



Women's College for Arts
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**The POSSIBLE MODULATORY EFFECT OF A
PROTECTIVE AGENT AGAINST METHYLMERCURY
INDUCED DEVELOPMENTAL NEUROTOXICITY IN
RATS**

A thesis submitted

For

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"وَعَلَّمَكَ مَا لَمْ تَكُن تَعْلَمُ وَكَانَ فَضْلُ اللَّهِ
عَلَيْكَ عَظِيمًا"

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LIST OF CONTENTS

LIST OF TABLES.....	v
LIST OF FIGURES.....	vii
LIST OF ABBREVIATIONS.....	xiv
I - INTRODUCTION AND AIM OF THE WORK.....	1
II – REVIEW OF THE LITERATURE.....	8
A- The toxic effect of Methylmercury (MMC).....	8
1-Growth rate, mortality rate and external symptoms.....	8
2-Congenital malformations of fetuses.....	10
3- Brain study.....	13
3.1- Effect of MMC on oxidative stress.....	13
3.1.1- Effect of MMC on malondialdehyde (MDA) levels.....	13
3.1.2- Effect of MMC on Nitric oxide (NO).....	16
3.1.3-Effect of MMC on Total antioxidant activity (TAA).....	17
3.2- Effect of MMC on Na ⁺ / K ⁺ ATPase.....	21
3.3- Effect of MMC on Acetylcholinesterase (AChE).....	21
3.4- Effect of MMC on Ca ²⁺ level.....	22
3.5- Effect of MMC on DNA	23
3.6- Effect of MMC on monoamines.....	25
3.7- Effect of MMC on amino acids.....	26
4- Effect of MMC on behavior tests in neonates.....	27
4.1 Effect of MMC on open field test.....	27
4.2 – Effect of MMC on Morris test.....	30
5- Histological studies.....	32
B- Protective effect of naringenin.....	33
1-Protective effect of naringenin on body weight.....	34
2-Protective effect of naringenin on MDA levels.....	34
3- Protective effect of naringenin on NO content.....	37
4- Protective effect of naringenin on Total antioxidant activity (TAA)...	38
5- Protective effect of naringenin on Na ⁺ / K ⁺ ATPase.....	40
6- Protective effect of naringenin on acetylcholinestrace activity (AChE).....	41
7- Protective effect of naringenin on DNA fragmentation.....	41
8- Protective effect of naringenin on Monoamines.....	42
9-Protective effect of naringenin on amino acids.....	43
10- Protective effect of naringenin on behavior tests.....	43
11-Protective effect of naringenin on histological disorders.....	43
III-MATERIALS AND METHODS.....	46
A- Experimental animals.....	46
B- Chemicals.....	46
a) Methyl mercuric (II) chloride (MMC).....	46
b) The protective agent Naringenin.....	46
C- Experimental design.....	47

The first main group (fetuses group).....	48
a- Maternal toxicity and external symptoms.....	49
b- Growth rate.....	49
c- Teratological examinations.....	49
d- Biochemical investigations.....	50
1- Oxidative stress evaluation.....	50
-Determination of Malondialdehyde (MDA).....	51
-Determination of nitric oxide (NO).....	53
-Determination of Total Antioxidant Activity (TAA).....	55
2- Evaluation of DNA fragmentation in fetal brain.....	58
e- Histological examination.....	62
The second main group (pups group).....	62
Maternal and pups general observations.....	63
Effect of treatment on developing brain.....	63
1-Neurobehavioral assessments.....	63
A -Morris water maze test.....	63
B - Open field test.....	66
2- Neurochemicals assessments.....	69
A-The concentrations of monoamines in cortex, hippocampus and Midbrain...	69
B-The concentrations of amino acids in frontal cortex.....	71
C-The activities of Na^+ / K^+ ATPase and acetyl cholinesterase (AChE) in cortex and brain stem and the total protein.....	72
-Determination of Na^+ / K^+ ATPase.....	72
-Determination of Acetyl cholinesterase (AChE) activity.....	74
-Total protein estimation.....	76
3-Oxidative stress assessment.....	77
4-Determination of Calcium (Ca) concentration.....	77
5- Evaluation of DNA fragmentation	78
6-Histological Examination.....	78
D- Statistical analysis.....	79
IV- RESULTS	80
I- Maternal toxicity.....	80
a- External symptoms.....	80
b- Maternal mortality.....	80
c- Body weight gain.....	80
II- Pregnancy outcome.....	84
a- Uterine weight.....	84
b- The number of implantation sites.....	84
c- Percentage of abortion.....	84
d- Post implantation loss index.....	85
III- Effect of MMC treatment on fetuses.....	91
a-Fetal growth rate.....	91
b- Litter weight.....	92
c- Fetal Sex distribution.....	92

