"We need awareness" so that medical practitioners and therapists are prepared to treat all who come to them with LE and LD. We need awareness to inspire communities around the world to embrace this as a common cause. We need awareness to encourage our greatest minds to engage in research and our institutions to then make a priority of funding this research. And then we need to make finding a cure a global priority. Miracles await if we dedicate ourselves to finding them."

—William Repicci, President and CEO, LE&RN

Cover photo: Kathy Bates is honored as LE&RN’s National Spokesperson at the 24 Hour Plays Broadway Gala, New York City, November 2019.
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THE MISSION

Our mission is to fight lymphatic diseases (LD) and lymphedema (LE) through education, research, and advocacy. We seek to accelerate prevention, treatments, and cures while bringing patients and medical professionals together to address unmet needs.

Lymphatic diseases, such as lymphedema (LE), affect an estimated 10 million Americans and 250 million worldwide. In addition, up to 17 million Americans are estimated to have the related lymphatic disease called lipedema, which affects mostly women. More Americans have LE than AIDS, Parkinson’s disease, Muscular Dystrophy, Multiple Sclerosis, and ALS—combined. Yet, these diseases remain invisible to most and are often misdiagnosed or left untreated while funding for research remains limited.

Since the Lymphatic Education & Research Network (LE&RN) was founded in 1998, this internationally recognized nonprofit organization and its worldwide members have been determined to change this and find cures through education, research and advocacy.

LE&RN is at the forefront of raising awareness of lymphatic diseases. In 2016, the U.S. Senate established March 6 as World Lymphedema Day® in response to a bill written by LE&RN. Through educational programs and with the help of notable celebrities, LE&RN has brought visibility to these diseases and is committed to advancing advocacy and inspiring patients to feel hopeful and empowered to fight for themselves.
“I had a double mastectomy and, as a result, I have lymphedema in both arms. If we want to win this fight, we need to stand together and educate the world.”

-Kathy Bates
Academy Award-winning Actor and LE&RN Spokesperson
OVERVIEW

The lymphatic system is a circulatory system which is critical to immune function and good health. When it is compromised and lymph flow is restricted, the physical impact to patients can be devastating, life altering, and can lead to shortened lifespan. LE is one such lymphatic disease. LE is a chronic, debilitating, and incurable swelling that can be a result of cancer treatment, inherited or genetic causes, damage to the lymphatic system from surgery or an accident, or from parasites as in lymphatic filariasis.

Up to 10 million Americans and an estimated 250 million worldwide suffer from LE and related lymphatic diseases. This includes up to 30% of breast cancer survivors, children born with lymphatic diseases, veterans who have suffered physical trauma, and tens of millions living with filariasis. Currently, there are no cures and few treatments for these diseases.

Following a history of neglect when it came to LD and LE, a group of advocates emerged in the 1990s determined to change the course of history. In 1998, the Lymphatic Education & Research Network (LE&RN) would be founded as a parent’s response to medical professionals and researchers who were unable to provide answers for her daughter suffering from LD and LE. With lymphatic science and research all but non-existent, LE&RN began providing annual grants to research fellows at the world’s leading universities, putting LD on the radar in the medical community, in academia, in government, and with pharmaceutical companies.

Advocacy with the National Institutes of Health (NIH) would lead to lymphatics becoming a focus of research there, as well as the founding of the Trans-NIH Coordinating Committee for Lymphatic Research and Medicine. LE&RN was instrumental in establishing the first, and still only, endowed university Chair in Lymphatic Research and Medicine; Dr. Stanley Rockson holds the Chair at Stanford University and is a driving force as Founding Chair of LE&RN’s Scientific and Medical Advisory Council.
LE&RN is committed to making LE a global health priority and is leading this effort internationally through a host of programs:

**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 specialized research activities.

**Network**—building an extensive network of participants and partners to advance advocacy.

In 2016, LE&RN addressed three obstacles facing LE awareness:

- Recognition of LE as a disease, rather than by the euphemism of calling it a condition.
- Making it easier for lymphedema to enter the national conversation by using the acronym LE.
- Moving the conversation beyond a search for treatments to the expectation that cures be demanded.

