



The mission of the National Lymphedema Network is to empower our community of lymphedema practitioners and patients by providing educational opportunities, networking opportunities, and interventional resources. We aim to inspire clinicians to provide world-class care for people with lymphedema by promoting best practice guidelines and standards of care.

EDUCATION

- **Patient Summit** - Annual one-day conference for all people living with lymphatic disease. Learn from experts in the field, network opportunities, learn from the industry partners on compression, devices, and more!
- **Lymphedema Treatment Symposium** - Annual two-day free virtual symposium that covers what lymphedema is and the treatment involved with Complete Decongestive Therapy.
- **Live Webinars** - One-hour educational sessions that are free to the community and offers the opportunity to ask questions to the speaker.
- **On-Demand Education Library** - Watch sponsored and past webinars offered to the community at any time for free!
- **Website** - The NLN provides information on our website about lymphedema and treatment.
- **Printed Materials** - A request for a printed informational booklet can be made on the NLN's website.

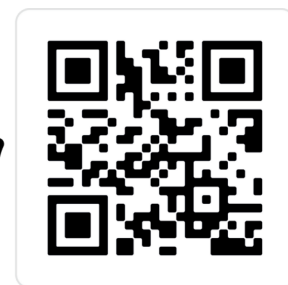
**Be on the lookout in 2026 for NEW materials!*

SUPPORT

- **Clinical Focus Groups** - Two discussion groups for NLN Members to ask questions about lymphedema, treatment, and creating healthy habits.
- **Support Team** - Call or email the NLN to ask questions and speak with certified therapists about any lymphedema related topic.
- **Lymphedema Therapy Directory** - Find a certified lymphedema therapist near you.
- **Lymphedema Sponsor & Dealer Directory** - Find a company near you who sells compression products or devices.

VOLUNTEER

- **Patient Committee** - Comprised of both patients and clinicians who help plan the annual in-person Summit and aide in the development of programs targeted for people living with lymphatic disease.



Become a Member



Introducing the Lymphedema Education, Advocacy, & Development (LEAD) program. This program is designed to educate and empower individuals with lymphedema to become confident advocates, raising awareness, influencing policy, and supporting others within the lymphedema community.

Participants will learn critical skills, gain valuable resources, and join a network of passionate advocates working together to make a meaningful impact.



Program to be launched in 2026!

Scan the QR code to express your interest and stay informed about the program launch!