

<i>Surname</i>	<i>First name</i>	<i>Title</i>	<i>Date ofbirth</i>	<i>Gender</i>
<b>Najmabadi</b>	<b>Hossein</b>	<b>Prof.</b>	<b>28.08.1957</b>	<b>Male</b>

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University of Social Welfare and Rehabilitation Sciences  
Kodakyar Ave., Daneshjo Blvd., Evin,  
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#### ACADEMIC EDUCATION

<i>Field ofstudy</i>	<i>College / University (Country)</i>	<i>Degree</i>
Biology (1979-1983)	University of North Texas, Denton, Texas	B.S. Biology
Medical Technology (1980-1984)	University of North Texas, Denton, Texas	B.S. Medical technology

#### SCIENTIFIC DEGREES

<i>Degree</i>	<i>Field ofstudy</i>	<i>College / University (Country)</i>	<i>Y. of Graduation</i>
PhD	Molecular Biology, Double minor in Human Genetics & Biochemistry	University of North Texas, Denton, Texas	1989

#### PROFESSIONAL EXPERIENCE

<i>Period</i>	<i>Position / Function</i>	<i>Institution</i>
2006-present	Professor, Head & Director of Genetics Research Center (GRC)	Genetics Research Center, University of Social Welfare & Rehabilitation Sciences, Tehran, Iran
2002 - 2006	Associate Professor, Head & Director of GRC	Genetics Research Center, University of Social Welfare & Rehabilitation Sciences, Tehran, Iran
1996-2002	Assistant Professor, Head & Director of GRC	Genetics Research Center, University of Social Welfare & Rehabilitation Sciences, Tehran, Iran
1995-1996	Assistant Professor of Medicine	Charles Drew University of Medicine & Science – UCLA, Los Angeles, CA,USA
1990-1995	Postdoctoral fellow	Harbor – UCLA Medical Center, Torrance, CA, USA

#### MISCELLANEOUS

2017- present	Establishment & Director, Iranian Population Database "Iranome ( <a href="http://www.iranome.com">www.iranome.com</a> )
2011 - present	Editorial Board, Archive of Iranian Medicine
2011- present	Editorial Board, Clinical Genetics
2011-Present	Educational Board (PhD Degree), University of Social Welfare and Rehabilitation Sciences
2010 -present	Member, Association for Molecular Pathology (AMP)
2006-Present	Member, European Society of Human Genetics (ESHG)
2003-Present	Establishment & Director, Iranian Human Gene Bank
1996 - Present	Director of Molecular Division, Kariminejad-NajmabadiPhatology and Genetics Center
1996 - present	Member, American Society of Human Genetics (ASHG)
1980 - present	Member, American Society of Clinical Pathology (ASCP)

