Cell interactions in Wound Biofilm and \textit{in vitro} Biofilm Revealed by Electron Microscopy

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Biofilms are aggregation of bacteria which are encapsulated in extracellular polymeric substance (EPS) and form complex structures. Biofilm associated chronic wound is a significant burden for patient and healthcare system [1-2]. Treatment of biofilm infection designed based on \textit{in vitro} culture rarely promote the chronic wound healing successfully. The principal limitation of applying results from \textit{in vitro} biofilm to clinical wound biofilm treatment is that the controlled culture condition cannot represent the real environment of wound biofilm. To understand the structural details of how bacteria survive and thrive in wound, and also compare the structural differences of wound biofilm and \textit{in vitro} biofilm, we applied Electron Microscopy (EM) techniques in structural study of porcine wound biofilm and \textit{in vitro} biofilm.

Wound biofilms were collected from porcine model with a clinically relevant mixed-species infection that established in our lab. Wound biofilm tissues were cut into 200µm thick sections. \textit{In vitro} pseudomonas aeruginosa PAO1 biofilms were cultured in agar plate at 37 °C for 48 hours. Wound biofilm sections and \textit{in vitro} biofilm discs were chemically fixed and \textit{en bloc} stained. After dehydration and infiltration, samples were embedded in durecupan resin and incubated at 60°C for 2 days. The resin embedded biofilm was processed by ultramicrotome. Electron microscopy data were collected on Helios Nanolab 600 DualBeam (FIB/SEM), Tecani F20 S/TEM, Probe Corrected Titan\textsuperscript{3TM} 80-300 S/TEM. Images were processed and visualized using IMOD [3], Chimera [4] and Avizo software packages.

Our EM result shows that wound biofilm seldom forms large aggregation on the surface of wound; instead bacteria actively interact with phagocytes, fibroblast, collagen and adipose cells (Figure 1). Bacteria niches in collagen and adipose tissues have various numbers of microorganism aggregation ranged from a few tens to a few hundreds. Occasionally isolated bacteria were found inside intact tissue, which may present a living state of bacteria that explore new niches and escape the chase of immunization cells. \textit{In vitro} biofilm forms organized layers which can extend to 200µm thick. EPS between bacteria forms web-like intercellular networking (Figure 2) which may facilitate the nutrient and information transfer, promote biofilm grow and disperse. We concluded that bacteria and host interaction is the main scenario of wound biofilm. \textit{In vitro} results can be applied to wound biofilm in limited local aggregation, but not to the whole wound environment. Effective wound biofilm treatment will need research work established on real wound model [5].

References:

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Figure 1. Cell interactions in wound biofilm. A. Bacteria and collagen. B. Bacteria and fibroblast; C. Bacteria and phagocyte. D. Bacteria and adipocyte. E. Surface rendering of 3D structure of bacteria and fibroblast. Bacteria were colored in green and purple, fibroblast was colored in peach. F. Bacteria and adipocyte. Scale bar in a-d: 1µm, in f: 5µm.

Figure 2. Extracellular Polymeric Substance (EPS) in in vitro Biofilm. A. STEM image of PAO1 biofilm. B. Volume rendering of PAO1 biofilm obtained from STEM tomography. Pseudo color was added. Scale bar: 500nm.