, homogeneous complex coacervate gels or heterogeneous fibrous precipitate gels are formed, respectively.

This chapter concerns the preparation and characterisation of polyelectrolyte complexes between CS and CRG by using different pH conditions. In previous studies, pH was found to be an important parameter affecting the complex formation between polyelectrolytes, where at least one of the macromolecules is a weak PE type (only stable in solution at narrow pH range). The pH can significantly influence the dissociation of the weak PE, and thereby modify the strength of electrostatic interaction (SEI) between PEs. This phenomenon is attributed to the acid dissociation constant (pKₐ) of the PEs. The pKₐ can be used to describe the strength of acids. The lower the pKₐ value, the stronger the acid. The pKₐ of carrageenan is 2, and the sulfate groups dissociate to become negatively-charged when the pH is higher than the pKₐ value. Hence, the pH range for CRG deprotonation is large and therefore the material is considered to be a strong PE i.e. the polyanion attains its fully ionised form in a wide pH-range. In contrast, the pKₐ of the glucosamine segments in chitosan is 6.5, and the material becomes positively-charged when the pH of the solution is below the pKₐ value. Therefore, the pH range for the protonation of CS is relatively narrow (< pH 6.5) and considered a weak PE.

The SEI of PEC gels can be assessed in various ways i.e. physically or chemically. Simple visual assessment can provide a good estimate of the physical properties of the gels but is not adequate for comparison. The potential difference (zeta potential) of PEs can also be measured quantitatively using a zetasizer, and the product of the zeta potential (ζ-potential) values of both molecules at each pH can be obtained to predict the SEI. Rheology measurements on PEC gels can be used to provide a direct quantitative comparison of the SEI. It is envisaged that the pH will significantly affect the SEI and therefore the viscosity of PEC gels. These results will inform the choice of fabrication pH and will provide a platform from which PEC-based films and scaffolds can be developed. The production and characterisation of films and scaffolds will be discussed in Chapter 5 and Chapter 6, respectively.
The objectives of this chapter were to investigate:

1. The charge density of CS and CRG at different pHs using zeta potential measurements to predict the strength of electrostatic interaction in PECs.
2. The preparation of CS-CRG PEC gels at different pHs.
3. The morphology of PEC gels using optical microscopy.
4. The yield and moisture content of CS-CRG PEC gels.
5. The effect of pH on PEC formation to control the degree of self-crosslinking and ultimately the viscoelastic properties of the gels using rheology.
6. The effect of pH on the resulting composition of the PECs using nitrogen, carbon, and sulfur (NCS) elemental analysis.
7. The confirmation of the electrostatic interaction between PEs using Fourier-transform infrared spectroscopy (FTIR) and X-ray diffraction (XRD).

4.2 Materials and Methods

4.2.1 Materials

Low molecular mass chitosan (72 mPa.s, 1 % w/v in 1 % v/v acetic acid at 25 °C) with 93.1 % degree of deacetylation (DDA) and κ-carrageenan (6.4 mPa.s, 0.3 % w/v in H₂O at 25 °C) were purchased from Sigma-Aldrich, UK. The chitosan was sourced from Pandalus borealis (cold water shrimp) shell in Iceland. The carrageenan was sourced from Chondrus crispus (red seaweed) in the Philippines. Hydrochloric acid (HCl) and sodium hydroxide (NaOH) were also purchased from Sigma-Aldrich, UK. All other chemical reagents were of analytical grade unless stated otherwise.

4.2.2 Thermogravimetric Analysis

TGA measurements were carried out in a TA Instruments Q600 (TA Instruments, England). The samples (2-4 mg) were placed in platinum pans and heated from 25 to 700 °C at a heating rate of 20 °C min⁻¹ and an air flow of 60 ml min⁻¹. This technique was used to analyse the humidity and degradation temperature of the materials and the amount of impurities present in PEs and PECs. The percentage impurities could then be accounted for during the preparation of the PECs. The weight loss of the materials is typically associated with the release of adsorbed moisture content and the actual degradation of the polymeric chains. The mass of the materials was considered 100 % at the onset of degradation. In other words, the amount of impurities in the samples were normalised and the average of three measurements was taken for each sample.
4.2.3 Ζeta Potential

The ζ-potential of CS (0.1 % w/v) and CRG (0.1 % w/v) were determined with a Zetasizer Nano ZS (Malvern Instruments Ltd., Malvern, UK) at 25 °C. A stock solution of 500 ml of each polyelectrolyte solution was made. Experiments were performed at pH values ranging from 2-12 at one pH-unit interval with pH resolution of ± 0.01 units. A wide range of pH values was tested to assess the possible interaction window and the strength of the electrostatic interaction was calculated (\(\zeta_{CS} \times \zeta_{CRG} = mV^2\)) according to Weinbreck et al. (2004)\(^{115}\). The pH of solutions was adjusted to the required level by the addition of 0.1, 1 and 5 M NaOH or 0.1, 1 and 2 M HCl. At each pH, the PE solution was allowed to reach equilibrium for a minimum of 60 s. Aliquots of 10 ml of the solution at each pH were retrieved and used for the ζ-potential measurements. The ζ-potential of the polyelectrolyte complexes was determined by diluting the PEC to a solid content of 0.1 % w/v. The PEC solid content was initially determined using Equation 4.9 (Section 4.2.8).

4.2.4 Molar Ratio Calculation

The polysaccharides used in this study possess the charged groups NH\(_3^+\) and OSO\(_3^-\) for CS and CRG, respectively. The number of charged groups present in each monomer unit was calculated from their corresponding molecular formulae. It was estimated that 1 mole of NH\(_3^+\) was present in 165 g of CS taking into account the degree of deacetylation (93.1 %) provided by the manufacturer. Similarly, it was calculated that 1 mole of OSO\(_3^-\) was present in 384 g of κ-CRG by considering one charged group per disaccharide unit. The amount of CS and CRG required to produce 5 g of PEC at CS-CRG 1:1 molar ratio, the following calculation was used:

\[
\frac{[NH_3^+]}{[OSO_3^-]} = 1
\] (4.1)

Since there is only one charged group per monomer unit in each polysaccharide, the molecular mass per charge in CS is 165 g/mol and in CRG is 384 g/mol:

\[
\frac{\text{The molecular mass per positive charge in CS}}{\text{The molecular mass per negative charge in CRG}} = \frac{165}{384} = 0.43
\] (4.2)

Required amounts of PEC produced was 5 g:

\[
CS + CRG = 5 \text{ g}
\] (4.3)

Combining (4.2) and (4.3):

\[
\frac{CS}{5 - CS} = 0.43
\] (4.4)
The impurities of CS and CRG was found with TGA analysis and the final amounts of CS and CRG required to produce a 1:1 molar ratio was calculated as below:

\[
\frac{1.5}{100 \% - \text{Impurity}\%} = CS \text{ g required to add} \quad (4.5)
\]

and

\[
\frac{3.5}{100 \% - \text{Impurity}\%} = CRG \text{ g required to add} \quad (4.6)
\]

### 4.2.5 Preparation of PEC Gels

CS and κ-CRG PEC gels were prepared based on the method first reported by Araujo et al. (2014) and a diagram of the experimental setup is shown below for visual representation (Fig. 4.1). For a unit molar ratio (UMR) of 1:1 CS-CRG, approximately 1.53 g of CS (0.38 % w/v) and 4.40 g of κ-CRG (0.63 % w/v) were dissolved in 400 ml of 0.16 M HCl and 700 ml of ultrapure type 1 water respectively for 24 hours under vigorous magnetic stirring. The pH of the CS solution was increased from 2 to 4 using approximately 10 ml of 5 M NaOH. The κ-CRG solution was heated to 60 °C before the mixing reaction using a water bath. The CS solution was kept at 20 °C and added to the CRG solution in a drop-wise fashion at a rate of 14 ml/min. The polyelectrolyte reaction was carried out inside a 2 l Pyrex media bottle (Fisher Scientific, UK) with vigorous stirring using both a magnetic flea and overhead stirrer at 5000 rpm. After complete addition of CS, the mixture was cooled using a water bath at RT for 15 min, followed by pH measurements. The pH of the polymer solutions and the PEC mixtures were measured with a pH-meter (Thermo Scientific Orion 3-star Benchtop, UK) at RT. This process was repeated to obtain PEC mixtures prepared at pH 3-12 at 1-unit intervals. The polyelectrolyte mixture was then poured into 50 ml centrifuge tubes and centrifuged (Hermle LaborTechnik Z300, Germany) at 4500 rpm (3222 x g) for 2 min and washed twice with ultrapure type 1 water to remove the dissociated counterions and unreacted CS and CRG. This washing and centrifuging protocol were then repeated. Subsequently, the supernatant was decanted, and the PEC gel was homogenised at 13.500 rpm for 1 min using a medium-sized homogenizer (VR, VDI 25, UK) while cooling using an ice box. The homogenised PECs were then centrifuged again to remove the excess water. All homogenised PECs were stored at 4 °C until further use. The PEC gels were photographed for visual assessment using a Canon EOS 5D Camera, Japan.
Figure 4.1 – A schematic diagram showing the preparation of PEC gels and final medical materials.
4.2.6 Optical Microscopy

Optical micrographs were taken using bright field illumination with an optical microscope equipped with dry lenses (GX Microscopes, L2000B HTG, UK) and a digital camera (Moticam, 3.0 MP, China). The PEC gels were gently mixed before analysis to ensure that they were homogeneous. A drop of PEC gel suspension (non-homogenised) was placed on a microscope glass slide and covered with a glass coverslip (22 mm diameter x 0.17 mm thickness) by ensuring no air gap is trapped between the sample and the coverslip. The scale of the micrographs was calibrated using a 1 cm graticule with 100 µm intersections.

4.2.7 Yield of PEC Gels

After PEC gel preparation, the total PEC mass was measured using an analytical balance (Sartorius, BP61 model, Germany) with an accuracy of ± 0.1 mg. Three samples of approximately 0.1-0.3 g were weighed before and after freeze drying. The samples were freeze-dried at -20 °C and 80 mTorr for 17.5 hours until completely dry, as was confirmed by TGA. The PEC yield was determined by mass using the following equation:

\[
Yield, \% = \frac{m_d}{m_i} \times 100
\]  

(4.7)

where \( m_i \) is the total pure PE powder mass used to make the PE solutions i.e. 5 g, and \( m_d \) is the mass of the dried PEC.

4.2.8 Moisture, Solid Content and pH of PEC Gels

The moisture content of PECs was analysed using the following equation:

\[
Moisture \, Content, \% = \frac{m_w - m_d}{m_d} \times 100
\]  

(4.8)

where \( m_w \) is the mass of the wet PEC sample and \( m_d \) is the mass of the dry PEC sample after freeze-drying. The final solid content (wt. %) of the centrifuged polyelectrolyte complexes was determined using the following equation:

\[
PEC \, Solid \, Content, \% = \frac{m_d}{m_w} \times 100
\]  

(4.9)

The results of the solid content of PECs were used for the production of solvent-cast films and can be found in Chapter 5 – PEC Films. The pH of the PEC gels before and after washing steps was measured when the PEC gel was in a solution property. Type 1 water was added to PEC gels to turn PEC gels into PEC solutions.
4.2.9 Nitrogen, Carbon and Sulfur Elemental Analysis

The composition of the freeze-dried complexes and individual materials was determined by measuring the amount of nitrogen present in PECs. The elements were measured using nitrogen, carbon, sulfur (NCS) element analyser (FlashEA 1112, Thermo Fischer Scientific, Italy). The sensitivity of the machine to sulfur concentration was low. Hence, only the % nitrogen data was used to determine the composition of the final complex. The instrument was calibrated using a sulphanilamide standard. Approximately, 1 to 2 mg of lyophilised samples for 17.5 hours at 80 mTorr were measured using a microbalance (Sartorius, Cubis-MSA, Germany) with a sensitivity to ± 0.1 µg. The materials were placed inside tin container catalysts and heated to 980 °C. The materials were combusted, and the elements present were converted into simple gases (CO₂, N₂ and SO₂). The gases were then homogenised, and the system controlled the exact conditions (pressure, temperature and volume). A column was used to separate the homogenised gases in a stepwise steady-state manner to detect the elements as a function of their thermal conductivities. The machine identified the total N and C to the precision of 0.1 µg. The NCS analysis measurements were undertaken by James Rolfe at the Department of Earth Science, University of Cambridge. The raw data of N measured by the machine was presented in mass % and were subsequently converted into mol % using the following equations:

\[
M_{CRG}^{PEC} = \frac{\%CS_p - \%PEC}{\%CS_p} \times M_t \tag{4.10}
\]

\[
M_{CS}^{PEC} = M_t - M_{CRG}^{PEC} \tag{4.11}
\]

\[
MolCRG_{PEC} = \frac{M_{CRG}^{PEC}}{CRG_{mwt}} \tag{4.12}
\]

\[
MolCS_{PEC} = \frac{M_{CS}^{PEC}}{CS_{mwt}} \tag{4.13}
\]

where \(M_{CRG}^{PEC}\) is the mass of CRG in PEC and \(M_{CS}^{PEC}\) is the mass of CS in PEC. The % CS₂ and % PEC denote the % N found in pure CS and the PEC samples, directly from the NCS analysis machine, respectively. \(M_t\) is the total pure mass of CS and CRG used (5 g). CRG\(_{mwt}\) and CS\(_{mwt}\) represent the molecular mass per disaccharide unit for both CS (165 g/mol) and CRG (384 g/mol), respectively. Finally, the molar ratio of CS and CRG were presented as \(MolCS_{PEC}\) and \(MolCRG_{PEC}\), respectively.

4.2.10 Fourier Transform Infrared Spectroscopy

The PEC gels produced at different pHs were freeze-dried for 17.5 hours at 80 mTorr before analysis. The PECs were analysed using an FTIR instrument (Spectrum 100 Perkin Elmer, USA) with attenuated total reflectance using a zinc selenide (ZnSe) crystal. The absorbance values of CS and CRG were
analysed independently. The non-complexed mixture of CS and CRG were also analysed and compared with the PECs to confirm complex formation by electrostatic interaction. Electrostatic interaction of PECs was confirmed using the absorbance values at 1592 cm\(^{-1}\) and 1216 cm\(^{-1}\). The data were averaged over 16 scans, using a resolution of 8 cm\(^{-1}\), in the spectral region of 1700 – 600 cm\(^{-1}\). The complexes were also confirmed after homogenisation to determine whether the homogenisation process has disrupted the electrostatic interactions.

### 4.2.11 X-Ray Diffraction

A vertical diffractometer (PW 1830P, Philips, Netherlands) was used to determine the crystallinity and identify possible crystalline contaminants and confirm complex formation. The individual PEs were analysed using a step size of 0.05° and a 4 s dwell time from 5-80°, while the PECs were analysed from 5-50°. The voltage and current used were 30 kV and 30 mA, respectively. Measured diffraction patterns were compared against powder diffraction files from the International Centre for Diffraction Database (ICDD) using PANalytical Highscore Plus software.

### 4.2.12 Rheology

Rheological measurements of PEC gels were performed to measure the SEI between PEs at varying pH values directly. Controlling the self-crosslinking of PECs may potentially be valuable for controlling the mechanics and stability of films and scaffolds produced. The viscoelastic properties of PECs were determined using a rheometer (Discovery Hybrid 2.0, TA Instruments, USA). The temperature during measurement was controlled at 20 °C using a Peltier element. A parallel plate (stainless steel) geometry (0°, 40 mm) was used with a gap of 500 µm between the flat surfaces of both elements. For each sample, approximately 1 ml of PEC gel sample was used. The samples were subjected to amplitude sweep (strain sweep) of 0.1-100 % strain (up) at a frequency of 10 rad s\(^{-1}\) to determine their linear viscoelastic region (LVR). The samples were put into a humidity trap to reduce the evaporation of the water situated inside the gels. The oscillating sweep measurement was carried out over an extended angular frequency domain of 0.1-100 (up) rad s\(^{-1}\) at a strain of 1 %, which was in the linear regime for all samples (as confirmed by amplitude sweep measurements). Finally, the flow curves were measured at increasing shear rate (flow sweep or rate sweep) from 0.01 s\(^{-1}\) to 100 s\(^{-1}\) to determine the shear properties such as the change in viscosity of the materials.
4.3 Results

4.3.1 Thermogravimetric Analysis

The TGA scans of CS, CRG and CS-CRG PECs prepared at different pHs are shown in Figure 4.2. The mass loss occurred in three distinct stages. The first stage showed approximately a 10 % step-drop due to water evaporation at temperatures from 25 to 150 °C. The second stage is the rapid degradation of CS and CRG with onset temperatures of 270 °C and 220 °C, respectively. The third stage is the complete degradation of the organic matter of the sample, and only the impurities were retained within the mixture. The temperature of this stage starts at 650 °C for CS and 500 °C for CRG with around ten times more impurities present in CRG (20.51 %) compared to CS (1.96 %). The amount of impurities (presumed to be inorganic) was calculated from the mean of three different sample measurements (Fig. 4.3). The onset temperature of PEC degradation was higher for the complexes prepared at higher pH. The pH 9 PEC contained the highest impurity (5.78 %) followed by the rest of the PECs with impurities not exceeding 2.43 %.

![Thermogravimetric analysis of CS, CRG and CS-CRG PECs prepared at different pHs.](image)

![Amount of impurities present in CS, CRG and CS-CRG PECs prepared at various pHs as determined by TGA (mean ± SD, n = 3).](image)
4.3.2 Ζeta Potential

In Figure 4.4, the ζ-potential of CS and CRG are presented for pH values ranging from 3-12. The ζ-potential of CS is shown to be at a maximum (+30 mV) at pH 2-5. As the pH was increased to pH 8 (with the addition of NaOH), the ζ-potential decreased gradually towards neutrality (0 mV). Further addition of NaOH resulted in a switchover of the ζ-potential to -7 mV. The ζ-potential of CRG was found to be strongly negative and remained relatively constant throughout the pH range studied from pH 3 (-60.6 mV) to pH 10 (-68.9 mV). The lowest and highest charge density of CRG was found at pH 2 (-50 mV) and pH 11 (-93.7 mV), respectively. The PECs at 1:1 molar ratio displayed a negative ζ-potential across the full range of pHs tested.

![Figure 4.4 – ζ-potential of CS, CRG and CS:CRG 1:1 molar ratio at different pH values. Mean ± SD, n = 3.](image)

Finally, the SEI of PECs at a particular pH was predicted by multiplying the CS charge with the CRG charge measured at the same pH (Fig. 4.5). It was predicted that the interaction between CS and CRG is the strongest at pH 4 and 5.

![Figure 4.5 - The strength of the electrostatic interaction (SEI) for CS-CRG. Mean ± SD, n = 3.](image)
4.3.3 Visual Assessment of PEC Gels

The immediate observation was that the pH strongly influenced the appearance and consistency of the polyelectrolyte complex gels. The newly prepared polyelectrolyte complex gels after the washing and centrifuging protocol are shown in Figure 4.6. After phase separation, a dense white precipitate was formed at $3 \leq \text{pH} \leq 7$ and a translucent watery complex coacervate at $7.4 \leq \text{pH} \leq 12$. The supernatant produced across the pH range showed very low turbidity suggesting that there might have been small amounts of unreacted PE and non-centrifuged PEC remaining.

Figure 4.6 – Visual assessment of CS-CRG PECs at pH 3-12 after washing and centrifugation protocol (non-homogenised).
4.3.4 Optical Microscopy

The optical micrographs in Figure 4.7 show that the PEC gels exhibited random fibrous structures. The PECs at low pH range displayed more fibres with larger fibre diameters. These fibres were formed through the coagulation and entanglement of the reacting PEs. However, at higher pH, a large number of these fibres were dissolved back into the solution. This can be observed by a significant number of thin fibres at high pH compared to the bundled fibres at low pH. At low pH, the fibres were tightly packed and difficult to resolve under the OM. However, when the PEC gels were diluted with water, the morphology and structure of the PECs became visible.

![Figure 4.7 - Optical micrographs of PEC gels prepared at different pHs.](image)

4.3.5 Yield, Moisture and Solid Content of PEC Gels

As predicted from the ζ-potential measurements shown in Figure 4.4, the SEI was highest at the lower pH range and lowest at the upper pH range (Fig. 4.5). Figure 4.8 corroborates this result where the maximum PEC yield of 85% occurred at pH 5. At the higher pH range, the yield decreased with the lowest being at pH 8 to 12. The PEC yield from pH 7 to pH 7.4 dropped by 43%. This shows that only a slight change in pH could lead to considerable change in physicochemical properties and that the boundary limit for polyelectrolyte complexing is between pH 7 and pH 7.4. The results in this section also corroborate the visual assessment results shown in Figure 4.6. It is worth noting that the yield dropped above pH 5 as the pH started to approach the pK_a of CS. In Figure 4.9, the results show that the moisture contents of PECs at higher pHs (range 9-12) were approximately eight times larger than the moisture contents present at lower pH range (3-7) PECs. The moisture content is a different representation of the solid content (Fig. 4.10).
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Figure 4.8 – Yield of PEC gels, produced at varying pH (3-12). Mean ± SD, n = 3.

Figure 4.9 - Moisture content of PEC gels produced at varying pH (3-12). Mean ± SD, n = 3.

Figure 4.10 - Solid content of PEC gels produced at varying pH (3-12). Mean ± SD, n = 3.
4.3.6 pH Measurements of PEC Gels

After the preparation of the PECs, the pH of the PEC gel solution was measured before and after the washing step with ultrapure type 1 water. It was found that the PECs produced at moderate pH (4 < pH < 8) only changed very slightly in pH after the washing step. On the other hand, the PEC gels produced at pH 3, and 9 changed by 1 pH unit and reached pH values closer to neutrality (Fig. 4.11).

![Figure 4.11 – pH of PEC gels before and after washing steps with ultrapure type 1 water.](image)

4.3.7 Nitrogen, Carbon and Sulfur Elemental Analysis

In Figure 4.12 below, the % nitrogen and % sulfur of all candidate materials is shown in molar %. The nitrogen content, which depicts the amount of CS in the PEC, increases at higher pH. Although the interaction strength reduces with increasing pH up to pH 6 (Fig. 4.4), the composition was unchanged over this range (Fig. 4.12). The PECs produced at the higher pH range (pH 9-12), contained approximately 30 % more CS compared to PECs produced at the lower pH range (pH 3-8).

![Figure 4.12 - The effect of pH on the molar composition of PECs. The result is based on the % nitrogen by using the formulae shown in the method in Section 4.2.9. Mean, n = 3.](image)
4.3.8 Fourier Transform Infrared Spectroscopy

In this section, the FTIR spectra were used to confirm PEC formation. The FTIR spectra of the individual PEs and PECs are shown in Figure 4.13. The characteristic absorption bands for CS were at 1650 and 1587 cm\(^{-1}\) and for CRG at 1230, 914, 844 cm\(^{-1}\). The sulfate group (1230 cm\(^{-1}\)) in CRG shifted to 1216 cm\(^{-1}\) during PEC formation. Also, the band in CRG at 1630 cm\(^{-1}\) and band in CS at 1650 cm\(^{-1}\) were shifted and combined to form a stronger peak at 1634 cm\(^{-1}\) after PEC formation. At pH 8 and above no PEC formation could be confirmed, as there was no trough present at 1529 cm\(^{-1}\) and 1216 cm\(^{-1}\). The absence of complex formation was also confirmed by mixing unreacted CS with CRG powder (dark blue curve), which appeared to have the same fingerprint region as the non-complexed PECs produced at pH 8-12.

![Figure 4.13 - FTIR data of CS, CRG, CS and CRG non-complexed mixture and CS-CRG complexes (pH 3-12). Red ellipses are highlighting the wavenumbers for the confirmation of PEC formation.](image-url)
4.3.9 X-ray Diffraction

X-ray diffraction peaks are presented for both the individual PEs and the PECs (Fig. 4.14). CS exhibited two diffraction peaks at 2θ = 10.6° and 19.9°. On the other hand, CRG showed five dominant sharp peaks at different 2θ (26°, 41°, 50°, 66° and 74°). All PECs presented only one broad peak at 2θ = 22°. It can also be observed that no crystalline peaks were found in the PECs. The only small sharp peak that can be noted in the PECs was at 2θ = 29° for pH 9, which may be due to some inorganic remnants.

