Celebrating 30 Years of Advocacy for the Rare Disease Community

NORD
The Yang family helped organize a Rare Disease Day event at the University of Alabama at Birmingham and joined NORD’s media campaign to promote awareness.
Dear NORD Members and Friends:

As we celebrated the 30th anniversary of NORD and the Orphan Drug Act, we honored NORD’s illustrious history and the pioneering individuals who established NORD and focused a national spotlight on the need for orphan drug legislation. Their dedicated and passionate advocacy has improved the lives of millions of Americans.

Our anniversary year was also a time to reassess the community’s needs and challenges, and to renew our commitment to work toward a world in which every life is lived to the fullest and every patient has a treatment or cure.

The year began – poignantly – with accolades to actor Jack Klugman, who played such a pivotal role in the Orphan Drug Act and the history of NORD. Jack died at his home in California on December 24, 2012, and NORD issued a statement honoring his life that was widely quoted in the press.

Friends of NORD from its earliest days remember Jack as NORD’s “Honorary Chairman” who had highlighted the need for the Orphan Drug Act in two episodes of his popular television show, Quincy, M.E.

Jack’s death came just one week before a major event hosted by the Food and Drug Administration (FDA) commemorating the 30th anniversary of the Orphan Drug Act. A central theme of that event was the crucial role played by the ad hoc coalition that became NORD in enactment of the Orphan Drug Act.

Today, NORD continues to provide leadership to the rare disease community. As you read the following brief summary of NORD’s 30th anniversary year, please be aware that these accomplishments were only possible because of the dedicated members and advocates who generously donated their time, expertise, and guidance to NORD. We thank, in particular, our board of directors, medical advisors, member organizations and Corporate Council for their ongoing and important support.

At this defining moment, when the science of rare diseases is advancing at an unparalleled pace, NORD remains dedicated – as it has been since 1983 – to improving the lives of the patients and families who so desperately need our help.

Sincerely,

Peter L. Saltonstall, President and CEO

E. Michael D. Scott, 2013 Board Chair
Highlights of our 30th Anniversary Year

Celebrating 30 Years of Progress

NORD’s 30th anniversary began on a bittersweet note, with the death of actor Jack Klugman, an early champion of the Orphan Drug Act and NORD. Through his popular TV show, Quincy M.E., Klugman focused national attention on the need for treatments for people with rare diseases. He took a personal interest in the establishment of NORD and continued to follow NORD’s progress through its early years.

On January 4th, the 30th anniversary of the signing of the Orphan Drug Act by President Ronald Reagan, the FDA hosted an event featuring a moving video interview with NORD founder Abbey S. Meyers and remarks by current NORD President and CEO Peter L. Saltonstall. We hosted our own celebration in May with children and young adults presenting medals to heroes of the rare disease movement.

Extending Our Voice Globally

NORD and the Japan Patients Association (JPA) formed a partnership in 2013 aimed at connecting rare disease patients in the two countries. The signing ceremony took place in Washington, DC, with CEO Peter L. Saltonstall representing NORD and Tateo Ito, CEO, representing JPA. Both emphasized the need for global partnership.

In signing the document, President Ito said the two organizations “would like to make use of our valuable knowledge and experiences for the mutual benefit of patients in both countries.” Saltonstall added, “Global collaboration will drive progress toward new treatments, improved public policies and better lives for people with rare diseases.”

Mentoring our Members for Growth and Impact

Membership continued to grow in 2013, with 13 new organizations joining the NORD community:

- Association for Creatine Deficiencies
- CADASIL Association
- Cutaneous Lymphoma Foundation
- Dup15 Alliance
- Genetic and Rare Disorders Organisation
- Global Foundation for Peroxisomal Disorders
- Mesothelioma Applied Research Foundation
- PCD/KS Foundation
- PRISMS - Parents and Researchers Interested in Smith-Magenis Syndrome
- Pulmonary Fibrosis Foundation
- Red Sanfillippo AC
- Rothmund-Thomson Syndrome Foundation
- SSADH Association

Helping our members grow and accomplish their goals is at the heart of our mission. In 2013, we significantly expanded member services to include a broad range of webinars, teleconferences and in-person regional meetings, all aimed at giving members an inside track on current issues and opportunities.

We also provided scholarships for 70 representatives from member organizations to attend NORD’s Annual Breakthrough Summit on Rare Diseases and Orphan Products, a great opportunity to interact and network with researchers, government officials, and industry leaders. To help prepare these patient leaders for the Summit, we invited them to submit brief essays about living with a rare disease. NORD consolidated the riveting pieces into an op-ed on behalf of the community, which was published in The Wall Street Journal.
Bringing Rare Disease Day to Communities Across America

NORD organized a Handprints Across America campaign for Rare Disease Day to encourage grassroots participation and support. Hundreds of people from communities across the U.S. submitted brief messages and photos of themselves holding the Rare Disease Day logo.

