

Curriculum Vitae

Personal information

First name / Surname: Abdelbasset Amara

Date and place of birth: 13/08/1984 in Sousse, Tunisia

Nationality: Tunisian

Gender: Male

Marital status: Married

Qualification: PhD in Biotechnology and Biological Sciences

Speciality: Human Genetics and Molecular Biology

Grade: Assistant Professor

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Address: 62 Sidi Bouraoui Street, 4000 Sousse - Tunisia.

Education

2015: PhD degree in Biotechnology and Biological Sciences. With a <u>very honourable</u> <u>appreciation.</u> University of Monastir, Tunisia

2009: Master degree in Genetics. With a very good appreciation. University of Monastir,

Tunisia

2007: Bachelor degree in Medical Biotechnology Sciences and Techniques. With good

<u>appreciation</u>. University of Monastir, Tunisia **2003: High School diploma.** Sousse, Tunisia

Personal statement

I was trained as a geneticist and a molecular biologist in Tunisia and France. My research studies were performed in the laboratories of cytogenetics and molecular genetics, the unit of molecular endocrinology in Farhat Hached university hospital in Sousse, Tunisia and the unit of genomics and metabolic diseases in Pasteur Institute of Lille in France. The aim of my studies was the identification of the genetic causes of several rare genetic diseases such as MODY diabetes and spinal muscular atrophy in Tunisia and Africa. Hence contributing in exploring the genetic characteristics of Tunisians and Africans and developing strategies for the molecular diagnosis of these rare genetic diseases.

Positions and employment

- > From 2016 to current
- Assistant professor, Faculty of applied medical sciences, Northern Border university, KSA
 - **>** 2015-2016

- Head of studies and trainings, Private Higher Institute of Nursing of Sousse, Tunisia.
- Assistant professor, Private Higher Institute of Nursing of Sousse, Tunisia.
 - > 2014
- Lecturer, Private Higher School of engineering and applied technology of Sousse, Tunisia.

Academic and administrative tasks

- Rapporteur of Medical laboratory technology department council, Faculty of medical applied sciences, University of Northern Border region, KSA.
- Member in the scientific research committee, Faculty of medical applied sciences, University of Northern Border region, KSA
- Member in the quality committee, Faculty of medical applied sciences, University of Northern Border region, KSA
- Head of studies and trainings in higher institute of nursing of Sousse, Tunisia
- Experience in teaching human genetics and molecular biology, medical technical skills, clinical biology, biochemistry, anatomy, physiology, bioinformatics and biomedical research
- Supervising and evaluating dissertations of undergraduate studies

Scientific interchange and workshops

- "Familial early-onset diabetes is not MODY in Tunisian population". 1st International Symposium of Young Researchers in Biology. Monastir, Tunisia – May 12-14, 2016. Oral presentation.
- "Familial early-onset diabetes is different from classical MODY in several Tunisian families".
 International forum of biological science and biotechnology, Sousse, Tunisia Mars 25-28, 2013.
 Poster.
- Participation at EGID symposium (European Genomic Institute for Diabetes) Lille, France, October 10-11, 2012.
- "Absence de mutations des gènes HNF1A, HNF4A et INS chez dix patients tunisiens avec un diabète de type MODY". Congress of the Francophone Society of Diabetes, Geneva, Switzerland Mars 22-25, 2011. Poster.
- "Effect of copy number variation of *SMN1* neighbouring genes on SMA phenotype in Tunisian patients". European Human Genetics Conference, **Gothenburg, Sweden June 12-15, 2010. Poster.**
- "Correlation of *SMN2*, *NAIP*, *p44*, *H4F5* and *occludin* gene copy number with spinal muscular atrophy phenotype in Tunisian patients". Eighth days of biotechnology, **Sousse**, **Tunisia December 2009**.

Poster.

• Participation in workshop called "Statistical analysis of data using SPSS software", Sousse-Tunisia 2009.

Technical skills and competences

- -Techniques of molecular biology such as: DNA extraction, simplex and multiplex PCR, Triple primers PCR, RFLP-PCR, MLPA, Southern blot, Sequencing, Gel and Capillary electrophoresis, FISH, Karyotype and Cell culture.
- -Techniques of next generation sequencing: Whole exome sequencing (Illumina platform), RainDance technology for targeted sequencing and whole genome genotyping arrays.
- -Techniques of immunoprecipitation and immunomarquage as ELISA.

Computer and bioinformatics skills

Microsoft Word, Excel, Power Point, Internet.

Bioinformatics tools and databases (Ensembl, UCSC genome browser, NCBI, Uniprot...), softwares (SPSS, SeqScape, GeneMarker, genome studio, alamut, genome browser...)

