



Faculty of Science
Chemistry Department

Synthesis and Characterization of Novel Magnetic Nano-Materials and Studying Their Potential Application in Recovery of Metal Ions

Thesis Presented By

Saber Ibrahim Mohamed Moussa

Assistant Lecturer

*Nuclear Chemistry Department – Hot Laboratories
Center – Atomic Energy Authority*

Submitted To

*Faculty of Science, Ain Shams University,
for The Degree of Ph.D. of Science (Chemistry)*

2013



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Supervised By

Prof. E. A. saad

*Prof. of Inorganic and
Radiation Chem. Faculty of
Sci.– Ain Shams University*

Prof. H.F. Ghoneimy

*late Prof. of Physical Chem.
Hot Labs. Center- Atomic
Energy Authority*

Prof. N. A. Tadrous

*Prof. of Radio Chem. Hot
Labs. Center - Atomic
Energy Authority*

Prof. R.R. Sheha

*Head of Nucl. Chem. Dept.
Hot Labs. Center - Atomic
Energy Authority*

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M. Sc. Chemistry, 2007

Thesis supervisors

Signature

1- Prof. E. A. saad

*Prof. of Inorganic and Radiation
Chemistry - Faculty of Sci.– Ain
Shams University*

.....

2- Prof. H.F. Ghoneimy

*late Prof. of Physical Chem. - Hot Labs.
Center- Atomic Energy Authority*

.....

3- Prof. N. A. Tadrus

*Prof. of Radio Chem. Hot Labs.
Center - Atomic Energy Authority*

.....

4- Prof. R.R. Sheha

*Head of Nuclear Chem. Dept. Hot Labs.
Center - Atomic Energy Authority*

.....

Head of the Chemistry Department

Prof. Dr. Maged Shafik Antonious Nakhla

Acknowledgment

First, I am deeply thankful to my God "Allah", by the grace of whom, the progress and success of this work was possible.

*I would like to express my deep gratitude to **Prof. Dr. Ebtisam Ahmed Saad**, prof. of inorganic Chemistry, Faculty of Science, Ain Shams University; for his sponsorship and moral support of this work,*

*I am greatly indebted to **Prof. Dr. Reda Rashad Sheha**, Head of Nuclear Chemistry Department, Radioisotopes Production and Radiation Sources Division, Hot Laboratories Center, Atomic Energy Authority (EAEA); for suggesting the topics of this thesis, planning of the experimental work, effective supervision, valuable discussions, sincere advices, and continuous encouragement during all phases of carrying this work,*

*I would like to express my sincere appreciation to **Prof. Dr. Hussein Fouad Ghoneimy**, late Prof. of Physical Chemistry, and former Head of the Radioisotopes Production and Radiation Sources Division, Hot Laboratories Center, Atomic Energy Authority (EAEA); for his supervision.*

*Thanks are also expressed for **prof. Dr. Nadia Abdo Tadrous**, Emeritus Prof., Nuclear Chemistry Department, Hot Laboratories Center, Atomic Energy Authority (EAEA); for his help during the experimental work,*

*Thanks are also expressed for **prof. Dr. Mohamed Kamal Shehata**, Emeritus Prof., Nuclear Chemistry Department, Hot Laboratories Center, Atomic Energy Authority (EAEA); for his fatherhood sensation, continuous encouragement and reviewing initial manuscript.*

*The assistance and support of **all staff members** of the Nuclear Chemistry Department, notably of **Dr. Mamdoh Refaat** are deeply appreciated.*

Saber Ibrahim

I would like to express my deep thanks to

*My mother, my wife
and my children
Ibrahim, Abd-Allah and Aya*

for their unlimited support, and looking after me.

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