



Dr Neda Mustafa Bogari

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| 2018 April | Associate Professor of Genetics , Department of Medical Genetics, Umm AlQura University, Makkah, Saudi Arabia |
| 2011-2018 | Academic coordinator - Department of Medical Genetics, Umm AlQura University, Makkah, Saudi Arabia |
| 2012 –2014 | Vice-Dean of Institute of Consulting Research and studies for scientific chairs, Umm AlQura University, Makkah, Saudi Arabia |
| 2009-2018 | Assistant Professor of Genetics , Department of Medical Genetics, Umm AlQura University, Makkah, Saudi Arabia |
| Personal Details | Date of Birth: 27/3/75 Nationality: Saudi, Place of Birth :Makkah |
| Education | |
| April 2002- January 2008 | PhD Human Genetics , University of Southampton, Subject: Cardiovascular Genetics, Project: A study of the relationship between ABCA1 genes polymorphisms, plasma lipid levels and risk of atherosclerosis |
| September- December 2007 Training (First Semester MSC Forensic Genetics) | DNA Profiling and population Genetics and Current Issues in Genomic Research / Forensic Genetics , Faculty of applied sciences, University of the west of England, Bristol. |
| 1994-2001 | Bachelor in Medical Sciences , College of Medicine, Umm AL-Qura University, Makkah, Saudi Arabia. |
| Publications | |
| Submission process | Next generation exome sequencing of paediatric asthma disease identifies 8 novel within 21 rare variants in candidate genes |
| 2019 | Whole exome sequencing detects novel variants in Saudi children diagnosed with eczema Neda M.Bogari, Amr A.Amin, Husni H.Rayes, AhmedAbdelmotelb, Faisal A. Al-Allaf, AnasDannoun, Hiba S. Al-Amodi, Anas A.Sedayo, HilalAlmalk, AmnaMoulana, RaniaBalkhair, FatmaJambi, FakhriahMadani, MwafaqAbutalib, Mohiuddin M.Taher, Abdellatif Bouazzaoui, AshwagAljohani, Mustafa N.Bogari, ImranAli Khan, J Infect Public Health. 2019 Jun 15 |
| 2019 | Exome Sequencing Studies for Kids with Non-Familial Food Allergy. Neda M Bogari, Amr A Amin, Husni H Rayes, Ahmed Abdelmotelb, Faisal A. Al- Allaf, Anas Dannoun, Mamdouh S Alhazmi, Saed A Basardah, Mohiuddin M Taher, Udaya Raja GK, Ashwag Aljohan, Soud Abdulraof A Khogeer, Mohamed Mahmoud Nour Eldein, Mohammad Adil, Mustafa Bogari, Imran Ali Khan, Khalid Khalaf Alharbi and Ahmed Fawzy Genet. Mol. Res. 18 (3) |
| 2019 | Genetic biomarkers predict susceptibility to autism spectrum disorder through interactive models of inheritance in a Saudi community , Nasser A. Elhawary, Mohammed T. Tayeb, Ikhlas A. Sindi, Nermeen Qutub, Mona Rashad, Ahmad Muftia, Arwa H. Arab, Asim Khogeer, Ezzeldin N. Elhawary, Anas Dannoun and Neda Bogari , Cogent Biology, Published: 16 April 2019 |
| 2019 | Whole Exome Sequencing Reveals Multiple Mutations in Uncommon Genes of Familial Hypercholesterolaemia , Mohammad Athar ^{1,2} , Faisal Amhed Al-Allaf ^{1,3} , Fahad Alnouri ⁴ , Mohiuddin Mohammed Taher ^{1,2} , Abdellatif Bouazzaoui ^{1,2} , Neda Mustafa Bogari , Wafa Mohammed Elbejrani ³ , Hani Saleh Faidah ⁵ , Zainularifeen Abduljaleel ^{1,2,*} J Cardiovasc Disease Res., 2018; 9(4):x-x |
| 2019 <u>eBook</u> | Vaccine-Drug interaction , <u>eBook- chapter 1</u> , El Rashedy AA, Bogari N , Maha A Hassan and Fawzy A, Humman Vaccines, avidscience. |
| 2019 | Whole Exome Sequencing: Novel Genetic Polymorphisms in Saudi Arabian Attention Deficit Hyperactivity Disorder (ADHD) Children. Neda Mostafa Bogari, Amr Ahmed Amin, Ashwag Aljohani, Ghida Dairi, Mahmoud Zaki El-Readi, Anas Dannoun, Udaya Raja, Mohammad Adil, Nermeen Qutub, Suhair Alhelfawi, Amal Alobaidi, Derar Alqudah, Hussain Banni, Safaa Yehia Eid, Huda Balto, Faisal Al-Allaf, Mohiuddin Taher, Hiba Saed Al-Amoodi, Ahmed Fawzy |
| 2019 | The Genetic Variant c.553G>T in the Lipoprotein A5 Effects on Lipid Profile Parameters Levels Neda M. Bogari, Ashwag Aljohani, Amr A. Amin, Faisal A. Al-Allaf, Anas Dannoun, Mohiuddin M. Taher, Atalla Elsayed, Dareen Ibrahim Rednah, Osama Elkhatee, Massimo Porqueddu, Francesco Alamanni, Ahmed Fawzy World Journal of Cardiovascular Diseases Vol.9 No.2 |
| 2019 | Identification of six novel factor viii gene variants using next generation sequencing and molecular dynamics simulation. Al-Allaf FA, Abduljaleel Z, Bogari NM , Owaidah TMA, Taher MM, Athar M, Elsendiony A, Abalkhail H, Abdellatif A, Elbejrani W, Bouazzaoui A. Acta Biochim Pol. 2019 Feb 22;66(1):23-31 |
| 2019 | A genetic variant c.553G > T (rs2075291) in the apolipoprotein A5 gene is associated with altered Triglycerides levels in Coronary artery disease (CAD) patients with lipid lowering drug Neda M. Bogari, Ashwag Aljohani, Amr A. Amin, Faisal A. Al-Allaf, Anas Dannoun, Mohiuddin M. Taher, Atalla |