## MOST IMPORTANT PUBLICATIONS

- 1- Neishabury M, Mehri M, Fattahi Z, **Najmabadi H**, Azarkeivan A. Novel variants in Iranian individuals suspected to have inherited red blood cell disorders, including bone marrow failure syndromes. *Haematologica* 2020; 105(1): e1–e4.
- 2- Mehrjoo Z, Fattahi Z, Beheshtian M, Mohseni M, Poustchi H, Ardalani F, Jalalvand K, Arzhanghi S, Mohammadi Z, Khoshbakht S, Najafi F, Nikuei P, Haddadi M, Zohrehvand E, Oladnabi M, Mohammadzadeh A, Jafari MH, Akhtarkhavari T, Gooshki ES, Haghdoost A, Najafipour R, Niestroj LM, Helwing B, Gossmann Y, Toliat MR, Malekzadeh R, Nürnberg P, Kahrizi K, **Najmabadi H**, Nothnagel M. Distinct genetic variation and heterogeneity of the Iranian population. *PLoS Genet.* 2019; 15(9):e1008385.
- 3- Fattahi Z, Beheshtian M, Mohseni M, Poustchi H, Sellars E, Nezhadi SH, Amini A, Arzhanghi S, Jalalvand K, Jamali P, Mohammadi Z, Davarnia B, Nikuei P, Oladnabi M, Mohammadzadeh A, Zohrehvand E, Nejatizadeh A, Shekari M, Bagherzadeh M, Shamsi-Gooshki E, Börno S, Timmermann B, Haghdoost A, Najafipour R, Khorram Khorshid HR, Kahrizi K, Malekzadeh R, Akbari MR, **Najmabadi H**. Iranome: A catalog of genomic variations in the Iranian population. *Hum Mutat.* 2019; 40(11): 1968-84.
- 4- Kahrizi K, Hu H, Hosseini M, Kalscheuer VM, Fattahi Z, Beheshtian M, Suckow V, Mohseni M, Lipkowitz B, Mehvari S, Mehrjoo Z, Akhtarkhavari T, Ghaderi Zh, Rahimi M, Arzhanghi S, Jamali P, Falahat Chian M, Nikuei P, Sabbagh Kermani F, Sadeghinia F, Jazayeri R, Tonekaboni SH, Khoshaeen A, Habibi H, Pourfatemi F, Mojahedi F, Khodaie-Ardakani MR, Najafipour R, Wienker TF, **Najmabadi H**, Ropers H-H. Effect of inbreeding on intellectual disability revisited by trio sequencing. *Clin Genet.* 2019; 95(1):151-9.
- 5- Inanloo Rahatloo K, Peymani F, Kahrizi K, **Najmabadi H**. Whole-transcriptome analysis reveals dysregulation of actin-cytoskeleton pathway in intellectual disability patients. *Neuroscience.* 2019; 404: 423-44.
- 6- Palmer EE, Stuhlmann T, Weinert S, Haan E, Van Esch H, Holvoet M, Boyle J, Leffler M, Raynaud M, Moraine C, Van Bokhoven H, Kleefstra T, Kahrizi K, **Najmabadi H**, Ropers H-H, Delgado MR, Sirsi D, Golla S, Sommer A, Pietryga MP, Chung WK, Wynn J, Rohena L, Bernardo E, Hamlin D, Faux BM, Grange DK, Manwaring L, Tolmie J, Joss S, Study DDD, Cobben JM, Duijkers FAM, Goehring JM, Challman TD, Hennig F, Fischer U, Grimme A, Suckow V, Musante L, Nicholl J, Shaw M, Lodh SP, Niu Z, Rosenfeld JA, Stankiewicz P, Jentsch TJ, Geetz J, Field M, Kalscheuer VM. De novo and inherited mutations in the X-linked gene CLCN4 are associated with syndromic intellectual disability and behavior and seizure disorders in males and females. *Molecular Psychiatry.