Infantile Neuroaxonal Dystrophy (INAD) Clinical Trial Ground Transportation Patient Assistance Program

For patients participating in Retrotrope Multicenter Clinical Trial Phase 2/3 RT001

What is the purpose of this program?

NORD’s Infantile Neuroaxonal Dystrophy Ground Transportation Patient Assistance Program provides patients participating in the Retrotrope RT001 study in children with Infantile Neuroaxonal Dystrophy with financial assistance for ground transportation related to participation in this study. Individuals receiving assistance in this Program are referred to NORD by Study Site Coordinators.

About the Program

NORD Patient Services representatives will interface directly with study site coordinators to help facilitate seamless coordination of ground transportation arrangements and provide support on behalf of the Study subjects.

The Study site will notify NORD when a new subject enters the Study so that our dedicated Patient Services Representatives may assist the new subject with ground transportation arrangements.

NORD will pre-book and pre-pay expenses for Study participants ground transportation between airport and lodging (hotel) and Study Site for one (1) Study Site visit.

Our goal is to ensure patients are supported and their burden is lessened with regards to the cost and arrangement of ground transportation to and from their study visit.

For additional information or, if you are in need of such assistance, please contact NORD at:

Phone: 877.291.7865
Fax: 203.405.0534
Email: INADTravel@rarediseases.org

NORD
The National Organization for Rare Disorders (NORD), a 501(c)(3) organization, is an independent charity dedicated to the identification, treatment and cure of rare “orphan” diseases such as Infantile Neuroaxonal Dystrophy through education, advocacy, research and patient service programs.

NORD was founded by families struggling to obtain access to treatments and whose advocacy for change led to the passage of the Orphan Drug Act in 1983.

Funding for NORD comes from a variety of sources including corporate donations, foundation grants, public contributions, and membership dues.

Learn more about NORD by visiting www.rarediseases.org

NORD does not recommend or endorse any particular medical treatment but encourages patients to seek the advice of their clinicians. Donations to NORD for this and other programs may be made by contacting NORD at rarediseases.org

©2018 NORD. All rights reserved. NORD and ‘Alone we are rare…’ are registered trademarks of The National Organization for Rare Disorders. NORD is a registered 501(c)(3) charity.