Elsayed, Dareen Ibrahim Rednah, Osama Elkhatee, Massimo Porqueddu, Francesco Alamanni, Soud Abdulraof A. Khogeer, and Ahmed Fawzy. BMC Cardiovascular Disorder, 2019; 19: 2

2018 **Pattern of Thyroid Lesions in Western Region of Saudi Arabia: A Retrospective Analysis and Literature Review.** Saeed MI, Hassan AA, Butt ME, Baniyaseen KA, Siddiqui MI, Bogari NM, Al-Allaf FA, Taher MM. J Clin Med Res. 2018 Feb;10(2):106-116

2018 **Identification of a Novel ATM Missense Mutation by Next Generation Sequencing in Choroid Plexus Papilloma**
Poster: AMP Europe 2018, Rotterdam, April 30-2 May

2018 **Next Generation Sequencing on Ion Proton for Mutation Detection in Brain Tumors: Development of Molecular Pathology Assays in the Kingdom of Saudi Arabia**
Poster: AMP Europe 2018, Rotterdam, April 30-2 May

2017 **Molecular Analysis of Factor VIII and Factor IX Genes in Hemophilia Patients: Identification of Novel Mutations and Molecular Dynamics Studies** ·Faisal A. Al-Allaf, Mohiuddin M. Taher, Zainularifeen Abduljaleel, Abdellatif Bouazzaoui, Mohammed Athar, **Neda M. Bogari**, Halah A. Abalkhail Tarek MA. Owaida ·Journal of Clinical Medicine Research 9,4, April 2017, 317-331

2016 **Antisense Oligonucleotide Tool in Therapy for Polyglutamine Neurodegenerative Disorders, eBook, Bogari N and Fawzy A,** Genome Engineering, smgbooks.

2016 **Drug Design Targeting RNA, eBook, El Rashedy AA, Bogari N and Fawzy A,** Genome Engineering, smgbooks.

2016 **Obesity is associated with cardiac disease and the apolipoprotein A5 gene a genetic variant c.553G > T in is associated with an increased risk of coronary artery disease, Poster,** Obesity and chronic diseases, 25-27 July 2016 Las Vegas USA.

2016 **VULNERABILITY OF GENETIC VARIANTS TO THE RISK OF AUTISM AMONG SAUDI CHILDREN,** Translational Genomics Human Genome Meeting, Poster, 28 February-2 March 2016. Houston, USA

2016 **Prediction of Hepatic Fibrosis in Patients with Chronic Hepatitis C Genotype 4: A Non-Invasive Biochemical Analysis.** Amr Amin, Mohammed Mukhtar, Essam Noor Eldin, Fahd Gethami, Fayed Hafez, Sameer Fatani, Neda Bogari, Mohammed Noor Eldin, Abdullatif Babakr, Soud Khogeer and Ahmed Fawzy. Biosciences Biotechnology Research Asia 13(1):287-297.