* 2018; 23(2):222-30.
- 7- Hu H, Kahrizi K, Musante L, Fattahi Z, Herwig R, Hosseini M, Oppitz C, Abedini SS, Suckow V, Larti F, Beheshtian M, Lipkowitz B, Akhtarkhavari T, Mehvari S, Otto S, Mohseni M, Arzhanghi S, Jamali P, Mojahedi F, Taghdiri M, Papari E, Soltani Banavandi MJ, Akbari S, Tonekaboni SH, Dehghani H, Ebrahimpour MR, Bader I, Davarnia B, Cohen M, Khodaei H, Albrecht B, Azimi S, Zirn B, Bastami M, Wiczorek D, Bahrami G, Keleman K, Vahid LN, Tzschach A, Gärtner J, Gillissen-Kaesbach G, Varaghchi JR, Timmermann B, Pourfatemi F, Jankhah A, Chen W, Nikuei P, Kalscheuer VM, Oladnabi M, Wienker TF, Ropers H-H, **Najmabadi H**. Genetics of intellectual disability in consanguineous families. *Molecular Psychiatry.* 2018; 24(7):1027-39.
- 8- **Najmabadi H**, Hu H, Garshasbi M, Zemojtel T, Abedini SS, Chen W, Hosseini M, Behjati F, Haas S, Jamali P, Zecha A, Mohseni M, Püttmann L, Vahid LN, Jensen C, Moheb LA, Bienek M, Larti F, Mueller I, Weissmann R, Darvish H, Wrogemann K, Hadavi V, Lipkowitz B, Esmaeeli-Nieh S, Wiczorek D, Kariminejad R, Firouzabadi SG, Cohen M, Fattahi Z, Rost I, Mojahedi F, Hertzberg C, Dehghan A, Rajab A, Banavandi MJS, Hoffer J, Falah M, Musante L, Kalscheuer V, Ullmann R, Kuss AW, Tzschach A, Kahrizi K, Ropers HH. Deep sequencing reveals 50 novel genes for recessive cognitive disorders. *Nature.* 2011; 478:57-63.
- 9- Pak C, Garshasbi M, Kahrizi K, Gross C, Apponi LH, Noto JJ, Kelly SM, Leung SW, Tzschach A, Behjati F, Abedini SS, Mohseni M, Jensen LR, Hu H, Huang B, Stahley SN, Liu G, Williams KR, Burdick S, Feng Y, Sanyal S, Bassell GJ, Ropers H-H, **Najmabadi H**, Corbett AH, Moberg KH, Kuss AW. Mutation of the conserved polyadenosine RNA binding protein, ZC3H14/dNab2, impairs neural function in Drosophila and humans. *PNAS.* 2011; 108:12390-5.
- 10- Mani A, Radhakrishnan J, Wang H, Mani A, Mani M-A, Nelson-Williams C, Carew KS, Mane S, **Najmabadi H**, Wu D, Lifton RP. LRP6 mutation in a family with early coronary disease and metabolic risk factors. *Science.* 2007; 315:1278-82.

