

5.1.3 Neurological Diseases, Neuroscience & Mental Health Area

Genetic Diseases

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DESCRIPTION

The Genetic Diseases Group is a multidisciplinary group with an important clinical component in its research within the Genetics Department of the Hospital de Sant Pau. Its activities also have a translational orientation facilitated by the clinical environment in which it performs the laboratory work.

The primary objective of the Genetic Diseases Group is the progress of the molecular diagnosis of genetic and rare diseases through genomic, transcriptomic, functional, and pharmacogenetic studies improving the knowledge of the molecular mechanisms of described genes and the identification of new ones. The results of these genetic studies allow us to establish in many cases different therapeutic options and to offer genetic counseling to the families. The fact that the group is made up entirely of Sant Pau Genetics Dept staff, supports the clinical translational effect of their research work.

MAIN LINES OF RESEARCH

- Genetics of congenital myopathies, congenital, and adult muscular dystrophies, including limb girdle muscle dystrophies, and congenital myasthenic syndromes.
- Spinal atrophy and smn genes.
- Genetics of hereditary cancer.
- Pharmacogenetics.
- Cardiogenetics.
- Genetics in pediatrics-obstetrics.



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SCIENTIFIC CHALLENGES

Congenital myopathies, congenital and adult muscular dystrophies, including limb girdle muscle dystrophies, and congenital myasthenic syndromes

- Identification of mutations and genetic variants.
- Studies of molecular pathology and disease mechanisms of neuromuscular diseases-related genes.
- Transcriptome analysis by RNA Seq technology.
- Functional studies on the DMD genes to determine the pathogenic effect of the identified variants.
- Long Read Sequencing by ONT technology to characterize long repeat expansions in size and composition.

Spinal atrophy and SMN genes

- To design new strategies for the study and improvement of the molecular diagnosis of atypical cases.
- To perform functional studies for the characterization of genetic variants of uncertain clinical effect to improve diagnosis.
- Studies of molecular pathology, disease mechanisms, and SMN gene expression.
- Identification of modifying genes.
- Study of biomarkers for validation processing in spinal muscular atrophy.
- Study of the neuromuscular junction in human development.

Hereditary cancer

- Development of Precision Medicine in hereditary syndromes of predisposition to cancer: an approach to the diagnosis, treatment and prevention of cancer through the identification of pathogenic variants in multigene assays.
- Development of functional studies for the characterization of genetic variants of uncertain clinical effects.
- Analysis of biomarkers in liquid biopsy-free circulating tumor DNA (cfDNA) as a predictor of response to the treatment of breast cancer.

- Contribution to the development of an open database of genetic variants included in Hereditary Cancer Genetic Testing Panels.

Pharmacogenetics

- Identification of biomarkers of response/toxicity to drugs used in cancer, rheumatology, inflammatory bowel diseases, cardiology, hematology.
- Congenital coagulopathies: molecular pathology of hemophilia.
- Duchenne and Becker muscular dystrophy: molecular pathology of DMD gene.
- Identification of the molecular alterations responsible for hereditary hearing loss.
- Studies of the molecular pathology causing ataxias.
- Clinical and Molecular diagnosis of complex diseases of pediatric-onset with special attention in Neurodevelopmental Disorders.

Cardiogenetics

- Clinical and Molecular studies of patients with cardiomyopathies (dilated, hypertrophic, arrhythmogenic, non-compacted, etc) to identify disease-associated genetic variants.
- Study of clinically discordant families with the same pathogenic genetic variant to identify phenotype modifiers in cardiomyopathies.
- Clinical and Molecular characterization of patients with aortic diseases (Aortic Aneurysms, Aortic Valvular Disease, Aortic Occlusive Disease, Traumatic Aortic Disease, Aortic Dissections).

Pediatrics-obstetrics

- Clinical and Molecular studies of syndromic patients:
- Identification of responsible variants and discovery of new genes
- Clinical and Molecular studies of fetal anomalies detected during pregnancy.

ACTIVE GRANTS

- Gonzalez Quereda, Lidia. Análisis integrado de datos genómicos y transcriptómicos para el avance en el conocimiento molecular y diagnóstico de las miopatías. PI22/01859. Instituto



