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Research Interests: Molecular Genetics, Molecular Evolution					
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Education					
Date	Degree	Duration	Institution	Country/City	Major
1994	M.D.	6.5	Shahid Beheshti University of Medical Sciences,	Iran	Medicine

2000	Ph.D.	4.5	University of London	UK	Molecular Biology
Faculty member					
Year	Position	Duration	Institution/Course		Location
2002	Assistant Prof.	5 years	University of Social Welfare and Rehabilitation Sciences,		Tehran Iran
2007	Associate Prof.	9 years	University of Social Welfare and Rehabilitation Sciences,		Tehran Iran
2016	Professor	-present	University of Social Welfare and Rehabilitation Sciences,		Tehran Iran
Field of Specialization					
- - - -					
Language Ability					
- English -					
Teaching Experience					
Immunogenetics					
Cancer Genetics					
Molecular Medicine					

Symposia/Programs/Workshops				
Year	Position	Duration	Institution/Course	Location
Research Experience				
Year	Position	Institution/Course	Location	
Molecular Genetics	Principle Investigator	University of Social Welfare and Rehabilitation Sciences	Tehran, Iran	
Administrative Responsibilities				
Year	Position	Institution/Event	Location	
2016-present	IRJ Editor-in-Chief	University of Social Welfare and Rehabilitation Sciences	Tehran, Iran.	
Academic and Scientific Responsibilities				
Year	Responsibility			
Faculty Member	2002-present			
Scientific Membership				
Year	Association, Society	Location		

Publications

Genome-wide prediction and prioritization of human aging genes by data fusion: a machine learning approach. Arabfard M, **Ohadi M**, Rezaei Tabar V, Delbari A, Kavousi K. BMC Genomics. 2019 Nov 9;20(1):832. doi: 10.1186/s12864-019-6140-0.

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Year	Journal name	Article title	Vol. & Page

2. Books

Year	Book title	Publisher	Location
2005	Dictionary of Gene Technology	Nashr e Jaame-eh Negar	Tehran, Iran

Journal Reviews / Editorial Board

From	Journal Title	Location
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Research Paper Presented in Conferences / Seminars			
Year	Conference/Seminar	Paper title	Location