## OTHER PUBLICATIONS

1. Mehrjoo Z, Kahrizi K, Mohseni M, Akbari M, Arzhanghi S, Jalalvand K, **Najmabadi H**, Farhadi M, Mohseni M, Asghari A, Mohebbi S, Daneshi A. Limbic system associated membrane protein mutation in an Iranian family diagnosed with Ménière's disease. *Arch Iran Med.* 2020; 23(5):319-325.
2. Taghizadeh S, Vazehan R, Beheshtian M, Sadeghinia F, Fattahi Z, Mohseni M, Arzhanghi S, Nafissi S, Kariminejad A, **Najmabadi H**, Kahrizi K. Molecular diagnosis of hereditary neuropathies by whole exome sequencing and expanding the phenotype spectrum. *Arch Iran Med.* 2020; 23(7):426-33.
3. Fatehi F, Okhovat AA, Nilipour Y, Mroczek M, Straub V, Töpf A, Palibrk A, Peric S, Rakocevic Stojanovic V, **Najmabadi H**, Nafissi S. Adult-onset very-long-chain acyl-CoA dehydrogenase deficiency (VLCADD). *Eur J Neurol.* 2020. doi: 10.1111/ene.14402.
4. Mehvari S, Larti F, Hu H, Fattahi Z, Beheshtian M, Abedini SS, Arzhanghi S, Ropers HH, Kalscheuer VM, Auld D, Kahrizi K, Riazalhosseini Y, **Najmabadi H**. Whole genome sequencing identifies a duplicated region encompassing Xq13.2q13.3 in a large Iranian family with intellectual disability. *Mol Genet Genomic Med.* 2020; 26:e1418.
5. Bowers M, Liang T, Gonzalez-Bohorquez D, Zocher S, Jaeger BN, Kovacs WJ, Röhrl C, Cramb KML, Winterer J, Kruse M, Dimitrieva S, Overall RW, Wegleiter T, **Najmabadi H**, Semenkovich CF, Kempermann G, Földy C, Jessberger S. FASN-dependent lipid metabolism links neurogenic stem/progenitor cell activity to learning and memory deficits. *Cell Stem Cell.* 2020; 27(1):98-109.e11.
6. Rahimi Bidgoli MM, Javanparast L, Rohani M, **Najmabadi H**, Zamani B, Alavi A. CAPN1 and hereditary spastic paraplegia: a novel variant in an Iranian family and overview of the genotype-phenotype correlation. *Int J Neurosci.* 2020:1-13. doi: 10.1080/00207454.2020.1763344.
7. Mohseni M, Akbari M, Booth KT, Babanejad M, Azaiez H, Ardalani F, Arzhanghi S, Jalalvand K, Nikzat N, Ghodrattpour F, Jamali P, Adeli OA, Habibi H, Kahrizi K, **Najmabadi H**. When transcripts matter: delineating between non-syndromic hearing loss DFNB32 and hearing impairment infertile male syndrome (HIIMS). *J Hum Genet.* 2020; 65(7):609-17.
8. Rostami P, Zendehele K, Shirkoohi R, Ebrahimi E, Ataei M, Imanian H, **Najmabadi H**, Akbari MR, Sanati MH. Gene panel testing in hereditary breast cancer. *Arch Iran Med.* 2020; 23(3):155-62.
9. Kariminejad A, Ghaderi-Sohi S, Keshavarz E, Hashemi SA, Parsimehr E, Szenker-Ravi E, Khatoo M, Faraji Zonooz M, Reversade B, **Najmabadi H**, Hennekam RC. A GLI3 variant leading to polydactyly in heterozygotes and Pallister-Hall-like syndrome in a homozygote. *Clin Genet.* 2020; 97(6):915-9.
10. Pashaei M, Rahimi Bidgoli MM, Zare-Abdollahi D, **Najmabadi H**, Haji-Seyed-Javadi R, Fatehi F, Alavi A. The second mutation of SYCE1 gene associated with autosomal recessive nonobstructive azoospermia. *J Assist Reprod Genet.* 2020; 37(2):451-8.
11. Cheraghi S, Moghbelinejad S, **Najmabadi H**, Kahrizi K, Najafipour R. A novel PTC mutation in the BTB domain of KLHL7 gene in two patients with Bohring-Opitz syndrome-like features. *Eur J Med Genet.* 2020; 63(4):103849.
12. Fadaee M, Mohseni M, Zeinali F, Shokouhi J, Fatahi Z, **Najmabadi H**, Kahrizi K. The second Iranian family with a known missense mutation in AP4M1, precise clinical characterization conducted to AP4 deficiency syndrome. (Submitted, Iranian Red Crescent Medical Journal)
13. Kariminejad A, Yazdan H, Rahimian E, Kalhor Z, Fattahi Z, Zonooz MF, **Najmabadi H**, Ashrafi M. SZT2 mutation in a boy with intellectual disability, seizures and autistic features. *Eur J Med Genet.* 2019; 62(9):103556.
14. Hosseini M, Fattahi Z, Abedini SS, Hu H, Ropers HH, Kalscheuer VM, **Najmabadi H**, Kahrizi K. GPR126: A novel candidate gene implicated in autosomal recessive intellectual disability. *Am J Med Genet A.* 2019; 179(1):13-9.
15. Mehregan H, Mohseni M, Jalalvand K, Arzhanghi S, Nikzat N, Banihashemi S, Kahrizi K, **Najmabadi H**. Novel mutations in MYTH4-FERM domains of myosin 15 are associated with autosomal recessive nonsyndromic hearing loss. *Int J Pediatr Otorhinolaryngol.* 2019; 117:115-26.
16. Beheshtian M, Fattahi Z, Fadaee M, Vazehan R, Jamali P, Parsimehr E, Kamgar M, Zonooz MF, Mahdavi SS, Kalhor Z, Arzhanghi S, Abedini SS, Kermani FS, Mojahedi F, Kalscheuer VM, Ropers HH, Kariminejad A, **Najmabadi H**, Kahrizi K. Identification of disease-causing variants in the EXOSC gene family underlying autosomal recessive intellectual disability in Iranian families. *Clin Genet.* 2019; 95(6):718-25.
17. Kahrizi K, Huber M, Galetzka D, Dewi S, Schröder J, Weis E, Kariminejad A, Fattahi Z, Ropers HH, Schweiger S, **Najmabadi H**, Winter J. Homozygous variants in the gene SCAPER cause syndromic intellectual disability. *Am J Med Genet A.* 2019; 179(7):1214-25.

