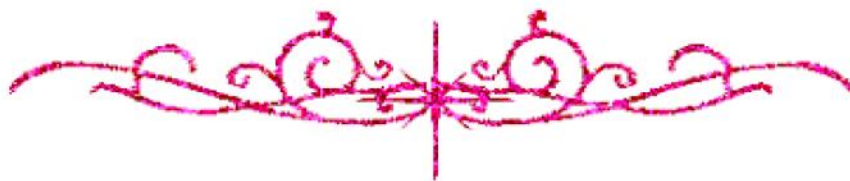


بسم الله الرحمن الرحيم





شبكة المعلومات الجامعية التوثيق الالكتروني والميكرو فيلم



جامعة عين شمس

التوثيق الإلكتروني والميكروفيلم

قسم

نقسم بالله العظيم أن المادة التي تم توثيقها وتسجيلها
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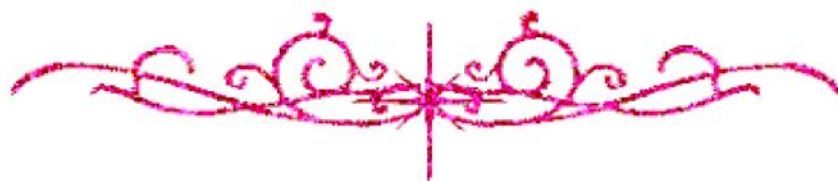


بعض الوثائق الأصلية تالفة





بالرسالة صفحات
لم ترد بالأصل





Biochemistry Department
Faculty of Science
Ain Shams University

Role of Vitamin D Receptor Polymorphism in Nephropathy of Type II Diabetes Mellitus patients

A thesis submitted for the degree of master in science as a partial fulfillment for requirements of the master degree in science

By

Noha Mohammed Sharkawy
(B.Sc. in Biochemistry/Chemistry, 2007)

Under Supervision of

Prof. Dr. Dina M. Seoudi
Professor of Biochemistry
Faculty of science - Ain Shams University

Dr. Abdel-Rahman B. Abdel-Ghaffar
Assistant Professor of Biochemistry
Faculty of science -Ain Shams University

Dr. Doaa Mostafa Gharib
Assistant Professor of Medical Biochemistry
and Molecular Biology
Faculty of Medicine- Cairo University

2020

بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ

وَعَلَيْكَ يَا كَلِيمُ

وَكَيْفَ كَانَ فَضْلُ اللَّهِ عَلَيْكَ عَظِيمًا

Dedication

At first, I praise and thank Allah for his greatness and for giving me the strength and courage to complete this thesis.

I would like to dedicate this work with all my deepest love and appreciation to every member of my faithful family, father, mother, brother, sister and my husband for their endless love, support and encouragement and for all my friends and those from whom I have learned, whatever and whenever they are.

Noha M. Sharkawy

Declaration

I declare that this thesis has been composed and the work recorded here has been done by me.

I have not been submitted for any other degree at thesis or any other university.

Noha M. Sharkawy

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Contents

Contents	
Title	Page
List of Abbreviations	I
List of Figures	IV
List of tables	VI
Abstract	VII
Chapter I : Introduction	
Introduction	1
Chapter II: Aim of work	
Aim of the work	4
Chapter III: Review of Literature	
1. Diabetes mellitus (DM)	5
1.1. Definition of DM	5
1.2. Classification of diabetes	6
1.2.1 Common types of diabetes	6
1.2.2. Type 2 diabetes mellitus (T2DM)	7
1.3. Epidemiology of T2 DM	8
1.3.1. Epidemiology of T2DM worldwide	8
1.3.2. Epidemiology of T2DM in Egypt	9
1.4. Diagnosis of diabetes	10
1.5. Risk factors for type 2 DM	12
1.6. Pathophysiology of T2DM	13
1.7. Diabetic complications	14
2. Kidney anatomy	15
3. Diabetic nephropathy (DN)	18
3.1. Definition of DN	18
3.2. Classification of DN	18
3.3. Risk factors for DN	20

