

A Pharmacognostical Study of
***Celtis australis* L. and *Celtis occidentalis* L.**
cultivated in Egypt

A Thesis Submitted by

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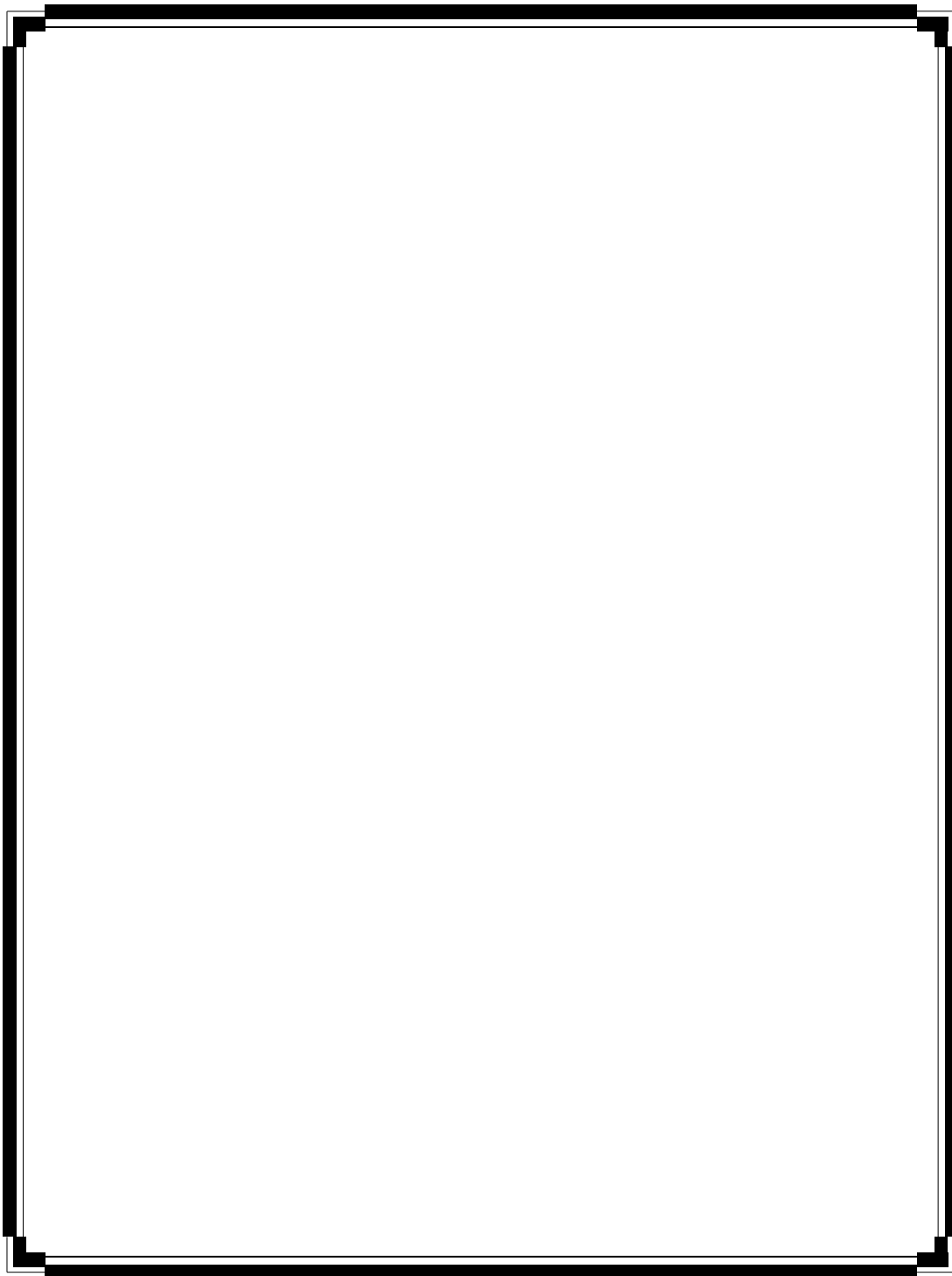
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بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ

وَمَا تَوْفِيقِي إِلَّا بِاللَّهِ عَلَيْهِ

تَوَكَّلْتُ وَإِلَيْهِ أُنِيبُ

صدق الله العظيم

(سورة هود آية ٨٨)

APPROVAL SHEET

**A Pharmacognostical study of *Celtis australis* L.
and *Celtis occidentalis* L. cultivated in Egypt**

Approved by:

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List of Abbreviations