![Figure 4.14 – XRD patterns for CS, CRG and CS-CRG PECs produced at different pHs. The red ellipse highlights PEC formation. The red ellipse is highlighting the 2theta angle for the confirmation of PEC formation.](image-url)
4.3.10 Rheology

Rheological measurements were carried out to describe the SEI of PECs quantitatively. A reasonable pH range (3-9) was selected from the visual assessment analysis to describe the effect of pH on the viscoelastic properties. Amplitude (strain) sweeps showed that the pH influenced the viscoelastic properties of wet PECs significantly (Fig. 4.15). The storage moduli (G’) and loss moduli (G’’) indicated that the linear viscoelastic region (LVR) of all PECs were well within 1% strain. Hence a constant strain of 1% was used for the frequency sweep measurements. The PEC gels produced at higher pH range (pH 6-9) induced greater strain limit (%) than the PEC precipitates prepared at the lower pH range (pH 4-5). In all cases, the G’ was higher than the G’’ at lower strains (0.1-10%). The order of the viscoelastic properties of the PECs with pH were as follows: G’ and G’’ = pH 5 > pH 4 > pH 3 > pH 6 > pH 7 > pH 7.4 > pH 8 > pH 9 (Fig. 4.15).

The frequency sweep curves of the PECs prepared at different pH are shown in Figure 4.16. All PECs displayed a moderate continuous increase in viscoelastic properties at higher angular frequencies. The storage moduli of pH 5 (~ 10000 Pa) was in the order of two magnitudes greater than the storage moduli of PEC produced at pH 9 (~ 100 Pa). The strength of PEC interaction with respect to pH is given in order from highest to lowest elastic moduli: pH 5 > pH 4 > pH 3 > pH 6 > pH 7 > pH 7.4 > pH 8 > pH 9 (Fig. 4.16).
Flow experiments were also carried out to further quantify the effect of pH on the rheological properties of CS-CRG PECs. As the shear rate was increased, the viscosity was decreased (Fig. 4.17). However, the PECs prepared at pH 3, 4 and 5 showed viscosities two orders of magnitude higher than those found in PECs prepared at higher pH values (6, 7, 7.4, 8 and 9).
4.4 Discussion

4.4.1 Thermal Degradation

4.4.1.1 Thermal Degradation Properties of CS, CRG and PECs

Figure 4.2 showed that the degradation temperature of CS was higher than that of CRG. This will be related to the temperature of oxidation of each material and may also be influenced by the higher crystallinity of CS. The PECs which have a mixture of CS and CRG showed degradation temperatures intermediate between the extremes. This intermediate degradation temperature found in PECs may be due to the disruption of the hydrogen-bonding caused by the electrostatic interaction among the polysaccharide chains. The hydrogen bonding is also responsible for the compact CS structure. In fact, the compactness within CS was previously shown to increase at higher pH neutralisation. This was due to the self-rearrangement and formation of more ordered regions within the network as was studied by wide-angle X-ray scattering analysis (WAXS). This was also supported by the increased thermal stability of CS and chondroitin sulfate (ChS) PECs prepared at pH > 6 and between CS and heparin PECs as illustrated by the higher degradation temperature for PECs prepared at higher pH.

4.4.2 Impurities

4.4.2.1 Amount of Impurities Present in PEs

The impurities remaining within the materials were confirmed to be an inorganic salt. This can be supported by the complete degradation of the organic material within approximately 500 °C. To confirm the inorganic material, XRD was carried out on CS, CRG and their PECs. The crystalline phases of CRG matched with the ICDD database for potassium chloride (KCl). The absence of crystalline peaks in CS and PECs may have been due to the relatively small amounts of impurities (~ 2 %) as was found with TGA.

4.4.2.2 The Cause of High Salt Content in CRG

A possible explanation for the high impurity in CRG can be related to the processing methods used. CRG can be extracted from red seaweed in two possible ways: an alcohol (isopropanol) method or, a more widely used, salt (KCl) precipitation method. The former method produces a purer form of CRG but is more expensive. The latter method involves the complexing of the sulfate groups present in CRG with the potassium ions present in KCl. The use of KCl, explains the reason for the high salt content in CRG.
4.4.2.3 The Cause of Low Salt Content in CS

On the other hand, dilute NaOH is typically used to deproteinate the washed crustacean shells to produce chitin. Chitin is then further treated with concentrated NaOH to deacetylate the chitin into CS. Therefore, the impurity present in CS is likely to be small amounts of sodium chloride (NaCl).

4.4.2.4 Absence of Crystalline Phases in PECs

The absence of sharp XRD crystalline peaks in PECs signifies that the crystalline contaminants have been removed almost entirely during the washing steps. Nevertheless, there was a small exception for PEC pH 9 showing a crystalline peak at 2θ = 29°, which could be due to the increased remnant at 5.71 wt. % as was confirmed with TGA. Finally, the amount of salt impurity in each sample was taken into account during the preparation of polyelectrolyte complexes to produce a 1:1 molar ratio of amino groups and sulfate groups present in each CS and CRG monomer, respectively.

4.4.3 Nature of Interactions

4.4.3.1 The pH is one of the Strongest Factors Influencing Charge Density

The pH is widely considered to be one of the strongest factors that affect the charge density of PEs. When the charge density changes, the three-dimensional configuration and flexibility of polyelectrolytes also changes. It is widely accepted that when the ζ-potential is ≤ -30 mV or ≥ +30 mV, then the PE solution is considered stable (dispersed) due to the high repulsion of like-charged PEs. This creates an extended polymer chain that is stiff. However, when the temperature increases, the chains of the PEs are still charged but conform to a random coil-like structure. PEs that are being neutralised will turn into a random-like conformation because the charge repulsions on polymer backbones are cancelled.

4.4.3.2 The Effect of pH on the CS Charge Density

The acid dissociation constant (pKₐ) of CS has been reported to be around 6.5. When CS powder was dissolved in acid, NH₂ groups were protonated by the hydrogen ions in the acid giving rise to positively-charged NH₃⁺ groups (Eq. 4.14). This was the state when CS was dissolved in acid. In contrast, the addition of a base e.g. NaOH deprotonates the NH₃⁺ groups back to NH₂ groups which result in the precipitation of CS (Eq. 4.15) and therefore exhibiting a weak polyelectrolyte behaviour. This was the state when CS was precipitated in alkali. CS was found to be 17 % positively-charged at pH 7.4, which was found to closely match with the results previously obtained by Denuziere et al. (1996), exhibiting a positive charge of 16 % at pH 7.4. In CS, the switchover to negative values in the pH range
8-13 (Eq. 4.16) is likely due to the screening of NH$_2$ groups of CS by the excess hydroxide ions from the added alkali.

\[\text{CS} + \text{NH}_2 + \text{HCl} \rightleftharpoons \text{CS} + \text{NH}_3^+ + \text{Cl}^-\] (4.14)

\[\text{CS} + \text{NH}_3^+ + \text{Cl}^- + \text{NaOH} \rightleftharpoons \text{CS} + \text{NH}_2 + \text{NaCl} + \text{H}_2\text{O}\] (4.15)

\[\text{CS} + \text{NH}_2 + \text{NaOH} \rightleftharpoons \text{CS} + \text{NH}_2 + \text{Na}^+ + \text{OH}^-\] (4.16)

### 4.4.3.3 The Effect of pH on the CRG Charge Density

In contrast, the negative ζ-potential values of CRG are the result of dissociation of OSO$_3$K giving rise to OSO$_3^-$ in CRG molecules (Eq. 4.17). This was the state when CRG was dissolved in water. The sulfate groups of CRG can also exist in conjugated form with calcium and sodium ions. Even though the pK$_a$ value of CRG was reported to be at pH 2$^{100}$, the ζ-potential for CRG was unexpectedly still very high at pH 2 (-28 mV). Therefore, the CRG molecules showed a strong PE behaviour. Similar ζ-potential values were previously found in CRG from pH 2-7 at a concentration of 0.5 wt.%$^{109}$. However, the ζ-potential values of CRG at pH 8-12 were not reported before. Reports from previous studies only showed significant swelling in CS-CRG PEC gels at pHs 11-12$^{7,145}$. The high swelling was thought to be due to the excess negative repulsive charge of CRG, and the results match with the ζ-potential at pH 11 and 12 found in the present study. The significant increase in Na$^+$ and OH$^-$ ions may have also led to the disruption of the CRG gel helices and thereby further exposing the negatively charged groups leading to strong negative ζ-potential$^{111,145}$.

\[\text{CRG} + \text{OSO}_3^- \text{K}^+ + \text{H}_2\text{O} \rightleftharpoons \text{CRG} + \text{OSO}_3^- + \text{KOH} + \text{H}^+\] (4.17)

### 4.4.3.4 Electrostatic Interaction between CS and CRG

Equation 4.18 shows the electrostatic reaction between CS and CRG and the release of counterions in the form of salt. Formation of polyelectrolyte complexes is the result of interactions between the amino groups and the sulfate groups present in CS and CRG, respectively$^{195}$. The PEC complex in this study displayed a negative ζ-potential across all pH range. This may have been caused by the dissolution of the CRG molecules into the solvent mixture and the higher opposite charge density.

\[\text{CRG} + \text{OSO}_3^- + \text{KOH} + \text{H}^+ + \text{CS} + \text{NH}_3^+ + \text{Cl}^- \rightleftharpoons \text{CRG} + \text{OSO}_3^- + \text{HN} + \text{CS} + \text{KCl} + \text{H}_2\text{O}\] (4.18)
4.4.3.5 Calculating the Strength of Electrostatic Interaction

The strength of the electrostatic interaction (SEI) was determined by multiplying the absolute ζ-potentials for both polyelectrolytes at each pH as was first applied by Weinbreck et al. (2004) and Espinosa-Andrews et al. (2013). The SEI is a theoretical assumption of the electrostatic strength between the PEs at various pH ranges. The ζ-potential measurements showed that the attraction between PEs (SEI) is strongest at pH 3-5 and weakest at pH 8-12 as was presented in Figure 4.5. Similar results were found previously, where the interaction between chitosan (CS) and gum-arabic (GA) was the highest between pH 3.5-5 and lower at pH 2 and 6 due to the protonation of GA below pH 3 and the deprotonation of CS above pH 5. The maximum SEI of the CS-CRG gels (1900 mV^2) were almost twice as high than the SEI of the CS-GA gels (1000 mV^2) and was reflected by the stronger viscoelastic properties of the gels in the CS-CRG PEC gels.

4.4.4 Physicochemical Properties of PEC Gels

4.4.4.1 PEC Preparation Conditions

During PEC formation, the pHs were chosen to optimally charge the PEs without subjecting them to an extreme pH environment. The use of extreme pH environments can result in acidic degradation of CS and CRG. The primary degradation mechanism of CS was previously shown to be due to acid hydrolysis, while it was stable against alkali degradation. Similarly, preparation of CRG solution at a pH less than 4 should be avoided. For these reasons, the pH of CS during PEC preparation was adjusted to pH 4 before it was added to the CRG solution. In addition, the elevated temperature (60 °C) used during PEC preparation was intended to increase the solubility and reduce the viscosity of the CRG solution. CRG molecules can change from a helical to a random coil-like conformation at higher temperatures (above the average gel-sol transition temperature of 30-40 °C). This enhances the flexibility of the macromolecular backbone. The combination of the factors above was used to increase the rate of PEC interaction and to obtain the highest yield possible.

4.4.4.2 The Effect of Interaction on the Physical Properties of PECs

To obtain homogeneous PEC gels, the SEI between the PEs needs to be at an optimal level. The PEC gels can change from a heterogeneous precipitate gel to a homogeneous complex coacervate gel at high and low SEI, respectively. In this study, PEC precipitate gel fibres were formed during the drop-wise addition due to the vigorous mixing coupled with the strong interaction at pH 4. Other studies investigating CS-CRG PECs have also observed precipitate formation due to the high SEI caused by the
strong sulfate groups present in CRG\textsuperscript{143,145}. Hugerth \textit{et al.} (1997)\textsuperscript{143} produced dense water-insoluble gel precipitates by reacting CS with varying DDA (73 \%, 84 \% and 93 \%) and different CRG types (κ-, λ-, ι-). Sakiyama \textit{et al.} (1993)\textsuperscript{145} also showed PEC precipitation when CS and κ-CRG interacted in the absence of NaCl. However, the addition of salt at 5.7 wt. \% during the interaction between CS and CRG was shown to produce homogeneous PEC gels\textsuperscript{13}. This is because the added salt can screen some of the charged groups present in the PEs and therefore reduce the electrostatic interaction. The advantage of obtaining homogeneous coacervate gels is that films and scaffolds can be produced with reproducible mechanical properties. Another method previously used to form homogeneous CS-CRG PEC gels was to add CS vigorously to the CRG and allowing the mixing to continue for 5-7 days at room temperature\textsuperscript{21}. Finally, homogeneous PEC gels can be formed through homogenisation, as the process was shown to be non-disruptive to the electrostatic interaction when confirmed with FTIR (data not shown). Homogeneous PEC gels can be used to form homogeneous films and scaffolds. The viscoelastic properties of the PEC gels were later quantitatively determined using rheology. Viscosity measurements were used to provide a direct relationship with the SEI as was previously proposed by Weinbreck \textit{et al.} (2004)\textsuperscript{199}.

\textbf{4.4.4.3 The Effect of pH on PEC Yield}

The PEC yield was used to determine the effect of pH on the efficiency of PEC formation between CS and CRG. At pH 5 the interaction between CS and CRG produced the highest complex yield of 85 \%. It may be that pH 5 the interaction is optimal for polyelectrolyte complex formation. Espinosa-Andrews \textit{et al.} (2007)\textsuperscript{200} previously found a similar result, where the maximum complex yield (92 \%) between GA and CS appeared to be at pH 5 regardless of the CS concentration used (0.25, 05 and 1 \% w/w). Similarly Huang \textit{et al.} (2014)\textsuperscript{28} showed the highest complex yield between GA and CS at pH 4.5. The reasons stated for the high complex yield at pH 5 was that the charge densities of the PEs were stoichiometrically balanced, allowing for greater complexation to occur with higher complex yield\textsuperscript{134,200}. However, the ζ-potential of CS-CRG PECs was at -23 mV which does not fully support this notion, unless the PEC was surrounded by excess CRG molecules which resulted in the formation of a negative charge which was previously proposed for chitosan-dextran sulfate PEC colloids\textsuperscript{110}. The low yield and high moisture content of PEC gels produced at high pH are the product of the neutralised CS. When CS becomes deprotonated, CS-CRG PEC gel dissolves which reduces the yield. Also, the strongly negatively-charged CRG increases the attraction of water molecules, and ultimately the moisture content in the PEC gels\textsuperscript{109}.
4.4.4.4 The Effect of Centrifugation on PEC Yield

The centrifugation step also played a significant role on the PEC yield produced. The yield was expected to be higher when higher speeds and longer centrifugation times were used. However, it was found that the centrifugation at maximum speed (4500 rpm) for 2 minutes was the optimal method used to recover most of the complex gels within the shortest time possible.

4.4.4.5 The pH of Prepared PEC Gels Before and After Washing Steps

Finally, the pH of the PEC gels was confirmed before and after washing steps with ultrapure type 1 water, to confirm whether the pH of the PEC gels remains constant after the washing steps. It can be suggested that negligible remnants of acids and alkalis were present in the PECs prepared between pH 4 and pH 8 since the pH of the PECs after washing remained approximately the same. However, the PEC gels produced at pH 3 and 9 exhibited 1 pH unit closer to neutrality when washed with ultrapure type 1 water. This suggests that the materials at pH 3 and 9 were not fully adjusted to the required pH level, and therefore more time should have been allowed to equilibrate the pH value of the PECs during PEC preparation.

4.4.5 Composition of PEC Gels

4.4.5.1 The Importance of Measuring the PEC Composition

Knowing the composition of PEC gels is necessary to understand and characterise the materials. Furthermore, from a regulatory point of view within the pharmaceutical and biomedical industry, there is a need to quantify the amounts of material fractions present in the complex. Since only small quantities of PEs remained in the supernatant after reacting at the low pH range, it can be assumed that most of the PEs have been consumed into the complexes. This was expected since both PEs were highly charged at pH 3-5. At higher pHs, the CS precipitates and combines with the PECs.

4.4.5.2 FTIR Absorbance Bands of PEs and PECs

Characteristic FTIR absorbance bands were present in CS at 1650 and 1587 cm\(^{-1}\) due to the secondary amide (C=O stretching) and amine (N-H bending), respectively. The bands present in pure CRG at 1230, 914, 844 cm\(^{-1}\) were due to the S-O bond of the sulfate group, 3,6-anhydro-D-galactose and D-galactose-4-sulfate, respectively. FTIR measurements were also carried out to confirm the presence of electrostatic interactions as this was not possible with the spectrophotometric or NCS elemental analysis technique. The broad absorbance region at 1200-1000 cm\(^{-1}\) is a standard fingerprint region for polysaccharides, and the absorbance was the highest at 1033-1026 cm\(^{-1}\) due to the C-O-C
stretches (glycosidic linkage) that is present in both CS and CRG \(^{149}\). The absorbance band at 1529 cm\(^{-1}\) in Figure 4.13 was only visible after PEC formation and was thought to be due to OSO\(_3^-\)-NH\(_3^+\) that is responsible for the electrostatic complexing \(^{23}\). Electrostatic PEC formation was only confirmed at pH 3 - 7.4 due to the existence of electrostatic interactions between PEs.

**4.4.5.3 Interrelationship between XRD and FTIR**

The relatively broad diffraction peaks in CS at 2θ = 10.6° and 19.9° are indicative of crystal form I and crystal form II, respectively \(^{119,205}\). Therefore CS is semi-crystalline as was previously reported in the literature \(^{132,206,207}\). On the other hand, CRG showed a broad XRD curve denoting an amorphous structure (Fig. 4.14). The broad peak observed in PECs at 2θ = 22° indicates that PECs were also amorphous in nature. This may have been due to the quick rearrangement of the PEs in the complex. Therefore the PEs did not have enough time to rearrange into a more ordered and crystalline structure \(^{208}\). The results from FTIR were corroborated with XRD, confirming PEC formation at pH 3 - 7.4. The suppression of the CS peaks in the XRD curve indicates good compatibility and strong interactions between CS and CRG in PECs. The CS peaks started to reappear in PECs produced at high pH (pH \(\geq 9\)), showing that the CS was slowly precipitating out from the complex. These peaks were absent in the complexes due to the disruption of the hydrogen bonds between amino and hydroxyl groups in CS \(^{208}\).

**4.4.6 Rheological Properties of PEC Gels**

**4.4.6.1 Direct Quantitative Measure of the Strength of Electrostatic Interaction of PEC Gels**

The main advantage of directly characterising the wet PEC gels is that their native binding state can be measured while their electrostatic complexing remains undisrupted. The direct measurement of the PEC gels with rheology can save time, avoid loss of structural information and therefore becomes fundamental to the understanding of PEC film and scaffold preparation and their physicochemical properties.

**4.4.6.2 Amplitude Sweep**

The linear viscoelastic region (LVR) is the area where the G’ and G” were independent of strain. This point was reached earlier for some PECs, but overall it was around 1 % (Fig. 4.15). Breakdown and reformation of PEC structure network occur when the strain is beyond the LVR \(^{38,109}\). For all the PECs, the G’ was higher than the G” indicating that the PECs were predominantly exhibiting an elastic behaviour. However, at higher strains (10-100 %), the crosslinks of the PEC structure were broken, resulting in a material with a Newtonian fluid-like behaviour.
4.4.6.3 Frequency Sweep

Previously it was shown that the extent of electrostatic complexing is highly influential on the viscoelastic properties. Weinbreck \(^{199}\) and Espinosa \(^{102}\) have also shown that electrostatic complexing of whey-protein/gum-arabic was the strongest at pH 4. The complex coacervates formed in the previous study were more viscous than elastic \((G'' > G')\) \(^{43}\). This may be due to the use of a weaker anionic PE (gum-arabic) compared with a strong anionic PE (κ-CRG) utilised in the present study. The mechanism behind the viscous behaviour may be attributable to the higher hydrophobic forces (polymer-polymer interaction) compared to the strong electrostatic interactions found in this study \(^{38,209}\). No cross-over point was observed for CS-CRG PEC gels at any oscillating frequency used, implying that the PECs were highly elastic and possessed characteristics of a gel-like structure \(^{186}\). Even at higher pH where the CS was fully deprotonated, some interaction remained between CS and CRG due to some induced electrostatic, hydrophobic and physical entanglements \(^{6}\). A previous study on CS with BSA reported a cross-over point between \(G'' > G'\) at higher frequencies showing a more viscous-like behaviour \(^{210}\).

4.4.6.4 Viscosity Measurements

From the viscosity measurements, it is apparent that the PEC gels displayed shear thinning properties. All PECs behaved as Bingham fluids (evidenced by the sharp decrease in viscosity upon shearing). This flow behaviour is a result of structure breakdown in PECs, which was recovered at rest within several minutes (data not shown). Again, this result was consistent with the pH, showing agreeable results between the viscosity and SEI of PECs in Figure 4.18. The only previous rheological measurements conducted on CS-CRG PECs were carried out by Shumilina et al. (2002) \(^{21}\). Measurements were performed at strains no larger than 0.01 % and in the oscillating regime at broader frequencies ranging from 0.001 to 40 Hz (0.0063 to 251 rad/s). They observed that the mechanical properties were significantly affected by the concentration of CS but not by the concentration of λ-carrageenan. In addition, the PEC gels produced from κ- and ι-CRG with CS were stronger but also more brittle than the CS/λ-CRG PEC gels. However, the κ-CRG and ι-CRG were more sensitive to temperature changes and showed a significant reduction in viscosity at higher temperatures. The sensitivity was caused by the change in structural conformation of CRG from helix-to-coil transition at 30-40 °C. Although in theory, λ-CRG has more sulfate groups per monomer of CRG to form more electrostatic interactions, the crosslinks induced by the helix in κ-CRG and ι-CRG appeared to induce a greater effect on the gel strengths than the effect of the number of electrostatic interactions did.
4.4.6.5 Desired PEC Viscosity for Producing Freeze-Dried Scaffolds

Florczyk et al. (2011) recommended that the viscosity of the chitosan-alginate PEC slurry should be below 300 Pa.s to produce uniform pores in scaffolds because the migration of growing ice crystals becomes difficult at high viscosity leading to large and irregular pores. The polyelectrolyte complexes produced at pH 3-7 showed the highest yield but did not have suitable viscosities for the production of uniform freeze-dried PEC scaffolds. According to the measured viscosities, the most appropriate pH candidates for uniform scaffold production of CS-CRG PEC gels would be those prepared at pH 7.4 and above. Hence, the best results to control porosity in freeze-dried scaffolds is to have the lowest viscosity possible without compromising on the mechanical properties of the final freeze-dried structure.

Lastly, the inter/intra-molecular crosslinking of the PEs is also another important parameter to consider for the stability and the mechanics of films and scaffolds. The freeze-dried PECs prepared at pH 3-5 used for FTIR, XRD and NCS analysis showed noticeably lower stability than those prepared at higher pH. This may have been due to the greater intra-crosslinking but low inter-crosslinking caused by the strong electrostatic interactions in low pH PECs. The formation and effect of inter and intra-crosslinking aspect will be described in greater detail in Chapter 5 and Chapter 6.

