



At this half year mark in 2018, we take a moment to reflect on what we have done, and what still lies ahead.

Our Mission, Our Goals

At ***the Foundation for Peripheral Neuropathy***, we look at our mission as four pillars:

AWARENESS



We believe by spreading the word about PN, we can help more people as well as attract more attention to the disease and more research dollars for a cure.

In the first half of 2018, we:

- Sponsored National Neuropathy Awareness Week.
- Increase our social media presence and developed posters, flyers, a video and other materials to raise awareness for PN.

We provide information to patients, caregivers, and healthcare providers on PN, and how to live with the condition so those affected can live better lives.

EDUCATION



In the first half of 2018, we:

- Held a Patient Conference in Washington DC reaching over 300 people either in attendance, through live-streaming or through website views.
- Provided over 50 tips and articles through our newsletters and emails.
- Updated our brochures on peripheral neuropathy and our website to continue to offer timely information to patients and their families.

SUPPORT



We provide avenues of support to patients and caregivers to make living with this condition more bearable.

In the first half of 2018, we:

- Provided one-on-one support to patients through our info line, addressing well over 1000 patients' concerns in the first half of the year.
- Organized what will be a regular virtual gathering of our Support Group leaders to provide dialogue and support for the more than 100 volunteer leaders providing support to others.

We believe that by facilitating research, we can play a part in the development of better treatments and, hopefully, a cure.

RESEARCH



In the first half of 2018:

- The Peripheral Neuropathy Research Registry continued to offer and gather data for research scientists. Patient registration reached over 1500.
- Through our ongoing Clinical Trial Awareness initiatives, 7 clinical trials or research studies were able to reach interested participants.

So much lies ahead!

- Plans are underway for the next patient conference in Fall 2018 as well as other educational programming.
- We see more activity in the research space and are looking for ways to increase our Clinical Trials Awareness Program to be a conduit between the scientific and the patient communities.
- The Peripheral Neuropathy Research Registry continues to expand both in the patient registrants and scientific attention.
- We continue to expand ways to provide much needed support to peripheral neuropathy patients, and gain awareness for peripheral neuropathy in both the scientific community and among the general populace.

Of course, all of this takes time, money and volunteers (as well as our very busy staff of three) to accomplish.

Your support is important to us. Together, we can look forward to an exciting second half of 2018 and beyond!

If you wish to continue to help support our mission, please visit our website, **www.foundationforpn.org**, or call (847) 883-9942.