



**Sistema Integral de Información Académica**  
**Coordinación de Planeación, Evaluación y**  
**Simplificación de la Gestión Institucional**  
**Reporte individual**



**ARIADNA ESTELA GONZALEZ DEL ANGEL**

## **Datos Generales**

**Nombre:** ARIADNA ESTELA GONZALEZ DEL ANGEL

**Máximo nivel de estudios:** DOCTORADO

**Antigüedad académica en la UNAM:** 12 años

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## **Nombramientos**

**Último:** PROFESOR ASIGNATURA A TP No Definitivo  
Facultad de Medicina  
Desde 01-01-2008 (fecha inicial de registros en el  
SIIA) hasta 15-05-2019

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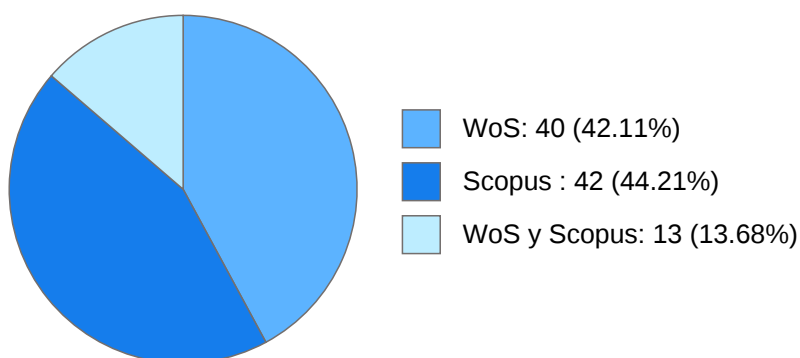
## **Estímulos, programas, premios y reconocimientos**

SNI II 2017 - 2023  
SNI I 2011 - 2016

**ARIADNA ESTELA GONZALEZ DEL ANGEL**

**DOCUMENTOS EN REVISTAS**

**Histórico de Documentos**



| # | Título                                                                                                                                                                                     | Autores                                                                                                  | Revista                                     | Año  |
|---|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------|---------------------------------------------|------|
| 1 | Arginase deficiency in Mexico: Insights from the experience of a metabolic reference center                                                                                                | CYNTHIA FERNANDEZ LAINEZ MIGUEL<br>ANGEL ALCANTARA ORTIGOZA<br>ARIADNA ESTELA GONZALEZ DEL ANGEL et al.  | Molecular Genetics and Metabolism Reports   | 2025 |
| 2 | Isolated methylmalonic acidemia in Mexico: Genotypic spectrum, report of two novel MMUT variants and a possible synergistic heterozygosity effect                                          | CYNTHIA FERNANDEZ LAINEZ MIRIAM<br>ERANDI REYNA FABIAN MIGUEL ANGEL ALCANTARA ORTIGOZA et al.            | Molecular Genetics and Metabolism Reports   | 2024 |
| 3 | Concordance Between Biochemical and Molecular Diagnosis Obtained by WES in Mexican Patients with Inborn Errors of Intermediary Metabolism: Utility for Therapeutic Management              | MIGUEL ANGEL ALCANTARA ORTIGOZA ARIADNA ESTELA GONZALEZ DEL ANGEL MIRIAM ERANDI REYNA FABIAN et al.      | INTERNATIONAL JOURNAL OF MOLECULAR SCIENCES | 2024 |
| 4 | In Silico Structural Protein Evaluation of the Phenylalanine Hydroxylase p.(Tyr77His) Variant Associated with Benign Hyperphenylalaninemia as Identified through Mexican Newborn Screening | MIGUEL ANGEL ALCANTARA ORTIGOZA ARIADNA ESTELA GONZALEZ DEL ANGEL ISABEL CRISTINA IBARRA GONZALEZ et al. | Children-Basel                              | 2023 |

