

The Peak Registry A Global Longitudinal Study of PK Deficiency

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An Unwavering Commitment to Patients with Pyruvate **Kinase Deficiency**

Introducing a study dedicated to advancing knowledge of:

- Natural history
- Disease burden
- Patient outcomes
- Current and historical treatment and outcomes

The Peak Registry is a global longitudinal study of patients diagnosed with pyruvate kinase deficiency

Participation in the Peak Registry benefits the community of patients and practitioners seeking information about disease burden, treatment outcomes and clinical practices. Benefits may include:

- Increased awareness among patients and practitioners of the signs, symptoms and diagnosis of PK deficiency
 - Knowledge of current supportive care options and opportunities to better manage the challenges faced by patients with PK deficiency
 - Collaboration with practitioners around the globe who are dedicated to their patients with PK deficiency
 - Opportunity to contribute to the medical discourse around PK deficiency



To learn more, or to remotely enroll a patient in the Peak Registry, please visit PeakRegistry.com or email medinfo@agios.com