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18. Mehregan H, Mohseni M, Akbari M, Jalalvand K, Arzhangi S, Nikzat N, Kahrizi K, **Najmabadi H**. Novel mutations in KCNQ4, LHFPL5 and COCH genes in Iranian families with hearing impairment. *Arch Iran Med*. 2019; 22(4):189-197.
  19. Najafi K, Gholami S, Moshtagh A, Bazrgar M, Sadatian N, Abbasi G, Rostami P, Khalili S, Babanejad M, Nourmohammadi B, Faramarzi Garous N, **Najmabadi H**, Kariminejad R. Chromosomal aberrations in pregnancy and fetal loss: Insight on the effect of consanguinity, review of 1625 cases. *Mol Genet Genomic Med*. 2019; 7(8):e820.
  20. Babanejad M, Zarandy MM, Nikzat N, Bazazzadegan N, Arzhangi S, Mohseni M, Kahrizi K, **Najmabadi H**. G130V de novo mutation in an Iranian pedigree with nonsyndromic hearing loss without palmoplantar keratoderma. *Int J Pediatr Otorhinolaryngol*. 2019; 126:109607.
  21. Ataei R, Khoshbakht S, Beheshtian M, Abedini SS, Behravan H, Esmaeili Dizghandi S, Godratpour F, Mirzaei S, Bahrami F, Akbari M, Keshavarzi F, Kahrizi K, **Najmabadi H**. Contribution of Iran in elucidating the genetic causes of autosomal recessive intellectual disability. *Arch Iran Med*. 2019; 22(8):461-71.
  22. Bazazzadegan N, Vazehan R, Fadaee M, Fattahi Z, Abolhassani A, Parsimehr E, Kalhor Z, Faraji Zonooz M, Ahangari F, Dehdahsi S, Samiee F, Jamali P, Habibi H, Nourizadeh Y, Mahdavi S, Beheshtian M, Kariminejad A, Smith RJ, **Najmabadi H**. Brief report of variants detected in hereditary hearing loss cases in Iran over a 3-year period. *Iran J Public Health*. 2019; 48(10):1910-5.
  23. Kariminejad A, Vahidnezhad H, Ghaderi-Sohi S, Ghannadan AR, Youssefian L, Parsimehr E, Faraji Zonooz M, Kariminejad MH, Uitto J, **Najmabadi H**, Hennekam RC. Widespread aplasia cutis congenita in sibs with PLEC1 and ITGB4 variants. *Am J Med Genet A*. 2019; 179(8):1547-55.
  24. Mohammadzadeh A, Akbaroghli S, Aghaei-Moghadam E, Mahdieh N, Badv RS, Jamali P, Kariminejad R, Chavoshzadeh Z, Ghasemi Firouzabadi S, Mansour Ghanaie R, Nozari A, Banihashemi S, Hadipour F, Hadipour Z, Kariminejad A, **Najmabadi H**, Shafeghati Y, Behjati F. Investigation of chromosomal abnormalities and microdeletion/ microduplication(s) in fifty Iranian patients with multiple congenital anomalies. *Cell J*. 2019; 21(3):337-49.
  25. Kariminejad A, Szenker-Ravi E, Lekszas C, Tajsharghi H, Moslemi AR, Naert T, Tran HT, Ahangari F, Rajaei M, Nasserli M, Haaf T, Azad A, Superti-Furga A, Maroofian R, Ghaderi-Sohi S, **Najmabadi H**, Abbaszadegan MR, Vleminckx K, Nikuei P, Reversade B. Homozygous null TBX4 mutations lead to posterior amelia with pelvic and pulmonary hypoplasia. *Am J Hum Genet*. 2019; 105(6):1294-301.
  26. Bekiesinska-Figatowska M, Hosseini M, Arzhangi S, **Najmabadi H**, Rosenfeld JA, Du H, Marafi D, Blaser S, Teitelbaum R, Silver R; Baylor-Hopkins Center for Mendelian Genomics, Posey JE, Ropers HH, Gibbs RA, Wiszniewski W, Lupski JR, Chitayat D, Kahrizi K, Gawlinski P. Bi-allelic pathogenic variants in TUBGCP2 cause microcephaly and lissencephaly spectrum disorders. *Am J Hum Genet*. 2019; 105(5):1005-15.
  27. Fattahi Z, Sheikh TI, Musante L, Rasheed M, Taskiran II, Harripaul R, Hu H, Kazeminasab S, Alam MR, Hosseini M, Larti F, Ghaderi Z, Celik A, Ayub M, Ansar M, Haddadi M, Wienker TF, Ropers HH, Kahrizi K, Vincent JB, **Najmabadi H**. Biallelic missense variants in ZBTB11 can cause intellectual disability in humans. *Hum Mol Genet*. 2018; 27(18):3177-88.
  28. Abedini SS, Kahrizi K, de Poupiana LR, **Najmabadi H**. tRNA methyltransferase defects and intellectual disability. *Arch Iran Med*. 2018; 21(10):478-85.
  29. Kazeminasab S, Taskiran II, Fattahi Z, Bazazzadegan N, Hosseini M, Rahimi M, Oladnabi M, Haddadi M, Celik A, Ropers HH, **Najmabadi H**, Kahrizi K. CNKSR1 gene defect can cause syndromic autosomal recessive intellectual disability. *Am J Med Genet B Neuropsychiatr Genet*. 2018; 177(8):691-9.