Today, it is recognized that beyond lymphatic diseases such as LE, lipedema and filariasis, lymphatic research is impacting research on cancer metastasis, heart disease, Alzheimer's, AIDS, Rheumatoid Arthritis, Multiple Sclerosis, Diabetes, obesity and a host of other diseases. LE&RN legislative action is at the forefront of educating politicians on the need to fund lymphatic research and address the insurance needs of those with these diseases. As a circulatory system that is part of the body’s immune response, the possibilities to impact public health globally through lymphatic research are infinite.

For more information on these lymphatic diseases, visit: WorldLymphedemaDay.org.
PROGRAMS

EDUCATION
LE&RN offers educational programs targeting academia, the medical and patient communities, and the public at large.

WHY EDUCATION?
Although the World Health Organization (WHO) estimates as many as 250 million people worldwide suffer with LE, and tens of millions more suffer from the lymphatic disease lipedema, these diseases remain virtually unknown to most. Those who seek treatments too often find that there are no local certified lymphedema therapists. With medical students receiving as little as 30 minutes of lymphatic system training during their education, physicians remain ill-equipped to diagnose and treat patients.

To change this situation, LE&RN has implemented programs that inform and engage patients, therapists, researchers, and the medical community. LE&RN broadcasts a monthly Livestream Symposium Series, which brings the world’s foremost authorities in LD and LE to patients, their families, and medical professionals. These symposia are permanently available for viewing on LE&RN’s website, which also offers a world-class library of information and resources accessible to anyone.

LE&RN sponsors International Scholarship Awards for young research investigators in lymphatics.

LE&RN created the first online 7-Credit Continuing Medical Education (CME) course for Physicians: Lymphatic-Vascular Disease Diagnosis & Treatment. The course features lectures by leaders in the field and addresses our mission to educate healthcare professionals about LD.

LE&RN established the first-ever standards for Centers of Excellence in the Diagnosis & Treatment of Lymphatic Diseases to guarantee that patients are able to find the best alternatives for comprehensive institutional care.

LE&RN sponsors the LE&RN/LymphNotes U.S. Lymphedema Therapist Scholarship to provide tuition support to therapists seeking certification in LE therapy.

“After years of relative neglect, the field of lymphatic research is finally getting the attention it deserves. Clearly, the biggest impact to this field has been the establishment of LE&RN. Now more than ever, there is great opportunity for scientific advancement. Importantly, lymphatic research will enhance the lives of millions, prevent future disease, and preserve the good health we all wish to enjoy. Research defines our future.”

—Stanley G. Rockson, MD, Center for Lymphatic and Venous Disorders, Stanford University School of Medicine
LE&RN publishes monthly educational newsletters and multimedia presentations, and uses social media to provide comprehensive updates of news in the field.

LE&RN offers a web-based Ask the Experts feature, which allows patients, their families, and professionals to directly ask questions to prominent authorities in the field.

LE&RN’s Virtual Expo is a web-based resource that brings together lymphatic-related manufacturers, distributors, schools, pharmaceutical companies, and accessory businesses in one place to offer solutions and perspectives for patients and therapists alike. It allows patients and professionals 24/7 access to available treatments and services.

LE&RN is at the forefront of making lymphatic research a priority with the National Institutes of Health and with both local and national politicians.

LE&RN has established an International Biennial Conference (the Lymphatic Forum) focused on lymphatic research, which debuted in Chicago in June 2017.

Educational Impact:

• 50 LE therapist scholarships
• Over 100 young investigator conference scholarships
• More than 300 patient questions answered in Ask the Experts
• Over 100 Livestream Symposiums
• More than 400 newsletters
• Up to 20 company profiles in LE&RN’s Virtual Expo

Travel Award Scholarship winners from around the world at Lymphatic Forum 2019, sponsored by NAVBO and LE&RN; Austin, Texas, May 30–June 1.
LE&RN acknowledges the ongoing and urgent need to educate medical professionals, and as such is creating learning modules that the organization is bringing to hospitals across the country. Only when hospitals, doctors, nurses, and therapists are knowledgeable, will they be able to diagnose and provide patients with LD with the right treatment.

**FINAL POINT**
Educating and increasing awareness of LD and LE among doctors, patients, and the public at large will lead to the commitment of resources and research necessary for discovering effective treatments and ultimately, cures.

