Since 1998, LE&RN has been dedicated to the fight against lymphedema and lymphatic disease. Lymphedema and other lymphatic diseases affect up to 10 million Americans—more than AIDS, Parkinson’s disease, Muscular Dystrophy, Multiple Sclerosis and ALS combined. Yet, lymphatic diseases routinely go undiagnosed and untreated, and research is grossly underfunded. Together, LE&RN and its members are determined to change this.

EDUCATION
- Free Livestream Symposium Series featuring the world’s leading authorities
- The LE&RN/Lymph Notes Lymphedema Therapist Scholarships
- Young Investigator Scholarship Awards
- Weekly Newsletter
- Free Ask the Experts web feature

RESEARCH
- Research Fellowship Awards
- The LE&RN National Lymphatic Disease & Lymphedema Patient Registry & Tissue Bank
- Lymphatic Research & Biology (the world’s only peer review lymphatic journal)
- NIH Collaboration & research funding initiatives

ADVOCACY
- The Face of Lymphedema Challenge
- Celebrity spokesperson engagement
- Annual Walk to Fight Lymphedema & Lymphatic Diseases
- State Chapters
- Political Awareness Initiative
- Lymphedema Treatment Act advocacy
- Activist social media
“I’m KATHY BATES. I had a double mastectomy two years ago and as a result I have lymphedema in both arms. If we want to win this fight against lymphedema and lymphatic disease, we need to stand together and educate the world. Join me. I urge you to become a member of LE&RN. Then, please take The Face of Lymphedema Challenge to promote awareness and research.”

We will never cure lymphedema and lymphatic disease if we stay invisible. We need to stand together and we need to have our friends and family in this fight as well. Kathy Bates can’t do this alone. We need to put a face on these diseases so that they become a national priority. The days of suffering in silence are behind us. Join Kathy and LE&RN. Become a Member of LE&RN today. Simply set up a recurring monthly donation at LymphaticNetwork.org. Just click on “Become a Member”—$60 a year, $5 a month.

Then, join the Face of Lymphedema Challenge. See Kathy’s video testimony on LE&RN’s YouTube channel. Then make your video saying, “I have lymphedema” or “Someone I love has a lymphatic disease” and send it to LE&RN at:

Challenge@LymphaticNetwork.org

A world to change—together!