**ARIADNA ESTELA GONZALEZ DEL ANGEL**

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|----|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------|------|
| 5  | Proposed clinical approach and imaging studies in families with oculo-auriculo-vertebral spectrum to assess variable expressivity                                                                                                                         | MIGUEL ANGEL ALCANTARA<br>ORTIGOZA ARIADNA ESTELA<br>GONZALEZ DEL ANGEL Bernardette<br>Estandia-Ortega et al.     | AMERICAN<br>JOURNAL OF<br>MEDICAL<br>GENETICS PART<br>A | 2022 |
| 6  | The Enigmatic Etiology of Oculo-Auriculo-Vertebral Spectrum (OAVS): An Exploratory Gene Variant Interaction Approach in Candidate Genes                                                                                                                   | MIRIAM ERANDI REYNA FABIAN JOSE<br>ANTONIO VELAZQUEZ ARAGON<br>ARIADNA ESTELA GONZALEZ DEL<br>ANGEL et al.        | LIFE-BASEL                                              | 2022 |
| 7  | Unusual Clinical Manifestations in a Mexican Patient with Sanfilippo B Syndrome                                                                                                                                                                           | MIRIAM ERANDI REYNA FABIAN<br>MIGUEL ANGEL ALCANTARA<br>ORTIGOZA LUIS LEONARDO FLORES<br>LAGUNES et al.           | Diagnostics                                             | 2022 |
| 8  | Functional characterization of the p.(Gln195His) or Tainan and novel p.(Ser184Cys) or Toluca glucose-6-phosphate dehydrogenase (G6PD) gene natural variants identified through Mexican newborn screening for glucose-6-phosphate dehydrogenase deficiency | MIGUEL ANGEL ALCANTARA<br>ORTIGOZA BEATRIZ HERNANDEZ<br>OCHOA ARIADNA ESTELA GONZALEZ<br>DEL ANGEL et al.         | CLINICAL<br>BIOCHEMISTRY                                | 2022 |
| 9  | Cleft Lip Palate in a Patient with 5q14.3 Deletion Syndrome: A Possible Unreported Feature?                                                                                                                                                               | MIGUEL ANGEL ALCANTARA<br>ORTIGOZA ARIADNA ESTELA<br>GONZALEZ DEL ANGEL Hernández L.F.<br>et al.                  | CYTOGENETIC<br>AND GENOME<br>RESEARCH                   | 2022 |
| 10 | Genotypic spectrum underlying tetrahydrobiopterin metabolism defects: Experience in a single Mexican reference center                                                                                                                                     | MIGUEL ANGEL ALCANTARA<br>ORTIGOZA ISABEL CRISTINA IBARRA<br>GONZALEZ ARIADNA ESTELA<br>GONZALEZ DEL ANGEL et al. | Frontiers in<br>Genetics                                | 2022 |
| 11 | Whole-genome amplification/preimplantation genetic testing for propionic acidemia of successful pregnancy in an obligate carrier Mexican couple: A case report                                                                                            | MIGUEL ANGEL ALCANTARA<br>ORTIGOZA ARIADNA ESTELA<br>GONZALEZ DEL ANGEL Neumann A.<br>et al.                      | World Journal<br>Of Clinical<br>Cases                   | 2021 |
| 12 | Further Evidence That Defects in Main Thyroid Dysgenesis-Related Genes Are an Uncommon Etiology for Primary Congenital Hypothyroidism in Mexican Patients: Report of Rare Variants in FOXE1, NKX2-5 and TSHR                                              | MIGUEL ANGEL ALCANTARA<br>ORTIGOZA SERGIO ENRIQUEZ FLORES<br>ARIADNA ESTELA GONZALEZ DEL<br>ANGEL et al.          | Children-Basel                                          | 2021 |