4.4.6.6 The Effect of SEI was greater than the Effect of Composition and Moisture Content on PEC Gel Viscosity

Overall, it can be concluded that the viscosities of the PEC gels were dominated by the electrostatic interactions rather than the composition and moisture contents of the PECs (see red ellipses in Fig. 4.18 and Fig. 4.19). The increase in pH from 5-8 led to a lower strength of electrostatic interaction and consequently a lower viscosity (Fig. 4.18). Although the interaction strength was reduced by increasing pH, the composition was unchanged over the pH range of 3-8 (Fig. 2.2). Unexpectedly, the moisture content appeared not to be dependent on the composition of the PECs prepared at pH 3-8. Nor did the moisture content show a significant effect on the viscosity. This can be confirmed at pH 5 and 6, where the viscosity of the PEC gel at pH 5 was about two orders of magnitude higher than the gel at pH 6, and the moisture contents and compositions were approximately the same. The PECs prepared at pH 9-12 exhibited an increase in the fraction of CS and the moisture content but changed the viscosity only very little when compared to the substantial viscosity change caused by the electrostatic interactions from pH 5 to 6. However, the SEI at pH 6 did not match with the viscosity measurement, and therefore it is important to note that the SEI results from the ζ-potential measurements can only provide an approximate prediction of the strength of PEC interactions. The solid content of the PEC gel measured at pH 6 was shown to be nearly as high as the PEC gels prepared at pH 3 and 4 (Fig. 4.10).
However, the viscosity of the PEC gels prepared at pH 3 and 4 were almost two orders of magnitude higher than the PEC prepared at pH 6. Nonetheless, it would be interesting to further investigate the effect of moisture and solid content on the viscoelastic properties by adjusting the water content of the PEC gels.

Figure 4.18 – Viscosities and strength of electrostatic interactions of CS-CRG PEC gels prepared at various pHs.

Figure 4.19 – Molar composition and moisture content of CS-CRG PEC gels prepared at various pHs.
Finally, a proposed structure of the PEC gels prepared at different pHs is illustrated below in Figure 4.20. The strong interaction between PEs when reacted at low pH results in very dense complexes with low moisture content and high fibrous structure which may restrict the sliding of the PEC units and ultimately increases the PEC viscosity. The proposed structure of PEC gels was devised from visual assessment, optical microscopy, moisture content and rheology.

![Figure 4.20 – Proposed structure of PEC gels devised from visual assessment, optical microscopy, moisture content and rheology.](image-url)
4.5 Conclusions

This chapter described the production and characterisation of crosslinker-free CS-CRG PEC gels. Various pHs (2-12) were investigated. Overall, it was shown that the pH affected the charge of CS significantly which affected the overall SEI between CS and CRG complex gels. The SEI was predicted using ζ-potential measurements and corroborated with the viscosity, which provided a direct measure of the strength of self-crosslinking. Overall it was found that the self-crosslinking was high at pH 3-5 and low at pH 6-12. The self-crosslinking of the PEC gels was found to be the highest at pH 5. This was confirmed by the highest SEI, yield, solid content, and viscosity measurements. The strength of interaction reduced when the pH was decreased or increased, due to the protonation of CRG and deprotonation of CS, respectively. From the results, it was shown that the viscosity was mainly influenced by the SEI and not by the composition or moisture content (3 ≤ pH ≤ 7). The high SEI at low pH resulted in the formation of fibrous precipitates caused by the strong self-crosslinking and the vigorous mixing. FTIR and XRD confirmed complex formation in PECs prepared at pH 3-7.4. The composition measured with NCS analysis was found to remain relatively the same from pH 3-8 at approximately 1:1 molar ratio between CS and CRG. In total, the amount of CS in PEC was found to be in excess, however the amount of CRG on the surface of the PEC was dominating. Therefore, the composition was unaffected when the SEI was reduced. The composition was only changed significantly from pH 9 and above, because of the inclusion of the neutralised CS to the PECs.

The next chapter will present the inter-relationship between the viscosity of the gels and the mechanics of films prepared at various pHs. Electrostatic self-crosslinking may be an ideal alternative to control the mechanics and stability of films used for medical applications. If the physicochemical properties of PEC films can be controlled, then the film materials may be used as promising crosslinker-free biomaterials.
Chapter 5 – Polyelectrolyte Complex Films

5.1 Introduction

A polyelectrolyte complex (PEC) film is a flexible two-dimensional (2-D) strip made from complexed macromolecules. The study of films is a useful way to determine the physicochemical and biological properties of the struts from complex three-dimensional (3-D) scaffolds. These strut models are essential for the development of new materials and complex structures with tailored surface properties. The measurements of intrinsic properties of different scaffold materials can be difficult. This is because the porosity and pore sizes in scaffolds can vary significantly from one material to another and could drastically influence the bulk properties.

In Chapter 4 it was shown that the composition of PECs and the strength of electrostatic interaction (SEI) affected the form of the PEC gels. Precipitate complex fibres were obtained when the CS and CRG PEs interacted in their fully ionic state i.e. at low pH (< pH 7). In contrast, a homogeneous complex coacervate was formed when CS was weakly charged i.e. at high pH (> pH 7).

The initial response of the body to the biomaterial will not only depend on the composition of the material but also on the topography of its surface. Surface properties may be modified using a layer-by-layer method to enhance the cell-material interactions. However, the limitation is that many dipping cycles are needed to fill the matrix material which makes the layer-by-layer method time consuming. For these reasons, polyelectrolyte complex (PEC) films have been proposed as materials which may be functional both from a mechanical perspective and have surface functionality.

In this chapter, the tensile properties of films produced from the various PECs produced at different pH were measured and correlated with the viscosity of gels. It is envisaged that the PECs prepared at various pHs will provide different PEC film stabilities, since the nature of the self-crosslinking will be dependent on the types of interactions present in the PECs, such as electrostatic (intra-crosslinking) or secondary interactions (inter-crosslinking). In this thesis, inter-crosslinking is defined as the interaction between the PEC molecules forming a network via entanglements, whereas, intra-crosslinking is the formation of PEC molecules without forming a network. Solvent casting PEC gels prepared at different pH conditions may provide tailorable PEC films that possess adequate stiffness and strength without the need for chemical crosslinking.
The objectives of this chapter were to investigate:

1. The preparation of CS-CRG PEC films produced at different pH conditions.
2. The water absorption capacity and stability of films.
3. The surface properties of films using SEM.
4. The mechanical properties of films using a tensile testing machine.

5.2 Materials and Methods

5.2.1 Preparation of CS, CRG and PEC Films

CS and CRG films were prepared based on the method reported by Carneiro et al. (2013)\(^\text{(147)}\). In brief, CS solution was produced by dissolving CS 2 % w/v in 0.16 M HCl for 24 hours. CRG solution was produced by dissolving CRG 2 % w/v in ultrapure type 1 water for 24 hours followed by 1 hour of heating at 60 °C. CS films were fabricated by casting 9 ml of the CS solution in round silicone moulds with a diameter of 54 mm using a 5-ml transfer pipette. CRG films were fabricated by casting 8 ml of CRG solution at 60 °C into round preheated Teflon\(^\text{®}\) moulds with a diameter of 50 mm. Teflon\(^\text{®}\) moulds were pre-heated in the oven to 60 °C to avoid early jellification of CRG and to ensure the production of uniform films. Furthermore, the polyelectrolyte solutions were pipetted slowly to prevent the formation of air bubbles within the films. PEC films were prepared by adding appropriate amounts of ultrapure type 1 water to the homogenised PEC gels developed in Chapter 4, to produce PEC suspensions of 2 wt. % prior to solvent casting. Also, the solid contents of the PEC gels can be found in Chapter 4. The pH of the diluted PEC gels was recorded, and 8 ml of PEC solution prepared at different pH values (3, 5, 7.4 and 9), was pipetted into round Teflon\(^\text{®}\) moulds with a diameter of 50 mm. The solutions were then dried in a humidified cell incubator (CO\(_2\) Incubator, BB 15, ThermoScientific, USA) for two weeks. The humidity in the cell incubator was maintained using 100 ml of ultrapure type 2 water inside a 100-ml glass beaker. After drying, the CS and CRG films were immersed in 1 M NaOH and 1 M KCl solutions respectively for 24 hours using a shaker at 100 rpm and subsequently washed with ultrapure type 1 water for 1 hour and 1 min respectively using a shaker at 100 rpm. The dried PEC films did not require further treatment. All films were ultimately left to dry in air for 24 hours followed by freeze-drying (VirTis, SP Scientific, AdvantEdge 2.0 BenchTop, UK) for 17.5 hours using the programme shown in Table 5.1.
5.2.2 Preparation of Collagen Films

Type I collagen films from bovine Achilles tendon (Sigma) were prepared based on the method reported by Grover et al. (2012)\(^{213,218}\). The solubilised and homogenised collagen slurry (1 % w/v) in 0.05 M acetic acid was placed in silicone moulds at a volume of 9 ml. The films were dried in a humidified incubator for two weeks. Some of the dried collagen films were crosslinked using 1.150 g 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride (EDC) and 0.276 g N-hydroxy-succinimide catalyst (NHS, both Sigma–Aldrich) in 70 % ethanol/water (v/v) per gram of film. The collagen films were crosslinked for 24 h and washed twice and thrice with 70 % ethanol and ultrapure type 1 water, respectively. The films were then dried overnight in a laminar flow hood followed by freeze-drying using the protocol presented in Table 5.1.

5.2.3 Swelling and Dissolution

The effect of pH on the PEC stability was determined using swelling and dissolution experiments. These experiments investigate the individual CS and CRG films and the PEC films prepared at pH 3, 5, 7.4 and 9. Circular films (ø 8 mm) were produced using a disposable biopsy punch (Kai, UK). Sample triplicates were submerged in PBS (0.01 M, pH 7.4) at a sample/PBS ratio of 2 mg/ml and incubated at 37 °C for 28 days. The mass of the samples was measured at different time intervals (1 h, 6 h, 24 h, 3 d, 7 d, 10 d, 14 d, 21 d and 28 d). The wet (swollen) mass \((m_w)\) of the films was measured after gently dabbing the wet films on tissue paper to remove any excess water on the surface. The mass loss was then found when the films were dried using a freeze-drier for 17.5 hours (Table 5.1) until a constant mass was reached \((m_d)\). The final dry mass was recorded using a microbalance (Mettler Toledo, XPE analytical, Switzerland) that is accurate to ± 0.1 mg. Swelling of films was reported in terms of water absorption:

### Table 5.1 – Freeze-drying protocol used for drying small-sized film materials.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Time (min)</th>
<th>Ramp/ Hold</th>
<th>Pressure (Torr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start</td>
<td>20</td>
<td>10</td>
<td>R</td>
</tr>
<tr>
<td>Ramp/ Cooling</td>
<td>-20</td>
<td>60</td>
<td>R</td>
</tr>
<tr>
<td>Anneal</td>
<td>-20</td>
<td>120</td>
<td>H</td>
</tr>
<tr>
<td>Dry</td>
<td>-20</td>
<td>60</td>
<td>H</td>
</tr>
<tr>
<td>Dry</td>
<td>0</td>
<td>900</td>
<td>H</td>
</tr>
<tr>
<td>Dry</td>
<td>20</td>
<td>30</td>
<td>R</td>
</tr>
<tr>
<td>Stop</td>
<td>20</td>
<td>60</td>
<td>H</td>
</tr>
</tbody>
</table>
Water Absorption, % = \frac{\text{Wet Mass (} m_w \text{) } - \text{Dry Mass (} m_d \text{)}}{\text{Dry Mass (} m_d \text{)}} \times 100 \hspace{1cm} (5.1)

The percentage mass loss was calculated using the equation below:

\text{Mass Loss, %} = \frac{\text{Initial Mass (} m_0 \text{) } - \text{Dry Mass (} m_d \text{)}}{\text{Initial Mass (} m_0 \text{)}} \times 100 \hspace{1cm} (5.2)

where \( m_0 \) is the initial dry mass of the sample and where \( m_d \) is the dried mass of the sample at a different time interval. The acidic or alkaline by-products of the films were also determined during the swelling and dissolution study, using pH measurements. Photographs of the dried films were taken at days 0 and 28 using a Canon EOS 5D camera, Japan.

5.2.4 Enzymatic Degradation

Cylindrical films of CS, CRG and PECs prepared at pH 3, 5, 7.4 and 9 were cut using a biopsy punch (Ø 8 mm). The films were submerged in PBS (0.01 M, pH 7.4) containing 1.5 µg/ml lysozyme (hen egg-white, 70,000 U/mg, Sigma-Aldrich, UK) and incubated at 37 °C for 28 days with shaking at 100 rpm (KS 4000 IC Control, IKA, Germany). The concentration of lysozyme was chosen to closely match the concentration of lysozyme present in human serum according to Porstmann et al. (1989). The lysozyme solution was refreshed every other day to ensure continuous enzyme activity. The mass of the samples was measured at different time intervals (24 h, 7 d, 14 d, 21 d and 28 d). The films were then rinsed with ultrapure type 1 water and the dry mass was found by removing excess water by lyophilization using the programme presented in Table 5.1. The final dry mass was recorded using a balance (XPE analytical, Mettler Toledo, Switzerland) that is sensitive to ± 0.1 mg. Photographs of the dried films were taken at days 0 and 28 using a Canon EOS 5D camera, Japan. The percentage mass loss was calculated using the equation below:

\text{Mass Loss, %} = \frac{\text{Initial Mass (} m_0 \text{) } - \text{Dry Mass (} m_d \text{)}}{\text{Initial Mass (} m_0 \text{)}} \times 100 \hspace{1cm} (5.3)

5.2.5 Scanning Electron Microscopy

Small sections of PEC films were mounted on 12 mm SEM stubs and sputter-coated (Emitech K575 sputter-coater, UK) with platinum for 50 seconds using a deposition current of 40 mA to increase the surface conductivities of the PEC films. The surface morphologies of the films were then imaged with secondary electrons (SE) using an SEM (FEI Nova Nano SEM 450, The Netherlands). The surface roughness of PE and PEC films was compared at 500 x magnification. A voltage of 5 kV and a working
distance of approximately 5 mm with a relatively small (2.5 nm) spot size (to increase resolution) were used to image the morphological structure of PEC films at 0 and 28 d dissolution time points.

### 5.2.6 Contact Angle Measurements

The droplet contact angle on film surfaces was measured using a sessile drop method. A face positioned HD Camera (Canon EOS 5D, Japan) with a 100-mm macro lens was positioned on the droplet. A 5 µl droplet of ultrapure type 1 water was placed on the horizontal surface of each film using a 20 µl pipette and observed with a camera positioned at a fixed horizontal angle and stable platform. The contact angle measurements of the water droplets on the film surfaces were obtained using a contact time of 60 seconds. The water contact angles were analysed using the protractor tool in SketchUp. The measurements were carried out in triplicate.

### 5.2.7 Mechanical Tensile Testing

The mechanical properties of films were evaluated using a Hounsfield tensile tester (Hounsfield, H5KS Materials Tester, USA) equipped with interchangeable load cells of 5 N and 250 N. The 5 N load cell was used for all the materials except for chitosan, XL collagen and nX collagen which were tested with a 250 N load cell. The ultimate tensile strength was used to determine the effect of pH on the extent of self-crosslinking of PECs. All film samples were immersed in PBS (0.01 M, pH 7.4) for 24 hours at RT and cut into rectangular samples of 26 mm in length and 7 mm in width using a scalpel. Subsequently, the films were rehydrated in PBS, and the sample thickness was measured five times along the gauge length (10 mm) of the film using a digital micrometer (Mitutoyo, Japan) to obtain a mean thickness value. The screw of the micrometer was only moderately tightened to prevent damage to the wet films. The average thickness of all films combined was at 0.24 ± 0.19 mm. Similarly, the grips of the Hounsfield tester holding the films were finger-tightened to avoid both slippage and breakage of the wet films at the grips. A preload of 0.01 N was used to align the films before the loading measurements were started. All tests were performed at room temperature, and a constant extension rate of 6 mm min⁻¹ was used. All samples were stretched until failure and materials that failed at the grips were discounted. The tensile strength (TS, MPa) was measured by dividing the maximum breaking force (N) by the cross-sectional area (mm²) of each film. The linear region of the stress-strain curves was used to calculate the Young’s modulus (E, GPa). The elongation at break (EB, %) was measured using a laser extensometer (Hounsfield 500L-270 with an error of 0.5 - 0.2 % with a strain range of 5 % - 85 % full-scale deflection (FSD). The EB % is the ratio between the extension of the film at the point of rupture to the initial length of the sample expressed in percentage. The reported data for each material was the average of five measurements.
5.3 Results

5.3.1 Swelling

The ability of the materials to swell during cell culture or implantation was monitored at regular time points in PBS buffer. The results showed that the swelling of all films increased with longer incubation times, but reached their maximum swelling within 24 hours (Fig. 5.1). This demonstrates a time-dependent nature of swelling from 0 h to 24 h for all materials. The water absorption within the first hour was around 100 % for CS, 2100 % for CRG, 110 % for pH 3 PEC and 160 % for pH 9 PEC. It was found that the amount of water absorbed by CRG was 21 times greater than the quantity absorbed by CS within the first hour of swelling. The extent of water absorption of the PEC films was changed and remained the same after 24 hours, showing the highest swelling at pH 3 (550 %) and lowest at pH 9 (200 %). PEC pH 3 absorbed about twice as much water than the PECs prepared at pH 5 and 7.4 and three times more than the PEC prepared at pH 9. The water absorption in CS films remained constant at 100 % after being submerged in PBS for 28 days, whereas the CRG film was fully dissolved within six hours.

![Figure 5.1 - Water absorption of films in PBS pH 7.4 over a 28-day period.](image)

5.3.2 Dissolution

For most of the materials, the maximum dissolution occurred within 24 hours (Fig. 5.2). CRG was fully dissolved after six hours, but only 2.9 % of the total mass was lost from CS within 24 hours. A slightly higher mass loss was recorded for the PECs compared to the CS. The largest mass loss in PECs within
28 days was found to be for the PEC pH 3 (18.3 %). The PECs prepared at pH 9 showed the lowest mass loss (~ 0 %) for the duration of the study.

The films used in the dissolution study were washed and freeze-dried (Fig. 5.3). It can be seen that the CRG and CS films were transparent and opaque at 0 days respectively, whereas the PEC films exhibited opaqueness. After 28 days of dissolution, the PEC films increased in size and became more opaque due to the surface porosity. The CS and PEC films did not change significantly in morphology after 28 days of dissolution. All films were highly flexible when swollen. The PEC films at lower pH (3 and 5) exhibited a higher macroscopic roughness than the CS and PEC films prepared at high pH (7.4 and 9).
5.3.3 PEC Films pH Measurements

The pH of the PBS buffer was measured during the dissolution study of the films. It was found that the pH of the PBS buffer did not change significantly for most of the materials tested (Fig. 5.4). The greatest change in pH was exhibited by the PEC films prepared at pH 3 and 5, where the pH of the PBS changed from pH 7.4 to pH 7.14 within 6 hours. The PEC film prepared at pH 7.4 showed a slight reduction in pH to 7.26 within 28 days. For the rest of the samples (PEC pH 9, CS and CRG), the pH values of the PBS remained almost unchanged for the duration of the study, reaching a plateau between pH 7.3 and pH 7.5 after 3 days. The pH of the PBS buffer above remained relatively constant between pH 7.4 and pH 7.5.

\[ \text{Figure 5.4 – Solution pH measurements of the PBS-buffer during the swelling and dissolution study of the film samples.} \]

5.3.4 Enzymatic Degradation

The degradation profile of CS and the PEC films prepared at pH 3, 5, 7.4 and 9 are shown in Figure 5.5. The PECs prepared at lower pH exhibited a higher rate of mass loss than those prepared at high pH. No mass loss in films was found on the first day, except for the PECs prepared at pH 3, with a mass loss of 25%. The CS and PEC films made at pH 7.4 and pH 9 exhibited the least mass loss (1%) throughout the duration of the study. The pH 3 and pH 5 PEC films exhibited the highest mass loss within 28 days at 70% and 30%, respectively.
Figure 5.5 – Enzymatic degradation profile of CS and PEC films prepared at pH 3, 5, 7.4 and 9 using PBS and lysozyme.

The films used in the enzymatic degradation study were washed and freeze-dried after 28 days of degradation (Fig. 5.6). Nearly all the materials changed significantly in morphology from a translucent appearance to a rough white surface after 28 days. It can be easily observed that the CRG films showed the largest reduction in size and changed from a white translucent appearance to a soluble polymer. The pH 3 PEC films showed a significant amount of enzymatic degradation and the films turned to orange-brown.

Figure 5.6 – Freeze-dried films after 0 and 28 days of degradation.
5.3.5 Scanning Electron Microscopy

The surface morphology of the as-cast PE and PEC films were imaged using an SEM (Fig. 5.7). Almost all films, except for CS, appear smooth and uniform microscopically. The CRG film displayed the clearest and smoothest surface.

Figure 5.7 – SEM micrographs of the film surfaces at 0-day dissolution. Materials from top left to bottom right: CRG, pH 3, pH 5, pH 7.4, pH 9 and CS. The micrographs are shown at a magnification of 500x, and the scale bar is 100 µm. The micrograph of CRG film is shown at a magnification of 1000x, and a scale bar is 50 µm.

The SEM micrographs of the PE and PEC films incubated in PBS for 28 days showed rougher surfaces than the as-cast materials (Fig. 5.8). The pH 3, pH 5 and CS films were rougher than the pH 7.4 and pH 9 films. It was also observed that the micro-fibres protruding from the CS film surface became more prominent after 28 days of dissolution.

Figure 5.8 – SEM micrographs of the film surfaces after 28 days of dissolution. Materials from top left to bottom right: CRG, pH 3, pH 5, pH 7.4, pH 9 and CS. The micrographs are shown at a magnification of 500x, and the scale bar is 100 µm.
5.3.6 Contact Angle Measurements

Glass and polystyrene (PS) surfaces were selected as a reference for hydrophilic and hydrophobic controls, respectively. Figure 5.9 shows that the contact angle measurements between the droplets and the surfaces of the films were the lowest for glass (31° ± 9°) and the highest for polystyrene (69° ± 3°). The maximum contact angle for the prepared films was found on CS (58° ± 14°) and lowest on CRG (38° ± 6°). All PEC films prepared were relatively similar in contact angles (40° - 50°) and within error.

![Figure 5.9](image-url)  
*Figure 5.9 – Contact angle measurements of water droplets on film samples. Mean ± SD, n = 3.*
5.3.7 Mechanical Tensile Testing

The stress-strain curves for all the materials were compared with the crosslinked (XL) and non-crosslinked (nX) collagen films (Fig. 5.10). Overall, the collagen films exhibited higher stiffness or Young’s modulus (E) and ultimate tensile strength (UTS) than the PE and PEC films. The crosslinked collagen showed the highest E and UTS of all materials but was three times less extensible than non-crosslinked collagen. With regards to the polysaccharides, CS films exhibited a higher E and UTS than the CRG and PEC films. The CRG films showed the lowest E and UTS. However, the elongation at break (EB) for CRG was rather high in comparison with the other materials tested. Overall, the mechanical properties of the PEC films were found to be in between the properties of CS and CRG films. The PECs prepared at higher pH showed greater E and UTS than the PECs prepared at lower pH.

Figure 5.10 - Tensile-strain curves of films using 5 N load cell with a constant extension rate of 6 mm min⁻¹. Note that the stiffest materials (CS, XL Coll and nX Coll) were tested with a 250 N load cell at identical conditions.
In Figure 5.11, the initial portions of the curves are expanded and the points at which the tangents are drawn to measure Young’s moduli, are shown.

Figure 5.11 - Expanded region of the initial part of the tensile-strain curves of films adapted from Fig. 5.10. The tangent of the initial part of the slope is used to measure the Young’s modulus.

The Young’s moduli of the film materials are presented in Figure 5.12. The Coll XL (128.9 MPa) was about three times stiffer than the Coll nX (46.2 MPa). The E of the Coll nX can be matched with the E of pH 9 PEC. The E remained relatively low for the PEC films prepared at the lower pH range (3-6), compared with the films prepared at high pH range (7-9).

