NAVIGATING THE PATH

AMERICAN LIVER FOUNDATION | 2019 ANNUAL REPORT
When my son was diagnosed with liver disease, we were lost. A nurse told us to call the American Liver Foundation. It’s been our compass ever since.

- Chelsea W.

Founded in 1976, the American Liver Foundation (ALF) is the nation’s largest patient advocacy organization for people impacted by liver disease. ALF reaches more than three million people each year with health information, education and support services via its national team, field offices, and an active online presence. Recognized as a trusted voice for those battling liver disease, ALF also operates a national toll-free HelpLine, educates patients, policymakers and the public, and provides grants to early-career researchers. For more information about ALF, please visit www.liverfoundation.org.
Number of elected officials ALF met with during our annual Awareness Day on Capitol Hill: **89**

Donated to cutting-edge liver research: **$275,000**

Steps taken at our Liver Life Walks: **2.5 BILLION**

Inquiries received by our HelpLine Specialists: **10,000**

2019

**BY THE NUMBERS**

ALF advocated for the Living Donor Protection Act (H.R. 1124/ S. 511). Key elements of this legislation were advanced by an Executive Order to enhance organ donation.

People served by our education and support services: **2,000,000**

Raised at our walks, challenges and culinary events to fight liver disease: **$5 MILLION**

We also advocated for the LIVER Act (H.R. 3016/S. 3074) which has gained cosponsors and raised awareness of the challenges and opportunities facing those impacted by liver diseases and associated conditions.

People connected on our online communities: **30,000**

Medical professionals viewed our educational webinars: **10,000**

Individuals visited the ALF website: **3 MILLION**

Community education Programs were held across the country: **367**
Dear Friends,

Welcome to the American Liver Foundation (ALF) 2019 Annual Report.

As we look back over the year, we’re proud to be part of an organization that is singularly focused on improving the lives of people with liver disease. If you or a family member has a serious liver condition, you know the patient journey is full of twists and turns. Nobody has to travel that road alone. ALF is there at every step.

NAVIGATING THE PATH

In 2019, our National Helpline responded to nearly 10,000 inquiries. We helped individuals with non-alcoholic steatohepatitis plan a more healthful diet. We connected those with hepatitis C and other conditions to medication assistance. We guided many on the transplant waiting list through the process of registering at two or more transplant hospitals. And we were there to listen when an exhausted mother called us because her only daughter was on at-home hospice care for end-stage liver disease. At the end of the emotional conversation, the mother told our Helpline Specialist: “You and the work you do are as important as the doctors and the nurses.”

A COMMUNITY TO LEAN ON

Approximately 30,000 people have joined one of the American Liver Foundation’s online communities. There, they share their stories and connect with others who have had similar experiences. They ask questions and find answers, gain new perspectives, and support one another.

The members of ALF’s National Patient Advocacy Committee share personal healthcare experiences to benefit a much broader audience. Our passionate advocates offer hope and support to others going through the challenges of liver disease.

MAKING A DIFFERENCE IN PATIENTS’ LIVES

ALF’s website welcomed 3 million visitors in 2019, including Henry, who sent us the following note:

“I got my lab tests back today. My heart was pounding. I was scared. I had no idea what they meant. Then I found a video on your website called ‘Understanding Your Liver Enzyme Test Results.’ It was exactly the information that I needed. Thank you and God bless.”

We are committed to making a difference for everyone battling liver disease and everyone at risk. Across the country, 367 community educational programs were held, often in high-risk, underserved communities. Over 300,000 people searched our comprehensive video library and viewed our informational videos and webinars.

To help empower the medical community to provide the best care for patients, professional education programs were conducted throughout the country. Additionally, almost 10,000 medical professionals viewed the webinars in our growing library of professional resources.

YOU CAN COUNT ON THE ALF

As our country and the world continue to navigate the challenges associated with COVID-19, ALF remains committed to supporting the liver community. We have created numerous resources to help guide liver patients through the pandemic.

We also wish to acknowledge the tremendous dedication and commitment of medical professionals across the country who have been on the frontlines for liver patients during this public health crisis. You are our heroes.

OUR SINCEREST THANKS

We are particularly grateful to Nicholas DeRoma, who is completing his term as National Board Chair and Thomas Nealon III, who has served as National Board Chair and then President and CEO. Thankfully, both will remain involved as Board members. You can read about them in this report.

We also admire all of you in our liver community. You have inspired us with your determination and perseverance – especially during these difficult times - and with the countless hours you’ve expended in support of ALF. Together, we will achieve our vision of a world without liver disease.

