Dear U.S. Pain community,

If you are among the estimated 100 million Americans living with some form of pain, you know all too well that it can have an enormous impact. It’s the leading cause of long-term disability, with back pain and headaches as the leading culprits. Studies show people with pain disproportionately experience depression, anxiety, and suicidal thoughts. Unsurprisingly, research shows that pain costs at least $635 billion annually in lost productivity, medical costs, and more.

Meanwhile, there is no single, proven treatment for pain; we don’t truly understand how pain works in the body. Despite this, the National Institutes of Health spends less than 2 percent of its research budget on pain, and doctors in medical school get less training on pain management than veterinarians.

People with pain deserve better. That’s why it’s so important for our organization to continue doing the work we do. Since our founding in 2011, we’ve have been able to reach even more people with pain, as well as their caregivers and clinicians. We now offer dozens of programs in education, support, awareness, and advocacy.

Whether we’re responding to a patient via email or hosting an educational workshop, our guiding principle is to offer empowerment, inspiration, and most of all, hope. We thank you for continuing to stand alongside us as we fight for positive change for people with pain.

Sincerely,

Nicole Hemmenway
Interim CEO,
U.S. Pain Foundation

“ Our guiding principle is to offer empowerment, inspiration, and most of all, hope. ”

MISSION
The mission of U.S. Pain Foundation is to empower, educate, connect, and advocate for people living with chronic conditions that cause pain.
**REACH**

- **12,500** monthly e-newsletter recipients
- **217,000** fans on Facebook
- **13,500** followers on Twitter
- **1,600** volunteers
- **960** provider offices

**EDUCATE & SUPPORT**

- **30,000** Invisible Project magazines distributed to hospital systems throughout the country
- **5** Twitter chats
- **8** webinars hosted
- **2** Support Group Leader trainings
- **Provided educational materials** more than **20,000 copies** to healthcare professionals nationwide
- **Hosted a Take Control of Your Pain Day in New York**
- **Mailed 1,200 resource kits**
- **Exhibited at 15 conferences**
- **Expanded support group network to more than 10 in-person groups and three monthly calls.**
INVITED TO SPEAK in front of major groups such as the U.S. Food and Drug Administration, Joint Commission, Harvard School of Public Health, and National Conference of State Legislatures

AMBASSADORS WORKED TO OBTAIN 52 PROCLAMATIONS declaring September as Pain Awareness Month and coordinated 60 LANDMARKS BEING LIT IN BLUE

Participated in more than 62 STATE and NATIONAL ADVOCACY coalitions, alliances, and task forces

CLOSE TO 100 TESTIMONIES and comments on patient issues at the state and national level

Featured in 45 NEWS ARTICLES AND/OR OP-EDS, including Morning Consult, The Atlantic, The Boston Globe, Newsday and Stateline

900 ADVOCATES engaged in online action campaigns

Conducted 5 SURVEYS on patient safety and access issues, which received more than 3,200 RESPONSES total

INVITED TO SPEAK in front of major groups such as the U.S. Food and Drug Administration, Joint Commission, Harvard School of Public Health, and National Conference of State Legislatures

Actively engaged in 80 LEGISLATIVE BILLS throughout the country
OUR PROGRAMS

Awareness
- Ambassador Network
- INvisible Project
- Pain Awareness Month
- Pain warrior bracelets

Advocacy
- Advocacy Network
- People with Pain Matter

Educational
- KNOWvember campaign
- Learn about Your Pain
- Medical Cannabis program
- The National Coalition of Chronic Pain Providers & Professionals
- Pain Medicine 411
- Take Control of Your Pain events

Support
- Pain Connection
- Pediatric Pain Warriors
- Veterans in Pain

Fundraising
- Real Hope, Real Heroes gala
- Points for Pain
- Fight Gone Bad
- Triumph Over Pain

“Pain Connection gave me my life back! It completely changed my outlook on how I view pain once I finished Filling in the Gaps 1 and 2 plus participating in their Pain Symposium… Am I free of pain? No. But, I am free to thrive with the newly learned techniques that I developed at my Pain Connection meetings. For that, I’m grateful!”

- ANNE SMITH, MARYLAND

“I’m so grateful that the INvisible Project allowed me the opportunity to share my story with the country. It was a life-changing experience for me. My hope is that it not only educated others about what it’s like to live with severe chronic pain, but mostly that it gave other pain warriors a glimmer of hope that life can still be good while living with pain.”

- MICHELE RICE, CALIFORNIA

“When the U.S. Pain Foundation reached out to us and asked if we’d team up with them to help fight the pain issues that our veterans are challenged with, we were quick to say yes. They have made a significant impact all over the country, and have been very effective with legislation, advocacy, and awareness.”

- ANTHONY AMEEN, ARIZONA

“When I was part of the inaugural INvisible Project in 2010, I had no idea that a year later, I would be helping the pain community through my advocacy efforts. Being a voice for the countless others who are living with chronic pain has given me a sense of inclusion and being part of something bigger than myself. I have witnessed firsthand how the patient voice is able to make a difference in state and federal legislation.”

- WENDY FOSTER, CONNECTICUT
THANK YOU!

The programs and services of U.S. Pain Foundation would not be possible without the support of our many donors, both private and corporate. Their contributions help advance our mission of improving the lives of people with pain. Detailed information about all of our donors can be found on our website.

We’d like to give special thanks to our 2018 Corporate Council members:

Platinum Members:

Gold Members: