Interactions between *Pseudomonas aeruginosa* Bacteria and Corneal Fibroblasts in Human Microbial Keratitis

by

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ABSTRACT

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INTERACTIONS BETWEEN PSEUDOMONAS AERUGINOSA BACTERIA AND CORNEAL FIBROBLASTS IN HUMAN MICROBIAL KERATITIS

Ahmad Fathy Elsahn

Microbial keratitis (MK) is a leading cause of blindness in the developing world. *Pseudomonas aeruginosa* (PA) is the most common pathogen isolated from contact lens related MK, and is usually associated with significant visual complications. The intact corneal epithelium forms a strong barrier to bacterial penetration into deeper tissue, and only when bacteria reach the stroma, through a breach in the epithelium, does a typical ulcer develop, with stromal suppuration and necrosis. Since corneal fibroblasts are one of the most prominent cellular components in the stroma, they probably have an important role in the pathogenesis of corneal ulceration.

The aim of this study was to establish an *in vitro* model of PA microbial keratitis in order to examine the microbiological, biochemical and ultrastructural changes associated with live infection of primary human corneal fibroblasts (hCF) with PA, and to assess the role of these cells in the initiation and progression of the pathogenesis of the disease.

Human corneal fibroblast monolayers were infected with bacteria, and bacterial association was assessed at 3, 6 and 9h, bacterial invasion was assessed at 9h and cytokine profiles were assessed at 9 and 24 h post challenge. Host cell death was assessed at 9h using a lactate dehydrogenase assay (LDH). Gentamicin was added at 3h to eliminate all extra cellular bacteria and limit pathogen induced effects, and bacterial replication and survival and its effect on host cell viability were assessed. Induction of cytokine and matrix metalloproteinase (MMP) production by hCF in response to PA infection was assessed by a sandwich immunoassay and a gelatine zymography.

To examine the interactions of PA with the cornea as an organ, donor corneal buttons remaining after DSEK were incubated in a medium inoculated with PA for 3-9h, then processed for scanning and transmission electron microscopy. Scanning and Transmission Electron Micrographs (SEM/TEM)
of bacterial association and invasion of corneal cells were acquired and showed that bacteria associated to superficial epithelial cells but only invaded and colonized the stroma in the presence of a mechanical breach to the epithelial basement membrane.

PAO1 bacteria caused monolayer disruption within 24h with all infective doses ($10^1$-$10^8$ CFU/mL). PAO1 associated to hCF in a dose- and time-dependant manner utilizing type IV pilus and flagella. Bacterial internalization was detected by the gentamicin assay, which also demonstrated the ability of PAO1 to survive and replicate within the fibroblasts, and was dependant on the SRC tyrosine kinase and the actin microfilament system, where internalisation was diminished when these systems were inhibited. Cytotoxicity was observed by 9h post challenge, and was reliant on bacterial type III secretion system and flagella, where mutant bacterial strains induced less cytotoxicity. The more virulent PA14 bacteria induced more cytotoxicity than PAO1. The presence of intracellular bacteria did not affect cell viability in the presence of gentamicin.

In response to PAO1 infection, hCF produced the pro-inflammatory IL-1β and GM-CSF, although GM-CSF seems to be released later than IL-1β as shown by protecting the cells with gentamicin 3h post challenge. Infection with PA14 caused an earlier and more potent cytokine production than PAO1, reflecting the pattern of cytotoxicity. Bacterial components did not have a significant effect on cytokine production by corneal fibroblasts. Cytokine production was partly reliant on bacterial type III secretion system and flagella, and wild type PA14 strains induced an earlier and more potent IL-1β production by infected corneal fibroblasts.

Gelatine zymography showed that hCF produce specific MMPs in response to live PA challenge for 24h, and these MMPs were not released in response to treatment with individual bacterial components and virulence factors. Bacteria alone produced alkaline protease, a bacterial MMP. Sandwich immunoassays showed that MMP-1, MMP-2, MMP-3 and MMP-9 were produced by infected fibroblasts to a level similar to uninfected cells. Gentamicin protected cells produced more MMPs than unprotected cells.
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PRESENTED WORK, PUBLICATIONS AND AWARDS

Posters and Presentations


April 2014  Bacterial Infection of the Cornea. Oral Presentation. Wessex Regional Study Day, Southampton Eye Unit. Southampton, UK


May 2013  Early Events of *Pseudomonas aeruginosa* Microbial Keratitis: An Electron Microscopic Study. Oral Presentation. 21st VISTA Scientific Corneal Conference. Seattle, WA, USA


May 2012  Cytopathic and Immunologic Effects of *Pseudomonas aeruginosa* Infection of Human Corneal Fibroblasts. Oral Presentation. 20th VISTA Scientific Corneal Conference. Fort Lauderdale, FL, USA

May 2012  Interactions of *Pseudomonas aeruginosa* with Human Corneal Fibroblasts *in-vitro*. Poster. Association for Research in Vision and Ophthalmology (ARVO) Annual meeting. Fort Lauderdale, FL, USA

Publications


Awards

- National Institute for Health Research Biomedical Research Centre for Ophthalmology grant

Upcoming Activities

- A poster presentation entitled ‘The role of type 4 pilus, flagella and type III secretion system in the interactions between Pseudomonas aeruginosa bacteria and corneal fibroblasts in human microbial keratitis’ has been accepted for presentation as a poster in the upcoming Association for Research in Vision and Ophthalmology (ARVO) Annual meeting on 1-5 May 2016 in Seattle, WA, USA
- An author’s manuscript entitled ‘Interactions between Pseudomonas aeruginosa bacteria and corneal fibroblasts in human microbial keratitis’ is being prepared for submission to the Investigative Ophthalmology & Visual Science (IOVS) journal
DECLARATION OF AUTHORSHIP

I, Ahmad Elsahn, declare that the thesis entitled “Interactions between *Pseudomonas aeruginosa* bacteria and corneal fibroblasts in human microbial keratitis” and the work presented in it are my own and has been generated by me as the result of my own original research.

I confirm that:

1. This work was done wholly or mainly while in candidature for a research degree at this University;

2. Where any part of this thesis has previously been submitted for a degree or any other qualification at this University or any other institution, this has been clearly stated;

3. Where I have consulted the published work of others, this is always clearly attributed;

4. Where I have quoted from the work of others, the source is always given. With the exception of such quotations, this thesis is entirely my own work;

5. I have acknowledged all main sources of help;

6. Where the thesis is based on work done by myself jointly with others, I have made clear exactly what was done by others and what I have contributed myself;

7. None of this work has been published before submission

Signed: ........................................................................................................................................................................

Date: ........................................................................................................................................................................
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**DEFINITIONS AND ABBREVIATIONS**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>BCA</td>
<td>Bicinchoninic acid</td>
</tr>
<tr>
<td>CD</td>
<td>Cytochalasin D</td>
</tr>
<tr>
<td>CF</td>
<td>Corneal fibroblasts</td>
</tr>
<tr>
<td>CFCM</td>
<td>Corneal fibroblast culture medium</td>
</tr>
<tr>
<td>CFIM</td>
<td>Corneal fibroblast infection medium</td>
</tr>
<tr>
<td>CFU</td>
<td>Colony forming unit</td>
</tr>
<tr>
<td>dFCS</td>
<td>Decomplemented foetal calf serum</td>
</tr>
<tr>
<td>DSEK</td>
<td>Descemet's stripping endothelial keratoplasty</td>
</tr>
<tr>
<td>GFP</td>
<td>Green fluorescent protein</td>
</tr>
<tr>
<td>GM-CSF</td>
<td>Granulocyte monocyte colony stimulating factor</td>
</tr>
<tr>
<td>hCFs</td>
<td>Human corneal fibroblasts</td>
</tr>
<tr>
<td>IL-1β</td>
<td>Interleukin 1-beta</td>
</tr>
<tr>
<td>LDH</td>
<td>Lactate dehydrogenase</td>
</tr>
<tr>
<td>MK</td>
<td>Microbial keratitis</td>
</tr>
<tr>
<td>MMP</td>
<td>Matrix metalloproteinase</td>
</tr>
<tr>
<td>MOI</td>
<td>Multiplicity of infection</td>
</tr>
<tr>
<td>PA</td>
<td><em>Pseudomonas aeruginosa</em></td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate buffered saline</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase chain reaction</td>
</tr>
<tr>
<td>SDS-PAGE</td>
<td>Sodium dodecyl sulphate-polyacrylamide gel electrophoresis</td>
</tr>
<tr>
<td>SEM</td>
<td>Scanning electron microscopy</td>
</tr>
<tr>
<td>TEM</td>
<td>Transmission electron microscopy</td>
</tr>
<tr>
<td>TFP</td>
<td>Type IV pilus</td>
</tr>
<tr>
<td>TNF-α</td>
<td>Tumour necrosis factor-alpha</td>
</tr>
<tr>
<td>TTSS</td>
<td>Type III secretion system</td>
</tr>
<tr>
<td>WT</td>
<td>Wild type bacteria</td>
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Chapter 1: INTRODUCTION
1.1 The global impact of corneal blindness

The World Health Organization (WHO) Prevention of Blindness and Deafness Programme reports in its most recent systematic search and review to obtain a global estimate of visual impairment that the estimated number of people visually impaired in the world is 285 million, 39 million of which are blind.\(^1\) Corneal opacities accounted for 7-15% of the world’s blind population, making it the 3rd most common cause of blindness.\(^2,3\) In some regions, such as Saudi Arabia and India, the prevalence of corneal blindness can be as high as 25%,\(^4-7\) and in Malawi\(^8\) and Bangladesh,\(^9\) corneal scarring was accountable for 39–55% of unilateral blindness. Unlike other major causes of blindness, like cataract, glaucoma and macular degeneration, corneal blindness frequently affects all age groups, including people of working age, making it a major public health problem, with significant long term socio-economic implications.\(^10\)

Corneal opacification is the third most common cause of childhood blindness worldwide, after non-corneal causes such as congenital cataract and glaucoma,\(^10\) and is often due to a single episode of infection, such as ophthalmia neonatorum resulting from Neisseria gonorrhoea and Chlamydia trachomatis infections.\(^2\) Surveys of child blindness in Africa have shown that approximately 70% of all visual disability in this group is caused by corneal opacification.\(^11\) Patients suffering corneal blindness endure long term loss of vision, reduced quality of life and decreased productivity. In addition, there are significant costs to patients and carers, such as lost income and distress from pain and discomfort.\(^12\) Blind children have a lifetime of increased morbidity ahead of them, and even increased mortality, with up to 60% of blind children dying within 1 year of becoming blind.\(^10\)

The aetiology of corneal blindness is diverse and varies according to ocular diseases endemic to each geographic location. Important diseases classically associated with corneal blindness include trachoma, onchocerciasis, leprosy, ophthalmia neonatorum, and xerophthalmia.\(^13\) Trachoma remains the world’s leading infectious cause of blindness and ocular morbidity. It is estimated by WHO that, at present, there are about 4.9 million people already blind from trachomatous corneal scarring and an additional 10 million suffering from trichiasis which puts them at risk of corneal blindness.\(^14\) Onchocerciasis is another major cause of blindness in the world, although the incidence of onchocerciasis-related blindness has been declining since the introduction of the onchocerciasis control programme.\(^15\) Leprosy causes corneal scarring from interstitial keratitis, exposure keratitis due to lagophthalmos and loss of corneal sensation. Despite remarkable progress in the management of leprosy in the past few decades, corneal complications are still a significant cause of corneal blindness, affecting 10-12 million people, with approximately 250,000 blind from the disease,
mostly in Africa and the southern India. Xerophthalmia and Ophthalmia neonatorum are important causes of blindness in children.

Although these diseases remain a big public health problem, the introduction of control programs by the WHO has been successful in decreasing their incidence. This, in addition to worldwide decline in the number of cases of trachoma, has drawn attention to other causes of corneal blindness. These include trauma, microbial keratitis and the use of traditional eye medicines, which are often contaminated and can provide a route for the spread of pathogens.

There is an estimated 1.5-2 million people that develop corneal ulceration every year in developing countries. In a recent study by Wang et al., infectious keratitis accounted for as much as 40% of childhood and 20% of adult corneal blindness in their study population in China, whereas Rapoza et al. and Bowman et al. reported corneal infections, including trachoma, were the leading cause of unilateral and bilateral corneal blindness in central Tanzania and Gambia, respectively. In the Nepal blindness survey, corneal trauma and ulceration were the second most common cause of unilateral visual loss after cataract, accounting for 7.9% of all blind eyes. All this clearly shows that corneal infection is a significant public health problem, yet it is alarmingly under-diagnosed and under-treated, that some consider it a ‘silent epidemic’ in the developing world.

Infectious keratitis can be caused by bacteria, viruses, fungi, protozoa, and parasites (Table 1.1). On a global level, common risk factors for the disease include ocular trauma, contact lens wear, recent ocular surgery, pre-existing ocular surface disease, dry eyes, lid deformity, corneal sensation impairment, chronic use of topical steroids, and systemic immunosuppression. These risk factors and causative organisms, however, vary tremendously with geographical location; for instance, non-surgical trauma to the eye accounted for 48.6–65.4% of all corneal ulcers in Nepal and India, but only accounted for 27% of all cases at a large county trauma referral centre in the United States.

Contact lens wear accounts for up to 50.3% of cases of microbial keratitis in the western world, and the estimated annual incidence of microbial keratitis per 10,000 contact lens wearers is up to 25.4 (95% CI, 14.6–29.5) for extended wear of silicone hydrogel lenses. Pseudomonas aeruginosa is the most commonly isolated causative organism for contact lens associated microbial keratitis, accounting for up to 68.8% of isolates. Pseudomonas aeruginosa microbial keratitis is not only one of the most significant types of the disease, but also one of the most devastating, causing significant morbidity and disability.
### Table 1.1 Causative organisms of microbial keratitis.

<table>
<thead>
<tr>
<th>BACTERIAL</th>
<th>Gram positive</th>
<th>Gram negative</th>
</tr>
</thead>
<tbody>
<tr>
<td>Staphylococcus spp (aureus, epidermidis)</td>
<td>Pseudomonas aeruginosa</td>
<td></td>
</tr>
<tr>
<td>Streptococcus pneumoniae</td>
<td>Moraxella spp</td>
<td></td>
</tr>
<tr>
<td>Streptococcus pyogenes</td>
<td>Klebsiella pneumoniae</td>
<td></td>
</tr>
<tr>
<td>Streptococcus viridans</td>
<td>Enterobacter aerogenes</td>
<td></td>
</tr>
<tr>
<td>Corynbacterium diphtheroides</td>
<td>Serratia spp (marcescens, liquefaciens)</td>
<td></td>
</tr>
<tr>
<td>Nocardia spp</td>
<td>Acinetobacter spp</td>
<td></td>
</tr>
<tr>
<td>Propionibacteria acnes</td>
<td>Enterococcus spp</td>
<td></td>
</tr>
<tr>
<td>Bacillus spp</td>
<td>Burkholderia cepacia</td>
<td></td>
</tr>
<tr>
<td>Mycobacterium spp</td>
<td>Escherichia coli</td>
<td></td>
</tr>
<tr>
<td>Micrococcus spp</td>
<td>Stenotrophomonas maltophilia</td>
<td></td>
</tr>
<tr>
<td>Actinomyces spp</td>
<td>Neisseria gonorrhoeae</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Haemophilus spp</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kingella kingae</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pseudomonas spp (non-aeruginosa)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Citrobacter spp</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Aeromonas spp</td>
<td></td>
</tr>
</tbody>
</table>

| CHLAMYDIAL                     | C trachomatis, psittaci                |                                                    |

| FUNGAL                         |                                        |                                                    |
| Fusarium spp                   |                                        |                                                    |
| Aspergillus spp (flavus, fumigatus, niger, terreus and nidulans) | |                                                    |
| Alternaria spp                 |                                        |                                                    |
| Cuvularia spp                  |                                        |                                                    |
| Lasiodiplodia theobromae       |                                        |                                                    |
| Paecilomyces spp               |                                        |                                                    |
| Penicillium spp                |                                        |                                                    |
| Scedosporium apiospermum       |                                        |                                                    |
| Cephalophora irregularis       |                                        |                                                    |
| Cladosporium cladosporoides    |                                        |                                                    |
| Cylindrocarpon spp             |                                        |                                                    |
| Exserohilum rostratum          |                                        |                                                    |
| Bipolaris spp                  |                                        |                                                    |
| Candida spp                    |                                        |                                                    |
| Pythium insidiosum             |                                        |                                                    |

| VIRAL                           |                                        |                                                    |
| Herpes simplex virus type 1    |                                        |                                                    |
| Varicella zoster virus          |                                        |                                                    |
| Human adenovirus (serotypes 8, 19, and 37) | |                                                    |
| Enterovirus type 70            |                                        |                                                    |
| Coxsackievirus type 24         |                                        |                                                    |
| Echovirus type 13              |                                        |                                                    |
| Poliovirus type 3              |                                        |                                                    |
| Cytomegalovirus                |                                        |                                                    |

| PARASITIC                      |                                        |                                                    |
| Acanthamoeba spp (castellani, polyphaga and culbertsoni) | |                                                    |
| Microsporidia spp              |                                        |                                                    |
| Hartmannella spp               |                                        |                                                    |
| Vahlkampfia spp                |                                        |                                                    |
| Dictyostelium polycephalum     |                                        |                                                    |
| Phthiriasis palpebrarum        |                                        |                                                    |
| Oestrus ovis                   |                                        |                                                    |

| NEMATODAL                      |                                        |                                                    |
| Loa loa                       |                                        |                                                    |
| Onchocerca volvulus            |                                        |                                                    |
1.2 The Host: Structure and function of the normal cornea

1.2.1 Macroscopic anatomy of cornea

The cornea is a transparent avascular tissue that is exposed to the external environment. The anterior corneal surface is covered by the tear film, and the posterior surface is in direct contact with the aqueous humor. The adult human cornea measures 11 to 12 mm horizontally and 9 to 11 mm vertically. It is approximately 0.5 mm thick at the centre, and its thickness increases gradually toward the periphery, where it is about 0.7 mm thick. The shape of the anterior corneal surface is convex, although the curvature along the corneal surface is not constant, and aspheric, where it is steeper and almost spherical at the centre, with a radius of curvature between 7.5 and 8.0 mm, and flatter towards the periphery. The refractive power of the cornea is 40 to 44 dioptres, which constitutes about two-thirds of the total refractive power of the eye.

The cornea is one of the few avascular tissues in the body. It is also one of the most heavily innervated tissues in the body, and the density of nerve endings in the cornea is about 300 to 400 times greater than that in the skin. Most of the sensory nerves in the cornea are derived from the long ciliary nerves of the ophthalmic branch of the trigeminal nerve, entering the cornea mostly as a continuation the nerves in the suprachoroidal space. Approximately 43 nerve bundles enter the cornea, predominantly in the mid and deep stroma, evenly distributed through the limbal circumference. Corneal nerves penetrate Bowman’s layer perpendicularly and vertically (i.e. from inferior to superior) in the mid-peripheral cornea, to form a sub basal nerve plexus. The cornea contains peptidergic, sympathetic, and parasympathetic nerve fibres, with various neurotransmitters, including substance P, calcitonin gene-related peptide, neuropeptide Y, vasoactive intestinal peptide, galanin, methionine-encephalin, catecholamines, and acetylcholine.

Corneal epithelial and endothelial cells are metabolically active and require adenosine triphosphate (ATP) as an energy source, which is obtained by glycolysis under aerobic conditions. A supply of glucose and oxygen is thus essential to maintain the normal metabolic functions of the cornea. Glucose is obtained by diffusion from the aqueous humor, whereas oxygen is obtained primarily by diffusion from the tear fluid, which absorbs oxygen from air, and to a lesser extent by diffusion from the aqueous humor and limbal circulation. Disruption of oxygen supply to the cornea, such as that associated with contact lens wear, can lead to corneal hypoxia and consequent stromal oedema. Closure of the eyelids during sleep also reduces the amount of oxygen that reaches the cornea, which can change corneal metabolism from aerobic to anaerobic, with consequent accumulation of lactate.
The corneal surface is covered by the tear fluid, which protects the cornea from dehydration and helps to maintain the epithelial surface smooth. It contains almost all of the biochemical components of blood, including albumin and globulins, but few of the cellular components. The thickness and volume of the tear film are about 7 µm and 6.5 ± 0.3 µl, respectively. The tear film consists of three layers: a superficial lipid layer (~0.1 µm), an aqueous layer (~7 µm), and a mucin layer (0.02 to 0.05 µm). The constituents of the superficial lipid layer of the tear film are secreted by Meibomian glands and other secretory glands of the eyelid, the aqueous layer is secreted by the main and accessory lacrimal glands, and the mucin layer is produced by goblet cells in the conjunctival epithelium and corneal epithelial cells. More than 98% of the total volume of the tear film is water, but it also contains many biologically important molecules, including electrolytes, glucose, immunoglobulins, lactoferrin, lysozyme, albumin, and oxygen, in addition to several biologically active substances such as histamine, prostaglandins, growth factors, and cytokines. Thus, the tear film serves not only as a lubricant and source of nutrients for the corneal epithelium but also as a source of regulatory factors required for epithelial maintenance and repair and modulation of corneal epithelial migration, proliferation, and differentiation. In pathological conditions such as allergic keratoconjunctivitis, corneal trauma, or corneal infection, tear fluid provides a path for various inflammatory cells, growth factors, and inflammatory cytokines derived from the palpebral or bulbar conjunctiva to reach the cornea. The tear fluid of individuals with corneal ulceration also contains matrix metalloproteinases (MMPs) that degrade extracellular matrix proteins such as collagen and basement membrane components.
1.2.2 Microscopic anatomy of the cornea

The structure of the cornea is relatively simple, consisting of five layers: the epithelium, Bowman’s layer, the stroma, Descemet’s membrane, and the endothelium (Figure 1.1 A). The epithelium, stroma and endothelium are cellular layers, whereas Bowman’s layer and Descemet’s membrane are acellular and act as interfaces. Epithelial cells are embryologically derived from the surface ectoderm, whereas stromal keratocytes (corneal fibroblasts) and endothelial cells are of neural crest (neuroectodermal) origin. Components of the cornea interact with each other to maintain its integrity and function.\[51\]

The optical properties of the cornea are determined by its transparency, surface smoothness, contour, and refractive index. Corneal transparency is mostly attributable to the arrangement of collagen fibres in the stroma.\[51\] Both the mean diameter of each collagen fibre and the mean distance between collagen fibres are relatively homogeneous and are less than half of the wavelength of visible light (400 to 700 nm). This anatomic arrangement is thought to be responsible for the fact that scattering of an incident ray of light by each collagen fibre is cancelled by interference from rays scattered by neighbouring collagen fibres,\[82\] allowing light to pass through the cornea with minimal scatter. If the diameter of, or the distance between, collagen fibres become disorganized (as occurs in fibrosis or oedema), incident rays are scattered randomly and the corneal transparency is significantly affected.

1.2.2.1 The epithelium

The corneal epithelium is composed of non-keratinized stratified squamous epithelial cells, spanning a thickness, which is constant over the entire corneal surface, of approximately 50 µm, about 10% of the total thickness of the cornea. There are 5-6 layers of three different types of epithelial cells (Figure 1.1 B), and although the corneal epithelium is normally non-keratinized, it may develop keratinization under certain pathological conditions, such as vitamin A deficiency.\[83\]

The surface of the corneal epithelium consists of 2-3 layers of terminally differentiated superficial cells. These cells are flat and polygonal with a diameter of 40-60 µm and a thickness of 2-6 µm. Their surface is covered with microvilli, which form microplicae\[84\] that increase the surface area of each cell and thereby promote the absorption of oxygen and nutrients from tear fluid.

Superficial cells are well differentiated, and so they do not proliferate, exhibit a low metabolic activity and contain fewer organelles and less RNA than do the other types of corneal epithelial cells. Numerous glycoprotein and glycolipid molecules are embedded in the cell membrane of epithelial cells and are collectively known as the glycocalyx. These molecules interact with the mucinous layer
of the tear film and help to maintain its structure. Two different types of superficial cells are present in the corneal epithelium: large, dark cells and small, light cells. The former are mature, have a dense coat of microvilli, and are about to undergo desquamation into the tear film; the latter have fewer microvilli and are thought to be younger.

Although superficial cell desquamation occurs in normal physiologic conditions, apoptosis and increased desquamation can occur under conditions of stress, such as ultraviolet radiation, hypoxia, infection or mechanical injury. Apoptosis contributes to stromal keratocyte cell turnover under normal conditions, but it is not clear whether it is also involved in normal epithelial cell turnover.

Beneath the superficial cells of the corneal epithelium lie 2-3 layers of wing cells, so called because of their characteristic wing-like shape. Wing cells are in an intermediate state of differentiation between basal and superficial cells and are rich in intracellular keratin tonofilaments, but have sparse cytoplasmic organelles. The cell membranes of adjacent wing cells interdigitate and have numerous desmosomes and gap junctions between them (see Figure 1.2).

Basal epithelial form a single layer resting on the basement membrane. Neighbouring cells are joined laterally by desmosomes, gap junctions, and adherens junctions (zonula adherens). They are the only epithelial cells capable of mitosis, giving rise to wing cells, which in turn differentiate into superficial flat cells. The latter desquamate into the tear film approximately 7-14 days after differentiation. Due to their mitotic activity, basal cells demonstrate numerous intracellular organelle, free ribosomes, rough endoplasmic reticulum, mitochondria, centrioles, microfilaments, microtubules, and glycogen granules, in contrast to wing cells or superficial cells, which have less organelles. The posterior surfaces of the basal cells lie flat over the basement membrane, and are adherent to it with hemidesmosomes and anchoring fibrils. The later consist of type VII collagen and penetrate the basement membrane into the stroma, where they, in addition to stromal type I collagen, form anchoring plaques. This arrangement is critical in maintaining the stability of the entire epithelium by ensuring the adhesion of the epithelial cells to the basement membrane and underlying stroma, and its disruption leads to recurrent epithelial erosions.
Figure 1.1 Microscopic anatomy of the cornea

A, Histology of the cornea showing the epithelium (Ep), Bowman’s layer (BL), the stroma (S), Descemet’s membrane (DM), and the endothelium (En). B, Scanning electron microscopy of the corneal epithelium, comprising a basal cell layer (B), 2-3 layers of wing cells (W), and 2-3 layers of superficial cells (S). (Reproduced with permission from Krachmer et al, Cornea. Mosby. 2005)
The basal epithelial cells synthesize and secrete type IV collagen and laminin, which are major components that form the basement membrane, along with fibronectin, heparan sulphate, type VII collagen, proteoglycans and fibrin.\textsuperscript{94} When examined by transmission electron microscopy, the basement membrane is 40-60 nm thick and is composed of a pale layer (the lamina lucida) immediately posterior to the cell membrane of the epithelial basal cells and an electron-dense layer (the lamina densa) adjacent to the stroma.

The presence of the basement membrane between the basal epithelium and the underlying stroma fixes the polarity of epithelial cells, which is important in the virulence of microbes and the pathogenesis of microbial keratitis.\textsuperscript{95-97} It provides a matrix on which cells can migrate and is important for maintenance of the stratified and well-organized structure of the corneal epithelium. It also appears to play a prominent role in epithelial wound healing, particularly fibronectin. Fibronectin is thought to provide a temporary matrix over which cells can migrate during the transition period between epithelial debridement and the formation of a new basement membrane, which can take more than a week, and although type IV collagen and laminin are essential for maintenance of the steady state of the corneal epithelium, fibronectin appears to play a more important role under conditions of acute injury to it.\textsuperscript{98}

After damage to the corneal epithelium, actively migrating epithelial cells no longer manifest gap junctions or desmosomes in the wounded region lacking a basement membrane. Reestablishment of the continuity of the corneal epithelium is accompanied by the synthesis and deposition of basement membrane proteins and by the reassembly of the various types of junctional apparatus, suggesting that the presence of the basement membrane may be required for reformation of cell-cell junctions in the corneal epithelium.\textsuperscript{99}

The epithelium functions as barrier to various external biological and chemical insults. This is achieved by a variety of junctional complexes between epithelial cells and the interdigitation of adjacent cell membranes, which, together, form an effective barrier against the influx of fluid and chemical molecules into the underlying stroma, thereby preventing corneal oedema and blurred vision. Superficial cells are bound together by tight junctions, which form the main component of the superficial epithelial barrier, whereas basal cells and wing cells have gap junctions that allow the passage of nutrients and different molecules between cells. Adherens junctions and desmosomes are present in all layers of the epithelium and help to maintain the structural integrity of the epithelium.\textsuperscript{51}
The epithelium also acts as a barrier to microbial penetration into deeper layers of the cornea, mainly through the enzymes and cytokines of the tear film, the desquamation of surface epithelial cells, the tight junctions between adjacent epithelial cells, and the epithelial basement membrane, with its pores that are too small to allow bacterial penetration into the stroma. Rapid renewal and the formation of intercellular junctions between corneal epithelial cells help to protect the cornea from microbial challenge. Both cell-cell and cell-matrix interactions are important for maintenance of the normal stratified structure and physiological functions of the corneal epithelium. The characteristics of the different types of intercellular junctional complexes present between the corneal epithelium are summarized in Table 1.2 and in Figure 1.2.

Cellular components of the epithelium also play an important role in corneal immunology. Dendritic Langerhans cells, bone marrow derived specialized macrophages involved in antigen processing, are abundant at the periphery of the corneal epithelium but are not present in the central region of the normal cornea. These cells express human lymphocyte antigen (HLA) class II molecules and are thought to function in the afferent arm of the ocular immune response by presenting antigens to T cells. In addition, their numbers are increased in conditions characterized by ocular inflammation and are reduced by treatment with corticosteroids. Injury to the central cornea results in the rapid migration of peripheral Langerhans cells to the damaged area.

The tear film covered epithelium largely contributes to maintaining the smooth optical surface necessary for the sharp focusing of light onto the retina. Loss of the smooth contour of the epithelium, like in epithelial defects or irregularities or in dry eyes, leads to scattering of light, degradation of image quality and consequent blurring of vision.
Figure 1.2 Types of intercellular junctional complexes in the corneal epithelium


### Table 1.2 Characteristics of corneal epithelial cells

<table>
<thead>
<tr>
<th></th>
<th>Shape</th>
<th>Layers</th>
<th>Size</th>
<th>Junctional Complexes</th>
<th>Cytoplasmic Organelles</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Superficial Cells</strong></td>
<td>Flat</td>
<td>2-4</td>
<td>Diameter: 40-60 µm</td>
<td>Desmosomes</td>
<td>Sparse</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Thickness: 2-6 µm</td>
<td>Tight Junctions (zonula occludens)</td>
<td></td>
</tr>
<tr>
<td><strong>Wing Cells</strong></td>
<td>Wing-like</td>
<td>2-3</td>
<td>Variable</td>
<td>Desmosomes</td>
<td>Sparse</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Gap junctions</td>
<td></td>
</tr>
<tr>
<td><strong>Basal Cells</strong></td>
<td>Columnar</td>
<td>1</td>
<td>Diameter: 8-10 µm</td>
<td>Desmosomes</td>
<td>Prominent</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Thickness: 18-20 µm</td>
<td>Gap junctions</td>
<td>mitochondria and Golgi</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Hemidesmosomes (zonula adherens)</td>
<td>apparatus</td>
</tr>
</tbody>
</table>

(Reproduced with permission from Krachmer et al, Cornea. Mosby. 2005)
1.2.2.2 Bowman’s layer

Lying directly posterior to the epithelial basement membrane is Bowman’s layer, an acellular layer of relatively randomly arranged collagen fibres and proteoglycans that is 12 µm thick. The collagen fibres of this layer are primarily collagen types I and III and measure 20-30 nm in diameter, smaller than stromal collagen fibres. It is considered to be the anterior portion of the corneal stroma, and its collagen fibres appear continuous with stromal collagen fibres and are synthesized by stromal fibroblasts.

The physiological role of Bowman’s layer is unclear. Many of the functions originally attributed to it are now known to be mediated by the epithelial basement membrane. Bowman’s layer does not regenerate after injury or Excimer laser surgery, and yet a normal epithelium is formed and maintained even in the absence of Bowman's layer. Furthermore, many mammals do not have a Bowman’s layer but still exhibit a well-organized epithelial structure.

1.2.2.3 The stroma

The corneal stroma constitutes more than 90% of the thickness of the cornea (Figure 1.1), and many of the properties of the cornea, such as transparency, shape stability and tensile strength are attributable to the anatomic and biochemical properties of the stroma. One of the most important functions of the stroma is maintenance of corneal transparency. This is achieved by the regular arrangement, uniform diameter and continuous slow production and degradation of its collagen fibres. In addition, the epithelial and endothelial cells play an important role in maintaining the biological activities of keratocytes and the arrangement of collagen fibres through their regulation of the water content of the stroma.

The corneal stroma consists of extracellular matrix, stromal cells and nerve fibres. The cellular components only comprise 2-3% of the total volume of the stroma, with the remaining portion mostly comprised of extracellular matrix containing collagen and glycosaminoglycans. Collagen constitutes more than 70% of the dry weight of the cornea and is mostly type I, in addition to smaller amounts of types III, V and VI.