d- Viable fetuses.....	92
e- External abnormalities.....	93
f- Skeletal examination.....	103
IV-Biochemical investigation on fetuses.....	121
a- Oxidative stress.....	121
1- Lipid peroxidation (Malondialdehyde) (MDA).....	121
2- NO level.....	121
3-Total antioxidant activity (TAA).....	122
b- Percentage of fragmented DNA.....	122
V- Histological examination of fetal cerebral cortex.....	126
Effect of treatments on neonates.....	135
A. The clinical signs of toxicity.....	135
B. Effect of treatments on developing brain.....	139
I- Neurobehavioral evaluations.....	139
a - Morris water maze test.....	139
b- The open field test.....	144
II- Neurochemicals assessments.....	150
a- The concentrations of monoamines (NE, DA & 5-HT) in different brain areas	150
1- The concentrations of monoamines in frontal cortex.....	150
Levels of NE	150
Levels of DA.....	151
Levels of 5-HT.....	151
2- The concentrations of monoamines in Hippocampus.....	156
Level of NE.....	156
Levels of DA.....	156
Levels of 5-HT.....	157
3- The concentrations of monoamines in Midbrain.....	161
Level of NE.....	161
Levels of DA.....	161
Levels of 5-HT.....	162
b- The concentrations of amino acids (aspartic acid, glutamic acid and GABA) in brain frontal cortex.....	166
1- Level of Aspartic acid.....	166
2-Level of glutamic acid.....	166
3-Level of GABA.....	167
c- Na ⁺ / K ⁺ ATPase and cholinesterase (AChE) activities and calcium content in cortex and brain stem.....	171
1- Na ⁺ / K ⁺ ATPase activity.....	171
In cortex.....	171
In brain stem.....	172
2- Acetyl cholinesterase (AChE) activity.....	172
In cortex.....	172

In brain stem.....	173
3- Calcium (Ca ⁺²) content.....	173
In cortex.....	173
In brain stem.....	174
III-Oxidative stress (MDA and NO levels and TAA) in cortex and brain stem.....	180
1- Malondialdehyde (MDA) content.....	180
In cortex.....	180
In brain stem.....	181
2- Nitric Oxide (NO) level.....	181
In cortex.....	181
In brain stem.....	182
3-Total Antioxidant Activity (TAA).....	183
In cortex.....	183
In brain stem.....	183
IV- The percentage of DNA fragmentation.....	190
V- Histological examination of cerebral cortex.....	193
V- DISCUSSION AND CONCLUSION.....	202
VI-SUMMARY.....	248
VII-REFERENCES.....	257
VIII- ARABIC SUMMARY	

LIST OF TABLES

Table		Page
Table (1):	The average maternal body weight gain (g) in different experimental groups (mean $\pm$ S.E.).	82
Table (2):	The mean average of uterine weights (g), no. of implantation sites /litter, % of partial abortion and post implantation loss in mothers of different groups. (Mean $\pm$ S.E.).	86
Table (3):	The fetal body weight (g), fetal body length (cm), the litter weight (g), fetal sex distribution, live and dead fetuses and the % of external abnormalities/ litter in the different groups, Mean $\pm$ S.E.	94
Table (4):	The levels of MDA, NO, TAA and the percentage of DNA fragmentation in the brain of fetuses maternally treated with methylmercuric chloride (1 mg /kg /day) without or with naringenin (50 mg /kg /day). mean $\pm$ S.E.	123
Table(5):	The average weekly body weight (g) of both male and female pups in different experimental groups (mean $\pm$ S.E.).	136
Table (6):	The average changes in Morris water maze 1 st , 2 nd , 3 rd , 4 th , 5 th , 6 th , 7 th , 8 th trials and the no. of crossing over the platform position for male pups maternally treated with MMC (1 mg /kg /day) without or with naringenin (50 mg /kg /day). Mean value $\pm$ SE.	140
Table (7):	The average changes in Morris water maze 1 st , 2 nd , 3 rd , 4 th , 5 th , 6 th , 7 th , 8 th trials and the no. of crossing over the platform position for female pups maternally treated with MMC (1 mg /kg /day) without or with naringenin (50 mg /kg /day). Mean value $\pm$ SE.	141
Table (8):	The average changes in open field parameters ; cross lines, rearing frequency, entrance to the central area and freezing times of male and female pups maternally treated with MMC (1 mg /kg /day) without or with naringenin (50 mg /kg /day). Values are expressed as mean $\pm$ S.E.	146
Table (9):	The monoamines levels in frontal cortex in both male and female pups maternally treated with MMC (1 mg /kg /day) without or with naringenin (50 mg /kg /day). Mean value $\pm$ SE.	153