With great appreciation,

Daniel E. Weil
National Board Chair

Lynn Gardiner Seim, MSN, RN
Executive Vice President and Chief Operating Officer

David Ticker
Executive Vice President and Chief Financial Officer
On his 50th birthday, Tom Nealon decided to run a marathon. Tom always dreams big. Soon he ran the New York City and Chicago marathons, and the annual Marine Corps Marathon in Washington D.C., as well as the Miami and Disney marathons. But he always hoped to run the Boston Marathon.

So, in 2002, he took a philanthropic route to the starting line. Agreeing to raise funds to help ALF end liver disease, Tom joined our Boston Marathon Team.

That is when he met one-year-old Zachary Rue, who was born with biliary atresia, a rare liver disease. ALF matches runners with patients. “From the moment we spoke, it was like we knew each other,” Zac’s mom Heather recalls of meeting Nealon. The feeling was mutual.

Zac’s strength and story motivated Tom to do even more for ALF. To this day, Tom holds the record as the person who has raised the most funds for ALF by competing in marathons: over $1,500,000.

As a lawyer and compassionate leader, Tom has helped build and shape the ALF. He has spent the last eight years in senior leadership roles, including national board chair and then President and CEO.

Under Tom’s leadership, ALF broadened its education programs for medical professionals and patients, expanded its federal advocacy efforts, secured new organizational and industry partnerships, provided critical support to young researchers, shared the patient voice at national conferences and the FDA, and provided life-saving information and services to an unprecedented number of individuals with liver disease — as well as their families, friends and those at risk.

Tom recently stepped down from the position of CEO and President, however, he remains an active Board Member and mentor to 18-year-old Zac. Thank you, Tom

Tom Nealon is motivated by a desire to do good in the world. This, combined with his integrity, tenacity and intelligence, is a powerful force. ALF and those with liver disease are the beneficiaries.

David Ticker
ALF EVP & Chief Financial Officer

Tom stepped into the CEO role during a challenging time at the Foundation. With his practical application of logic and his well-mannered temperament, he guided the organization to stability and expanded the impact of ALF both with our patients and in the medical community.

Michael Kerr
ALF Board of Directors
“Nick has been an anchor for the ALF for so many years. His leadership experience coupled with his kind and caring nature has kept the ALF grounded in even the roughest of waters.”

Lynn Seim
ALF EVP & Chief Operating Officer

All of us at ALF owe Nick a tremendous debt of gratitude. His dedication, compassion, mentorship and extensive attention to detail have immeasurably benefitted the ALF and liver patients everywhere.”

Daniel Weil, Chair
ALF Board of Directors

Nick DeRoma has great leadership talents. With corporate experience in executive positions, he brings both practical application of proven solutions and wisdom to ALF. As a student of knowledge, his plain-spoken manner and anecdotes help make his points. He sees problems clearly and can fashion solutions which enable him to motivate others to action and to focus on the goal.

Michael Kerr
ALF Board of Directors

To describe Nick in one sentence is an insurmountable task - so instead - I thank you for all you have done for liver patients, their families, caregivers and me. Over the years, you have mentored me, guided me and encouraged me to become the best version of myself.

Alicia Eppich
Director, CT Division

We thank Nick DeRoma for serving as Board Chair from 2018 – 2020 and continuing to remain a vital part of our Board of Directors.

Nick is an attorney and retired from the practice of law. During his extensive international legal career as in-house counsel, he served as Executive Vice President and Chief Legal Officer, Alcoa, Inc.; Senior Vice President and Chief Legal Officer, Nortel Networks Corp.; Vice President & General Counsel, IBM Europe, Middle East, Africa; General Counsel, IBM North America; General Counsel, IBM Asia; and Managing Attorney, IBM Corp.

He is a graduate of William & Mary Law School and the University of Connecticut, and has been awarded an Executive MBA from the Harvard Business School.

Nick has been a powerful force for assisting people with liver disease. Always humble, Nick seldom praises himself. So today, we want to do it ourselves.
Every year, the American Liver Foundation supports early-career researchers whose efforts bring us closer to finding cures for liver disease.

This year, we awarded five incredibly talented early-career scientists who are raising the bar on cutting-edge liver research.

