



شبكة المعلومات الجامعية

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شبكة المعلومات الجامعية
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شبكة المعلومات الجامعية التوثيق الالكتروني والميكروفيلم



شبكة المعلومات الجامعية

جامعة عين شمس

التوثيق الالكتروني والميكرو فيلم

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2044
2044

**LASER PHOTOCHEMICAL TARGETING OF LIPOSOME
ENCAPSULATED DRUG**

A THESIS

Submitted

By

Moustafa Shaaban Abdel-Moneim Abu Hamed

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**For The Degree of
MASTER OF SCIENCE**

**In Biophysics
(Molecular Biophysics)**

**Biophysics Department
Faculty of Science, Cairo University**

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APPROVAL SHEET

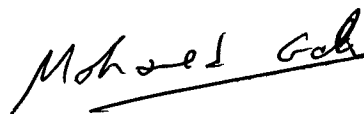
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Name of the candidate
Moustafa Shaaban Abdel-Moneim Abu Hamed

Submitted to the
Biophysics Department, Faculty of Science, Cairo University

Supervision Committee

1- Prof. Dr. Mohamed Hasanin Gaber
Professor of Biophysics
Faculty of Science, Cairo University

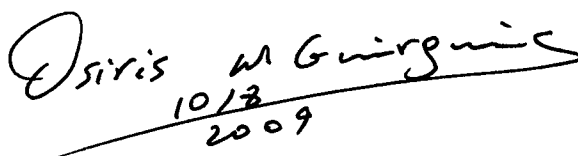


2- Dr. Maha Fadel Mohamed
Assistant professor of Pharmaceutical Technology
National Institute of Laser Enhanced Sciences, Cairo University



Head of Biophysics Department

Prof. Dr. Osiris Wanis Guirguis



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2009

بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ



سُبْحَانَكَ يَا مَنْ لَا إِلَهَ إِلَّا أَنْتَ الْإِلهُ الْوَاحِدُ الْقَهَّارُ
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صَلَّى الْعَظِيمِ

(فصلت 53)

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To...

Our beloved, Great Prophet

Mohamed

Who learned us and learned the entire world...

How to be human beings

MY MOTHER,

MY FATHER,

MY WIFE,

MY CHILDREN,

AND

ALL MY FAMILY

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