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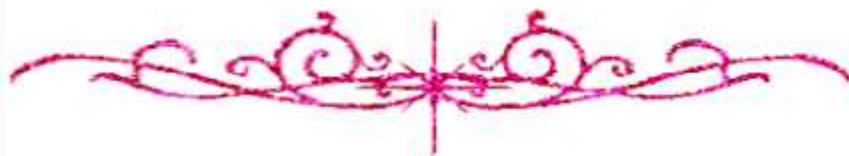
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نم بحمد الله





Ain Shams University
Faculty of Women
For Arts, Science & Education
Botany Department

Bio-agents stress as regulators of virulence and apoptosis quorum sensing in pathogenic *Candida* species.

A Thesis

Submitted in Partial Fulfillment of the Requirements for the
Degree of

M.Sc. Microbiology

By

Rahma Mostafa Mohammed Ibrahim El-Gazzar

B.Sc. (Microbiology and Chemistry, 2012)

Supervised By

Prof. Mehreshan Taha El Mokadem

Professor of Microbiology, Botany department at Women Faculty
For Arts, Science and Education

Dr. Samia Hassan Abou-Zekry

Lecturer of Microbiology, Botany department at Women Faculty
For Arts, Science and Education

Prof. Amal Ahmed Ibrahim Mekkawy

Professor of Mycology, The Regional Center of Mycology and
Biotechnology, Al-Azhar University.

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Ain Shams University
Faculty of Women
For Arts, Science & Education
Botany Department

Approval Sheet

Title: Bio-agents stress as regulators of virulence and apoptosis quorum sensing in pathogenic *Candida* species

Name: Rahma Mostafa Mohamed Ibrahim El-Gazzar

B.Sc. (Microbiology and Chemistry, 2012) Date: / /

Advisory committee

Prof. Mehreshan Taha EL-Mokadem

Professor of Microbiology, Botany Department, Women Faculty For Arts, Science and Education

Prof. Yossria Mohamed Hassan Sheteh

Professor and head of Microbiology Department , Science Faculty, Ain shams University

Prof. Nagwa Mahmoud Sedky Osman

Professor of Microbiology and biotechnology, Science Faculty, Al-Azhar University

Prof. Amal Ahmed Ibrahim Mekkawy

Professor of Mycology, The Regional Center of Mycology and Biotechnology, Al-Azhar University



Ain Shams University
Faculty of Women
For Arts, Science & Education
Botany Department

Title: Bio-agents stress as regulators of virulence and apoptosis quorum sensing in pathogenic *Candida* species.

Name: Rahma Mostafa Mohammed Ibrahim El-Gazzar
B.Sc. (Microbiology and Chemistry, 2012) Date: / /

SUPERVISION COMITTEE

Prof. Mehreshan Taha El-Mokkadem

Prof. of Microbiology
Faculty of Women for Arts, Science and Education
Ain Shams University

Dr. Samia Hassan Abou-Zekry

Lecturer of Microbiology
Faculty of Women for Arts, Science and Education
Ain Shams University

Prof. Amal Ahmed Ibrahim Mekkawy

Professor of Mycology, The Regional Center of Mycology and Biotechnology, Al-Azhar University

Announcement

**This thesis has not been previously,
submitted for any degree at this or at any
other university.**

Signature

Rahma Mostafa Mohammed Ibrahim El-Gazzar

DEDICATION

*To my father`s soul, my
mother for their endless Love,
Support & Encouragement, To
My lovely Husband Mostafa who
was always with me and has
been source of strength through
this entire experience, I hope I
have made you proud.*

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