Connecting Patients

WE ARE COMMITTED TO BRINGING THE BEST NF CARE TO PATIENTS.

NF CLINIC NETWORK
More than 15,000 patients a year visit our NF Clinic Network.

NF REGISTRY
Join the thousands of people who have enrolled in the NF Registry, a critical tool in matching NF patients to clinical trials.

To join or learn more, go to nfregistry.org.

RESOURCE LIBRARY
The CTF Resource Library connects patients with the education and information they need through patient brochures, educational webinars, and recorded symposia.

Connecting Researchers

WE ENSURE THAT THE BEST NF PROFESSIONALS FROM AROUND THE WORLD ARE COLLABORATING ON THE LATEST NF RESEARCH.

NF CONFERENCE
Our annual gathering of NF researchers and clinicians advances collaboration, research, and care for people with NF.

NF RESEARCH
In addition to numerous research grant opportunities, CTF initiates and funds scientific collaborations and industry partnerships that accelerate NF research and drug development.

To make a donation, become involved, or learn about Foundation activities in your area, please visit our website at ctf.org.

ABOUT THE CHILDREN'S TUMOR FOUNDATION

The CTF Resource Library connects patients with the education and information they need through patient brochures, educational webinars, and recorded symposia.

www.ctf.org | info@ctf.org
1-800-323-7938
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ACROSS THE COUNTRY, THE CHILDREN’S TUMOR FOUNDATION PROVIDES SUPPORT TO NF PATIENTS AND FAMILIES, RAISING NF AWARENESS AND FUNDS FOR RESEARCH.

About NF
NF refers to a group of genetic conditions that collectively affect 1 in every 2,000 births. NF includes neurofibromatosis type 1 (NF1) and all forms of schwannomatosis (SWN), including NF2-related schwannomatosis (NF2), formerly called neurofibromatosis type 2. These conditions cause tumors to grow on nerves throughout the body, and can lead to deafness, blindness, disfigurement, bone abnormalities, learning disabilities, disabling pain, and cancer. NF is genetic, which means that it is not the result of anything a parent did or did not do, and it is not contagious. NF occurs in all populations equally, and there is not yet a cure.

About Children’s Tumor Foundation
Founded in 1978, the Children’s Tumor Foundation (CTF) began as the first grassroots organization solely dedicated to finding treatments for NF. Today, CTF is a highly recognized global nonprofit foundation, the leading force in the fight to end NF, and a model for other innovative research endeavors.

Our Mission
Drive research, expand knowledge, and advance care for the NF community.

Our Vision
End NF.

Connecting Communities

SHINE A LIGHT NF WALK
Shine a Light NF Walks bring families and friends together to raise funds and build community.

FIGHT NF YOUR WAY
Make a difference in the fight to end NF by creating a fundraiser in your community, or joining one of our many annual special events.

CUPID'S UNDIE RUN
Cupid’s is a mile(ish) fun run in which brightly adorned, underwear-clad team members run outdoors for NF awareness.

NF PATIENT SYMPOSIA
Whether at our national NF Summit or a local NF clinic symposium, families hear the latest in NF research and connect with other NF families.

NF CAMP
Young people with NF gather in the summer for fun, friendship, and a lifetime of memories.

NF ENDURANCE
Professional and amateur athletes run, bike, and swim in endurance events to raise awareness and funds for NF research.