# Stanford



# Christina (Christy) Tise, MD, PhD

Assistant Professor of Pediatrics (genetics)
Pediatrics - Medical Genetics

# **CLINICAL OFFICE (PRIMARY)**

• Center for Academic Medicine- Genetics

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# Bio

# BIO

Dr. Christina (Christy) Tise is a physician scientist and Assistant Professor in the Division of Medical Genetics at Stanford with subspecialty training in Clinical Biochemical Genetics. Dr. Tise has developed multiple research projects focused on the clinical impact of biochemical genetic conditions in pregnancy and newborn health, including a project focused on unforeseen diagnoses in individuals initially identified through state newborn screening which has resulted in a number of publications.

Dr. Tise also researches the genetic etiologies of recurrent pregnancy loss and the impact of inherited metabolic conditions on human reproduction. She is involved in several research initiatives including contributing to the development of TRIOS, a multi-site, NIH-funded research study to evaluate the genetic causes of recurrent pregnancy loss. In serving as the primary research mentor for a recent Masters of Genetic Counseling graduate, Dr. Tise's research on carrier and newborn screening has highlighted areas of ancestry-related healthcare inequities specific to the field of Medical Genetics.

Dr. Tise's primary academic and advocacy interests are embodied in this work, specifically the overlap between biochemical and molecular analysis, and the clinical utility of innovative technologies for diagnosis and treatment of genetic disease. This is an unbelievably thrilling time for the field of Medical Genetics, as it promises immense progress and opportunity for all fields of medicine, and Dr. Tise is determined, honored, and incredibly excited to be a part of it!

Research interests: newborn screening, carrier screening, prenatal screening, genetics of recurrent pregnancy loss, biochemical genetics, novel gene discovery, variant interpretation, founder populations, diagnostic genetic testing, bioethics, GWAS/ExWAS

# **CLINICAL FOCUS**

- Biochemical Genetics
- Newborn Screening
- Reproductive Genetics
- Perinatal Genetics
- Clinical Genetics and Genomics

#### ACADEMIC APPOINTMENTS

- Assistant Professor University Medical Line, Pediatrics Medical Genetics
- Member, Maternal & Child Health Research Institute (MCHRI)

#### HONORS AND AWARDS

- Golden Helix Faculty Teaching Award, Medical Genetics Residency Program (2023)
- Pfizer/ACMGF Next Generation Clinical Laboratory Biochemical Genetics Fellowship Award, American College of Medical Genetics (2021)
- Henry Christian Award, American Federation for Medical Research (AFMR) (2020)
- Outstanding Recent Biochemistry Undergraduate Alumni Award, Virginia Tech (2018)
- J. Edmund and Kathryn S. Bradley Award for Excellence in Pediatrics, University of Maryland School of Medicine (2018)
- Commencement Speaker for Epidemiology and Human Genetics Graduate Program, University of Maryland School of Medicine (2016)
- 2016 PhD Thesis of the Year Award, Graduate Program in Life Sciences, University of Maryland School of Medicine (2016)

#### BOARDS, ADVISORY COMMITTEES, PROFESSIONAL ORGANIZATIONS

- Member, Stanford MCHRI Education Committee (2022 present)
- Member, American College of Medical Genetics Membership Committee (2021 present)

#### PROFESSIONAL EDUCATION

- Board Certification: Clinical Genetics and Genomics, American Board of Medical Genetics and Genomics (2021)
- Board Certification: Clinical Biochemical Genetics, American Board of Medical Genetics and Genomics (2023)
- MD, University of Maryland School of Medicine, Medicine (2018)
- PhD, University of Maryland School of Medicine , Human Genetics and Genomic Medicine (2016)
- Fellowship: Stanford University Clinical Biochemical Genetics Fellowship (2022) CA
- Residency: Stanford University Division of Medical Genetics (2021) CA
- Internship: Stanford University Pediatric Residency at Lucile Packard Children's Hospital (2019) CA
- Medical Education: University of Maryland Office of the Registrar (2018) MD
- BS, Virginia Tech, Biochemistry (2009)

# Research & Scholarship

# **CLINICAL TRIALS**

· Pregnancy and Developmental Outcomes After Transfer of Reportedly Aneuploid or Mosaic Embryos, Recruiting

# **Teaching**

# **COURSES**

#### 2023-24

• Introduction to Medical Genetics: GENE 272 (Aut)

#### 2022-23

• Introduction to Medical Genetics: GENE 272 (Aut)