  30. Abedini SS, Forouzesheh Pour F, Karimi K, Ghaderi Z, Farashi S, Tavakoli Koudehi A, Javadi Pirouz H, Mobini Nejad SB, Azarkeivan A, **Najmabadi H**. Frequency of  $\alpha$ -globin gene triplications and coinheritance with  $\beta$ -globin gene mutations in the Iranian population. *Hemoglobin*. 2018; 42(4):252-6.
  31. Alavi A, Esmaeili S, Nafissi S, Kahrizi K, **Najmabadi H**. Genotype and phenotype analysis of 43 Iranian facioscapulohumeral muscular dystrophy patients; Evidence for anticipation. *Neuromuscul Disord*. 2018; 28(4):303-14.
  32. Babanejad M, Adeli OA, Nikzat N, Beheshtian M, Azarafra H, Sadeghnia F, Mohseni M, **Najmabadi H**, Kahrizi K. SLC52A2 mutations cause SCABD2 phenotype: A second report. *Int J Pediatr Otorhinolaryngol*. 2018; 104:195-9.
  33. Booth KT, Azaiez H, Kahrizi K, Wang D, Zhang Y, Frees K, Nishimura C, **Najmabadi H**, Smith RJ. Exonic mutations and exon skipping: Lessons learned from DFNA5. *Hum Mutat*. 2018; 39(3):433-40.
  34. Booth KT, Kahrizi K, Babanejad M, Daghighi H, Bademci G, Arzhangi S, Zareabdollahi D, Duman D, El-Amraoui A, Tekin M, **Najmabadi H**, Azaiez H, Smith RJ. Variants in CIB2 cause DFNB48 and not USH1J.
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- Clin Genet. 2018; 93(4):812-21.
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  36. Hosseini M, **Najmabadi H**, Kahrizi K. Calpains: Diverse functions but enigmatic. *Arch Iran Med.* 2018; 21(4):170-9.
  37. Imtiaz A, Belyantseva IA, Beirl AJ, Fenollar-Ferrer C, Bashir R, Bukhari I, Bouzid A, Shaukat U, Azaiez H, Booth KT, Kahrizi K, **Najmabadi H**, Maqsood A, Wilson EA, Fitzgerald TS, Tlili A, Olszewski R, Lund M, Chaudhry T, Rehman AU, Starost MF, Waryah AM, Hoa M, Dong L, Morell RJ, Smith RJH, Riazuddin S, Masmoudi S, Kindt KS, Naz S, Friedman TB. CDC14A phosphatase is essential for hearing and male fertility in mouse and human. *Hum Mol Genet.* 2018; 27(5):780-98.
  38. Kazeminasab S, **Najmabadi H**, Kahrizi K. Intellectual disability and ataxia: Genetic collisions. *Arch Iran Med.* 2018; 21(1):29-40.
  39. Mehri M, Zarin M, Ardalani F, **Najmabadi H**, Azarkeivan A, Neishabury M. Novel mutations in mitochondrial carrier family gene SLC25A38, causing congenital sideroblastic anemia in Iranian families, identified by whole exome sequencing. *Blood Cells Mol Dis.* 2018; 71:39-44.
  40. Nozari A, Aghaei-Moghadam E, Zeinaloo A, Mollazadeh R, Majnoon MT, Alavi A, Ghasemi Firouzabadi S, Mohammadzadeh A, Banihashemi S, Nikzaban M, **Najmabadi H**, Behjati F. A novel splicing variant in FLNC gene responsible for a highly penetrant familial dilated cardiomyopathy in an extended Iranian family. *Gene.* 2018; 659:160-7.
  41. Charzewska A, Maiwald R, Kahrizi K, Oehl-Jaschkowitz B, Dufke A, Lemke JR, Enders H, **Najmabadi H**, Tzschach A, Hachmann W, Jensen C, Bienek M, Poznański J, Nawara M, Chilarska T, Obersztyn E, Hoffman-Zacharska D, Gos M, Bal J, Kalscheuer VM. The power of the Mediator complex-Expanding the genetic architecture and phenotypic spectrum of MED12-related disorders. *Clin Genet.* 2018; 94(5):450-6.
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  43. Fazeli Z, Ghaderian SMH, **Najmabadi H**, Omrani MD. High expression of miR-510 was associated with CGG expansion located at upstream of FMR1 into full mutation. *J Cell Biochem.* 2018. doi: 10.1002/jcb.27505. Online ahead of print.PMID: 30160796
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  45. Kariminejad A, Dahl-Halvarsson M, Ravenscroft G, Afroozan F, Keshavarz E, Goullée H, Davis MR, Faraji Zonooz M, **Najmabadi H**, Laing NG, Tajsharghi H. TOR1A variants cause a severe arthrogryposis with developmental delay, strabismus and tremor. *Brain.* 2017; 140(11):2851-9.
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