2017 ANNUAL REPORT
LE&RN is at the forefront of raising awareness of lymphatic diseases, such as lymphedema and lipedema. Founded in 1998, LE&RN provides annual grants to Research Fellows at the world’s leading universities, putting lymphatic diseases on the radar in the medical community, in academia, in government, and with pharmaceutical companies.

Through LE&RN’s education programs, such as our Livestream Symposium Series, medical practitioners, therapists, patients, and family members can benefit from hearing presentations given by the world’s leading lymphatic and lymphedema experts.

LE&RN’s International Patient Registry and Biorepository provides researchers with access to information and bio-samples from patients with lymphatic diseases. The quarterly journal Lymphatic Research and Biology delivers the latest developments and advancements in lymphatic biology and pathology from the world’s leading biomedical investigators.

Through events such as our countrywide Walk to Fight Lymphedema & Lymphatic Diseases and through legislative advocacy efforts, LE&RN is building an extensive network of people seeking new treatment options and cures.

Lymphedema and other lymphatic diseases impact the lives of an estimated 10 million Americans and more than 250 million worldwide. Yet, these diseases remain invisible to most and are often misdiagnosed or left untreated, while funding for research remains limited. In 2017 LE&RN addressed three strategies: First, inaugurating the Lymphatic Forum conference to spur lymphatic research; Second, partnering with Harvard University Medical Schools to present the Lymphedema Symposium for medical practitioners; and Third, expanding global awareness through media and recognition through Research America.

Lymphatic—uncovering the links between lymphatic function, lymphatic diseases, and wellness
Education—serving as an essential educational resource for doctors, patients, and the public at large
& Research—searching for treatments and cures through our specialized research activities
Network—building an extensive network of participants and partners to advance advocacy
LEADING THE FIGHT AGAINST LYMPHATIC DISEASE AND LYMPHEDEMA

LE&RN is a not-for-profit corporation, founded in 1998, under the laws of the State of New York.

The Organization’s mission is to fight lymphatic diseases such as lymphedema and lipedema through education, research and advocacy. LE&RN seeks to accelerate the prevention, treatment, and cure of lymphatic diseases while bringing patients and medical professionals together to address the unmet needs surrounding lymphatic disorders.

Some activities of the organization during 2017 include the following:

- LE&RN expanded the International Patient Registry and Biorepository at Stanford University as a resource to stimulate research and support clinical trials and drug development.
- LE&RN continued its grant fellowship award program.
- LE&RN continued its educational and awareness outreach programs targeting academia, government, industry, and the medical and patient communities.
- LE&RN continued its bimonthly live-stream symposium series, which brings the world’s foremost authorities in lymphatic disease and lymphedema to patients, their families, and professionals.
- LE&RN continued expanding its website and social media to provide comprehensive updates of news in the field.
- LE&RN produced four issues of Lymphatic Research and Biology, an international, peer-reviewed biomedical journal providing the forum for the exchange of cutting-edge scientific developments in lymphatic science and medicine.
- LE&RN secured ongoing Congressional and National Institutes of Health support for lymphatic research.
- LE&RN continued the LE&RN/LymphNotes Lymphedema Scholarship Program to provide tuition support to therapists seeking certification in lymphedema therapy.
- LE&RN continued offering a web-based “Ask the Experts” feature that allows patients, their families, and professionals to directly ask questions to the most prominent authorities in the field.
- LE&RN spokesperson, Academy Award Winner Kathy Bates continued her advocacy and was honored by Research America.
- LE&RN expanded its Honorary Board, headed by Kathy Bates and including notable celebrities who support LE&RN’s mission. Dame Judy Dench joined.
- LE&RN’s Virtual Expo was updated to allow patients and professionals 24/7 access to available treatments and services.
- World Lymphedema Day celebrated its 2nd year with events across the globe.
- LE&RN outreach expanded to national audiences with pieces on CBS Sunday Morning, Dr. Oz, USA Today, NIH Medline Plus and a host of other outlets.
- LE&RN members met with the co-presidents of the Chan-Zuckerberg BioHub to discuss the need for research.
## LE&RN Timeline

### 2017
- LE&RN Spokesperson **Kathy Bates** honored by Research America with “Impact on Public Opinion Award”
- LE&RN co-organizes the inaugural biennial researcher event **Lymphatic Forum: Exploring the Lymphatic Continuum** in Chicago
- LE&RN partners with Harvard University Medical Schools for the first **Lymphedema Symposium** in Boston

### 2016
- March 6th officially established as **World Lymphedema Day** by U.S. Senate, New York legislature, and countries around the world
- **International Chapters** inaugurated
- LE&RN spearheads **$70M US Senate Appropriations** request sponsored by Senator Charles Schumer (NY)
- Spokesperson **Kathy Bates takes LE&RN's mission center stage** on a multitude of network television shows and printcover stories

### 2015
- LE&RN/Fat Disorder Research Society **Lipedema Postdoctoral Fellowship Awards** established
- LE&RN/NAVBO partnership to establish biennial conference in lymphatics
- **State Chapters** inaugurated

### 2014
- “**Ask the Experts**” web feature established to connect patients to leading authorities
- The LE&RN/LymphNotes Lymphedema Therapist Scholarship Program is founded
- **NYC Walk to Fight Lymphedema and Lymphatic Diseases** crosses Brooklyn Bridge for the first time
- The Face of Lymphedema Challenge and Membership Campaign are kicked off
- Academy Award-winning actress **Kathy Bates becomes LE&RN’s spokesperson**
- New York State Legislature awards funding for **National Patient Registry & Tissue Bank**

### 2013
- LRF becomes LE&RN with an expanded mission and opens New York City office
- Live-streamed **Medical & Research Symposium Series** established

### 2012
- LE&RN **National Lymphatic Disease and Lymphedema Tissue Bank** established
- **Breakthrough study**, blood test for diagnosis of lymphedema published—Stanford University School of Medicine Endowed Chair

### 2010
- First annual **Walk to Fight Lymphedema & Lymphatic Diseases**
- LE&RN **National Lymphatic Disease and Lymphedema Patient Registry** created

### 2008
- First-ever **Endowed Chair of Lymphatic Research and Medicine** at Stanford University School of Medicine established

### 2006
- **Postdoctoral Fellowship Awards** program inaugurated

### 2005
- **Grants and Awards Programs** established

### 2004
- The Biennial Gordon Research Conference Series, Molecular Mechanisms in Lymphatic Function and Disease established

### 2003
- **Lymphatic Research and Biology**, an international peer-reviewed journal, has its first printing

### 2002
- Congressional and National Institutes of Health (NIH) support for lymphatic research and lymphatic diseases secured

### 1998
- LE&RN is founded under the name **Lymphatic Research Foundation** to promote research, treatments and cures
LYMPHATIC EDUCATION & RESEARCH NETWORK

LYMPHATIC EDUCATION & RESEARCH NETWORK

• Philip Braginsky (Chair)
• Ken Cerini (Treasurer)
• Debbie Miles (Secretary)
• Dr. Joseph Dayan
• Dr. Riku Rautsola
• Dr. Sheila Ridner
• Dr. Doron Ilan

President and CEO
• William Repicci

Accountants
• Cerini & Associates, LLP

2017 FINANCIALS

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