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- de Salud Carlos III (ISCIII). Duration: 2023-2025. 87.120,00 €.
- Riera Armengol, Pau. Integración de la farmacogenética y la telemedicina mediante salud móvil para el manejo de los efectos adversos en pacientes con cáncer colorrectal. (ePGx study). PI21/01913. Instituto de Salud Carlos III (ISCIII). Duration: 2022-2024. 63.525,00 €.
 - Riera Armengol, Pau. Integració de la farmàcia hospitalària i comunitària per a la implementació de la farmacogenètica a la pràctica assistencial. JOAQUIM BONAL 2019. Fundació Acadèmia de Ciències Mèdiques i de la Salut de Catalunya i Balears. Duration: 2019-2023. 14.000,40 €.
 - Riera Armengol, Pau. Integración de la farmacogenética y la telemedicina mediante salud móvil para el manejo de los efectos adversos en pacientes con cáncer colorrectal. (ePGx study). FEFH 2022. Fundación Española de Farmacia Hospitalaria. Duration: 2021-2023. 15.400,00 €.
- ### SCIENTIFIC PRODUCTION
- Abuli A, Costa M, Codina M, Valenzuela I, Leno J, Rovira E, Cueto A, Fernández P, García E, Cusco I, Tizzano EF. Experience using singleton exome sequencing of probands as an approach to preconception carrier screening in consanguineous couples. JOURNAL OF MEDICAL GENETICS. 2023; 60(6)DOI:10.1136/jmg-2022-108607. PMID:36600615. IF:4,000 (Q2/3D). Document type: Article.
 - Arqueros C, Salazar J, Gallardo A, Andrés M, Tibau A, Bell OL, Artigas A, Las A, Cajal T, Lerma E, Barnadas A. Secreted Protein Acidic and Rich in Cysteine (<i>SPARC</i>) Polymorphisms in Response to Neoadjuvant Chemotherapy in HER2-Negative Breast Cancer Patients. BIOMEDICINES. 2023; 11(12):3231. DOI:10.3390/biomedicines11123231. PMID:38137452. IF:4,700 (Q1/3D). Document type: Article.
 - Bermejo L, Fernández CPD, González L, Segarra A, Nedkova V, Gallano P, Martín P, Hernández A, Olivé M, Arteche A, Domínguez C. Distal myopathy due to digenic inheritance of <i>TIA1</i> and <i>SQSTM1</i> variants in two unrelated Spanish patients. NEUROMUSCULAR DISORDERS. 2023; 33(12)DOI:10.1016/j.nmd.2023.10.016. PMID:38016875. IF:2,800 (Q3/6D). Document type: Article.
 - Codina M, Trujillano L, Abuli A, Rovira E, Muñoz P, Campos B, Fernández P, Palau D, Carrasco E, Valenzuela I, Cueto AM, Las A, Limeres J, Leno J, Costa M, Moles A, Balmana J, Diez O, Cusco I, García E, Tizzano EF. An spanish study of secondary findings in families affected with mendelian disorders: choices, prevalence and family history. EUROPEAN JOURNAL OF HUMAN GENETICS. 2023; 31(2)DOI:10.1038/s41431-022-01240-5. PMID:36446894. IF:5,200 (Q1/2D). Document type: Article.
 - Collet R, Olmedo G, Ruiz I, Martínez A, Rodríguez B, Bernal S, Kulisevsky J, Pagonabarraga J. Late-Onset Beta-Propeller Protein-Associated Neurodegeneration: A Case Report. Movement Disorders Clinical Practice. 2023; 10(8)DOI:10.1002/mdc3.13811. PMID:37635772. IF:4,000 (Q2/4D). Document type: Article.
 - De Pourcq JT, Riera P. Critical Commentary on the paper: Compatibility of prolonged infusion antibiotics during Y-site administration with parenteral nutrition. Nursing in Critical Care. 2023; 28(6)DOI:10.1111/nicc.12910. PMID:37232328. IF:3,000 (Q1/2D). Document type: Editorial Material.
 - Esmel R, Valenzuela I, Riaza L, Rodríguez B, Rosés F, Boronat S, Sabaté A. Arterial tortuosity syndrome: Phenotypic features and cardiovascular manifestations in 4 newly identified patients. European Journal of Medical Genetics. 2023; 66(9):104823. DOI:10.1016/j.ejmg.2023.104823. PMID:37619836. IF:1,900 (Q4/8D). Document type: Article.
 - Iznardo H, Bernal S, Boronat S, Roé E. Sclerotic Bone Lesions as a Clue in the Diagnosis of Three Generations of Tuberous Sclerosis Complex: Case Report and Review of Literature. PEDIATRIC NEUROLOGY. 2023; 148DOI:10.1016/j.pediatr-neurol.2023.07.022. PMID:37634327. IF:3,800 (Q1/2D). Document type: Article.
 - Miarons M, Gordon AM, Riera P, Nicolas FG, Spanish Soc Hosp Pharm S. Allelic Frequency of DPYD Genetic Variants in Patients With Cancer in Spain: The PhotoDPYD Study. ONCOLOGIST. 2023; 28(5):e304-e308. DOI:10.1093/oncolo/oyad077. PMID:37014829. IF:5,800 (Q1/3D). Document type: Article.