Figure 5.12 – Young’s moduli of film samples. Mean ± SD, n = 5.
Both Coll XL and Coll nX films showed significantly higher UTS than the individual PE and PEC films (Fig. 5.13).

The extensibility of the materials ranged from 15 to 35 % (Fig. 5.14). Overall the PEC films exhibited higher elongations to break than the individual PEs. However, for most of the films tested, the average extensibility of all the materials may be rounded to 25 % within error.
5.4 Discussion

5.4.1 Structural Properties

5.4.1.1 PEC Films Crosslinking: Inter/Intra-Crosslinking

Chemical crosslinking has a significant effect on the swelling and stability of materials. Typically, a higher degree of crosslinking results in a lower degree of swelling and greater stability in the material \(^{36,220}\). In this study, it was found that the swelling and stability of the PEC materials were mainly influenced by the composition and the crosslinking type. The crosslinking type can be divided into inter and intra-crosslinking. The intra-crosslinking is the strong and direct electrostatic binding between the PEs, whereas the inter-crosslinking is the weaker secondary interactions and entanglement between the PEC units \(^{189}\). However, it is the latter type that provides the overall stability of the solid material since it keeps the macromolecules interconnected. The diagram in Figure 5.15 shows the effect of pH on the formation of inter and intramolecular interactions. Four possible structure scenarios can occur when the PEC films are produced at various pHs.

5.4.1.2 Interaction at pH 2-3

At pH 2-3, the CS and CRG macromolecules are known to be degraded by the acidic conditions which result in the reduction of the molecular weight of PE chains \(^{196,221}\). The degradation of the individual polymers was not measured with GPC in this thesis, but the degradation of PECs prepared at pH 3 was indirectly observed by the browning effect of the films in the degradation study. The decrease in PE chain lengths may reduce the number of entanglements and inter-crosslinking which results in unstable structures (Fig. 5.15 (a)) \(^{185}\). Also, the formation of more interstitial gaps in the PEC structure results in greater water absorption in the film.

5.4.1.3 Interaction at pH 4-5

At pH 4-5, the macromolecules are likely to be longer than at pH 2-3 due to the less extreme pH conditions that result in the degradation of the molecules. The macromolecular chains are likely to be extended when the solid PE is dissolved in the aqueous solution due to the high electrostatic repulsion caused by the high charge density within each PE (Fig. 5.15 (b)) \(^{200,222}\). This extension of chains was also indirectly observed by the increase in solution viscosities during the dissolution of the individual PEs. Since the pH range is between the pK\(_a\) values of both PEs, the materials were fully dissociated. Therefore, the interaction between the oppositely-charged materials is stronger which may result in more intra-crosslinking than inter-crosslinking as evidenced by the non-stable PEC precipitates.
formed. Such PEC structures formed are known as ladder-like and can lead to the formation of heterogeneous complex precipitate gels. However, some intermolecular binding is likely to exist, since entanglements between PEs are inevitable even at highly ionised states. In Chapter 4 it was shown that the strong strength of electrostatic interaction (SEI) in the PECs prepared at pH 2-5 resulted in the formation of heterogeneous fibrous PEC gel precipitates (Fig. 4.6 and 4.7). Hence, the solvent-cast PEC films prepared at low pH were inherently rough in macroscale (Fig. 5.3). The strong intra-crosslinking of PEs at low pH may have resulted in the exclusion of water upon drying which led to the reduction of the volume occupied by the polymers in the films and the creation of local ridges and pores. However, the individual PE films, especially CRG (Fig. 5.3) exhibited transparent and smooth surfaces like those produced by Lima et al. (2013). CRG films were transparent because they are amorphous and the CS films were opaque due to their semi-crystalline nature. The PEC films were mostly transparent due to their relatively amorphous structure as was depicted from XRD results (Fig. 4.14). Freeze-drying has also induced porosity in the films, turning them opaque. Previous results also confirmed that CS films were slightly yellow in appearance and were found to be robust and rubbery when hydrated.

5.4.1.4 Interaction at pH 6-8

At pH 6-8, CS becomes deprotonated as was derived with zeta potential measurements (Fig. 4.4) and therefore loses its high charge density which is likely to form a random coil-like structure (Fig. 5.15 (c)). Nonetheless, some charges remain within CS up to pH 8 as was previously proved by the zeta potential measurements (Fig. 4.4). As a result, the electrostatic interactions between the PEs decreases and the entanglements and secondary forces such as hydrogen bonding between hydroxyl groups dominate. Therefore, the inter-crosslinking of the materials increases which results in the formation of more stable PEC films. The PEC structure formed is known as the scrambled egg-like model and may give rise to the formation of homogeneous complex coacervate gels. The weak electrostatic interaction led to strong inter-crosslinking at high pH. The inter-crosslinks are responsible for the stable structure that resists uniaxial tension. Therefore, the PEC films prepared at high pH range were higher in stiffness and strength compared to the PECs prepared at lower pH.

5.4.1.5 Interaction at pH 9-12

At pH 9-12, the electrostatic interaction between CS and CRG is completely diminished due to the complete deprotonation of CS. The inability of CRG to bind to the CS reduces the complex yield as was previously shown in Figure 4.8. It has been demonstrated that when the CS becomes fully neutralised, it precipitates on top of any remaining complex (Fig. 5.15 (d)). This precipitation was observed by the
NCS elemental analysis which showed the increase in the fraction of CS at higher pH (Fig. 4.13) and the inability to see the electrostatic complexing with FTIR (Fig. 4.14). The increase in CS content also reduces the water absorption, because CS is less hydrophilic than CRG. The PEC gels and films produced at this pH are more homogeneous than the PECs produced at low pH.

In summary, the PECs produced at higher pH range were more inter-crosslinked than the PECs prepared at lower pH. This can be supported by various experimental observations such as the low stability (Fig. 5.2), stiffness (Fig. 5.12) and dense fibrous precipitate structure formed in PECs prepared at low pH (Fig. 4.7). In contrast, the PEC films prepared at high pH contained a higher fraction of CS and were more stable. The high pH PEC films also absorbed less water, exhibiting homogeneous complex coacervate gels and film surfaces.
5.4.2 Stability

5.4.2.1 Swelling of Films

The swelling of PEC films within 24 hours ranged between 200 % and 550 % for pH 9-3. The water absorption of CS and PEC pH 7.4 films was found to be at 100 % and 250 %, respectively. In comparison with the materials tested in this study, crosslinked collagen scaffold material previously prepared by Davidenko et al. (2010) was found to swell by 400 %. CRG is known to be very hydrophilic (Fig. 5.1 - 2100 % water absorption within 1 hour) due to the repulsion of its strongly negatively-charged sulfate groups present on the macromolecules which result in an increase in chain distances. As a result, the free volume of the molecules becomes larger, and the material becomes more permeable to water. The driving force of water absorption in films was due to the difference in water potential between the macromolecules and the aqueous phase.

5.4.2.2 Hydrophilicity of Film Surfaces

Contact angle measurements were carried out to find the hydrophilicity of the materials and to support the swelling results. A surface is considered hydrophilic and hydrophobic when the water contact angle is smaller and larger than 90°, respectively. The contact angles in this study confirmed the hydrophobicity of the treated tissue culture plate polystyrene (69° ± 2.2°) and the hydrophilicity of the glass surface (31° ± 9°). These values are in good agreement with the previous studies found for treated polystyrene (65.5°) and glass surfaces (24°). The contact angle for non-treated PS has been recorded at 90°. The CS (58° ± 14°) films exhibited higher drop contact angles compared with the CRG films (38° ± 6°). Other researchers have found similar results for CS (50°) and CRG (40°). It can be confirmed that the hydrophilicity of CRG is higher than the hydrophilicity of CS as can be corroborated by the swelling study (Fig. 5.1) and contact angle measurements (Fig. 5.9). The differences of contact angles seen in the PEC materials were approximately within the uncertainty of the measurements.

5.4.2.3 Stability of Films

The stability of PEC films within 28 days in the non-enzymatic media ranged between -1 % and 18 % for pH 9 to pH 3 (Fig. 5.16). The non-enzymatic mass loss of PEC pH 7.4 and CS films was found to be at 2 % and 1 %, respectively. In comparison with stability testing of collagen scaffolds in non-enzymatic media, it was previously shown by Davidenko et al. (2015) that the non-crosslinked and crosslinked types dissolved about 80 % and 20 %, respectively within 28 days. Although the collagen samples were
produced as scaffolds, it seems that the PEC materials are as stable as the crosslinked collagen. The CRG film exhibited the lowest stability due to its high water solubility ($pK_a = 2$) 100.

![Figure 5.16 – Enzymatic and non-enzymatic (dissolution) mass loss in PBS with and without lysozyme after 28 days, respectively.]

### 5.4.2.4 The Effects of CS Parameters on PEC Stability

The negligible mass loss of the CS films in both the enzyme and non-enzyme media shows that the materials are stable against degradation and dissolution. Firstly, this may be because the amino groups of CS are deprotonated above pH 6 and thus become insoluble. For that reason, the PECs prepared at pH 9 showed higher stability than the PEC prepared at pH 3, due to the higher CS content in pH 9 PEC. Secondly, the CS used in this study is highly stable due to its high degree of deacetylation (DDA = 93.1 %). It was previously shown that CS was highly stable against enzymatic degradation due to the formation of highly crystallised structures at high DDA (> 90 %) 61. A similar effect was found in CS produced with a small DDA (< 10 %). In contrast, CS with an intermediate DDA (50 %) is highly water-soluble, and the degradation can be accelerated using a low molar mass CS 235. The high solubility at intermediate DDA was thought to be due to the large amorphous fraction in the polymer caused by the destruction of the secondary structure 236.

The overall mass loss in the films was found to be greater when lysozyme was used, compared with the PBS solution only. This is because the hexameric binding sites in lysozyme hydrolyse the glycosidic bonds present between the monosaccharide units in CS 61,237. The hexasaccharide units containing 3-4 or more acetylated units in CS contribute mainly to the initial degradation of the polysaccharide 238. In addition, the higher soluble CRG content in the PECs prepared at lower pH explains the large initial mass loss found in the pH 3 PEC film. The low degradation rate at later time points may have been due to the diminished $N$-acetylglucosamine residues.
5.4.2.5 Degradation of Films

Although the degradation study used was the closest method to simulate the *in vivo* environment, it is by no means a full replication of the physiological conditions. Therefore, all that can reasonably be achieved with this data is the ranking of PEC mass loss in the presence and absence of lysozyme.

5.4.2.6 Release of Acidic or Alkaline By-Products from PECs

During the dissolution study, pH measurements were carried out to measure the release of acidic or alkaline by-products from the films. The slight pH reduction of the PBS media with pH 3 and pH 5 PEC films showed that there was slight release of excess hydrogen ions due to the acidic production conditions. Nonetheless, the pH of the PBS remained relatively constant for the other samples, and therefore no significant acidic or alkaline by-products were released. It is therefore concluded that the PECs prepared at various pH conditions did not adversely affect the pH and that the PECs may be tolerable in biomedical applications in respect to the pH without further treatment.

5.4.3 Mechanical Properties

The films were placed in the tensile testing machine grips and pre-loaded at 0.01 N to align the films prior to testing. The 0.01 N pre-load was used since the error of the 5 N load cell is at around 0.2 %. Therefore, any loads below 0.01 N was considered as noise. The Young’s moduli of films were found from the linear regression part of the stress-strain curve in tensile testing. The point at which the tangent was drawn in the curve is shown in Figure 5.11. The linear part of the curve prior to yield stress was chosen for each material for measuring the Young’s Modulus. It was found that for each material and repeat a different strain range was used, depending on the linear regression curve. The problem with the current method used is that breakage of the films can occur at the edge of the grips. Perhaps on next occasion, dumbbell-shaped films may need to be used or a notch should be made in the middle of the gauge length to allow for breakage in the middle of the film. With regards to the Young’s modulus, the unloading part of the curve may be used, but this requires the films not to be ruptured i.e. only elastically deformed. However, since all the films were tested in the same way, the initial part of the loading curve is adequate for comparison of Young’s moduli of different materials. The stress-strain curves were also truncated to remove the toe region as much as possible and start with the linear region to measure the Young’s modulus. However, for some of the materials curves, the toe region is still present and therefore the linear region is taken that follows the toe region.
5.4.3.1 Mechanical Properties of Collagen from this Study and Grover’s Study

The mechanical properties of the PE and PEC films were compared with the collagen films because collagen is widely used in tissue engineering applications. The tensile stress-strain curves of the PECs and PEs were typical of polysaccharides \(^{239}\) and similar to those found for CS-gelatin nanofibers \(^{119}\). The non-crosslinked collagen showed a significantly lower E than the crosslinked collagen films since there were no interfibre covalent bonds present to prevent polymer chain sliding in the non-crosslinked collagen \(^{213}\). The Young’s modulus (E) and ultimate tensile strength (UTS) of the collagen films in this study were found to be higher than those reported by Grover et al. (2012) \(^{218}\). This may have been due to several reasons; the collagen films prepared by Grover and co-workers was submerged overnight in PBS at 37 °C rather than at 20 °C for 1 hour for this study. They also used a specialised rig to keep the films wet during tensile testing and therefore the materials were more hydrated during testing and potentially more compliant. In addition, Grover’s samples were crosslinked for 2 h \(^{218}\) rather than 24 h in this study.

5.4.3.2 The Viscoelastic Properties of Wet PEC Gels do not Associate with Mechanical Properties of PEC films

The results in Chapter 4 suggest that the PECs prepared at lower pH formed stronger electrostatic interactions as was indicated by the higher viscoelastic properties and viscosities (Section 4.3.10). It was therefore hypothesised that the PEC films prepared at lower pH could also result in a higher E and UTS. However, this notion was found not to be true.

5.4.3.3 The Effect of CS Structure on the Mechanical Properties

The higher CS content in the PECs also played a crucial role towards the higher E and UTS, indicating a dominant effect of composition on the mechanics of the films. The macromolecules of CS result in the formation of hydrogen bonds between hydroxyl groups and amino groups between the CS macromolecules, resulting in greater crystallinity and stability \(^{240}\). It can also be confirmed that the PECs with the higher swelling, were identified with lower E and UTS due to the greater plasticising effect caused by the water molecules \(^{241}\). Other important factors include the molecular weight (MW) of CS and the acid type used to solubilise the CS. In a previous study, the UTS of CS varied from 7-150 MPa when the MW of CS was increased and when malic acid was used to solubilise CS \(^{240}\). The elongation at break (EB) of the CS film was reported to be around 14.6-18.7 % when dissolved in acetic acid \(^{224}\) and 31.9-104.9 % when dissolved in lactic acid \(^{240}\). It was also found that at high DDA, the MW did not have an effect on the tensile properties of the CS film \(^{242}\). However, when the MW of CS was
small, and the DDA was between 10-90 %, the UTS of the films was low due to the fewer entanglements and hydrogen bonds 61.

5.4.3.4 The cause of Low Mechanical Properties in CRG

CS and CRG films have been tested before by other researchers under tensile stress in dry conditions, and it was found that the CRG films had higher UTS (30-38 MPa) and a lower EB (4.6-7.7 %) than the CS films with a UTS and EB values of 3.2-4.7 MPa and 62.7-84.2 %, respectively 148. The results from this chapter showed that the low tensile properties found in wet CRG might be due to its higher water absorption capacity than CS. This increases the polymer chain movement 34 and weakens the intermolecular forces between adjacent polymer chains 240. A similar result was previously found by Yin et al. (1999) 241, showing lower UTS properties in gelatin than in chitosan (3 kPa) 241 due to the higher swelling capacity.

5.4.3.5 Stiffness of Biological Tissues

Overall, it was shown that the PEC films were relatively stable and that chemical crosslinking was not necessary. The E of the PEC films were in the range of 2.9 – 13.9 MPa. In contrast, the collagen films were at 128.9 MPa (Coll-XL) and at 46.2 MPa (Coll-nX). The stiffness of most soft tissues is shown below the mechanical properties of the PECs and the collagen films (Table 5.2). However, since the PECs were lower in E than the collagen films, they may be more suitable for soft tissue implants from the perspective of the stiffness. The E of articular cartilage and skin were the closest match in E to the PECs. According to Chahine et al. (2004) 243, the E of cartilage was found to be between 3-13 MPa, which closely resembles the E of PECs at 2.9 – 13.9 MPa. According to Pal (2014) 244, the UTS of cartilage was found to be at 3 MPa, which most closely resembles the UTS of PEC pH 9 (2.4 MPa) and CS (3.4 MPa). The PECs prepared in this study may be most suitable as blank slates for the repair of cartilage tissues because of the closely related mechanical properties and the polysaccharide composition. The PECs largely resemble the glycosaminoglycans (GAGs) present in cartilage tissues such as hyaluronic acid, heparin sulfates, chondroitin sulfates and keratan sulfate which modulate the lubrication and mechanics of cartilage 72. Nonetheless, it is possible that chondrocyte-like cells may not adhere and proliferate well on the surface and therefore, protein surface modification may be carried out to improve the biocompatibility. The biological response of the films will be discussed in Chapter 7.
Table 5.2 - Tensile stiffness values of biological tissues.

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<th>Soft Tissue Type</th>
<th>Stiffness, MPa</th>
<th>Reference</th>
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<tr>
<td>Brain</td>
<td>$0.15-0.3 \times 10^{-3}$</td>
<td>40</td>
</tr>
<tr>
<td>Liver</td>
<td>$0.4-0.6 \times 10^{-3}$</td>
<td>40</td>
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<tr>
<td>Adipose</td>
<td>$0.3-15.6 \times 10^{-3}$</td>
<td>246</td>
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<tr>
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<td>245</td>
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</table>

5.4.3.6 Young’s Modulus of Films and Storage Modulus of Gels vs. SEI of Gels

Figure 5.17 shows the relationship of Young’s moduli of PEC films versus the SEI of PEC gels as previously presented in Chapter 4. It can be observed that the film stiffness increases when the SEI reduces. The high electrostatic interaction in PECs prepared at low pH resulted in strong intracrosslinking, which made the materials even less stable under uniaxial tensile testing. Therefore, the stiffness of PEC films at low pH (3-6) was much lower than those at high pH (7-9). The opposite was observed in rheology, where the viscoelastic behaviour of PEC gels was greater in PECs prepared at lower pH due to the strong electrostatic interaction (Fig. 5.18). A storage modulus of PEC gels higher than 3.2 kPa result in unstable PEC films with lower Young’s moduli. The data suggests that there is a boundary pH, that results in the formation of stable films. This pH boundary is likely to be between pH 6 and 7 and where the electrostatic interaction is between 700-1500 mV² (Fig 5.17 – green dotted lines).
Chapter 5 – Polyelectrolyte Complex Films

Figure 5.17 – The relationship between the Young’s modulus of PEC films and the SEI of the PEC gels.

Figure 5.18 – The relationship between Young’s moduli of PEC films and storage moduli of PEC gels at an angular frequency of 1 rad/s.
5.5 Conclusions

In summary, the properties of thin films were investigated and used to model the scaffold struts to understand the microscale properties of these biomaterials. Relatively stable PEC films were produced from CS-CRG PEC gels prepared at pH 7 and above. It was shown that the nature of crosslinking (inter/intra-crosslinking) between CS and CRG were affected by different pH regimes. The changing composition of PECs resulted in various swelling, stability, surface roughness and mechanical properties of PEC films. The preparation of PECs at various pH conditions did not lead to the release of acidic or alkaline by-products to affect the pH of the PBS media. From the results, it can be observed that the PECs may have advantages over collagen due to the lower E in PECs compared with collagen and the lack of the need for chemical crosslinkers which are usually highly toxic. The PECs may also have advantages over CS because the PECs are softer, more stable at low pH values. The lower E and UTS of PECs may be of greater advantage for specific soft-tissue engineering applications compared with the higher E and UTS materials such as CS and collagen.

From the stiffness perspective, the most suitable applications for the PEC films would be for soft tissue implants with moderate E such as cartilage. The next chapter will investigate whether mechanically stable scaffolds can be produced from PECs prepared at a wide range of pH conditions. Chapter 7 will introduce the cell-material interaction to determine the biological response of the PEC films.
Chapter 6 – Polyelectrolyte Complex Scaffolds

6.1 Introduction

In the context of tissue engineering, a porous scaffold is an interconnected structure used to regenerate damaged tissue by providing mechanical support to guide the growth of cells while allowing cell and nutrient infiltration, and vascularization to occur. Successful scaffolds require optimised structural and physical properties and can offer appropriate biological cues. Polyelectrolyte complex (PEC) scaffolds have been used for repair in the liver, cartilage, bone, and for drug delivery and stem cell delivery for the treatment of cardiac ischemia. Anti-fouling materials were also produced from scaffolds based on chitosan with oxazoline oligomers which were shown to repel protein adhesion and kill bacteria upon contact. The latter strategy was thought to be useful for the prevention of micro-organism resistance in the context of biomedical applications.

In Chapter 5, it was shown that the mechanical properties of the PEC films were mainly affected by the inter/intra-crosslinking which were shown to dramatically change the stability, swelling and mechanics of the PECs in their dry and wet states. The preparation of PECs at low pH conditions was inducing a higher level of electrostatic interaction (SEI) between PEs which resulted in stronger intra-crosslinking but weaker inter-crosslinking in PEC films. It was the inter-crosslinking formed by secondary interactions and entanglements that provided stability and elasticity to the films. It is therefore envisaged that a similar effect would take place with the freeze-dried scaffold swelling, stability and mechanics. The aim of this chapter is to find whether stable scaffolds could be produced from CS, CRG and PECs at different pHs and to potentially tailor the stiffness and strength of biomaterials without chemical crosslinking. The mechanical behaviour of PEC scaffolds will be compared to the mechanical properties of films and the viscoelastic properties of PEC gels. A scaffold with bulk properties most suitable for soft tissues will be identified. In this chapter, PEC scaffolds are prepared by freeze-drying homogenised PEC gels. The swelling properties, pore structure and mechanical properties of the scaffolds are then studied.
The objectives of this study were to investigate:

1. Whether stable PEC scaffolds can be produced at varying pH conditions.
2. The water uptake of the struts and pores of the scaffold.
3. The effect of production pH on the resulting structure and morphology of PEC scaffolds using a scanning electron microscope (SEM).
4. The pore size and pore size distribution using X-ray microtomography (Micro-CT).
5. The mechanical properties of scaffolds using a mechanical compression machine.

6.2 Materials and Methods

6.2.1 Preparation of Scaffolds

CS solutions were prepared by dissolving CS 1 % w/v in 0.16 M HCl for 24 hours. CRG solutions were prepared by dissolving CRG 1 % w/v for 24 hours followed by 1 hour of heating at 60 °C until fully dissolved. All materials were stored at 4 °C before use. CS and CRG solutions (1 ml) were cast in 48 well-plates (14 mm diameter and 20 mm depth) made of treated polystyrene (Corning® Costar®, Sigma-Aldrich, UK). The solutions were then freeze-dried in a VirTis AdVantage 2.0 benchtop freeze-dryer (Biopharma Process Systems, UK) using the protocol shown below in Table 6.1 and Figure 6.1. In summary, the solutions were frozen to -20 °C at a constant cooling rate of 1 °C min⁻¹. The temperature was then held constant at -20 °C for 8 hours. The ice phase was sublimed under vacuum (0.08 Torr) at 0 °C for 24 hours. The freeze-dried scaffolds were then immersed in 1 M NaOH and 1 M KCl solutions respectively for 24 hours using a shaker at 100 rpm. The scaffolds were subsequently washed with ultrapure type 1 water for 1 hour and 1 min respectively using a shaker at 100 rpm. The wet scaffolds were then freeze-dried again using the same freeze-drying protocol mentioned above. On the other hand, the PEC gels were homogenised, and 1 g of the PEC gel was placed into 48-well plates. The PEC gels were freeze-dried using the same protocol. Next, the PEC scaffolds were photographed using a Canon EOS 5D camera, Japan.
Table 6.1 – Freeze-drying protocol for small-sized scaffolds.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Time (min)</th>
<th>Ramp/ Hold</th>
<th>Pressure (Torr)</th>
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<td>20</td>
<td>5</td>
<td>H</td>
</tr>
<tr>
<td>Ramp</td>
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<td>40</td>
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<td>R</td>
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<td>1440</td>
<td>H</td>
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<tr>
<td>Stop</td>
<td>20</td>
<td>60</td>
<td>H</td>
</tr>
</tbody>
</table>

Figure 6.1 – The programmed protocol for the preparation of freeze-dried PEC scaffolds. This figure is intended to summarise Table 6.1. Note that the temperature and pressure do not represent the real conditions of the samples.
6.2.2 Swelling

Cylindrical scaffolds (14 mm diameter and varying heights) were submerged in 5 ml ultrapure type 1 water at RT for 14 hours. The swollen weight of the scaffolds (water uptake of pores and struts) was found by gently blotting the wet samples with tissue paper to remove the excess water. The percentage water uptake by the scaffolds was calculated using the equation below:

\[
\text{Water Uptake Scaffold, \%} = \frac{\text{Wet Mass} (m_w) - \text{Initial Mass} (m_o)}{\text{Initial Mass} (m_o)} \times 100
\]  

(6.1)

where \( m_w \) is the mass of the wet scaffold at a specific time point, and \( m_o \) is the initial mass of the dry scaffold.