**ARIADNA ESTELA GONZALEZ DEL ANGEL**

|    |                                                                                                                                                            |                                                                                                         |                                                                              |      |
|----|------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------|------|
| 13 | Genetic spectrum and clinical early natural history of glucose-6-phosphate dehydrogenase deficiency in Mexican children detected through newborn screening | MIGUEL ANGEL ALCANTARA<br>ORTIGOZA ARIADNA ESTELA<br>GONZALEZ DEL ANGEL CARLOS LOPEZ<br>CANDIANI et al. | ORPHANET<br>JOURNAL OF<br>RARE DISEASES                                      | 2021 |
| 14 | Hair pigment distribution changes after haematopoietic stem cell transplantation in Griscelli syndrome type 2                                              | MARCO ANTONIO YAMAZAKI<br>NAKASHIMADA RODRIGO ROLDAN<br>MARIN SONIA TOUSSAINT CAIRE et al.              | JOURNAL OF<br>THE EUROPEAN<br>ACADEMY OF<br>DERMATOLOGY<br>AND<br>VENEREOLGY | 2021 |
| 15 | First comprehensive TSC1/TSC2 mutational analysis in Mexican patients with Tuberous Sclerosis Complex reveals numerous novel pathogenic variants           | MIRIAM ERANDI REYNA FABIAN<br>MIGUEL ANGEL ALCANTARA<br>ORTIGOZA SERGIO ENRIQUEZ FLORES<br>et al.       | SCIENTIFIC<br>REPORTS                                                        | 2020 |
| 16 | Nonmosaic Trisomy 19p13.3p13.2 Resulting from a Rare Unbalanced t(Y;19)(q12;p13.2) Translocation in a Patient with Pachygyria and Polymicrogyria           | ARIADNA ESTELA GONZALEZ DEL<br>ANGEL MIGUEL ANGEL ALCANTARA<br>ORTIGOZA Martínez Anaya D. et al.        | CYTOGENETIC<br>AND GENOME<br>RESEARCH                                        | 2020 |
| 17 | TSC2/PKD1 contiguous gene syndrome, with emphasis on a case with an atypical mild polycystic kidney phenotype and a novel genetic variant                  | MIRIAM ERANDI REYNA FABIAN JAIME<br>BERUMEN CAMPOS ARIADNA ESTELA<br>GONZALEZ DEL ANGEL et al.          | Nefrologia                                                                   | 2020 |
| 18 | Tumor de cordón sexual con túbulos anulares y cistadenoma mucinoso de ovario en una adolescente con síndrome de Peutz-Jeghers                              | ROSALIA GARZA ELIZONDO ARIADNA<br>ESTELA GONZALEZ DEL ANGEL<br>EDUARDO LOPEZ CORELLA et al.             | ACTA<br>PEDIATRICA DE<br>MEXICO                                              | 2020 |
| 19 | Molecular analysis provides further evidence that Chitayat syndrome is caused by the recurrent p.(Tyr89Cys) pathogenic variant in the ERF gene             | MIGUEL ANGEL ALCANTARA<br>ORTIGOZA ARIADNA ESTELA<br>GONZALEZ DEL ANGEL<br>Caro-Contreras A. et al.     | AMERICAN<br>JOURNAL OF<br>MEDICAL<br>GENETICS PART<br>A                      | 2019 |
| 20 | Report of a patient with a de novo non-recurrent duplication of 17p11.2p12 and Yq11 deletion                                                               | MIGUEL ANGEL ALCANTARA<br>ORTIGOZA BERTHA MOLINA ALVAREZ<br>ARIADNA ESTELA GONZALEZ DEL<br>ANGEL et al. | MOLECULAR<br>CYTOGENETICS                                                    | 2019 |
| 21 | Diagnosis of Laron syndrome using monoplex-polymerase chain reaction technology with a whole-genome amplification template: A case report                  | MIGUEL ANGEL ALCANTARA<br>ORTIGOZA ARIADNA ESTELA<br>GONZALEZ DEL ANGEL Neumann A.<br>et al.            | World Journal<br>Of Clinical<br>Cases                                        | 2019 |