Keratocytes are the predominant cellular components of the corneal stroma and are thought to turn over every 2-3 years. The spindle-shaped keratocytes are scattered between the lamellae of the stroma and extend long processes that are connected to neighbouring cells at their tips by gap junctions (Figure 1.3). The detection of fluorescent dye transfer between neighbouring keratocytes suggests that the gap junctions in these cells are functional and serve to transmit intercellular signals. Damage to the corneal stroma results in a loss of the network connections of
keratocytes near the site of injury; this leads to activation of keratocytes, which play an essential role in corneal stromal wound healing as well as in the pathogenesis of corneal ulceration.\textsuperscript{[119]}

Keratocytes possess an extensive intracellular cytoskeleton, including prominent actin filaments, which allows them to contract, and may be responsible for the maintenance of corneal shape and the packed structure of collagen in the stroma. The shape and function of keratocytes are regulated by the extracellular environment. Keratocytes cultured in a collagen gel become less active mitotically, appear similar to keratocytes in situ and provide a model system for studying keratocyte biology.\textsuperscript{[120, 121]} Keratocytes in the normal cornea are inactive but are rapidly activated to fibroblast and myofibroblasts phenotypes by various types of insult to the stroma. Activated fibroblasts at the site of a persistent epithelial defect may contribute to stromal dissolution by increasing their synthesis and secretion of collagen-degrading enzymes, like matrix metalloproteinases (MMPs), and by exhibiting phagocytic activity for foreign material.\textsuperscript{[119]}

The collagen fibres in the stroma are highly uniform in diameter (22.5 to 35 nm), and the distances between collagen fibres are also highly uniform (41.4 ± 0.5 nm).\textsuperscript{[122, 123]} This regular arrangement is of paramount importance in determining corneal transparency, and any disturbance in the uniformity of the inter-fibre distance, like what occurs during stromal oedema or scarring, can result in a loss of corneal transparency. In the corneal stroma, the collagen fibres form about 300 lamellae, each lamella courses parallel to the surface of the cornea from one limbus to the other, and then run along the limbal circumference. The turnover of collagen molecules in the cornea is slow, requiring 2-3 years.\textsuperscript{[124]}

Various glycosaminoglycans are present between collagen fibres in the corneal stroma and they have the ability to absorb and retain large amounts of water. All of these glycosaminoglycans bind to core proteins to form proteoglycans, with the exception of hyaluronan (formerly known as hyaluronic acid). Keratan sulphate is the main glycosaminoglycan in the cornea, constituting about 65% of the total glycosaminoglycan content. Other glycosaminoglycans include chondroitin sulphate and dermatan sulphate.\textsuperscript{[125]} The major glycosaminoglycan in the embryonic eye is hyaluronan, which is present during morphogenesis of the corneal stroma but its amount is reduced after the stroma has formed. The reduction of hyaluronan as the structure of the cornea is established during embryonic development is accompanied by an increase in the amounts of keratan sulphate, chondroitin sulphate, and dermatan sulphate, giving rise to the glycosaminoglycan composition of the adult stroma.\textsuperscript{[126]}
Figure 1.3 Transmission electron microscopy of the corneal stroma

1.2.2.4 Descemet's membrane

Descemet's membrane comprises the basement membrane of the corneal endothelial cells, and measures about 8-10 µm thick. It is mostly composed of type IV collagen and laminin\(^\text{[127]}\) but also contains fibronectin.\(^\text{[128,129]}\) Although its collagen fibres are not continuous with those of the stroma, unlike Bowman's layer, Descemet's membrane adheres tightly to the posterior surface of the corneal stroma and reflects any change in the shape of the stroma, and if the corneal stroma swells, the folding of Descemet's membrane can be observed clinically.

Descemet's membrane is tough and resistant to enzymatic degradation by MMPs, however, it can be torn easily on exposure to shearing stress. In cases of corneal ulceration, Descemet's membrane remains intact but protrudes outwards as a descemetocele as a result of the intraocular pressure and dissolution of the overlying stroma.\(^\text{[51]}\)

1.2.2.5 The endothelium

The endothelial cells form a single layer posterior to Descemet's membrane in a characteristic well-arranged mosaic pattern. These cells have a uniform polygonal (mostly hexagonal) shape and a uniform size, with 5 µm in thickness and 20 µm in diameter.\(^\text{[130]}\) An increase in the variability of cell area is termed polymegathism, whereas deviation from hexagonality of shape is referred to as pleomorphism. Both polymegathism and pleomorphism are important markers of endothelial damage, where normal endothelial cells fill the gaps left behind by lost cells. Although corneal endothelial cells proliferate in cell culture conditions, they do not proliferate in vivo in humans, suggesting that despite their ability to undergo mitosis, there may be factors in aqueous humor or other components in their surrounding environment may inhibit their proliferation.\(^\text{[131,132]}\)

Endothelial cell density in the normal, healthy cornea decreases with age.\(^\text{[133]}\)

Endothelial cells contain a large nucleus and abundant cytoplasmic organelles, including mitochondria, endoplasmic reticulum, free ribosomes, and Golgi apparatus, indicating that the cells are metabolically active and have secretory functions.\(^\text{[130]}\) The anterior surface of endothelial cells is flat and directly adjacent to Descemet's membrane, whereas the posterior, free surface of the cells forms microvilli and marginal folds that protrude into the anterior chamber, thereby maximizing the surface area exposed to aqueous humor.\(^\text{[134]}\) The endothelial cells interdigitate\(^\text{[135]}\) and contain various junctional complexes, including zonula occludens, macula occludens, and macula adherens. They do not possess desmosomes, but do have gap junctions which allow the transfer of small molecules and electrolytes between the endothelial cells; this arrangement provides a leaky barrier to aqueous humour.\(^\text{[136]}\) Loss or damage to corneal endothelial cells results in corneal oedema.
1.2.3 Maintenance of normal corneal integrity

The normal anatomical and biochemical structure of the extracellular matrix (ECM) of the corneal stroma is a prerequisite for a normally functioning cornea. Keratocytes maintain the stromal ECM by continually synthesising new collagen and glycosaminoglycans and degrading older, damaged components using ECM degrading enzymes, like MMPs, in a highly regulated process that is essential for the maintenance of corneal homeostasis.\cite{137} On the other hand, although keratocytes grow relatively quickly in cell cultures, they seem to have a lower mitotic activity in vivo, indicating that their growth may be regulated by the surrounding collagen and ECM components.\cite{121}

Collagen degradation by corneal keratocytes is regulated by various growth factors and cytokines. Under certain conditions of insult, such as injury, infections, and immunologic reactions, resident stromal keratocytes lose their quiescence and attain an activated phenotype, enter into the cell cycle and their cell size and organelle content increase. They begin to exhibit a fusiform shape, multiple nucleoli and a lack of cytoplasmic granules, and subsequently migrate to the site of injury.\cite{137} A number of growth factors and cytokines contribute to the activation of keratocytes, including, for example, IL-1α, which is expressed by the corneal epithelium and released into the stroma upon injury, and has been identified as an important mediator for the expression of the collagenase gene in activated fibroblasts.\cite{119}
1.3 The Pathogen: Biology of *Pseudomonas aeruginosa*

1.3.1 Background

*Pseudomonas aeruginosa* is a gram-negative, asporogenic, monoflagellated, rod-shaped bacterium that measures about 1-5 µm long and 0.5-1.0 µm wide. It belongs to phylum Proteobacteria, class Gamma proteobacteria, order Pseudomonadales, family pseudomonadaceae, genus pseudomonas, and species *Pseudomonas aeruginosa*. The word *Pseudomonas* means "false unit", from the Greek *pseudo* (Greek: false) and *monas* (Greek: a single unit). The stem word *mon* was used early in the history of microbiology to refer to germs, e.g., Kingdom Monera. The species name *aeruginosa* is a Latin word meaning "copper rust", which describes the blue-green bacterial pigment seen in laboratory cultures of the species.[138]

*P. aeruginosa* was first described as a distinct bacterial species at the end of the nineteenth century, after the development of sterile culture media by Pasteur. In 1882, a pharmacist named Carle Gessard published the first scientific study on *P. aeruginosa*, entitled “On the blue and green coloration of bandages”. This study showed *P. aeruginosa*’s characteristic water-soluble pigmentation which, on exposure to ultraviolet light, fluoresced blue-green light.[139] This was later attributed to the siderophores pyoverdin and pyocyanin (from "pyocyaneus" referring to "blue pus" characteristic of suppurative infections caused by *P. aeruginosa*), which also reflected the organism’s old names; *Bacillus pyocyaneus*, *Bakterium aeruginosa*, *Pseudomonas polycolor*, and *Pseudomonas pyocyaneus*.

*P. aeruginosa* is a very ubiquitous organism,[140] and it catabolizes a wide range of organic molecules, including organic compounds like benzoate,[141] among many other organic compounds for growth. On the other hand, organic growth factors are not essential for its growth, and it is even occasionally observed growing in distilled water,[142] which clearly indicates its minimal nutritional requirements. In the laboratory, the simplest medium for growth of *Pseudomonas aeruginosa* consists of acetate as a source of carbon and ammonium sulphate as a source of nitrogen. Its optimum temperature for growth is 37°C, but it is able to grow at temperatures as high as 42°C.[143] Its tolerance to a wide variety of physical conditions, including temperature, contributes to its ecological success as an opportunistic pathogen. It has a natural predilection to exist in soil and water, which is demonstrated in its predilection to grow in moist environments.[144]

Although the organism primarily utilizes aerobic respiration for energy production, it is also a facultative anaerobe, and, in the absence of oxygen, can utilize nitrate and other terminal electron acceptors for respiration.[145] Even in the absence of those, it is still able to ferment arginine.
by substrate-level phosphorylation.\cite{146} As a result, _P. aeruginosa_ has been found in a wide range of different environments such as soil, water, humans, animals, plants, sewage, and hospitals. It is also capable of growth in diesel and jet fuel, where it is known as a hydrocarbon-using microorganism, causing microbial corrosion.\cite{147,148}

The typical _Pseudomonas_ bacterium in nature might be found in a biofilm, attached to some surface or substrate, or in a planktonic form, as a single cell actively motile by means of polar flagella. It is commonly found in soil and water, and is a very important soil bacterium that is capable of breaking down polycyclic aromatic hydrocarbons and making rhamnolipids, quinolones, hydrogen cyanide, phenazines, and lectins.\cite{149} It is a commensal of the human gut, and is the predominant inhabitant of oligotrophic aquatic ecosystems, containing high-dissolved oxygen content but low plant nutrients throughout, making it the most abundant organism on earth.\cite{140}

There are several strains of _Pseudomonas aeruginosa_; these include strain PAO1, strain PA7, strain PA14, and strain 2192. Most of these were isolated based on their distinctive grapelike odour of aminoacetophenone, pyocyanin production, and the colonies’ structure on agar media.\cite{150}

### 1.3.2 Culture Morphology and Cell structure

_P. aeruginosa_ isolates may produce several colony types (Figure 1.6 A and B). Natural isolates from soil or water typically produce a small, rough colony. Clinical samples, in general, demonstrate two smooth colony types. One type has a fried-egg appearance, which is large, smooth, with flat edges and an elevated appearance. Another type, frequently obtained from respiratory and urinary tract secretions, has a mucoid appearance, possibly due to the production of alginate slime. The smooth and mucoid colonies are presumed to play a role in colonization and virulence.\cite{151}

#### 1.3.2.1. Cell surface and outer membrane:

_P. aeruginosa_ is a Gram-negative microbe, with an outer membrane and a cell wall. Its outer membrane contains Protein F (OprF), which functions both as a structural protein, maintaining the bacterial cell shape, and as a porin, allowing certain molecules and ions to come into the cells. Protein F provides _P. aeruginosa_ outer membrane with an exclusion limit of 500 Da, thus lowering the permeability of the outer membrane, which serves the organism well since this would decrease the intake of harmful substances into the cell and give _P. aeruginosa_ a high resistance to antibiotics.\cite{152}
Lipopolysaccharides belong to a group of polysaccharides that are present on the bacterial cell surface and serve as a barrier between the cell wall and the surrounding environment. They form structural components of biofilms, play an important role in maintaining outer membrane structural integrity and are important mediators for host-pathogen interactions and bacterial virulence. In *P. aeruginosa* infection during late stage cystic fibrosis, the exopolysaccharide alginate is responsible for the mucoid phenotype of the isolated offending strains.\(^{153, 154}\)

1.3.2.1 Flagellum and Pili:

*P. aeruginosa* utilize pili and polar flagella for motility and for association to target cells. The flagellum is a sheathless long appendage extending from one pole of the bacterium that consists of various combinations of at least 10 heat labile antigens. These antigens, together with the heat stable somatic O-antigen, form the basis of serological classification of the organism.\(^{155}\) In addition to using it for motility, *P. aeruginosa* uses its single polar flagellum to also display chemotaxis to sugars and other useful molecules in the surrounding environment. According to the size and the antigenicity of the flagellin subunit, *P. aeruginosa* strains are classified to have either A-type or B-type flagella. The flagellum is very important during the early stages of infection, as it can attach to and invade host tissues.\(^{156}\)

The bacterial flagellum can be subdivided into three components (Figure 1.4). The first is the basal body, which anchors the flagellum to the cell membrane, provides the power for rotation, and secretes the other two components. The second component is the hook, which serves as a flexible universal joint changing the angle of flagellar rotation. The third is the helical filament, which is composed primarily of the protein flagellin. The junction between the hook and the basal body has the core rotor structure, or the switch complex, which consists of an axial rod, the FliF ring, and the C ring, which is composed of the conserved multi-domain proteins FliG, FliM, and FliN.\(^{157}\) FliG lies close to the membrane and is involved in torque generation. FliM lies in the middle of the rotor and interacts with phosphorylated CheY (a signalling protein). FliN is located at the cytoplasmic end of the rotor and is essential for flagellar assembly and switching.\(^{158}\) Bacterial locomotion is achieved by rotating the flagella, and forward propulsion is achieved by counter clockwise (CCW) rotation, whereas a change in direction is achieved by switching to clockwise (CW) rotation. It appears that the binding of CheY-P to FliM (and possibly FliN) promotes a conformational change within the rotor to change its rotation sense from CCW to CW.\(^{158}\)
Bacteria use flagella for locomotion and, equally, for association to inert surfaces or target cells. This would make one wonder; how do bacteria know when to ‘swim-or-stick’? How does an individual bacterium recognize the proximity to a target surface and switch from swimming freely in its environment to sticking to a surface? How to switch from a motile to a sessile lifestyle? This can be explained by surface sensing, a sensory transduction mechanism that determines the phase of the switch and often involves the bacterial flagellum.\textsuperscript{[159]} As such, flagella are not only important in propulsion and locomotion of bacteria, but also have a critical mechano-sensory role in the initial stages of bacterial association to target surfaces.\textsuperscript{[157]} The swim-or-stick switch involves the inhibition of flagellar synthesis and rotation coupled with increased synthesis of the polymers and structures that are required for long-term attachment to a surface, such as pili.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{flagellum_4.png}
\caption{Structure of the flagellum}
\end{figure}

Shematic representation of the main components of the flagellum. The motor is composed of the stator (MotA and MotB proteins) and the rotor (C ring, composed of FliG, FliM, and FliN). Ion (H\textsuperscript{+} or Na\textsuperscript{+}) flow through the MotAB channel provides the power to rotate the flagellum. The left side of the diagram represents a schematic of the Gram-negative structure with the YcgR brake. IM, inner membrane; OM, outer membrane; PG, peptidoglycan. Modified with permission from Belas et al., 2014.
Pili, or fimbriae, are another type of polar filaments that extend from the surface of the bacterium (Figure 1.6 D). Most clinical isolates of *P. aeruginosa* are palliated, and it has been suggested that it is the tips of the pili, which can be either N-methyl-phenyl-alanine (NMePhe) or type IV pili (TFP), that enhance the organism’s association to host cells.\(^{160}\) Pili are made up of homopolymers from the protein pilin, which is encoded by the pilA gene. In *Neisseria gonorrhoea*, which also has type IV pili, it has been shown that pili are markers of virulence and that they may have antiphagocytotic properties.\(^{161}\) *P. aeruginosa* also uses its pili to achieve a special type of motility, termed twitching motility, in which the pilus is extended, attached to the target surface, then retracted, pulling the organism in the desired direction of travel.\(^{162}\)

The structure of *P. aeruginosa* TFP has been proposed by Watts *et al.* and Folkhard *et al.* (Figure 1.5). Pilin, or PilA, protein subunits form a hollow cylindrical structure by means of a helical array arrangement. The outer diameter of this structure is about 5 nm and that of the central channel is about 1.2 nm.\(^{163,164}\) The *P. aeruginosa* TFP displays pilin subunits with tip-base differentiation and a structural domain of pilin which is only accessible at the tip of the pilus, and only this part is responsible for the adhesion function.\(^{161}\) Functionally, the assembly of pilin subunit promotes the extension of TFP, and disassembly promotes retraction.\(^{165}\) This process of assembly/disassembly is driven by the AAA-ATPases PilF (extension) and PilT1/PilT2 (retraction) (Figure 1.5). The outer membrane channel of the TFP machinery is formed by PilQ, whereas PilM, PilN, and PilO, are involved in pilus assembly and may couple the cytoplasmic and periplasmic sides of the pilus.\(^{166}\)

Overall, *P. aeruginosa* flagellum and pili have similar functionality (for attachment) and structure (both are filamentous structures on the surface of the cell), and their motility is controlled by RpoN, especially during initial attachment to the human host and under low nutrient conditions.\(^{167,168}\) The main difference between them is that TFP bind to host N-glycans at the apical surfaces of polarized epithelial cells, whereas flagella bind to the heparan sulphate chains of heparan sulphate proteoglycans at the basolateral surfaces of the polarized epithelium.\(^{96}\)
Figure 1.5 Structure of the type-4 pilus

Schematic representation of the TFP structure in *Thermus thermophiles*, which is representative of the pilus. The type IV pilus machinery is formed from at least 10 different proteins. The PilQ (orange) forms a channel in the outer membrane for secretion of PilA4 (green). PilW (light orange) is involved in DNA transport, PilQ assembly, and pilus extension. PilC (red) is located in the inner membrane and is essential for pilus formation. PilM (light brown), PilN (dark brown), and PilO (beige) possibly connect the periplasmic and cytoplasmic sides of the pilus. The cytoplasmic ATPases PilF (bright yellow) and PilT1/PilT2 (pale yellow) drive pilus extension and retraction (red arrows). Modified with permission from Gold et al., 2015
Figure 1.6 Structural characteristics and virulence factors of *P. aeruginosa*

1.3.2.2 Genomic structure:

*P. aeruginosa* has a unique genomic structure. It possesses a single and supercoiled circular chromosome in the cytoplasm in addition to numerous chromosome-mobilizing plasmids that are instrumental in its pathogenicity. Among these plasmids, the TEM, OXA, and PSE plasmids encode beta lactamase production, which is a key factor in its antibiotic resistance. The genome is relatively large containing 5.3-7.0 Mega base pairs, and the first complete genome sequencing was performed for strain PAO1,\(^\text{[169]}\) derived from an Australian wound isolate from the 1950s. The PAO1 strain has been, and still is, the most common reference for genetic and functional studies on *P. aeruginosa*. The PAO1 genome consists of a circular chromosome containing 6,264,403 base pairs that encode for 5,570 predicted open reading frames and protein sequences, making it the largest bacterial genome to ever be sequenced, with almost the same genetic complexity as simple eukaryotes.\(^\text{[169]}\)

The second *P. aeruginosa* genome sequence was published for strain PA14,\(^\text{[170]}\) a clinical isolate displaying higher virulence than PAO1. In cystic fibrosis (CF) patients, strain LESB58 was found to be highly transmissible and can potentially also cause severe infections in non-CF human hosts.\(^\text{[170, 171]}\) Strain PA7 is a clinical isolate from Argentina with an unusual antimicrobial multi-resistance pattern. It shares only 93.5% nucleotide identity in the core genome with the other sequenced strains, making it a taxonomic outlier within the species. Some of the numerous differences between PA7 and PAO1, including the complete absence of exoS, exoU, exoT and exoY of the type 3 secretion system, are probably the result of a frame shift mutation in the mvfR quorum sensing regulatory gene.\(^\text{[172]}\)

1.3.3 Pathogenesis and Virulence

The pathogenesis of *Pseudomonas* infections is probably multifactorial, as suggested by the number and the diversity of virulence factors possessed by the bacterium, which is to be expected in the wide range of diseases caused by *Pseudomonas aeruginosa*. The virulence of *P. aeruginosa* can be attributed to several factors; Adhesins, in the form of fimbriae, flagella, glycocalyx and alginate may help in association of the bacterium to the epithelium. Invasins, such as elastase, alkaline protease, phospholipase, lecithinase, leukocidin and siderophores all help in deeper invasion and dissemination. Toxins, like Exotoxin A and Exoenzyme S cause tissue destruction and cell death, thus enhancing dissemination. The capsule, lipopolysaccharides and biofilm slime layer all help in evading phagocytosis and host immune responses.
1.3.3.1 **Association and locomotion:**

*P. aeruginosa* is an opportunistic pathogen that rarely affects healthy individuals or breach intact host defences. It usually requires a substantial breach in first-line defences and epithelial barriers in order to initiate infection. Such a breach can result from a bypass of normal cutaneous or mucosal barriers (e.g., trauma, surgery, serious burns, or indwelling devices), disruption of the protective balance of normal mucosal flora by broad-spectrum antibiotics, or alteration of the immunologic defence mechanisms.\textsuperscript{173}

The first step in *P. aeruginosa* infections is colonization of the altered epithelium. Association is primarily mediated by TFP (Figure 1.6 C and Figure 1.5), but several other non-pilus adhesins that may also be responsible for the binding to mucin have been described; their role in the infection process, however, remains unclear. Flagella (Figure 1.6 C and D and Figure 1.4), which are primarily responsible for motility, also act as adhesins to epithelial cells.\textsuperscript{156, 160}

1.3.3.2 **Invasion and dissemination:**

The ability of *P. aeruginosa* to invade tissues depends on its resistance to phagocytosis and host immune defences, and the extracellular enzymes and toxins it produces that break down physical barriers and contribute to bacterial invasion. The bacterial capsule and slime layer effectively protect the bacteria from opsonisation by antibodies, complement deposition, and phagocytosis.\textsuperscript{174}

Proteases, haemolysins, cytotoxins and siderophores produced by *P. aeruginosa* can cause extensive tissue damage, bloodstream invasion, and dissemination.

Several proteases that are released by *P. aeruginosa*, including alkaline protease, LasA elastase, LasB elastase, protease IV and *P. aeruginosa* small protease (PASP), play a major role during acute *P. aeruginosa* infection.\textsuperscript{175-183} Alkaline protease is a 50.4-kDa zinc-dependent metalloprotease of *P. aeruginosa* that utilizes type I secretion for its release.\textsuperscript{184, 185} LasA elastase is a 20-kDa zinc metalloendopeptidase that is synthesized as a pre-proenzyme with a 22-kDa amino-terminal propeptide, but little is known about its maturation and secretion.\textsuperscript{186, 187} It degrades elastin to render it sensitive to degradation by other proteases such as LasB elastase, alkaline protease, and neutrophil elastase. LasA protease action on proteins, including elastin, is limited and directed mainly to Gly3 and Gly2-Ala sequences that are uncommon in most proteins.\textsuperscript{188}

LasB elastase is a type II secreted, zinc-dependent metalloprotease that is initially released as a 52-kDa precursor protein which undergoes auto catalytic cleavage to produce the 33-kDa mature protease and an 18-kDa propeptide.\textsuperscript{177, 188, 189} Premature activation of the enzyme is prevented by the propeptide region, which remains non-covalently associated to the elastase until interrupted by
LasB elastase acts on a number of proteins including elastin, fibrin and collagen. It also lyses fibronectin to expose receptors for bacterial attachment on the surface epithelium. Studies in animal models show that *P. aeruginosa* mutants defective in LasB elastase production are less virulent than their parent strains, which supports the role of LasB elastase as a virulence factor. LasB elastase can inactivate substances such as human immunoglobulin G and A, airway lysozyme and complement components, suggesting that LasB elastase probably interferes with host defence mechanisms. In microbial keratitis, elastase and alkaline protease act together to destroy the ground substance of the cornea and other supporting structures composed of fibrin and elastin, and to inactivate gamma Interferon and tumour necrosis factor.

Cytotoxins, such as the pore-forming protein Leukocidin, are important virulence factor. Although originally named leukocidin because of its effect on neutrophils, it appears to be cytotoxic to most eukaryotic cells, including epithelial cells. This protein probably causes cytotoxicity by increasing cell membrane permeability, ultimately leading to cell death.

Haemolysins, like Phospholipase C, rhamnolipid and lecithinase, appear to act synergistically to break down lipids and lecithin, thereby contributing to tissue invasion through their cytotoxic effect. Rhamnolipid, a rhamnose-containing glycolipid biosurfactant, inhibits the mucociliary transport and ciliary function of human respiratory epithelium, and is believed to solubilize the phospholipids of lung surfactant, making them more accessible to cleavage by phospholipase C, by its detergent like structure.

Siderophores are iron chelating agents that maintain adequate intracellular levels of iron by scavenging and importing it from the extracellular environment. During infections, *P. aeruginosa* produces yellow-green, water-soluble compounds known as pyoverdins, which are considered to be essential virulence factors, since they strongly compete with host cells to obtain the iron essential for their metabolism. Pyocyanin is a blue secondary metabolite produced by *P. aeruginosa* that impairs the normal function of human nasal cilia, disrupts the epithelium and exerts a proinflammatory effect on phagocytes. Pyochelin, a derivative of pyocyanin, sequesters iron from the environment to support the growth of the pathogen.
1.3.3.3 Type III secretion system:

*P. aeruginosa* uses complex, needle-like structures on its surface that function in a highly regulated manner to transport effector proteins into host cells (Figure 1.6 E). These complex processes are collectively known as the type III secretion system (TTSS), and they constitute an important component of the organism’s virulence. Thirty-six genes that are clustered together in the *P. aeruginosa* chromosome are involved in the biogenesis and regulation of the type III secretion system and encode for its complex structure. It is composed of five parts; first, proteins constituting the needle complex that transports substrates from the bacterial cytosol to the extracellular environment; second, proteins that translocate secreted proteins into host cells; third, proteins that regulate the secretion process; fourth, chaperone proteins that facilitate secretion of their cognate partners; and finally, the so called effector proteins that are injected into the host cells.\(^{[203]}\)

The needle complex is 60–120 nm long and 6–10 nm wide, and is responsible for the transport of proteins through the bacterial cytoplastic membrane, the peptidoglycan layer and the outer membrane.\(^{[204]-[206]}\) The needle-like filament, which comprises subunits of the protein PscF, is thought to be a passageway through which secreted factors move and may also serve as a sensor for host cell contact.\(^{[207], [208]}\)

Translocation proteins are specialized proteins that, upon contact with target host cells, accumulate on the tip of the TTSS needle and induce the formation of pores on host cell membranes. This creates a continuation between the bacterial and the host cell cytoplasms, allowing the transfer of bacterial effector proteins into the host cell.\(^{[209]}\) There are three types of translocator proteins, two TTSS loci-encoded membrane hydrophobic translocators and one hydrophilic partner. The hydrophilic translocator PcrV does not directly associate to host cell membranes; rather it provides a platform to enhance the membrane pore formation by the hydrophobic translocators. The major hydrophobic translocator PopB and the minor hydrophobic translocator PopD have been shown to lyse cholesterol rich host cell membranes when associated with their common chaperones PcrH.\(^{[210]}\)

The process of effector protein delivery across the host cell membrane is so efficient that less than 0.1% of secreted effector proteins escape to the extracellular environment.\(^{[211]}\) Host cell death can be caused either directly, through pore-mediated increases in membrane permeability,\(^{[212]-[215]}\) or indirectly, through the activation of broad cellular defence responses.\(^{[216]}\) *P. aeruginosa* type III secretion and translocation apparatus also triggers the inflammasome-mediated activation of caspase 1 through Nod-like receptor ice protease activating factor (IPAF; also known as NLRD4),\(^{[217]-[220]}\) which results in the production of the interleukins IL-1β and IL-18 and in pyroptosis.\(^{[212], [218], [219]}\)
Some effector proteins bind to chaperones before they are secreted, which facilitate their bacterial cytosol storage and the appropriate delivery to the secretion apparatus.\textsuperscript{[221]} SpcS (previously known as orf1) serves as a chaperone for both exoS and exoT and is required for their maximal secretion,\textsuperscript{[222, 223]} while SpcU is a chaperone for exoU.\textsuperscript{[224]}

Four effector proteins of the \textit{P. aeruginosa} TTSS have been identified: ExoS, ExoT, ExoU and ExoY. In isolates from acute infections, the exoS gene is found in 58–72\% of isolates, the exoU gene in 28–42\%, the exoY gene in 89\% and the exoT gene in 92–100\%. Nearly all strains have either the exoS or the exoU gene but not both. The secreted effector proteins define the behaviour of a particular strain during infection, so exoS-secreting strains cause delayed cell death with features of apoptosis, whereas exoU secretion induces rapid host cell lysis.\textsuperscript{[225, 226]}

\textbf{ExoS} is a bifunctional toxin that has both GTPase activating protein (GAP) activity and ADP ribosyl transferase (ADPRT) activity. The GAP domain targets the small GTPases Rho, Rac and cell division cycle 42 (CDC42), which maintain the organization of the host cell actin cytoskeleton. These regulatory GTPases normally switch between an active, GTP-bound form and an inactive, GDP-bound form, but the ExoS GAP domain largely maintains these enzymes in the GDP-bound, inactive form, leading to the disruption of the host cell actin cytoskeleton.\textsuperscript{[227, 228]} This leads to host cell rounding and decreased internalization of \textit{P. aeruginosa}, suggesting that the GAP domain of ExoS may have a role in preventing phagocytosis.\textsuperscript{[229-231]}

\textit{P. aeruginosa} has a clever mechanism in place to prevent ExoS from turning against it. It requires the binding of a eukaryotic host cofactor to bind to the ADPRT domain for the activation of ADPRT activity but not for GAP activity.\textsuperscript{[232, 233]} This requirement ensures that ExoS is not activated until after it is injected into the host cell and bound to the host cell cofactor. Following its injection and activation, the ADPRT activity of ExoS causes an irreversible actin cytoskeleton disruption and inhibition of DNA synthesis, vesicular trafficking and endocytosis and cell death. This possibly contributes to a reduction in cell–cell adherence, which can facilitate \textit{P. aeruginosa} penetration through epithelial barriers into deeper layers.\textsuperscript{[234-237]}

In \textit{P. aeruginosa} infections, there is rapid killing of host immune cells. This could either be the result of the direct cytotoxic effect of ExoS on these cells, or it may be a host cells response to the presence of ExoS to try and eliminate intoxicated cells.\textsuperscript{[203]} The cell death that is induced has features of both apoptosis and necrosis.\textsuperscript{[235, 238]}

\textbf{ExoT} shares 76\% amino acid identity with ExoS and is also a bifunctional toxin with N-terminal GAP activity and carboxy-terminal ADPRT activity. Similar to ExoS, ExoT GAP activity towards Rac, Rho
and CDC42 causes disruption of the actin cytoskeleton that manifests as cell rounding, cell detachment and inhibition of cell migration and phagocytosis.\textsuperscript{[230, 231, 239, 240]} ExoT ADPRT domain also requires binding to a host cell cofactor for activation.\textsuperscript{[241]} However, unlike ExoS, ExoT ADP ribosylates CRKI at Arg20, which interferes with CRK signalling, causing the disruption of signalling to RAC1 and the blockage of cell division at the cytokinesis stage.\textsuperscript{[242, 243]}

The GAP and ADPRT activities of ExoT work together to alter the host actin cytoskeleton and to inhibit cell migration, adhesion and proliferation. This blocks phagocytosis, disrupts epithelial barriers, and thus facilitates bacterial dissemination.\textsuperscript{[230, 243]} ExoT has also been linked to delays in wound healing,\textsuperscript{[244, 245]} and causes apoptosis-like cell death through its ADPRT activity, possibly due to the inhibition of CRK signalling.\textsuperscript{[246]} The contribution of ExoT to disease in mammalian models is modest when compared to that of ExoU and ExoS.\textsuperscript{[230, 243]}

**ExoU** is a potent phospholipase that is capable of causing rapid cell death in eukaryotic cells. It has a chaperone, SpcU, and requires a eukaryotic host cofactor for activation.\textsuperscript{[247, 248]} ExoU possesses a patatin-like domain that confers phospholipase A2 (PLA2) activity, which hydrolyses the ester bond of the acyl group of phospholipids, resulting in the release of free fatty acids and lysophospholipids. This results in host cell death, characterized by rapid loss of plasma membrane integrity, within one to two hours, which is consistent with necrosis.\textsuperscript{[249, 250]} ExoU killing is directed against phagocytes and epithelial cells, thereby promoting bacterial persistence and dissemination.\textsuperscript{[243, 251]}

**ExoY** is a secreted adenyl cyclase that has two extracellular adenyl cyclase (CyaA) domains which act together to bind ATP. It requires a host cell cofactor for full enzymatic activity.\textsuperscript{[252]} Injection of ExoY into mammalian cells results in an elevation of intracellular cAMP concentration,\textsuperscript{[253, 254]} which leads to the disruption of the actin cytoskeleton\textsuperscript{[253, 255]}, inhibition of bacterial uptake by host cells\textsuperscript{[256]} and increased endothelial permeability.\textsuperscript{[252, 257-260]} Such activities would be predicted to lead to more severe disease but this has not been uniformly observed,\textsuperscript{[214, 261]} and the significance of ExoY in infection remains unclear.\textsuperscript{[203]}
1.3.3.4 Quorum sensing:

Bacteria are able to sense their environment, process information, and react appropriately; but they can also sense their own cell density, communicate with each other, and behave as a population instead of individual cells. This phenomenon is called quorum sensing or cell-to-cell signalling and is a common phenomenon described in many bacteria. Both Gram-positive and Gram-negative species utilize various signal molecules to mediate cell-to-cell communication.\cite{253, 254, 262}

Quorum sensing (QS) is a global regulatory system that controls the expression of numerous genes and phenotypes, and is governed by the secretion and detection of signal molecules. When these signals reach a threshold concentration, they bind to specific receptor proteins, which then initiate transcription of the QS-controlled genes. This enables the majority of the bacterial population to simultaneously express a specific phenotype.\cite{255}

Quorum sensing controls the expression of somewhere between 3 and 10% of the genome of \textit{P. aeruginosa};\cite{256, 263} many of these genes either directly mediate the pathogenicity of \textit{P. aeruginosa}, or play significant roles in the persistence of the organism during infection, mainly by contributing to the formation of biofilms that are able to withstand the actions of the immune system. The primary QS system in \textit{P. aeruginosa} is the Las system, which encodes the proteins LasI and LasR.\cite{264} The LasI protein catalyses the production of the AHL molecule, whereas LasR controls virulence factor production, including LasB elastase, LasA elastase (staphylolysin), alkaline protease, exotoxin A and LasI itself.\cite{265-268} Another important QS system in \textit{P. aeruginosa} is the Pseudomonas quinolone signal (PQS), which plays an important role in controlling the production of pyocyanin and rhamnolipid virulence factors.\cite{269} It is therefore clear that QS controls a significant proportion of the virulence factors used by \textit{P. aeruginosa} to mediate infection.