The percentage water uptake by the struts was obtained by compressing the scaffolds between tissue paper to remove all the water present within the pores of the scaffolds. It was calculated using the equation below:

\[
\text{Water Uptake Struts, \%} = \frac{\text{Wet Mass Struts} (m_{ws}) - \text{Initial Mass} (m_o)}{\text{Initial Mass} (m_o)} \times 100
\]  

(6.2)

where \( m_{ws} \) is the mass of the wet struts of the scaffold.

The percentage water uptake by the pores in the scaffold was calculated using the equation below:

\[
\text{Water Uptake Pores, \%} = \text{Water Uptake Scaffold, \%} - \text{Water Uptake Struts, \%}
\]  

(6.3)

6.2.3 Scanning Electron Microscopy

A thin cross-section of the middle part (transverse plane) of the cylindrical scaffolds was mounted on 12 mm stubs using double-sided carbon black tape. The materials were sputter-coated (Emitech K575 sputter-coater, England) with platinum for 50 s using a deposition current of 40 mA to increase their surface conductivities. Then the surface morphology of the scaffolds was observed using an SEM (FEI Nova Nano SEM 450, Netherlands) with secondary electrons. A voltage of 5 kV and a working distance of approximately 5 mm with a relatively small (2.5 nm) spot size (to increase resolution) were used to view the morphology of polyelectrolyte scaffolds at a magnification of 50 x, 100 x and 500 x for each sample.

6.2.4 X-Ray Micro-Computed Tomography

Cylindrical scaffolds prepared in a 48-well plate (14 mm diameter and 20 mm depth) were analysed using a micro-CT (SkyScan 1172 X-ray micro-computed tomographer, SkyScan NV, Belgium). An X-ray source potential of 25 kV and an X-ray source current of 130 μA at an exposure time of 1.8 s and a
rotation step of 0.2°. High-resolution images (4.87 µm per pixel) were obtained using an 11 MP camera. Image analysis was performed using NRecon (Skyscan NV, Belgium) for clipping 300 cross-section slices as the volume of interest from the inner middle part of the scaffold. Three-dimensional reconstructions of scaffolds were obtained using DataViewer (Skyscan NV, Belgium). Pore diameter measurements were performed using ImageJ (NIH, USA) by training the image software on identifying the pores and the struts of the scaffolds. The pictures of the scaffolds were turned into binary (black and white) pixels. The average pore size was then calculated, followed by the pore size distribution of the scaffolds. The porosity of the scaffolds was measured according to the method reported by Bruker® using CTAn software (Skyscan NV, Belgium). The software reports a percent object volume which is the volume of solid in the selected region of interest divided by the total region of interest. Porosity is then calculated as 100 - percent object volume.

6.2.5 Mechanical Compression Testing

Compression stress-strain analysis of wet scaffolds was performed parallel to the plane of the vertical cylindrical scaffold using a mechanical testing machine (Hounsfield Low Load Electric Screw, UK). Scaffolds with diameters of 14 mm were immersed in ultrapure type 1 water at RT for 14 hours. The dimensions of swollen scaffolds were measured using a Vernier calliper. The compression plates were lowered to contact the sample and produce a small, but detectable load (0.002N). The scaffolds were then tested in compression with a 5 N load cell and a crosshead speed of 6 mm min⁻¹ until a sharp increase in stress was detected (densification of the sample) before the tests were terminated. The Young’s modulus (E) was obtained via a linear regression of the initial linear region of the stress-strain curve. The stress values were obtained at 5 % and 20 % strain. Five replicate measurements were performed for each scaffold material.
6.3 Results

6.3.1 Stability and Swelling

The stability and swelling behaviour of the scaffolds were investigated in PBS to assess their suitability as implants for soft tissues. The fluid uptake of the scaffolds by the pores and struts of the materials could provide a measure of the physiologically relevant buffers to be taken up during cell culture or in vivo studies. Dry porous scaffolds were obtained after freeze-drying the PEC gels in 48-well plates (Fig. 6.2). The dry scaffolds prepared at lower pH range (3 \( \leq \) pH \( \leq \) 6) were more brittle and more prone to disintegration during handling than the PEC scaffolds prepared at higher pH range (7 \( \leq \) pH \( \leq \) 9). The neutralised CS was very robust and stable during handling while the CRG was yielding and breaking to the touch.

![Dry porous scaffolds obtained from freeze-dried carrageenan (CRG), chitosan (CS) and PEC gels prepared at pH 3, 4, 5, 6, 7, 7.4, 8 and 9.](image)

---

*Figure 6.2 – Dry porous scaffolds obtained from freeze-dried carrageenan (CRG), chitosan (CS) and PEC gels prepared at pH 3, 4, 5, 6, 7, 7.4, 8 and 9.*
The stability of the scaffolds before and after swelling are shown in Figure 6.3. The CRG and the PECs prepared at pH 3 and 5 were more prone to disintegration, whereas the CS and PECs prepared at pH 7.4 and 9 remained intact and stable after swelling.

Figure 6.3 – Appearance of carrageenan (CRG), chitosan (CS) and PEC scaffolds prepared at pH 3, 5, 7.4 and 9 before and after swelling. Scaffolds were submerged in PBS at RT for 14 hours. Unstable scaffolds: CRG, PEC pH 3 and PEC pH 5. Stable scaffolds: PEC pH 7.4, PEC pH 9 and CS.
Figure 6.4 shows the water uptake of the scaffolds, struts and pores of the materials over a 14-hour period. It can be observed that the water uptake increases for the materials prepared at higher pH. The maximum water uptake was observed for pH 9 PEC which could hold 28 times its original weight. The lowest water uptake was observed for pH 3 PEC which could hold only four times its original weight. The chitosan scaffolds exhibited twice as much water uptake than the hydrophilic CRG scaffolds. Most of the water uptake can be seen to arise from the pores of the scaffold rather than the struts. The strut water uptake was approximately the same for the PECs prepared at pH 7.4, pH 9 and CS scaffolds. They were also able to return to the non-swollen state and maintain their stability when incubated in water after being compressed between tissue paper. The PECs prepared at pH 3, pH 5 and CRG disintegrated in the swollen state during handling. For that reason, the water uptake of the struts and pores could not be determined for the low pH PECs and CRG samples.

Figure 6.4 – Water uptake of carrageenan (CRG), chitosan (CS) and PEC scaffolds prepared at pH 3, 4, 5, 6, 7, 7.4, 8 and 9 when submerged in ultra-pure type 1 water at RT for 14 hours.
6.3.2 Scanning Electron Microscopy

In this section, SEM was used to determine the surface morphology and structure of porous scaffolds. The micrographs of the freeze-dried individual PEs and PECs are shown in Figure 6.5. During sample preparation and SEM viewing, it was observed that the scaffolds possessed different structures and pore sizes throughout their transverse sections. The pores in the scaffolds were small at the bottom and gradually increased in size higher up in the scaffold. For this reason, the homogeneous central region was chosen for the comparison of the structures and morphologies for different scaffold materials. The CS and PEC scaffolds prepared at pH 7.4 and pH 9 possessed a highly porous and homogeneous structure with no phase separation or crystal deposition whereas the CRG and PEC scaffolds prepared at pH 3 and pH 5 were harder to image because they were very brittle and crumbled during sample preparation.
Figure 6.5 – SEM micrographs of carrageenan (CRG), chitosan (CS) and PEC scaffolds prepared at pH 3, 5, 7.4, and 9. Samples were imaged at 50 x, 100 x and 500 x magnifications from left to right.
6.3.3 X-Ray Micro-Computed Tomography

Micro-CT can be used to obtain pore size distribution, interconnectivity, and 3-D models of the scaffolds. Due to the scope of this study only detailed 3-D images of the scaffolds and pore size distributions were measured (Fig. 6.6 - 6.9). Nearly all scaffolds exhibited an interconnected and homogeneous structure, except for the CRG and PECs prepared at pH 3 and 5, which possessed a dense and fibrous heterogeneous structure (Fig. 6.5 and Fig. 6.6).

![X-ray microtomography images](image)

**Figure 6.6** – X-ray microtomography images (side views) of the carrageenan (CRG), chitosan (CS) and PEC scaffolds prepared at pH 3, 5, 7.4, and 9.

CS scaffolds were shown to be the most homogeneous throughout the whole structure with highly dense struts resulting in small pore sizes (Fig. 6.7). The CRG scaffold formed a thin skin on the outside due to the dissolution of the CRG material during KCl crosslinking. However, overall, the CRG scaffold was porous with large pore sizes. The PECs prepared at pH 7.4 and 9 showed a relatively homogeneous porous structure with high interconnectivity and inter-crosslinking.
Figure 6.7 – X-ray microtomography images of carrageenan (CRG), chitosan (CS) and PEC scaffolds prepared at pH 3, 5, 7.4, and 9. The scaffolds with cross-section and transverse section are displayed on top and bottom, respectively.

The average pore sizes of the individual PE scaffolds (85 µm) was shown to be lower than those found in the PEC scaffolds (150 µm) (Fig. 6.8). The PEs also presented narrower pore size distribution with more pores present in the smaller range. However, the CRG scaffolds contained more pores in the larger range. On the other hand, the PEC scaffolds exhibited a more wide-ranging pore size distribution. The histograms of the PEC scaffolds prepared at pH 3 and 5 were similar. The same was true for the PEC scaffolds prepared at pH 7.4 and 9 (Fig. 6.9). The porosity of the scaffolds was also measured and it appears to show that the PECs exhibited higher porosity than the individual PEs with the highest porosity in PECs pH 7.4 and 9 (Fig. 6.9).

Figure 6.8 – Average pore size and pore size distribution of carrageenan (CRG), chitosan (CS) and PEC scaffolds prepared at pH 3, 5, 7.4, and 9.
Figure 6.9 - Histogram illustrating the pore size distribution and porosity of carrageenan (CRG), chitosan (CS) and PEC scaffolds prepared at pH 3, 5, 7.4, and 9. Pore sizes are presented in micrometres.
6.3.4 Mechanical Compression Testing

Compression tests were carried out on scaffolds to predict the behaviour of the materials in wet conditions. The compression stress-strain curves of swollen scaffolds are shown in Figure 6.10 (a). These curves represent one of the replicate measurements for each scaffold material.

![Stress-strain curves of carrageenan (CRG), chitosan (CS) and PEC scaffolds prepared at pH 3, 4, 5, 6, 7, 7.4, 8 and 9. The scaffolds were swollen in ultrapure type 1 water for 14 h and tested under uniaxial compression.](image)

The tangent of the curve for the Young’s Modulus calculation in compression is drawn below in Figure 6.10 (b). The Young’s modulus of the scaffolds was taken from the initial part of the slope. The initial part of the curve is magnified to separate the overlapping of the stress-strain curves. All Young’s moduli measurements were taken below 10 % strain which follows with the standard porous materials stress-strain curves.
The stiffness of the PE scaffolds was found to be lower than the PEC scaffolds (Fig. 6.11). The maximum stiffness in PEC scaffolds was found to be at pH 7 (20.6 kPa), and the lowest stiffness was at pH 6 (10.17 kPa). The stiffness of the CS and CRG scaffold was at 8.7 kPa and 6.38 kPa, respectively.

The compressive stress of the scaffolds at 5% strain was in the range of 0.5-1 MPa for all the scaffold materials. However, at higher strain (20%) the PEC pH 7 scaffold was among the strongest (4.36 MPa). The compressive stress of the PEC scaffolds prepared at pH 3, 4, 5, 6, 8 and 9 was lower than the individual CS scaffold (Fig. 6.12). However, the PECs prepared at pH 7 and 7.4 were the strongest and most stable.
Figure 6.12 – The compressive stress of carrageenan (CRG), chitosan (CS) and PEC scaffolds prepared at pH 3, 4, 5, 6, 7, 7.4, 8 and 9. Testing was carried out under uniaxial compression at a strain of 5 and 20%.
6.4 Discussion

6.4.1 Stability and Swelling

6.4.1.1 Crosslinking of Scaffold

As discussed in Chapter 5, the reason for the lower stability in PECs prepared at low pH was because the PECs were mainly intra-crosslinked while PECs prepared at high pH were mainly inter-crosslinked which provided the higher stability. The stability is formed by the entanglements and secondary forces such as van der Waals, hydrogen bonds and hydrophobic interactions between adjacent PEC particles.

6.4.1.2 Water Absorption and Stability of Scaffolds

The amount of water taken up by the scaffold depends on the material hydrophilicity and on the capability of pores to retain fluid. In fact, on average the pores in the scaffold were taking up six times more water than the PEC strut materials. Therefore, scaffolds with higher porosity exhibit higher water uptake. It can be pointed out from the struts and the pores that the PEC prepared at pH 9 absorbed the highest amount of water of all the materials used. This was likely to be due to a more rigid pore structure that could hold higher amounts of water which made the scaffold structure more mechanically resistant. In addition, the porosity was high because the freeze-dried PEC prepared at pH 9 contained the lowest amount of solid content of all the PEC materials (Chapter 4). The lower amount of solid content means that more water was present per gram of gel during freeze-drying. Therefore, the higher water content meant that more water is crystallised during freezing which results in a more porous structure after sublimation. The CRG scaffolds exhibited lower swelling than the CS scaffolds because the CRG started to dissolve and form a thin skin on the outside of the scaffold which lowered the diffusion and absorption of water into the scaffold.

6.4.2 Pore Structure and Pore Size Analysis

6.4.2.1 PECs Prepared at Higher pH were more Inter-Crosslinked

For cell migration and nutrient/oxygen transport to occur, scaffolds need to be highly porous. Moreover, the pores should be interconnected to promote the homogeneous regeneration of the tissue in a 3-D manner. CS and PECs prepared at higher pH appear to possess these properties and are therefore expected to be more suitable for tissue engineering applications.
6.4.2.2 Freeze-Drying Scaffolds and Scaffold Topography

It is well established that phase separation occurs after freezing a water-soluble polymer\(^\text{14}\). When the CS and CRG solutions are freeze-dried, the water freezes and nucleates into ice crystals\(^\text{13}\). This pushes any polymer to the side resulting in phase separation. After lowering the pressure to 80 mTorr, a porous scaffold is obtained due to the sublimation of the ice\(^\text{258,259}\). This forms an inter-crosslinked scaffold that is highly porous and interconnected. For a PEC scaffold, however, this does not always occur, because PEC gels can be a precipitate rather than complex coacervates. Precipitation occurs particularly when the solid content of the PEC gel is high (> 2 wt. %). The PEC precipitates produced at low pH (< pH 7) were mainly intra-crosslinked. The significant gaps in PEC pH 3 and pH 5 scaffolds suggest that the micro and macrostructure of the struts were not inter-crosslinked. Thus, when the electrostatic interaction between PEs was stronger; denser complex precipitates were produced which were harder to control during freeze-drying\(^\text{199}\). This corroborates the results previously obtained for the high swelling and low stability found in PEC films prepared at pH 3 and 5 (Chapter 5). The low pH range PECs are comparable to the micrographs found by Rajkumar \textit{et al.} (2013)\(^\text{260}\) which show a typical densely packed powder like scaffold which may have been crushed when the scaffold was mildly pressed onto the carbon black tape during SEM preparation. This proves that the scaffolds were only intra-crosslinked and not inter-crosslinked at low pH production.

6.4.2.3 PECs Prepared at Higher pH were more Stable

On the other hand, the PECs prepared at a high pH range are both inter and intra-crosslinked, creating a more stable scaffold. The structure of the PECs prepared at higher pH range was found to be similar to the CS-CRG PECs obtained by Araujo \textit{et al.} (2014)\(^\text{6}\). Araujo investigated scaffolds produced at different molar ratios but only at pH 7.4. On the other hand, Li \textit{et al.} (2013)\(^\text{23}\) investigated bovine serum albumin (BSA) loaded CS-CRG PECs and showed a similar dense heterogeneous pore structure to the PECs produced in this study in the low pH range.

6.4.2.4 Average Pore Size and Porosity of PEC Scaffolds

The average pore sizes of the CS-CRG PEC scaffolds were found to be higher than 100 µm, which according to previous research would make the scaffolds suitable for cartilage tissue engineering\(^\text{261}\). The individual CS and CRG scaffolds have higher porosity because they have a larger number of small pores which have a greater surface-to-volume ratio than large pores.
6.4.3 Mechanical Properties

6.4.3.1 Stress-Strain Curve Explanation

A typical compression stress-strain curve of wet porous scaffolds with low density, open-cell foams, with an interconnected network of struts consists of three distinct stages (Fig. 6.13) \(^{226,262}\). The stages are explained in more detail by Gibson and Ashby (1999) \(^{262}\). In brief, the linear elastic zone is controlled by the struts bending and this is the regime where the Young’s modulus of the hydrated scaffold is measured. The collapse plateau and the densification regime are the beginning and the completion of pore collapse within the scaffold, respectively \(^{262}\). The stresses of the scaffolds were found at 5 % and 20 % strain.

For clarity, the initial portion of the loading curve is expanded to distinguish the overlapping of the curves (Fig. 6.10 (b)). The accuracy of the results (linear regression) was determined by successively adding the points in the data to make sure the squared R values was > 0.95. The mechanical properties of scaffolds depend highly on the density of the material and on the porosity of scaffolds. For the mechanical properties of the scaffolds to be compared directly, it would be necessary to measure the pore content of each scaffold material. The porosity of the PEC scaffolds was approximately the same. However, the porosity of CS and CRG were much lower than the PEC scaffolds porosity (Fig. 6.9).
6.4.3.2 The Effect of Inter-Crosslinking, Composition, Pore Size Distribution on E and UTS

The inter-crosslinking and pore size distribution of the scaffolds seemed to have a greater impact on the mechanical properties of the materials than the composition. This can be supported by the lower stress and stiffness of PECs prepared at pH 9 compared to the PECs produced at pH 7.4 (Fig. 6.11 and 6.12) which contained more CS as was observed from Chapter 4 (Fig. 4.12). Conflicting conclusions were made in previous research by Araujo et al. (2014)⁶; they found that CS or CRG content had a greater impact on the mechanical properties compared to the porosity and pore size of scaffolds. This may have been because the composition was changed directly by adding different molar ratios of CS and CRG and by keeping the pH constant at pH 7.4 which did not affect the inter-crosslinking. The E of the PECs measured in this study is comparable to the E of PECs found in Araujo’s study. However, there was no clear pattern of the stiffness and stress measured in the present study, indicating that there may be more parameters responsible for the mechanical properties of scaffolds, such as the solid content of the PEC gels that were freeze-dried into scaffolds. Then PEC films have been formed by diluting the solid content of the PEC gels to 2 wt. %. This was not the case with the scaffolds, in which the PEC gels were freeze-dried in their native solid contents, because the PEC coacervate or most likely the PEC precipitate formed at lower pH is very dense and sediments when diluted which creates a heterogeneous scaffold that is very dense at the bottom and thin at the top. The pH 7 and pH 7.4 PECs were shown to be the most stable with the highest stiffness and elasticity which makes them suitable for soft tissue engineering applications.

6.4.3.3 The Effect of pH Preparation Conditions on Scaffold Pore Sizes

The PECs prepared at low pH contained a greater number of pores that are of larger size. This can make the structure of the scaffold less stable as was shown for PEC pH 3 and pH 5 compared to PEC pH 7.4 and pH 9. The pH 5 PEC pore sizes are shown to be higher than pH 9, solely because the former was freeze-dried at a solid content of 5 wt. % whereas the latter was formed at 1 wt. %. The larger pore structure in PECs prepared at low pH can be observed in the micro-CT images (Fig. 6.7) and the pore-size distributions (Fig. 6.9).

The PECs prepared at low pH (pH 3 and 5) possessed a high solid content of 4.5 and 5.5 wt. % respectively, whereas the PEC gels prepared at pH 7.4 and 9 possessed a solid content of 2.2 % and 1 %, respectively. Nonetheless, the PEC scaffold prepared pH 7.4 was high in stiffness and stress due to the higher inter-crosslinking of the scaffold.
6.4.3.4 Comparison of PEC Scaffolds to Collagen Scaffolds

The PEC scaffolds were found to have a higher Young’s modulus than the collagen and collagen-hyaluronic acid scaffolds prepared for adipose tissue engineering by 2-5 folds. This may be because the collagen scaffolds were prepared at 1 wt. % whereas the PEC scaffolds were between 1 wt. % and 3 wt. %.

6.4.3.5 Stiffness of Biological Tissues

An ideal scaffold should present similar mechanical properties (stiffness, strength, and extensibility) to the native tissues that are intended to be repaired. The mechanical properties of the scaffold can be sensed by the mechanical sensors of the cells and can, therefore, be essential to the guidance and stimulation of cells. The stiffness of some soft tissues has been measured before by other researchers and shown to be in the range of 0.16-949 kPa (Table 6.2). It was also previously found that neurons/epithelial cells, myoblasts and chondrocytes grew best on gel surfaces with the stiffness of 0.1, 1 and 10 kPa, respectively. From the perspective of the stiffness of the tissues, the stable PEC scaffolds prepared at pH 7 and above may be used as implants for neural, adipose and skeletal muscle applications.

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</table>

6.4.3.6 Critical pH Conditions on Scaffold Physicochemical Properties – A Summary

As was observed previously with the mechanical properties of PEC films, it can be suggested that there is a critical pH value between pH 6 and 7 where the mechanical properties of the scaffolds change significantly (Figure 6.14). This can be related to the high SEI and solid content in PECs prepared at pH 3-6 and abruptly changes at pH 7-9. The graph shows that the SEI and solid content of PECs can dramatically affect the Young’s modulus of PEC scaffolds. There may be a critical SEI value at around 1000 mV², where the interaction switches from intra-crosslinking to inter-crosslinking.
6.4.3.7 Interrelationship of PEC Films and Scaffolds Stiffness

Figure 6.14 illustrates the interrelationship of the Young’s moduli of PEC films (as presented in Chapter 5) and PEC scaffolds. Since the PEC films were prepared at the same solid content, it is likely that only the inter-crosslinking was predominantly affecting the stiffness of films (Fig. 6.16). On the other hand, the scaffolds were prepared at the native solid content, and the Young’s modulus was therefore affected not only by the composition but also by the pore structure and solid content. Hence, it would be difficult to compare the mechanical properties of films with those of scaffolds. Nevertheless, a trend exists that the PECs prepared at higher pH were more stable than those prepared at low pH for both films and scaffolds.
Figure 6.15 – A comparison between the Young’s modulus of films and scaffolds obtained using tensile and compressive testing, respectively.

