



Characterization of multidrug-resistant and extended-spectrum β -lactamase-producing Gram negative bacteria in an intensive care unit

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ANNOUNCEMENT

This thesis has not been previously submitted for any degree.

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بسم الله الرحمن الرحيم

"قالوا سبحانك لا علم لنا إلا ما علمتنا
إنك أنت العليم الحكيم"

صدق الله العظيم

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