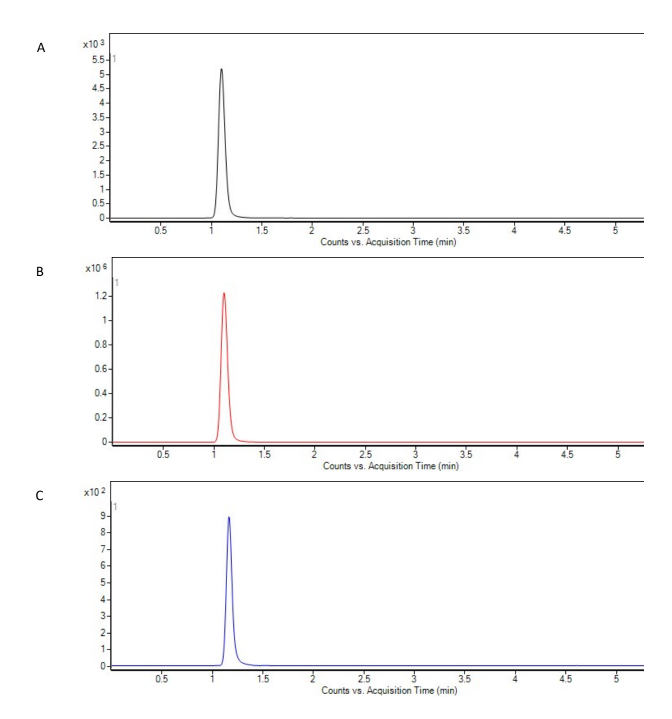
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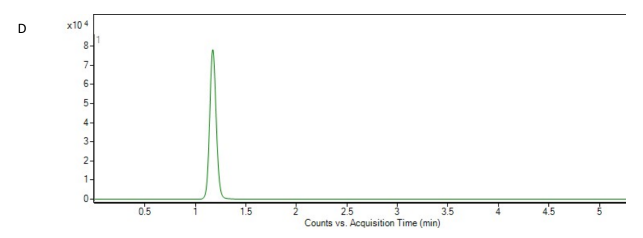


Figure 1S. Chromatograms in the serum sample spiked with meropenem, IS (A), meropenem(B), cefiderocol, IS (C), cefiderocol (D)

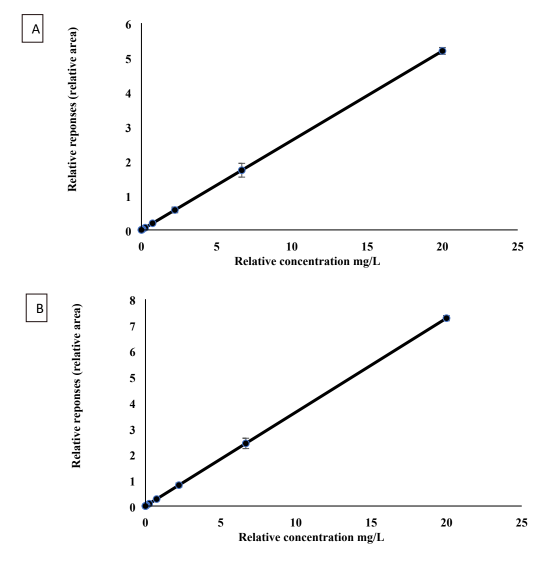


Fig. 2S The LC/MS calibration curve showing the relative area of analyte (A: cefiderocol. B: meropenem) to internal standard plotted against relative concentration analyte to internal standard. Each data point represents the average of

six separately prepared spiked standards.

a Weighting was 1/x for all analytes. R², coefficient of determination.