Chelator-Based Catheter Lock Solutions in Eradicating Organisms in Biofilm

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Two different chelator-based antimicrobial catheter lock solutions, methylene blue-citrate-parabens (MB-CIT) and minocycline-EDTA-25% ethanol (M-EDTA-25% ETOH), were compared in 2-h biofilm eradication experiments. Eradication of both mature and immature Gram-positive, Gram-negative, and fungal biofilms was assessed. M-EDTA-25% ETOH was able to fully eradicate all biofilms within 2 h. MB-CIT was only effective against immature biofilms but was unable to fully eradicate most of the mature biofilms tested.

The lumen of central venous catheters (CVC) is an important route for central line-associated bloodstream infections (CLABSI) and is usually locked with heparin or flushed with saline. However, heparin has recently been shown to stimulate Staphylococcus aureus biofilm adherence to the catheter surface (1, 2). Chelators (such as citrate or EDTA), on the other hand, have been shown to have anticoagulant/antithrombotic activity equivalent to that of heparin in the advantage of disrupting biofilm and enhancing antimicrobial agents that penetrate biofilm, such as minocycline (3–5). Chelator-based catheter lock solutions consisting of either methylene blue with citrate (MB-CIT) (Zuragen; Ash Access Technology, Lafayette, IN) or minocycline with EDTA (M-EDTA) have been shown to have activity against organisms embedded in biofilm and were also found to be effective in preventing a CLABSI after a dwell time of at least 24 h (6–10). However, a dwell time of 24 h is not practical as it cannot be accomplished in sick, hospitalized patients requiring different intravenous agents and blood products through various lumens. Previously, we have shown that adding 25% ethanol (ETOH) to M-EDTA would enhance its activity and rapidly eradicate methicillin-resistant Staphylococcus aureus (MRSA) and Candida parapsilosis within 2 h in immature biofilm (11). In the current study, we compare the activities of M-EDTA-25% ETOH and MB-CIT against various resistant bacterial and fungal (Candida species) pathogens in immature and mature biofilm after 2 h of exposure.

Biofilm was grown on sterile silicone discs (1-cm diameter) following a modified Kuhn’s method (12). Silicone discs were placed into a 24-well tissue culture plate and incubated overnight at 37°C with 1 ml of human plasma. The plasma was removed and replaced with 1 ml of a 5.5 × 10⁵ CFU/ml inoculum of various organisms. For immature biofilm, the plates were incubated for 24 h at 37°C. For mature biofilm, the plates were incubated for 48 h, with the medium being replaced after 24 h. The inoculum was then removed, and the discs were washed by shaking for 30 min in 0.9% sterile saline. After washing, the discs were placed in 1 ml of lock solution and incubated at 37°C for 2 h. The discs were then removed and placed in 5 ml of 0.9% sterile saline and sonicated to disrupt any remaining biofilm. The resulting solution was then quantitatively cultured by making serial dilutions in 0.9% sterile saline and plating on agar plates (tryptic soy agar [TSA] plus 5% sheep blood for all bacterial organisms and Sabouraud dextrose agar for yeasts). The challenge organisms were methicillin-resistant Staphylococcus aureus (MRSA), vancomycin-resistant Enterococcus faecium (VRE), multidrug-resistant (MDR) Pseudomonas aeruginosa (PS), MDR Acinetobacter baumannii (An), Candida albicans (CA), and a clinical strain of Candida glabrata. The lock solutions tested were the quadruple combination MB-CIT (0.05% methylene blue, 7.0% sodium citrate, 0.15% methylparaben, and 0.015% propylparaben) and the triple combination M-EDTA-25% ETOH (0.1% minocycline, 3% calcium EDTA, and 25% ethanol). No lock solution (broth) was applied to the controls.

Quantitative recoveries (CFU/disk) from biofilms following exposure to the lock solutions for 2 h are presented in Fig. 1A to C for the Gram-positive, Gram-negative, and fungal organisms, respectively. Reductions in recoveries for MB-CIT versus controls were significant (P < 0.05) for A. baumannii at 24 and 48 h, P. aeruginosa at 24 and 48 h, VRE E. faecium at 24 and 48 h, MRSA at 24 h, C. glabrata at 24 h, and C. albicans at 24 and 48 h. There were no significant reductions (P > 0.05) for C. albicans at 48 h and MRSA at 48 h. M-EDTA-25% ETOH gave significant reductions versus the results for controls (P < 0.01) against both 24- and 48-h biofilms of all organisms. The reductions in viable organisms with M-EDTA-25% ETOH treatment versus the reductions with MB-CIT treatment were significant (P < 0.01) for C. albicans at 24 and 48 h, C. glabrata at 48 h, and MRSA at 24 and 48 h. Trends were present but underpowered to show significance for VRE E. faecium at 48 h (P = 0.07) and A. baumannii at 48 h (P = 0.07). It is notable that the number of organisms recovered in some of the assays was below the lower limit of detection in the serial dilution and quantitative cultures.

Our data show that M-EDTA-25% ETOH and MB-CIT were equally effective in eradicating VRE E. faecium and MDR P. aeruginosa within 2 h in immature biofilm. In addition, these two lock solutions were equally efficacious in rapidly eradicating Acinetobacter in immature biofilm. However, M-EDTA-25% ETOH
isms causing CLABSI (13), MB-CIT might have a limited role in prevention. However, based on our data, MB-CIT might be useful as salvage therapy in CLABSI caused by VRE E. faecium or Pseudomonas aeruginosa.

On the other hand, M-EDTA-25% ETOH was highly effective in eradicating all biofilm-embedded organisms tested in this study within 2 h of exposure. Its broad spectrum and rapid cidal activity potentially make it a useful agent for prevention of CLABSI in high-risk patients that require heavy usage of the catheter and concurrent short lock time. In addition, M-EDTA-25% ETOH may be used in the salvage of indwelling CVC in the setting of CLABSI caused by any organism where there is a mature biofilm that has already been formed. Patients with CLABSI are often hospitalized and require various intravenous products, and in that setting, a daily hour-long catheter lock with M-EDTA-25% ETOH may provide a rapid and effective salvage solution to the indwelling CVC.

In conclusion, the two chelator-based lock solutions tested are highly effective in eradicating VRE E. faecium and P. aeruginosa embedded in immature and mature biofilm within 2 h. For eradication of the majority of organisms that cause CLABSI, such as Staphylococcus, Acinetobacter, and Candida species, through a short lock time period, the M-EDTA-25% ETOH lock was highly effective and should be studied further in clinical trials to verify its utility in the prevention and treatment of CLABSI. It also remains to be verified that the complete M-EDTA-25% ethanol lock composition does not impair the functional (mechanical) performance of polymethane catheters within which it will indwell.

**REFERENCES**


MB-CIT has been shown to have activity against organisms embedded in biofilm after 24 h of exposure, and Maki et al. have shown that this solution, when locked in for at least 48 h in hemodialysis patients, does significantly decrease the risk of CLABSI (8, 10). However, even in that hemodialysis clinical trial, it was shown that there were breakthrough infections with Staphylococcus aureus and Gram-negative Bacillus organisms in the arm that used MB-CIT, which is consistent with our data. Since staphylococci, Gram-negative organisms (such as Klebsiella spp., P. aeruginosa, Enterobacter spp., and Escherichia coli), and Candida species represent more than 70% of organisms causing CLABSI (13), MB-CIT might have a limited role in prevention. However, based on our data, MB-CIT might be useful as salvage therapy in CLABSI caused by VRE E. faecium or Pseudomonas aeruginosa.