In order for \textit{P. aeruginosa} to overcome host defences, the bacteria need to secrete extracellular virulence factors at high levels, and this is only attainable when the bacterial population reaches a certain density. Quorum sensing allows \textit{P. aeruginosa} to coordinate the expression of virulence factors across the entire bacterial population through the regulated expression of their virulence genes. It can stimulate a host immune response\cite{270} or disrupt the epithelial barrier function.\cite{271} In microbial keratitis, because the host immune response usually results in severe corneal damage, including perforation of the globe and blindness;\cite{272, 273} and because an intact barrier function is essential to defend against \textit{P. aeruginosa} infection,\cite{274, 275} QS could represent a virulence factor in their own right.\cite{255}
1.3.3.5 Biofilm formation

The exceptional ability to survive and thrive in a hostile environment by colonizing surfaces is widespread among bacteria and is due to the sessile mode of growth and the formation of biofilms (Figure 1.6 F). The biofilm mode of growth is an ancient strategy that is extremely resistant to many hostile chemicals in the environment, including disinfectants and antibiotics like aminoglycosides, beta lactams and fluoroquinolones. Biofilms also provide bacteria the ability to avoid or overcome the host innate and active immune defence mechanisms.\(^{[276]}\)

Biofilms are hydrophilic, and water is the predominant constituent of the biofilm. They consist of bacterial microcolonies, constituting 10–20% of the volume of the biofilm, surrounded by exopolysaccharide alginate produced by the bacteria, penetrated by minute anastomosing water channels that form a primitive circulatory system.\(^{[277]}\) One-micrometer latex particles can migrate through these channels by convection, whereas soluble materials are transported by molecular diffusion through the water-filled matrix of the biofilm. An oxygen gradient is present from top to bottom in biofilms so that the bacteria at the base of the biofilm are surrounded by an anaerobic milieu inside the biofilm matrix, whereas the water-filled channels contain dissolved oxygen.\(^{[276]}\)

The thicknesses of biofilms are variable and uneven (13–60 μm), and the growth rate of bacteria in biofilms is reduced compared with that of planktonic bacteria.\(^{[278]}\) Coordinated activity of single cells inside the biofilm can be detected, and is caused by quorum sensing between cells of a similar nature, as can be seen between eukaryotic cells.\(^{[279]}\) Bacterial biofilms are therefore dynamic masses of bacteria, and the proportion between the sessile biofilm and the detached planktonic bacteria depends on unspecific and immunologically specific defence mechanisms in addition to the presence of antibiotics.\(^{[276]}\)
1.3.3.6 Antibiotic Resistance

*Pseudomonas aeruginosa* is notorious for its resistance to antibiotics and disinfectants[^280] and the antibiotic resistance of *P. aeruginosa* strains isolated from cystic fibrosis (CF) patients is significantly higher than strains isolated from non-CF patients.[^281] The organism is naturally resistant to many antibiotics due to several factors. Firstly, it is intrinsically resistant to antimicrobial agents due to a combination of restricted permeability of its outer membrane and the efficient removal of antibiotic molecules that do penetrate by the action of efflux pumps. Secondly, it has the genetic capacity to express a wide array of resistance mechanisms, including mutation in chromosomal genes that regulate resistance. Finally, it can acquire additional resistance genes from other organisms (such as bacilli, actinomycetes and moulds) via plasmids (both R-factors and RTFs), horizontal gene transfer (mainly transduction and conjugation), transposons, and bacteriophages.[^281]

*Pseudomonas aeruginosa* tends to colonize surfaces in the anionic polysaccharide alginate biofilm form, and this makes the bacteria resistant to therapeutic concentrations of antibiotics. The high bacterial cell density and the physical exclusion of the antibiotic inside the biofilm matrix are also factors that aid in the resistance of biofilms to antibiotics. A general stress response within the biofilm leads to physiological changes in the bacterial cells, where protective mechanisms and induced key metabolic processes are shut down.[^282] The overall resistance is dependent on the interaction between the entire population of cells within the biofilm. This population is heterogeneous, containing fast- and slow-growing cells, with different modes of antibiotic resistance. Some of these bacteria are resistant through expression of inactivating enzymes and efflux pumps, others through non-expression of these systems.[^283]
1.3.4 Clinical Significance

Most pseudomonads are known to be pathogens of plants rather than animals, but three *Pseudomonas* species are regularly pathogenic to humans: *Pseudomonas mallei*, *Pseudomonas pseudomallei* and *Pseudomonas aeruginosa*. *Pseudomonas mallei*, also known as *Burkholderia mallei*, causes a disease in horses known as glanders. *Pseudomonas pseudomallei*, also known as *Burkholderia pseudomallei*, is the agent of melioidosis, a highly fatal tropical disease of humans and other mammals, and is also an opportunistic pathogen contracted through the contamination of wounds with mud or soil.\[284\]

*Pseudomonas aeruginosa* is an opportunistic human pathogen, and it rarely infects healthy individuals. It often colonizes immunocompromised patients, like those with cystic fibrosis, cancer, or AIDS. It complicates 90% of cystic fibrosis cases and affects up to two thirds of critically-ill hospitalized patients, usually causing more devastating disease. It is also one of the leading Gram negative nosocomial infections, leading to a 40-60% mortality rate. Within the hospital, *P. aeruginosa* has numerous reservoirs, including disinfectants, respiratory equipment, food, sinks, taps, toilets, showers and mops. Furthermore, it is constantly reintroduced into the hospital environment on fruits, plants, vegetables, as well by visitors and patients transferred from other facilities. Spread occurs from patient to patient on the hands of hospital personnel, by direct patient contact with contaminated reservoirs, and by the ingestion of contaminated foods and water. The organism, however, also colonizes the oropharynx of up to 6% and is recovered from the faeces of 3-24% of healthy individuals.\[140, 285\]
Table 1.3 Diseases caused by Pseudomonas aeruginosa

<table>
<thead>
<tr>
<th>Disease</th>
<th>Role of <em>P. aeruginosa</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacteraemia and septicaemia</td>
<td><em>Pseudomonas aeruginosa</em> causes bacteremia primarily in immunocompromised patients. Predisposing conditions include hematologic malignancies, immunodeficiency relating to AIDS, neutropenia, diabetes mellitus, and severe burns. Most <em>Pseudomonas</em> bacteremia is acquired in hospitals and nursing homes. <em>Pseudomonas</em> accounts for about 25% of all hospital acquired Gram-negative bacteremias. <em>P. aeruginosa</em> septicaemia is rare, but it can be associated with high morbidity and mortality, as a result of the marked toxaemia, antibiotic resistance and severity of underlying disease.</td>
</tr>
<tr>
<td>Bone and joint infections</td>
<td><em>Pseudomonas</em> infections of bones and joints result from direct inoculation of the bacteria or the haematogenous spread of the bacteria from other primary sites of infection. Blood-borne infections are most often seen in IV drug users and in conjunction with urinary tract or pelvic infections. <em>P. aeruginosa</em> has a particular predilection for fibro-cartilaginous joints of the axial skeleton. It is the most common pathogen isolated in osteochondritis after puncture wounds of the foot, and causes chronic contiguous osteomyelitis, usually through direct inoculation of the bone.</td>
</tr>
<tr>
<td>Central nervous system infections</td>
<td><em>Pseudomonas aeruginosa</em> causes meningitis and brain abscesses. The organism can invade the CNS from a contiguous structure such as the inner ear or a paranasal sinus. It can be directly inoculated during head trauma, surgery or invasive diagnostic procedures, or it can spread from a distant site of infection such as the urinary tract.</td>
</tr>
<tr>
<td>Ear infections including otitis externa</td>
<td><em>Pseudomonas aeruginosa</em> is the predominant bacterial pathogen in otitis externa, including swimmer's ear. The bacterium is infrequently found in the normal ear, but often colonizes the external auditory canal in association with injury, maceration, inflammation, or simply wet and humid conditions.</td>
</tr>
<tr>
<td>Endocarditis</td>
<td><em>Pseudomonas aeruginosa</em> infects heart valves of IV drug users and prosthetic heart valves. The organism establishes itself on the endocardium by direct invasion from the blood stream.</td>
</tr>
<tr>
<td>Eye infections</td>
<td><em>Pseudomonas aeruginosa</em> can cause devastating infections in the eye. It is one of the most common causes of bacterial keratitis and it is the most common cause of contact lens associated microbial keratitis. It is one of the causative agents of infective endophthalmitis and has also been isolated as one of the etiologic agents of neonatal ophthalmia.</td>
</tr>
<tr>
<td>Gastrointestinal infections</td>
<td><em>Pseudomonas aeruginosa</em> can affect any part of the gastrointestinal tract from the oropharynx to the rectum, primarily in immunocompromised individuals. The organism has been implicated in perirectal infections, paediatric diarrhoea, typical gastroenteritis, and necrotizing enterocolitis. The gastrointestinal tract is also an important portal of entry in <em>P. aeruginosa</em> septicemia and bacteremia.</td>
</tr>
<tr>
<td>Respiratory infections</td>
<td>Respiratory infections caused by <em>Pseudomonas aeruginosa</em> largely occur in individuals with a compromised lower respiratory tract or a compromised immunity. Primary pneumonia occurs in patients with chronic lung disease and congestive heart failure, whereas bacteraemic pneumonia commonly occurs in neutropenic cancer patients undergoing chemotherapy. Lower respiratory tract colonization of cystic fibrosis patients by mucoid strains of <em>P. aeruginosa</em> is common and almost impossible to eradicate.</td>
</tr>
<tr>
<td>Skin and soft tissue infections</td>
<td><em>Pseudomonas aeruginosa</em> can cause a variety of skin infections, both localized and diffuse. Common predisposing factors include burns, trauma, dermatitis, high moisture conditions such as those found in the toe webs of athletes, hikers and combat troops, in the perineal region and under diapers of infants, and on the skin of whirlpool and hot tub users. Individuals with AIDS can be easily infected. <em>P. aeruginosa</em> has also been implicated in folliculitis and in some resistant forms of acne vulgaris.</td>
</tr>
<tr>
<td>Urinary tract infections</td>
<td><em>Pseudomonas aeruginosa</em> is the third leading cause of hospital-acquired UTIs, accounting for about 12% of all infections of this type. Urinary tract infections caused by <em>P. aeruginosa</em> are usually hospital-acquired and can occur via ascending or descending routes, usually secondary to urinary tract catheterization, instrumentation or surgery.</td>
</tr>
</tbody>
</table>
1.4 The Barrier: Innate corneal defences against infection

1.4.1 Background

The normal inflammatory processes that occur in response to injury or infection to re-establish the integrity of the affected tissue can also cause significant injury to host tissue. This is because they depend on degradation of the affected tissue, proliferation of fibroblasts and new blood vessels to replace damaged structures, and tissue remodelling to restore the strength of the affected site. In most body surface tissues, this damage and scarring are acceptable outcomes, as they do not interfere with the primary function of the tissue. In the cornea, however, repair can result in significant loss of function, refractive distortion and opacity, leading to significant reduction in visual function, despite the full restoration of structural integrity. As a result, the cornea has evolved several complex adaptations to limit and regulate inflammation. At the same time, it has also developed formidable defences to prevent infection, and, in the event of an infection, additional adaptations that initiate robust immune and inflammatory responses to preserve ocular integrity.

1.4.2 The ocular immune privilege

Several sites in the body normally require tight regulation of inflammatory and immune responses to prevent potentially damaging consequences of these responses to delicate anatomic structures. These sites include the eye, the brain and the pregnant uterus and are collectively termed immune-privileged sites.[305, 306] The ocular immune privilege entails unique anatomic, physiologic, and immunoregulatory properties of immune and parenchymal cells, functioning as an integrated unit to develop a mechanism that suppresses the intense local immunogenic inflammation that would have otherwise developed in its absence.[306] The ocular immune privilege is provided by several physiological and regulatory phenomena, including the anterior chamber associated immune deviation (ACAID) and soluble and cell membrane-bound immuno-suppressive factors in the anterior segment (Table 1.4). Ocular resident cells, such Müller cells in the retina, directly inhibit immune cells by secreting soluble factors and by contact-dependent mechanisms. The iris and ciliary body pigment epithelial cells, and the retinal pigment epithelium (RPE) inhibit T cells and even induce them to become T regulatory cells.[307] In addition, the conjunctiva-associated lymphoid tissue (CALT) is important in the induction of tolerance to ubiquitous non-pathogenic antigens.[308] The cornea and the retina are devoid of lymphatics in the native state, although lymphangiogenesis is induced in the inflamed cornea,[309] and this also contributes to the dampened immune responses in the quiescent eye.
Table 1.4 Factors contributing to ocular immune privilege

<table>
<thead>
<tr>
<th>Factor</th>
<th>Location</th>
<th>Immune privilege effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood ocular barrier</td>
<td>Tight junctions of epithelial cells, iris, ciliary body, vascular endothelial cells</td>
<td>Exclusion of macromolecules and inflammatory cells</td>
</tr>
<tr>
<td>TGF-β2</td>
<td>Aqueous humor</td>
<td>Inhibition of T-cell proliferation</td>
</tr>
<tr>
<td>VIP</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>Müller cells</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>Pigmented epithelial cells</td>
<td>Retina and uveal tract</td>
<td></td>
</tr>
<tr>
<td>TGF-β2</td>
<td>Aqueous humor</td>
<td>Inhibition of IFN-γ release by Th1 cells</td>
</tr>
<tr>
<td>VIP</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>α-MSH</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>Pigment epithelium</td>
<td>Retina and uveal tract</td>
<td></td>
</tr>
<tr>
<td>TGF-β2</td>
<td>Aqueous humor</td>
<td>Inhibition of pro-inflammatory factor release by macrophages</td>
</tr>
<tr>
<td>CGRP</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>α-MSH</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>TGF-β2</td>
<td>Aqueous humor</td>
<td>Inhibition of NK-cell activity</td>
</tr>
<tr>
<td>MIF</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>Qa-2</td>
<td>Cell membranes</td>
<td></td>
</tr>
<tr>
<td>ACAID</td>
<td>Anterior chamber, vitreous</td>
<td>Inhibition of DTH</td>
</tr>
<tr>
<td>TGF-β2</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>CGRP</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>α-MSH</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>VIP</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>FasL</td>
<td>Cornea, retina, iris, ciliary body</td>
<td>Deletion of infiltrating inflammatory cells</td>
</tr>
<tr>
<td>Membrane-bound complement regulatory proteins</td>
<td>Cornea</td>
<td>Inhibition of complement cascade</td>
</tr>
<tr>
<td>CD55</td>
<td>Corneal epithelium, stroma</td>
<td></td>
</tr>
<tr>
<td>Crry</td>
<td>Ciliary body</td>
<td></td>
</tr>
<tr>
<td>Soluble complement-regulatory proteins</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>CD55</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>CD59</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>CD46</td>
<td>Aqueous humor</td>
<td></td>
</tr>
<tr>
<td>&lt;1.3 kDa inhibitor of C1q</td>
<td>Aqueous humor</td>
<td></td>
</tr>
</tbody>
</table>


ACAITD, anterior chamber associated immune deviation; CGRP, calcitonin gene-related peptide; Crry, cell surface regulator of complement; DTH, delayed type hypersensitivity; FasL, Fas ligand; MSH, melanocyte stimulating hormone; TGF-β, transforming growth factor β; VIP, vasoactive intestinal peptide.
1.4.2.1 The Anterior Chamber Associated Immune Deviation

The Anterior Chamber Associated Immune Deviation (ACAID) is an unusual systemic immune response, whereby following the introduction of a foreign antigen into the anterior chamber of the eye, a signal is produced that communicates with the immune system through the spleen. This leads to a series of events with several important features: inhibition of systemic delayed-type hypersensitivity, inhibition of complement-fixing antibody responses, maintenance of a normal cytotoxic T-cell and humoral immune response, and transfer of ACAID to immunologically naive recipients through antigen-specific splenic suppressor CD4+ and CD8+ T-cells.

There are three phases of ACAID. The first is the ocular phase, which involves the generation of an antigen-specific signal that is generated in the eye and delivered to the systemic immune system. This signal is produced by the F4/80+ antigen-presenting cells (APC) in the anterior chamber, which are exposed to a variety of immunosuppressive agents (Table 1.4). This leads to a reduced capacity to Th1 inducing IL-12, an enhanced production of the Th2 cytokine IL-10, a reduced expression of CD40 co-stimulatory molecule and the autocrine production of transforming growth factor-beta (TGF-β). In the second thymic phase, F4/80+ APC migrate to the thymus where they generate NK1+, CD8+, CD4- efferent cells, which suppress the delayed-type hypersensitivity. Finally, in the third splenic phase, which spans at least 7 days in the spleen, the F4/80+ APC migrate to the spleen where they generate CD8+ regulatory suppressor cells.

Two cell populations are responsible for ACAID. The first are CD4+ cells that produce increased amounts of IL-10 and decreased amounts of IFN-γ. These Th1 type cells are called afferent suppressor cells and are required to generate a second population of CD8+ cells, called the efferent suppressor cells, which in turn inhibit the expression of delayed-type hypersensitivity responses. T-cells bearing the γδ T-cell receptor and B cells are also required for ACAID.

In addition to ACAID, many other immunoregulatory phenomena contribute to the ocular immune privilege (Table 6.6). The blood-ocular barrier, located at the tight junctions between the cells of the ciliary epithelium, physically provides a barrier to cellular infiltration. In addition, soluble factors inhibit a variety of immunological processes, including T-cell proliferation, IFN-γ production by Th1 cells, proinflammatory factors secreted by macrophages, natural killer (NK) cell activity and delayed hypersensitivity responses. There are also both bound and soluble complement regulatory proteins that inhibit the complement cascade.
1.4.3 Corneal barriers against infection

Despite the continuous exposure of the cornea to ubiquitous microorganisms that naturally inhabit its surroundings, invasive infection rarely happens. Even when intact mouse or rat corneas were inoculated by extremely large inocula of both invasive and cytotoxic strains of *P. aeruginosa*, the ocular surface was rapidly cleared from offending bacteria without pathology.\[^{313}\] In contrast, corneal epithelial cells cultured in vivo, interestingly, are easily invaded and killed by most clinical and laboratory isolates of *P. aeruginosa*.\[^{314, 315}\] This suggests that the healthy eye possesses certain defence mechanisms against infection that are absent from laboratory cell culture conditions. Since these mechanisms are always present, even when the cornea is healthy, they are likely different from the host immune responses that are activated when an infection dose occur.\[^{173}\]

Firstly, the conjunctiva lies immediately adjacent to the cornea, and it is provided with a system that detects antigens and induces an immune response by the direct action of the lymphatic cells or the secretion of soluble antibodies termed the conjunctiva-associated lymphoid tissue (CALT).\[^{308}\] This system of lymphoid tissue was mainly observed in the palpebral conjunctiva, occurring in the form of a diffuse lymphoid layer of lymphocytes and plasma cells in the lamina propria and organized follicular lymphoid accumulations composed of B lymphocytes apically covered by lymphoepithelium. Thus, this system contains all components necessary for a complete immune response, which can serve an important function in the early protection against corneal infections.

An important in vivo factor missing in culture conditions is tear fluid. Although many *P. aeruginosa* strains grow readily in undiluted tear fluid despite its antimicrobial components, such as lactoferrin and lysozyme,\[^{316, 317}\] tear fluid can protect both cultured corneal epithelial cells *in vitro* and injured and healing mouse corneas *in vivo* from infection by invasive and cytotoxic strains of *P. aeruginosa*.\[^{318, 319}\] The tear fluid physically contributes to the cleansing of the ocular surface through the mechanical action of blinking, but it is also thought to directly act on corneal epithelial cells to make them more resistant to invasion and cytotoxicity by *P. aeruginosa*.\[^{313}\] In one study, human corneal epithelial cells pre-treated with human tears were more resistant to invasion and cell death caused by *P. aeruginosa*,\[^{320}\] and demonstrated an upregulation of nuclear factor kB (NF-kB) and activating protein 1 (AP-1), along with the up- and downregulation of genes encoding for cytokines, transcription factors and junctional proteins.\[^{320}\]
In another study, tear fluid increased the barrier function of corneal epithelial cells in vitro,\textsuperscript{[319]} which can help the multi-layered corneal epithelium protect itself against microbial traversal. This protective function of tear fluid may replace the well-established roles believed to be played by commensal microbes in modulating ocular homeostasis and innate immune defence.\textsuperscript{[173]} Although tear fluid does not directly affect bacterial viability, it contains several factors that can bind microbes and suppress bacterial virulence factors, like mucins, secretory immunoglobulin A, and surfactant protein D, and this can ultimately alter the interactions between \textit{P. aeruginosa} and corneal epithelial cells.\textsuperscript{[321-323]}

In many host systems where there is interaction between bacteria and epithelial cells, bacteria tend to avoid the apical surfaces of epithelial cells. This has been observed in the gastrointestinal system, for instance, where gut bacteria, despite their enormous numbers, keep a zone that is about 50 µm wide free of microbes above the epithelial cell surface, and this is thought to be due to mucins and other antimicrobial factors present on the cell surface.\textsuperscript{[324, 325]} A similar phenomenon has also been shown by live cell imaging of the corneal epithelium, where \textit{P. aeruginosa} bacteria avoided the apical surfaces of corneal epithelial cells, and maintained a safe distance from the surface although always remaining within range of them.\textsuperscript{[315]} This may be due to the presence of an environment similar to that of the gut, where mucins and other novel antimicrobial compounds are expressed from the surface.\textsuperscript{[326, 327]} The apical surfaces of corneal epithelium cells are covered by a dense glyocalyx that is formed of MUC1, MUC4, and MUC16 mucins, and these are thought to be important in preventing association of bacteria, such as \textit{P. aeruginosa} and \textit{S. aureus}, to corneal epithelial cells.\textsuperscript{[328, 329]} Lysates of human corneal epithelial cells contain keratin-derived antimicrobial peptides (KDAMPs), which rapidly kill several clinical isolates of \textit{P. aeruginosa}, \textit{S. pyogenes} and \textit{S. aureus}. In mutant mice deficient in cytokeratin 6A, from which KDAMPs are derived, the antimicrobial activity of the corneal epithelial cell lysates was reduced, and the corneal epithelium was significantly more susceptible to bacterial association \textit{in vivo}.\textsuperscript{[326]}

Another factor that is different \textit{in vivo} from culture conditions is the three dimensional arrangement of epithelial cells in the epithelium as a tissue, with all the junctional complexes and interactions between cells in different layers. The intact corneal epithelium is a formidable barrier to infection, and most animal models designed to study corneal infections bypass the epithelium by design, either by scarifying the epithelium, or by bypassing it all together and directly injecting the pathogens into the stroma. The pathology that occurs during microbial keratitis requires bacterial entry into the stroma, which then leads to the activation of inflammatory and immune responses and their subsequent damaging sequelae.\textsuperscript{[330, 331]}
The resistance of the intact corneal epithelium to bacterial traversal can be attributed to a multitude of factors. As a first line of defence, superficial epithelial cells desquamate to reduce the microbial load of the offending inoculum. When *P. aeruginosa* is inoculated onto a healthy rat cornea, most internalized bacteria are found in cells that are readily shed from the eye with rinsing.\(^{[314]}\) This supports the theory that internalization followed by desquamation is a mechanism for clearing bacteria that manage to adhere to the surface. Tight junctions between superficial cells are another major contributing factor, but other factors are also likely to be important. In a study where tissue paper blotting of the corneal surface was used to disrupt superficial tight junctions,\(^{[332]}\) *P. aeruginosa* could adhere to superficial cells, but could not penetrate beyond the epithelial surface, despite the loss of barrier function to fluorescein. Thus, the loss of tight junctions promoted bacterial association to superficial cells, but prevented adherent bacteria from penetrating, suggesting that the corneal epithelial defences to these two initial steps of infection are separable, and involve multiple different factors.

In corneas treated with the calcium chelator ethylene glycol tetra-acetic acid (EGTA), bacteria were able to penetrate the epithelium, even after tissue paper blotting. This could be attributable to disruption of other types of cell-to-cell junctions in the absence of calcium, since their integrity is generally calcium-dependent,\(^{[332]}\) or to other defence mechanisms of the corneal epithelial surface, such as surfactant protein D, whose activity is also calcium dependent.\(^{[333]}\) This is supported by the observation that *P. aeruginosa* could partially traverse the tissue paper–blotted corneal epithelium of SP-D knockout mice *in vivo*.\(^{[332]}\)

Other factors may be involved in epithelial defence against *P. aeruginosa* traversal. For instance, the corneas of mice deficient in myeloid differentiation primary response protein 88 (MyD88) are susceptible to *P. aeruginosa* penetration without the need for tissue paper blotting or EGTA treatment.\(^{[334]}\) MyD88 is an essential component of toll-like receptor (TLR) signalling and IL-1R signalling, and its activity leads to activation of cytokines and chemokines, secretion of antimicrobial peptides, and recruitment of phagocytic cells.\(^{[335-338]}\) It is important in the regulation of expression of antimicrobial peptides, like human beta defensin 2 (hBD-2) and cathelicidin LL-37, both of which are expressed by corneal epithelial cells stimulated by TLR or IL-1R agonists, and this may also demonstrate its involvement in corneal epithelial defence against bacterial association and traversal.\(^{[338,339]}\) In fact, hBD-2 has already been shown to be important in protecting the corneal epithelium against *P. aeruginosa* colonization.\(^{[340]}\)
If *P. aeruginosa* bacteria do manage to traverse the multi-layered corneal epithelium and all of its defences, the epithelial basement membrane (basal lamina), composed mostly of extracellular matrix proteins, like laminin and type IV collagen, prevents them from actually entering the corneal stroma. This is because the basal lamina acts as a physical filter, as its pores are smaller than the size of most bacteria.\[^{341}\] This may explain why even when the corneal epithelium is made susceptible to bacterial association and traversal, using either EGTA or MyD88 knockout mice, this does not always lead to microbial keratitis.\[^{332,334}\] Basal lamina extracellular matrix proteins can also form a chemical barrier to bacterial passage.\[^{100}\] In addition, they improve the barrier function of the epithelial cells growing on top of them, possibly through the effects on junctional integrity or mucin and antimicrobial peptide expression of the cells.\[^{173}\]
1.5 The Challenge: Pathogenesis of microbial keratitis

1.5.1 Background

When the bacteria do reach the corneal stroma, they are able to start colonizing the tissue, and a corneal ulcer develops. An ulcer is defined as a local epithelial defect with excavation of tissue and infiltration by immune cells. Break down of the corneal epithelium is a prerequisite for development of stromal melting and tissue loss, and a healthy corneal epithelium prevents stromal degradation and is essential for stromal healing, as demonstrated by several studies. Most corneal ulcers are associated with a marked inflammatory response which typically involves dense infiltration of neutrophils and other leukocytes, and physical blockade or depletion of neutrophils and other leukocytes in animal models prevents ulceration.

Most corneal ulcers are associated with a marked inflammatory response which typically involves dense infiltration of neutrophils and other leukocytes, and physical blockade or depletion of neutrophils and other leukocytes in animal models prevents ulceration. The typical stromal necrosis and tissue loss seen in microbial keratitis are caused by enzymatic degradation of extracellular matrix by pathogen and host released proteases. These involve the interplay of several host and pathogen factors that initiate and develop the progress of the pathogenesis. Although the infection is not initiated until the epithelial defences are overcome or bypassed, the cornea has additional innate defences and immune mechanisms that help to fight the infection. These mechanisms alert the immune system and recruit immune cells to the area, which in turn combat the offending microbes by phagocytosis and secretion of lytic enzymes, and by secreting cytokines and chemokines to recruit more immune cells to help with clearing out the pathogens.

1.5.2 Pathogen recognition and innate immunity in the corneal epithelium

One of the initial steps in the inflammatory process in relation to microbes is activation of the Toll-like receptor (TLR) family of pathogen recognition molecules. The TLR family is a class of single membrane-spanning non-catalytic receptors that recognize structurally conserved molecules derived from microbes, namely pathogen associated molecular patterns (PAMPs), once they have breached physical barriers, such as the corneal epithelium, and activate immune cell responses. Thirteen TLRs have been identified in humans and mice, which respond to lipids, proteins or nucleic acids.

TLR4, the most complex member of this receptor family, recognizes lipopolysaccharide (LPS), but it must be associated with the co-stimulatory molecules LPS binding protein (LBP), CD14, and MD-2 to recognise it. MD-2 is a bifunctional, 25-kD protein that is essential for TLR4 signalling by coupling endotoxin recognition to TLR4 activation. LBP and CD14 combine to extract single endotoxin molecules from endotoxin aggregates or from bacterial outer membrane and form monomeric endotoxin:CD14 complexes. LPS is then transferred from CD14 to MD-2, which,
together with the other accessory molecules, facilitate recognition of the endotoxin at very low concentrations.\textsuperscript{348, 349}

In \textit{P. aeruginosa}, LPS and flagellin are recognised by TLR4 and TLR5 receptors, respectively,\textsuperscript{350} initiating a pathway that involves recruiting the adaptor molecule MyD88, which mediates signalling by all TLRs except TLR3, and also mediates signalling by interleukin (IL)-1R and IL-18R. MyD88 recruits IL-1 receptor-associated kinase 1 (IRAK1) and IRAK4, and leads to activation of transforming growth factor (TGF)-\(\beta\)-associated kinase 1 (TAK1), the mitogen-activated protein (MAP) kinases p38MAPK and JNK8.\textsuperscript{335} These MAP kinases mediate the phosphorylation of transcription factors NF\(\kappa\)B and AP-1, and the transcription of proinflammatory and chemotactic cytokines.\textsuperscript{351-354} TLR3 and TLR4 lead to activation of the TRIF-dependent pathway, and this leads to the production of type I interferons associated with viral infection. Thus, TLR4 activates cells through two independent signalling pathways, leading to a diverse and amplified response.\textsuperscript{335}

Activation of the TLR4 and TLR3 receptors leads to recruitment of the adaptor molecule TIR domain-containing adaptor inducing IFN-\(\beta\) (TRIF),\textsuperscript{355} which signals to the NF-kB and JNK pathways through RIP1 and TRAF6. TRIF also activates TRAF3, which leads to phosphorylation of IRF3 through TBK1 and IKK, and stimulates the production of type I interferons.\textsuperscript{354, 356} The presence of TLR4 in \textit{P. aeruginosa} corneal infection has also been associated with the release of antimicrobial factors such as inducible nitric oxide synthase (iNOS) and hBD-2.\textsuperscript{357}

In cell cultures, human corneal epithelial cells respond to TLR2, TLR3, and TLR5 ligands by producing antimicrobial peptides, such as \(\beta\)-defensins and cathelicidin,\textsuperscript{327, 338, 358} but, interestingly, are unresponsive to LPS.\textsuperscript{359, 360} This is because, although TLR4 is constitutively expressed on the surface of human corneal epithelial cells, it is the MD-2 that is not expressed, and thus LPS responsiveness is only demonstrated in the presence of either exogenous MD-2 or by transfection with an MD-2 expressing plasmid.\textsuperscript{360-362} This is supported by the finding that incubation with IFN-\(\gamma\), which activates the Jak2/STAT1 pathway and the binding of p-STAT1 to IFN Gamma Activating Sites (GAS) on the MD-2 promoter, results in MD-2 gene and surface expression, and LPS responsiveness.\textsuperscript{363} As IFN-\(\gamma\) is produced by infiltrating natural killer cells during \textit{P. aeruginosa} corneal infection,\textsuperscript{363, 364} it is likely that MD-2 expression and LPS responsiveness by corneal epithelial cells, that further contributes to the inflammatory response during active bacterial infection by producing proinflammatory and chemotactic cytokines, is secondary to that of myeloid immune cells.\textsuperscript{365}
1.5.3 Recruitment of immune cells into the cornea

Recruitment of adaptor molecules MyD88 to TLR4 and TLR5, and recruitment of TIRAP and TRIF to TLR4, all lead to NF-κB translocation to the nucleus, and expression of pro-inflammatory and chemotactic cytokines. These include CXCL1/KC, which recruits neutrophils from limbal capillaries to the corneal stroma, and IL-1α and IL-1β, which then activate the IL-1R1/MyD88 pathway in macrophages and resident corneal epithelial cells and keratocytes.\textsuperscript{350}

Interleukin-1β (IL-1β) is a proinflammatory cytokine produced by activated macrophages and monocytes. It is involved in the generation of systemic and local responses to infection, injury, and immunological challenges and is the primary cause of chronic and acute inflammation. It is also a key player in the febrile response.\textsuperscript{366} IL-1β is produced as an inactive cytoplasmic precursor (proIL-1β, p35) that must be cleaved to generate the mature active form (p17). The IL-1β-converting enzyme, better known as caspase-1\textsuperscript{367-369} is required for this cleavage. Caspase-1 is a member of a family of inflammatory caspases, including human caspase-4 and -5, and mouse caspase-11 and -12, that contains an N-terminal caspase recruitment domain (CARD).\textsuperscript{370}

Production of the mature IL-1β is a two-step process. Initially, the transcription of the 31kDa pro-form is induced, followed by post-translational cleavage of the preform to the 17kDa mature form.\textsuperscript{371, 372} The assembly of a multiprotein complex containing one or more Nod-like receptors (NLRs), called inflammasomes, activates Caspase-1. Normally expressed in macrophages, the NLRC4 inflammasome, also termed IPAF, is activated by flagellin or type III secretion proteins, leading to caspase-1 recruitment and IL-1β cleavage.\textsuperscript{365} The 31kD pro-IL-1β can be cleaved at different sites to produce the 18kD mature form of IL-1β by either caspase-1 or elastase, resulting in different sized cleavage products.\textsuperscript{373, 374} It is thought that, during the early stages of infection, IL-1β processing by neutrophils plays an important role, while inflammasome activity by macrophage, and possibly neutrophil, contribute at a later stage.\textsuperscript{365}

The recruitment of neutrophils from limbal blood vessels to the site of infection in the corneal stroma, and the subsequent bacterial killing, is likely due to the effect of both IL-1α and IL-1β acting together. When recruited, neutrophils generate reactive oxygen species (ROS) utilizing NADPH oxidase.\textsuperscript{375} This production of ROS and MMPs, including collagenases, disrupts the extracellular matrix and prevents microbial dissemination.\textsuperscript{376} In the cornea, this disruption of the collagen matrix results in loss of corneal clarity and impaired vision.
The normal corneal stroma was always believed to be devoid of myeloid immune cells, but it is now known that this is not entirely accurate. It is generally acknowledged that a number of corneal stimuli, including, infection, trauma, and cauterization, result in the presence of MHC class II+ dendritic cells (DC) in the central cornea, and this was, until recently, presumed to be entirely a result of de novo infiltration of these cells into the cornea. \[377, 378\] Recent studies, however, demonstrate that the central stroma contains exclusively MHC class II+ CD80+ CD86+ dendritic cells. Furthermore, corneas stained for CD14, an “immature” cell-surface marker reported to be associated with lack of differentiation of DC, demonstrate a large number of CD14+ cells in the stroma. This indicates that although myeloid cells are present in the normal stroma, they are undifferentiated monocytic precursor cells distinct from DC and macrophage populations present elsewhere. Thus, in contrast to other organs, where terminally differentiated populations of resident DC and/or macrophages outnumber colonizing precursors, large numbers of DC within the cornea remain in an undifferentiated state, and these cells only mature to a DC or a macrophage phenotype in response to inciting corneal stimuli. \[379\] Since corneal fibroblasts are the principal cell type present in the stroma, they may well have a role in immune activation when exposed to pathogens, perhaps as a first line of defence against progressive corneal ulceration. This has been described in other systems, and fibroblasts are regarded by some as resident sentinel cells capable of synthesizing chemokines, that initiate wound healing and clearance of invading microorganisms, and condition the cellular and cytokine environment in areas of inflammation by an array of factors they express. \[380\] In addition, fibroblasts can be activated to control extracellular matrix synthesis and produce cytokines and chemokines \[381, 382\] which is analogous to cells of the immune system, such as macrophages and B and T lymphocytes.