2016 **Impact of next generation sequencing in glucose-6-phosphate dehydrogenase deficiency studies. Neda Bogari.** Bioinformatics, 2016

2015 **Molecular Genetic Diagnosis for a Family with Type I Spinal Muscular Atrophy (SMA) via analysis of Survival Motor Neuron (SMN) Gene. Neda M Bogari,** Fareed R Bogari, Husni H Rayes, Noha M Alqassimi, Huda M Balto, Anas I Dannoun, Raneem H Abushanab, Amr Ahmed Amin, Soud Abdulraof AKhogeer, Rami Nassir, Azza M Abdel-Latif, Ahmed Fawzy. Journal of Rare Disorders: Diagnosis & Therapy. 2015;1 3:21

2015 **A novel SNP in 3' UTR of INS gene: A case report of neonatal diabetes mellitus. Bogari NM,** Rayes HH, Mostafa F, Abdel-Latif AM, Ramadan A, Al-Allaf FA, Taher MM, Fawzy A. Diabetes Res Clin Pract. 2015 Jul 10

- 2015 **No association of apolipoprotein B gene polymorphism and blood lipids in obese Egyptian subjects.** Bogari NM, Abdel-Latif AM, Hassan MA, Ramadan A, Fawzy A.J Negat Results Biomed, 2015 Mar 18;14(1):7 PMID: 25889118
- 2014 **Apolipoprotein B (XbaI) allele frequencies in an Egyptian Population: impact on blood lipids.** Neda M. Bogari ,Azza M. Abdel-Latif and Maha A. Hassan , Ahmed Fawzy,Int J Biol Med Res. 2014; 5(2): 3981-3987
- 2014 **Transporter TAP1-637G and Immunoproteasome PSM89-60H Variants Influence the Risk of Developing Vitiligo in the Saudi Population** Nasser AttiaElhawary, **Neda Bogari**, EssamHussienJiffri, Mona Rashad, AbdulhamidFatani, Mohammed Tayeb Dis Markers. 2014; 2014: 260732. Published online 2014 December 7. doi: 10.1155/2014/260732PMCID: PMC4273470
- 2013 **Genetic Polymorphisms of Glutathione S-Transferase Genes and Risk of Colorectal Cancer in the Saudi Population**Mohammed T. Tayeb1, Ahmad O. Babalghith 1, Ahmed Fawzy2, Suleiman Jastaniah3, **Neda M. Bogari**UQU Med. J Vol. 4, No.2, pp. 61-71 (2013)
- 2013 **The MTHFR 677T Allele May Influence the Severity and Biochemical Risk Factors of Alzheimer's Disease in an Egyptian Population.**Elhawary NA, Hewedi D, Arab A, Teama S, Shaibah H, Tayeb MT, **Bogari N.** PMID: 24223459
- 2011 **Null genetic risk of ACE gene polymorphisms with nephropathy in type 1 diabetes among Egyptian population.** Elhawary N, **Bogari N**, Rashad M, Tayeb M. The Egyptian journal of Medical Human Genetics 2011 ;12: 187-192.
- 2010 **Molecular updating of β -thalassemia mutations in the Upper Egyptian population.** Jiffri EH, **Bogari N**, Zidan KH, Teama S, Elhawary NA. Hemoglobin. 2010;34(6):538-47.
- 2007 **Functional polymorphism in ABCA1 influences age of symptom onset in coronary artery disease patients, Human molecular genetics,** Kyriakou T1, Pontefract DE, Viturro E, Hodgkinson CP, Laxton RC, **Bogari N**, Cooper G, Davies M, Giblett J, Day IN, Simpson IA, Albrecht C, Ye S. 2007 Jun 15;16(12):1412-22PMID: 17412755
- Grand**
- 2016 Biochemical and Genetic study of Attention Deficit Hyperactivity Disorder (ADHD) in Saudi Arabia children by **Deanship of Scientific Research UQU**
- 2014 *Genetic determinants for allergy and asthma in a Saudi Arabian population* by **King Abd-Elaziz for science and technology**
- 2013 *Susceptibility of candidate genes to the risk of autism among Saudi children* by **King Abd-Elaziz for science and technology**
- 2012 A study of the association of the R219K polymorphism in the ATP-binding cassette transporter A1 (ABCA1) gene with atherosclerosis, coronary artery disease and hyperlipidaemia in a Saudi Arabian population by **Institute of Scientific research, Umm-Alqura university**
- 2012 Gene polymorphisms associated with vitiligo in Saudi community by **Institute of Scientific research, Umm-Alqura university**
