Letter from the President
Looking Toward the Next Ten Years and Beyond

Your support in 2018 proved that we are stronger together. With your support, we introduced novel new programs, such as the Star Sisters webinar series and the Tina Talks Turner’s children’s book, which you can read more about on page five. With your generous contributions, we attended the 2019 Pediatric Endocrinology Nursing Society (PENS) Conference, where we educated the professionals who stand at the front lines of diagnosing Turner Syndrome. We are pleased to share that a TSF Professional Member spoke at the conference on Specific Care Considerations for Patients with Turner Syndrome. Thanks to you, we made great strides to fulfill our mission in 2019.

But our work didn’t begin there and it is not where it will end. This mission began a decade ago, when the message of Turner Syndrome lacked hope. The status quo saw Turner Syndrome as a lifetime of doctor’s visits and medical expenses for those lucky enough to survive – and receive a timely diagnosis. Fortunately, through greater awareness and education, we’re happy to report a movement toward acceptance, understanding, and positivity.

Over the past ten years, we’ve seen an increase in TS research and professional interest. Yet, there is still so much that remains to be accomplished. Too many girls are diagnosed in adolescence or later, too late for important medical interventions. Too many lives are lost to cardiac complications associated with Turner Syndrome due to a lack of professional knowledge. Too many girls and women spend their lives feeling alone without the chance to meet another like them.

The good news is that these problems can be solved. If we continue working together for the next ten years and beyond, we can be sure to see real change. Together we can build a community of support, education, and awareness. Join us in supporting this mission today – every life touched by Turner Syndrome is counting on you.

Your partner in change,

Laura Fasciano
Director & Founding President

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TSF Timeline
A Decade of Service

2009
TSF is registered as a 501(c)3 organization.
The Patient Registry is introduced to understand the TS community's needs and history.

2010
The first patient education workshop is held at a local library. Workshops have grown to hospital and university settings.
Team TSF, a national awareness movement, runs in its first race, the NJ Marathon.

2011
TSF hosts the first "Educating the Educators" conference in collaboration with Kean University.

2012
Importance of Early Diagnosis video airs in medical centers across the U.S., reaching over 8 million viewers.
Turner Syndrome Diaries video series is created to raise awareness and showcase positive, hopeful TS journeys.

2013
CME activity, Identifying and Managing Care of Girls with Turner Syndrome, reaches more than 27,000 health care professionals.
TS Awareness Month legislation is passed in NJ.

2014
TSF contributes to the Hormone Health Network's Transition Task Force which developed resources to promote a smooth transition from pediatric to adult care.

2015
The Council on Infertility (COI) publishes guidelines on fertility preservation in TS.

2016
The Tina Talks Turner's video and companion book is created to help young girls understand and come to terms with their TS diagnosis.

2017
The Turner Syndrome Research eXchange (TSRX) is introduced, a de-identified patient centered registry that informs research.

2018
The Council on Cardiology (COC) is introduced with a goal of lowering high cardiac mortality and morbidity rates associated with TS.
The inaugural Autumn Retreat camp weekend for TS families is held in collaboration with the YMCA.

2019
The TS Patient Handbook is developed to help individuals stay organized, manage their care, and reflect on their TS journey. Stay tuned!
Welcome New Board Members!

Lori Kobular is a veteran volunteer of the Turner Syndrome Foundation. She has taken her experience leading charity events and love for crafting and turned them into something impactful – organizing multiple successful fundraising events. Lori is proud to volunteer in honor of her daughter, Julie, who was diagnosed with TS at birth. Lori is interested in contributing to research initiatives and advocacy efforts to improve care standards throughout the lifespan.

Janis Elwell has been a Turner Syndrome Foundation volunteer since January. Her degree in Computer Science and experience in Executive Administrative roles have contributed greatly to data management in 2019. As a longtime volunteer in her community, Janis brings the perspective of our invaluable volunteers to the TSF Board. Janis aims to raise awareness of Turner Syndrome throughout her community and beyond.