**ARIADNA ESTELA GONZALEZ DEL ANGEL**

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|----|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------|------|
| 22 | Mutational spectrum of Mexican patients with tyrosinemia type 1: In silico modeling and predicted pathogenic effect of a novel missense FAH variant                                                                           | ISABEL CRISTINA IBARRA GONZALEZ<br>MIGUEL ANGEL ALCANTARA<br>ORTIGOZA ARIADNA ESTELA<br>GONZALEZ DEL ANGEL et al.    | Molecular<br>Genetics &<br>Genomic<br>Medicine                               | 2019 |
| 23 | Clinical characterization and identification of five novel FOXL2 pathogenic variants in a cohort of 12 Mexican subjects with the syndrome of blepharophimosis-ptosis-epicanthus inversus                                      | OSCAR FRANCISCO CHACON<br>CAMACHO ANGEL NAVA CASTAÑEDA<br>MARIA DE LOURDES RODRIGUEZ<br>CABRERA et al.               | Gene                                                                         | 2019 |
| 24 | Predominance of dystrophinopathy genotypes in mexican male patients presenting as muscular dystrophy with a normal multiplex polymerase chain reaction DMD gene result: A study including targeted next-generation sequencing | MIGUEL ANGEL ALCANTARA<br>ORTIGOZA MIRIAM ERANDI REYNA<br>FABIAN ARIADNA ESTELA GONZALEZ<br>DEL ANGEL et al.         | GENES                                                                        | 2019 |
| 25 | Genetic and clinical characterization of 73 Pigmentary Mosaicism patients: Revealing the genetic basis of clinical manifestations                                                                                             | VICTORIA DEL CASTILLO RUIZ<br>LOURDES CAROLA DURAN MC<br>KINSTER ARIADNA ESTELA GONZALEZ<br>DEL ANGEL et al.         | ORPHANET<br>JOURNAL OF<br>RARE DISEASES                                      | 2019 |
| 26 | Expanding the clinical features of autoinflammation and phospholipase C $\beta$ 2-associated antibody deficiency and immune dysregulation by description of a novel patient                                                   | MARCO ANTONIO YAMAZAKI<br>NAKASHIMADA MIGUEL ANGEL<br>ALCANTARA ORTIGOZA ARIADNA<br>ESTELA GONZALEZ DEL ANGEL et al. | JOURNAL OF<br>THE EUROPEAN<br>ACADEMY OF<br>DERMATOLOGY<br>AND<br>VENEREOLGY | 2019 |
| 27 | Novel Phenotypes and Cardiac Involvement Associated With DNA2 Genetic Variants                                                                                                                                                | ARIADNA ESTELA GONZALEZ DEL<br>ANGEL Bisciglia M. Vargas-Cañas S.<br>et al.                                          | FRONTIERS IN<br>NEUROLOGY                                                    | 2019 |
| 28 | Identification of a novel SLC12A6 pathogenic variant associated with hereditary motor and sensory neuropathy with agenesis of the corpus callosum (HMSN/ACC) in a non-French-Canadian family                                  | ARIADNA ESTELA GONZALEZ DEL<br>ANGEL JOSE ANTONIO VELAZQUEZ<br>ARAGON MIGUEL ANGEL ALCANTARA<br>ORTIGOZA et al.      | NEUROLOGY<br>INDIA                                                           | 2018 |
| 29 | Unique association of hypochondroplasia with craniosynostosis and cleft palate in a Mexican family                                                                                                                            | ARIADNA ESTELA GONZALEZ DEL<br>ANGEL MIGUEL ANGEL ALCANTARA<br>ORTIGOZA Caro-Contreras A. et al.                     | AMERICAN<br>JOURNAL OF<br>MEDICAL<br>GENETICS PART<br>A                      | 2018 |

**ARIADNA ESTELA GONZALEZ DEL ANGEL**

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|----|-----------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------|------------------------------------------------|------|
| 30 | Further delineation of achondroplasia?hypochondroplasia complex with long-term survival                                                                         | ARIADNA ESTELA GONZALEZ DEL ANGEL MIGUEL ANGEL ALCANTARA ORTIGOZA VICTORIA DEL CASTILLO RUIZ et al.      | AMERICAN JOURNAL OF MEDICAL GENETICS PART A    | 2018 |
| 31 | Mutational spectrum of PTS gene and in silico pathological assessment of a novel variant in Mexico                                                              | ISABEL CRISTINA IBARRA GONZALEZ MIGUEL ANGEL ALCANTARA ORTIGOZA SERGIO ENRIQUEZ FLORES et al.            | BRAIN & DEVELOPMENT                            | 2018 |
| 32 | Newborn cystic fibrosis screening in southeastern Mexico: Birth prevalence and novel CFTR gene variants                                                         | ISABEL CRISTINA IBARRA GONZALEZ MIGUEL ANGEL ALCANTARA ORTIGOZA ARIADNA ESTELA GONZALEZ DEL ANGEL et al. | JOURNAL OF MEDICAL SCREENING                   | 2018 |
| 33 | Clinical characterization of patients with autosomal dominant short stature due to aggrecan mutations                                                           | ARIADNA ESTELA GONZALEZ DEL ANGEL Gkourogianni A. Andrew M. et al.                                       | JOURNAL OF CLINICAL ENDOCRINOLOGY & METABOLISM | 2017 |
| 34 | An uncommon inheritance pattern in Niemann-Pick disease type C: Identification of probable paternal germline mosaicism in a Mexican family                      | MIGUEL ANGEL ALCANTARA ORTIGOZA ARIADNA ESTELA GONZALEZ DEL ANGEL Cervera-Gaviria M. et al.              | BMC NEUROLOGY                                  | 2016 |
| 35 | Expansion of the variable expression of Muenke syndrome: Hydrocephalus without craniosynostosis                                                                 | ARIADNA ESTELA GONZALEZ DEL ANGEL MIGUEL ANGEL ALCANTARA ORTIGOZA Estandía-Ortega B. et al.              | AMERICAN JOURNAL OF MEDICAL GENETICS PART A    | 2016 |
| 36 | Germinal mosaicism in a sample of families with duchenne/becker muscular dystrophy with Partial Deletions in the DMD Gene                                       | ARIADNA ESTELA GONZALEZ DEL ANGEL MIGUEL ANGEL ALCANTARA ORTIGOZA Bermúdez-López C. et al.               | GENETIC TESTING AND MOLECULAR BIOMARKERS       | 2014 |
| 37 | A patient with trisomy 13 mosaicism with an unusual skin pigmentary pattern and prolonged survival                                                              | ARIADNA ESTELA GONZALEZ DEL ANGEL Estandía-Ortega B. Gaviño-Vergara A. et al.                            | PEDIATRIC DERMATOLOGY                          | 2014 |
| 38 | CTNS gene analysis emphasizes diagnostic value of eye examination in patients with cystinosis                                                                   | MIGUEL ANGEL ALCANTARA ORTIGOZA ARIADNA ESTELA GONZALEZ DEL ANGEL Martínez-Bernal A.B. et al.            | Journal Of Pediatric Genetics                  | 2013 |
| 39 | Molecular analysis of the PAX8 gene in a sample of Mexican patients with primary congenital hypothyroidism: Identification of the recurrent p.Arg31His mutation | MIGUEL ANGEL ALCANTARA ORTIGOZA ARIADNA ESTELA GONZALEZ DEL ANGEL Martínez-Cruz V. et al.                | CLINICAL ENDOCRINOLOGY                         | 2012 |

