The Role of Biofilms in Pathology of the Ocular Surface

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Over the past few years, we have witnessed an increasing awareness of the importance of treating ocular surface disease (OSD). Diagnoses such as dry eye and blepharitis, which were historically under-appreciated, now command attention from the ophthalmologist and investment from the ocular pharmaceutical industry. Proper treatment of the ocular surface is paramount to the health of our patients because the morbidity is real and the scope of the problem is large. Dry eye affects 30 million or more people in the USA, and up to half of patients presenting to eye-care specialists may suffer from blepharitis. In support of this, we have witnessed the commitment of big pharma and biotech alike to these causes, ensuring that both our armamentarium and its deployment will look distinctly different in the future. Moreover, the support of academia in the publication of treatment algorithms such as DEWS II and CEDARS has underscored the complexity of these pathologies and offered guidance to the treating physician. As we begin to unravel the nature of OSD and specifically blepharitis, novel therapeutic targets come into focus. One such target at the forefront of antimicrobial research is the bacterial biofilm. Recent interest in a “biofilm theory of dry eye disease” as the natural sequelae of decades of chronic blepharitis, highlights the attention the role of biofilm in OSD has garnered.

Biofilms have been around since the beginning of time. They are well-documented components of primordial slime and deep-sea hyperthermoc vents. Eculidating their role in humans, and especially ocular disease, has come at slow pace. When understanding the biofilm, we must acknowledge that bacteria behave differently under certain conditions. A planktonic or solitary bacterium is distinct from a sessile network of other bacteria as components in a biofilm. Biofilms comprise an extracellular polymeric substance matrix secreted by bacteria on a living or abiotic surface, which facilitates the development of a specialized community capable of communication, pooling of resources and protection. These capabilities enable bacteria to carry out a variety of functions not accessible to planktonic bacterial systems, which can lead to the overwhelming of host defences and infection. It is in this networked microbiological milieu that the answer to some difficult questions about how antibiotic resistance develops and how otherwise benign commensal microorganisms become pathogenic may lie.

To better understand the biofilm, it is crucial to examine its components. Its formation is known to occur in distinct stages. The first involves attachment of the planktonic bacteria to a surface. This is followed by adhesion, the formation of microcolonies, maturation and eventual detachment of the cells. The ultrastructure of a biofilm is understood through imaging with techniques such as confocal or electron microscopy. It contains polymeric substances, such as polysaccharides, nucleic acids and proteins, which are built up over a substrate and result in a three-dimensional biomass. Within this structure, a delicate homeostasis is achieved among similar or diverse species of bacteria. The organization of the matrix provides for water channels and varying gradients of nutrients, oxygen and pH, which all contribute to the formation of heterogeneous microcolonies and phenotypes within the biofilm. These microcolonies may take certain shapes, such as pillar or mushroom-shaped structures. Those in close proximity to the nutrients may be more metabolically active, while others that are more deeply embedded may lie in a state of
It has traditionally been taught that biofilms in ophthalmology are responsible for a wide variety of pathologies, including endothelialitis, keratitis, lacrimal system disease and periocular infections. Most of these manifestations are thought to develop from the introduction of foreign materials, such as intraocular lenses, contact lenses, scleral buckles, keratoprosthesis, orbital implants and punctal plugs, which may provide a charged surface on which bacteria initially attach. There is, however, a growing body of evidence that the eye itself, specifically the lid margin, is subject to the development of this biofilm naturally and without the prerequisite of foreign material introduction. Biofilms are typically found at the junction of mucous membranes, and it has been suggested that biofilm builds up year after year, until the bioburden results in vast over-colonization and eventual pathology in the form of blepharitis and even dry eye. It has also been reported that biofilms of the eyelids and eyelashes may specifically contribute to the difficulty of treating infectious blepharitis. Microorganisms ensconced in a biofilm matrix may be up to 1,000 times more resistant to antibiotics.

The further examination of bacterial species, such as Staphylococcus spp., reveals prerequisites of gene expression necessary for biofilm formation. For example, strains of S. epidermidis and coagulate-negative Staphylococi contain the ica operon responsible for the production of a homoglycan called polysaccharide intercellular adhesin. This molecule is responsible for the critical process of initial bacterial attachment, along with biofilm resistance and inhibition of neutrophil-dependent killing. Once this first stage of biofilm formation is complete, the process of quorum sensing is a cornerstone to intercellular communication. Homoserine lactones along with potassium-mediated ionic signals aid in this cell-to-cell signalling and may be responsible for an eventual change in gene expression. With respect to virulence, it is generally accepted that coagulate-negative Staphylococci encode fewer putative factors than S. aureus. Operons encoding for chemical signalling and expression of virulence factors, such as enterotoxins, leucocidins, alpha-toxin and protein A, play an important role in the development of S. aureus pathology.

The question of how to best combat robust biofilms responsible for human disease remains in play. It is prevailing thought that preventing the initial step of adhesion or destroying the mature biofilm represent good options. This can be accomplished by modifying surface topography of an implanted device or utilizing biocidal molecules. Some of the most well-regarded biofilm-destroying agents include small molecules, silver ions, nanoparticles, antiseptics, detergents, naturally occurring plant compounds and certain antibiotics. In vitro assays have also been developed, such as the Calgary Biofilm Device (MBEC Biofilms Technology Ltd., Calgary Alberta, Canada), microtitre plates, flow cell model, or suspended substratum reactor, which may assist in rapidly determining anti-biofilm susceptibilities.

In the field of ophthalmology, promising research has been conducted with azithromycin ophthalmic solution (Azasite®; Insite Vision, Inc., Alameda, CA, USA), the BlephEx® device (Scope Ophthalmics, Crawley, UK), hypochlorous acid-containing compounds, and most recently, novel formulations of dilute povidone-iodine (1.2, 2.2). Although mechanical-type debridement at the lid margin to improve meibomian gland function and reduce dry eye symptoms is a good start, patients may be treated more comprehensively through chemical means that not only address the biofilm but also reduce oxidative stress and provide neurostimulatory effects. Novel formulations of dilute povidone-iodine offer this possibility and have demonstrated both in vitro effectiveness against established biofilms of multidrug-resistant S. aureus, Klebsiella pneumoniae, Pseudomonas aeruginosa and Candida albicans, as well as clinical effectiveness with both subjective and objective improvement in clinical signs and symptoms of OSD.

The biofilm plays an important role in ocular surface pathology and homeostasis, as evidenced by its age-old, nuanced complexity and ability to occur around mucous membrane junctions of the eye. It is part of the reason why, as we age, we are more likely to encounter chronic, recalcitrant OSD. Furthermore, it explains why many therapies fail when treating these common conditions; they simply do not reach their intended target. It is the hearty and self-propagating extracellular matrix that protects these phenotypically diverse microorganisms. The single planktonic bacterium is much less fortunate, much less protected and much more susceptible to conventional antibiotic therapy. We hope that some of the newer therapeutic agents currently under investigation have a definitive role in the care of our patients. The evidence is encouraging and suggests that novel antibiotic therapies will be coming to ophthalmology clinics in the near future.