**PROGRAMS**

**RESEARCH**
LE&RN seeks to accelerate prevention, treatments, and cures through research.

**WHY RESEARCH?**
The cures and treatments of tomorrow are tied to the research we do today. However, lymphatic diseases such as lymphedema and lipedema are under-researched, which leads to few treatments. As a result, medical schools do not properly educate future doctors about lymphatic diseases, which leads to many patients with these diseases going undiagnosed and suffering from a lifetime without proper treatments or hope for a cure. This is propagated by medical researchers who continue to focus their efforts elsewhere as long as there is limited research funding prioritized on lymphatic diseases.

One of LE&RN's first research initiatives and successes was the establishment of the biennial Gordon Research Conference Series devoted to Molecular Mechanisms in Lymphatic Function and Disease. In 2016, LE&RN’s Spokesperson, Academy Award-winning actress Kathy Bates, was a featured speaker on the first patient panel in the history of the GRC.

LE&RN has awarded dozens of postdoctoral research fellowships to the world's leading universities for over a decade, putting lymphatic diseases on the radar in the medical community, in academia, in government, and with pharmaceutical companies.

"Thank you for your dedicated efforts, as well as those of Kathy Bates on behalf of LE&RN, to promote our common objective for advancing research on the lymphatic system and related diseases. **We look forward to continuing future work with LE&RN as a strong research partner.**"

–Francis S. Collins, M.D., Ph.D.
Director of the National Institutes of Health
Dr. Stanley Rockson was instrumental in establishing the International Lymphatic Disease and Lymphedema Registry & Biorepository at Stanford University as a resource to stimulate research and support clinical trials and drug development. He also launched LE&RN’s journal, *Lymphatic Research and Biology*, the only peer-reviewed journal in the field, providing the forum for the exchange of cutting-edge scientific developments in lymphatic science and medicine.

In 2019, LE&RN established a lymphatic researcher database that is accessible and searchable by all 27 NIH Institutes and Centers.

LE&RN secured ongoing Congressional and National Institutes of Health (NIH) support for lymphatic research.

Research Impact:
- 22 research fellowships
- 44 young investigator research poster awards
- Over 100 researcher travel scholarships to conferences
- Over 600 research articles published in LE&RN’s peer-reviewed journal
- 2000 biorepository samples
- 1000 Patient Registry members

**WHERE WE ARE GOING/WHO WILL BENEFIT**

As interest in lymphatic research is expanding, government funding is being curtailed due to budget cuts. LE&RN has the ambition to expand its fellowship program to ensure that it is cultivating a new generation of researchers that will revolutionize medicine in the future. These breakthroughs will ultimately lead to the prevention, treatments, and cures that patients worldwide need and deserve.

*Dr. Anthony Fauci, NIAID Director (National Institutes of Health), Kathy Bates and LE&RN CEO William Repicci at the 2017 Research!America Awards where Dr. Fauci and Ms. Bates were both honored, Washington, DC.*
FINAL POINT
Lymphatic diseases such as LE and LI are life-sentences. Only research can provide hope that quality of life can be restored. The possibilities to impact public health globally through lymphatic research are vast.

PROGRAMS

ADVOCACY AND NETWORKING
Through several outreach programs and through legislative advocacy efforts, LE&RN is building an extensive network of participants and partners to advance advocacy.

WHY ADVOCACY AND NETWORKING?
Up to 10 million Americans, and hundreds of millions worldwide have LE. Millions more suffer from LI. Yet, most people have never heard of these lymphatic diseases, including those who suffer from them. Doctors often don’t recognize or properly diagnose these diseases and patients often don’t know that they are at risk or know the symptoms of their disease. Currently, insurance coverage is too limited.

Successes in bringing LD to the forefront of global attention are tied to two events: unrelenting advocacy on the part of the patients and their networks; and the availability of treatments that give the medical community a reason to invest themselves in the disease.

To this end, LE&RN has established U.S. State Chapters and International Chapters. Each Chapter is the gateway to the lymphatic community and serves as a platform where patients can connect with one another through support groups, with therapists, and at treatment centers.

LE&RN has engaged celebrities to help lead and bring visibility to this fight. Academy Award-winning actor Kathy Bates became LE&RN’s National Spokesperson in 2015. Ms. Bates has LE.
in both arms as a result of a double mastectomy. LE affects up to 30% of breast cancer survivors. Since she joined LE&RN, Ms. Bates has lectured at conferences including at the National Institutes of Health, American College of Phlebology, and American Cancer Society. She has also told the story of LD in countless print publications.