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- Mir JF, Rodríguez C, Estrada M, de Gamarra EF, Mangues MA, Bagaria G, Riera P. An Integrated Multidisciplinary Circuit Led by Hospital and Community Pharmacists to Implement Clopidogrel Pharmacogenetics in Clinical Practice. *Pharmacy*. 2023; 11(2):76. DOI:10.3390/pharmacy11020076. PMID:37104082. Document type: Article.
- Palones E, Curto E, Plaza V, Gonzalez L, Segarra A, Querol L, Bertoletti F, Rodríguez MJ, Gallano P, Crespo A. Clinical and functional characteristics, possible causes, and impact of chronic cough in patients with cerebellar ataxia, neuropathy, and bilateral vestibular areflexia syndrome (CANVAS). *JOURNAL OF NEUROLOGY*. 2023; DOI:10.1007/s00415-023-12001-9. PMID:37917234. IF:6,000 (Q1/2D). Document type: Article.
- Panque M, Guimarães L, Bengoa J, Pasalodos S, Cordier C, Esteban I, Lemos C, Moldovan R, Serra C. An European overview of genetic counselling supervision provision. *European Journal of Medical Genetics*. 2023; 66(4):104710. DOI:10.1016/j.ejmg.2023.104710. PMID:36731744. IF:1,900 (Q4/8D). Document type: Article.
- Rodríguez PM, Ravenscroft G, Natera D, Carr A, Manzur A, Liu WW, Vella NR, Jericó I, Gonzalez L, Gallano P, Montalto SA, Davis MR, Lamont PJ, Laing NG, Bourque P, Nascimento A, Muntoni F, Polavarapu K, Lochmüller H, Palace J, Beeson D. A novel phenotype of AChR-deficiency syndrome with predominant facial and distal weakness resulting from the inclusion of an evolutionary alternatively-spliced exon in CHRNA1. *NEUROMUSCULAR DISORDERS*. 2023; 33(2) DOI:10.1016/j.nmd.2022.12.011. PMID:36634413. IF:2,800 (Q3/6D). Document type: Article.
- Sainz L, Riera P, Moya P, Bernal S, Casademont J, Díaz C, Millán AM, Park HS, Lasa A, Corominas H. Clinical Value of IL6R Gene Variants as Predictive Biomarkers for Toxicity to Tocilizumab in Patients with Rheumatoid Arthritis. *Journal of Personalized Medicine*. 2023; 13(1):61. DOI:10.3390/jpm13010061. PMID:36675722. IF:3,508 (Q2/4D). Document type: Article.
- Sainz L, Riera P, Moya P, Bernal S, Casademont J, Diaz C, Millan AM, Park HS, Lasa A, Corominas H. Impact of IL6R genetic variants on treatment efficacy and toxicity response to sarilumab in rheumatoid arthritis. *ARTHRITIS RESEARCH & THERAPY*. 2023; 25(1):226. DOI:10.1186/s13075-023-03209-1. PMID:38001504. IF:4,900 (Q2/4D). Document type: Article.
- Sarkozy A, Bourke JP, Ferlini A, Barthélémy I, Cripe LH, Reuben E, Evangelista T et al. 263rd ENMC International Workshop: Focus on female carriers of dystrophinopathy: refining recommendations for prevention, diagnosis, surveillance, and treatment. Hoofddorp, The Netherlands, 13–15 May 2022. THE BOULEVARD, LANGFORD LANE, KIDLINGTON, OXFORD OX5 1GB, ENGLAND:PERGAMON-ELSEVIER SCIENCE LTD. 2023. p.p. 274-284. Document type: Conference Paper.
- Segarra A, Collet R, Gonzalez L, Vesperinas A, Caballero M, Carbayo A, Diaz J, Rodríguez MJ, Gállo E, Gallano P, Olive M. A new homozygous missense variant in LMOD3 gene causing mild nemaline myopathy with prominent facial weakness. *NEUROMUSCULAR DISORDERS*. 2023; 33(4) DOI:10.1016/j.nmd.2023.02.006. PMID:36893608. IF:2,800 (Q3/6D). Document type: Article.
- Segarra A, Dominguez C, Hernandez A, Sánchez MT, Camacho A, Rivas E, Campo A, Madruga M, Ortez C, Natera D, Nascimento A, Codina A, Rodríguez MJ, Gallano P, Gonzalez L. Genetic diagnosis of Duchenne and Becker muscular dystrophy through mRNA analysis: new splicing events. *JOURNAL OF MEDICAL GENETICS*. 2023; 60(6) DOI:10.1136/jmg-2022-108828. PMID:36535754. IF:4,000 (Q2/3D). Document type: Article.
- Trifunov S, Natera D, Carrera L, Codina A, Expósito J, Ortez C, Medina J, Alcalá ST, Bernal S, Alias L, Badosa C, Balsells S, Alcolea D, Nascimento A, Jimenez C. Full-Length SMN Transcript in Extracellular Vesicles as Biomarker in Individuals with Spinal Muscular Atrophy Type 2 Treated with Nusinersen. *Journal Of Neuromuscular Diseases*. 2023; 10(4) DOI:10.3233/JND-230012. PMID:37038823. IF:3,300 (Q2/5D). Document type: Article.