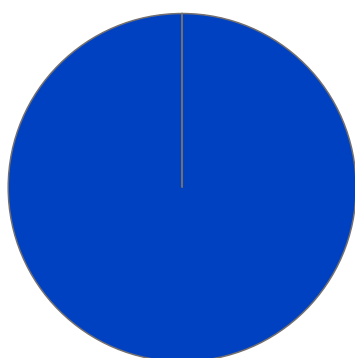
**ARIADNA ESTELA GONZALEZ DEL ANGEL**

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|----|------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------|---------------------------------------------|------|
| 40 | Phenotype?genotype discrepancy due to a 5.5-kb deletion in the GALT gene                                                           | ARIADNA ESTELA GONZALEZ DEL ANGEL JOSE ANTONIO VELAZQUEZ ARAGON MIGUEL ANGEL ALCANTARA ORTIGOZA et al. | Jimd Reports                                | 2012 |
| 41 | Trisomy of the short arm of chromosome 5 due to a de novo inversion and duplication (5)(p15.3 p13.3)                               | MIGUEL ANGEL ALCANTARA ORTIGOZA ALESSANDRA CARNEVALE CANTONI ARIADNA ESTELA GONZALEZ DEL ANGEL et al.  | AMERICAN JOURNAL OF MEDICAL GENETICS PART A | 2005 |
| 42 | Carrier detection and prenatal molecular diagnosis in a duchenne muscular dystrophy family without any affected relative available | MIGUEL ANGEL ALCANTARA ORTIGOZA ARIADNA ESTELA GONZALEZ DEL ANGEL ALESSANDRA CARNEVALE CANTONI et al.  | ANN GENET-PARIS                             | 2001 |

**ARIADNA ESTELA GONZALEZ DEL ANGEL**

**LIBROS Y CAPITULOS CON ISBN**

**Obras con registro ISBN**



■ Libros completos: 1 (100.00%)

| # | Título                                             | Autores                                                                                         | Alcance           | Año  | ISBN          |
|---|----------------------------------------------------|-------------------------------------------------------------------------------------------------|-------------------|------|---------------|
| 1 | Síndrome de down de la genética a la neurobiología | SARA FRIAS VAZQUEZ<br>VICTORIA DEL CASTILLO<br>RUIZ ARIADNA ESTELA<br>GONZALEZ DEL ANGEL et al. | Libro<br>Completo | 2019 | 9786073013666 |