- 2011 Study of genetic mutations tumor necrosis gene for pilgrims in Mecca and the holy sites by **Custodian of the Two Holy Mosques Institute for Hajj Research.**
- Relevant Industrial Experience**
- 2016 **One Year University of Southampton, Southampton, UK Professor John Holloway Lab**
- 18 June 2013 **One month Johns-Hopkins Medical School, Baltimore, MD, USA Professor Barbara Sollner-Webb Lab**
- 18 June 2011 **Two month Johns-Hopkins Medical School, Baltimore, MD, USA Professor Barbara Sollner-Webb Lab**
- 2011-2018 **Academic Coordinator** of Genetic department of the first year foundation, Umm AlQura University, Makkah, Saudi Arabia
- 2011-2017 **Academic Coordinator** of Genetic department, Umm AlQura University, Makkah, Saudi Arabia
- 2010-2015 **Quality Coordinator**, School of Medicine, Umm AlQura University, Makkah, Saudi Arabia
- 2001 Three month AL Hada Armed Forces Hospital
- 2001 Three month Hera General Hospital
- 2000 Two month Toxicology Center Makkah
- 1999 Two month Toxicology Center Makkah
- Conference, Workshops & Courses**
- 25-27/07/2016 **Obesity is associated with cardiac disease and the apolipoprotein A5 gene a genetic variant c.553G > T in is associated with an increased risk of coronary artery disease,** Obesity and Chronic Diseases, Las Vegas USA
- 28/02/2016 **VULNERABILITY OF GENETIC VARIANTS TO THE RISK OF AUTISM AMONG SAUDI CHILDREN,** Human Genome Meeting (HGM 2016), Huston USA
- 17-18 Sep 2009 **Genetics of Complex Diseases,** British Atherosclerosis Society, Queens College, Cambridge, UK
- 25- 27 July 2006 **Introductory Statistics for Biologists,** University of Oxford
- 17- 20 July 2006 **Techniques & Applications of Molecular Biology** A Course for Medical Practitioners, University of Warwick
- 23-24 June 2006 **High Density Lipoproteins:** from basic science to therapeutic applications, the XIV international Symposium on Atherosclerosis. Parma, Italy
- 18- 22 June 2006 **The XIV international Symposium on Atherosclerosis,** Rome, Italy
- 15 May 2006 **Genetic advances in adult and pediatric cardiology,** The Royal Society of Medicine, London
- 18-19 April 2005 **Clinical Molecular Genetics Society Conference,** Post graduate Medical Centre, Salisbury District Hospital, R/N:148
- 14 April 2005 **Statistical Tools for Chemists,** LGC, London
- 1-2 February 2005 **Professional Skills,** Biological Sciences University of Southampton

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| 25-26 May 2004 | Presentation in Faculty Postgraduate Conference, Southampton General Hospital, (A study of ABCA1 gene variations in relation to coronary artery disease) |
| 26 th June 2003 | Poster in Faculty Postgraduate Conference, Southampton General Hospital, (Variation in the ABCA1 gene and atherosclerosis) |
| 2001 | Total Quality Management Basic Concepts, AL Hada Armed, Forces Hospital, KSA |
| 2001 | Total Quality Management in Health Care , AL Hada Armed, Forces Hospital, KSA |
| 2001 | Rift Valley Fever , AL Hada Armed, Forces Hospital, KSA |
| Working | |
| 2006-2007 | Head teacher , Saudi Student School, Southampton |
| 1997-2001 | South Asia Institute of Hajj |
| Membership | |
| Since 2002 | British Society for Genetic Medicine, Membership number: 23123 |
| Since 2002 | Association for Clinical Genomic Science, Membership number: P0001353 |
| Since 2009 | European Society of Human Genetics, Membership number: 3591 |
| Since 2017 | The American Society of Human Genetics, Membership number: 354862 |
| Since 2018 | Genetics Society, Membership number: P0004915 |