Figure 6.16 – The proposed molecular structure of the scaffolds. The stiffness of the scaffolds was found to be lower at lower PEC preparation pH. The molecular structure of the PEC units prepared at higher pH is intertwined, forming an entangled network which results in higher overall scaffold stability.
6.5 Conclusions

This chapter described the production and characterisation of PEC scaffolds without the need for a crosslinking agent. It was found that the PEC scaffolds prepared at high pH exhibited higher inter-crosslinking and homogeneity. Thus, the scaffolds were more stable with more uniform pore size distribution. The composition influenced the properties of scaffolds, but other factors played a greater role in scaffolds such as the inter-crosslinking, PEC solid content, and pore size distribution of the scaffold. These factors can dramatically change the final properties of scaffolds which make the interpretation of the factors on the intrinsic properties of PEC scaffold materials more difficult than in PEC films. The PECs prepared at higher pH contained more pores that were of a smaller size inducing more water uptake. The scaffolds prepared at low pH were more densely packed and less inter-crosslinked than the high pH PECs. It can be concluded that the stress of scaffolds increased with the increase in the preparation pH of PEC gels. The mechanical properties alone cannot be used to draw a conclusion on the potential of the materials as implants. Therefore, the next chapter will present the biological properties of the materials tested in the form of 2-D films.
Chapter 7 – Biological Response of Polyelectrolyte Complexes

7.1 Introduction

At present, there is no clear consensus on the cell-binding properties of CS and CRG materials. While some researchers have found that polysaccharides are cell-compatible 272, others have found them to be non-biofouling as presented in the review by Junter et al. (2016) 27. Therefore, this chapter addresses the question of whether the CS, CRG and PEC materials support the adhesion of cells (fibroblasts, fibrosarcoma and chondrosarcoma cells) and a highly adhesive protein (tropoelastin).

There are two opposing strategies for modulating material interactions 273. The first is aimed at enhancing the cell-material interaction and in doing so promoting the attachment, migration, proliferation and differentiation of cells. Example applications include controlled drug delivery systems 274 and the regeneration of various tissues 272 such as peripheral nerves 61, liver 275, skin 16, cartilage 72 and bone 276. The second strategy involves creating non-adhesive surfaces for proteins and cells to prevent the activation of the immune response, blood coagulation (thrombosis), bacterial biofilm or extracellular matrix deposition. Examples of applications for this strategy can include coatings for catheters (urinary 277 and intravascular 278), mechanical heart valves 279, pacemakers 280, coronary stents 281, glucose sensors 282 and contact lenses 27,81,283,284.

When materials are implanted into the body, they are likely to come in contact with connective tissues. The most common cell type present in the connective tissue are fibroblasts. The main functions of fibroblasts are the production of the extracellular matrix (ECM) and wound closure 285. The ECM is a network of extracellular molecules such as collagens, proteoglycans, glycosaminoglycans, elastin, fibronectin, laminin, and several other glycoproteins secreted by cells that provide structural (three-dimensional) and biochemical support to the surrounding cells 286,287. Fibroblasts are anchorage-dependent cells in that they require integrin-mediated interactions to ECM molecules for their survival 288. Such interactions are linked to a wide range of cellular processes such as proliferation, differentiation and apoptosis. In some instances, the fibroblasts can differentiate into contractile myofibroblasts 285. This transition can be observed by the expression of alpha-smooth muscle actin and large focal adhesion points within the cell 289. Cell differentiation of fibroblasts into myofibroblasts is dependent on cytokines, the most potent of which is the transforming growth factor beta (TGF-β), adhesiveness and stiffness of the underlying material 290. When fibroblasts differentiate
into myofibroblasts, they excrete excessive ECM which can result in fibrotic diseases and host responses including the fibrous encapsulation of implants. In this work, three different cell lines (fibrosarcoma, fibroblasts and chondrosarcoma) have been examined to elucidate the biological properties of CS and CRG complexes. Preliminary cell studies were conducted with HT1080 human fibrosarcoma cells as this cell line is simple to grow and is widely used to study the in vitro adhesive properties of collagen due to its well characterised collagen-binding integrin, α2β1 to the GFOGER motif in collagen. Further work was conducted with primary human dermal fibroblasts to confirm whether these cells can adhere to CS, CRG and their complexes in a non-specific manner. These cells were chosen as they are highly prolific and adhere to a wide range of surfaces. Furthermore, they act as a model cell line to probe the potential for scar tissue formation which can be minimised when collagen-I and collagen-III forming fibroblasts cannot attach to the material’s surfaces. Chondrosarcoma (SW1353) cells are derived from a malignant bone neoplasm which form hyaline cartilaginous ECM. As such, chondrosarcomas were used here as a model line for chondrocyte function due to their highly proliferative activity and the ability to keep a consistent phenotype. The serum may bridge between the cell and the surface and this is integrin-mediated binding. In addition, GAG-like mediated signalling from the surface may provide non-integrin mediated cell adhesion. It is known that the primary integrin utilised by chondrocytes is α5β1 which binds to fibronectin (FN) that could adsorb onto the surface of a material from the surrounding blood serum. For the SW1353 chondrosarcoma line, this may be different, as they express integrin α5β3, the receptor for vitronectin, which is important for cell adhesion and migration. It was also previously shown that blocking integrin α6β1 binding to laminin can reduce the adhesion of SW1353, demonstrating that they utilise this integrin for adhesion.
Chapter 7 – Biological Response of Polyelectrolyte Complexes

7.2 Literature Review

7.2.1 Factors Affecting Cell Adhesion

7.2.1.1 Parameters Affecting Cell attachment

Cell attachment is a complex process as cells can respond to a variety of material properties such as the composition and the availability of charged side groups and the physical properties such as stiffness and roughness. Other parameters such as hydrophilicity/hydrophobicity and pore architecture (micropore, microfibre and nanofibre scaffold) can also play a significant role in cell binding. Biochemical factors such as cell types, genes regulation/expression, protein deposition, the presence of integrin recognition motifs and growth factors are fundamental to the fate of the cell.

7.2.1.2 Hydrophilicity/Hydrophobicity Effect on Cell Attachment

Hydrophobicity is complex with many contributions from chemical groups, both polar and non-polar, surface roughness and surface swelling. Similarly, cell adhesion is complex, sharing many of the influences as hydrophobicity such as surface roughness, charged and polar groups, topography and surface swelling and mobility. Despite this complexity, some general trends can be observed, however, these are often specific to the cell type being studied. It has been previously shown that surfaces with a moderate hydrophilicity can be beneficial to cell attachment with decreased attachment on strongly hydrophilic or hydrophobic materials. For example, extreme hydrophobicity and hydrophilicity can inhibit the adhesion of some cells. This is exemplified by unmodified polytetrafluoroethylene (PTFE) surfaces that are too hydrophobic while polyethylene glycol (PEG) surfaces that are too hydrophilic to allow direct cell adhesion. Murine fibroblasts have maximum adhesion on surfaces with a contact angle of around 70° which decreased significantly at higher and lower contact angles. However, it should be noted that for many studies, the change in hydrophobicity was controlled by using different materials such as PTFE for the most hydrophobic surface (116 ± 3°), PS for a hydrophobic/hydrophilic surface (75 ± 2°) and cellulose for a hydrophilic surface (18 ± 5°). Other materials including but not limited to polypropylene (PP) (92 ± 3°), polyvinyl alcohol (PVA) (42 ± 5°) and glass (SiO₂) (30 ± 2°) were used to fill the contact angle gap. It is clear that these samples do not solely differ in their hydrophobicity as they possess differing chemistry, topography, surface swelling and mobility. As such attributing cell adhesion to contact angle in isolation is often meaningless. Despite this, a general pattern could be found between fibroblast adhesion and contact angle using the materials above. Even here, the relationship between water contact angle and cell adhesion is not complete as the adhesion of the fibroblasts on glass was as high as the PS surface while the PVA surface exhibited...
lower cell adhesion than the cellulose surface all of which do not correlate with the water contact angle/cell adhesion theory. Accordingly, it is clear that other factors, such as surface chemical groups, roughness, charge and swelling are important for the adhesion of cells and that hydrophilicity or hydrophobicity alone cannot be used to predict cell attachment fully. To add further complexity, materials surfaces can adhere to active cell-adhesive molecules such as peptides and proteins. These possess specific sequences, for example, RGD that tether cells. Often the display of these sequences is influenced by the underlying material properties and so the cellular effect may be “indirect” via such molecules.

### 7.2.1.3 Hydrophilicity/Hydrophobicity Effect on Protein Attachment

Tamada & Ikada (1993) have also found that maximal protein (bovine serum albumin (BSA), bovine-y-globulin and fibronectin) adsorption was observed on different material surfaces with a contact angle of approximately 70°. They also found that preadsorption of BSA and FN, prevented and enhanced fibroblast adhesion to substrates, respectively. Similarly, according to Xu & Siedlecki (2007), proteins like BSA, fibrinogen and human factor XII adhere preferentially to surfaces with contact angles above 60° compared to surfaces with a contact angle below 60°. Here the wettability of the low-density polyethylene (LDPE) surfaces was controlled by glow discharge plasma which introduces surface chemical groups when exposed to air (oxidised to make it hydrophilic).

### 7.2.1.4 Preadsorption of Biomolecules (Proteins and Sugars) on Films

The preadsorption of protein onto a material surface is a major factor in determining the nature of cell interaction. For this reason, researchers have modified the surface of PEC samples with proteins such as collagen, fibronectin and laminin or cell adhesive motifs such as arginyl-glycyl-aspartic acid (RGD) to improve the viability and attachment of cells. In a previous study, chitosan-alginate (CS-Alg) PEC scaffolds were surface modified with specific peptides and proteins to enhance liver repair as measured by the production of albumin. It was shown that albumin release was increased in the following order when using the coatings RGD > collagen > fibronectin or laminin > unmodified scaffolds > tissue culture plate polystyrene. Other biologically active non-protein materials have been surface tethered to enhance the interaction between the cells and the surfaces. These include the attachment of sugar molecules including mannose, lactose and galactose molecules which bind to the lectins in cells and allow cell-cell interactions in hepatocytes. The active adhesion of cells through integrin-peptide binding sites can be distinguished from the passive cell adhesion where cells are adsorbed on surfaces through hydrophobic, Coulombic, and van der Waals forces. This passive adhesion usually occurs through multiple weak bonds and is generally lower in strength than the active adhesion found between cell-ECM ligands or peptides derived from ECM proteins e.g. RGD.
7.2.1.5 Effect of Surface Roughness on Cell Adhesion

Surface roughness is another important parameter that can influence the adhesion of cells \(^{314}\). The roughness can be divided into macroroughness (> 100 um), microroughness (100 nm – 100 um), and nanoroughness (< 100 nm) \(^{315}\). The response of cells to roughness has been found to depend on the cell type. For example, neural cells were more viable when cultured on nanoscale rough surfaces (6.25-49.37 nm) compared to microscale rough surfaces (87.2-200 nm) \(^{316}\). The neurons were less developed on the micro-rough surface with a round-shaped soma and poorly branched processes. Additionally, when MG63 osteoblast cells were cultured on polycarbonate surfaces with different roughness (micropore sizes ranging from 0.2 – 8 µm), higher cell adhesion and proliferation was noted with lower scale roughness \(^{317}\). On the other hand, it has also been demonstrated that non-regularly patterned surfaces with defined nanoroughness were shown to decrease cell adhesion \(^{318}\). In a previous study by Carneiro et al. (2013) \(^{147}\) 3T3 fibroblasts were found to have lower cell adhesion on rougher CS-CRG complexes compared to the smooth individual CS and CRG surfaces. However, it is not clear whether the increase in roughness resulted in the lower cell adhesion or if this is also due to the charges on these molecules that may provide steric hindrance to the cells on the molecular scale. However, the hydrophilicity was unlikely to be the main cause of lower cell adhesion on the CS-CRG PECs, since the CRG surface is more hydrophilic than the CS-CRG PECs and the number of cells adhering is relatively good. Nonetheless, the adsorption of fibrinogen was found to be higher on the PEC surfaces than on the individual PE surfaces. Moreover, the literature does not provide a consistent picture of the effect of roughness on cell adhesion because of the different definitions of surface topography at different length scale used by different authors. It is therefore difficult to determine or compare the effect of roughness on cell adhesion directly.

7.2.1.6 Effect of Substrate Structure on Cell Adhesion

Araujo et al. (2013) \(^{28}\) showed that osteoblasts could be successfully cultivated in CS-CRG PEC scaffolds both with and without hydroxyapatite and dexamethasone addition. However, such scaffold structures may enhance cell adhesion over flat 2-D films since these three-dimensional structures more closely mimic the native environment of the tissue due to the macro, micro and nanofibre scaffold structures that were observed. These factors potentially provide more contact points with the cells \(^{31}\) and a close match to the complex multi-variant structure of native tissues \(^{319}\). In addition, scaffolds can also entrap the cells making it difficult to remove them through washing steps. As such, their complex three-dimensional (3-D) structures can result in ambiguous consequences such as entrapment of the cells that were not specifically bound to the surface which can be washed away on 2-D films \(^{30}\). Therefore, films allow the measurement of cell adhesion in the absence of complex
multivariance of 3-D scaffolds. Also, the different physical properties (e.g. pore sizes and interconnectivity) of different materials can make the comparison of their intrinsic properties more complicated. For these reasons, in this thesis the scaffold struts have been modelled as thin films. In this way, the fundamental interaction of cells with the material can be investigated while avoiding the effects of structural parameters found in scaffolds.

7.2.2 Biological Properties of CS, CRG and PECs

7.2.2.1 Analogous Structure of CS and CRG to GAGs

The analogous structure of CS and CRG to the native tissue glycosaminoglycans (GAGs) may make the CS-CRG PECs potentially suitable as temporary tissue substitutes. The CRG structure was previously found to mimic the bioactivity of tissue-derived chondroitin-4-sulfate (C-4-S) and dermatan sulfate (DS) while the N-acetyl-glucosamine group in CS was found to interact with certain growth factors, receptors and adhesion proteins.

7.2.2.2 Biological Properties of CS

*In vitro* studies have shown that CS is non-cytotoxic towards myocardial endothelial cells, human microvascular endothelial cells, neurones, hepatocytes, chondrocytes, fibroblasts and keratinocytes. *In vivo* studies have also shown that CS possessed low cytotoxicity in human tissues such as skin and cartilage. It was previously found that low molecular weight (MW) CS exhibited a higher viability of L929 mouse connective tissue fibroblasts compared with the high MW CS. This was thought to be due to the synergistic effect created by the interaction between low-MW CS and the proteins secreted by the cells which maintain an optimal environment for cell growth. In addition, a high degree of deacetylation (DDA) enhanced attachment and proliferation of keratinocytes and attachment of fibroblasts. At high DDA, cells like the dorsal root ganglion and neurones attached and proliferated. This may have been due to the higher amino content in the higher DDA sample which results in a higher electrostatic interaction between the positive CS and the native negative surface charge of the cells which is due to the cell membrane phosphatidylcholine liposomes content. However, when the DDA of CS was around 50%, the dorsal root ganglion (DRG) did not attach and proliferate on the surface of the material. Instead, the DRG rearranged into cell clusters, through cell-cell rather than cell-material interaction. This may have been due to the high swelling of the CS at 50% DDA, which potentially makes the surfaces very hydrophilic, rough and with a high degree of surface mobility that could affect cell adhesion. Human microvascular endothelial cells (HMEC) were found to adhere well to CS and CS-heparin surface. The CS-Hep material could simultaneously prevent the adhesion of platelets, making them suitable for blood-contacting
Ultra-thin anti-fouling, anticoagulant and antibacterial coatings were also produced by using simultaneously positively-charged chitosan and negatively-charged sulfated chitosan through a layer-by-layer approach. It was shown that these coatings on glass or silicon were resistant to non-specific adsorption of proteins such as negatively-charged bovine serum albumin and positively-charged TGF-β1, and to the coagulation of human blood. The same coatings were also resistant to the formation of bacterial (Staphylococcus aureus) biofilm which is known to adhere and grow on polystyrene (PS), glass, Teflon and other materials.

### 7.2.2.3 Biological Properties of CRG

It was previously found that CRG gels were non-toxic to L929 cells and that no inflammatory response was observed after subcutaneous implantation in rats for one and two weeks. Similarly, McKim et al. (2016) have recently found that CRG does not induce cytotoxicity, or the expression of proinflammatory genes and does not increase cellular oxidative stress. CRG has also been shown to improve cell adhesion and viability, but its high water solubility renders it impractical for use as a cell substrate without crosslinking. Previous work on chondrocyte encapsulation using CRG hydrogels showed good cellular viability and proliferation for 21 days, and human adipose-derived stem cells (hASCs) were found to proliferate rapidly and express high levels of cartilage-specific markers. These results indicate that CRG may be a suitable substitute for cartilage repair.

Photocrosslinkable CRG was also used to encapsulate NIH-3T3 fibroblast cells, MC3T3 E1-4 preosteoblast cells, and human mesenchymal stem cells (hMSCs). Preliminary results indicated that these encapsulated cells possessed high cell viability (~75%) within the CRG gel for long periods (up to 21 days).

### 7.2.2.4 Biological Properties of PECs

The biological properties of the CS-CRG complexes have also been determined. It has been previously shown that CS-CRG complexes impose no cytotoxic effects in vitro and they offer potential as medical materials. Carneiro et al. (2013) have demonstrated that fibroblasts were more adherent on the individual CS and CRG surfaces compared to the CS-CRG PEC surfaces. This was thought to be due to some loss of interaction and non-specific binding sites at the PEC-cell surfaces. This was corroborated by another study where 3T3 mouse fibroblast cell attachment was higher on CS and CRG compared to the CS-CRG PECs. Normal human keratinocytes were found to adhere and spread preferentially on the CS surface as opposed to the chitosan-hyaluronic acid (CS-HyA) or chitosan-chondroitin sulfate (CS-ChS) surfaces. The biocompatibility of CS-CRG films has also been studied by Corvaglia et al. (2016) using a model tumour cell line (HeLa). These showed poor cell adhesion, as highlighted by the spheroid shape of the cells and cell cluster formation as opposed to...
the flattened cell shape typically observed on bioadhesive surfaces such as a PS (polystyrene) control. Nonetheless, the matrices were not detrimental to the viability of the cells. The reason for such results was thought to be due to the superficial charge (complexing of the charged groups between CS and CRG), surface morphology (steric hindrance) and matrix super hydrophilicity (reducing protein adhesion). Because of these properties, tumour selective cell growth was proposed as a potential application, since some tumours are anchorage-independent cells and can proliferate as cell clusters.

7.2.3 Biofouling and Anti-Adhesive Surfaces

7.2.3.1 Biofouling and Examples of Applications

In this thesis, the term "biofouling" is used for the non-specific attachment of unwanted proteins and cells on biomaterial surfaces that impair their function. The problems include associated implant rejection, malfunction of biosensors and the spread of infectious diseases. The most widely reported infections are in applications where the medical device's bridge between the inside and the outside of the body. Examples include the urinary catheter which is the most commonly used medical device and the second highest cause of infections after ventilator-associated infections.

7.2.3.2 Biofouling Process

Proteins responsible for cell attachment can adsorb onto the surface of materials through physical or chemical means such as wettability, electrical charge, surface roughness, pH and chemical groups such as carbon, amine and oxygen. The modes of protein and cell attachment to a surface are shown below in Figure 7.1. Cells normally require protein adhesion before they can attach to the surface. In an ionic solution, such as cell media, first an electric double layer is established at the surface of the material which in turn modulates the adsorption and conformation of proteins. These proteins, such as fibronectin, once attached to the surface can mediate cell adhesion through integrins in the cell membrane. When a cell is bound to the surface, extracellular matrix can be produced which provides further cues for cell spreading and proliferation. In other cases, the adhesion of a small amount of protein on a surface can reduce unwanted fouling. For example, platelet adhesion was inhibited by surfaces that were coated with small amounts (5 ng/cm²) of fibrinogen that provides biomaterials with anti-thrombogenic properties for good blood compatibility.
Figure 7.1 - Schematic representation of the dynamic proliferative fouling process which takes place over many length scales. Adapted from Magin et al. (2010) 333. The adhesion mechanism of protein and cell is annotated in the figure.

7.2.3.3 Anti-Adhesive Surface as a Non-Biofouling Strategy

Various strategies have been employed to control biofouling on materials surfaces. The main strategy used to control biofouling is through anti-adhesive surfaces 337. Often these anti-adhesive strategies involve the steric repulsion and the formation of a hydration layer which resist biofouling 338. During steric repulsion, the foulant (e.g. protein) compresses the polymer chains on the surface, which restrains the polymer chains. This creates unfavourable entropic loss which makes the adsorption of the protein entropically unfavourable 339. The formation of hydration layers is demonstrated by the highly organised surface-water (hydration layer) created in polyethylene glycol (PEG) via hydrogen bonding. The disruption of this bound water layer by protein association is energetically and kinetically unfavourable which in turn prevents protein adhesion 340. Zwitterionic molecules attract a thicker hydration layer (more water molecules per monomer unit) compared to PEG 341. This is due to the strong attraction of electrostatic interactions and the greater mobility of the water molecules in the hydration layer. Hence, zwitterionic molecules such as carboxybetaine (CB) or sulfobetaine (SB) are more fouling-resistant compared to PEG 335,342. These anti-fouling zwitterionic coatings have been applied to silicone urinary catheters to reduce biofilm formation by about 80 % when compared to
that of unmodified silicone catheters. It is likely that PEs and PECs utilise the same mechanism that zwitterionic molecules use due to their large macromolecular and oppositely-charged chains.

### 7.2.3.4 Polysaccharides as Anti-Adhesive Surfaces

In a manner that is similar to PEG, anionic polysaccharides possess potential anti-adhesive characteristics due to their surface topography (roughness), physiochemistry (surface free energy, hydrophilic or hydrophobic, cationic or anionic behaviour), polysaccharide molecular weights and experimental immobilisation conditions. In addition, negatively-charged polysaccharides have been considered to be important for the electrostatic repulsion of cells due to the negatively-charged cell surface glycosaminoglycans such as hyaluronic acid present around the cell membranes. Hyaluronic acid coatings on glass slides have been successfully used to reduce the adhesion of *S. epidermidis* and *E. coli* by several orders of magnitude compared to unmodified glass slide. Commercial products of hyaluronic acid coatings are currently used to minimise post-surgery adhesion (tissue attachments on implants) including Hydak® from Biocoat Inc, Horsham, Pa. and Incert-S® from Anika Therapeutics Inc., Bedford, Mass. Heparin is another widely-used polysaccharide for the coating of coronary stents, however to date this has not been found to reduce thrombosis and restenosis over non-coated coronary stents. Heparin has been blended with CS to form films with antithrombotic properties that prevent platelet adhesion and thrombosis. Commercial products of heparin as antithrombotic coatings include Bioline Coating® from Maquet Cardiopulmonary GmbH, Rastatt, Germany, Bioactive Surface CBAS® from Carmeda AB, Upplands Väsby, and Trillium® biosurface from Medtronic, Inc., Minneapolis, Minn. Dermatan sulfate has been polymerised with polyurethane at different degrees of substitution and was found to reduce the in vitro adhesion of platelets, red blood cells, macrophages and bacterial cells (*E. coli*) on all copolymer films compared to the unmodified polyurethane. An interesting study carried out by Bratskaya et al. (2007) showed the antibacterial and anti-adhesive properties of CS-CRG-based coatings against two *Enterococcus faecalis* strains that were isolated from infected biliary stents. The results showed that the multilayers were better at reducing bacterial deposition in comparison with the glass control and covalently grafted CS. The adhesion of negatively-charged bacteria was higher on CS terminated multilayers compared to the CRG terminated multilayers. However, the CRG terminated multilayers showed no antibacterial activity.
7.2.4 Aim and Objectives

In this chapter, the bioactivity of PECs was determined using three different types of cells and the highly adhesive protein tropoelastin. The effect of pH preparation on the biological properties of PECs is discussed. The overall aim is to examine the cell and protein interactive properties of CS, CRG and PECs. The objectives of this chapter were to investigate:

1. The cytotoxicity of the materials (CS, CRG and PECs) on the human fibrosarcoma (HT1080) cell line.
2. How the composition of CS, CRG and PEC film coatings can affect the adhesion and spreading of fibrosarcoma (HT1080), human dermal fibroblast and chondrosarcoma (SW1353) cells.
3. The adhesion of proteins using the model molecule tropoelastin to CS, CRG and PEC films.
7.3 Materials and Methods

7.3.1 Materials

All chemicals were of analytical grade and purchased from Sigma-Aldrich, UK unless stated otherwise. Low molecular weight chitosan powder with a 93.1 % degree of deacetylation, κ-carrageen and type I collagen from bovine Achilles tendon were purchased for making the films. Glutaraldehyde (GA) was purchased for fixing the cells in films. DAPI (4′,6-diamidino-2-phenylindole) and rhodamine phalloidin (Thermo Fisher Scientific, USA) were obtained to stain the cells. Dulbecco’s modified Eagle’s medium (DMEM), ethylenediaminetetraacetic acid (EDTA), foetal bovine serum (FBS) and 100 units penicillin and 0.1 mg streptomycin per 1 ml of media were used for the cell studies. CytoOne treated tissue culture plate polystyrene (TCPP) of 24, 48, 96-well plate sizes were purchased from Star lab and were used for cell studies. Tropoelastin was kindly provided by Professor AS Weiss at the University of Sydney for the protein adhesion study.