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**Reporte individual**



**ARIADNA ESTELA GONZALEZ DEL ANGEL**

**PARTICIPACIÓN EN PROYECTOS**

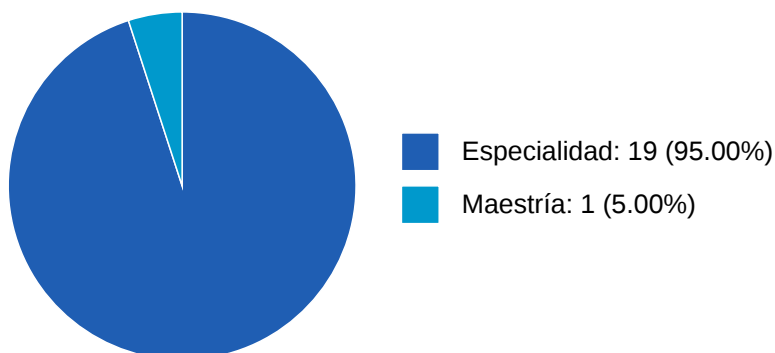
**No se encuentran registros en la base de datos de SISEPRO asociados a:**

**ARIADNA ESTELA GONZALEZ DEL ANGEL**

**ARIADNA ESTELA GONZALEZ DEL ANGEL**

**PARTICIPACIÓN EN TESIS**

**Histórico de Colaboraciones en Tesis**



| # | Título del documento                                                                                                                                                                    | Tipo de Tesis         | Sinodales                          | Autores                                                     | Entidad                                                             | Año  |
|---|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------|------------------------------------|-------------------------------------------------------------|---------------------------------------------------------------------|------|
| 1 | Caracterización molecular de pacientes con complejo de esclerosis tuberosa mediante amplificación múltiple dependiente de ligamiento (MLPA)                                             | Tesis de Especialidad | ARIADNA ESTELA GONZALEZ DEL ANGEL, | MIRIAM ERANDI REYNA FABIAN, Lechuga Becerra, Lorena,        | Facultad de Medicina,                                               | 2018 |
| 2 | Prevalencia de mutaciones patogénicas en el gen IRF6 responsables del síndrome de Van der Woude en pacientes mexicanos con diagnóstico clínico de labio paladar hendido no sindromático | Tesis de Especialidad | ARIADNA ESTELA GONZALEZ DEL ANGEL, | JOSE ANTONIO VELAZQUEZ ARAGON, Apam Garduño, David Alfonso, | Escuela Nacional de Enfermería y Obstetricia, Facultad de Medicina, | 2017 |
| 3 | Estudio molecular del gen candidato HOXA2 en una muestra de pacientes mexicanos con espectro facio-aurículo-vertebral (EFAV)                                                            | Tesis de Maestría     | ARIADNA ESTELA GONZALEZ DEL ANGEL, | Estandía Ortega, Bernardette,                               | Facultad de Medicina,                                               | 2017 |

**ARIADNA ESTELA GONZALEZ DEL ANGEL**

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|---|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------|----------------------------------|----------------------------------------------------------------------|-----------------------|------|
| 4 | Análisis molecular de los genes FOXE1 y NKX2.5 en pacientes con hipotiroidismo congénito por disgenesia atendidos en el Instituto Nacional de Pediatría : reporte preliminar de mutaciones germinales en el gen TSHR | Tesis de Especialidad | MIGUEL ANGEL ALCANTARA ORTIGOZA, | ARIADNA ESTELA GONZALEZ DEL ANGEL, González Núñez, Aidy,             | Facultad de Medicina, | 2016 |
| 5 | Mutaciones germinales en el gen NKX 2-1 (TTF-1) : ¿una causa frecuente de hipotiroidismo congénito por disgenesia?                                                                                                   | Tesis de Especialidad | MIGUEL ANGEL ALCANTARA ORTIGOZA, | ARIADNA ESTELA GONZALEZ DEL ANGEL, González Núñez, Aidy,             | Facultad de Medicina, | 2015 |
| 6 | CTNS gene study emphasizes diagnostic value of eye examination in cystinosis                                                                                                                                         | Tesis de Especialidad | MIGUEL ANGEL ALCANTARA ORTIGOZA, | ARIADNA ESTELA GONZALEZ DEL ANGEL, Martínez Bernal, Astrid Berenice, | Facultad de Medicina, | 2013 |
| 7 | Caracterización de mutaciones y polimorfismos en el gen nkx2.5 en una muestra de pacientes mexicanos con Síndrome de Down y defectos de septación cardiaca                                                           | Tesis de Especialidad | MIGUEL ANGEL ALCANTARA ORTIGOZA, | ARIADNA ESTELA GONZALEZ DEL ANGEL, Morales Toquero, Rodrigo,         | Facultad de Medicina, | 2012 |
| 8 | Estudio de asociación de labio y paladar hendido con polimorfismos en el gen MTHFR en pacientes mexicanos                                                                                                            | Tesis de Especialidad | MIGUEL ANGEL ALCANTARA ORTIGOZA, | ARIADNA ESTELA GONZALEZ DEL ANGEL, Estandia Ortega, Bernardette,     | Facultad de Medicina, | 2011 |

