

PHYTOCHEMICAL AND BIOLOGICAL STUDY ON CERTAIN PLANTS BELONGING TO FAMILY IRIDACEAE

Thesis submitted to

Faculty of Pharmacy
Ain Shams University

In partial fulfillment of the requirements
For the degree of
Doctor of philosophy in Pharmaceutical Sciences
(In Pharmacognosy)

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ACKNOWLEDGEMENT

First and foremost, I would like to thank God, the Almighty, for His guidance, providence and endless blessings that enabled me to proceed successfully throughout my research work.

I would like to express my profound gratitude and appreciation to the members of the advisory committee;

Prof. Dr. Abdel Nasser B. Singab, Professor of Pharmacognosy, Dean of Faculty of Pharmacy, Ain Shams University, who designed my research topic and closely followed up this work to its completion. I am very grateful for his patience and his willingness to help me despite his tremendous duties as a dean of the faculty and his teaching duties; he never turned me down when I asked for help. His active supervision, constructive criticism, insightful comments and constant guidance made this work possible. The manner in which Dr. Singab mentored me during my PhD work has challenged me to become a better scientist and to always seek quality rather than quantity research. I am really proud to be one of his students.

Dr. Mohamed M. El-Shazly, Lecturer of Pharmacognosy, Faculty of Pharmacy, Ain Shams University, for his devotion and kind supervision, conveying the true spirit of research. No words can express my deep gratitude to him for his valuable support and sincere guidance throughout my work. The brain-storming sessions and scientific discussions we had, helped me a lot throughout my work. His critical review has contributed a lot to the output of my write up. I aspire to become as successful, determined and ambitious as he is.

I wish to extend my immense appreciation and thanks to **Dr. Mohamed L. Ashour**, Lecturer of Pharmacognosy, Faculty of Pharmacy, Ain Shams University, for his continuous support, encouragement and valuable advice throughout my work and for hosting the GC-MS and GC-FID analysis in the Institute of Pharmacy and Molecular Biotechnology, Heidelberg University, Heidelberg, Germany.

Special thanks are due to **Prof. Dr. Hesham El-Beshbishy**, Professor of Biochemistry, Faculty of Applied Medical Sciences, Taibah University, Al-Madinah Al-Munawarah, Saudi Arabia, for hosting the *in vivo* hepatoprotective and antidiabetic studies; **Prof. Dr. Fang-Rong Chang**, Professor of Pharmacognosy and Director of the Graduate Institute of Natural Products, College of Pharmacy, Kaohsiung Medical University, Kaohsiung, Taiwan, for hosting the NMR analysis as well as the *in vitro* anti-allergic assay; **Prof. Dr. Michael Wink**, Professor of Biology, Head of Biology Department, Institute of Pharmacy and Molecular Biotechnology, Heidelberg University, Heidelberg, Germany, for hosting the GC-MS and GC-FID analysis; **Prof. Dr. Mei-Chin Lu**, Graduate Institute of Marine Biotechnology, National Dong Hwa University, Pingtung, Taiwan; the National

Museum of Marine Biology & Aquarium, Pingtung, Taiwan, for hosting the cytotoxicity studies.

My profound gratitude also goes to my colleague **Mohamed Saeed**, Assistant Lecturer of Pharmacognosy, Faculty of Pharmacy, Ain Shams University for carrying out the ESI-MS measurements at the Institute for Pharmaceutical Biology and Biotechnology, Heinrich-Heine University of Dusseldorf, Germany.

I would like to extend my profound gratitude and appreciation to my dear professors in the Department of Pharmacognosy, Faculty of Pharmacy, Ain Shams University;

No words can express my deep gratitude to **Prof. Dr. Mohamed M. Al-Azizi**, for setting an example to how a dedicated professor should be. His immense knowledge and unique character had a great impact on me in both academic and life aspects. His enthusiasm, dedication and commitment have always inspired me throughout my career.

Special thanks are due to **Assoc. Prof. Dr. Sherweit El-Ahmady**, Acting Head of Department, her spirit, devotion, passion and the utter desire to help and support have always been an outstanding inspiration to me; **Assoc. prof. Dr Omayama El-Dahashan** and **Dr. Rola Milad**, for their valuable advice, encouragement and friendly help; **Dr. Sherif Ebada**, **Dr. Eman Kamal** and **Dr. Haidy Gad** for their kind support and cooperation.

My deep gratitude goes to my friend and colleague **Dr. Fadia Youssef**, my lab mate, for her constant support and encouragement. Many thanks and appreciation is due to all my colleagues in the Department of Pharmacognosy, Faculty of Pharmacy, Ain Shams University, for their cooperation, support and the friendship we share.

I wish to extend my love and gratitude to my parents, **Fr. Shenouda Ayoub** and **Eng. Mona Bassilious** for their endless love, unconditional support and all the sacrifices they have made on my behalf. Their prayers for me were what sustained me that far. They have always inquired about my progress and encouraged me to do what I was most passionate about, scientific research, without their nurturing love, I would not have been here in the first place. I am also very grateful to my loving sister **Mariam Ayoub** for her loving support and continuous encouragement. If we ever had a family motto that would have been – *If there's a will, there's a way* – a philosophy of life I have been carrying with me every day. Special thanks are due to my grandmother for her endless love and unlimited kindness.

No words can express my deep gratitude and appreciation for my beloved husband **Dr. John Samir Abdou**. His immeasurable assistance, patience and understanding allowed me to focus my efforts on my work. Without his constant support and continuous encouragement none of this work would have been possible.

I am mostly thankful to my lovely daughters **Angelina** and **Karen**. Though now they are still very young to really comprehend how grateful I am to them, no matter how long my working days were, their cheering big smiles and loving hugs would melt away any exhaustion or disappointment, like they never existed. They have always been my source of everlasting joy.

“The LORD has done great things for us, and we are filled with joy.”

(Psalms 126:3)

Iriny M. Ayoub

Cairo, 2015

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