LE&RN’s outreach expands to national audiences and it has aired stories on “CBS Sunday Morning,” “Dr Oz,” “USA Today,” “NIH Medline Plus,” “People Magazine,” WebMD,” “The Doctors,” “The Wendy Williams Show,” and a host of other media outlets.

As a result, LE&RN has established The Face of Lymphedema Campaign that has brought thousands of new advocates to this cause. It has also resulted in LE&RN establishing an Honorary Board that now includes Dame Judi Dench, actors Patricia Clarkson and Steve Guttenberg, Bravo Network’s Andy Cohen, and Olympic Gold Figure Skating Champion Sarah Hughes.

Events such as LE&RN’s annual Walk to Fight Lymphedema & Lymphatic Diseases, which occur across the U.S., connect patients and their families with supporters and raise awareness with the public at large.

With the support of the U.S. Senate, LE&RN established an annual World Lymphedema Day™ on March 6 to be a time when events, lectures, and legislative proclamations put these diseases center stage.

Network and Advocacy Impact:
- National Walks to include New York; Colorado; Washington, DC; Massachusetts; and California
- Over 100 patient stories told
- 20,000 Facebook fans
- 34 corporate sponsors
- Over 20 State and International chapters

WHERE WE ARE GOING/ WHO WILL BENEFIT
LE&RN has petitioned the World Health Organization to make LE the focus of World Health Day.

In the closing weeks of 2019, Congress passed a final appropriations bill for fiscal-year (FY) 2020, marking a major milestone in LE&RN’s goal of establishing a National Commission on Lymphatic Diseases NIH. The bill encourages the Director of NIH “To establish a National Commission on Lymphatic Diseases to explore and make recommendations on the ongoing expansion and coordination of lymphatic disease research NIH-wide.” This victory is the culmination of an intense year of lobbying by LE&RN and many committed activists, particularly the tireless work of LE&RN National Spokesperson Kathy Bates.

“My mom has lymphedema and so, like all affected families, I feel that I have lymphedema. And that’s why I support LE&RN in creating the awareness that will lead to a cure.”

–Steve Guttenberg, Actor
FINAL POINT
To succeed at making LD a global priority it will require a sustained commitment by those in the political arena, the medical profession, as well as the community ranging from patients and their families to advocates and philanthropic supporters. Together they can increase awareness, inform and influence policies and practices, and fund research to discover new treatments and cures.

RECENT TIMELINE
In 2013, LE&RN embarked on a monumental plan. Capitalizing on the success of the first fifteen years, the organization rebranded itself to reflect an expanded mission. Its new name made it clear that education and advocacy became equal partners with its research agenda. From a field led by a few lone voices, a movement started that would change the world.

2020 & BEYOND
Moving forward, the focus is on building worldwide networks that foster cure-focused research, physician education, and Centers of Excellence so that no case of lymphatic disease goes untreated.

EDUCATION
For patient care to evolve, physicians need to become educated about these diseases and their treatments. LE&RN will need dedicated staff to continue to promote its Physician CME in Lymphatic-Vascular Disease Diagnosis & Treatment, with collaboration with AVLS (American Society & Lymphatic Society), ASBrS (American Society of Breast Surgeons.), American Venous Forum, and numerous other aligned physician groups.

LE&RN’s Centers of Excellence in the Diagnosis and Treatment of Lymphatic Diseases is a transformational program, which is establishing care centers around the world to meet the need of those living with lymphatic diseases. Staff is needed to coordinate this program and the patient chapters associated with each Center.

“For years I have suffered alone, in silence, with this disease. I feel embarrassed and deeply ashamed of what lymphedema has done to my body. I am beginning to accept, with the help of those closest to me, that I have nothing to be ashamed of. I feel so moved that a community of people with lymphedema has surfaced and is demanding to be heard—putting voices and faces to a disease people know so little about. I know that part of me will always maintain the hope for answers. Awareness is a very critical step in that journey forward.”

– Dara M., Patient, Buffalo, NY
RESEARCH
Cures come from research. Research requires funding. LE&RN is committed to awarding fellowships to post-doctoral researchers from the world's leading lymphatic laboratories. In addition, to bring the best and brightest young researchers into the field, LE&RN seeks to expand its scholarship program.