Linkage analysis, Genotyping data analysis

Languages

Arabic Native speaker

French very good level in speaking and writing

English very good level in speaking and writing

Spanish medium level in speaking and writing

Membership in societies

Member in the ATDOCS: "The Tunisian society of PhD and PhD students"

Member in the ATSB: "Tunisian society of Biological Science"

List of Publications

☐ Correlation of epigenetic markers with glioblastoma survival and molecular subtypes. Trabelsi S, Amara A et al. submitted 2022
☐ Clinical and Genetic Heterogeneity of Familial Early-Onset Diabetes: Case of Six Tunisian Patients Suspected of Maturity-Onset Diabetes of the Young 12. Amara A, Trabelsi S et al. International journal of medical research and health sciences. 2020, 9(2): 25-31.
□ Difficulties in the diagnosis aetiology of A- β- ketosis-prone diabetes in a North-African adult. Slim I, Amara A, Baba A et al. Saudi Journal of Biomedical Research. 2016; 1 (1): 1-5.
☐ Multiple Self-healing Palmoplantar Carcinoma (MSPC): a Familial Predisposition to Skin Cancer with Primary Palmoplantar and Conjunctival Lesions. Mamaï O, Boussofara L, Denguezli M, Escande-Beillard N, Kraeim W, Merriman B, Ben Charfeddine I, Stevanin G, Bouraoui S, Amara A, Mili A, Nouira R, H'mida D, Sriha B, Gribaa M, Saad A, Reversade B. The Journal of investigative dermatology 2015; (135): 304–308.
☐ Genetics of low spinal muscular atrophy carrier frequency in sub-Saharan Africa. Sangaré M, Hendrickson B, Sango HA, Chen K, Nofziger J, Amara A, Dutra A, Schindler AB et al. Ann Neurol. 2014; 75 (4): 525–532.
□ Familial early-onset diabetes is not a typical MODY in several Tunisian patients. Amara A, Chadli-Chaieb M, Ghezaiel H, Philippe J, Brahem R, Dechaume A, Saad A, Chaieb L, Frogue P, Gribaa M, Vaxillaire M. Tunis Med. 2012; 90(12):882-7.
□Correlation of SMN2, NAIP, p44, H4F5 and Occludin genes copy number with spinal muscular atrophy phenotype in Tunisian patients. A. Amara, L. Adala, I. Ben Charfeddine, O. Mamaï, A. Mili, T. Ben lazreg, D. H'mida, F. Amri, N. Salem, L. Boughamoura, A. Saad, M. Gribaa. Eur J Paediatr Neuro. 2012; 16(2):167-74.

□A c.3216_3217delGA mutation in AGL gene in Tunisian patients with a glycogen storage disease type III: evidence of a founder effect. A. Mili, I. Ben Charfeddine, A. Amara, O.

Mamaï, L. Adala, T. Ben Lazreg, J. Bouguila, A. Saad, K. Limem and M. Gribaa. Clinical genetics. 2012; 82(6):534-9.

□Reduction of palmoplantar keratoderma Buschke-Fischer-Brauer locus to only 0.967

Mb. Mamaï O, Boussofara L, Adala L, Amara A, Ben Charfeddine I, Ghariani N, Sriha B,

Denguezli M, Mili A, Belazreg T, Saad A, Gribaa M. J Dermatol Sci. 2012; 67(3):2102.

□Two novel CYP11B1 mutations in congenital adrenal hyperplasia due to steroid 11β hydroxylase deficiency in a Tunisian family. Ben Charfeddine I, Riepe FG, Kahloul N, Kulle AE, Adala L, Mamaï O, Amara A, Mili A, Amri F, Saad A, Holterhus PM, Gribaa M. Gen Comp Endocrinol. 2012; 175(3):514-8.

□Molecular and biochemical characterization of Tunisian patients with glycogen storage disease type III. Mili A, Ben Charfeddine I, Mamaï O, Cherif W, Adala L, Amara A, Pagliarani S, Lucchiarri S, Ayadi A, Tebib N, Harbi A, Bouguila J, H'Mida D, Saad A, Limem K, Comi GP, Gribaa M. J Hum Genet. 2012; 57(3):170-5.

□Cytogenetic and Molecular Aspects of Absolute Teratozoospermia: Comparison between Polymorphic and Monomorphic Forms. S. Brahem, H. Elghezal, H. Ghédir, H. Landolsi, A. Amara, S. Ibala, M. Gribaa, A. Saad and M. Mehdi. Urology 2011;78(6):1313-9

□*Molecular characterization of piebaldism in a Tunisian family*. Kerkeni E, Boubaker S, Sfar S, Bizid M, Besbes H, Bouaziz S, Ghedira N, <u>Amara A</u>, Manoubi W, Gribaa M, Monastiri K. **Pathologie-biologie** 2015; (In press)

□Genetic study in a Tunisian family revealed IVS1+1G>A mutation in the CHM gene. Ben Charfeddine I, Ben Lazreg T, Ben Rayana N, Amara A, Mamaï O, Knani L, Mili A, M'sakni A, Saad A, Ben Hadj Hamida F, Gribaa M. Ann Biol Clin 2015; (In press).

□Multiplex Minisequencing of the HBB Gene: A Rapid Strategy to Confirm the Most Frequent b-Thalassemia Mutations in the Tunisian Population. Ben Charfeddine I, Ben Lazreg T, M'sakni A, Amara A, Mlika A, et al. Hemoglobin 2015; (In press).