### 1.5.4 Proteases involved in the pathogenesis of corneal ulceration

The progressive tissue destruction and necrosis seen in infective corneal ulceration is caused by several enzymes liberated by offending bacteria, as well as enzymes released by local host epithelial and stromal cells and infiltrating immune cells. \[383-389\] This process is thought to be sequential, initiated by bacterial exoenzymes, like alkaline protease and elastase, but maintained and exacerbated by proteases released by resident corneal cells \[390\] and stimulated inflammatory cells. \[272, 391, 392\] Three types of proteases appear to be important in the pathogenesis of stromal necrosis:

- The plasmin system, the matrix metalloproteases (MMPs), and the cysteine proteinases and cathepsins released during phagocytosis.

Plasmin is a serine protease that is mainly used by body tissues to dissolve fibrin clots, although it can hydrolyse other substrates, like fibronectin. It normally exists as the proenzyme plasminogen in
blood, tears, cornea, and aqueous humor, and is converted into active plasmin by tissue plasminogen activator (t-PA) and urokinase plasminogen activator (u-PA). Plasminogen activators are present in many tissues, including the cornea, and on bacterial surfaces, where they facilitate bacterial penetration and migration by activating plasmin and dissolving fibrin clots. In the cornea, plasmin plays an important role in normal wound healing, where it dissolves the fibrin clot that fills the gap at the injury site, and allows epithelial cells to migrate and heal the defect. Plasminogen is constitutively expressed in the cornea, and becomes activated by t-PA in the event of injury. If the epithelial healing is not achieved, however, the continuous expression of t-PA leads to persistent activation of plasmin, which can exacerbate stromal degradation, either by its direct proteolytic action, or indirectly through activation of collagenase and other matrix metalloproteinases. Corneal injury also leads to elevated plasmin levels in tears, which may promote stromal necrosis even further.

Matrix metalloproteinases (MMP) are proteases that depend on zinc as an essential coenzyme and are important in degradation of the extracellular matrix. There are at least 17 different MMPs (Table 1.5) and they can be categorized into five groups: gelatines (MMP-2 and 9), collagenases (MMP-1, 8, 13 and 18), stromelysins (MMP-3, 10 and 11) membrane-type MMPs, and others. Matrix metalloproteinases act on a variety of extracellular matrix molecules including interstitial collagens and proteoglycans. MMPs are synthesized and secreted as inactive pro enzymes and are activated by cleavage of their N-terminus, through a series of complex cascades, in the extracellular space. The activation MMP procollagenase is triggered by several factors. Initially, the conversion of plasminogen to plasmin after the secretion of t-PA partially activates procollagenase. This is then potentiated by the activity of other MMPs produced by activated fibroblasts. Collagenase activation is also triggered by pathogen factors, like LPS, and host factors, like inflammatory cytokines, like IL-1, and growth factors, like epidermal growth factor (EGF), platelet aggregating factor (PAF), and platelet-derived growth factor (PDGF).

Because of their strong proteolytic activity, MMPs production, activation and degradation are tightly regulated. They are typically only released when tissue remodelling is required, and once that is achieved, non-specific and specific protease inhibitors control their activity. One example of nonspecific protease inhibitors is α-2 macroglobulin, typically found in blood and interstitial fluids. Of the specific MMP inhibitors, termed tissue inhibitors of metalloproteinases (TIMPs), at least four have been described. These proteins are typically produced and secreted by resident tissue cells, although some of them are found in plasma and in interstitial fluids.
The cornea is capable of producing at least four MMPs; gelatinase A, gelatinase B, collagenase-1 and stromelysin-1, although only gelatinase A has been isolated in the quiescent human and rodent cornea. Gelatinase A can degrade fibronectin, denatured collagens, and basement membrane. It is primarily present in the stroma, and because there is minimal stromal remodelling in normal cornea, it normally exists in its proenzyme form, associated with its inhibitor TIMP-2. In the event of injury, stromelysin, collagenase and gelatinase B are produced in the cornea. Stromelysin, which can degrade proteoglycans, can be detected in the cornea within 24 hours. Gelatinase B is maximally elevated early following injury, and it seems to be important for remodelling of damaged basement membrane, and its down-regulation is required for normal re-epithelialization. Collagenase levels increase 1 to 4 weeks after injury and persist for at least 7 to 9 months. This difference in timing of the expression of specific MMPs suggests that they have specific roles during the wound healing process.

Matrix metalloproteinases are thought to have significant roles in most corneal pathologies, including infectious keratitis. *P. aeruginosa*, for instance, produce MMPs that are inhibited by the MMP inhibitor galardin, and in *P. aeruginosa* keratitis, systemic administration of antibodies against MMP inhibitors significantly exacerbated the ulcers in mice. Thus, MMPs produced by *P. aeruginosa*, and neutralization of host inhibitors of MMPs, increase the virulence of *P. aeruginosa*. The development of new specific inhibitors of MMPs may prove to be beneficial in the treatment of infectious and non-infectious corneal ulcers and persistent epithelial defects.

A final group of proteases are those released during phagocytosis. These proteinases, namely cysteine proteinases and cathepsins, act primarily within lysosomes, and some of them, along with their inhibitors, have been identified within corneal cells, like cathepsin V, which is highly expressed in corneal epithelium. On the other hand, numerous acid hydrolases are produced by phagocytic neutrophils and monocytes. During their maturation within the bone, neutrophils produce a group of serine proteases, including leukocyte elastase and cathepsin G that are active against many extracellular components, and store them in cytoplasmic granules until released at target tissues. Monocytes, on the other hand, are capable of producing specific proteases tailored to the conditions of the site of inflammation. Both neutrophils and monocytes produce MMPs, and monocytes produce collagenase-1, stromelysin, gelatinease A, gelatinease B, and matrilysin, in addition to the unique MMP metalloelastase. In monocytes, these enzymes are not stored, but are newly synthesized at the site of inflammation as dictated by the surrounding inflammatory milieu. The role of lysosomal proteinases in ulcer formation in infectious keratitis is not fully understood. Although the presence of active cathepsin B, D, and L has been demonstrated...
in normal mouse corneal epithelium, and although their activities increase significantly during murine \textit{P. aeruginosa} microbial keratitis\textsuperscript{[422]}, the specific contribution of these proteases in corneal stromal loss has yet to be elucidated.

Table 1.5 Classes and types of matrix metalloproteinases (MMPs)

<table>
<thead>
<tr>
<th>MMP</th>
<th>Substrate</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Collagenases</strong></td>
<td></td>
</tr>
<tr>
<td>MMP-1</td>
<td>Native type I, II, III collagens</td>
</tr>
<tr>
<td>MMP-8</td>
<td></td>
</tr>
<tr>
<td>MMP-13</td>
<td></td>
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<tr>
<td>MMP-18</td>
<td></td>
</tr>
<tr>
<td><strong>Gelatinases</strong></td>
<td></td>
</tr>
<tr>
<td>MMP-2 (Gelatinease A)</td>
<td>Native type IV, V, VII collagens and gelatines</td>
</tr>
<tr>
<td>MMP-9 (Gelatinease B)</td>
<td></td>
</tr>
<tr>
<td><strong>Stomelysins</strong></td>
<td></td>
</tr>
<tr>
<td>MMP-3</td>
<td>Proteoglycans, fibronectin</td>
</tr>
<tr>
<td>MMP-10</td>
<td>Serine protease inhibitors</td>
</tr>
<tr>
<td>MMP-11</td>
<td></td>
</tr>
<tr>
<td><strong>Matrilysins</strong></td>
<td></td>
</tr>
<tr>
<td>MMP-7</td>
<td>Gelatines, fibronectin</td>
</tr>
<tr>
<td>MMP-12</td>
<td>Gelatines, fibronectin</td>
</tr>
<tr>
<td>MMP-19</td>
<td>Elastin</td>
</tr>
<tr>
<td>MMP-20</td>
<td></td>
</tr>
<tr>
<td><strong>Membrane-type MMPs</strong></td>
<td>Activation of progelatinease A</td>
</tr>
<tr>
<td>MMP-14</td>
<td></td>
</tr>
<tr>
<td>MMP-15</td>
<td></td>
</tr>
<tr>
<td>MMP-16</td>
<td></td>
</tr>
<tr>
<td>MMP-17</td>
<td></td>
</tr>
</tbody>
</table>

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1.6 Hypothesis and Objectives

1.6.1 Hypothesis

In microbial keratitis, corneal fibroblasts interact with *P. aeruginosa* bacteria and are involved in the early pathogen recognition and immune signalling that lead to recruitment of immune cells and the start of the immune cascades necessary for clearance of the offending pathogens.

1.6.2 Objectives

- To determine whether or not *P. aeruginosa* interact with human corneal fibroblasts.
- To examine the dynamics of host-pathogen interactions in microbial keratitis with respect to bacterial association and internalization to corneal fibroblasts.
- To determine the cytotoxic effects of *P. aeruginosa* on corneal fibroblasts.
- To assess the host cell response to infection with respect to matrix metalloproteinase and pro-inflammatory cytokine release by corneal fibroblast at different time points.
- To determine the role of the different bacterial virulence factors during host-pathogen interactions in human microbial keratitis.

We were aware of the different challenges of trying to design and implement this model, and we had to utilise and optimise cross-disciplinary experience of immunology, microbiology and clinical ophthalmology in order to examine such a complex pathology.
Chapter 2: MATERIALS and METHODS
2.1 General preparation of bacteria and host tissue

2.1.1 Bacteria

*Pseudomonas aeruginosa* wild type invasive strain PAO1 Holloway1C Stanier131 was obtained from NCIMB (Aberdeen, UK). Mutant *P. aeruginosa* strain PA14ΔpopB deficient in the PopB gene necessary for type III secretion system (TTSS), mutant strain PA14ΔflgK deficient in the FlgK gene necessary for normal flagellin and the parent wild type *P. aeruginosa* strain PA14 (PA14WT) were provided by Professor George O’Toole, University of Dartmouth, USA. Mutant green fluorescent protein (GFP) expressing *P. aeruginosa* PAO1 strain FliM tn7-GFP deficient in FliM gene necessary for normal flagella, mutant strain PilA tn7-GFP deficient in the PilA gene necessary for normal type IV pilus, mutant strain FliM PilA tn7-GFP deficient in both FliM and PilA gene and their parent strain PA tn7-GFP were acquired from Professor Tim Tolker-Nielsen, University of Copenhagen, Denmark.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAO1 Holloway1C Stanier131</td>
<td>NCIMB, Aberdeen, UK</td>
</tr>
<tr>
<td>PA14WT</td>
<td>Professor George O’Toole, University of Dartmouth, USA</td>
</tr>
<tr>
<td>PA14ΔflgK</td>
<td></td>
</tr>
<tr>
<td>PA14ΔpopB</td>
<td></td>
</tr>
<tr>
<td>PA tn7-GFP</td>
<td>Professor Tim Tolker-Nielsen, University of Copenhagen, Denmark</td>
</tr>
<tr>
<td>FliM tn7-GFP</td>
<td></td>
</tr>
<tr>
<td>PilA tn7-GFP</td>
<td></td>
</tr>
<tr>
<td>FliM PilA tn7-GFP</td>
<td></td>
</tr>
</tbody>
</table>

A master stock was made by adding a loopful of bacteria to sterile PBS to make a suspension, and this was stored in a liquid nitrogen tank. On the night before each experiment, PAO1 (Holloway1C Stanier131) and PA14 (WT, Δpopb and ΔflgK) were inoculated on nutrient agar (Oxoid, Basingstoke, UK) plates and grown over night, whereas PAO1 tnf-GFP (WT, FliM, PilA and FliM PilA) were grown on Luria-Bertani agar containing 60 µg/mL gentamicin (Appendix 7.1.1). Subsequent stocks were made and stored in a -80°C freezer using the Microbank system (Pro-Lab Diagnostics, Merseyside, UK). Inocula were prepared by suspending bacteria in corneal fibroblast infection medium (CFIM; Appendix 7.2.2).
2.1.2 Cell culture model

2.1.2.1 Isolation and culture of primary human keratocyte-derived corneal fibroblasts

Donor corneo-scleral rims remaining after corneal transplant procedures were obtained after the appropriate consent was taken from the donor. Five millilitres of corneal fibroblast culture medium (CFCM; Appendix 7.2.1) were placed into 30 ml universal container and 250 µl of 1 mg/ml collagenase solution in PBS was added to them. Under sterile conditions, the epithelium was scraped off the donor sample using a No. 15 scalp knife and the donor rim was divided into 4 equal sections. For each section, the transparent corneal tissue was cut out, placed into universal container containing CFCM and collagenase and incubated with gentle shaking for 3h at 37°C, after which the suspension was centrifuged at 1000 rpm (72 G) for 1 min. Under sterile conditions, the supernatant was carefully removed and discarded and the pellet was suspended in 7 ml of corneal culture medium and centrifuged at 2000 rpm (286 G) for 2 min. After repeating this process two more times, the pellet was suspended in 7 ml corneal culture medium, and the cell suspension was placed into 25 ml tissue culture flask. The flask was then incubated for 24 h at 37°C, 5% (v/v) CO₂, checking cells for adherence and removing dead cells and replacing the culture medium under sterile conditions. The cells were incubated until the cells were >90% confluent, replacing culture medium every 3 days.

2.1.2.2 Expansion of corneal fibroblasts:

Once the cells were confluent, the medium was removed from the flask and 10 ml of sterile PBS was added to the flask, gently rocking, until the cells were thoroughly rinsed, after which the PBS was removed and replaced with 5 ml of Trypsin/EDTA solution (Sigma-Aldrich, Dorset, UK). The flask was incubated at 37°C for 5 minutes, and then observed under a dissecting microscope until the complete dissociation of cells. Next, 7 ml of corneal culture medium were added to make a cell suspension, and the solution was centrifuged at 1000 rpm (72 G) for 4 minutes. The supernatant was discarded and the pellet was suspended in 9 ml of fresh medium. In a new flask, 3 ml of cell suspension were added to 7 ml of fresh culture medium and the flask was incubated for 24 h, after which adherence was checked under a dissecting microscope. Cells passaged between 3 and 6 times (P3-P6) were used in order to limit differentiation, as we noticed in our pilot experiments that cells at P7 or higher may be morphologically different. All cell culture preparation was done in a Class II safety cabinet.
2.1.2.3 Polymerase chain reaction (PCR) for detection of *Mycoplasma*

Corneal fibroblast cell cultures were tested for *Mycoplasma* contamination. Samples were prepared by taking 1 ml of culture medium from a confluent cell culture and centrifuging at 500g for 5 min. The supernatant was transferred to a fresh 1.5 ml Eppendorf tube and centrifuged at 15000g for 15 min, then the supernatant was decanted and the pellet suspended in 100 µl of fresh culture medium. For PCR setup, the PCR *Mycoplasma* test kit I/C (PromoCell, Heidelberg, Germany) was used. In each experiment, the kit negative and positive controls were included. For the sample mix, 2 µl of suspended pellet samples, 22 µl of rehydration buffer and 1 µl of DNA polymerase were added to dehydrated PCR tubes. A PCR programme was established according to the manufacturer’s recommended settings (first cycle: 94°C for 2 min; then 38 cycles: 94°C for 30 s, 55°C for 30 s, 72°C for 40 s, then final cooling to 4-8°C) and the reaction was run on a T3 thermal cycler (Biometra, Goettingen, Germany). After PCR, 10 µl of each control was run on a 2% (w/v) agarose gel (Appendix7.4.1) using a 5 mm combo, with electrophoresis at 100V for 40 min. The internal control band (479 base pairs) and the positive reaction band (265-278 base pairs) were visualised using a 2UV Transilluminator (BioDoc-It Systems, Upland, USA).

2.1.3 Whole tissue model

Donor corneal buttons left over from Descemet’s stripping endothelial keratoplasty (DSEK) were collected after appropriate ethical approvals were obtained. Donor information from the eye bank was checked to ensure that they agreed for their donated corneas to be utilized for research purposes as well as transplant procedures. The buttons were transferred from eye theatres to our laboratories in the storage medium provided by the eye bank and stored at 4°C until used in the infection experiments, within 72h of collection. The average age of donors was 78.8 years (range 65-88) with causes of death excluding those that would potentially affect our infection experiments (e.g. septicaemia).
2.2 Challenge of human corneal tissue with bacteria

2.2.1 Preparation of bacteria

An ampoule containing *P. aeruginosa* bacteria was transferred from the Microbank system to a Class II safety cabinet. Without allowing contents to thaw, a sterile plastic loop was used to capture one of the bacteria-laden pellets and transfer it to an agar plate, where the pellet was gently moved across the surface of the agar to spread the inoculum, after which the plate was incubated overnight in an incubator at 37°C, 5% (v/v) CO₂. On the following day, a bacterial suspension was prepared by suspending bacteria, picked from the agar plate using a sterile loop, in 1-2 ml of sterile PBS. In order to prepare suspensions with known concentrations of bacteria, the optical density (OD) of an initial bacterial suspension was measured in 1% (w/v) SDS/0.1% (w/v) NaOH solution at a wavelength 260 nm using a Hitachi U-1100 spectrophotometer (Hitachi, Tokyo, Japan) in UV transparent cuvettes (Starstedt, Nümbrecht, Germany). The optical density of the suspension was used to calculate the number of bacteria in the suspension and 10 fold serial dilutions were made by adding 100 µl of bacterial suspension to 900 µl sterile PBS, until the desired infecting doses were reached. Viable counting was performed by spreading 25 µl of the final bacterial suspension on the surface of the agar (adjusting for the total volume in the final calculation of bacterial counts), and was used to confirm the number of bacteria in the suspensions.

2.2.2 Infection of corneal donor buttons

In most cases, we used two donor buttons for our experiments. On the day of the experiment, the buttons were washed 4 times in warm sterile PBS, and one button was scarified by a 27 gauge needle to produce 4-5 linear epithelial scores that were deep enough to bypass the epithelium and score the underlying stroma. The buttons were then both bisected to yield 4 halves, and these were placed in a 24-well cell culture plate.

Bacterial suspensions were prepared to an infective dose of 10⁷ CFU/mL (as above), and 1 mL was added to each tissue-containing well. The plates were incubated in an atmosphere of 5% (v/v) CO₂ at 37°C for 3 or 9h. At these time points, the buttons were washed 4 times in warm sterile PBS and transferred to a container containing a fixative solution [3% (v/v) glutaraldehyde and 4% (v/v) formaldehyde in 0.1 M PIPES (1,4-Piperazinediethanesulfonic acid) buffer pH 7.2], fixed for a minimum of 1 hour, then processed for scanning or transmission electron microscopy, according to experiment. (See section2.3.2 for the methodology of electron microscopy).
2.2.3 Infection of corneal fibroblasts

Corneal fibroblasts were cultured to confluence in 24-well cell culture plates (Greiner Bio-One, Solingen, Germany). Cell quantification was carried out using the improved Neubauer cell counting chamber and trypan blue stain. The average cell count in a 24-well plate was $9.7 \times 10^5$/well ($n=4$ measurements). Before bacterial challenge, the CFCM in the wells was removed and the cell monolayers were gently rinsed 4 times in warm sterile phosphate buffered saline (PBS; Sigma-Aldrich, Dorset, UK) to remove any residual antibiotics.

Confluent corneal fibroblast monolayers were challenged in triplicate wells with bacterial suspension prepared by adding the bacteria to CFIM in 1ml volumes/well. Bacterial association was assessed at 3, 6, 9 and 24 hours. The concentration range of bacteria initially used was from low ($10^1$ and $10^2$ CFU/mL) to intermediate ($10^4$ and $10^6$ CFU/mL) to high ($10^8$ CFU/mL) and the corresponding multiplicity of infection (MOI) is shown in Table 2.1. Preliminary infection experiments demonstrated that high bacterial concentrations (>10⁶ CFU/mL) led to an almost total destruction of the monolayers. Consequently, only bacterial concentrations of $10^1$, $10^2$, $10^3$ and $10^4$ were used.

Table 2.1 Bacterial inocula and corresponding multiplicity of infection (MOI)

<table>
<thead>
<tr>
<th>Bacterial inoculum (CFU/mL)</th>
<th>Multiplicity of infection (bacteria/cell)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$10^1$</td>
<td>0.0001</td>
</tr>
<tr>
<td>$10^2$</td>
<td>0.001</td>
</tr>
<tr>
<td>$10^3$</td>
<td>0.1</td>
</tr>
<tr>
<td>$10^4$</td>
<td>1</td>
</tr>
<tr>
<td>$10^5$</td>
<td>10</td>
</tr>
<tr>
<td>$10^6$</td>
<td>100</td>
</tr>
<tr>
<td>$10^7$</td>
<td>1000</td>
</tr>
</tbody>
</table>
2.3 Measurement of total bacterial association

2.3.1 Viable counting assays

Corneal fibroblast monolayers were incubated for up to 24h and at each time point examined, the plate was transferred to the cabinet, the infection medium was removed and the cells were gently rinsed 4 times with warm sterile PBS to remove all non-associated bacteria. For the 9h and 24h time points, the infection medium was transferred to 1.5 ml Eppendorf tubes, centrifuged at 2200xg for 6 min, and the supernatants stored at -20°C for cytokine analysis (section 2.6.1). At each time point, fibroblasts were lysed by adding 250 µl of a saponin lysis solution (Appendix 7.3.3) to each well and incubating the plate for 15-20 minutes. Complete cell lysis was confirmed by observation under a light microscope, and a sterile plastic tip was used to mechanically aid in lysing the cells if necessary. Bacteria were quantified by viable counting, diluting the lysates in sterile PBS, and appropriate dilutions were inoculated onto nutrient agar plates (25 µl/plate) in triplicate and incubated overnight in an atmosphere of 5% (v/v) CO₂ at 37°C. Colony counts were quantified using a ProtoCOL model 60000 automated colony counter (Synoptics Ltd, Cambridge, UK).

2.3.2 Scanning Electron Microscopy (SEM)

Human corneal fibroblasts were grown to confluence on manufacturer collagen pre-coated transwells (ThinCerts™, transparent pore size 0.4 urn; Greiner Bio-one, Stonehouse, UK), challenged with a bacterial infective dose of 10⁴ and 10⁷ CFU/mL (MOI 0.1 and 100, respectively) and incubated in an atmosphere of 5% (v/v) CO₂ at 37°C for 3, 6 and 9h. Uninfected cells were included as negative controls. At each time point, the cells were washed 4 times with warm sterile PBS and then processed for SEM. Transwells were fixed for a minimum of 1h using a fixative solution [3% (v/v) glutaraldehyde and 4% (v/v) formaldehyde in 0.1 M PIPES (1,4-Piperazinediethanesulfonic acid) buffer pH 7.2] and were rinsed twice in buffer solution (0.1 M PIPES buffer) for 10min each. Then, a gradual concentration of ethanol [from 30%, 50%, 70%, 95% and up to 100% (v/v)] was used to dehydrate the samples (10min for each step and 20min for the final step). Cutting of the transwell membranes was done during the ethanol dehydration steps. Next, the cut membranes were critical-point dried and mounted onto metal stubs for the final sputter coating step. Coated specimens were viewed using the Quanta 200 scanning electron microscope (FEI, Hillsboro, USA) and several images were obtained for each specimen at magnifications of x3200 and up to x24000.
2.3.3 Assessment of the role of TFP and flagella in bacterial association to fibroblasts

In order to investigate the role of bacterial flagella (flg) and type IV pilus (TFP) in bacterial association to corneal fibroblasts, confluent fibroblast monolayers were challenged in triplicate with $10^7$ CFU/mL (MOI 100) CFIM and incubated in an atmosphere of 5% (v/v) CO$_2$ at 37°C for 3h. One triplicate was challenged with wild type PAO1 bacteria (PA tn7-GFP), another with mutant PAO1 strain FliM tn7-GFP deficient in flg, a third with mutant PAO1 strain PilA tn7-GFP deficient in TFP and a forth with mutant PAO1 strain FliM PilA tn7-GFP deficient in both flg and TFP (see section 2.1.1 for more details about bacterial strains). Viable counting assays were performed as described in section 2.3.1.

2.3.4 Laser scanning confocal microscopy

The bacterial strains used in this set of experiments were green fluorescent protein (GFP) expressing PAO1 strains (section 2.1.1), which allowed for direct visualization of bacterial association to corneal fibroblasts. Confluent fibroblast monolayers were challenged in triplicate with $10^7$ CFU/mL (MOI 100) CFIM and incubated in an atmosphere of 5% (v/v) CO$_2$ at 37°C for 3h. At the given time point (3h), the cell monolayers were washed gently 4 times with warm PBS, and then fixed with paraformaldehyde 4% (w/v) in PBS for 30 min. The cells were washed with 50 mM ammonium bicarbonate, and then washed with warm PBS. Evans Blue 0.0025% (v/v) in PBS was then added to the cells and left for 20 min at 37°C. The cell monolayers were washed thoroughly to remove all residual fixative, and were examined immediately after staining using the Leica TCS SPS on the DM6000 inverted microscope stand. The field of view was 246 $\mu$m$^3$, using a HCX PlanoApo CS 63x 1.3 NA glycerol emersion objective lens=63x. The excitation laser had a wavelength of 488 and 561 nm, and the detection laser had wavelength ranges of 500-555 nm and 587-750 nm.
2.4 Measurement of bacterial invasion

2.4.1 Gentamicin protection assays

Confluent corneal fibroblast monolayers were challenged in triplicate with bacterial suspensions of 10^2 (MOI 0.001) and 10^4 CFU/mL (MOI 0.1) and incubated in an atmosphere of 5% (v/v) CO_2 at 37°C for 9h. A higher bacterial infective dose (10^7 CFU/mL; MOI 100) was also used to challenge cell monolayers for a shorter time period (3h). In order to inhibit intracellular actin microfilament activity, monolayers were first pre-incubated with CFIM containing 2 µg/ml cytochalasin D (Appendix 7.3.2) for 30 min before adding the bacterial suspensions. At given time points, the cell monolayers were washed gently 4 times with warm PBS and then 1ml of CFIM containing 200 µg/ml of gentamicin (Sigma-Aldrich, Dorset, UK) was added to the wells and incubated at 37°C for 90min. This was sufficient to destroy all extracellular bacteria (confirmed by sampling the medium and viable counting after incubating on nutrient agar, ensuring >99% bacterial killing) and allowed for quantification of internalized bacteria. Monolayers were then washed 4 times with warm PBS and lysed using saponin, and bacteria were quantified by viable counting on nutrient agar plates, as described above.

2.4.2 Transmission Electron Microscopy (TEM)

Human corneal fibroblasts were grown to confluence on collagen pre-coated transwells (ThinCerts™, transparent pore size 0.4 urn; Greiner Bio-one, Stonehouse, UK), challenged with a bacterial infective dose of 10^7 CFU/mL (MOI 100) and incubated in an atmosphere of 5% (v/v) CO_2 at 37°C for 9h. Uninfected cells were used as negative controls. At the time point, the monolayers were washed 4 times with warm sterile PBS and then processed for TEM. Transwells were fixed for a minimum of 1h using a fixative solution [3% (v/v) glutaraldehyde and 4% (v/v) formaldehyde in 0.1 M PIPES (1,4-Piperazinediethanesulfonic acid) buffer pH 7.2], rinsed twice in buffer solution (0.1 M PIPES buffer) for 10min each and then fixed using post fixative [1% (w/v) osmium tetroxide in 0.1 M PIPES buffer, pH 7.2] for 1hr. Samples were briefly rinsed twice in PBS, and then a 2% (w/v) uranyl acetate solution was used for staining for 30min. A series of gradual concentrations of ethanol [from 30%, 50%, 70%, 95% and up to 100% (v/v)] was used to dehydrate the samples (10min for each step and 20min for the final step). Samples were incubated overnight, on a rotor (2rpm), in spur resin at a 50:50 ratio with acetonitrile. After that, the samples were placed in fresh spur resin and polymerised in an electric oven at 60°C for 20-24h. In preparation for microscopy, sample blocks were cut into ultrathin sections (about 90 nm thick) using glass knives on a Leica Reichert Ultracut E microtome (Leica, Wetzlar, Germany). A number of sections was stained with toluidine blue and checked under
light microscopy for the presence and positioning of cells, and then, when satisfactory, several sections per sample were fixed onto copper grids. Grids were stained using lead nitrate for 5min in the presence of NaOH capsules to absorb excess CO$_2$. Then, grids were briefly rinsed in distilled water and air dried. Sample viewing was done using the H7000 transmission electron microscope (Hitachi, Japan) at 75Kv and under different magnifications ranging from x3000 to x20000.

2.4.3 Assessment of the effect of ciprofloxacin on intracellular bacteria

Confluent fibroblast monolayers were challenged with PAO1 bacteria at a dose of $10^7$ CFU/mL (MOI 100) for 3h, and then gently washed 4 times in warm sterile PBS. CFIM (1ml) containing 200 µg/ml of gentamicin was added to designated wells and ciprofloxacin (Sigma-Aldrich, Dorset, UK) at doses of 10, 50 and 200 µg/mL were added to separate wells and incubated at 37°C for 90min. Monolayers were then washed 4 times with warm PBS and lysed using saponin, and bacteria were quantified by viable counting on nutrient agar plates, as described above.

2.4.4 Measurement of bacterial survival and replication inside invaded fibroblasts

Confluent corneal fibroblast monolayers were challenged in triplicate with bacterial suspensions of $10^7$ CFU/mL (MOI 100) and incubated in an atmosphere of 5% (v/v) CO$_2$ at 37ºC for 24h. Infected CFIM was removed from the wells at 3h post challenge and the cell monolayers were washed gently 4 times with warm sterile PBS, after which 1ml of CFIM containing 200 µg/ml of gentamicin was added to the wells and the plates were returned to the incubator. At 24h, fibroblast monolayers were washed 4 times with warm PBS and lysed using saponin, and bacteria were quantified by viable counting on nutrient agar plates, as described above. Sterility was confirmed by sampling the extracellular medium and viable counting after incubating on nutrient agar overnight. Bacterial invasion was assessed at 90 min, 3h, 6h and 21h after adding the gentamicin using the method described above.
2.4.5  Assessment of the role of the SRC kinase system in bacterial internalization

To investigate the role of SRC tyrosine kinase in bacterial internalization, two sets of experiments were performed: a gentamicin protection assay to determine the direct effect of tyrosine kinase inhibitors on the number of internalized bacteria into corneal fibroblasts, and SDS-PAGE and western blot to detect the expression and upregulation of the enzymes during live infection.

2.4.5.1  Gentamicin protection assays

Corneal fibroblasts (CF) were cultured to confluence in 24-well cell culture plates, and triplicate monolayers were then pre-treated for 30 min with the tyrosine kinase inhibitors Genistein (GST – 10 µg/mL) and PP2 (3 µg/mL) and with the actin microfilament inhibitor cytochalasin D (CD – 10 µg/mL), one triplicate per inhibitor. One triplicate was not pre-treated with any inhibitors and was used as control. Infective medium was prepared by adding inhibitors to CFIM in the same concentration used for pre-treating the fibroblasts so that each pre-treated triplicate was infected by CFIM with the same inhibitor originally used (for example, monolayers pre-treated with 10 µg/mL CD were infected with CFIM containing 10 µg/mL CD).

Fibroblasts monolayers were challenged in triplicate with $10^7$ CFU/mL (MOI 100) CFIM and incubated in an atmosphere of 5% (v/v) CO$_2$ at 37°C for 3h. To determine the effect of the inhibitors on the rate of bacterial growth, CFIM containing inhibitors was added to wells containing no fibroblasts, and the triplicates were incubated with the infected fibroblasts and treated in a similar manner.

At time point (3h), bacterial internalization was assessed as described in section 2.4.1. For the triplicate wells containing CFIM with inhibitors and no fibroblasts, bacteria were quantified by viable counting on nutrient agar plates immediately after the 3h incubation (Figure 2.1).

![Figure 2.1 Corneal fibroblast pre-treatment and infection using SRC kinase inhibitors](image-url)
2.4.5.2 Sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE)

In order to assess the presence and upregulation of SRC tyrosine kinase, SDS-PAGE with a 12% (w/v) acrylamide gel was performed in a second-dimension mini format electrophoresis system (Mini-PROTEAN; Bio-Rad, Hercules, CA, USA). Confluent corneal fibroblasts were challenged with $10^7$ CFU/mL (MOI 100) CFIM and incubated in an atmosphere of 5% (v/v) CO$_2$ at 37°C for 3h. At time point (3h), cell monolayers were washed 4 times with warm PBS, and then 100 µL of CytoBuster™ Protein Extraction Reagent (Fisher, Loughborough, UK) was added for 5 min, and the wells were mechanically scrapped by a plastic tip to ensure complete lysis of cells. The supernatants were stored in a -20°C freezer.

A. Preparation of the gels

A separating gel (Appendix 7.7.1) was prepared between the 2 glass panes of the system after placing on a rubber spacer and fixing by clips to avoid leakage. After the separating gel was set, a stacking gel (Appendix 7.7.2) was introduced using a plastic liquid transfer pipette, a comb was placed on top to form the wells and the stacking gel was left to set for 1h at room temperature.

B. Determination of protein concentration

In order to determine and standardize the protein concentration in culture supernatant samples, a Pierce™ BCA Protein Assay Kit (Life Technologies, Paisley, UK) was used (Appendix 7.7.7). In this assay, the presence of proteins reduces Cu$^{2+}$ to Cu$^{1+}$ in an alkaline solution, and one molecule of the resultant Cu$^{1+}$ chelates two molecules of bicinchoninic acid (BCA), resulting in an intense purple-coloured reaction product. Because of the highly sensitive and selective nature of the reaction, protein quantity can be determined by colorimetric detection.

Reagent A and Reagent B were mixed in a ratio of 50:1 to form BCA reagent, of which, a 200 µl aliquot was added to 25 µl of albumin standard (Appendix 07.7.8) or undiluted culture supernatant samples in a Nunc™ 96-well flat-bottomed polystyrene microtitre plate (Thermo-Scientific, Rockford, IL, USA). The plate was then incubated at 37°C for 30 min, and the absorbance at 570 nm was read in an iMark™ microplate reader (Bio-Rad, Hercules, CA, USA). The absorbance value of the albumin standards was plotted as a standard curve and the protein concentration of the samples was calculated by comparing the absorbance value with the standard curve.
C. Preparation and loading of the samples

After determining the protein concentration in each sample, all samples were diluted with distilled water so that they had similar protein concentrations. Twenty-five microliters of sample buffer (Appendix 7.7.3) were added to 100 µl of the adjusted samples, and the samples were boiled for 8 minutes. After carefully removing the comb and un polymerized gel, the gels were placed into the electrophoresis system, and the tank was filled with running buffer (Appendix 7.7.4). SeeBlue® Plus2 Pre-stained Protein Standard (Life Technologies, Paisley, UK) (7 µl ) and 20 µl of the boiled samples were added to the wells, and the gels were left to run at 170 V for 90 min or until all the bands have reached the bottom of the gels.