**ARIADNA ESTELA GONZALEZ DEL ANGEL**

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|----|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------|------------------------------------|--------------------------------------------------------------------|-----------------------|------|
| 9  | Análisis molecular del gen TTF-2 en pacientes con hipotiroidismo congénito por disgenesia                                                                                                      | Tesis de Especialidad | MIGUEL ANGEL ALCANTARA ORTIGOZA,   | ARIADNA ESTELA GONZALEZ DEL ANGEL, Godínez Chaparro, Juan Alberto, | Facultad de Medicina, | 2011 |
| 10 | Caracterización de mutaciones y polimorfismos en el gen CRELD1 en una muestra de pacientes mexicanos con síndrome de Down y defectos de septación cardiaca : estudio preliminar en 5 pacientes | Tesis de Especialidad | ARIADNA ESTELA GONZALEZ DEL ANGEL, | Álvarez Gómez, Rosa María,                                         | Facultad de Medicina, | 2011 |
| 11 | Análisis molecular de la delección de 5.5 kb en el gen galt en pacientes con galactosemia                                                                                                      | Tesis de Especialidad | ARIADNA ESTELA GONZALEZ DEL ANGEL, | JOSE ANTONIO VELAZQUEZ ARAGON, Frank Márquez, Nadine,              | Facultad de Medicina, | 2010 |
| 12 | Descripción clínica y estudio molecular de una familia con esclerosis tuberosa con expresividad variable                                                                                       | Tesis de Especialidad | ARIADNA ESTELA GONZALEZ DEL ANGEL, | NANCY HERNANDEZ MARTINEZ, Cervantes Blanco, Jorge Mauricio,        | Facultad de Medicina, | 2009 |
| 13 | Síndrome de Bruck : reporte de un caso en el Instituto Nacional de Pediatría y revisión de la literatura                                                                                       | Tesis de Especialidad | ROSALIA GARZA ELIZONDO,            | ARIADNA ESTELA GONZALEZ DEL ANGEL, Lagos Córdova, Elda Yara,       | Facultad de Medicina, | 2009 |
| 14 | Descripción clínica y análisis molecular de 15 exones del gen TSC2 en pacientes con esclerosis tuberosa                                                                                        | Tesis de Especialidad | ARIADNA ESTELA GONZALEZ DEL ANGEL, | Medina Luna, Paola,                                                | Facultad de Medicina, | 2008 |

