Register Today and Help Advance Research

Join thousands of patients living with Immune Thrombocytopenia (ITP) from across the globe and participate in the Platelet Disorder Support Association (PDSA) ITP Natural History Study Registry to advance research and improve the quality of life for ITP patients.

Enroll Today!

Enroll today in the ITP Natural History Study Registry at itpstudy.org!

For more information call PDSA at 1-877-PLATELET 877-528-3538 (toll free) or email research@pdsa.org.

Please rest assured that your identity will be protected in this secure, confidential HIPAA compliant registry, and there is no cost for you to participate.

Thank you for Making the World of ITP a more manageable place to be!
What is the ITP Registry?

The ITP Natural History Study Registry is an international patient-consented registry of individuals with ITP. The registry aims to collect data on the natural progression of ITP, enabling PDSA to gather data on diagnosis and treatment, management of care, quality of life, and clinician reporting. The registry is administered by PDSA and overseen by the National Organization for Rare Disorders, the U.S. Food and Drug Administration, and a committee of leading hematologists, ITP patients, and caregivers.

Why Register?

The ITP Registry will provide health care professionals and researchers with first-hand information about Immune Thrombocytopenia (ITP), individually and as a disease population, to increase our understanding of the progression of ITP over time. Natural history studies yield vital information essential to clinical trial design, such as biomarkers, demographics, important clinical symptoms, genetic and environmental variables, and patient perspectives.

Help Advance Research and Make a Difference

Your participation in the ITP Natural History Study Registry will increase what we know about ITP, help healthcare professionals improve treatments, and allow researchers to design better clinical trials, including development and testing of new treatments. By contributing your ITP story, you’re taking the first step to help accelerate research into new therapies for ITP and potentially a cure.