Medical Advisory Board

Henry Anhalt, DO, FAAP, FACOP, FACE
Karen Berkowitz, MD
Richard Bronson, M.D.
Pasquale Casale, M.D.
Allan J. Fisher, MD, FACOG, FACMG
Robert Korwin, DMD, MICOI, MAGD
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Scott Rivkees, M.D.
Judith Ross, M.D.
Dr. Karen Rubin, M.D.
Paul Saenger, M.D.
Michelle Schweiger, DO, MPH
Martin Weinstein, D.M.D., M.S.
Sunil Wimalawansa, MD, PhD, MBA, FACE, FACP, FRCP, DSc
Serving the TS Community

Program Updates in 2019

Turner Syndrome Council on Cardiology (COC)
The COC was established to improve cardiac care outcomes to save lives. The goals of this initiative are to promote early diagnosis to improve treatment of congenital cardiovascular disease, raise clinical awareness of cardiac care guidelines, emphasize preventative care, and develop a crisis plan for emergency cardiac care. The Council has made great strides in developing a professional education module to fulfill these goals, which TSF aims to release to physicians in 2020.

Thank you to Dr. Darlene Field, TSF Medical Director, and Dr. Sheetal Patel, COC Chairperson, for leading this life saving initiative, and to the members of the COC for their hard work and dedication:

Dr. Allan J. Fischer, MD, FACOG, FACMG
Dr. Khoa N. Nhuyen, MD
Dr. Karen Rubin, MD

Dr. Helen Binns, MD, MPH
Dr. Luciana T. Young
Dr. Pankaj Madan

“Congenital or acquired cardiovascular diseases cause increased mortality and morbidity in Turner Syndrome patients. Improving their cardiovascular health through collaboration, research, awareness and advocacy can improve their overall outcomes.”

Autumn Retreat
Imagine being diagnosed with a condition that your family, friends, even your doctors are not familiar with. Unfortunately, this is the reality for many in the TS community. The Autumn Retreat provides peer support – an important, and often neglected, aspect of care – by bringing women and girls together. The result is connections and friendships that last long after the camp weekend ends.

“It was an incredible weekend. Presentations and activities were phenomenal. It was great to also have down time to bond and share experiences.”

Star Sisters Webinar Series
Education is the cornerstone of TSF’s mission. With knowledge, patients can advocate for themselves and receive proper care. Patient education began with workshops at local libraries, expanded to medical centers, and has now grown to online webinars. This new platform allows TSF to reach individuals nationwide on their own schedule. To date, physicians representing various specialties have spoke on fertility, lifelong care milestones, research, and more.
Research Initiatives

Research is a team effort, recruiting patients, researchers, students, and more to accomplish a common goal. In 2019, we are proud to partner with Coriell Institute for Medical Research, Rutgers School of Public Health, and Emory University Department of Anthropology on various studies focused on Turner Syndrome. In addition, TSF’s own research registry, the Turner Syndrome Research eXchange (TSRX) has doubled in number of participants. TSRX is a patient-centered registry that connects the global research community to improve treatments and understanding of TS.

“Turner Syndrome Foundation helps improve lives every day with their commitment to research initiatives. The Turner Syndrome Research eXchange is to their credit...The objective, give everyone a chance to attain good health and educational outcomes.”

Tina Talks Turner’s

TSF has developed a series of videos that highlight the stories of women and girls of all ages, bringing awareness to many aspects of living with TS. Most notably, the “Importance of Early Diagnosis” video has been played in doctor’s offices across the U.S., reaching more than 8 million viewers. In 2016, TSF added “Tina Talks Turner’s” to the repertoire. Studies show that sharing a diagnosis with a child leads to improved health outcomes. “Tina Talks Turner’s” provides parents the tools to share a diagnosis. In 2019, the video was developed into a companion book that features journal pages to facilitate discussion and help a young girl cope with her diagnosis.

Professional Membership

The TSF Professional Membership is a network of providers dedicated to providing the best care to the TS community through a collaborative approach, development opportunities, and a transfer of knowledge. The goal of the Professional Membership is to highlight and connect the providers specializing in the treatment of TS. TSF invited Dr. B. Michelle Schweiger, pediatric endocrinologist at Cedars-Sinai Medical Center and TSF Professional Member, to speak at the 2019 PENS Conference in Long Beach, CA on Specific Care Considerations for Patients with Turner Syndrome.

“I was so impressed, as a nurse, that TSF offers education, at no charge, for patients, parents, educators, and health care providers. There is so much information available on the web site to support the families.”
The goal of the Turner Syndrome Foundation (TSF) is to support research initiatives and facilitate educational programs to increase professional awareness and enhance medical care of those affected by Turner Syndrome. Early diagnosis and comprehensive treatments over the lifespan may lead to a brighter and healthier future for all young girls and women with Turner Syndrome.

