



شبكة المعلومات الجامعية
التوثيق الإلكتروني والميكروفيلم

بسم الله الرحمن الرحيم



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التوثيق الإلكتروني والميكروفيلم



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التوثيق الإلكتروني والميكروفيلم

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**Phenotypic and Genotypic Characterization of *Staphylococcus epidermidis*
Isolated from Clinical Samples in Gaza Strip**

A thesis

Submitted in Partial Fulfillment of the Requirements for the Doctor of
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