# **Publications**

# **PUBLICATIONS**

• Molecular Testing in Newborn Screening: VUS Burden Among True Positives and Secondary Reproductive Limitations via Expanded Carrier Screening Panels. Genetics in medicine: official journal of the American College of Medical Genetics

Cook, S., Dunn, E., Kornish, J., Calderwood, L., Campion, M., Cusmano-Ozog, K. P., Tise, C. G. 2023; 101055

- MT-ATP6 mitochondrial disease identified by newborn screening reveals a distinct biochemical phenotype. American journal of medical genetics. Part A
  Tise, C. G., Verscaj, C. P., Mendelsohn, B. A., Woods, J., Lee, C. U., Enns, G. M., Stander, Z., Hall, P. L., Cowan, T. M., Cusmano-Ozog, K. P.
  2023
- Neonatal lupus is a novel cause of positive newborn screening for X-linked adrenoleukodystrophy. American journal of medical genetics. Part A
  Niehaus, A. D., Mendelsohn, B. A., Zimmerman, B., Lee, C. U., Manning, M. A., Cusmano-Ozog, K. P., Tise, C. G.
  2023
- Medical genetics training in the COVID-19 era: A resident's perspective. American journal of medical genetics. Part A
  Tise, C. G.

2021

• Unexpected diagnoses in patients with abnormal newborn screening

Tise, C., Velez-Bartolomei, F., Morales, J., Lee, C., Bernstein, J., Enns, G. ACADEMIC PRESS INC ELSEVIER SCIENCE.2021: S354

• Aicardi-Goutières syndrome may present with positive newborn screen for X-linked adrenoleukodystrophy. American journal of medical genetics. Part A Tise, C. G., Morales, J. A., Lee, A. S., Velez-Bartolomei, F. n., Floyd, B. J., Levy, R. J., Cusmano-Ozog, K. P., Feigenbaum, A. S., Ruzhnikov, M. R., Lee, C. U., Enns, G. M.

2021

• Genetics of recurrent pregnancy loss: a review. Current opinion in obstetrics & gynecology

Tise, C. G., Byers, H. M.

2021

 From Genotype to Phenotype: Nonsense Variants in SLC13A1 Are Associated with Decreased Serum Sulfate and Increased Serum Aminotransferases G3-GENES GENOMES GENETICS

Tise, C. G., Perry, J. A., Anforth, L. E., Pavlovich, M. A., Backman, J. D., Ryan, K. A., Lewis, J. P., O'Connell, J. R., Yerges-Armstrong, L. M., Shuldiner, A. R. 2016; 6 (9): 2909-2918

• Monozygotic twins discordant for a congenital cranial dysinnervation disorder with features of Moebius syndrome. American journal of medical genetics.

Gates, R. W., Webb, B. D., Stevenson, D. A., Jabs, E. W., DeFilippo, C., Ruzhnikov, M. R., Tise, C. G. 2023

- A homozygous Gly470Ala variant in PEX6 causes severe Zellweger spectrum disorder. American journal of medical genetics. Part A Galarreta, C. I., Wong, K., Carmichael, J., Woods, J., Tise, C. G., Niehaus, A. D., Schildt, A. J., Verscaj, C. P., Cusmano-Ozog, K. P. 2023
- BIOCHEMICAL, MOLECULAR, AND CLINICAL CHARACTERISTICS OF PEROXISOMAL DISORDERS DETECTED BY CALIFORNIA NEWBORN SCREENING (NBS) PROGRAM

Beltran, C., Abdenur, J., Chang, R., Barrick, R., Spongberg, R., Tise, C. G., Niehaus, A. D., Enns, G. ACADEMIC PRESS INC ELSEVIER SCIENCE.2023: 72

 Creatine Transporter Deficiency Presenting as Failure to Thrive: A Case Report of a Novel SLC6A8 Variant Causing a Treatable but Likely Underdiagnosed Genetic Disorder. Journal of investigative medicine high impact case reports

Tise, C. G., Palma, M. J., Cusmano-Ozog, K. P., Matalon, D. R.

2023; 11; 23247096231154438

Recent tPA administration can cause pseudo-hyperargininemia and may mimic arginase deficiency or arginine supplementation. JIMD reports
Cusmano-Ozog, K. P., Renck, A. K., Tise, C. G.

