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شبكة المعلومات الجامعية التوثيق الالكتروني والميكرو فيلم



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جامعة عين شمس

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SYNTHESIS OF CERTAIN ACRIDINE DERIVATIVES OF ANTICIPATED ANTITUMOR AND ANTI-HIV-1 ACTIVITIES

THESIS
Presented by

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Submitted in partial fulfilment for the
MASTER DEGREE
in
Pharmaceutical Sciences
(Pharmaceutical Chemistry)

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1997

بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ

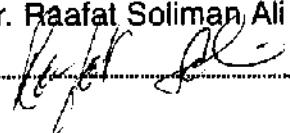
APPROVAL SHEET

Title of Thesis : Synthesis of Certain Acridine Derivatives of Anticipated Antitumor
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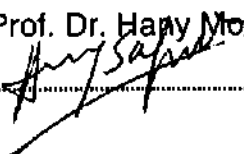
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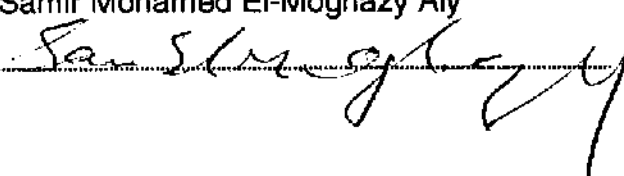
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Besides the work presented in this thesis, the candidate has attended prerequisite postgraduate course for one year which includes the following topics :

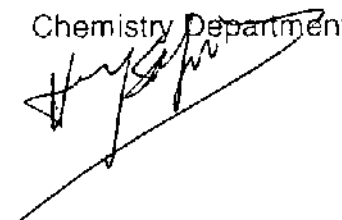
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- 2) Physical Chemistry .
- 3) Mathematics .
- 4) Statistics .
- 5) Special Course in Pharmaceutical Chemistry .

The candidate has successfully passed an examination in these topics .

Prof . Dr. Hany M. Safwat

Professor and Head of Pharmaceutical

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A handwritten signature in black ink, appearing to read 'Hany M. Safwat', is written over a diagonal line that extends from the bottom left towards the top right.

**To the memory of my father
To my mother and my sisters
To my husband and my son**

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