### Who We Serve

<table>
<thead>
<tr>
<th>TS Women &amp; Girls</th>
<th>3,000+</th>
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<tbody>
<tr>
<td>Family Members</td>
<td>8,100+</td>
</tr>
<tr>
<td>Medical Professionals</td>
<td>5,900+</td>
</tr>
<tr>
<td>Donors &amp; Volunteers</td>
<td>2,900+</td>
</tr>
</tbody>
</table>

### Program Results

<table>
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<tr>
<th>One-on-One Support to Women &amp; Caregivers</th>
<th>2,000+</th>
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<tbody>
<tr>
<td>TSRX Participants</td>
<td>100+</td>
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<tr>
<td>Awareness Month Petition Signatures</td>
<td>250+</td>
</tr>
<tr>
<td>Tina Talks Turner’s Viewers</td>
<td>800+</td>
</tr>
<tr>
<td>Webinar &amp; Workshop Participants</td>
<td>200+</td>
</tr>
<tr>
<td>Monthly Website Visitors</td>
<td>10,000+</td>
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What do these numbers mean? That thousands of women, girls, caregivers, family members, and physicians are better equipped to manage their Turner Syndrome journey.
What People Are Saying

Client Testimonies

A couple years ago I was looking to get more involved with the TS community, and I couldn't be happier or more grateful that I came across the Turner Syndrome Foundation. From the start I have felt apart of such a loving, caring community, I am in awe of all the hard work everyone involved with the Foundation contributes in terms of advocacy, raising awareness, research initiatives, and making sure that every girl and woman affected by TS has the best quality of life possible. The Foundation does so much to improve the lives of every girl, and woman affected by TS and I couldn't be more honored to be a small part of it all. I am looking forward to getting even more involved in the near future.

Sarah S.

I must admit out of any other educational site on TS, the Foundation I find to be personally the best. There is always a person on the other end trying to help you in anyway they can, and it’s very much appreciated. I’d definitely recommend the Foundation if you need any form of help. Jean M.

I was diagnosed with TS at age 8. I never had any support system and was never around others with TS until recently. I am now 30. It has made for some trying times, but I have overcome those trying times and am now so ecstatic to be a part of such an amazing organization. I have made some amazing TS friends, and hope to continue with new adventures and continue to make new relationships with other women with Turners. [TSF] events truly inspire me, they help me realize and know I am not alone in this fight. That is exactly what TS is everyday, a fight. In this fight though we have each other, and together we are always Turner strong. This helps me make it each day in this life.

Samantha O.

I have been involved with TSF and been a donor for several years. It is amazing what we can do with great volunteers and hard earned funding. TSF has great ideas to help women and families dealing with TS. Unfortunately, we have to choose which ones we can fund! I wish TSF could attract the type of funding and attention that the big charities get. Imagine the things we could do. Dawn P.
Financials

Budget

Only 10% of TSF’s budget is used for administrative expenses.

- Awareness (25%)
- Advocacy (25%)
- Education (25%)
- Research (15%)
- Administrative (10%)

Revenue

- Individual Contributions (35%)
- Fees for Service (23%)
- Foundation & Trust Grants (16%)
- Corporate & Business Grants (15%)
- Other (11%)

Full financial information available on GuideStar.org.

Top Corporate Sponsors

Pfizer  Genentech  Genentech

Columbia Bank Foundation

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Make a Difference  
An Opportunity for Everyone

No matter your experience or your connection to this cause, there is an opportunity for you here. From canvassing in your neighborhood to hosting a team building fundraiser in your office, we will work with you to develop a volunteer opportunity that’s right for you – and help you along the way. Get started today by contacting volunteer@tsfusa.org.

Multiply Your Impact

One of the most valuable ways to support TSF today is to give an outright donation. When you give, you’re helping TSF advocate, deliver physician and patient education, and support new research initiatives.

To multiply the impact of your donation, consider becoming a monthly donor, designating TSF in your will or estate, or obtaining a matching gift from your employer. At TSF, we offer all of our programs and resources free of charge to the TS community. With your support, we can keep it that for the next decade and beyond. To learn how you can give today, visit www.TurnerSyndromeFoundation.org or contact philanthropy@tsfusa.org.
Thank you to our generous donors & volunteers!

Special thank you to our Professional Members:

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Seattle Children’s Hospital

Jan Foote, DNP, ARNP, CPNP, FAANP
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