7.3.2 Film Preparation for Cell Culture

The PEC gels fabricated at various pH values (3, 5, 7.4 and 9) in Chapter 4 were homogenised and diluted with type 1 water to a concentration of 0.5 % w/v. For each sample, 200 µl of the diluted PEC suspension was added to each well in a 48-well plate and dried at room temperature in a laminar flow hood for 24 hours. Similarly, 200 µl of CS and CRG solutions prepared at 0.5 % w/v were solvent-cast in a 48-well plate for 24 hours. The dried CS films were then neutralised with 1 ml of 1 M NaOH overnight while shaken at 100 rpm using a rotating flatbed (TiMix, Labortechnik, Germany). The CS films were then washed twice with 1 ml of type 1 water followed by soaking in water for 1 hour to remove any excess acid or alkali remaining from the preparation process. For the cell studies on HT1080 cells, a positive control was prepared by adding 300 µl of soluble bovine type 1 collagen to the wells at a concentration of 10 µg/ml and incubated overnight in the fridge to form a collagen monolayer on the surface of the wells. Bovine serum albumin (BSA) was used as a negative control since it prevents the adhesion of the HT1080 cells onto the well surface during short-term (2 h) culture. Non-specific binding to the well surface not covered by the collagen monolayer was blocked with BSA by adding a volume of 300 µl BSA (1 % w/v) solution for 1 h then washing gently twice with 300 µl phosphate buffer solution (PBS) prior to cell/media addition. For long-term cell studies (2.5 d), the materials were pre-sterilised by spraying with 70 % ethanol (EtOH) followed by drying overnight in a sterile laminar flow hood.
7.3.3 Fibrosarcoma Cells

7.3.3.1 Cell Culture

Human fibrosarcoma (HT1080) cells were obtained from the European Collection of Animal Cell Cultures (ECACC), Porton Down, UK. Cells were seeded at a density of $2.5 \times 10^5$ cells per 75 cm$^2$ tissue culture flask in DMEM supplemented with 10 % (v/v) FBS and 100 units penicillin and 0.1 mg streptomycin per 1 ml of media. The cells were grown at 37 °C in a humidified incubator with 5 % CO$_2$ and 95 % air. The cells were passaged every 2-3 days before reaching 80 % confluency. For passaging, the medium was removed, and the cells were rinsed with PBS, followed by incubation in 5 ml of 0.5 g trypsin/0.2 g EDTA in PBS at 37 °C for 3 min until cells started to detach from the tissue culture flask. DMEM culture medium containing 10 % (v/v) FBS was added to inactivate the trypsin. After centrifugation at 1000 rpm for 3 min (Hermle Z300K, Germany), the supernatant solution was removed, and the cell pellet was resuspended in fresh medium supplemented with 10 % FBS and added at the required dilution to new flasks containing 18 ml of media.

7.3.3.2 Cytotoxicity Study

The CS, CRG, CS-CRG PECs prepared at pH 3, 5, 7.4 and 9, PTFE and PP films with a diameter of 8 mm and an approximate weight of 8.6 mg were pre-sterilised with 0.5 ml of 70 % v/v ethanol and left overnight in a sterilised laminar flow hood for the ethanol to evaporate. The films were then incubated in 1 ml of media supplemented with 10 % FBS for 3 days. Simultaneously, a 24-well plate was seeded with $1.5 \times 10^5$ HT1080 cells in 1 ml of media supplemented with 10 % FBS and 100 units penicillin and 0.1 mg streptomycin per 1 ml of media. The cells were incubated at 37 °C for 24 hours in a humidified incubator with 5 % CO$_2$ and 95 % air to form a cell monolayer. Subsequently, the media present in the cell culture wells was replaced with the film medium extract cultured for an additional 3 days. Ethanol was used as the positive (cytotoxic) control by adding 200 µl of 70 % v/v ethanol to each well to a final ethanol concentration of 12.7 % v/v. PP samples obtained from Falcon tubes were used as the negative controls due to their high inertness and non-cytotoxicity towards the cells. The media were removed at the end of the incubation period, and the cells were rinsed gently with PBS. Phase-contrast micrographs of the bound cells were taken for each material as detailed in Section 7.3.3.3. The cell number was determined using the pNPP assay as described in Section 7.3.3.3. The pH is important for cell viability, and it is therefore crucial to measure this change in the media since the PECs were prepared at different pH conditions. The pH of the medium was also found by using the absorbance values at 550 nm of phenol red present in the cell media and calibrating against a standard curve concluded from the absorbance values at 550 nm of phenol red against pH in the range of 4 – 8. All
sample measurements were carried out in triplicate and values were reported as mean ± standard deviation.

7.3.3.3 Short-Term Adhesion Study

Before cell adhesion analysis, HT1080s were detached from the cell culture flask with 0.5 g trypsin/0.2 g EDTA for 2 min, centrifuged and resuspended to a density of 2.5 x 10^5 cells/ml in DMEM without FBS. Then 200 µl of HT1080 cell suspension was added to each film for 90 min in DMEM either in the presence or absence of 10 % FBS. The cells were fixed with 5 % glutaraldehyde then viewed using an inverted phase-contrast microscope (ZEISS observer Z1, Germany) fitted with a ZEISS camera (Axiocam 503 mono, Germany).

Identification of the cells followed the same protocol except that loosely bound cells were removed with 2 x 200 µl DMEM washes. Bound cells were detected using the p-nitrophenyl phosphate (pNPP) substrate (Sigma) by adding 300 µl of lysis buffer containing 81 mM trisodium citrate, 31 mM citric acid, 0.1 % v/v Triton X-100, 1.85 mg/ml pNPP substrate (pH 5.4) to each well containing the cells for 24 hours at 4 °C. Subsequently, 200 µl of 2 M NaOH was added. A separate 96-well plate was used for the absorbance reading at a wavelength of 405 nm using a SpectroStar Nano microplate reader (BMG Labtech, Germany). The fibrosarcoma cell adhesion assays were performed in quintuplicate, and the values were reported as a means ± standard deviation.

7.3.3.4 Long-Term Adhesion Study

As for short-term cell analysis, the cells were imaged using microscopy and separately quantified with the pNPP assay. HT1080 cells were prepared as for short-term analysis except that they were resuspended at a density of 1.5 x 10^5 cells/ml in DMEM with 10 % (v/v) FBS. Then 200 µl of cells were added per well for 2.5 days at 37 °C, 5 % CO₂ and 95 % air. A separate well plate was used for the quantitative measure of cell viability using the pNPP assay as described in Section 7.3.3.3. All sample measurements were carried out in quintuplicate and values were reported as mean ± standard deviation. In addition to phase-contrast microscopy, the cells were stained and viewed using fluorescence microscopy. The cells were fixed by adding 50 µl of 25 % (w/v) glutaraldehyde directly to the cells to a final concentration of 5 % (v/v). After fixing the cells with glutaraldehyde, the cells were permeabilized in 200 µl 0.5 % w/v Triton X-100 in PBS for 5 min and washed three times with 1 ml of PBS. The cells were fluorescently stained to make them visible against the swollen rough film surfaces. Rhodamine conjugated to phalloidin was diluted 1:10000 (i.e. 2 µl rhodamine-phalloidin (stock 200 units, 6.6 µM/ml) in 20 ml in PBS) and added at 200 µl per well for 45 min. The samples were subsequently washed three times with 1 ml of PBS. The cells were also stained with 1:5000 diluted
4',6-diamidino-2-phenylindole (DAPI) in water using 200 µl of DAPI per well for 5 min. The samples were then washed three times with 1 ml of water. The wells were then viewed using an inverted fluorescent microscope (ZEISS Axio Observer, Germany) fitted with a ZEISS Axio cam 503 mono camera. DAPI stains the DNA in the nuclei of the cells showing as blue and the rhodamine phalloidin stains the actin in the cytoskeleton of the cells showing as red.

7.3.4 Primary Human Dermal Fibroblast Cells

7.3.4.1 Cell Culture

Primary human dermal fibroblast cells were obtained from ECACC, Porton Down, UK. Cells were seeded at a density of 2 x 10⁶ cells per 75 cm² tissue culture flask in DMEM supplemented with 10 % FBS and 100 units penicillin and 0.1 mg streptomycin per 1 ml of media. The cells were grown at 37 °C in a humidified incubator with 5 % CO₂ and 95 % air. The cells were passaged to a ratio of 1:3 every 4-5 d before reaching 80 % confluency. The cells were subcultured as detailed for fibrosarcoma cells in Section 7.3.3.1.

7.3.4.2 Long-Term Adhesion Study

Before cell adhesion/spreading experiments, fibroblasts were detached from the cell culture flasks centrifuged and resuspended as for HT1080 cells. After film preparation, 200 µl of fibroblast cells were added to the wells at a density of 0.5 x 10⁵ cells/ml per well for 3 days. The positive and negative controls were TCPP and 12.7 % ethanol, respectively. The viable cells were quantified using the pNPP assay and the cell morphology/structure imaged using fluorescent microscopy as described in Section 7.3.3.3 and 7.3.3.4, respectively. All sample measurements were carried out in quadruplicate and values were reported as means ± standard deviations.

7.3.5 Chondrosarcoma Cells

7.3.5.1 Cell Culture

Human chondrosarcoma (SW1353) cells from the American Type Culture Collection (ATCC) were grown in Dulbecco’s Modified Eagle Medium/Nutrient Mixture F-12 GlutaMAX™ supplement with 10 % (v/v) foetal bovine serum and 100 units penicillin and 0.1 mg streptomycin per 1 ml of media. Cells were seeded at a density of 3.75 x 10⁵ per 75 cm² tissue culture flask. The cells were grown at 37 °C in a humidified incubator with 5 % CO₂ and 95 % air. The cells were passaged every 3-4 days before reaching 80-90 % confluency. For passaging, the same method was used as presented earlier in Section 7.3.3.1.
7.3.5.2 Long-Term Cell Adhesion

The films were sterilised with several sprays of 70 % ethanol and left to evaporate for 1 hour, then washed with PBS and hydrated overnight using 0.5 ml of media containing 10 % (v/v) FBS. 200 µl of cells were seeded onto each sample at $5 \times 10^3$ cells/cm$^2$ in a 48-well plate. Cells were maintained under standard cell culture conditions (5 % CO$_2$, 95 % humidity and 37 °C) for 4 days. After culturing the cells, the media from the wells was removed, and the wells were washed twice with 1 ml of PBS to remove any loosely bound or non-attached cells. The cells were lysed by adding 100 µl of buffer containing 2 % v/v Triton X-100 in distilled water per well for 90 min at RT. Subsequently, 100 µl of LDH detection substrate (Sigma) prepared according to manufacturer instruction was added and incubated for 20 min until colour had developed. From each well, 100 µl was transferred to a new 96-well plate before the absorbance was read at 490 nm using a SpectroStar Nano microplate reader (BMG Labtech). Cell adhesion on films was performed in triplicate and values were reported as mean ± standard deviation.

7.3.6 Protein Adhesion

7.3.6.1 ELISA of Tropoelastin

The method was modified from Bax et al. (2011) $^{310}$. Briefly, discs of 8 mm diameter of CS, CRG, PECs (prepared at pH 3, 5, 7.4 and 9), PTFE and PS were placed in a 48-well plate. A 1 mg/ml stock solution of tropoelastin solution was diluted with PBS to a concentration of 20 µl/ml, and 1 ml of the diluted solution was added to each well for 1 h at RT. Unbound tropoelastin was removed by aspiration, and the samples were washed with 3 x 0.4 ml aliquots of wash buffer (PBS with 0.1 % w/v BSA). Non-specific antibody binding to the samples was blocked with 5 % (w/v) BSA in PBS for 1 h at RT. Following BSA blocking the samples were washed twice with 2 x 0.4 ml wash buffer, followed by incubation with 0.3 ml of 1:1500 diluted mouse anti-elastin antibody (BA-4) in wash buffer for 1 h at RT. The antibody solution was removed and the samples were washed 3 x with wash buffer before incubation with 0.3 ml of 1:5000 diluted goat anti-mouse secondary antibody coupled to horseradish peroxidase (HRP) (DAKO, UK) for 30 min at RT. The secondary antibody solution was removed and samples were washed with 3 x 0.4 ml wash buffer for 5 min per wash and then 3 x 0.4 ml wash buffer for 15 min. The samples were transferred to a new 48-well plate and 0.3 ml 3,3′,5,5′-tetramethylbenzidine (TMB) (ThermoFischer Scientific) solution added. After 30 min, the plates were agitated to develop a blue colour. Aliquots of 0.2 ml were transferred to a 96-well plate before the absorbances were read at 652 nm. A negative control that did not include tropoelastin coating was included. All sample measurements were carried out in triplicate and values were reported as mean ± standard deviation.
7.4 Results

7.4.1 Fibrosarcoma Cells

7.4.1.1 Cytotoxicity

In this study, the cytotoxicity towards HT1080 cells of media incubated with film materials was investigated. It was shown that all polysaccharide films did not adversely affect the viability of a preformed cell monolayer (Fig. 7.2). The media incubated with the negative control PP was within the standard error of the toxicity of the polysaccharide materials. In contrast, the positive control including 12 % EtOH clearly reduced the cell number due to its cytotoxic effect.

Figure 7.2 – Cytotoxicity of media pre-incubated with films towards an HT1080 monolayer. The materials studied include carrageenan (CRG), chitosan (CS), the negative control (non-toxic) polypropylene (PP), the positive control (toxic) ethanol (EtOH) and polyelectrolyte complexes (PECs) prepared at pH 3, 5, 7.4 and 9.

It is possible that the resultant PEC film pH may alter the pH of the DMEM which in turn could adversely affect the cell viability. As such, the change in pH of the cell-media was measured using the change in the absorbance of the phenol red present in the DMEM. The pH of the medium films extracts ranged from 7.6 to 8.1 for the materials tested (Table 7.1).

Table 7.1 – Phenol red absorbance at 550 nm of DMEM incubated with the disc materials. These were used to calculate the pH of the film medium extract used in the cytotoxicity study.

<table>
<thead>
<tr>
<th>Material</th>
<th>CRG</th>
<th>pH 3</th>
<th>pH 5</th>
<th>pH 7.4</th>
<th>pH 9</th>
<th>CS</th>
<th>PP</th>
<th>EtOH</th>
</tr>
</thead>
<tbody>
<tr>
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<td>7.84</td>
<td>7.6</td>
<td>7.7</td>
<td>7.8</td>
<td>7.84</td>
<td>8</td>
<td>8.1</td>
<td>8.1</td>
</tr>
</tbody>
</table>
7.4.1.2 Adhesion Analysis

Cell adhesion analysis showed no significant cell attachment to both the PEC and individual PE films. In fact, the cell adhesion on the polysaccharides was lower than the cell adhesion on the negative control film surface (BSA) (Fig. 7.3). Phase-contrast optical microscopy images of films showed that cells spread readily to both collagen (positive control) and TCPP after 90 min showing phase dark cells (Fig. 7.4). Conversely, no spreading was observed on any of the films composed of CS, CRG, PECs and BSA with rounded phase bright cells present.

Figure 7.3 – Attachment of HT1080 cells as measured using a pNPP assay. The materials studied include carrageenan (CRG), chitosan (CS), the positive control collagen (Coll), the negative control bovine serum albumin (BSA), tissue culture plate polystyrene (TCPP) and polyelectrolyte complexes (PECs) prepared at pH 3, 5, 7.4 and 9. The cells were cultured on the films for 90 min (short-term cell study).
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Figure 7.4 – Phase-contrast optical micrographs of HT1080 cells at a magnification of 200 x. The cells were cultured on the films for 90 min (short-term cell study).
Cell analysis was also conducted after long-term culture (2.5 d) (Fig. 7.5). The cell-derived absorbance in the long-term study also matches to the results found for the cells in the short-term study presented in Figure 7.3. Indeed after 2.5 days there was still minimal cell adhesion to all PECs, CS and CRG. The only difference is that in the long-term study, the cells exhibited a similar cell-derived absorbance on the collagen, BSA and TCPP surfaces. Low cell-derived absorbance was measured on CS, CRG and all PECs compared with the positive controls (Coll. and TCPP). Some cell-derived absorbance was noted on pH 7.4 and pH 9 but is much lower than the positive controls.

![Figure 7.5 - Measurement of HT1080 cells using the pNPP assay after 2.5 days (long-term cell study) in culture.](image)

The surface morphology of the cells and films are shown in the phase-contrast micrographs presented in Figure 7.6. It can be observed that the films tested in the long-term adhesion study were rough which can make it more difficult to distinguish between the cells and the material. To improve the resolution of the cells they were stained using fluorescent dyes (Fig. 7.7).
Figure 7.6 – Optical micrographs of HT1080 cells on films at a magnification of 100 x. The cells were cultured on the films for a duration of 2.5 d (long-term cell study). Arrows in red are pointing towards cells present on rough surfaces.
Figure 7.7 presents the fluorescent micrographs of DAPI stained nuclei of the cell. It was observed that the cells were distributed homogeneously on the collagen, TCPP and BSA surfaces. Conversely, the cells were clustered together on the CS, CRG and PEC films formed at all pHs.

Figure 7.7 – Fluorescent micrographs of DAPI stained HT1080 cells. The cells were magnified at 100 x. The cells were cultured on the films for 2.5 d (long-term cell study. Representative photos from three replicates are shown for each material.
Figure 7.8 presents the fluorescent micrograph of DAPI (stains the nuclei of the cell blue) and rhodamine phalloidin (stains the cytoskeleton of the cell orange). The staining of the nuclei can show that a cell is present, but it will not show the cell morphology. Therefore, the staining of the cytoskeleton can indicate the extent of cell spreading. It was noted that the cells cultured on the collagen, TCPP and BSA surfaces were more spread with cellular projections compared with the accumulation of rounded cells on the polysaccharide films.

Figure 7.8 – Fluorescent micrographs of HT1080 cells labelled with DAPI (blue) and rhodamine phalloidin (orange) at 100 x magnification. The cells were cultured on the films for a duration of 2.5 d (long-term cell study).
7.4.2 Primary Human Dermal Fibroblast Cells

7.4.2.1 Adhesion Analysis

Primary human dermal fibroblast cells were cultured for 4 days on CRG, PEC pH (3, 5, 7.4 and 9), CS, TCPP and TCPP containing 12 % ethanol were quantified using a pNPP assay (Figure 7.9). The CS, CRG and PEC films prepared at pH 3 and 5 exhibited slightly higher cell number than the PEC films prepared at pH 7.4 and 9. However, the cell-derived absorbance of all PEC films were approximately within experimental error of the cell-derived absorbance of ethanol (yellow line). The positive control (TCPP) exhibited approximately twice as many cells than on the polysaccharide films. The cell media without the cells was used to provide a background absorbance value (red line).

When stained with DAPI and rhodamine phalloidin, the cells found on the polysaccharide surfaces were exhibiting a clustered round morphology presumably due to greater cell-cell affinity compared to the cell-surface affinity (Fig. 7.10). The cells were small and round in the negative control wells. In contrast, the cells on the TCPP surfaces were flattened and possessed an extensive cytoskeleton.
Figure 7.10 – Fluorescent micrographs of primary human dermal fibroblasts labelled with DAPI (blue) and rhodamine phalloidin (orange) at 100 x magnification. The cells were cultured on the films for a duration of 4 d (long-term cell study).
7.4.3 Chondrosarcoma Cells

7.4.3.1 Adhesion Analysis

Unlike other cell-line experiments used in this thesis, the films used for the chondrosarcoma study were preincubated with serum prior to cell incubation.

Figure 7.11 clearly shows a low number of chondrosarcoma on the polysaccharide surfaces after 4 days in culture. There were approximately 20 x more cells present on the TCPP surface compared to the PECs. The addition of ethanol reduced the cell-derived absorbance to background (no cell) levels.

Optical micrographs of the chondrosarcoma are consistent with the absorbance values. The cells on the CRG films were found to be completely spherical and clustered. As shown before in Figure 7.6, the PEC films swell in the media in the long-term studies which make the cells difficult to distinguish from the surfaces of the films. Nevertheless, the cells on the PEC films were also found to be spherical and clustered with little spreading evident. The cells on the CS films were phase dark and spread although far fewer cells were present compared to the TCPP surface. The cells on the positive control (TCPP) were phase dark and spread, whereas the cells on the negative control (TCPP/ethanol) surface were small, round and phase bright.
Figure 7.12 – Phase-contrast micrographs of chondrosarcoma cells cultured for 4 d (long-term cell study) on films and viewed at a magnification of 100 x. Arrows in red are pointing towards cells present on the rough surfaces.
7.4.4 Protein Adhesion

7.4.4.1 ELISA of Tropoelastin

As cell adhesion to the PECs was low in the presence of serum, the ability of these materials to adsorb proteins was examined. To this end, enzyme-linked immunosorbent assays (ELISA) are used to detect the presence of the adhesive protein tropoelastin bound to the materials surfaces. Figure 7.13 shows that the PEC materials display limited physisorption of tropoelastin when compared to the individual PEs, PTFE and PS controls. The positive and negative controls include and exclude the presence of tropoelastin, respectively. However, in both controls, the secondary anti-mouse antibody and TMB substrate are included. Therefore, absorbance in the absence of tropoelastin indicates non-specific absorption to the material by either the primary or secondary antibody. The PECs showed the lowest tropoelastin adsorption compared with the rest of the materials tested (Fig. 7.14). The CRG material exhibited around twice as much absorbance compared to the CS and PTFE materials in the presence of tropoelastin, however this persisted in the absence of tropoelastin. Presumably, this is due to the swelling and disintegration of the CRG, which allows the diffusion of the tropoelastin, secondary anti-mouse antibody and TMB substrate into the material. This is particularly evident from Figure 7.14 and Figure 7.15 showing a pronounced blue colouration of the CRG film under the positive and negative conditions. The PS material exhibited the highest level of specific tropoelastin adsorption followed by CS and PTFE (Fig. 7.13). The PECs prepared at pH 5, 7.4 and 9 absorbed more tropoelastin through diffusion than the pH 3 PEC and CS discs as shown by higher absorbance values in the positive compared with negative condition (Fig. 7.13 and 7.15). This may be due to the higher swelling that was observed in PECs prepared at high pH within the first hour compared with the PECs prepared at low pH (Fig. 5.1). The negative controls (without tropoelastin) gave absorbances that were similar to the empty wells for almost all the materials except for CRG. It can be concluded that the tropoelastin strongly adhered to the CS, PTFE and PS surfaces but not on the PEC surfaces, showing that these are highly non-biofouling for this protein, particularly evident in the pH 3 PEC material.
Figure 7.13 – The absorbance of TMB on materials with tropoelastin (positive control) and without tropoelastin coating (negative control). The materials tested for tropoelastin adhesion include carrageenan (CRG), chitosan (CS), polytetrafluoroethylene (PTFE), polystyrene (PS), polyelectrolyte complexes (PECs) prepared at pH 3, 5, 7.4 and 9 and empty (no sample).