2.4.5.3 Western blotting

After the electrophoresis was finished, the SDS-PAGE gels containing the separated protein samples were immersed in transfer buffer (Appendix 07.7.5) and left on a gentle shaker for 15 min. For each gel, 6 layers of thick filter paper (3M, Bracknell, UK) and a piece of Amersham Hybond P 0.45 PVDF blotting membrane (GE Healthcare, Hatfield, UK) were cut to the same size as the gels and submerged in methanol for 5 min. A blotting sandwich was made by initially laying 3 layers of the filter paper on the bottom, followed by the blotting membrane, followed by the gel, and finally, the remaining 3 layers of filter paper. The sandwich was placed in a Trans-Blot SD semi-dry transfer cell (Bio-Rad, Hercules, CA, USA) at 10 V for 45 min. The membranes were then immersed in a blocking buffer (Appendix 7.7.6) and left on a gentle shaker for 1h, after which the buffer was replaced by 10 mL blocking buffer containing 18 µl anti-SRC antibody (Abcam, Cambridge, UK) and left, tightly covered, on a gentle shaker at room temperature overnight. After incubation, the membranes were washed 3x in PBS-Tween (Appendix 7.8.1), immersed in 10 mL PBS-Tween containing 4 µl goat anti-rabbit IgG HRP antibody, and left on a gentle shaker for 1h. The membranes were then washed 3x in PBS-Tween and read using a VersaDoc™MP 4000 imager (Bio-Rad, Hercules, CA, USA). A developing buffer was prepared using a Novex® ECL Chemiluminescent Substrate Reagent kit (Life Technologies, Paisley, UK) and a few drops of it were instilled on top of the membranes prior to reading.
2.5 Assessment of bacteria-induced cytotoxicity

2.5.1 Assessment of fibroblast cytotoxicity associated with live infection

Human corneal fibroblasts were grown to confluence in 96-well tissue culture plates (Greiner Bio-one, Stonehouse, UK). Cell quantification was carried out and the average count was $2.1 \times 10^4$ cells/well (n=4 measurements). Confluent monolayers were maintained in CFCM. Bacterial inocula were prepared in CFIM at similar MOIs to those of previous infection experiments (see Table 2.1). The monolayers were gently rinsed with warm sterile PBS and then 200 µl/well of infective CFIM was added to triplicate wells. Uninfected control wells were also included for measurements of spontaneous LDH release and maximum LDH release by lysed uninfected cells as well as blank wells, containing only CFIM, for background measurements. Plates were incubated in an atmosphere of 5% (v/v) CO$_2$ at 37°C for 9h. LDH measurements were carried out using the CytoTox 96® Non-Radioactive Cytotoxicity assay kit (Promega, Madison, WI, USA). The proprietary lysis solution (20 µl/well) was added to maximum LDH release wells 45min prior to the final time point to completely lyse the fibroblasts. Supernatants from each well were transferred to individual 1.5ml Eppendorf tubes and centrifuged at 2200g for 6 min to pellet the bacteria and dead cells. Next, 50 µl of supernatant from each tube was transferred to a 96-well measuring plate (Camlab, Cambridge, UK) to which 50 µl of the proprietary substrate mixture was added, followed by incubation in the dark at room temperature for 30min. A stop solution (50 µl/well) was added to all wells and the absorbance was read at 492nm using an iMark Absorbance Reader (Bio-Rad, USA). Cytotoxicity levels were calculated by dividing the average LDH release value of test sample by the average maximum LDH release value from lysed, uninfected cells using the following formula:

\[
LDH \text{ release of test (\%)} = \frac{LDH \text{ release of test}}{maximum \text{ LDH release}} \times 100
\]

2.5.2 Assessment of the effect of intracellular bacteria on fibroblast viability

To investigate the effect of intracellular bacteria on fibroblast viability, corneal fibroblasts were grown to confluence in a 24-well plate, and monolayers were challenged in triplicate with bacterial suspensions of $10^7$ CFU/mL (MOI 100) and incubated in an atmosphere of 5% (v/v) CO$_2$ at 37°C for 9 and 24h. At 3h post challenge, infected and uninfected monolayers (n=3) were washed 4 times in warm PBS, and 1 mL of CFIM containing 200 µg/ml of gentamicin was added to each well and left for the remainder of the 9 or 24h period. LDH measurements were carried out at 9 and 24h as described above but with the addition of 100 µl/well of the proprietary lysis solution to maximum LDH release wells, 45min prior to the final time point (to compensate for the larger number of cells and the larger volume of the wells).
2.5.3 Assessment of the effect of bacterial virulence on cytotoxicity

In order to assess the impact of bacterial virulence on corneal fibroblast cytotoxicity, corneal fibroblasts were challenged with two different strains of *P. aeruginosa*: the ‘invasive’ strain PAO1 and the ‘cytotoxic’ strain PA14. Confluent fibroblast monolayers were challenged in triplicate with $10^7$ CFU/mL (MOI 100) CFIM of each strain and incubated in an atmosphere of 5% (v/v) CO$_2$ at 37°C for 9h. LDH measurement was carried out at 9h as described in section 2.5.

2.5.4 Assessment of the role of bacterial virulence factors on cytotoxicity

In order to investigate the role of bacterial flagella (flg) and type III secretion system (TTSS) in bacterial cytotoxic effects on corneal fibroblasts, an LDH cytotoxicity assay was carried out. Corneal fibroblasts were cultured to confluence in 24-well cell culture plates, and then challenged in triplicate with $10^7$ CFU/mL (MOI 100) CFIM and incubated in an atmosphere of 5% (v/v) CO$_2$ at 37°C for 9h. Triplicate wells were challenged with wild type PA14 bacteria (PA14WT), mutant PA14 bacteria deficient in TTSS (PA14ΔPopb) and mutant PA14 bacteria deficient in flagella (PA14ΔFlgK) (see section 2.1.1 for bacterial strains). LDH measurement was carried out at 9h as described in section 2.5.1.
2.6  Assessment of host cell response to bacterial infection

2.6.1  Assessment of corneal fibroblast cytokine release associated with live infection

2.6.1.1  Detection of cytokine release by a sandwich immunoassay technique

Confluent corneal fibroblast monolayers were challenged in triplicate with a bacterial infective dose of \(10^1, 10^2, 10^3\) and \(10^4\) CFU/mL (MOI 0.0001, 0.001, 0.01, 0.1, respectively). A subset of fibroblast monolayers was infected with a higher infective dose of bacteria \(10^7\) CFU/mL; MOI 100) and treated with gentamicin 3h post-challenge to destroy all extracellular bacteria (as described above). This allowed for the study of the effect of internalized bacteria on cytokine production with the preservation of fibroblasts until the end of the 9 or 24h time period. The plates were incubated in an atmosphere of 5% (v/v) CO\(_2\) at 37°C. Supernatants were collected at 9 and 24h post-challenge and centrifuged at 2200g for 6 min, and the samples were stored at -20°C. Cytokine analyses were carried out using the Human Pro-Inflammatory 9-Plex Tissue Culture Kit (MSD, Rockville, MD, USA) which employs a sandwich immunosassay format where capture antibodies are coated in a patterned array on the bottom of the wells of the plate. Initially, 150 µL of the 1% (w/v) Proprietary Blocker B solution were dispensed into each well, and the plate was incubated with vigorous shaking (300-1000 rpm) for 1 hour at room temperature. Thereafter, calibrator solutions were prepared (Appendix 7.5.1) and 25 µL were added per duplicate wells to the first 16 wells. In the remaining wells, 25 µL/well of each undiluted sample was added. The plate was sealed with an adhesive plate seal and incubated with vigorous shaking (300-1000 rpm) for a further 2h at room temperature, after which all wells were washed 3X with PBS + 0.05% (v/v) Tween-20. Thereafter, 25 µL of detection antibody solution (Appendix 7.5.1.2) were added to all wells, and the plate was sealed again with an adhesive plate seal and incubated for 1 to 2 hours with vigorous shaking (300-1000 rpm) at room temperature. The plate was then washed 3x in PBS, after which 150 µL of 2X Read Buffer T (Appendix7.5.1.3) were added, and the plate was immediately analysed using the SECTOR® Imager (MSD, Rockville, MD, USA).
2.6.1.2 Assessment of the effect of bacterial virulence on cytokine production

Corneal fibroblasts were challenged with *P. aeruginosa* PA14 strain and with PAO1 strain. Confluent fibroblast monolayers were challenged in triplicate with $10^7$ CFU/mL (MOI 100) CFIM of each strain and incubated in an atmosphere of 5% (v/v) CO$_2$ at 37°C for 24h. Supernatants were collected at 9 and 24h post challenge and spun at 2200xg for 6 min, and the samples were stored at -20°C. Cytokine analysis was carried out using the V-PLEX Human IL-1β Kit (MSD, Rockville, MD, USA) which employs a sandwich immunoassay format where capture antibodies are coated on independent and well-defined spots on the bottom of the wells of the plate. Initially, calibrator solutions were prepared (Appendix 7.5.2.1) and 50 µL/well were added in duplicate to the first 16 wells. In the remaining wells, 25 µL/well of the proprietary Diluent 2 and 25 µL/well of each undiluted sample were added. The plate was sealed with an adhesive plate seal and incubated with vigorous shaking (300-1000 rpm) for a further 2h at room temperature, after which all wells were washed 3X with PBS-Tween (Appendix 7.8.1). Thereafter, 25 µL of detection antibody solution (Appendix 7.5.1.27.5.2.2) were added to all wells, and the plate was sealed again with an adhesive plate seal and incubated for 1 to 2 hours with vigorous shaking (300-1000 rpm) at room temperature. The plate was then washed 3x in PBS-Tween, after which 150 µL of Read Buffer T (Appendix 7.5.1.3) were added, and the plate was immediately analysed using the SECTOR® Imager (MSD, Rockville, MD, USA).

2.6.1.3 Assessment of the role of bacterial virulence factors on cytokine production

Corneal fibroblasts were cultured to confluence in 24-well cell culture plates, and then challenged in triplicate with $10^7$ CFU/mL (MOI 100) CFIM and incubated in an atmosphere of 5% (v/v) CO2 at 37oC for 9h. Similar to section 2.5.3, fibroblast triplicates were challenged with PA14WT, PA14ΔPopb and PA14ΔFlgK, and the cytokine analysis was carried out using the V-PLEX Human IL-1β Kit (MSD, Rockville, MD, USA) as described in section 0.

2.6.1.4 Assessment of the role of bacterial components on cytokine production

Confluent fibroblast monolayers were challenged in triplicates with live bacteria, heat killed bacteria, freeze-thaw killed bacteria and LPS, and controls included uninfected cells and lysed uninfected cells (see section 2.6.2.1). Culture supernatants were collected at 9 and 24h, and cytokine analysis was carried out using the V-PLEX Human IL-1β Kit (MSD, Rockville, MD, USA) as described in section 2.6.1.1.
2.6.2 Assessment of corneal fibroblast MMP release associated with live infection

2.6.2.1 Challenge of corneal fibroblast

Confluent corneal fibroblast monolayers were challenged in triplicate with live bacteria, heat killed (HK) bacteria and freeze-thaw (FT) killed bacteria, all with an infective dose equivalent to $10^7$ CFU/mL. In addition, triplicate cell monolayers were challenged with 100 ng/mL of pure *Pseudomonas aeruginosa* lipopolysaccharide (LPS), and another set of monolayers was challenged with $10^7$ CFU/mL (MOI 100) of live bacteria and then treated with 200 µg/mL gentamicin 3h post challenge. Negative controls included unchallenged cell monolayers and cells challenged only with 200 µg/mL gentamicin added 3h after start of experiment. Positive controls included live bacteria at a dose of $10^7$ CFU/mL, suspended in medium with no fibroblasts; and fibroblast monolayers lysed with 100 µL of proprietary lysis solution (Promega, Madison, WI, USA) 45 minutes prior to the desired time point. Serum free CFIM was used in all wells.

Heat killed bacteria were prepared by heating a stock solution of $10^8$ CFU/mL bacteria for 1h at 60°C using a water bath. The stock solution was then stored at 2-8°C, and a diluent of $10^7$ CFU/mL (MOI 100) was used on the day of each experiment. Freeze-thaw killed bacteria were prepared by freezing a stock solution of $10^9$ CFU/mL bacteria in a -80°C freezer overnight, then thawing the bacteria to 37°C in a water bath and repeating the freeze-thaw cycle until all bacteria were killed. Sterility of both heat-killed and freeze-thawed samples was verified by incubating a sample of each stock solution on nutrient agar plates overnight; this was repeated for freeze-thawing cycles until no bacterial growth was observed on the agar plates (8 cycles in total).

The plates were incubated in an atmosphere of 5% (v/v) CO$_2$ at 37°C. Supernatants were collected at 9 and 24h post challenge and centrifuged at 2200g for 6 min, and the samples were stored at -20°C. Analyses were done within 2-4 weeks of collecting the supernatants using a sandwich immunoassay technique and a standard gelatine zymography.
2.6.2.2 Gelatin zymography

Culture supernatants were thawed on ice to room temperature, and 15 µL of each sample was added to 15µL of zymogram sample buffer (Appendix 7.6.1). Precast 10% (w/v) polyacrylamide gelatine 10-well gels (Bio-Rad, Hercules, CA, USA) were placed into a second-dimension mini format electrophoresis system (Mini-PROTEAN; Bio-Rad, Hercules, CA, USA), and the electrophoresis tank was filled with a zymogram running buffer (Appendix 7.6.2). On each gel, the first two wells were used for the protein ladder (Thermo-Scientific, Rockford, IL, USA) and for the MMP positive control (Sigma-Aldrich, Dorset, UK). No samples were added to the third well, and 10 µL of each sample was added to each of the remaining wells. The gels were left to run at 90 V for 2h or until all the bands have reached the bottom of the gels. The gels were then washed four times with a zymogram renaturation buffer (Bio-Rad, Hercules, CA, USA), gently shaking for 15 min. The gels were transferred to separate containers and washed once with a zymogram development buffer (Bio-Rad, Hercules, CA, USA), gently shaking for 30 min, and then incubated at 37°C for 40 h in development buffer containing 1% (w/v) sodium azide. After incubation, the gels were washed three times in distilled water, soaked in a zymogram staining buffer (Appendix 7.6.3) for 1h, and washed several times in a zymogram destaining buffer (Appendix 7.6.4), with gently shaking, until satisfactory destaining was achieved.

2.6.2.3 Detection of MMP release by a sandwich immunoassay technique

In order to detect the release of MMP-1, MMP-2, MMP-3 and MMP-9, the Human MMP-2 Ultra-Sensitive kit and the Human MMP 3-Plex Ultra-Sensitive kit (MSD, Rockville, MD, USA) were used. The method for both assays is identical, using the specific detection antibody for each kit.

Initially, 25 µL/well of the proprietary Diluent 2 were added to each well, and the plate was sealed with an adhesive plate seal and incubated with vigorous shaking (300-1000 rpm) for 30 min at room temperature. Calibrator solutions were prepared (Appendix 7.5.3.1) and 25 µL/well were added in duplicate to the first 16 wells, and, 25 µL/well of each undiluted sample were added to the remaining wells. The plate was sealed with an adhesive plate seal and incubated with vigorous shaking (300-1000 rpm) for 2h at room temperature, after which all wells were washed 3x with PBS-Tween (Appendix 7.8.1). Thereafter, 25 µL of detection antibody solution (Appendix 7.5.3.2) were added to all wells, and the plate was sealed again with an adhesive plate seal and incubated for 1 to 2 hours with vigorous shaking (300-1000 rpm) at room temperature. The plate was then washed 3x in PBS-Tween, after which 150 µL of Read Buffer T (Appendix 7.5.1.3) were added, and the plate was immediately analysed using the SECTOR® Imager (MSD, Rockville, MD, USA).
2.6.2.4 Assessment of the effect of bacterial virulence on MMP production by fibroblasts

In order to assess the impact of bacterial virulence on MMP release by corneal fibroblast in response to bacterial infection, corneal fibroblasts were challenged with the PA14 and the PAO1 strains. Confluent fibroblast monolayers were challenged in triplicate with 10^7 CFU/mL (MOI 100) CFIM of each strain and incubated in an atmosphere of 5% (v/v) CO₂ at 37°C for 24h. Supernatants were collected at 24h post challenge and spun at 2200xg for 6 min, and the samples were stored at -20°C. Detection of MMP-1, MMP-2, MMP-3 and MMP-9 release was carried out using the the Human MMP-2 Ultra-Sensitive kit and the Human MMP 3-Plex Ultra-Sensitive kit (MSD, Rockville, MD, USA) as described in section 2.6.2.30.

2.6.2.5 Assessment of the role of bacterial virulence factors on MMP production by fibroblasts

To investigate the role of bacterial flagella (flg) and type III secretion system (TTSS) in influencing MMP production by corneal fibroblasts, a sandwich immunoassay technique was carried out. Similar to section 2.5.3, fibroblast triplicates were challenged with PA14ΔPopb and PA14ΔFlgK, and the detection of MMP-1, MMP-2, MMP-3 and MMP-9 release was carried out using the the Human MMP-2 Ultra-Sensitive kit and the Human MMP 3-Plex Ultra-Sensitive kit (MSD, Rockville, MD, USA) as described in section 02.6.2.3.

2.7 Statistical analysis and graphs

Statistical analysis was done using SPSS version 22.0 (IBM, Armonk, NY, USA). Graphs were made using GraphPad Prism 6 (GraphPad, La Jolla, CA, USA).
Chapter 3: **RESULTS**

**INTERACTIONS OF PSEUDOMONAS AERUGINOSA WITH CORNEAL FIBROBLASTS**
3.1 **Bacterial association to corneal fibroblasts**

3.1.1 **Total association**

Primary human corneal fibroblasts (hCFs) confluent in tissue culture plates were challenged with various concentrations of PAO1 bacteria and bacterial association was measured and plotted against time and initial infective dose. Measurement of total association was done at different time points (3-24h), depending on the viability of the cells as demonstrated by visual inspection of fibroblast monolayers at each time point.

Results were obtained from 8 independent experiments with primary cell lines obtained from three different donor samples. PAO1 bacteria demonstrated a dose-dependent association with cell monolayers over time. Infection with $10^6$ and $10^8$ cfu/monolayer (MOI 10 and 1000, respectively) of bacteria led to a rapid saturation of the monolayers ([Figure 3.1 A](#)) and monolayer destruction by 9-24h. On the other hand, all monolayers were visually intact at 9h after infection with lower doses of bacteria [$10^1$, $10^2$, $10^3$ and $10^4$ cfu/monolayer (MOI 0.0001, 0.001, 0.01 and 0.1, respectively). For these lower MOI, the dose- and time-dependent association of PAO1 bacteria with CF was linear ([Figure 3.1 B](#)). Monolayer destruction was observed at 24h with all infecting doses.

---

**Figure 3.1 Total bacterial association of PAO1 bacteria to human corneal fibroblasts.**

Corneal fibroblasts were challenged with $10^7$ CFU/mL PAO1 for 3, 6, 9 and 24h. The association of PAO1 bacteria to corneal fibroblasts was dose- and time-dependant. The data symbols represent the mean number of associated bacteria and the error bars the standard error of the mean of 4 independent experiments. Where no error bars are visible, they fall within the symbol.
3.1.2 Electron microscopy

Viable counting assays demonstrated that PAO1 bacteria associated to human corneal fibroblasts *in-vitro*. To visually confirm the association of the bacteria to the surface of the fibroblasts, scanning electron microscopy (SEM) was used. Figure 3.2A shows uninfected fibroblasts with the characteristic fusiform shape and smooth surface with cellular projections, whereas Figure 3.2B shows the association of PAO1 bacteria to the surface of corneal fibroblasts at 3h post challenge with an infective dose of $10^4$ CFU/mL (MOI 0.1). With a higher dose of infection ($10^7$ CFU/mL; MOI 100), a greater number of bacteria was observed associated to the surface of the fibroblasts (Figure 3.2C), with evidence of some bacteria penetrating beyond the cell membrane (Figure 3.2D). This supports the viable count findings from the association experiments, where PAO1 bacteria demonstrated a time-dependant association to human corneal fibroblasts, and suggests that bacteria may also invade the fibroblasts. At later time points, there was an almost complete destruction of fibroblast monolayers, and only cell debris and clusters of bacteria were observed scattered over the surface of the Transwell membrane (images not shown), suggesting that the classically invasive PAO1 bacteria can also be cytotoxic to corneal fibroblasts at high doses of infection.
Figure 3.2 Scanning electron micrograph of bacterial association to human corneal fibroblasts.

Corneal fibroblasts were challenged with various doses of PAO1 and assessed at 3h. Associated bacteria can be seen associated to fibroblasts (B) as compared to uninfected fibroblasts with no bacteria (A). The arrows point to associated bacteria (B and C) and to bacteria seemingly penetrating the surface of the fibroblasts (D). [Original magnification= x4000 (A,B and C) and 24000x (D)].
3.1.3 The role of pili and flagella in PA association to corneal fibroblasts

*Pseudomonas aeruginosa* utilize type 4 pilus (TFP)\(^{161}\) and flagella\(^{156}\) to associate to target host cells. We sought to investigate the role of these adhesins in our microbial keratitis model by comparing the rate of bacterial association to corneal fibroblasts between wild type PAO1 bacteria and mutant strains deficient in TFP and/or flagella. Initially, we sought to ensure that all strains had the same rate of growth so that any difference in association would not be attributable to a difference in growth rates. We found that all strains had similar rates of growth (Figure 3.3).

Regarding the rate of bacterial association to corneal fibroblasts, we found that mutant strains had a significantly lower rate of association to corneal fibroblasts than wild type strains, suggesting that TFP and flagella play an important role in bacterial association to these cells. This was visually confirmed by scanning laser fluorescence microscopy (Figure 3.5) of interactions between GFP-expressing bacteria and hCFs, which showed an evident reduction in the number of associated mutant bacteria to corneal fibroblasts compared to wild type strains. There was no statistically significant difference in the rate of bacterial association between double mutant strain mutant strain FliM PilA \(tn7\)-GFP and either mutant strain FliM \(tn7\)-GFP or mutant strain PilA \(tn7\)-GFP (Table 3.1).

**Figure 3.3 Rate of growth of wild type and mutant PAO1 strains**

Wild type and mutant bacteria were left to grow in culture medium for 5h, and samples for viable counts were collected at 3 and 5h. The rate of growth of the wild type PAO1 (WT) strain, mutant strain FliM \(tn7\)-GFP deficient in FliM gene necessary for normal flagellin function, mutant strain PilA \(tn7\)-GFP deficient in the PilA gene necessary for normal type 4 pilus and mutant strain FliM PilA \(tn7\)-GFP deficient in both FliM and PilA gene was similar. The symbols represent the mean number of bacteria and the error represent the standard error of the mean of 3 independent experiments. Where no error bars are visible, they fall within the point.
Corneal fibroblasts were challenged separately with $10^7$ CFU/mL GFP-expressing wild type PAO1 and with mutant strains FliM, PilA and FliM PilA for 3h. Mutant PAO1 strains deficient in TFP and/or flagella demonstrated lower rates of association to corneal fibroblasts than wild type bacteria, suggesting that TFP and flagella play an important role in bacterial association to those cells. The bars represent the mean number of associated bacteria and the error bars the standard error of the mean of 3 independent experiments.

Table 3.1 Difference in bacterial association between wild type and mutant PAO1 strain

<table>
<thead>
<tr>
<th>Bacterial strains</th>
<th>Mean difference</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAO1WT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PAO1ΔfliM</td>
<td>31539999.67*</td>
<td>0.000</td>
</tr>
<tr>
<td>PAO1ΔpilA</td>
<td>32871333*</td>
<td>0.000</td>
</tr>
<tr>
<td>PAO1ΔfliM pilA</td>
<td>34624444.33*</td>
<td>0.000</td>
</tr>
<tr>
<td>PAO1ΔfliM pilA</td>
<td>-3084444.67</td>
<td>0.419</td>
</tr>
<tr>
<td>PAO1ΔpilA</td>
<td>-1753111.33</td>
<td>0.641</td>
</tr>
</tbody>
</table>

* The mean difference in bacterial count between the mutant strains and the wild type is significant at the 0.05 level.
Corneal fibroblasts were challenged separately with $10^7$ CFU/mL GFP-expressing wild type PAO1 and with mutant strains FliM, PilA and FliM PilA for 3h. The number of mutant GFP expressing PAO1 bacteria (green dots) deficient in TFP (B), Flagella (C) and both TFP and flagella (C) associated to corneal fibroblasts (red) was evidently less than the number of associated wild type bacteria (A). [Field of view=246 µm$^2$; Objective lens=63x]
3.2 Bacterial invasion of corneal fibroblasts

3.2.1 Gentamicin protection assays

The SEM experiments above (Section 3.1.2) suggest that PAO1 may be capable of invading corneal fibroblasts. In order to investigate this further, a group of corneal fibroblast monolayers was infected with $10^2$ and $10^4$ CFU/mL bacteria (MOI 0.001 and 0.1, respectively) and another group was infected with $10^7$ CFU/mL (MOI 100). Cellular invasion over time was quantified using the gentamicin assay, with the high infective dose assessed at 3h and the lower doses at 9h. Gentamicin was the antibiotic of choice since it effectively killed concentrations of PAO1 bacteria at growth levels expected in the culture medium after 3 and 9h (>99% bacterial killing, data not shown) and also because it is unable to penetrate the cells and kill intracellular bacteria.\textsuperscript{[423]} Cytochalasin D (CD) was added to the monolayers to investigate whether the internalization of PAO1 bacteria by corneal fibroblasts occurred by an actin microfilament-dependent process.

Cellular invasion was detectable by 9h post challenge with the lower infective doses. The number of internalized bacteria recovered from cell monolayers treated with cytochalasin (CD+G) was significantly lower than untreated cells (p<0.01, Table 3.2), suggesting that these bacteria were internalized by fibroblasts using an actin microfilament dependent process (Figure 3.6 A and B).

With the higher infective dose ($10^7$ CFU/mL; MOI 100), PAO1 bacteria invaded corneal fibroblasts by 3h post challenge, and the number of internalized bacteria recovered from cell monolayers treated with cytochalasin D was also significantly lower than untreated cells (p<0.01), similar to lower infective doses and longer duration (Figure 3.6 C and Table 3.2).

On the other hand, cytochalasin D had no significant effect on the number of bacteria adhering to fibroblasts in any of the infective doses or time points (p>0.1, Table 3.2), suggesting that the reduced number of recovered bacteria seen with cytochalasin D is not related to a reduction number in association.

In order to further analyze these data, an invasion index was calculated and expressed as a percentage of the total number of associated bacteria. The rate of invasion was higher with higher infecting dose and shorter duration, with a geometric mean level of invasion (±95% confidence limits) of 1.4% (0.68,2.87%); than with lower infecting dose and longer duration, with 0.3% (0.04,2.61%). However, this difference was not statistically significant (P>0.05).
Figure 3.6 Gentamicin protection assay of bacterial invasion of human corneal fibroblasts.

A group of corneal fibroblasts was challenged with low infective doses ($10^2$ and $10^4$ CFU/mL) and assessed at 9h post challenge (A and B), and another group was challenged with a higher infective dose ($10^7$ CFU/mL) and assessed at 3h post challenge (C). Cytochalasin D decreases bacterial internalization by inhibiting actin microfilaments. The bars represent the number of recovered bacteria with and without gentamicin treatment and with and without pre-treatment with cytochalasin D, and the error bars represent the standard error of the mean from 8 independent experiments. The asterisks indicate statistically significant differences (p<0.05)

Table 3.2 Comparison between invasion rates in fibroblasts pre-treated with cytochalasin D and untreated cells.

<table>
<thead>
<tr>
<th>Inoculum</th>
<th>Without Cytochalasin D</th>
<th>With Cytochalasin D</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>SEM</td>
<td>Mean</td>
</tr>
<tr>
<td>Invasion</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$10^2$ CFU/mL</td>
<td>1913</td>
<td>268</td>
<td>395</td>
</tr>
<tr>
<td>$10^4$ CFU/mL</td>
<td>39446</td>
<td>10300</td>
<td>2189</td>
</tr>
<tr>
<td>$10^7$ CFU/mL</td>
<td>294778</td>
<td>51378</td>
<td>46200</td>
</tr>
<tr>
<td>Association</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$10^2$ CFU/mL</td>
<td>1218000</td>
<td>300340</td>
<td>1463077</td>
</tr>
<tr>
<td>$10^4$ CFU/mL</td>
<td>1482000</td>
<td>3820312</td>
<td>17082667</td>
</tr>
<tr>
<td>$10^7$ CFU/mL</td>
<td>29011111</td>
<td>6720335</td>
<td>16511111</td>
</tr>
</tbody>
</table>

*The mean difference is significant at the 0.05 level.
3.2.2 Electron microscopy

Results from the gentamicin assay suggested that PAO1 bacteria may possibly be internalized by corneal fibroblasts during bacterial challenge and transmission electron microscopy (TEM) was used to visually confirm the presence of bacteria inside the fibroblasts. After examining several sections of fibroblasts monolayers infected with PAO1, bacterial internalization was observed occurring at different stages (Figures 3.7A-E). PAO1 bacteria were detected in close proximity to the surface of corneal fibroblasts (Figures 3.7A and B), with a cup-like invagination from the surface of the cell, where PAO1 bacteria were observed to be inside (Figures 3.7C). More advanced stages of invasion were also observed with PAO1 bacteria within cell-bound vacuoles completely inside host cells (Figures 3.7D). As a result, the ability of PAO1 to invade corneal fibroblasts was confirmed by TEM, although invasion was likely to be a rare event.

3.2.3 Effect of ciprofloxacin on internalized bacteria

Ciprofloxacin has been shown to penetrate the intact cornea and into the anterior chamber,[424] and is considered the gold standard for treatment of clinical P. aeruginosa microbial keratitis.[425] We sought to investigate the effect of ciprofloxacin on intracellular bacteria for two reasons; firstly, to confirm that the recovered bacteria from the gentamicin assays are truly intracellular and not from extracellular compartments/folds inaccessible to gentamicin; secondly, to test the ability of ciprofloxacin to accumulate inside corneal fibroblasts. The doses were chosen either to reflect MIC ranges used clinically or to match the dose of gentamicin we used.

The number of recovered bacteria from ciprofloxacin-treated infected fibroblasts was significantly less than that of gentamicin-treated cells (Table 3.3), and was inversely proportional to the dose of ciprofloxacin, indicating that ciprofloxacin accumulates inside corneal fibroblasts in a dose dependant manner, killing intracellular susceptible bacteria. The number of bacteria recovered from gentamicin-treated cells was approximately $10^5$ CFU, whereas that from ciprofloxacin-treated cells was between $10^3$ and $10^4$ CFU (Figure 3.8).
Figure 3.7 Transmission electron micrograph of PAO1 bacteria invading corneal fibroblasts.

The arrows point to bacteria adhering to the cell (A), internalized by endocytosis through a cup like invagination of the fibroblast cell membrane (B), residing inside a cytoplasmic endocytotic vesicle (C), and possibly externalizing out of the fibroblast (D). [Magnification= x45000 (A), x18500 (B) and ...]
Figure 3.8 Gentamicin-ciprofloxacin invasion assay

Infected fibroblasts were treated with gentamicin (200 µg/mL) and ciprofloxacin (10, 50 and 200 µg/mL). The bars represent the number of recovered bacteria without antibiotics and with gentamicin and ciprofloxacin treatment, and the error bars represent the standard error of the mean from 8 independent experiments. The asterisks indicate statistically significant differences (p<0.05).

Table 3.3 Difference between the number of recovered bacteria in gentamicin-treated and ciprofloxacin-treated fibroblasts

<table>
<thead>
<tr>
<th>Antibiotic</th>
<th>Mean difference</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gentamicin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ciprofloxacin10</td>
<td>119281*</td>
<td>0.000</td>
</tr>
<tr>
<td>Ciprofloxacin50</td>
<td>126582*</td>
<td>0.000</td>
</tr>
<tr>
<td>Ciprofloxacin200</td>
<td>128749*</td>
<td>0.000</td>
</tr>
</tbody>
</table>

*The mean difference is significant at the 0.05 level.

The number of recovered bacteria from ciprofloxacin treated fibroblasts was significantly lower than that from gentamicin treated cells.
3.2.4 Bacterial survival and replication inside invaded fibroblasts

Because higher bacterial doses caused almost complete destruction of cell monolayers, which hindered accurate assessment of bacterial association and invasion of fibroblasts at the 24h time point, a subset of fibroblast monolayers was treated with gentamicin to destroy all extracellular bacteria and hence protect the fibroblasts from the bacteria replicating freely in the culture medium, changing its pH and expressing exotoxins and other virulence factors that eventually destroy the cells. This allowed us to assess the survival and replication of bacteria that had already invaded the fibroblasts prior to the addition of gentamicin, and at 24h, cell monolayers were still visually mostly intact. The number of recovered bacteria increased significantly in a time dependent manner (Table 3.4), suggesting that PAO1 bacteria survived and replicated inside cornea fibroblasts (Figure 3.9). In uninfected wells, monolayers treated with gentamicin were also intact, suggesting that gentamicin alone doesn’t have a detrimental effect on cultured corneal fibroblasts, which is consistent with evidence in other cell lines.[426] Table 3.4 shows the statistical analysis of the difference between the numbers of intracellular bacteria recovered at 3, 6, 9 and 24h. There was no statistically significant difference between the number of intracellular bacteria recovered at 3h and at 6h, or between the number at 6h and at 9h. There was, however, a statistically significant difference between the numbers of intracellular bacteria at 3 and 9h (P <0.05), and at 3 and 24h (P <0.005). The difference between the numbers of intracellular bacteria was significantly higher at 24h than at all other time points.
Figure 3.9 Bacterial survival and replication inside corneal fibroblasts.

Gentamicin was added 3h post challenge to preserve the cells for the 24h duration. The bars represent the number of retrieved bacteria, and the error bars represent the standard error of the mean from 3 independent experiments. The asterisks indicate statistically significant differences (p<0.05)

Table 3.4 Difference between intracellular bacterial counts at different time points.

<table>
<thead>
<tr>
<th>Time</th>
<th>Mean Difference</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>3h</td>
<td>6h</td>
<td>214056</td>
</tr>
<tr>
<td></td>
<td>9h</td>
<td>221900</td>
</tr>
<tr>
<td></td>
<td>24h</td>
<td>681878*</td>
</tr>
<tr>
<td>6h</td>
<td>3h</td>
<td>214056</td>
</tr>
<tr>
<td></td>
<td>9h</td>
<td>7844</td>
</tr>
<tr>
<td></td>
<td>24h</td>
<td>467822*</td>
</tr>
<tr>
<td>9h</td>
<td>3h</td>
<td>221900*</td>
</tr>
<tr>
<td></td>
<td>6h</td>
<td>7844</td>
</tr>
<tr>
<td></td>
<td>24h</td>
<td>4599778*</td>
</tr>
<tr>
<td>24h</td>
<td>3h</td>
<td>6818778</td>
</tr>
<tr>
<td></td>
<td>6h</td>
<td>467822*</td>
</tr>
<tr>
<td></td>
<td>9h</td>
<td>459978*</td>
</tr>
</tbody>
</table>

* The mean difference is significant at the 0.05 level.