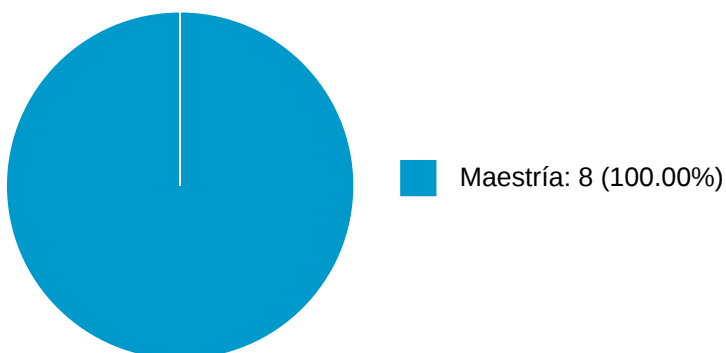
**ARIADNA ESTELA GONZALEZ DEL ANGEL**

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|----|-------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------|------------------------------------|---------------------------------------------------------|-----------------------|------|
| 15 | Diagnostico molecular de las mutaciones Q188R, K285N, N314D y S135L en pacientes con galactosemia clasica en el Instituto Nacional de Pediatría | Tesis de Especialidad | ARIADNA ESTELA GONZALEZ DEL ANGEL, | Monroy Santoyo, Susana,                                 | Facultad de Medicina, | 2008 |
| 16 | Sindrome de Kabuki : reporte de un caso en Mexico y revision de la literatura                                                                   | Tesis de Especialidad | ARIADNA ESTELA GONZALEZ DEL ANGEL, | Tinajero Esquivel, Maria Magdalena,                     |                       | 2007 |
| 17 | Presentacion de una adolescente mexicana con sindrome de Meier-Gorlin (Oreja-rotula-talla baja) y revision de la literatura                     | Tesis de Especialidad | ARIADNA ESTELA GONZALEZ DEL ANGEL, | Villarroel Cortés, Camilo Ernesto,                      |                       | 2004 |
| 18 | Estudio de la edad osea en pacientes con acondroplasia en el Instituto Nacional de Pediatría                                                    | Tesis de Especialidad | ARIADNA ESTELA GONZALEZ DEL ANGEL, | Arteaga Alcaraz, Maria Georgina,                        |                       | 2004 |
| 19 | Frecuencia de la mutacion PRO250ARG en el gen FGFR3 en pacientes con craneosinostosis                                                           | Tesis de Especialidad | ARIADNA ESTELA GONZALEZ DEL ANGEL, | LORENA SOFIA OROZCO OROZCO, Rasmussen Almaraz, Astrid,  |                       | 2000 |
| 20 | Estudio clinico y genetico de la microtia                                                                                                       | Tesis de Especialidad | ALESSANDRA CARNEVALE CANTONI,      | ARIADNA ESTELA GONZALEZ DEL ANGEL, Llano Rivas, Isabel, |                       | 1997 |

**ARIADNA ESTELA GONZALEZ DEL ANGEL**

**DOCENCIA IMPARTIDA**

**Histórico de docencia**



| # | Nivel titulación | Asignatura                          | Entidad              | Alumnos | Semestre |
|---|------------------|-------------------------------------|----------------------|---------|----------|
| 1 | Maestría         | TRABAJO DE INVESTIGACIÓN IV         | Facultad de Ciencias | 1       | 2017-2   |
| 2 | Maestría         | TRABAJO DE INVESTIGACION III-393972 | Facultad de Ciencias | 1       | 2017-1   |
| 3 | Maestría         | TRABAJO DE INVESTIGACION II         | Facultad de Ciencias | 1       | 2016-2   |
| 4 | Maestría         | TRABAJO DE INVESTIGACION I          | Facultad de Ciencias | 1       | 2016-1   |
| 5 | Maestría         | TRABAJO DE INVESTIGACION IV         | Facultad de Medicina | 1       | 2011-1   |
| 6 | Maestría         | TRABAJO DE INVESTIGACION III        | Facultad de Medicina | 1       | 2010-2   |
| 7 | Maestría         | TRABAJO DE INVESTIGACION II         | Facultad de Medicina | 2       | 2010-1   |
| 8 | Maestría         | TRABAJO DE INVESTIGACION            | Facultad de Medicina | 1       | 2009-2   |



**Sistema Integral de Información Académica**  
**Coordinación de Planeación, Evaluación y**  
**Simplificación de la Gestión Institucional**  
**Reporte individual**



**ARIADNA ESTELA GONZALEZ DEL ANGEL**

**PATENTES**

**No se encuentran registros en la base de datos de patentes asociados a:**

**ARIADNA ESTELA GONZALEZ DEL ANGEL**

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**FUENTES DE INFORMACIÓN**

**Internos**

| # | Información                                                          | Fuente | Sistema     | Periodo   |
|---|----------------------------------------------------------------------|--------|-------------|-----------|
| 1 | Grupos ordinarios y resumen de historias académicas                  | DGAE   | SIAE        | 2008-2025 |
| 2 | Nombramientos, datos generales, estímulos, premios y reconocimientos | DGAPA  | RUPA        | 2008-2025 |
| 3 | Producción Académica                                                 | CH     | Humanindex  | 2008-2021 |
| 4 | Producción Académica                                                 | CIC    | SCIC        | 2000-2017 |
| 5 | Proyectos                                                            | DGPO   | SISEPRO     | 2018-2022 |
| 6 | Tesis                                                                | DGB    | TESIUNAM    | 2008-2025 |
| 7 | Tutorías en Posgrado                                                 | CGEP   | SIIPosgrado | 2008-2021 |

**Externos**

| #  | Información             | Fuente          | Sistema      | Periodo   |
|----|-------------------------|-----------------|--------------|-----------|
| 8  | Documentos Indexados    | Elsevier        | Scopus       | 2008-2025 |
| 9  | Documentos Indexados    | Thomson Reuters | WoS          | 2008-2025 |
| 10 | Obras con registro ISBN | INDAUTOR        | Agencia ISBN | 2008-2025 |
| 11 | Patentes                | IMPI            | SIGA         | 2008-2024 |