**2019 POSTDOCTORAL RESEARCH FELLOWSHIP AWARDS:**

**BILIARY ATRESIA FUND FOR THE CURE AWARD**
Jessica Llewellyn, PhD - University of Pennsylvania
Project: Identification and characterization of fibrogenic cell of the extrahepatic bile duct
Mentor: Rebecca G. Wells, MD

**HANS POPPER MEMORIAL AWARD**
Omar Haque, MD - Massachusetts General Hospital, Harvard Medical School
Project: Ex-Vivo Liver Graft Regeneration via Normothermic Machine Perfusion in a Two-Thirds Partial Hepatectomy Model
Mentors: Korkut Uygun, PhD & James F. Markmann, MD, PhD

**ALEXANDER M. WHITE, III MEMORIAL AWARD**
Debanjali Dasgupta, PhD - Mayo Clinic, Rochester
Project: Hepatic IRE1alpha activation induces EV release, leading to hepatic injury
Mentor: Harmeet Malhi, MBBS

**IRWIN M. ARIAS, MD AWARD**
Taylor M. Coe, MD - Massachusetts General Hospital
Project: Role of IdeS in Depletion of IgG to Reduce Antibody Mediated Graft Damage in Non-Human Primate Liver Xenotransplantation
Mentor: James F. Markmann, MD, PhD

In 2019, our Liver Scholar Award went to Hien Dang, PhD, who is making a difference in the battle against liver cancer - one of the fastest-rising cancers in the U.S.

Dr. Dang is part of a research team that discovered a mechanism that affects the DNA of more than one-third of liver cancer patients. With her ALF grant, she is working to develop a biomarker that will help identify tumors driven by this mechanism. Ultimately, her goal is to create more precise therapies for liver cancer and save lives.

When you donate to ALF, your gift supports our research program and helps ensure that bright minds remain in the field of liver research.

By doing your part, you help scientists do theirs.
I am one of over 40,000 people diagnosed with liver cancer each year.

Cancer makes you take stock of your life and what matters most to you. I always tell people: “There will be challenges and bad days. But your inner strength will surprise you.”

When I learned I needed a transplant, I thought, “Okay, I can get through this”. But the transplant team didn’t agree. It wasn’t because of my general health. It was because I’m single with no kids and my relatives live too far away.

The team said I wouldn’t have enough support post-transplant. I was scared. But I wasn’t ready to give up. Neither was the American Liver Foundation. Reaching out to them was one of the smartest things I ever did. ALF staff helped to identify hospitals that might be open to my situation. I traveled by car, train and plane to meet with those medical centers. After every appointment, ALF staff talked to me. And they lifted my spirits every time I was told: “You aren’t a viable transplant candidate.”

Five hospitals and 4,000 miles later, I found my transplant team!

When my surgery was over, the ALF staff were the first people I called. Thanks to the American Liver Foundation, I never had to carry the burden of liver cancer alone.

The American Liver Foundation was my advocate. They are a voice for all of us - every adult and child - living with liver disease. We need that voice now more than ever because liver disease is on the rise. Liver cancer is the fastest-increasing cause of cancer death in the United States. And - because so many of us eat too much and don’t exercise enough - as many as 100 million Americans are at risk for unhealthy fat build-up in their livers.

I like that ALF is there for people who are living with liver disease now and supports research and policies that will one day end liver disease. Please join me in supporting ALF so they can continue to provide a brighter future for all those impacted by liver disease.
Our efforts were made possible because of the generosity of the individuals, foundations and corporations listed here.* For the millions of people served by ALF, we thank you.

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“Going through the testing and diagnosis process can be just as frightening as getting used to a new diagnosis, but, we are not alone!”

- Julie M. Autoimmune Hepatitis
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*If we inadvertently omitted a contributor, we sincerely apologize and would like to know. Kindly call us at (646) 737-9405.

Thank You
Gifts to the American Liver Foundation help improve life for people with liver disease through advocacy, education, support services and research. We are committed to being the best possible stewards of donated funds, honoring donor intent and practicing financial transparency.

WHERE YOUR MONEY GOES

PROGRAM SERVICES
- PUBLIC EDUCATION: $1,790,670 (26.20%)
- RESEARCH: $377,122 (5.50%)
- PROFESSIONAL EDUCATION: $1,507,357 (22.10%)
- PATIENT SERVICES: $1,677,847 (24.50%)
- COMMUNITY SERVICES: $1,480,641 (21.70%)

TOTAL PROGRAM SERVICES: $6,833,637 (100.00%)

SUPPORT & PROGRAM EXPENSES
- PROGRAM SERVICES: $6,833,637 (78.2%)
- MANAGEMENT: $823,259 (9.4%)
- FUNDRAISING: $1,080,670 (12.4%)