The number of recovered intracellular bacteria at 24h was significantly higher than the numbers at 3, 6 and 9h.
3.2.5 The role of the SRC kinase system in bacterial internalization

The SRC tyrosine kinase family of enzymes\(^{[427]}\) and the actin microfilament system\(^{[314]}\) play an important role in the internalization of *P. aeruginosa* into eukaryotic cells. We sought to investigate the role of these systems in bacterial internalization into corneal fibroblasts by inhibiting SRC tyrosine kinase and the actin microfilament system in corneal fibroblasts and then assessing the rate of bacterial invasion using a standard gentamicin protection assay.

Initially, to rule out the possibility that the observed inhibition was due to inefficient association of *P. aeruginosa* to pretreated corneal fibroblasts, bacterial association was assessed. We found that none of the tested inhibitors had a significant effect on bacterial association to corneal fibroblasts (Table 3.5 and Figure 3.10).

![Figure 3.10 Bacterial association to corneal fibroblasts pre-treated with inhibitors](image)

**Figure 3.10 Bacterial association to corneal fibroblasts pre-treated with inhibitors**

Corneal fibroblasts were pre-treated separately with genistein, PP2, and cytochalasin D for 30 min, and then challenged with \(10^7\) CFU/mL PAO1 bacteria for 3h, and gentamicin was added for a further 90 min. Bacterial association was not affected by pre-treatment of corneal fibroblasts with genistein, PP2 or cytochalasin D. The bars represent the mean number of bacteria and the error bars the standard error of the mean of 3 independent experiments.

**Table 3.5 Difference in bacterial association in the presence and absence of inhibitors**

<table>
<thead>
<tr>
<th>Inhibitors</th>
<th>Mean difference</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>No inhibitors</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cytochalasin D</td>
<td>-1078888888.9</td>
<td>0.419</td>
</tr>
<tr>
<td>Genistein</td>
<td>-720333333.3</td>
<td>0.589</td>
</tr>
<tr>
<td>PP2</td>
<td>1265555555.6</td>
<td>0.924</td>
</tr>
</tbody>
</table>

* The mean difference is significant at the 0.05 level.

There was no significant difference in bacterial association between pre-treated and untreated fibroblasts.
In the gentamicin protection assay, the number of bacteria recovered from corneal fibroblasts pre-treated with the SRC tyrosine kinase general inhibitors genistein and PP2 was significantly less than that from untreated cells (Figure 3.11), and the number of bacteria recovered from corneal fibroblasts pre-treated with cytochalasin D was significantly less than that from untreated cells (Table 3.6), suggesting that the SRC tyrosine kinase system and the actin-microfilament system also plays an important role in bacterial internalization in our model of microbial keratitis (Figure 3.11A).

Upon investigating the presence and upregulation of SRC tyrosine kinase by SDS-PAGE and Western blotting, we found that corneal fibroblasts expressed a higher level of the enzyme compared to uninfected cells, suggesting that the expression of SRC tyrosine kinase is upregulated in response to infection to facilitate bacterial internalization into the cells (Figure 3.11B). Beta actin expression was tested and was equal across all samples, confirming equal expression of proteins in all cells.

![Figure 3.11 Role of SRC tyrosine kinase and the actin microfilament system in bacterial internalisation](image)

Corneal fibroblasts were pre-treated separately with genistein, PP2, and cytochalasin D for 30 min, and then challenged with $10^7$ CFU/mL PAO1 bacteria for 3h, and gentamicin was added for a further 90 min. For the Western blot, fibroblasts were challenged with $10^7$ CFU/mL PAO1 bacteria for 3h, and then lysed, and the supernatants were run. Genistein, PP2, and cytochalasin D significantly reduced bacterial internalisation into corneal fibroblasts (A), and infected cells demonstrated heightened expression of SRC tyrosine kinase (B). The bars represent the mean number of recovered internalized bacteria and the error bars the standard error of the mean of 3 independent experiments.

<table>
<thead>
<tr>
<th>Inhibitors</th>
<th>Mean difference</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>No inhibitors</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cytochalasin D</td>
<td>48244.4*</td>
<td>0.000</td>
</tr>
<tr>
<td>Genistein</td>
<td>41911.1*</td>
<td>0.000</td>
</tr>
<tr>
<td>PP2</td>
<td>33455.6*</td>
<td>0.000</td>
</tr>
</tbody>
</table>

*The mean difference in recovered bacterial counts is significant at the 0.05 level.

The number of recovered bacteria from untreated fibroblasts was significantly higher than that from cells pre-treated with cytochalasin D, Genistein and PP2.
3.3 Bacterial cytotoxic effect on corneal fibroblasts

3.3.1 Lactate dehydrogenase (LDH) Assays

In order to further investigate the apparent death and destruction of cell monolayers induced by PAO1 bacteria, the release of LDH from corneal fibroblasts was quantified. In our experiments, corneal fibroblasts monolayers were infected with bacteria at a MOI ranging from 0.0001 to 1000 to assess the capacity of PAO1 bacteria to induce cell injury. Measurement of LDH release obtained for each bacterial inoculum was converted to a percentage of the maximum LDH release value that was measured from lysed uninfected cells. Measurements were taken at 9h post infection as it was demonstrated from infection experiments (Section 3.1.1) that cell death was most likely to occur at or after this time, and visual inspection of fibroblast monolayers showed them to be mostly intact at 9h. In addition, we found in our pilot experiments that there was no induction of LDH release at 3h and 6h post challenge, and cytotoxicity was not assessed at these time points thereafter. LDH release was detectable at 9h post challenge, and the maximum LDH release occurred at an MOI of 1, after which LDH release declined (Figure 3.12).

3.3.2 Effect of intracellular bacteria on fibroblast viability

Results from invasion experiments suggest that PAO1 bacteria survive and replicate inside gentamicin protected cells (section 3.1.3). We sought to investigate whether this presence had a significant effect on corneal fibroblast viability to gain a better understanding of whether bacterial internalization is a part of the pathogenesis, or if the bacteria invade fibroblasts for a different purpose, for example, as a reservoir for re-infection or to avoid contact with immune cells.

We found that, contrary to infected unprotected fibroblasts, which released significantly higher levels of LDH than uninfected cells, fibroblasts protected with gentamicin, and containing intracellular bacteria, only released LDH in amounts similar to spontaneous release by uninfected cells (Figure 3.13A). There was no significant difference between cytotoxicity in fibroblasts containing intracellular bacteria and uninfected cells at 9 or 24h (Table 3.7), suggesting that cell viability was not affected by the presence of replicating bacteria inside them. This agrees with other evidence, and in one study of intestinal epithelial cells, *L. monocytogenes* was found to multiply readily intracellularly without affecting intestinal epithelial cell viability.\[428\]
Figure 3.12 LDH release from infected fibroblasts as a percentage of maximum release by complete lysis of cells.

Corneal fibroblasts were challenged with various doses ($10^1$-$10^9$ CFU/mL) of PAO1 bacteria for 9h. The cytopathic effect was elucidated at 3, 6 and 9h post challenge. LDH released by infected corneal fibroblasts at 3 and 6h were similar to constitutive levels released by uninfected cells, so cytotoxicity was not assessed at these time points after the pilot experiments. The data symbols represent the mean value of LDH release percentage and the error bars in the 9h curve represent the standard error of the mean of 3 independent experiments. Where no error bars are visible, they fall within the symbol.
Figure 3.13 Effect of intracellular bacteria on LDH release

Corneal fibroblasts were challenged with $10^7$ CFU/mL of PAO1 bacteria for 9 and 24h and gentamicin was added at 3h post challenge to preserve the fibroblasts for the duration of the experiment. LDH release in protected fibroblasts was similar to spontaneous release by unchallenged cells at 9h (A) and at 24h (B). The bars represent the percentage of LDH release, and the error bars represent the standard error of the mean from 3 independent experiments. The asterisks indicate statistically significant differences.

Table 3.7 Comparison between LDH release percentage in infected and uninfected fibroblasts with and without gentamicin protection

<table>
<thead>
<tr>
<th>Cytotoxicity at 9h</th>
<th>Mean Difference</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infected</td>
<td>Uninfected</td>
<td>35.86677</td>
</tr>
<tr>
<td>Infected+Gentamicin</td>
<td>Uninfected</td>
<td>8.6667</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Cytotoxicity at 24h</th>
<th>Mean Difference</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infected</td>
<td>Uninfected</td>
<td>129.30000</td>
</tr>
<tr>
<td>Infected+Gentamicin</td>
<td>Uninfected</td>
<td>17.90000</td>
</tr>
</tbody>
</table>

* The mean difference is significant at the 0.05 level.

Infected fibroblasts produced significantly higher rise in percentage of LDH release compared to uninfected cells. With gentamicin protection, there was no significant difference in the percentage of LDH release between infected and uninfected cells.
3.3.3 Effect of bacterial virulence on cytotoxicity

*Pseudomonas aeruginosa* PA14 is a highly virulent strain and is known to be more virulent than the moderately virulent PAO1 strain.[429] We sought to investigate whether higher bacterial virulence impacts the cytotoxic effect of these bacterial strains by comparing LDH release by corneal fibroblasts when infected by those 2 strains.

We found that infection by PA14 caused significantly more LDH release by corneal fibroblasts than PAO1 (Table 3.8) suggesting that higher bacterial virulence induces more cytotoxicity in infected corneal fibroblasts (Figure 3.14).

![Figure 3.14 Effect of bacterial virulence on LDH release](image)

Corneal fibroblasts were separately challenged with $10^7$ CFU/mL PAO1 and PA14 for 9h. Corneal fibroblasts infected with PA14 released significantly more LDH than those infected with PAO1, suggesting that cytotoxicity may be correlated with bacterial virulence. The bars represent the percentage of LDH release, and the error bars represent the standard error of the mean from 3 independent experiments. The asterisks indicate statistically significant differences ($p<0.05$).

<table>
<thead>
<tr>
<th>Bacterial strains</th>
<th>Mean Difference</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAO1</td>
<td>PA14</td>
<td>80.56*</td>
</tr>
</tbody>
</table>

* The mean difference is significant at the 0.05 level.

Fibroblasts infected with PA14 produced a significantly higher rise in percentage of LDH release compared to those infected with PAO1.
To rule out the possibility that the difference in cytotoxicity noticed was due to a variation in bacterial association to fibroblasts, bacterial association was assessed. We found that there was no statistically significant difference in bacterial association between the two strains (Table 3.9), indicating that the difference in cytotoxic effect noted was not because of a difference in the efficiency of association between the two bacterial strains (Figure 3.15).

![Bacterial association to corneal fibroblasts infected with PAO1 and PA14](image)

Figure 3.15 Bacterial association to corneal fibroblasts infected with PAO1 and PA14

Corneal fibroblasts were infected with $10^7$ CFU/mL of PAO1 and PA14, and bacterial association was assessed at 3h. There was no difference in bacterial association between the 2 strains. The bars represent the number of bacteria, and the error bars represent the standard error of the mean from 3 independent experiments.

<table>
<thead>
<tr>
<th>Bacterial strains</th>
<th>Mean Difference</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAO1</td>
<td>PA14</td>
<td>415611111.7</td>
</tr>
</tbody>
</table>

Table 3.9 Difference in association between PA14 and PAO1 strains

* The mean difference in bacterial counts is significant at the 0.05 level.

There was no significant difference in bacterial association between PAO1 and PA14
3.3.4 The role of flagella and type III secretion system in cytotoxicity

*Pseudomonas aeruginosa* utilize flagella for motility and association to target cells,[156] and then uses the type III secretion system (TTSS) to inject its toxins into host cells.[213] We sought to investigate the role of these virulence factors in inducing cytotoxicity is corneal fibroblasts during acute microbial keratitis. Our experiments on the effect of bacterial virulence on cytotoxicity showed that the more virulent PA14 causes more cytotoxicity to fibroblasts, so we used a wild type PA14 strain and compared it to mutant strains PA14Δpopb deficient in the PopB translocator protein essential for normal TTSS function, and PA14ΔflgK deficient in flagella.

We found that corneal fibroblasts challenged with mutant strains PA14Δpopb and PA14ΔflgK released significantly less LDH than fibroblasts challenged with wild type PA14, suggesting that the cytotoxic effect of *P. aeruginosa* is highly dependent on their flagella and type III secretion systems (Figure 3.16). The difference between cytotoxicity induced by PA14Δpopb and by PA14ΔflgK was not statistically significant (Table 3.10).

![Figure 3.16 The role of bacterial flagella and type III secretion system in bacterial cytotoxic effects](image)

Corneal fibroblasts were separately challenged with 10^7 CFU/mL wild type PA14 and mutant strains PA14Δpopb and PA14ΔflgK for 9h. Corneal fibroblasts infected with mutant PA14 deficient in flagella and TTSS released less LDH than those infected with wild type PA14, suggesting that these virulence factors play an important role in inducing cytotoxicity. The bars represent the percentage of LDH release, and the error bars represent the standard error of the mean from 3 independent experiments. The asterisks indicate statistically significant differences (p<0.05).

<table>
<thead>
<tr>
<th>Bacterial strains</th>
<th>Mean difference</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>PA14 WT</td>
<td>PA14Δpopb</td>
<td>112.33*</td>
</tr>
<tr>
<td></td>
<td>PA14ΔflgK</td>
<td>92.67*</td>
</tr>
<tr>
<td>PA14ΔflgK</td>
<td>PA14Δpopb</td>
<td>19.67</td>
</tr>
</tbody>
</table>

* The mean difference in percentage of LDH release is significant at the 0.05 level.
To rule out the possibility that the difference in cytotoxicity noticed was due to a variation in bacterial association to fibroblasts, bacterial association was assessed. We found that, although associated bacteria were numerically less in PA14FlgK than the other strains (Figure 3.17), there was no statistically significant difference in bacterial association between the three mutant strains (Table 3.11). This might be explained by the presence of other intact bacterial association mechanisms, such as the pilus, which is normal in all strains, so the bacteria can associate to some extent to the fibroblasts, but cannot cause cytotoxicity to the same extent as the wild type strain, suggesting that cytotoxicity might be flagellum dependent as well.

![Figure 3.17 Bacterial Association to fibroblasts infected with wild type and mutant PA14](image)

**Corneal fibroblasts were infected with 10⁷ CFU/mL of wild type PA14 and mutant strains PA14Δpopb and PA14ΔflgK, and bacterial association was assessed at 3h. There was no difference in bacterial association between the 3 strains. The bars represent the number of bacteria, and the error bars represent the standard error of the mean from 3 independent experiments.**

**Table 3.11 Difference in association between wild type and mutant PA14 strains**

<table>
<thead>
<tr>
<th>Bacterial strains</th>
<th>Mean difference</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>PA14 WT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PA14Δpopb</td>
<td>-170122222.6</td>
<td>0.693</td>
</tr>
<tr>
<td>PA14ΔflgK</td>
<td>318255555.3</td>
<td>0.467</td>
</tr>
</tbody>
</table>

* The mean difference in bacterial counts is significant at the 0.05 level.

There was no significant difference in bacterial association between wild type and mutant PA14 strains.
Chapter 4: RESULTS

CORNEAL FIBROBLAST RESPONSE TO PSEUDOMONAS AERUGINOSA INFECTION
4.1 Cytokine production by corneal fibroblasts

4.1.1 Cytokine production in response to live PAO1 infection

Culture supernatants from all infection experiments were collected and analysed for pro-inflammatory cytokine release by corneal fibroblasts. Cytokines tested included Interleukin-1 beta (IL-1β), Granulocyte macrophage-colony stimulating factor (GM-CSF), Interferon gamma (IFN-γ), Interleukin-2 (IL-2), Interleukin-6 (IL-6), Interleukin-8 (IL-8), Interleukin-10 (IL-10), Interleukin-12 (IL-12) and Tumour necrosis factor alpha (TNF-α). In the initial experiments, supernatants were collected at 24h, by which point complete destruction of cell monolayers was observed for all infecting doses. Results showed that with the exception of IL-1β, which showed significantly higher levels of release (P = 0.004), all tested cytokines showed lower levels than constitutive levels released by uninfected cells (Figure 4.1).

4.1.2 Cytokine production by infected corneal fibroblasts protected with gentamicin

We hypothesized that the above results were due to destruction of corneal fibroblasts before they had a chance to release the cytokines, and that IL-1β was possibly produced early before fibroblast destruction was significant. To test this hypothesis, we challenged the fibroblasts with bacteria for 3h, then added gentamicin to kill all extracellular bacteria and protect the fibroblasts for the remainder of the 24h period. We found that cytokine levels were higher, to varying degrees, in infected cells than controls for most tested cytokines (Figure 4.2); however, the differences were not statistically significant, with the exception of IL-1β and GM-CSF, which showed significantly higher levels (Table 4.1). This may indicate that, when given enough time, corneal fibroblasts upregulate their release of pro-inflammatory cytokines in response to live infection by PAO1 bacteria.

4.1.3 Early cytokine production by infected corneal fibroblasts

To test whether IL-1β was released early shortly after challenge, culture supernatants were collected at 9h post challenge, with and without the addition of gentamicin at 3h. We found that IL-1β release by infected fibroblasts was higher than that by uninfected controls (Figure 4.3), although the difference was not statistically significant (Table 4.2). All other tested cytokines showed lower levels than uninfected controls with and without gentamicin protection (data not shown), which may merely reflect the amount of cell loss caused by 9h of bacterial challenge.
Figure 4.1 Cytokine production with live uninterrupted infection of corneal fibroblasts.

Corneal fibroblasts were infected with $10^7$ CFU/mL of live bacteria for 24h, and supernatants were tested for cytokine levels by a sandwich immune technique. All tested cytokines demonstrated lower levels than constitutive release by uninfected cells, with the exception of IL-1β, which was higher. The bars represent the mean level of cytokine release, and the error bars represent the standard error of the mean from 3 independent experiments. The asterisk represents statistically significant difference ($p<0.05$).
Corneal fibroblasts were infected with $10^7$ CFU/mL of live bacteria for 24h, and gentamicin was added at 3h post challenge to preserve the fibroblasts for the duration of the experiment. Infected protected fibroblasts produced significantly higher levels of IL-1β and GM-CSF than uninfected cells, although the difference was not statistically significant with the other tested cytokines. The bars represent the level of cytokine release, and the error bars represent the standard error of the mean from 3 independent experiments. The asterisks represent statistically significant differences (p<0.05).
Table 4.1 Comparison between cytokine levels released by uninfected cells and corneal fibroblasts protected with gentamicin after 24h of bacterial challenge.

<table>
<thead>
<tr>
<th>Cytokine</th>
<th>Infection</th>
<th>Mean difference</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>IL-1β</td>
<td>Infected</td>
<td>34.68*</td>
<td>0.001</td>
</tr>
<tr>
<td>GM-CSF</td>
<td>Infected</td>
<td>66.33*</td>
<td>0.041</td>
</tr>
<tr>
<td>IFN-γ</td>
<td>Infected</td>
<td>10.67</td>
<td>0.337</td>
</tr>
<tr>
<td>IL-10</td>
<td>Infected</td>
<td>1.33</td>
<td>0.433</td>
</tr>
<tr>
<td>IL-12</td>
<td>Infected</td>
<td>0.67</td>
<td>0.719</td>
</tr>
<tr>
<td>IL-2</td>
<td>Infected</td>
<td>3.00</td>
<td>0.286</td>
</tr>
<tr>
<td>IL-6</td>
<td>Infected</td>
<td>1845.00</td>
<td>0.243</td>
</tr>
<tr>
<td>IL-8</td>
<td>Infected</td>
<td>322.00</td>
<td>0.644</td>
</tr>
<tr>
<td>TNF-α</td>
<td>Infected</td>
<td>2.67</td>
<td>0.324</td>
</tr>
</tbody>
</table>

* The mean difference (pg/mL) is significant at the 0.05 level.

The level of cytokines released by infected fibroblasts was significantly higher than that by uninfected cells.
Figure 4.3. IL-1ß production by infected corneal fibroblasts at 9 and 24h post challenge as compared to uninfected controls.

Corneal fibroblasts were infected with $10^7$ CFU/mL of live bacteria for 9 and 24h, and gentamicin was added at 3h post challenge to preserve the fibroblasts for the duration of the experiment. Infected fibroblasts produced a higher level of IL-1ß than uninfected cells. The bars represent the level of cytokine production, and the error bars represent the standard error of the mean from 3 independent experiments. The asterisks represent statistically significant differences (p<0.05).

Table 4.2 Comparison between IL-1ß release by corneal fibroblasts protected with gentamicin and uninfected cells after 9 and 24h of bacterial challenge.

<table>
<thead>
<tr>
<th>Time</th>
<th>No gentamicin protection</th>
<th>Mean difference</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>9h</td>
<td>Infected</td>
<td>Uninfected</td>
<td>2.32</td>
</tr>
<tr>
<td></td>
<td>*P=0.010</td>
<td></td>
<td></td>
</tr>
<tr>
<td>24h</td>
<td>Infected</td>
<td>Uninfected</td>
<td>29.90*</td>
</tr>
<tr>
<td></td>
<td>*P=0.001</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time</th>
<th>Gentamicin protection</th>
<th>Mean difference</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>9h</td>
<td>Infected</td>
<td>Uninfected</td>
<td>0.23</td>
</tr>
<tr>
<td></td>
<td>*P=0.001</td>
<td></td>
<td></td>
</tr>
<tr>
<td>24h</td>
<td>Infected</td>
<td>Uninfected</td>
<td>34.68*</td>
</tr>
</tbody>
</table>

* The mean difference is significant at the 0.05 level.

The level of IL-1ß released by infected fibroblasts was significantly higher than that by uninfected cells at 24h, regardless of gentamicin protection.
4.1.4  The role of individual bacterial cellular components on cytokine production

To investigate whether corneal fibroblasts produced cytokines in response to individual bacterial components to an extent similar to their response to infection by live bacteria, we tested the levels of IL-1β production. Similar to section 4.1.4, we used 3 different cell lines from 3 different donors and although there was some inter-donor variability, we found that, while fibroblasts challenged with live bacteria produced significantly higher levels of IL-1β (Table 4.3), fibroblasts challenged with heat killed bacteria, freeze-thaw killed bacteria and lipopolysaccharide produced IL-1β in levels similar to constitutive levels produced by unchallenged cells, suggesting that IL-1β production is related to active interaction between the fibroblasts and the live bacteria rather than in response to certain exotoxins or bacterial cellular components (Figure 4.4). Lysed uninfected fibroblasts Released slightly higher IL-1β level than uninfected cells (Figure 4.4), and the difference was statistically significant in one out of the three donors, but not significant in the other two (Table 4.3).

Figure 4.4 Interleukin-1β production by corneal fibroblasts in response to challenge with live bacteria and various exotoxins and bacterial cellular components

Corneal fibroblasts were challenged with with $10^7$ CFU/mL of live bacteria (I), heat killed bacteria (HK), freeze-thaw killed bacteria (FT) and lipopolysaccharide (LPS). Controls included uninfected cells (C) and lysed uninfected cells (S). At 24h, infected fibroblasts produced higher levels of IL-1β levels than uninfected or bacterial component-challenged cells. The bars represent the level of cytokine release, and the error bars represent the standard error of the mean from 3 independent experiments. S1,S2 and S3 are cell lines from different donor
Table 4.3 Difference between IL-1β production by lysed uninfected corneal fibroblasts, and cells challenged with live bacteria and various exotoxins and bacterial cellular components

<table>
<thead>
<tr>
<th>Cell Line</th>
<th>IL-1β concentration (pg/mL)</th>
<th>Mean difference</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>S1</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infected</td>
<td>Heat killed</td>
<td>8.33333*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Freeze-thaw killed</td>
<td>8.35333*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Lipopolysaccharide</td>
<td>8.35000*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Lysed uninfected</td>
<td>8.07000*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Uninfected</td>
<td>8.32667*</td>
<td>0.000</td>
</tr>
<tr>
<td>Uninfected</td>
<td>Heat killed</td>
<td>0.00667</td>
<td>0.992</td>
</tr>
<tr>
<td></td>
<td>Freeze-thaw killed</td>
<td>0.02667</td>
<td>0.966</td>
</tr>
<tr>
<td></td>
<td>Lipopolysaccharide</td>
<td>0.02333</td>
<td>0.971</td>
</tr>
<tr>
<td></td>
<td>Lysed uninfected</td>
<td>-0.25667</td>
<td>0.685</td>
</tr>
<tr>
<td><strong>S2</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infected</td>
<td>Heat killed</td>
<td>81.14000*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Freeze-thaw killed</td>
<td>81.57000*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Lipopolysaccharide</td>
<td>81.99000*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Lysed uninfected</td>
<td>51.22667*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Uninfected</td>
<td>82.14667*</td>
<td>0.000</td>
</tr>
<tr>
<td>Uninfected</td>
<td>Heat killed</td>
<td>-1.00667</td>
<td>0.928</td>
</tr>
<tr>
<td></td>
<td>Freeze-thaw killed</td>
<td>-0.57667</td>
<td>0.958</td>
</tr>
<tr>
<td></td>
<td>Lipopolysaccharide</td>
<td>-0.15667</td>
<td>0.989</td>
</tr>
<tr>
<td></td>
<td>Lysed uninfected</td>
<td>-30.92000*</td>
<td>0.013</td>
</tr>
<tr>
<td><strong>S3</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infected</td>
<td>Heat killed</td>
<td>2.35795*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Freeze-thaw killed</td>
<td>2.35376*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Lipopolysaccharide</td>
<td>2.35599*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Lysed uninfected</td>
<td>2.25525*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Uninfected</td>
<td>2.35163*</td>
<td>0.000</td>
</tr>
<tr>
<td>Uninfected</td>
<td>Heat killed</td>
<td>0.00632</td>
<td>0.923</td>
</tr>
<tr>
<td></td>
<td>Freeze-thaw killed</td>
<td>0.00213</td>
<td>0.974</td>
</tr>
<tr>
<td></td>
<td>Lipopolysaccharide</td>
<td>0.00436</td>
<td>0.947</td>
</tr>
<tr>
<td></td>
<td>Lysed uninfected</td>
<td>-0.09638</td>
<td>0.154</td>
</tr>
</tbody>
</table>

* The mean difference is significant at the 0.05 level.
4.1.5 Effect of bacterial virulence on cytokine production

As discussed above, *P. aeruginosa* PA14 is more virulent than PAO1, and our results show that this higher virulence is associated with a more potent cytotoxic effect on corneal fibroblasts. We sought to investigate whether higher bacterial virulence also impacts IL-1ß production by corneal fibroblasts, as we hypothesized that a more virulent strain may be associated with more damage to the cornea, and hence, the amount of cytokine produced might be released earlier and in higher amounts to stop the offending pathogen.

We used 3 different cell lines from 3 different donors, and analyzed IL-1ß production at 9 and 24h. Although we noticed some variability in cytokine production between the different donors, we found that at 9h, infection by PA14 caused significantly more IL-1ß production by corneal fibroblasts than PAO1 for each cell line (Table 4.4). At 24h, however, IL-1ß production was higher with infection by PAO1 than with infection by PA14, possibly due to earlier destruction of fibroblasts by the more virulent PA14 (Figure 4.5), which also reflects the higher pattern of cytotoxicity seen with PA14 (section 3.3.3). This suggests that higher bacterial virulence induces an earlier and more potent IL-1ß production, which is in agreement with our hypothesis.

4.1.6 The role of flagellin and type III secretion system in cytokine production

Our experiments on the effect of bacterial virulence factors on its cytotoxic effects on corneal fibroblasts demonstrate that mutant bacteria deficient in the type III secretion system and flagella induce significantly less cytotoxic effects on corneal fibroblasts than wild type bacteria (Section 3.3.4). We sought to investigate if these same virulence factors had a similar effect on IL-1ß production by corneal fibroblasts; especially that higher bacterial virulence was associated with an earlier and more potent IL-1ß production (as described above).

Similar to section 4.1.4, we used three different cell lines from three different donors, and analyzed IL-1ß production at 9 and 24h. Although we noticed the same variability in cytokine production between the different donors, we found that at 9h, infection by wild type PA14 caused significantly more IL-1ß production by corneal fibroblasts than mutant PA14 strains PA14Δpopb and PA14ΔflgK for each cell line (Table 4.5). At 24h, however, similar to the pattern observed in section 4.1.4, IL-1ß production was higher with infection by mutant PA14 strains PA14Δpopb and PA14ΔflgK than with infection by wild type PA14, although the difference was occasionally not statistically significant (Figure 4.6 and Table 4.5).
Figure 4.5 Effect of bacterial virulence on cytokine production
Corneal fibroblasts were infected with $10^7$ CFU/mL of PAO1 and PA14 for 24h and supernatants were collected at 9 and 24h. Cytokine production was compared between fibroblasts infected with PAO1 bacteria (PAO1), PA14 bacteria (PA14) and uninfected cells (control). At 9h, corneal fibroblasts infected with PA14 produced more IL-1ß than those infected with PAO1, whereas at 24h, this pattern was reversed. The bars represent the level of cytokine release, and the error bars represent the standard deviation from 3 independent experiments (n=3).

Table 4.4 Difference in IL-1ß production between fibroblasts challenged with PA14 and with PAO1 across 3 different cell lines

<table>
<thead>
<tr>
<th>Time</th>
<th>Cell Line</th>
<th>IL-1ß concentration (pg/mL)</th>
<th>Mean difference</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>PAO1</td>
<td>PA14</td>
<td></td>
</tr>
<tr>
<td>9h</td>
<td>S1</td>
<td></td>
<td></td>
<td>-23.8*</td>
</tr>
<tr>
<td></td>
<td>S2</td>
<td></td>
<td></td>
<td>-173.9*</td>
</tr>
<tr>
<td></td>
<td>S3</td>
<td></td>
<td></td>
<td>-4.3*</td>
</tr>
<tr>
<td>24h</td>
<td>S1</td>
<td></td>
<td></td>
<td>5.7*</td>
</tr>
<tr>
<td></td>
<td>S2</td>
<td></td>
<td></td>
<td>42.3</td>
</tr>
<tr>
<td></td>
<td>S3</td>
<td></td>
<td></td>
<td>1.7*</td>
</tr>
</tbody>
</table>

* The mean difference in IL-1ß levels (pg/mL) is significant at the 0.05 level.
Corneal fibroblasts were infected with $10^7$ CFU/mL of wild type PA14 and mutant strains PA14Δpopb and PA14ΔflgK for 24h and supernatants were collected at 9 and 24h. Cytokine production was compared between fibroblasts infected with different PA14 strains (PA14WT, PA14Δpopb and PA14ΔflgK) and uninfected cells (control). At 9h, corneal fibroblasts infected with wild type PA14 produced more IL-1ß than those infected with mutant PA14 strains PA14Δpopb and PA14ΔflgK, whereas at 24h, this pattern was reversed, although the difference was mostly not statistically significant. The bars represent the level of cytokine release, and the error bars represent the standard deviation from 3 independent experiments.

Table 4.5 Difference in cytokine production between corneal fibroblasts infected with wild type PA14 and with mutant PA14 strains PA14Δpopb and PA14ΔflgK

<table>
<thead>
<tr>
<th>Time</th>
<th>Cell Line</th>
<th>IL-1ß concentration (pg/mL)</th>
<th>Mean difference</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>9h</td>
<td>S1</td>
<td>PA14 WT PA14Δpopb</td>
<td>49.4*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PA14ΔflgK</td>
<td>23.7*</td>
<td>0.001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PA14ΔflgK PA14Δpopb</td>
<td>25.7*</td>
<td>0.001</td>
</tr>
<tr>
<td></td>
<td>S2</td>
<td>PA14 WT PA14Δpopb</td>
<td>189.4*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PA14ΔflgK</td>
<td>185.2*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PA14ΔflgK PA14Δpopb</td>
<td>4.2</td>
<td>0.832</td>
</tr>
<tr>
<td></td>
<td>S3</td>
<td>PA14 WT PA14Δpopb</td>
<td>4.5*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PA14ΔflgK</td>
<td>1.4*</td>
<td>0.039</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PA14ΔflgK PA14Δpopb</td>
<td>3.1*</td>
<td>0.001</td>
</tr>
<tr>
<td>24h</td>
<td>S1</td>
<td>PA14 WT PA14Δpopb</td>
<td>-32.0*</td>
<td>0.003</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PA14ΔflgK</td>
<td>-54.8*</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PA14ΔflgK PA14Δpopb</td>
<td>22.8*</td>
<td>0.013</td>
</tr>
<tr>
<td></td>
<td>S2</td>
<td>PA14 WT PA14Δpopb</td>
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<td>0.412</td>
</tr>
<tr>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>PA14ΔflgK</td>
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<td>0.291</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PA14ΔflgK PA14Δpopb</td>
<td>-0.00259</td>
<td>0.999</td>
</tr>
</tbody>
</table>

* The mean difference in IL-1ß levels (pg/mL) is significant at the 0.05 level.
4.2 Matrix Metalloprotease (MMP) Production by Corneal Fibroblasts

4.2.1 Gelatine Zymography

In order to assess whether MMPs were produced by corneal fibroblasts in response to bacterial components or endotoxins, or whether it was related to exotoxins released by live bacteria, our live infection model was tested against different controls with isolated bacterial virulence factors and cellular components. These controls included heat killed (HK) bacteria, which preserved the cellular integrity of the bacteria with all the cell wall and membrane associated virulence factors; freeze-thaw (FT) killed bacteria, which destroyed the cell wall and membrane exposing all intracellular components and virulence factors without denaturing them, and pure *Pseudomonas aeruginosa* lipopolysaccharide (LPS). Our positive controls included bacteria alone with no cells and lysed uninfected fibroblasts. A subset of fibroblasts was treated with gentamicin 3h post challenge to test whether intracellular bacteria had a different effect on MMP production. Culture supernatants were collected at 9 and 24h to test whether there was a difference in MMP production between early and late stages of infection.

We found that fibroblasts challenged with HK bacteria, FT bacteria and LPS produced a constitutive 65 kDa gelatinase, possibly inactive MMP-2, similar to unchallenged cells. The same was true for cells treated with gentamicin after 3h of infection. Bacteria alone produced a 54 kDa gelatinase, possibly alkaline protease, whereas fibroblasts challenged by live bacteria released several gelatinases that were specific to the interaction between them and live bacteria. These could be other bacterial gelatinases only released upon contact with host cells, other host gelatinases, degradation products of MMP-2 and MMP-9 or a combination of the above. Our pilot experiments showed that there was no change in MMP production from constitutive levels at 9h for all challenging factors, and MMP production was not tested at this time point thereafter.
Figure 4.7 Gelatine zymogram showing MMP release by corneal fibroblasts.