Contents

3.4. Epidemiology of DN	21
3.4.1. Epidemiology o DN worldwide	21
3.4.2. Epidemiology of DN in Egypt	21
3.5. Diagnosis of DN	22
3.5.1. Used biomarkers and their predictive values	23
3.6. Molecular mechanisms involved in DN pathogenesis	24
3.6.1. Increased polyol pathway flux	25
3.6.2. Accumulations of AGEs	27
3.6.3. PKC pathway	28
3.6.4. Increased flux through the hexosamine pathway	30
3.6.5. Relationship of these pathways	31
4. Vitamin D	32
4.1. Sources of vitamin D	32
4.2. Epidemiology of vitamin D deficiency	33
4.3. Metabolism of vitamin D	34
4.4. Molecular action of vitamin D	35
4.4.1. Action of Vitamin D on T2DM	36
4.4.2. Mechanism of action of vitamin D on DN	42
5. Vitamin D receptor (VDR)	45
5.1. VDR gene location	45
5.2. Polymorphisms in VDR gene	46
5.2.1. Main types of VDR gene polymorphisms	47
Chapter IV: Subjects and Methods	
1. Subjects	50
1.1. Inclusion criteria	51
1.2. Exclusion criteria	51

Contents

2. Samples collection	52
2.1.Blood samples	52
2.2.Urine sample	52
3. Biochemical examinations	53
3.1.Quantitative determination of Glucose	53
3.2.Quantitative determination of Glycosylated Hemoglobin (HbA1C)	55
3.3.Measurement of HOMA-IR	58
3.4.Quantitative determination of Urea in serum	62
3.5. Quantitative determination of serum Creatinine	64
3.6. Quantitative determination of Microalbuminuria	66
4. Genetic investigations	68
4.1.DNA extraction	68
4.1.1.Spin-column technique of DNA extraction	68
4.1.2.Quantitative assessment of DNA	71
4.1.3.Qualitative assessment of DNA using gel electrophoresis	72
4.2.DNA amplification using PCR	72
4.3.Detection of PCR products using gel electrophoresis	75
4.4.Genotyping via PCR-RFLP	79
5. Statistical Analyses	80
Chapter V: Results	
1. Demographic and clinical characteristics of the studied subjects	81
2. Descriptive analysis of biochemical parameters	82

Contents

2.1.Determination of HbA1C	82
2.2.Determination of plasma Glucose	84
2.3.Determination of fasting plasma Insulin	86
2.4.Determination of HOMA-IR	88
3. Descriptive analysis of kidney function tests in the studied groups	89
3.1.Determination of Urea in serum	89
3.2. Determination of serum Creatinine	90
3.3. Determination of Microalbumin in urine	91
4. Genetic assay	92
4.1.DNA assessment and quantification	92
4.2.Genotyping of exon 9 gene polymorphism of VDR	93
4.2.1.Amplification of exon 9 of VDR gene by PCR	93
4.2.2. <i>Taq I</i> PCR-RFLP	94
4.3. <i>Taq I</i> genotypic and allelic distribution	95
Chapter VI: Discussion	101
Summary	123
Conclusion	125
References	126
المستخلص العربي	1
الملخص العربي	3

List of Abbreviations

Abbreviation	Full name
A	Absorbance
ACE	Angiotensin converting enzyme
ACR	Albumin to Creatinine ratio
AGEs	Advanced glycation end products
ANG I	Angiotensin I
ANOVA	One way analysis of variance
ATP	Adenosine triphosphate
BUN	Blood Urea Nitrogen
Conc.	Concentration
CKD	Chronic kidney disease
DAG	Diacylglycerol
DM	Diabetes mellitus
DNA	Deoxyribonucleic acid
dNTPs	Deoxynucleotide triphosphates
DN	Diabetic nephropathy
ECM	Extracellular matrix
EDTA	Ethylene diamine tetra acetic acid
ELISA	enzyme-linked immunosorbent assay
ESRD	End stage renal disease
ET-1	Endothelin-1
EtBr	Etidium Bromide
Fig.	Figure
G	Guanine
GDPH	glyceraldehyde-3-phosphate dehydrogenase
GFR	Glomerular filtration rate
GHb	Glycosylated Hb
GLUT4	Glucose transporter 4
HbA1C	Glycated Hemoglobin