Figure 7.14 – Image of the 48-well plate with the films and substrates inside the wells. The top half of the plate contains materials coated with tropoelastin (positive control) and the bottom half was not coated with tropoelastin (negative control).
Figure 7.15 – Images of the washed films after tropoelastin detection showing the difference between the absorption and adhesion of the tropoelastin to the films. The carrageenan (CRG) sample is not present due to disintegration.
7.5 Discussion

7.5.1 Cell Studies

7.5.1.1 Cytotoxicity

Before conducting cell adhesion studies the cytotoxicity of the materials was first established using the fibrosarcoma cell line. Using PEC conditioned media, the materials were found to be non-toxic towards fibrosarcoma cells. This agrees with a previous study, where CS-CRG complexes ranging from 0.1 and 3 mg/ml was shown to be non-cytotoxic to the 3T3 fibroblast cell line. By comparison, a CS-CRG PEC concentration of around 9 mg was used in the current study. Therefore, given that this elevated concentration of PECs did not exhibit a cytotoxic effect on the fibrosarcoma cell line, it can be concluded that CS-CRG PECs are non-toxic at the concentration studied.

7.5.1.2 pH Measurements

The cytotoxicity results also showed that the film extract media did not reach extremely low pH values that could be potentially detrimental to the cells. By using the phenol red as the pH indicator that is present in DMEM, the media incubated with PECs prepared at pH 3-9 were within the pH range of 7.6 to 8.1 (Table 7.1). This appears relatively high compared to the DMEM pH of 7.4-7.7 but may reflect the increase in pH when the well plate was removed from the incubator. The incubator contains 5 % CO₂ compared to the 0.04 % CO₂ in normal atmosphere. Therefore, it is possible that the CO₂ was released from the media, resulting in an increase in pH prior to the absorbance reading. Despite this limitation, it was shown that the PECs did not fundamentally alter the pH of the media and so this should not influence cell behaviour.

7.5.1.3 Fibrosarcoma and Fibroblast Adhesion

Cell adhesion analysis was conducted to establish whether CS, CRG and CS-CRG PEC films could support cell attachment. In this study fibrosarcoma cells from connective tissue and fibroblast cells from human dermal tissue were used to test the surface compatibility of CS, CRG and CS-CRG PEC films. Fibrosarcoma and fibroblasts possess a wide variety of receptors and attach to a variety of materials. Therefore, these cells were chosen as an initial test platform. Unlike many matrix proteins, such as collagen, CS and CRG are devoid of specific amino acid sequences (motifs) that interact with specific integrins on the above-mentioned cells. For example, there are no specific motifs such as GFOGER on the film surface that interact with integrin α2β1 that is present on HT1080 cells or RGD motif that interact with integrins α5β1 and α5β3 that are present on fibroblast cells. However, cells
can bind through “passive” interactions, that is through multiple weak non-specific bonds such as van der Waals, salt bridges, water exclusion, right surface roughness, topography, atoms or chemical functional groups such as carbon, amine or oxygen groups, hydrophobic and Coulombic forces. To compare this potential ‘passive’ interactions and ‘active’ interaction, BSA and collagen were used as a negative and positive control surface, respectively. Collagen is a structural protein in human tissues and is widely used in tissue engineering applications because of its excellent mechanical and cell-adhesive function, while BSA is a serum protein that prevents the adhesion of cells to the substrates. These controls allow comparison to native matrix interactions with those on the polysaccharide films.

7.5.1.4 Short-Term Study of Fibrosarcoma (HT1080 cells) with and without Serum Proteins Present in Media

In the short-term HT1080 cell study, the cell-derived absorbance was low (Fig. 7.3), with the cells also exhibiting a spherical cell morphology (Fig. 7.4). These data indicate that the cells bind weakly to the surfaces of the polysaccharide materials. The experiment was conducted both in the presence and absence of 10% FBS present in the cell media showing that surface adsorption of serum proteins did not influence the adhesion of these cells. Many sera-derived proteins for example fibronectin and vitronectin are involved in the adhesion and proliferation of cells. Therefore, the lack of cell binding suggests that the proteins present in FBS did not adsorb onto the surfaces and therefore could not act as intermediary adhesion molecules that bridge between the surface of the materials and the cells.

7.5.1.5 Long-Term Study of HT1080 Cells

In the longer-term HT1080 cell study, the films absorbed some of the cell media resulting in a rough surface (Fig. 7.6). This roughness was likely to have been caused by the fibrous PEC gel precipitates owing to the strong intra-crosslinking and high CRG content. A similar observation was found by Carneiro et al. (2013) where they concluded that the higher roughness in PEC films was caused by the increased heterogeneous fraction of CRG in the PEC. This surface roughness meant that the cell morphology could not be distinguished from the surface of the material using phase-contrast microscopy. Instead, fluorescent markers were used to visualise the cells against the rough surface (Figs. 7.7 and 7.8). This showed that HT1080 cells accumulated in cell clusters with little cell spreading on the PECs. The rounded cell morphology could be due to the critical dependence on adhesion for these anchorage-dependent cells. Therefore, the affinity of cell-cell interaction may have been greater than the cell-surface interaction, which resulted in cell cluster formation. This is indicative of surfaces that are non-adherent to the cells as was previously shown with CS-CRG films using HeLa model tumour cell lines. Interestingly in the long-term adhesion assay using HT1080 cells (Fig. 7.5),
the BSA showed extensive cell interaction. The high cell adhesion to BSA coated surfaces after long-term culture could be due to the enzymatic degradation of the BSA by the cells which ultimately exposes the underlying TCP surface. In addition, small amounts of fibronectin and vitronectin present in the serum can displace the albumin due to their larger surface areas, which also contributes to higher cell adhesion. Therefore, ethanol was used instead of BSA as the negative control. In summary, the cells were capable of binding to the positive controls, confirming that the polysaccharide surfaces were non-adhesive substrates.

### 7.5.1.6 Roughness Affecting Cell Adhesion

Although the roughness of the PEC films in the current study was not measured in either dry or wet condition with AFM, the macro-roughness was visually observed with the naked eye, and it was found that the PEC films produced at lower pH (pH 3 and 5) were rougher at the macroscale than those prepared at higher pH (pH 7.4 and 9). Others have found that HeLa (human cervical cancer cell line) cell adhesion was limited on the rough CS-CRG PEC surfaces (227 ± 26 to 309 ± 34 nm) compared with the smooth CRG films (140 ± 8 nm). Therefore it is possible that the macro-roughness of the PECs produced at lower pH may have reduced the adhesion of fibrosarcoma cells. Alternatively, this reduction in cell adhesion may have been due to the higher steric hindrance and/or high hydrophilic nature of PEC films (Fig. 5.1).

### 7.5.1.7 Chondrosarcoma Cell Adhesion

Chondrosarcoma cells were chosen as a model for chondrocytes as they occupy highly glycosylated cell niche in vivo. As such it was hypothesised that the similarities of CS and CRG to chondroitin sulfates and hyaluronic acid might provide the chondrosarcoma cells with similar receptor sites for adhesion and proliferation. However, like the other previous cell lines, chondrosarcoma cells showed very low adhesion to the polysaccharide surfaces when compared with the TCPP surfaces. Lower cell numbers were observed on the PECs than the CS or CRG films which may be due to the neutral charge and zwitterionic nature which provides a highly hydrophilic environment that may be entropically unfavourable for adhesion. Another potential explanation for the low adhesion on the polysaccharide films is the steric and electrostatic repulsion between the cell surface macromolecules and the macromolecules on the surface of the film. Previous reports show that chondrocyte cells bind to collagen but not hyaluronic acid (HyA) substrates. Chondrocytes synthesise their own hyaluronic acid (HyA), and it was found that only the cells on the collagen films produced their own extracellular matrix. For this reason, collagen is widely used as a cell adhesion enhancer. These authors proposed that the inhibition of adhesion and spreading of the chondrosarcoma cells on HyA coated surfaces was due to the excess HyA which results in steric exclusion and/or electrostatic
repulsion between the two hydrated layers (HyA from the cell membrane and the HyA present on the surface). This is consistent with the adhesion study of A6 frog kidney epithelial cells on a HyA film surface \(^{343}\). Like CS and CRG, HyA is a large, linear glycosaminoglycan polyelectrolyte composed of repeating disaccharide units (glucuronic acid and \(N\)-acetylglucosamine) with molecular weights ranging from \(10^6\) to \(10^7\) Da and extended lengths of 2-25 µm \(^{360}\). When HyA is present on the surface in large amounts, it was proposed that a layer of flexible molecule extending in the solution becomes repulsive. Therefore, interactions between the membrane of the cell and the surface can occur at a considerable distance from the adhesive surface (> 100-fold larger than the integrin-mediated focal adhesions) \(^{343}\). Due to the similarity between the PECs and HyA it may be that the same mechanism is taking place between the CS, CRG and CS-CRG surfaces and the glycosaminoglycans from the cells. Another study concluded that the HCS-2/8 human chondrosarcoma cells could not adhere to anti-adhesive poly-L-lysine-hyaluronic acid (PLL-HyA) films prepared by a layer-by-layer method \(^{361}\). The anti-adhesive properties of the films to the cells were thought to be attributed to the low rigidity, high water content and the gel-like character of PLL and HyA materials. This is consistent with the PECs produced in this study where the CS-CRG PECs were found to be soft, swollen and gel-like in character (Chapter 5). Therefore it appears that the PECs produced in this thesis are non-cell adhesive in a similar manner to native glycosaminoglycan molecules.

7.5.2 Protein Study

7.5.2.1 Tropoelastin Functions and Structure

Tropoelastin was used for protein binding assays due to its strong surface binding abilities \(^{310,362–364}\). Tropoelastin is a major elastic constituent of elastic tissues such as skin, vocal fold, lung, and elastic cartilage \(^{310}\). Tropoelastin is known to adhere very strongly to various surfaces due to the presence of both hydrophobic and hydrophilic domains within the same protein \(^{365}\). The hydrophobic domains consist of non-polar residues glycine, valine and proline, whereas the hydrophilic domains consist of lysine and alanine \(^{366}\). Tropoelastin is an asymmetric protein with an appended C-terminus cell binding foot and a long spring-like N-terminal open to intermolecular crosslinking \(^{367}\).
7.5.2.2 Measurement of Tropoelastin Adhesion

The amount of tropoelastin adsorbed on the surfaces can be measured spectrophotometrically by using an ELISA assay (Fig. 7.13). These are well established being extensively used to measure protein deposition onto surfaces. During this assay, when tropoelastin adheres to the surface, the primary antibody couples to the tropoelastin, followed by a coupling of the secondary antibody (Fig. 7.16). This secondary antibody is conjugated to the enzyme HRP which catalyses a TMB substrate to a coloured product. Elisa analysis showed that very little tropoelastin was adsorbed to the surfaces of the PEC materials.

3,3',5,5'-tetramethylbenzidine (TMB) → Blue Colour (Abs: 652 nm)

Coupled Horseradish Peroxidase (HRP)

2' Antibody (Goat Anti Mouse)

1' Antibody (Mouse Anti Elastin (BA-4))

Tropoelastin → Surface

*Figure 7.16 – A schematic diagram of the detection of tropoelastin when adsorbed onto surfaces showing the substrate and the mechanism for colour change when tropoelastin is present.*

This supports the cell adhesion analysis in the presence of serum where it was concluded that the PEC materials are highly non-biofouling. This is particularly so since tropoelastin is known to be highly surface adhesive. Indeed, tropoelastin obtained from the same supplier was previously shown to adhere to a wide range of surfaces, such as PTFE, PS and stainless steel. Hence the inability of tropoelastin to adhere to the polysaccharide surfaces indicates that the PEC materials are potentially non-adhesive to a wide range of protein-based biomolecules. Nonetheless, the slight colour change observed in the tropoelastin coated PEC films prepared at pH 5, 7.4 and 9 after the washing (Fig. 7.15) indicates that some tropoelastin may have adhered to or diffused into the films specifically as is schematically drawn in Figure 7.17. No colour change was observed in the absence of tropoelastin coating. CRG films differed from the other materials as they swelled and disintegrated significantly thereby entrapping the proteins and antibodies, resulting in a large colour change that was independent of tropoelastin adhesion. Tropoelastin was found to adhere to the surfaces of CS, PTFE and PS.
Figure 7.17 – A schematic diagram detailing the tropoelastin, primary antibody and secondary antibody interaction processes for each material. Adsorption of tropoelastin on chitosan (CS), polytetrafluoroethylene (PTFE) and polystyrene (PS) can be observed. PEC pH 3 film exhibited no adsorption or diffusion of tropoelastin within 1 hour. The other polyelectrolyte complexes (PECs) prepared at pH 5, 7.4 and 9 exhibited little diffusion or adhesion of tropoelastin. The CRG sample absorbed large quantities of tropoelastin and antibodies due to the swelling and disintegration of the sample.

7.5.2.3 Mechanisms of the Non-Adhesive Surfaces

The adsorption of many proteins to materials surfaces is energetically favourable due to the release of energetically unfavourable surface adsorbed water molecules and counterions associated with both the protein and material-solvent interface which provides the entropic driving force. The low adherence of tropoelastin to the PEC surfaces may be due to the highly hydrophilic, rough and mobile nature of surfaces where the interaction with surface associated water may be more energetically favourable than the interaction with the domains of the proteins to the hydrophilic surfaces (hydrophilic and hydrophobic incompatibility) (Fig. 7.18 and Fig. 7.19). For the same reason, polyethylene glycol (PEG) has been widely used as a non-biofouling material which may be due to its high swelling, hydrophilic and chemical properties. In recent years, polyacrylic acid (PAA), like PEG, has also been widely used as a grafted polymer to prevent adhesion of proteins due to the steric repulsion and excluded volume effects. In addition, PAA polyelectrolytes have shown exceptional protein resistant surfaces in high salt concentrations as these conditions suppress the “counterion evaporation” from proteins which is the major driving force for adhesion of proteins to PEs. Therefore non-fouling properties of the PECs developed in this thesis may be due to similar water-binding, steric repulsion and counterion evaporation mechanism.
7.5.2.4 Complete Summary of the Non-Adhesive Surface of PEC

Figure 7.19 summarises the current potential non-biofouling mechanisms of the CS-CRG PECs schematically. The reason for the highly anti-adhesive surfaces of CS-CRG PECs is potentially caused by one or more of the factors i.e. hydrophilicity, steric repulsion, excluded volume effects and charge repulsion.

- (1) Hydrophilic Surface - (Adsorption Unfavourable)
- (2) Steric Repulsion (Small and Large Macromolecular Chains)
- (3) Excluded Volume Effect (Space Already Occupied)
- (4) Negative Charge Repulsion from CRG

Figure 7.19 – A schematic diagram representing the non-biofouling mechanisms that may be involved in CS-CRG PECs. The blue arrow represents the desorption of cells and proteins.
7.5.2.5 Potential Applications

The strong anti-adhesive properties of PECs has good potential for a wide variety of applications such as the prevention of bacterial adhesion and the prevention of platelet adhesion to biomaterials which is the cause of undesirable biofilm formation and blood clotting, respectively. Other researchers have also investigated the adhesion of plasmatic (globular) proteins such as fibrinogen and bovine serum albumin on CS, CRG and CS-CRG PEC films. These proteins were tested because they are known to be involved in the first event of the adsorption process that occurs when a material comes into contact with blood. Both studies mentioned above have shown that heparin or sulfonated chitosan can reduce the attachment of the plasma proteins when compared to the unmodified chitosan surface. Plasma proteins are known to attach to the CS surfaces during thrombosis. This is consistent with the protein binding properties of CS films in this study as evidenced by adsorbing tropoelastin from the solution. Highly sulfated heparin or molecules with negatively-charged groups that are similar to CRG are known to provide a repulsive force to the negatively-charged proteins which can reduce the adsorption of proteins due to electrostatic interactions with the material. Other researchers have also shown that sulfated CS and CRG decrease the adsorption of proteins and were thought to be useful as anti-thrombogenic materials for blood contacting applications. Therefore, further study to examine biofilm and thrombus formation on the materials produced in this study represents an exciting area of future research.

7.6 Conclusions

Overall, the materials were found to be non-toxic towards the fibrosarcoma cells. The pH of the cell medium was not significantly affected by the PEC films produced at lower pH range. Thus, the PEC films prepared at various pH values were non-adherent rather than cytotoxic. Three general cells (fibrosarcoma, fibroblasts and chondrosarcoma cells) showed little cell adhesion on all polysaccharide materials. Cell clusters were formed on all the polysaccharide films with many cells exhibiting characteristics of apoptosis with a round cell morphology. The use of three different types of cells showing the same effect means that the low adhesion observed was not due to a one-off outcome but instead appear to be a general effect. Finally, the PEC films were highly non-adherent to the adhesive tropoelastin which indicates the potential for a wide variety of non-biofouling applications.
Chapter 8 – Overall Conclusions and Future Work

8.1 General Conclusions

The aim of this thesis was to explore the effect of pH on polyelectrolyte complex formation to control the degree of self-crosslinking of materials by avoiding the use of toxic chemical crosslinkers. The pH was found to have a significant influence on the viscoelastic properties of PEC gels and the mechanical properties of PEC films and scaffolds. The physicochemical properties of PEC gels were mainly affected by the strength of electrostatic interaction (SEI) whereas dry PEC films were mainly affected by the inter/intra-crosslinking (to some extent reliant on SEI) and composition. In addition to the composition and inter/intra-crosslinking, the mechanical properties of PEC scaffolds were also affected by the pore sizes.

The preparation pH controlled the degree of electrostatic interaction in the PEC gel formation. At lower pH ranges (2-6) the SEI in PEC gels was higher than the SEI of PECs prepared at higher pH ranges (7-12). This resulted in stronger self-crosslinking in the PEC gels prepared at pH 2-6. The strong interactions led to the formation of higher yield, solid content, viscosity and fibre content in PEC gels. The high fibre content in PEC gel precipitates prepared at low pH led to a rougher PEC film surface. The strong electrostatic bond interaction between NH$_3^+$ and OSO$_3^-$ was confirmed for pH 3-7.4 with FTIR at wavenumber, 1529 cm$^{-1}$. The FTIR wavenumber band was absent at higher pH due to the weaker electrostatic interaction caused by the deprotonation of CS. As measured with nitrogen, carbon and sulfur (NCS) elemental analysis, the PECs prepared at pH range 3-8 contained a higher fraction of CRG than the PECs prepared at pH range 9-12. The higher CRG content resulted in an increase in the hydrophilicity of the PEC films, inducing greater swelling in films prepared at lower pH. In addition, the strong electrostatic interaction at pH 3 and 5 resulted in the formation of strong intra-crosslinked and weakly inter-crosslinked PEC films and scaffolds. Therefore the PEC films prepared at lower pH range were less connected as a network, exhibiting lower strength and stiffness and, greater mass loss and swelling. A summary of the physicochemical properties of PECs produced at low pH is presented schematically in Figure 8.1.
The biological properties of the materials were tested in the form of films. The results show that the CS, CRG, CS-CRG PEC films exhibited highly non-biofouling properties. All polysaccharide materials used were found to be non-cytotoxic when tested with fibrosarcoma cells. However, the low cell viability, round cell morphology and cluster formation of all the cells (fibrosarcoma, fibroblast and chondrosarcoma) on the polysaccharide films, indicated that the materials were highly non-adhesive. Nonetheless, the lower adhesion of tropoelastin on the PEC films compared with the CS and CRG films shows that the PEC materials may possess high non-biofouling properties. The highly non-adhesive surface of PECs may have been caused due to several factors including high hydrophilicity, swelling, roughness, surface mobility and chemical groups which made it energetically unfavourable for both hydrophobic and hydrophilic molecules to attach readily to the surface.

The main advantage of the PEC system lies in its ability to control the physicochemical properties of materials by changing a broad range of reaction parameters. Furthermore, no auxiliary chemicals are used which may be beneficial to both the environment and the long-term effects upon the body. In addition, covalently crosslinked materials may be too stiff for some soft tissues, where the adhesion of cells is not desired. Examples of applications may be to use the PEC materials in the form of film or coating in brain, adipose, skin or articular cartilage implants due to their appropriate mechanical properties. It is, therefore, envisaged that the self-crosslinked CS-CRG PEC gels, films, and scaffolds would have high potential as crosslinker-free anti-fouling biomaterials.
8.2 Recommendations for Future Work

Further research upon PEs might extend their characterisation by using gel permeation chromatography/size exclusion chromatography-multi-angle light scattering detection (GPC/SEC-MALS) to find the molecular weights (MW) of CS and CRG. Circular dichroism (CD) and small angle X-ray diffraction and spectroscopy (SAXS) could be used to determine the structural conformation of PEs and PECs. The enthalpic and entropic change during complexation of CS and CRG may also be measured using isothermal titration calorimetry (ITC). Finally, atomic force microscopy (AFM) can be used to show the nanoscale surface roughness of films in both dry and wet conditions.

An interesting study may be to test the drug releasing properties of the PEC films prepared at different pHs. This is because the materials are self-crosslinked and the viability of the drug may not be affected as much as in the systems that require chemical crosslinking. Also many drugs are sensitive to elevated temperatures and PEC systems do not require heat during preparation; except when the concentration of the PEs is high which makes it less soluble. Furthermore, the electrostatic environment may be suitable for the slow and controlled release of charged drugs. However, the hypothesis is that the PECs produced at lower pH might elicit higher drug delivery rates due to the lower inter-crosslinking.

The potential anti-fouling properties of the materials may require further biological characterisation. It would be interesting to study the adhesion of different proteins (fibrinogen, laminin, vitronectin, albumin) and cells (muscle cells, neural cells, monocytes and neutrophils) on the surface of PEC films. This may provide a complete picture of the non-biofouling properties of the PEC materials. Experiments may be conducted on bacterial adhesion, using Gram-positive and Gram-negative bacteria for applications such as urinary catheters. Platelet adhesion and whole blood clotting studies may also be carried out to find applications for blood contacting devices such as arterial stents.

The materials may also be used for tissue engineering applications as blank slates. These blank slate PEC materials may be surface modified by incorporating signalling molecules such as peptides, arginyl-glycyl-aspartic acid (RGD) or collagen which can alter the interaction between the material and cell surface. Other functionalisation techniques have included layer-by-layer using heparin (Hep) and hyaluronic acid (HyA) to specifically target integrin binding sites present on cells, to encourage the growth of the desired cells, and discourage the growth of unwanted cells. It would also be interesting to make PECs from proteins (polyampholytes) such as collagen, fibronectin, gelatin, silk fibroins, soybean-based materials or a combination of those with CS and other polysaccharides. Finally, blending proteins such as collagen to the PEC materials without covalent crosslinking may be used to
improve the initial attachment of cells. It is envisaged that the future work would potentially be of great usability to treat targeted diseases such as skin ulcers (hard to heal chronic open wounds) and degenerative joint diseases (deteriorated articular cartilage). Clinically, the PEC materials may be used in one form or a combination of the forms such as PECs gels, film and scaffolds for wound healing applications (Fig. 8.2).

![Effect of pH on self-crosslinking](image)

**Figure 8.2 – Creating versatile, application tailored systems for wound healing applications. The medical device may be used as a form of gel, film, scaffold or in the combination of the mentioned forms.**

In the future, it may be worth investigating the biological properties of PEC scaffolds. There may be a different response between cell attachment on films and scaffolds due to the different surface and structural parameters.

In the search for effective, sustainable and economical biomaterials it is anticipated that these future studies will enable the properties and functions of CS-CRG PEC gels, films and scaffolds to be fully characterised for both non-biofouling and tissue engineering applications.
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