TOTAL EXPENSES: $8,737,566 (100.00%)
## BALANCE SHEET

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<th>Assets</th>
<th>2019</th>
<th>2018</th>
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<td>Cash &amp; Cash Equivalents</td>
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<td>Investments</td>
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<td>Contributions Receivable</td>
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<td>Prepaid Expenses &amp; Other Assets</td>
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<tr>
<td>Property &amp; Equipment, Net</td>
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<tr>
<td><strong>Total Assets</strong></td>
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<td><strong>$5,260,328</strong></td>
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<table>
<thead>
<tr>
<th>Liabilities &amp; Net Assets</th>
<th>2019</th>
<th>2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liabilities:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Accrued Expenses</td>
<td>$264,817</td>
<td>$364,152</td>
</tr>
<tr>
<td>Payroll &amp; Related Liabilities</td>
<td>$308,085</td>
<td>$304,462</td>
</tr>
<tr>
<td>Funds Held on Behalf of Others</td>
<td>$774,204</td>
<td>$793,980</td>
</tr>
<tr>
<td>Awards and Grants Payable</td>
<td>$137,500</td>
<td>$237,500</td>
</tr>
<tr>
<td>Deferred Rent Payable</td>
<td>$81,638</td>
<td>$133,507</td>
</tr>
<tr>
<td>Loan Payable</td>
<td>---</td>
<td>$99,509</td>
</tr>
<tr>
<td><strong>Total Liabilities</strong></td>
<td><strong>$1,566,244</strong></td>
<td><strong>$1,933,110</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Net Assets Surplus (Deficit)</th>
<th>2019</th>
<th>2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Without Donor Restriction</td>
<td>$614,131</td>
<td>$(474,617)</td>
</tr>
<tr>
<td>With Donor Restriction</td>
<td>$3,086,822</td>
<td>$3,801,835</td>
</tr>
<tr>
<td><strong>Total Net Assets</strong></td>
<td><strong>$3,700,953</strong></td>
<td><strong>$3,327,218</strong></td>
</tr>
</tbody>
</table>

| Total Liabilities & Net Assets | $5,267,197 | $5,260,328 |

## STATEMENT OF ACTIVITIES

### Public Support & Revenues

<table>
<thead>
<tr>
<th>Description</th>
<th>2019</th>
<th>2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Contributions</td>
<td>$4,141,294</td>
<td>$4,317,775</td>
</tr>
<tr>
<td>Contributed Services</td>
<td>$525,462</td>
<td>$268,000</td>
</tr>
<tr>
<td>Investment Income Net</td>
<td>$349,305</td>
<td>$(121,217)</td>
</tr>
<tr>
<td>Other</td>
<td>$35,890</td>
<td>$25,653</td>
</tr>
<tr>
<td>Special Events (Net of Direct Expenses)</td>
<td>$4,570,320</td>
<td>$4,739,542</td>
</tr>
<tr>
<td><strong>Total Public Support &amp; Revenues</strong></td>
<td><strong>$9,622,271</strong></td>
<td><strong>$9,229,753</strong></td>
</tr>
</tbody>
</table>

### Program Services

<table>
<thead>
<tr>
<th>Description</th>
<th>2019</th>
<th>2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Public Education</td>
<td>$1,790,670</td>
<td>$1,859,566</td>
</tr>
<tr>
<td>Research</td>
<td>$377,122</td>
<td>$587,528</td>
</tr>
<tr>
<td>Professional Education</td>
<td>$1,507,357</td>
<td>$1,374,807</td>
</tr>
<tr>
<td>Patient Services</td>
<td>$1,677,847</td>
<td>$1,725,794</td>
</tr>
<tr>
<td>Community Services</td>
<td>$1,480,641</td>
<td>$1,844,091</td>
</tr>
<tr>
<td><strong>Total Program Services</strong></td>
<td><strong>$6,833,637</strong></td>
<td><strong>$7,391,786</strong></td>
</tr>
</tbody>
</table>

### Supporting Services

<table>
<thead>
<tr>
<th>Description</th>
<th>2019</th>
<th>2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Management &amp; General</td>
<td>$1,334,229</td>
<td>$1,197,107</td>
</tr>
<tr>
<td>Fundraising</td>
<td>$1,080,670</td>
<td>$959,680</td>
</tr>
<tr>
<td><strong>Total Support Services</strong></td>
<td><strong>$2,414,899</strong></td>
<td><strong>$2,156,787</strong></td>
</tr>
<tr>
<td><strong>Total Cost of Program &amp; Supporting Services</strong></td>
<td><strong>$9,248,536</strong></td>
<td><strong>$9,548,573</strong></td>
</tr>
</tbody>
</table>

| Change in Net Assets                       | $373,735    | $(318,820)  |
| **Net Assets, Beginning of Year**          | $3,327,218  | $3,646,038  |
| **Net Assets, End of Year**                | **$3,700,953** | **$3,327,218** |

## Financial Statements

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A world without liver disease.

OUR MISSION.
ALF’s mission is to promote education, advocacy, support services and research for the prevention, treatment and cure of liver disease.