Corneal fibroblasts were challenged with live bacteria and controls for 24h. In the upper gel, the two well on the left contain the protein ladder (PL) and the MMP positive control (C+). The third well had no samples (E). B represents bacteria alone, I is cells challenged with live bacteria, S is lysed uninfected cells, H is cells challenged with heat killed bacteria, F is cells challenged with freeze/thaw killed bacteria, L is cells challenged with LPS, and C is unchallenged cells. In the lower 2 gels, IG is infected cells treated with gentamicin 3h post challenge, and CG is uninfected treated with gentamicin 3h post challenge. The light coloured bands represent areas in the gelatin gel digested by the MMPs as they deposit according to their weight during the gel electrophoresis. The arrows point to MMPs released in response to live infection.
4.2.2 Sandwich immunoassay

4.2.2.1 Effect of individual cellular components on MMP production

The results of our gelatine zymography show that infected corneal fibroblasts released specific gelatinases only in response to challenge by live bacteria. We sought to further identify those gelatinases by specifically investigating the presence of MMP-1, MMP-2, MMP-3 and MMP-9 in culture supernatants. MMP-1, also known as interstitial collagenase, MMP-2 and MMP-9, also known as gelatinase A and B, respectively, and MMP-3, also known as stromelysin-1, are all zinc dependant endopeptidases that play important roles in extracellular matrix degradation, leukocyte migration and immune modulation associated with acute infections.\(^{430}\) By using a sandwich immunoassay technique, we were able to detect the levels of expression of these MMPs in our model of microbial keratitis.

We found that infected corneal fibroblasts produced a higher level of MMP-1 and MMP-9 than constitutive secretion by uninfected cells, although the difference was not statistically significant, and individual cellular components produced MMP levels similar to uninfected cells, which supports our zymography results.

With regard to MMP-2 and MMP-3, infected fibroblasts produced a lower level of MMPs than uninfected cells, whereas individual bacterial cellular components produced MMP-2 levels similar to uninfected cells and MMP-3 levels lower than uninfected cells. This may indicate that challenge of corneal fibroblasts by live bacteria or individual cellular components can inhibit the production of, or destroy the already produced, MMP-2 and MMP-3.

4.2.2.2 Effect of gentamicin protection on MMP production

Because we noticed a difference in cytokine production between corneal fibroblasts protected with gentamicin and unprotected fibroblasts (section 4.1.2), we sought to investigate whether gentamicin protection would have a similar effect on MMP production. We found that with gentamicin protection, the level of MMP-1, MMP-3 and MMP-9 produced by corneal fibroblasts was significantly higher than with no protection (Table 4.6), although the difference in production between infected and uninfected gentamicin protected fibroblasts was not statistically significant (Figure 4.9). For MMP-2, however, gentamicin protection did not have any effect on MMP production by fibroblasts, and MMP-2 production by infected fibroblasts was consistently lower than constituent levels. We hypothesized that this may be related to the destruction or inhibition of production of the MMP by the bacteria. Lysed uninfected cells released MMP levels slightly higher than constitutive levels, with the exception of MMP-3 where the level was lower than constitutive.
Figure 4.8 MMP production by corneal fibroblasts in response to challenge with live bacteria and individual bacterial components

Corneal fibroblasts were infected with 10^7 CFU/mL of live bacteria (I), 10^7 CFU/mL of live bacteria with gentamicin protection (IG), heat killed bacteria (HK), freeze-thaw killed bacteria (FT) and lipopolysaccharide (LPS). Controls included uninfected cells (C) and lysed uninfected cells (S), and MMP production was assessed by a sandwich immunoassay technique at 24h. The levels of MMP-1 and MMP-9 produced by infected fibroblasts were higher than those produced by uninfected cells, whereas the levels of MMP-2 and MMP-3 were lower than constitutive levels. Individual cellular components induced a similar level of production of MMPs by corneal fibroblasts, except for MMP-3, where the levels were higher. The bars represent the level of MMP production, and the error bars represent the standard error of the mean from 3 independent experiments.
Figure 4.9 MMP production by fibroblasts protected by gentamicin and unprotected cells

Corneal fibroblasts were infected with $10^7$ CFU/mL of PAO1 and gentamicin was added at 3h post challenge to preserve the fibroblasts for the duration of the experiment. Corneal fibroblasts protected with gentamicin produced more MMPs than uninfected cells, with the exception of MMP-2, where the level of production by infected cells was consistently lower than constitutive levels. The bars represent the level of MMP production, and the error bars represent the standard error of the mean from 3 independent experiments.

Table 4.6 MMP production by infected gentamicin protected and unprotected corneal fibroblasts

<table>
<thead>
<tr>
<th>MMP</th>
<th>Gentamicin protection</th>
<th>Mean difference</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>MMP-1</td>
<td>Protected</td>
<td>Unprotected</td>
<td>64265*</td>
</tr>
<tr>
<td>MMP-2</td>
<td>Protected</td>
<td>Unprotected</td>
<td>1915.1</td>
</tr>
<tr>
<td>MMP-3</td>
<td>Protected</td>
<td>Unprotected</td>
<td>9419.4*</td>
</tr>
<tr>
<td>MMP-9</td>
<td>Protected</td>
<td>Unprotected</td>
<td>631.6*</td>
</tr>
</tbody>
</table>

* The mean difference is significant at the 0.05 level.

Table 4.7 MMP production by infected and uninfected gentamicin protected fibroblasts

<table>
<thead>
<tr>
<th>MMP</th>
<th>Infection</th>
<th>Mean difference</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>MMP-1</td>
<td>Infected</td>
<td>Uninfected</td>
<td>40351.5</td>
</tr>
<tr>
<td>MMP-2</td>
<td>Infected</td>
<td>Uninfected</td>
<td>-22795.9*</td>
</tr>
<tr>
<td>MMP-3</td>
<td>Infected</td>
<td>Uninfected</td>
<td>3667.3</td>
</tr>
<tr>
<td>MMP-9</td>
<td>Infected</td>
<td>Uninfected</td>
<td>215.5</td>
</tr>
</tbody>
</table>

* The mean difference is significant at the 0.05 level.
4.2.2.3 Effect of bacterial virulence on MMP production

As discussed above, *P. aeruginosa* PA14 is more virulent than PAO1, and our results show that this higher virulence is associated with a higher level of cytokine production and a more potent cytotoxic effect on corneal fibroblasts. We sought to investigate whether higher bacterial virulence also impacts MMP production by corneal fibroblasts, and whether there was a difference in production between the different MMPs. Because we observed a higher cytokine production at 9h (Section 4.1.4), we decided to also assess MMP production at 9h as well as 24h.

We found that, contrary to cytotoxicity and cytokine production, infection of corneal fibroblasts with the more virulent PA14 did not result in production of higher levels of MMPs at 9h (Figure 4.10). This is consistent with the results of our earlier experiments that showed that there was no significant MMP production at 9h (Section 4.1.4) and the even lower detected level of MMPs may be secondary to a faster destruction of corneal fibroblasts by PA14 compared to PAO1 before they were able to produce the MMPs.

Similarly, at 24h, the level of MMP production by corneal fibroblasts infected with PA14 was lower than the level by fibroblasts infected with PAO1. This is also probably due to the faster destruction of corneal fibroblasts by PA14 before they were able to produce the MMPs.

4.2.2.4 The role of flagellin and type III secretion system in MMP production

Our experiments on the effect of bacterial virulence factors on corneal fibroblast cytotoxicity and cytokine production demonstrate that mutant bacteria deficient in the type III secretion system and flagella induce significantly less cytotoxic effects on corneal fibroblasts and less cytokine production than wild type bacteria (Sections 4.1.5 and 4.1.6). We sought to investigate if these virulence factors had an impact on MMP production by corneal fibroblasts, and whether there was a difference between different time points, so we investigated MMP production at 9 and 24h.

We found that at both 9 and 24h, corneal fibroblasts infected with wild type PA14 produced less MMPs than mutant PA14 strains PA14Δpopb and PA14ΔflgK (Figure 4.12 and Figure 4.13), although the difference was mostly statistically significant at 9h and non-significant at 24h (Table 4.10 and Table 4.11). We hypothesized that this reflects the pattern of cytotoxicity, where the cells are destroyed earlier by the more virulent wild type strain before they were able to produce the MMPs.
Corneal fibroblasts were infected with $10^7$ CFU/mL of PA14 and PAO1 for 9h. Infection with PA14 did not induce a higher level of MMP production by corneal fibroblasts than infection with PAO1. The bars represent the level of MMP production, and the error bars represent the standard error of the mean from 3 different experiments.

Table 4.8 Difference in MMP production between fibroblasts infected with PAO1 and PA14 at 9h

<table>
<thead>
<tr>
<th>MMP</th>
<th>MMP concentration (pg/mL)</th>
<th>Mean difference</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>MMP-1</td>
<td>PA01</td>
<td>PA14</td>
<td>4981.6</td>
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<tr>
<td>MMP-2</td>
<td>PA01</td>
<td>PA14</td>
<td>1413.3</td>
</tr>
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<td>MMP-3</td>
<td>PA01</td>
<td>PA14</td>
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<tr>
<td>MMP-9</td>
<td>PA01</td>
<td>PA14</td>
<td>40.4*</td>
</tr>
</tbody>
</table>

* The mean difference is significant at the 0.05 level.

Figure 4.10 MMP production by corneal fibroblasts infected with PAO1 and PA14 at 9h
Table 4.9 Difference in MMP production between fibroblasts infected with PAO1 and PA14 at 24h

<table>
<thead>
<tr>
<th>MMP</th>
<th>MMP concentration (pg/mL)</th>
<th>Mean difference</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>MMP-1</td>
<td>PAO1</td>
<td>PA14</td>
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</tr>
<tr>
<td>MMP-2</td>
<td>PAO1</td>
<td>PA14</td>
<td>4812.6</td>
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<td>MMP-3</td>
<td>PAO1</td>
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<tr>
<td>MMP-9</td>
<td>PAO1</td>
<td>PA14</td>
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</table>

* The mean difference is significant at the 0.05 level.

Figure 4.11 MMP production by corneal fibroblasts infected with PAO1 and PA14 at 24h
Corneal fibroblasts were infected with $10^7$ CFU/mL of PA14 and PAO1 for 24h. Infection with PA14 did not induce a higher level of MMP production by corneal fibroblasts than infection with PAO1. The bars represent the level of MMP production, and the error bars represent the standard error of the mean from 3 different experiments.
Corneal fibroblasts were infected with wild type PA14 and mutant PA14 strains PA14Δpopb and PA14ΔflgK for 9h, and MMP production was assessed. Fibroblasts infected with wild type PA14 produced less MMPs than those infected with mutant strains PA14Δpopb and PA14ΔflgK. The bars represent the level of MMP production, and the error bars represent the standard deviation of the mean from 3 different experiments.

### Table 4.10 Difference in MMP production between corneal fibroblasts infected for 9h with wild type and mutant PA14 strains

<table>
<thead>
<tr>
<th>MMP</th>
<th>MMP concentration (pg/mL)</th>
<th>Mean difference</th>
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</thead>
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<tr>
<td>MMP-1</td>
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<td></td>
<td>PA14Δpopb</td>
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<td>PA14ΔflgK</td>
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<td>PA14Δpopb</td>
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* The mean difference is significant at the 0.05 level.
Figure 4.13 MMP production by corneal fibroblasts infected with wild type and mutant PA14 strains at 24h

Fibroblasts infected with wild type PA14 produced less MMPs than those infected with mutant strains PA14Δpopb and PA14ΔflgK. The bars represent the level of MMP production, and the error bars represent the standard deviation of the mean from 3 different experiments.

Table 4.11 Difference in MMP production between corneal fibroblasts infected for 24h with wild type and mutant PA14 strains

<table>
<thead>
<tr>
<th>MMP</th>
<th>MMP concentration (pg/mL)</th>
<th>Mean difference</th>
<th>P-Value</th>
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* The mean difference is significant at the 0.05 level.
Chapter 5: **RESULTS**

**INTERACTIONS OF PSEUDOMONAS AERUGINOSA WITH THE CORNEA AS AN ORGAN**
5.1 Model Design

In Descemet’s stripping endothelial keratoplasty (DSEK) partial corneal transplants procedures, only the donor corneal endothelium and a small part of the posterior stroma are used in the procedure [Figure 5.1 (A)], leaving behind the majority of the stroma, Bowman’s layer and the epithelium [Figure 5.1 (B)] to be discarded.

After obtaining the necessary ethical approvals, the left over donor DSEK buttons were collected and used for the experiments. These buttons were live and contained all the anatomical components of the cornea involved in early microbial keratitis, and so, when infected with bacteria in a controlled environment, they provided a good model for human corneal infection, albeit without the tears, nerves, blood vessels and immune cells. Such arrangement is probably more representative of infectious crystalline keratopathy (ICK), where the infection is associated with little or no corneal or the anterior segment inflammation, usually due to a local or systemic immunocompromised state.\textsuperscript{431, 432} In these cases, the absence of the host immune response allows for the visualization of the formation and spread of the infection through the transparent cornea.\textsuperscript{432}

![Figure 5.1 Human corneal DSEK button model](image)

The anterior part of the cornea (A), comprising the epithelium and the majority of the stroma, is used in our model. The posterior part (B) is used in corneal endothelial transplant procedures (DSEK)
5.2 Bacterial association to the corneal epithelium

5.2.1 Scanning electron microscopy

*Pseudomonas aeruginosa* bacteria were shown to associate to and penetrate the superficial layers of the murine corneal epithelium up to the anterior basement membrane. We sought to investigate the association and penetration of PAO1 bacteria with whole-layer human corneal epithelium by SEM. Figure 5.2 A and B show the appearance of the normal corneal epithelium with the characteristic smooth surface and the normal minimal exfoliation of superficial epithelial cells.

When infected with PAO1 bacteria for 3h, the epithelial surface appeared much rougher, and the number of exfoliating cells increased dramatically (Figure 5.2 C and D). With higher magnifications, the bacteria appeared to associate to the surface of the superficial epithelial cells, and some appeared to be invading those cells (Figure 5.2 E and F). This mechanism of exfoliating superficial epithelial cells is an important defence mechanism whereby the bacterial load is reduced and thus minimizing the number of bacteria progressing further through the tissue.

In order to investigate the penetration of bacteria in the event of an epithelial breech, the epithelial barrier was overcome by scoring the surface of the buttons before infecting them. PAO1 bacteria appeared to associate to superficial and deeper cells within the score, possibly penetrating to deeper tissue (See Figure 5.4).
Figure 5.2 Scanning electron micrograph of bacterial association to the intact corneal epithelium.

Whole corneal buttons were infected with $10^7$ CFU/mL of bacteria for 3h before being fixed and processed for SEM. A and B are scans of uninfected buttons; C, D, E and F are infected buttons with increasing magnification. The arrows point to bacteria adhering to (E) and penetrating the surface of (F) superficial cells [Original magnification; A and C= x200, B and D= x1600, E= x5000 and F= x24000].
5.3 Bacterial penetration into corneal stroma

5.3.1 Light Microscopy

In studies of murine models of microbial keratitis, bacteria could not penetrate the intact epithelial basement membrane, mainly because the size of its pores is too small for bacteria to pass through.\(^\text{[173]}\) In our model, vertical scores were made so that they bypass the epithelial basement membrane and extend deep into the stroma, and we used both scarified and non-scarified corneal buttons to test the ability of bacteria to penetrate deeper into the stroma in the presence or absence of an intact basement membrane.

Figure 5.3 shows a light microscopic section of a scored button, showing the depth of the scores. The scores were made so that they bypass the epithelial basement membrane and into the stroma. Epithelial cells were seen to migrate to fill the gap as part of the normal wound healing response, which demonstrates that these corneal buttons are viable, with normally active cells. This is important, as it means that this model closely resembles a clinical situation, and the interactions seen here between bacteria and live corneal epithelial and stromal cells may be very similar to what occurs in a real-life human microbial keratitis.

5.3.2 Transmission electron microscopy (TEM)

In our SEM examination of scored corneal buttons, bacteria were seen to possibly associate to deeper cells within the scores, but the extent of the penetration could not be assessed due to the limitation of SEM (Figure 5.4). To further assess the penetration of bacteria into the stroma, TEM was used to examine sections of the infected corneal buttons taken at the level of the epithelial scores. In the button with an intact basement membrane, although the bacteria could penetrate through the layers of epithelium to reach the basement membrane, they were seen to line up on its epithelial side, but no bacteria were seen penetrating into the adjacent stroma (Figure 5.5 A). In the scarified button, however, the area of the epithelial score showed epithelial cells migrating to fill it, and PAO1 bacteria could be seen within the score and gaining access into the adjacent stroma (Figure 5.5 B, C and D). At later time points, bacteria were seen to start aggregating into a colony, seemingly establishing the infection within the stroma (Figure 5.5 E and F).
Figure 5.3 Light micrograph of human corneal button.
Whole corneal buttons were infected with $10^7$ CFU/mL of bacteria for 3h. The corneal epithelium was scarified by a needle to bypass the basement membrane and allow the bacteria to penetrate into the stroma (A, arrow). Epithelial cell can be seen migrating to fill the gap in a classical wound healing response (B, arrow).

Figure 5.4 Scanning electron micrograph of bacterial association to scarified corneal epithelium
Whole corneal buttons were infected with $10^7$ CFU/mL of bacteria for 3h. Bacteria (arrows) associated to the surface of superficial cells (A, original magnification= x3200), and possibly deeper cells within the score (B, original magnification= x6000).
Figure 5.5 Transmission electron micrograph of bacterial penetration into the corneal stroma.

Corneal buttons were challenged $10^7$ CFU/mL of bacteria for 3 and 9h before fixation and processing. The arrows point to bacteria penetrating into the stroma (S), and epithelial cells (EC) are seen to be migrating into the score. [Original magnification; A and C= x200, B and D= x1600, E= x5000 and F= x24000]
Chapter 6: DISCUSSION
6.1 Host-pathogen interactions in microbial keratitis

On the cellular level, we hypothesized that corneal fibroblasts do have a role in immune activation, they probably interact with bacteria on different levels, and that this interaction possibly leads to the release of cytokines and/or other mediators that modulate the immune system and possibly recruit immune cells. There are several main interactions between pathogens and host cells. The first essential step for further pathogen/host interactions is association of the bacteria to the host cells;\cite{433} then there is invasion and dissemination, which enables bacteria to propagate within the tissue and reach new sites; and cytotoxicity, which is related to the virulence and pathogenicity of the offending pathogen. Host cell response is a subsequent important interaction that normally follows some or all of the other pathogen/host interactions.

6.1.1 *P. aeruginosa* bacteria associate to corneal fibroblasts

Association of bacteria to cell surfaces is a primary essential step in bacterial colonization of the tissue at the onset of infection.\cite{434, 435} In our model, when the initial multiplicity of infection (MOI) was lower than or equal to 0.1 CFU/mL, PAO1 bacteria associated to primary corneal fibroblasts in a linear dose- and time-dependent manner up to 9h post challenge, which is consistent with other systems with well-established host-pathogen interactions.\cite{436} When the MOI was higher, however, the association was almost at its highest at 3h post challenge and did not increase much on subsequent time points. This is probably due to saturation of fibroblast monolayers with bacteria, and also due to cell destruction, which we have noted to start at 9h with MOI of 1 CFU/mL or higher. At 24h post challenge, severe cytotoxicity and complete monolayer destruction was noted with all infecting doses, indicating that cytotoxicity is indeed another important interaction in this model.

We also demonstrated that mutant PAO1 bacteria deficient in PilA, responsible for normal TFP functions, could not associate efficiently to corneal fibroblasts, neither could bacteria deficient in FliM, essential for normal flagellar functions, which confirms the role of these two important adhesins in the pathogenesis of microbial keratitis, and paves the way for investigating possible therapeutic strategies targeting them. This is consistent with infection models in other systems. In a murine pulmonary infection model, mutant *P. aeruginosa* PAK strains deficient in PilA and FliC led to significantly lower rates of pneumonia and mortality in infected mice.\cite{155} Mutant PAO1 strains deficient in PilA and FliM demonstrated deficient cap formations in their biofilms.\cite{437}

The association of bacteria to host cells involves specialized interactions between bacterial adhesins and their specific receptors on the surface of target cells.\cite{161} *P. aeruginosa* adhesins include alginate,\cite{438} surface-associated exoenzyme S,\cite{439} the flagellum,\cite{156} the pilus,\cite{160} and outer
membrane proteins. The pilus is believed to be involved in the initial contact between the bacteria and the target cell, mainly due to its extension from the cell surface, and type IV pilus (TFP) accounts for about 90% of the association capability of *P. aeruginosa* to human lung cells.

Integrins and cadherins are among the known cellular receptors for bacterial association. Integrins are a family of cell surface receptors that recognize specific ECM submolecular structures, and bacteria were found to utilize integrins to associate to many extracellular matrix molecules, including fibronectin, collagens, laminin, elastin, heparin, heparan sulphate, and chondroitin sulphate. Saccharide epitopes of globo series cell membrane glycolipids, and mannosylated integral membrane glycoproteins (uroplakins) have been shown to be involved in bacterial association in *E.coli* cystitis. Piliated bacteria can also bind to immunoglobulins, such as CD147, which was shown to be a vascular endothelial cell receptor for *N. meningitides* infection. Glycoconjugates, such as glycolipids, glycosylated proteins, and proteoglycans, act as receptors for *P. aeruginosa* binding in respiratory infections, and they demonstrate distinct apical and basolateral localization for the polarized airway epithelial cells.

6.1.2 *P. aeruginosa* bacteria invade corneal fibroblasts

Another important host-pathogen interaction is bacterial internalization into host cells. In our model, *P. aeruginosa* invasion of corneal fibroblasts could be detected by 3h post challenge, although it was a relatively rare event, occurring in about 2% of the time. This is probably because in a clinical microbial keratitis, *P. aeruginosa* bacteria do not particularly need to invade corneal fibroblasts in order to traverse the stroma. The stroma is mostly formed of collagen and extracellular matrix proteoglycans with a relatively small number of fibroblasts, and bacteria produce proteases that digest these proteins to help them propagate through the tissue. On the other hand, bacteria may use this internalization to evade host defences and antibiotics, and this is noticed clinically when reactivation of the pathology occurs despite antibiotic treatment. Our results show that PAO1 survived and even multiplied inside invaded corneal fibroblasts. This is not surprising if the main purpose of invading the corneal fibroblasts was to avoid the host immune cells, and is consistent with observations in other systems. Many bacteria replicate within the host cytosol, like *Salmonella* that reside in a cytosolic vacuole to avoid phagolysosomal fusion, or *Shigella* and some pathogenic *Burkholderia* spp. that live freely in the host cytosol.

In our TEM studies, PAO1 bacteria were noticed to enter the cells by endocytosis, where they reside within membrane-bound vacuoles. In a previous study, Fleiszig and co-authors predicted that bacterial entry to cells was an endocytic process when they observed the presence of bacteria in
intracellular vacuoles.\textsuperscript{[448]} Our TEM photographs show the invagination of the cell membrane and the actual formation of the endocytic vesicle around a bacterium during internalization.

When bacteria encounter host cells, different strains behave in a different manner, according to the type of the host cell. Some bacteria evolved specialised strategies to invade initial cellular barriers and disseminate through tissues, like for example \textit{N.meningitidis} which invade meningeal cells cross the blood-cerebrospinal fluid barrier to enter the subarachnoid space.\textsuperscript{[449]} On the other hand, several bacterial species classically known to be extracellular pathogens have evolved molecular strategies to actively induce their entry into target cells for replication and/or dissemination to other host tissues, and many have been shown to internalise into host cells, probably using intracellular compartments for persistence in target tissues.\textsuperscript{[441]}

Some bacteria inhibit internalization into professional immune cells, which phagocytise bacteria and kill them by lytic enzymes secreted into the bacteria-containing phagosomes, by utilising the ExsA-regulated effector molecules exoenzyme T (ExoT), which causes epithelial cell rounding and inhibition of bacterial internalization.\textsuperscript{[231, 258]} These strains are generally branded as cytotoxic because of their ability to kill host cells through their type III secreted toxins. Invasive strains, on the other hand, invade and multiply within host cells.\textsuperscript{[314]} Bacterial invasion may contribute to the pathogenesis or to the progression of the infection by allowing the bacteria to evade host immune effectors and induce changes in host cell function\textsuperscript{[450]}, or evading antibiotics used clinically whenever these agents are unable to penetrate the host cell membrane.\textsuperscript{[451]}

Bacteria use several mechanisms for internalization utilizing the existing signalling pathways of the host cells, particularly tyrosine kinases and the induction of tyrosine phosphorylation in target cells. Among the different tyrosine kinase systems, the family of Src tyrosine kinases seems to be significantly involved in the induction of bacterial internalization by cellular tyrosine phosphorylation in several bacterial species.\textsuperscript{[452]} In our model, pre-treating corneal fibroblasts with the Src tyrosine kinase inhibitors significantly reduced bacterial internalization into these cells. Additionally, corneal fibroblasts upregulated their expression of Src tyrosine kinase when infected with bacteria. Inhibition of the actin microfilament system also reduced bacterial internalization into corneal fibroblasts. This suggests that corneal fibroblasts utilize the Src kinase and the actin microfilament systems for bacterial internalization, much like corneal epithelial cells and other cellular environments around the body. On the other hand, bacterial association was not affected by the inhibition of Src tyrosine kinase, indicating that this system is not required for bacterial association, which is consistent with other evidence.\textsuperscript{[453]}

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The Src family of protein tyrosine kinases plays important roles in the regulation of signal transduction by a large number of cell surface receptors in multiple cellular environments. They play an important role in regulating fundamental cellular processes, including cell growth, differentiation, cell shape, migration and survival, and specialized cell signals. \(^4\) All Src tyrosine kinases consist of an N-terminal SH3 and SH2 domain, the kinase domain, and a regulatory C-terminal tyrosine residue; and are generally kept in an inactive state by tyrosine phosphorylation of the C-terminal regulatory tyrosine. Dephosphorylation of the tyrosine residue results in conformational changes in the protein that opens the kinase domain, and this results in auto-phosphorylation of a stimulatory tyrosine in the kinase domain (Tyr-416) and full activation of the kinase. \(^5\) It has been previously shown that inhibition of tyrosine kinases prevents the invasion of rabbit corneal epithelial cells by \textit{P. aeruginosa}, indicating that tyrosine kinases play a role in the internalization of by \textit{P. aeruginosa} into those cells. \(^6\) There is other evidence of the important role of Src tyrosine kinases for \textit{P. aeruginosa} internalization by mammalian cells. For instance, infection of different types of cells with various \textit{P. aeruginosa} strains results in rapid activation of Src tyrosine kinase, detectable as early as 5-10 minutes after initiation of infection, preceding the internalization of the bacteria which happens at 15 minutes. In addition, activation of Src-like tyrosine kinases results in tyrosine phosphorylation of several cellular proteins. \(^7\) The activation of Src tyrosine kinases has been elucidated in the internalization of several other bacterial species, including \textit{Yersinia enterocolitica}, \textit{S. flexneri}, \textit{L. monocytogenes}, and \textit{Neisseria gonorrhoeae}. \(^8\) This may indicate that the processes of bacterial internalization into cells are conserved, and different bacteria may utilize similar signalling pathways for it. \(^7\)

### 6.1.3 \textit{P. aeruginosa} bacteria destroy corneal fibroblasts

Although PAO1 is an invasive strain, it does retain some cytotoxic properties on human cells. \(^9\) In our bacterial association experiments, all fibroblast monolayers were completely destroyed at 24h for all infective doses, even those as low as 10 bacteria per monolayer. We investigated that by quantifying lactate dehydrogenase (LDH) released by corneal fibroblasts and comparing it to the maximum LDH release that was measured from lysed uninfected cells. Our results showed that cell death was noted at 9h post challenge, with 50% of LDH release detected with a MOI of 0.1 CFU/cell, and a maximum release with an MOI of 1 CFU/cell. With higher MOIs, the level of LDH declines, and this is probably due to the rapid cell death and hence the reduced amount of overall LDH released by dying “leaky” fibroblasts into the medium. It could also be due to degradation of LDH already released by the lytic enzymes released with cellular apoptosis or pyroptosis.
In our prolonged gentamicin protection studies, we demonstrated that PAO1 bacteria were able to survive and replicate inside corneal fibroblasts, and in spite of that, cell monolayers were visually fairly intact for up to 24h post challenge. This led us to hypothesize that the presence of replicating bacteria inside fibroblasts did not kill them, and to test that hypothesis, we measured the LDH released by infected cells as compared to that released by gentamicin protected cells containing bacteria. We found that fibroblasts with intracellular bacteria only released LDH in an amount similar to spontaneous release by uninfected cells, suggesting that cell viability was not affected by the presence of replicating bacteria inside the cells. This supports the assumption that *P. aeruginosa* use corneal fibroblasts as reservoirs for infection, and hence they need to preserve the cells within which they hide.

*P. aeruginosa* has a number of different strains that employ different strategies to effect their pathogenicity. Some strains are predominantly invasive, like PAO1, but do retain some cytotoxic effects in mammalian cells,[460] while cytotoxic strains are inherently capable of invasion, and low levels of invasion are detectable in corneal epithelial cells before cytotoxicity starts, and their invasive properties are still detectable even if cytotoxicity is disabled by gene mutation.[455] Invasive and cytotoxic strains differ in certain genes that are regulated by the transcriptional activator ExsA, encoded by *exsA*. Only invasive strains have the 49-kDa form of exoenzyme S (*exoS*) whereas *ExoU* (*exoU*) is present only in cytotoxic strains.[250] Although invasion and cytotoxicity were originally thought to be sequential events, where cytotoxicity was thought to be dependent on invasion, it has been shown that that invasion and cytotoxicity are not sequential, and that they are independent events, although they both require protein tyrosine kinase.[455]

*P. aeruginosa* isolates demonstrate a high degree of variability in virulence, ranging from moderately virulent strains, like PAO1, which belongs to a relatively uncommon clonal group, to highly virulent strains like PA14 which belongs to one of the most common clonal groups isolated.[461] The PA14 genome contains two pathogenicity islands that are absent in PAO1 which carry several genes implicated in virulence, like *exoU* and the Rcs/Pvr systems, and the deletion of these pathogenicity islands impacts the virulence of PA14 in several murine infection models.[462-464] In our model, PA14 demonstrated significantly higher cytotoxic effects on corneal fibroblasts than PAO1, which is consistent with the other evidence.

*P. aeruginosa* utilize the highly regulated type III secretion system to inject its effector proteins into target cells. These proteins cause host cells death either directly, through pore-mediated increases in membrane permeability;[212-215] or indirectly, through the activation of broad cellular defence responses.[216] *P. aeruginosa* type III secretion system triggers the inflammasome-mediated
activation of caspase 1, resulting in the production of the interleukins IL-1β and IL-18 and in pyroptosis. The type III secretion system is composed of a needle complex, translocator proteins, regulator proteins, chaperone proteins and effector proteins. It is an extremely efficient system, but it requires the full function of all its components to function properly.

In our experiments, we chose to use wild type and mutant PA14 strains to investigate the role of individual virulence factors, because PA14 is predominantly cytotoxic and would have the full machinery required for cytotoxicity. We used a mutant PA14 strain deficient in the popB translocator hydrophobic protein, which is essential for target cell pore formation critical for a normal type III secretion system function. This allowed us to assess the role of this system in bacterial cytotoxicity in our model. We also used PA14 deficient in flgK, one of the flagellar hook associated proteins essential for its function, and this allowed us to assess the significance of bacterial association as a prerequisite for further interactions, including cytotoxicity. Our results showed that wild type PA14 bacteria induced significantly more cytotoxicity on corneal fibroblasts than both mutant strains, confirming the important role of type III secretion system in bacterial cytotoxic effects on corneal fibroblasts. Our results also suggest that cytotoxicity is flagellum dependent, demonstrated by the significant reduction of the cytotoxic effect of flagella-deficient mutant to a similar level to TTSS deficient mutants, despite the latter being intact in that strain.

6.1.4 Corneal fibroblasts produce cytokines in response to *P. aeruginosa* infection

All the above studies and results suggested to us that there are indeed active interactions between PAO1 bacteria and primary human corneal fibroblasts. We hence sought to investigate whether corneal fibroblasts respond to these interactions by releasing cytokines and other mediators that modulate the pathogenesis of microbial keratitis. We were particularly interested in two classes of mediators: pro-inflammatory cytokines and matrix metalloproteases (MMP).

In our model, we measured the level of 9 pro-inflammatory cytokines in culture supernatants from our early infection experiments, but found that IL-1β was the only cytokine that showed a significantly higher level than constitutive levels released by uninfected cells at 24h post challenge. We hypothesized that this may be due to destruction of corneal fibroblasts before the transcription and release of those cytokines, and that IL-1β was possibly released early before fibroblast destruction was significant. IL-1β is produced as an inactive cytoplasmic precursor that is cleaved to generate the mature active form by caspase-1 on activation of the inflammasome pathway, so it is plausible that its activation and release occurs much earlier than all the other cytokines. This was indeed confirmed when we tested cytokine release after 9h of infection, when IL-1β level was also higher than constitutive levels. Moreover, when gentamicin was used to protect the fibroblasts for
the duration of the challenge, most tested cytokines showed a higher level than constitutive release, suggesting that corneal fibroblasts do have a role in immune activation.

Classically, when pathogen recognition receptors, such as TLRs, are stimulated by offending pathogens, inflammatory pathways are initiated, and this typically involves recruitment of MyD88, leading to NF-κB translocation to the nucleus and expression of pro-inflammatory and chemotactic cytokines. These include CXCL1/KC, IL-1α and IL-1β which recruit neutrophils from limbal capillaries to the corneal stroma, and activate macrophages and resident corneal epithelial cells and keratocytes.IL-1β in particular is involved in the generation of systemic and local responses to infection, injury, and immunological challenges and is the primary cause of chronic and acute inflammation. It is also likely that neutrophil mediated IL-1β processing has a broad role in early stage infection, with macrophage and possibly neutrophil inflammasome activity contributing at a later stage. The fact that IL-1β was released early by corneal fibroblasts in our model in response to infection by P. aeruginosa supports the assumption that these cells are not simple bystanders but are in fact sentinel cells that actively modulate immune pathways.

Our group has shown previously that activated corneal fibroblasts isolated from microbial keratitis patients upregulate their TLR4 receptors and release several pro-inflammatory cytokines when treated with pure LPS. Naïve corneal fibroblasts isolated from normal corneas, however, did not behave in the same manner. We tried to identify whether cytokine production was related to the active infection process by live bacteria itself, or whether it could also be induced by bacterial cellular components and endotoxins. We challenged fibroblasts with heat killed bacteria that have preserved cell wall and external components, freeze-thaw killed bacteria that have ruptured cell walls exposing intracellular components and pure P. aeruginosa LPS which activate pathogen recognition receptors like TLR4. Our results show that only fibroblasts challenged with live cells produced significantly higher levels of cytokines than unchallenged cells, and cells challenged with bacterial components alone produced similar levels of cytokines as constitutive levels. This suggests that it is the interaction between live bacteria and the fibroblasts that induces this response, and since intracellular bacteria in fibroblasts protected with gentamicin produced IL-1β in a level similar to unprotected cells, we concluded that cytokine production is probably impacted by a combination of the association and the type III secretion system interactions.