Advancing Diagnosis

NORD published three physician guides in 2013, on Gaucher disease, Pompe disease, and infantile spasms. Written by medical experts, these guides are free and available online at www.nordphysicianguides.org. The purpose is to promote early, accurate diagnosis and appropriate treatment for patients.
Training Tomorrow’s Advocates

“Having a child with a rare disease makes you feel helpless… powerless. At least, that’s how I felt until I found NORD,” says Kelli Foster. Kelli and her daughter, Brooke, both live with a rare disease known as mastocytosis.

Kelli attended a NORD conference a few years ago and found that she was greatly empowered by that experience. Connecting with senior officials from NIH and FDA, along with rare disease medical experts and pharmaceutical industry leaders, showed Kelli that she and her family were not alone but part of a community.

That experience gave Kelli the courage to become a more active and vocal advocate. And ultimately, she was able to help her own daughter and others within her patient organization access treatment that had previously not been available to them.

Brooke also benefited from her mother’s awakening to advocacy. At a 2013 Rare Disease Day event in New Jersey to educate members of that state’s legislature about rare disease challenges, Brooke gave a presentation that was one of the highlights of the event.

“The NORD conference taught me that every voice is important,” says Kelli. And Brooke echoed that sentiment in a NORD video to encourage young advocates. “If you want to get involved,” she said, “just DO it!”

If you want to get involved, just DO it!
SINCE 1983 NORD HAS PROVIDED LEADERSHIP AND PATIENT REPRESENTATION IN PUBLIC POLICY.
Providing Lifesaving Patient Services

When Jon and Amanda Miller heard that NORD was hosting a Rare Disease Day event near their home in New Jersey, they decided to come and say thank you. That’s because a NORD patient assistance program helped the Millers at a time of crisis in their lives.

Their son, Evan, was just three months old when he went into a medical crisis and was diagnosed with tyrosinemia type 1, a rare metabolic disease. The good news for the Millers was that there is an FDA-approved treatment for that disease.

The bad news was that the family did not have insurance that would cover the cost of the drug, and the hospital wouldn’t release Evan without the medication that was his lifeline.

“It was very stressful,” Amanda says. “It was bad enough to have to worry about Evan’s medical condition. But to also have to worry about how we were going to get his medication was almost overwhelming.”

Then a friend told Jon about NORD and its patient assistance programs. Jon called NORD and – even though it was the day before Thanksgiving – within one day, NORD had arranged for the medication to be delivered to the hospital, and little Evan and his parents were soon on their way home.

Today, Evan is doing well and Jon and Amanda are grateful that a safety net was in place to help them access the lifesaving therapy their baby needed.

“NORD’s Patient Assistance Program provided Evan with the lifesaving medication he needed.”
NORD pioneered patient assistance programs for medications & continues to provide leadership in that field.
Stretching Dollars to Save Lives

When 14-year-old Jacob Maren wanted to do something to help others as a bar mitzvah and school leadership project, he and his parents visited the NORD offices near his home in Connecticut. Jacob was particularly interested in educating physicians about a rare, potentially very serious condition known as infantile spasms that had touched his life briefly when he was very young.

Ultimately, Jacob raised $10,000 to make it possible for NORD to create and publish a Physician Guide to Infantile Spasms. The guide is available free to physicians around the world at www.nordphysicianguides.org.

NORD created the physician guide website a few years ago as part of its campaign to reduce the time to diagnosis and treatment for patients. The guides are written by medical experts, and each time a new guide is published, NORD conducts outreach to physicians in targeted practice fields to promote awareness of the new resource.

Since the American Academy of Neurology (AAN) had published new treatment guidelines for infantile spasms around the time Jacob first approached NORD, the educational team at NORD worked with AAN and arranged for two physicians who had been involved in the new guidelines to write the NORD guide.

Through the years, NORD has demonstrated its ability to gain maximum benefit for patients and their families from donations of all sizes.

“Jacob raised funds for NORD to create a physician guide on a rare disease affecting infants where early diagnosis is critical.”
CLINICIANS FROM MORE THAN 160 COUNTRIES HAVE ACCESSED NORD’S NEW ONLINE PHYSICIAN GUIDES TO PROMOTE DIAGNOSIS AND TREATMENT FOR PATIENTS
2013 Financial Position

Revenue

Program Grants. ................................................................. $13,090,825
Patient Assistance and Research Program Fees. ...................... $1,418,853
Special Events Revenue ...................................................... $776,744
Membership Dues ............................................................... $751,040
Contributions and Bequests ................................................ $617,280
Royalties and Other ........................................................... $146,154
Investment Income ............................................................ $18,998

Total Revenue ........................................................................ $16,819,894
Patient Services .................................. $12,459,521
Advocacy ............................................. $385,505
Research and Medical Scientific Affairs. ............................................. $620,136
Membership and Education. ............................................. $708,174
General and Administrative ............................................. $2,117,125
Development and Communications ............................................. $1,308,380

Total Expenses ............................................. $17,598,841
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For more than 30 years, NORD has been helping families like the Millers meet the challenge of living with a rare disease.
Alone we are rare. Together we are strong.™