Researchers need tools to find cures. A new Director position is needed to oversee the International Patient Registry & Biorepository Coordinator. This is necessary to grow its number of bio samples, get patients registered, and link to labs internationally.

ADVOCACY & NETWORKING
LE&RN has established World Lymphedema Day®, which is now a global phenomenon. A Project Director with a dedicated budget is needed to promote and coordinate this event worldwide.

Connecting the world means expanding LE&RN's Chapters globally. It also means establishing State and Regional Chapters in the U.S. Capitalizing on this momentum requires a dedicated budget and staff-associate.

LE&RN's Lobby Days now bring hundreds of advocates from around the world to Washington, DC where they meet with their legislators and where LE&RN's agenda is presented to the National Institutes of Health (NIH), Centers for Disease Control and Prevention (CDC), Veterans Administration (VA), and others. It is critical that we have funding to maintain and grow this activity and the expense of a DC Government Relations Firm.

2020–2022
EDUCATION
- Hire Director of Centers of Excellence Program
- Expand and promote LE&RN's CME in Lymphatic-Vascular Disease Diagnosis & Treatment
- Produce 20 Scientific Livestream Symposiums
- Publish 24 newsletters and dozens of breaking news e-blasts

RESEARCH
- Award 10 two-year post-doctoral research fellowships
- Acquire up to 200 new biorepository samples and eexpand the Patient Registry by 200 patients
- Expand research staff to 2 full-time positions
- Award 20 scholarships and eight poster awards for young investigators attending key lymphatic conferences

Dr. James Madara, CEO of the American Medical Association (AMA), and LE&RN Spokesperson Kathy Bates at 2016 LE&RN/AMA Roundtable, Chicago, IL.
ADVOCACY & NETWORKING

• Award up to 40 institutions the LE&RN Centers of Excellence in LD designation and establish patient oversight chapters at each institution.
• Expand Washington, DC Lobby Days and Rally to include thousands of advocates who meet with our legislators
• Roll out PSA Ad Campaign with Kathy Bates to expand awareness
• Establish a 24-hour patient call line with dedicated staff
• Expand International Chapters with full-time staff support
• Hire a Director of Development and Outreach Associate

LE&RN Youth Ambassador Emma Detlefsen introduced at 2017 Research!America Awards, Washington, DC.
THE OPPORTUNITY

Listed here are LE&RN’s fundraising priorities for the next two years, which will enable LE&RN to realize its strategic growth plan in the U.S. and internationally.

<table>
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<th>Two-Year Statement of Need</th>
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<tr>
<td>Ongoing Program Support</td>
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<td>Education</td>
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<td>Update CME Physician Certification Program</td>
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<td>PSA Ad Campaign With Kathy Bates</td>
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"Events like the Lymphatic Forum are essential. They provide the opportunity to hear and learn about a wide variety of different aspects of the lymphatic field. This enables us researchers to share ideas, to have fruitful discussions and to learn to look at our own project from different perspectives. During this meeting you can feel that you are part of a well-connected and supporting community, unified in trying to elucidate one by one, the missing pieces of knowledge of the lymphatic world."

— Samia Bachmann, Institute of Pharmaceutical Sciences, Zurich
WHO IS INVOLVED

For a full listing of LE&RN’s Leadership Team, please visit: lymphaticnetwork.org/about/leadership-team/

Stanley G. Rockson, M.D., FACP, FACC
Stanford University
Founding Chair of Scientific & Medical Advisory Council

Kari Alitalo, M.D., Ph.D.
Finnish Academy of Sciences
Scientific & Medical Advisory Council

Peter S. Mortimer, M.D
St. George’s Hospital, London
Scientific & Medical Advisory Council

Mihaela Skobe, Ph. D.
Mount Sinai Hospital
Scientific & Medical Advisory Council

Joseph Dayan, M.D.
Memorial Sloan Kettering
Board of Directors

Sheila H. Ridner, Ph.D., R.N.
Vanderbilt University
Board of Directors

William Repicci
President & CEO

Kathy Bates
National Spokesperson and Chair of the Honorary Board

Dame Judi Dench
Honorary Board

Steve Guttenberg
Honorary Board

Dave McDowell
AOL/Yahoo
Leadership Council

Nancy Gray, Ph.D.
Gordon Research Conferences
Leadership Council