Table(10):	The monoamines levels in the hippocampus of male and female pups maternally treated with MMC (1 mg /kg /day) without or with naringenin (50 mg /kg /day). Mean value $\pm$ SE	158
Table (11) :	The monoamines levels in midbrain area in both male and female pups maternally treated with MMC (1 mg /kg /day) without or with naringenin (50 mg /kg /day). Mean value $\pm$ SE.	163
Table(12):	The amino acids levels (μ g/ g tissue) in frontal cortex in both male and female pups maternally treated with MMC (1 mg /kg /day) without or with naringenin (50 mg /kg /day). Mean value $\pm$ SE.	168
Table(13):	Na ⁺ K ⁺ ATPase and acetyl cholinesterase activities and Ca ⁺² content in cortex in both male and female pups maternally treated with MMC (1 mg /kg /day) without or with naringenin (50 mg /kg /day). Mean value $\pm$ SE.	175
Table(14):	Na ⁺ K ⁺ ATPase and acetyl cholinesterase activities and Ca ⁺² content in brain stem of both male and female pups maternall treated with MMC (1 mg /kg /day) without or with naringenin (50 mg /kg /day). Mean value $\pm$ SE.	176
Table(15):	The MDA and NO contents and TAA in cortex in both male and female pups maternally treated with MMC (1 mg /kg /day) without or with naringenin (50 mg /kg /day). Mean value $\pm$ SE.	185
Table (16):	The MDA and NO contents and TAA in brain stem in both male and female pups maternally treated with MMC (1 mg /kg /day) without or with naringenin (50 mg /kg /day). Mean value $\pm$ SE.	186
Table(17):	The percentage of DNA fragmentation in both male and female pups maternally treated with MMC (1 mg /kg /day) without or with naringenin (50 mg /kg /day). Mean value $\pm$ SE.	191

LIST OF FIGURES

Figure		page
Fig.(1) :	Histogram showing the average increase of maternal body weight gain for rats of control, naringenin, methylmercuric chloride, and methylmercuric chloride + naringenin groups.	83
Fig.(2) :	Histogram showing the uterine weight (g) of pregnant rats of control, naringenin, methylmercuric chloride, and methylmercuric chloride + naringenin groups.	87
Fig.(3) :	Histogram showing the implantation sites/ litter of rats of control, naringenin, methylmercuric chloride, and methylmercuric chloride + naringenin groups.	87
Fig.(4):	Histogram showing the percentage of abortion in rats of control, naringenin, methylmercuric chloride, and methylmercuric chloride + naringenin groups.	88
Fig.(5):	Histogram showing the post implantation loss in rats of control, naringenin, methylmercuric chloride, and methylmercuric chloride + naringenin groups.	88
Fig.(6):	Photograph of uteri of control, naringenin, methylmercuric chloride, and methylmercuric chloride + naringenin groups showing growth retardation and improvement of growth rate.	90
Fig.(7):	Photograph of uteri of control and methylmercuric chloride treated rat showing early resorbed fetus and partial abortion.	90
Fig.(8):	Histogram showing fetal body weight (g) of control, naringenin, methylmercuric chloride, and methylmercuric chloride + naringenin groups.	95
Fig.(9):	Histogram showing the fetal body length (cm) of control, naringenin, methylmercuric chloride, and methylmercuric chloride + naringenin groups.	95
Figs.(10):	Histogram showing the litter weight (g) of control, naringenin, methylmercuric chloride, and methylmercuric chloride + naringenin groups.	96
Fig.(11):	Histogram showing the percentage of external abnormalities/ litter of control, naringenin, methylmercuric chloride, and methylmercuric chloride + naringenin groups.	96