2022; 63 (6): 563-567

 Short Bones, Renal Stones, and Diagnostic Moans: Hypercalcemia in a Girl Found to Have Coffin-Lowry Syndrome. Journal of investigative medicine high impact case reports

Tise, C. G., Matalon, D. R., Manning, M. A., Byers, H. M., Grover, M.

2022; 10: 23247096221101844

 MITOCHONDRIAL-ATP6-ASSOCIATED DISEASE PRESENTS WITH DISTINCT PATTERN ON NEWBORN SCREENING: SHOULD IT BE INCLUDED AS A SECONDARY CONDITION?

Tise, C., Mendelsohn, B., Lee, C., Woods, J., Hall, P., Tang, H., Rinaldo, P., Cowan, T., Cusmano-Ozog, K.

ACADEMIC PRESS INC ELSEVIER SCIENCE.2022: 247-248

• SUCCESSFUL CARDIAC TRANSPLANTATION AND LONG-TERM FOLLOW-UP IN DNAJC19-ASSOCIATED DILATED CARDIOMYOPATHY WITH ATAXIA

Tahata, S., Tise, C., Floyd, B., Cusmano-Ozog, K., Ruzhnikov, M., Enns, G.

ACADEMIC PRESS INC ELSEVIER SCIENCE.2022: 302

• Biochem for the Win! The added value of biochemical genetic testing for diagnosis and variant interpretation in the genomic era

Tise, C., Grand, K., Corado, J., Gates, R., Graham, J., Enns, G., Gomez-Ospina, N., Mak, J., Cowan, T., Cusmano-Ozog, K. ELSEVIER SCIENCE INC.2022: S24

Clinical characterization of a new individual with mild SC4MOL deficiency: diagnostic and therapeutic implications JOURNAL OF TRANSLATIONAL
GENETICS AND GENOMICS

Morales, J., Curry, C. J., Tise, C. G., Kratz, L., Enns, G. M.

2022; 6 (2): 257-265

 Profound neonatal lactic acidosis and renal tubulopathy in a patient with glycogen storage disease type IX#2 secondary to a de novo pathogenic variant in PHKA2. Molecular genetics and metabolism reports

Morales, J. A., Tise, C. G., Narang, A., Grimm, P. C., Enns, G. M., Lee, C. U.

2021; 27: 100765

• Case 1: Rapidly Rising Bilirubin Level in a 3-day-old Term Infant. NeoReviews

Tise, C. G., Joshi, N. S., Erice-Taganas, A. D., Blecharczyk, E. M.

2020; 21 (10): e687-e690

• nonsense variants on DHEA homeostasis. Molecular genetics and metabolism reports

Tise, C. G., Anforth, L. E., Zhou, A. E., Perry, J. A., McArdle, P. F., Streeten, E. A., Shuldiner, A. R., Yerges-Armstrong, L. M. 2017; 10: 84-91

• Are Patients with Psychogenic Movement Disorders More Likely to be Healthcare Workers? MOVEMENT DISORDERS CLINICAL PRACTICE

Perry, C. G., Holmes, K. G., Gruber-Baldini, A. L., Anderson, K. E., Shulman, L. M., Weiner, W. J., Reich, S. G.

2017; 4 (1): 62–67

• Educational Innovations in Clinical Pharmacogenomics CLINICAL PHARMACOLOGY & THERAPEUTICS

Perry, C. G., Maloney, K. A., Beitelshees, A. L., Jeng, L. J., Ambulos, N. P., Shuldiner, A. R., Blitzer, M. G.

2016; 99 (6): 582-584

Pharmacometabolomics reveals that serotonin is implicated in aspirin response variability. CPT: pharmacometrics & systems pharmacology

Ellero-Simatos, S., Lewis, J. P., Georgiades, A., Yerges-Armstrong, L. M., Beitelshees, A. L., Horenstein, R. B., Dane, A., Harms, A. C., Ramaker, R., Vreeken, R. J., Perry, C. G., Zhu, H., Sànchez, et al

2014; 3

• Pharmacogenomics of Anti-platelet and Anti-coagulation Therapy CURRENT CARDIOLOGY REPORTS

Fisch, A. S., Perry, C. G., Stephens, S. H., Horenstein, R. B., Shuldiner, A. R.

2013; 15 (7)

Pharmacogenomics of anti-platelet therapy: how much evidence is enough for clinical implementation? JOURNAL OF HUMAN GENETICS

Perry, C. G., Shuldiner, A. R.

2013; 58 (6): 339-345