This led us to investigate the effect of bacterial virulence on cytokine production. We hypothesized that since PA14 is more virulent than PAO1, it may induce a stronger response from corneal fibroblasts. Our results show that PA14 infection induced a significantly higher level of IL-1β by corneal fibroblasts than PAO1 at least at 9h post challenge, but this was reversed at 24h. This
suggests that corneal fibroblasts respond earlier and more potently to infection by highly virulent strains than they do to less virulent strains, possibly to start the immune cascade earlier and recruit more immune cells to the site of infection. The finding that the pattern of IL-1β production was reversed at 24h, where the level of the cytokine was higher with infection by PAO1 than by PA14 suggests that corneal fibroblasts were possibly destroyed earlier in the process and hence IL-1β level stopped increasing at that time compared to the other fibroblasts that continued their production for a longer period so that the level climbed to a higher level. Another explanation is that bacteria are capable of destroying the already secreted cytokine, and that the more virulent PA14 have a higher capacity of destroying the cytokine than PAO1. This could be verified in future studies by measuring the levels of IL-1β by the 2 strains at more time points.

Interleukin-1β production through the inflammasome pathway has been previously demonstrated. NLRC4 activates caspase-1-dependent-IL-1β release, and is believed to be a general innate immune sensor that mediates caspase-1 activation during bacterial infection. It was concluded that the most likely triggers for NLRC4 activation are cytosolic flagellin and the inner rod protein of TTSS. Interestingly, both flagellin and the rod protein share a similar structural organization characterized by two long parallel α helices, and mutation in the conserved hydrophobic residues of these α helices disrupts the triggering of NLRC4 inflammasome activation activity of both flagellin and rod proteins. Similar to flagellin, the TTSS rod protein and the needle protein form a hollow structure as part of their secretion apparatus. Notably the needle protein and the rod are very similar in size and in three-dimensional structure. The rod protein, however, is erroneously secreted out of the bacteria by the TTSS, whereas the needle protein is essential for the function of the TTSS needle complex. The biochemical activity of the TTSS rod protein and the TTSS needle protein were demonstrated in activating the NLRC4 inflammasome, both in murine and human macrophages. The NLRC4 inflammasome, therefore, serves as an effective cytosolic surveillance system to detect bacterial flagella and/or TTSS.

The TTSS is believed to have evolved from the flagellum. They both have many structural resemblances, like the resemblance of the base of the TTSS to the basal body of the flagellum, and the needle to the flagellar hook. In addition, many of the structural proteins in the TTSS and flagellum systems bear significant sequence homology; the two systems display a highly similar three-dimensional architecture. When we investigated the role of TTSS and flagellum in cytokine production, we found that wild type PA14 bacteria induced significantly higher production of IL-1β than mutant strains. We have also showed that these mutant strains are less virulent and induced less cytotoxicity than the wild type strain. It is not merely the presence of the bacteria or their
cellular components that stimulate the cells and initiate the pathology. The loss of either the flagellum or the TTSS in the PA14 mutant strains caused significant reduction in both the cytotoxicity and the IL-1β production by corneal fibroblasts in our model.

6.1.5 Corneal fibroblasts produce MMPs in response to P. aeruginosa infection

The other class of mediators we were interested in is the MMPs. Matrix metalloproteinases (MMPs) are a family of proteolytic enzymes that have multiple roles in extracellular matrix (ECM) remodelling and immune pathways. In health, they have roles in normal development, menstruation and bone remodelling, and in disease, they are implicated in processes involving abnormal ECM turnover, like arthritis, atherosclerosis and neoplasia [472]. They also have important roles in immune modulation, facilitating white blood cell recruitment, cytokine and chemokine cleavage and defensin activation. [430] MMPs are important for successful resolution of acute infection; they facilitate the migration of immune cells, elimination of the offending pathogen, resolution of inflammation and eventually remodelling of the ECM. MMPs also activate pro-IL-1β, [473] cleave activated IL-1β, [474] and modulate chemokine activity. Tissue destruction by MMPs during excessive inflammation, however, may favour pathogen dissemination or persistence, by breaking down barriers or creating areas that are poorly accessed by host immune cells.

Corneal fibroblasts maintain the structural homeostasis of the corneal stroma by regulating synthesis and degradation of ECM components using MMPs. [137] In clinical microbial keratitis, host tissue destruction is caused by lytic enzymes released from the pathogen, corneal cells and immune cells. Bacterial proteases can be implemented in the early events of microbial keratitis, but further tissue destruction is mainly caused by proteases released by resident corneal cells [390] and infiltrating leukocytes. [372]

In our model, we demonstrated by gelatine zymography that corneal fibroblasts released several MMPs in response to challenge by live bacteria that were not released when fibroblasts were challenged with heat or freeze/thaw killed bacteria or with LPS, nor were they secreted when bacteria were protected inside the cells or when the cells were completely lysed. This suggests that these MMPs are specific to the interaction between the bacteria and the cells and are only released with live infection. Thus, as these MMPs secreted by corneal fibroblasts contribute to the tissue necrosis seen in microbial keratitis, this further supports the notion that fibroblasts have an active local role in combating the infection.

The cornea mainly produces four MMPs; MMP-1 (collagenase-1), MMP-2 (gelatinase A), MMP-3 (stromelysin-1) and MMP-9 (gelatinase B). [406] In normal corneas, however, only MMP-2 is secreted
MMP-1 mainly degrades type I collagen, the most abundant protein in the body, so it is widely expressed during several physiological and pathological tissue remodelling. The cornea is composed mainly of type I collagen, so MMP-1 is commonly isolated from the cornea. MMP-2 mainly degrades type IV collagen, and may be important in ECM remodelling in the late stages of microbial keratitis. MMP-3 degrades collagen types II, III, IV, IX, and X, proteoglycans, fibronectin, laminin, and elastin, in addition to activating MMP-1, MMP-7, and MMP-9. It is also implicated in immune cell migration and MMP-3-deficient mice have deficient neutrophil migration to infected tissue. MMP-9 degrades type IV collagen, and can upregulate chemotactic cytokines/chemokines IL-1β and MIP-2. Activated MMP-2 and MMP-9 were detected in the tear fluid of microbial keratitis patients, and it was concluded that their expression may be related to the severity of ulceration.

In our model, infected unprotected corneal fibroblasts produced a consistently lower level of all tested MMPs which probably indicates the destruction of the fibroblasts before the transcription and activation of the MMPs took place. This was supported by the finding that the levels of MMP-1, MMP-3 and MMP-9 produced by corneal fibroblasts were significantly higher with gentamicin protection than with no protection. Infected gentamicin-protected cells produced slightly higher levels of MMP-1 and MMP-9 than constitutive secretion by uninfected cells, although the difference was not statistically significant, suggesting that corneal fibroblasts may upregulate the production of these MMPs if given enough time. Lower levels of MMP-2 and MMP-3, however, were detected in infected protected fibroblasts than uninfected cells, which may indicate that challenge of corneal fibroblasts by live bacteria can inhibit the production of, or destroy the already produced, MMP-2 and MMP-3.

When investigating MMP levels in corneal fibroblasts challenged with individual bacterial cellular components, the MMP levels produced by infected protected fibroblasts were similar those in uninfected cells, suggesting that this process may be related to certain virulence factors only expressed by live bacteria. This led us to investigate the role of bacterial virulence and virulence factors, specifically flagella and type III secretion system, in this process, as they had a significant role in IL-1β production. We found that fibroblasts challenged with more virulent bacteria always produced less MMPs, which merely reflects the pattern of increased cytotoxicity associated with these strains. It seems that infection with less virulent strains, like PAO1 or mutant PA14 strains, allowed the fibroblasts more time to produce higher levels of MMPS than cells infected with more virulent strains, like wild type PA14. Although IL-1β has been shown to induce the expression of
MMPs in corneal epithelial cells and fibroblasts, we could not identify a correlation in our model. This is probably because the fibroblasts were killed before the effect of IL-1β was elucidated.

Our strategy for protecting the fibroblasts long enough to investigate the production of different mediators has worked well when the relatively short initial challenge with bacteria was enough to activate the immune pathways in the cells. On the other hand, removing all extracellular bacteria may have stopped the activation of other pathways by externally associated bacteria and their type III secretion system, thereby limiting the scope of studying these pathways in our model. Lower infective doses or different infection strategies may be used in future studies to avoid these aforementioned limitations.

6.2 A model of microbial keratitis

We have developed a model of human microbial keratitis that has 2 components: a whole tissue model of microbial keratitis, where we used whole human corneal tissue to examine the interactions in an environment that closely resembles a clinical situation, and this allowed us to study the pathology from a global tissue stand point; and a tissue culture model of human corneal fibroblasts that allowed us to examine the cellular and molecular components of the pathology.

Although the human corneal fibroblast monolayer model of microbial keratitis provides critical information about the pathogenesis of the disease, we felt it was also crucial to design another model, where the interactions of bacteria with the cornea as an organ can also be studied in detail to mimic a real life clinical situation more closely. Many investigators designed similar models in mice and other animals to study the disease, and there is, indeed, a wealth of information on the topic. But animal models, although usefully representative of the pathology, are still limited by the anatomical differences between human corneas and other mammalian corneas. The mouse cornea, for example, is much smaller than a human cornea, and mouse cells express different mediators to human cells, and yet murine microbial keratitis remains the most heavily utilized model.

We sought to investigate the sequence of events that occurs in the early stages of microbial keratitis, so we developed a human microbial keratitis model utilizing live human corneal tissue remaining after corneal transplant procedures. In a real life situation, clinical P. aeruginosa microbial keratitis rarely occurs with an intact epithelium; rather, it occurs when there is some form of epithelial injury. This can be surprising if one considers the fact that the corneal epithelium is a part of the external body surface that is continuously exposed to pathogens and offending agents. But the epithelium, like other body surfaces, has an entire arsenal of defence mechanisms that protects against a full blown pathology. In our whole tissue model, we studied the corneal tissue after infection with a high
dose of bacteria (MOI 100) for 3h by SEM and TEM. We showed that *P. aeruginosa* bacteria successfully associated to corneal epithelial cells, and even managed to penetrate the entire epithelium but could not penetrate further into the stroma unless there was a breach in the epithelial basement membrane. When there was a breach, bacteria were able to penetrate and colonize the stroma quite efficiently. This is consistent with studies utilizing mouse models, in which *P. aeruginosa* bacteria were unable to penetrate beyond the intact epithelial membrane into the stroma, but could reach the stroma through the discontinuous basement membrane.\(^{[100]}\)

The corneal epithelium has several defence mechanisms to combat infection. An important mechanical mechanism is the desquamation of superficial epithelial cells containing offending bacteria into tears. In a rat model, when *P. aeruginosa* was inoculated onto a healthy cornea, most internalized bacteria were found in cells that were readily desquamated from the cornea with rinsing,\(^{[314]}\) which led the investigators to conclude that internalization coupled with desquamation is a mechanism for clearing bacteria that manage to associate to the surface. In our model, infected corneal buttons showed a rougher surface and more apparently desquamating cells than uninfected cells by scanning electron microscopy. This probably indicates that the internalization/desquamation sequence is a defence mechanism that the human cornea employs as well, suggesting that this mechanism may be conserved across the different species.

### 6.2.1 Sequence of events in early microbial keratitis

We propose that in the healthy cornea, bacterial inocula are readily eliminated from the surface of the eye by the formidable epithelial defenses. Only in the presence of a breach are bacteria able to penetrate through the layers of the epithelial and the discontinuous basement membrane into the corneal stroma, where they can colonize and multiply. Because of the immune privilege of the eye, and the lack of mature immune cells to maintain visual function in the cornea, corneal fibroblasts are equipped to detect the presence of, and interact with, offending bacteria. Corneal fibroblasts are then able to produce IL-1β, which activates the resident CD14+ undifferentiated monocyte precursor cells to mature dendritic cells and macrophages, which in turn secrete cytokines and chemokines that attract neutrophils and other leukocytes, starting the immune cascade. Corneal fibroblasts also secrete MMPs to aid in leukocyte migration, basement membrane degradation and ECM remodelling.
In the event of an epithelial breach, bacteria are able to penetrate to the stroma, where they interact with stromal fibroblasts. These cells are equipped to be able to produce IL-1β in response to this interaction, which in turn activates the resident CD14+ undifferentiated monocyte precursor cells to mature dendritic cells and macrophages. These cells then start the immune cascade by secreting cytokines and chemokines that attract neutrophils and other leukocytes to the cornea, and these secrete proteolytic enzymes that ultimately lead to the stromal necrosis seen in progressive corneal ulceration.
6.3 Limitations and future work

6.3.1 Comparing different tissue fibroblasts to keratocyte-derived fibroblasts

Keratocytes embryologically develop from the neural crest, unlike other fibroblasts, like skin fibroblasts that develop from the mesenchyme. When keratocyte are activated, like in the event of trauma or infection, they attain a classical fibroblast phenotype, but they may behave differently from other tissue fibroblasts. In future studies, host-pathogen interactions can be studied in skin or gut fibroblasts and compared to the interactions we observed in this work.

6.3.2 Assessment of bacterial association and invasion in different strains

In our experiments, we tested for variations in bacterial association between wild type and mutant strains, and examined the effect of the absence of TFP and flagella on bacterial association early in the process, up to 3h. The total association in those strains, however, was not characterized in detail as it was in our early experiments with wild type PAO1 bacteria, and that was because our main focus was the early events of the process. In future studies, a more detailed characterisation can be pursued, and total bacterial association can be measured over more time points and with more infective doses to determine whether or not it was dose and time dependant in those mutant strains as it was in wild type strains. Different wild type strains, including isolates from clinical cases of microbial keratitis, can be used as well for that purpose, studying, for example, the difference between clinical isolates and laboratory strains in that regard. Bacterial typing of clinical isolates and correlating their virulence characteristics with the clinical picture of the corresponding patient may provide some very useful information about the prognosis of the disease and may help devise new strategies for treatment.

Similarly, bacterial invasion can be assessed and compared over more wild type and mutant strains and clinical isolates to better characterise the difference between those strains, in terms of bacterial internalisation. In addition, the role of the different virulence factors in bacterial internalisation, like TFP, flagella and TTSS, can also be assessed in future studies.

6.3.3 Removing infective medium to maintain steady infective dose

Our strategy for protecting the fibroblasts long enough to investigate the production of different mediators has worked well when the relatively short initial challenge with bacteria was enough to activate the immune pathways in the cells. On the other hand, removing all extracellular bacteria may have stopped the activation of other pathways by externally associated cells and their type III secretion system, thereby limiting the scope of studying these pathways in our model. Lower
infective doses or different infection strategies may be used in future studies to avoid these aforementioned limitations. For example, regularly replacing the infective medium with fresh medium containing known bacterial inocula may avoid the limitation of the uncontrolled exponential growth of bacteria in the culture medium over the course of the experiment.

6.3.4 Testing the gene expression of cytokines and MMPs in our experiments

In our experiments to test cytokine release by corneal fibroblasts in response to infection by different bacterial strains, and by mutant and wild type strains, we observed that the amount of cytokines seemed to decrease with time, and we proposed that this could either be because the fibroblasts are dying before producing the cytokines, or because the bacteria are destroying the cytokines after their release. One strategy to differentiate would be to test the gene expression of cytokines in our experiments; if the gene expression was similar, then this would suggest that the bacteria are destroying the already released cytokines. If, on the other hand, the gene expression detection was reduced, this may mean that the fibroblasts are destroyed before producing the cytokines or MMPs.

6.3.5 Assessment of cytokines and MMP release at more time points

In our experiments, cytokine production was assessed at 9 and 24h, mainly to allow enough time for gene transcription, translation and protein production. IL-1β release was assessed at 9h when its levels were elevated despite the destruction of the fibroblasts at 24h, but the levels at earlier time points were not measured. This can be explored in future studies to try to determine a more accurate timeline for the activation and release of IL-1β from corneal fibroblasts.

6.3.6 Testing the global cytokines and MMP release in the whole tissue model

Our main focus in our whole tissue model was to use this model to visually confirm our findings using electron microscopy, so the corneal buttons were fixed and processed immediately after the infective challenge. In future studies, and to assess the global tissue response to bacterial challenge, infective culture medium can be collected before fixing the buttons, and cytokine and MMP levels can be measured. This would ideally involve longer and more numerous time points to accurately assess the timelines of cytokine and MMP production.

6.3.7 Assessment of naïve corneal fibroblast response to a conditioned medium

In a study by Wong et al, activated corneal fibroblasts isolated from microbial keratitis patients upregulate their TLR4 receptors and release several pro-inflammatory cytokines when treated with pure LPS. In contrast, naïve corneal fibroblasts isolated from normal corneas, did not behave in the
same manner. Initially we thought that these cells were only challenged with LPS, so live infection may show a similar effect, but this was not the case. It might be possible that these fibroblasts were exposed to a different array of cytokines and other mediators within the milieu of a clinical microbial keratitis that led to their “activation” and their increased propensity to upregulate their inflammatory signalling. One way to examine that is to expose naïve corneal fibroblasts to a conditioned medium. In this medium, known doses of bacteria would be added to known numbers of neutrophils for specified times, and then the supernatant is added to naïve corneal fibroblasts for different intervals of time. The original and the resulting supernatants can be tested for cytokine production, and the results compared to estimate the induced cytokine production. These fibroblasts can also be tested for further activation and TLR4 upregulation when challenged with LPS or live bacteria.

6.3.8 Assessment of leukocyte response to cytokines released by infected fibroblasts

Supernatants resulting from live infection of corneal fibroblasts with *P. aeruginosa* can be added to polymorphonuclear leukocytes (PNL) and other immune cells to assess the effect of the liberated cytokines on immune cell activation, comparing resting cytokine levels to levels after exposure to the supernatant. This can also be assessed and compared among different wild type and mutant strains.

6.3.9 Introduction of leukocytes to the whole tissue model

In the whole tissue model, PNLs and other leukocytes can be added to the medium before and after bacterial challenge, and the resulting supernatant can be tested for cytokine levels. The buttons can also be fixed and examined by scanning and transmission electron microscopy at different time points to try to gain a better understanding of the sequence of events of microbial keratitis in this model. Scanning laser fluorescence microscopy can also be used for the same purpose using GFP expressing bacteria.

6.3.10 Introduction of epithelial cells into the tissue culture model

One of the main limitations of our tissue culture model is that it is devoid of epithelial cells. The interactions between epithelial cells and the underlying fibroblasts are very important in the pathogenesis and progression of microbial keratitis in a real life clinical situation, so the introduction of epithelial cells into the model can be pursued in future studies, and the host cell response can be studied in more detail and compared to our current data. The host-pathogen interactions between epithelial cells and *P. aeruginosa* has been extensively studied previously, so we elected not to study them again. However, these interactions can be studied as part of the introduction of epithelial cells.
into our model, in order to further dissect and understand the early sequence of events in microbial keratitis.
Chapter 7: APPENDICES
7.1 Reagents for Bacterial Culture

7.1.1 Luria-Bertani agar with 60µg/mL gentamicin

- Used to grow PAO1 tn7-GFP strains.
- To prepare 500 mL, add ingredients shown in Table 7.1 to 500 mL distilled water.

Table 7.1 Luria-Bertani agar with gentamicin

<table>
<thead>
<tr>
<th>Product</th>
<th>Source</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tryptone</td>
<td>Oxoid, Basingstoke, UK</td>
<td>5 g</td>
</tr>
<tr>
<td>Yeast extract</td>
<td>Bectod-Dickinson, Sparks, MD, USA</td>
<td>2.5 g</td>
</tr>
<tr>
<td>Sodium chloride</td>
<td>Fischer, Loughborough, UK</td>
<td>5 g</td>
</tr>
<tr>
<td>Bacterial agar</td>
<td>Oxoid, Basingstoke, UK</td>
<td>7.5 g</td>
</tr>
<tr>
<td>Gentamicin</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>4 mL</td>
</tr>
</tbody>
</table>

- Sterilize by autoclaving at 120°C, 2.7Kg cm²
- Add gentamicin when temperature drops to 55-60°C
7.2 Reagents for Cell Culture and Infection

7.2.1 Corneal fibroblast culture medium (CFCM)

- Used for isolating and growing corneal fibroblasts in preparation for infection experiments.
- To prepare 500 mL, ingredients were added as shown in Table 7.2.

<table>
<thead>
<tr>
<th>Product</th>
<th>Source</th>
<th>Concentration</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dulbecco’s Modified Eagle’s Medium/Nutrient Mixture F-12 Ham</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>-</td>
<td>500 mL</td>
</tr>
<tr>
<td>Decomplemented foetal bovine serum (dFBS)</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>10%</td>
<td>50 mL</td>
</tr>
<tr>
<td>Penicillin/Streptomycin A</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>2%</td>
<td>10 mL</td>
</tr>
<tr>
<td>Gentamicin</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>5 µg/ml</td>
<td>250 µL</td>
</tr>
<tr>
<td>Amphotericin B</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>0.2%</td>
<td>1 mL</td>
</tr>
<tr>
<td>Cholera toxin</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>0.1 µg/ml</td>
<td>50 µL</td>
</tr>
<tr>
<td>Insulin</td>
<td>Invitrogen, Paisley, UK</td>
<td>5 µg/ml</td>
<td>625 µL</td>
</tr>
<tr>
<td>Epidermal Growth Factor</td>
<td>Invitrogen, Paisley, UK</td>
<td>10 ng/ml</td>
<td>500 µL</td>
</tr>
</tbody>
</table>

- Sterilize ingredients using polysulfone membrane filter (pore size 0.2 m, Pall Life Sciences, Portsmouth, UK) before addition to medium base.

7.2.2 Corneal fibroblast infection medium (CFIM)

- Used to prepare bacterial suspensions in corneal fibroblasts infection experiments.
- To prepare 500 mL, add ingredients as shown in Table 7.3. Sterilize as above.

<table>
<thead>
<tr>
<th>Product</th>
<th>Source</th>
<th>Concentration</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dulbecco’s Modified Eagle’s Medium/Nutrient Mixture F-12 Ham</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>-</td>
<td>500 mL</td>
</tr>
<tr>
<td>Decomplemented foetal bovine serum (dFBS)*</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>1%</td>
<td>5 mL</td>
</tr>
<tr>
<td>Cholera toxin</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>0.1 µg/ml</td>
<td>50 µL</td>
</tr>
<tr>
<td>Insulin</td>
<td>Invitrogen, Paisley, UK</td>
<td>5 µg/ml</td>
<td>625 µL</td>
</tr>
<tr>
<td>Epidermal Growth Factor</td>
<td>Invitrogen, Paisley, UK</td>
<td>10 ng/ml</td>
<td>500 µL</td>
</tr>
</tbody>
</table>

*When testing for proteins, such as in gelatin zymography or in SDS-PAGE, the FCS was omitted from CFIM to improve the accuracy of the results.
7.3 Reagents for cell culture

7.3.1 Decomplemented foetal bovine serum (dFBS)
- Used in cell culture media.
- Prepared by placing the sealed 500 mL FBS bottle (Sigma-Aldrich, Dorset, UK) into water bath at 56°C for 1h.
- Store in aliquots in a -20°C freezer.

7.3.2 Cytochalasin D (CD) solution
- Used to block bacterial internalization in invasion assays.
- A working stock (10µg/mL) is prepared by adding 100 µL of CD neat stock solution (1mg/mL; Sigma-Aldrich, Dorset, UK) to 9.9 mL CFIM.
- For an experiment, this is further diluted 1/5 to a final concentration of 2µg/mL.
- Store in aliquots in a -20°C freezer.

7.3.3 Saponin lysis solution
- Used to lyse corneal fibroblasts infection experiments.
- To prepare 10 mL, add ingredients as shown in Table 7.4

<table>
<thead>
<tr>
<th>Product</th>
<th>Source</th>
<th>Concentration</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phosphate buffered saline (PBS)</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>-</td>
<td>10 mL</td>
</tr>
<tr>
<td>Saponin powder</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>1% (w/v)</td>
<td>0.1 g</td>
</tr>
<tr>
<td>Decomplemented foetal bovine serum</td>
<td>Sigma-Aldrich, Dorset, UK</td>
<td>1% (v/v)</td>
<td>100 µL</td>
</tr>
</tbody>
</table>

- Mix well until completely dissolved and sterilize using polysulfone membrane filter (pore size 0.2 µm, Pall Life Sciences, Portsmouth, UK).
7.4 Reagents used in polymerase chain reaction (PCR)

7.4.1 Agarose gel

- Used for running gel in PCR experiments.
- Prepared by adding 1g of Hi-res agarose (Geneflow, Elmhurst, UK) to 50 mL of Tris-acetate-EDTA buffer (40 mM Tris-acetate, 1 mM EDTA, pH 8.0)
- Mixture is dissolved by placing in microwave for 4 min on medium power.
- While hot, the dissolved gel is poured into a gel tray with a 5 mm comb in place.
- Gel is left to set for 30 min.

7.5 Reagents used in sandwich immunoassays

7.5.1 Reagents used in the Pro-Inflammatory 9-Plex assay

7.5.1.1 Calibrator solutions

- Used to prepare calibration curves.
- Add 1 mL to calibrator stock powder and leave for 5 min.
- From this 1µg/mL, prepare an 8-point calibration curve with solutions of 10000, 2500, 625, 156, 39, 9.8 and 0 pg/mL.
- Discard after use.

7.5.1.2 Detection antibody solution

- Used for the detection step of the assay.
- Add 3 mL Diluent 100 to Detection Antibody stock.
- Discard after use.

7.5.1.3 Reading Buffer

- Used for reading the plates.
- Add 10 mL of Read Buffer T stock to 10 mL deionized water.
- Discard after use.
7.5.2  Reagents used in the V-PLEX Human IL-1β assay

7.5.2.1  Calibrator solutions

- Used to prepare calibration curves.
- Add 1000 µL of proprietary Diluent 2 to lyophilized calibrator vial, vortex mix well and leave for 5 min. This becomes the highest calibrator.
- Prepare 7 Eppendorf tubes with 300 µL Diluent 2 in each tube
- Transfer 100 µL from the highest calibrator to the next tube.
- Repeat 4-fold serial dilution 5 more times.
- Use the last tube with only 300 µL Diluent 2 as the blank calibrator (Figure 7.1).

Figure 7.1 Serial dilution for calibrator solutions

7.5.2.2  Detection antibody solution

- Used for the detection step of the assay.
- Add 60 µL of SULFO-TAG stock antibody to 2.94 mL proprietary Diluent 3.
- Discard after use.
7.5.3 Reagents used in the Human MMP Ultra-Sensitive assays

7.5.3.1 Calibrator solutions

- Used to prepare calibration curves.
- Prepare 8 Eppendorf tubes marked SD-01 to SD-08.
- Prepare the highest calibrator (SD-01) by adding 15 µL of the Human MMP Calibrator Blend to 135 µL Diluent 2.
- Prepare SD-02 by adding 40 µL of SD-01 to 120 µL Diluent 2.
- Repeat 4-fold serial dilution 5 more times.
- Use the last tube with only 120 µL Diluent 2 as the blank calibrator.

7.5.3.2 Detection antibody solution

- Used for the detection step of the assay.
- Add 60 µL of the stock antibody blend to 2.94 mL proprietary Diluent 3.
- Discard after use.
7.6 Reagents used in Zymography

7.6.1 Zymogram sample buffer (2X):
- Added to samples prior to adding to the wells on the gel.
- To prepare 10 mL, add ingredients as shown in Table 7.5 Zymogram sample buffer.

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5 M Tris-HCl pH6.8</td>
<td>2.5 mL</td>
</tr>
<tr>
<td>100% Glycerol</td>
<td>2.0 mL</td>
</tr>
<tr>
<td>10% SDS</td>
<td>4.0 mL</td>
</tr>
<tr>
<td>0.1% Bromophenol blue</td>
<td>0.5 mL</td>
</tr>
<tr>
<td>Deionized water</td>
<td>1.0 mL</td>
</tr>
</tbody>
</table>

- Store in aliquots of 500 µL in a -20°C freezer.

7.6.2 Zymogram running buffer (5X):
- Used to fill the electrophoresis tank alongside the gels.
- To prepare 1L, add ingredients as shown in Table 7.6.

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tris</td>
<td>15g</td>
</tr>
<tr>
<td>Glycine</td>
<td>72g</td>
</tr>
<tr>
<td>SDS</td>
<td>5g</td>
</tr>
<tr>
<td>Deionized water</td>
<td>1L</td>
</tr>
</tbody>
</table>

- For an experiment, this is further diluted 1/5.
- Store at room temperature.

7.6.3 Zymogram staining buffer:
- Used to stain the gels.
- To prepare 1L, add ingredients as shown in Table 7.7 Zymogram staining buffer

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coomassie Blue R-250</td>
<td>5 mL</td>
</tr>
<tr>
<td>Methanol</td>
<td>300 mL</td>
</tr>
<tr>
<td>Acetic acid</td>
<td>100 mL</td>
</tr>
</tbody>
</table>

- Store protected from light at room temperature
- Can be recycled after use.
7.6.4 Zymogram de-staining buffer:

- Used to de-stain the gels.
- To prepare 1L, add ingredients as shown in Table 7.8.

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methanol</td>
<td>300 mL</td>
</tr>
<tr>
<td>Acetic acid</td>
<td>100 mL</td>
</tr>
<tr>
<td>Deionized water</td>
<td>600 mL</td>
</tr>
</tbody>
</table>

- Store at room temperature
7.7 Reagents used in SDS-PAGE and Western blots

7.7.1 Separating (resolving) gel

- Used to run the proteins in the gel electrophoresis.
- To prepare 4 gels, add ingredients as shown in Table 7.9

Table 7.9 Separating gel for SDS-PAGE

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>30% Acrylamide/0.8% bisacrylamide</td>
<td>6.0 mL</td>
</tr>
<tr>
<td>Separating buffer (1.5 M Tris-HCl, pH 8.8)</td>
<td>3.75 mL</td>
</tr>
<tr>
<td>Deionized water</td>
<td>5.25 mL</td>
</tr>
<tr>
<td>Ammonium persulfate 10% (w/v)</td>
<td>50 µL</td>
</tr>
<tr>
<td>TEMED</td>
<td>10 µL</td>
</tr>
</tbody>
</table>

7.7.2 Stacking gel

- Used to concentrate and pack proteins in the gel electrophoresis.
- To prepare 4 gels, add ingredients as shown in Table 7.10

Table 7.10 Stacking gel for SDS-PAGE

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>30% Acrylamide/0.8% bisacrylamide</td>
<td>0.65 mL</td>
</tr>
<tr>
<td>Stacking buffer (0.5 M Tris-HCl, pH 6.8)</td>
<td>1.25 mL</td>
</tr>
<tr>
<td>Deionized water</td>
<td>3.05 mL</td>
</tr>
<tr>
<td>Ammonium persulfate 10% (w/v)</td>
<td>25 µL</td>
</tr>
<tr>
<td>TEMED</td>
<td>5 µL</td>
</tr>
</tbody>
</table>

7.7.3 Sample buffer

- Used to prepare and load samples in the gel electrophoresis.
- To prepare a 4x solution, add ingredients as shown in Table 7.11

Table 7.11 Sample buffer for SDS-PAGE

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5 M Tris-HCl pH6.8</td>
<td>4.8 mL</td>
</tr>
<tr>
<td>100% Glycerol</td>
<td>4.0 mL</td>
</tr>
<tr>
<td>SDS</td>
<td>0.8 g</td>
</tr>
<tr>
<td>Bromophenol blue</td>
<td>5.0 mg</td>
</tr>
<tr>
<td>Deionized water</td>
<td>0.7 mL</td>
</tr>
</tbody>
</table>
7.7.4 Running buffer

- Used to fill the electrophoresis tank alongside the gels.
- To prepare a 10x solution, add ingredients as shown in Table 7.12

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tris-HCl pH 8.3</td>
<td>30.3 g</td>
</tr>
<tr>
<td>Glycine</td>
<td>144 g</td>
</tr>
<tr>
<td>SDS</td>
<td>10 g</td>
</tr>
<tr>
<td>Deionized water</td>
<td>1L</td>
</tr>
</tbody>
</table>

7.7.5 Transfer (blotting) buffer

- Used to blot the proteins from the gels onto the membranes.
- To prepare 250 mL, add 50 mL methanol to 200 mL running buffer (Table 7.12).

7.7.6 Blocking buffer

- Used to block all unbound protein.
- To prepare 20 mL, add 2 g powdered skimmed milk to 20 mL PBS-Tween (Appendix 7.8.1).
- Use 20 mL per gel.

7.7.7 Pierce™ BCA Protein Assay

- Used to quantify proteins in samples.
- Pierce™ BCA Protein Assay Kit components are shown in Table 7.13

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Constituents</th>
<th>Solvent</th>
</tr>
</thead>
<tbody>
<tr>
<td>BCA reagent A</td>
<td>Sodium carbonate</td>
<td>0.1 M sodium hydroxide</td>
</tr>
<tr>
<td></td>
<td>Sodium bicarbonate</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Bicinchoninic acid</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sodium tartrate</td>
<td></td>
</tr>
<tr>
<td>BCA reagent B</td>
<td>4% (w/v) cupric sulfate</td>
<td></td>
</tr>
<tr>
<td>Albumin standard</td>
<td>0.2 mg/mL bovine serum albumin</td>
<td>0.9% (w/v) saline</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.05% (w/v) sodium azide</td>
</tr>
</tbody>
</table>
### 7.7.8 Serial dilutions of albumin standard

- Albumin standards were used in the BCA protein assay
- The concentrations were prepared as shown in Table 7.14

<table>
<thead>
<tr>
<th>Label</th>
<th>Volume of diluent</th>
<th>Volume and source of BSA</th>
<th>Final concentration of BSA (µg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0</td>
<td>300 µL from stock</td>
<td>2000</td>
</tr>
<tr>
<td>B</td>
<td>125 µL</td>
<td>375 µL from stock</td>
<td>1500</td>
</tr>
<tr>
<td>C</td>
<td>325 µL</td>
<td>325 µL from stock</td>
<td>1000</td>
</tr>
<tr>
<td>D</td>
<td>175 µL</td>
<td>175 µL from B</td>
<td>750</td>
</tr>
<tr>
<td>E</td>
<td>325 µL</td>
<td>325 µL from C</td>
<td>500</td>
</tr>
<tr>
<td>F</td>
<td>325 µL</td>
<td>325 µL from E</td>
<td>250</td>
</tr>
<tr>
<td>G</td>
<td>325 µL</td>
<td>325 µL from F</td>
<td>125</td>
</tr>
<tr>
<td>H</td>
<td>400 µL</td>
<td>100 µL from G</td>
<td>25</td>
</tr>
<tr>
<td>I</td>
<td>400 µL</td>
<td>0</td>
<td>0 (Blank)</td>
</tr>
</tbody>
</table>

### 7.8 Miscellaneous reagents

#### 7.8.1 PBS-Tween

- Used to wash plates during SDS-PAGE and cytokine and MMP analysis.
- To prepare, add 250 µl tween to 500 mL PBS.
LIST OF REFERENCES


