Dear Friends of PDSA,

The Power of PDSA. That is the theme of this year’s Annual Report. When I took over the leadership reigns of PDSA in January of 2009, the PDSA Web site had 20,000 to 30,000 unique visitors per month. By 2014 that number increased to 50,000 to 60,000 unique visitors per month. The number of people being diagnosed with ITP has not dramatically increased, rather, more people are finding PDSA due to the collective efforts of the PDSA staff, board members, medical advisors, our industry partners, and you — our ITP awareness champions. Whether you organized a fundraiser, facilitated or attended a support group meeting, displayed your ITP awareness items, contacted the media, or sported purple for platelets, our ITP members were a huge part of our success in making the world of ITP a more manageable place to be!

I am proud to present our 5th Annual Report, which provides a snapshot of activities and major accomplishments. This past year PDSA and its network of volunteers served the greatest number of people with ITP in the history of the organization. We are proud of our many accomplishments — through social media; public awareness initiatives; and increased engagement with patients, family members and the health care community. We achieved record-breaking visibility, yet we know there is still much work to be done.

PDSA receives no federal funding. It never has. That is why the support of our individual and corporate donors, along with the efforts of our fundraisers and volunteers is crucial in helping PDSA to fulfill its mission. I am grateful to, and inspired by, the commitment of those who work to make this possible every day — those who give their time, energy and resources to make a difference in the lives of people who struggle with ITP and other platelet disorders. One look at this “year in review” will reveal the significant impact of their support and involvement.

PDSA was founded on the premise that informed patients are in a better position to understand their disease, have more meaningful dialogue with their doctors, and potentially have an improved opportunity to heal. We continue to be true to our mission and are heartened by messages from our members indicating that our information and support have been truly helpful, and in some cases life-changing and life-saving, in their journey toward better health. Whatever role you play within this dynamic community, your efforts help to ease the burden of those living with this disease. More importantly, your support helps us travel down the road toward better treatments and the hope of a cure.

Thank you,

Caroline Kruse
Executive Director
PDSA provided a wide variety of online and printed information to increase knowledge of ITP, treatments, and wellness practices.

**Newsletter**

Our quarterly newsletter, *The Platelet News*, offered the latest research and recommendations for treatment and improved health for ITP patients. These issues included reports from our annual meeting, advocacy meetings, and annual hematology conferences. They provided updates on PDSA support groups around the U.S., fundraising news, patients’ letters and stories, and other pertinent news.

**E-News**

Our monthly complimentary online newsletter was distributed to more than 17,000 on our mailing list. Each issue summarized 4-7 recent, relevant news articles related to ITP or general health, and provided resources to patients and families.

**Patient Booklets**

PDSA has published 21 educational booklets, including nine of our most popular booklets available in Spanish and French. In 2014 we added a new illustrated children’s booklet “Understanding ITP.” All of our booklets are distributed free at local support group meetings, medical meetings and conferences and are available by mail. Members are encouraged to take copies to their hematologists to reach ITP patients and increase awareness of ITP. PDSA partnered with Health Monitor Network on our 6th patient Guide to ITP. Featuring an ITP patient on the cover and PDSA members throughout the Guide, 150,000 copies were distributed to medical institutions and hematology offices across the U.S.

**Web Site**

Our Web site (www.pdsa.org), with more than 200 pages of FREE information, was updated throughout the year with the latest research and news. During 2014 the PDSA Web site received 50,000 – 60,000 unique visitors per month from 130 countries.

**Social Networking Sites**

PDSA’s social networking pages continue to grow in popularity. We now have more than 10,000 Facebook friends and Twitter followers. Our Inside Stories patient and medical expert videos have been viewed more than 12,000 times on YouTube and Vimeo.
Advocacy & Research

At industry and government meetings, PDSA serves as the voice of the ITP community. It represents and shows the strength of our community. In 2014 PDSA achieved its goal of encouraging advocacy and research for the treatment of ITP and other platelet disorders in the following ways:

We participated in meetings of the American Plasma Users Coalition (A-PLUS), an organization devoted to assuring quality IVIg access for all who need it. In 2014 this group addressed and commented on the following issues: Biologics Safety and Access, 21st Century Cures for Rare Diseases, Essential Health Benefits, MSM Blood Safety.

PDSA staff attended the following meetings:
The American Society of Hematology (ASH), European Hematology Association (EHA), Immunoglobulin Nursing Society (IgNS), National Patient Advocacy Leaders Summit (PALS), Ig Academy, Plasma Protein Forum, 7th Annual Rare Disease Day, 4th World Orphan Drug Congress, Role of Investigational New Drugs in Patient Care.

Meetings

To meet our goal of supporting the exchange of information between patients, researchers, physicians and other members of the wellness community, PDSA organized and hosted its 14th ITP Conference in 2014.

Annual Conference

ITP Conference 2014 was held July 11-13 in Manhattan Beach, CA and was attended by more than 200 patients, presenters and associates. Ten of the world’s most renowned ITP clinicians and researchers presented programs and met in small groups or one-on-one with patients and caregivers. The conference offered an advocacy program for Canadian patients, a special teen program, and our 2nd annual Kids Kamp. The conference included a Saturday evening dinner program, family fun event with a live band and a fundraising membership raffle. Scholarships were awarded to patients in need.
Support

In 2014 PDSA provided support services that promoted patient communication with other patients, physicians, and medical providers.

Support Groups
PDSA’s local support group program continues to grow each year with a total of 38 groups now actively engaged in major cities around the U.S. and in Canada. Our teleconference support group for parents of children with ITP celebrated its fifth anniversary with over 50 families participating and our teleconference support group for Canadian patients and family members continues to add more members.

Support Group Facilitators’ Workshop
At its annual conference, PDSA offered a workshop for its group facilitators, who provide important non-medical assistance to ITP patients and their families.

Online Discussion Group
PDSA provided online discussion groups for adults with ITP, teens with ITP, parents of children with ITP, and ITP and pregnancy. The site receives thousands of visits each day.

Personal Counseling to Members and Non-members
Our staff responded to more than 1,000 emails, letters and phone calls requesting information and answers about platelet disorders.

Name Exchange Program
We have more than 900 participants in our Name Exchange Program, developed for patients who want to communicate one-on-one with another patient.

Audi Gerstein Scholarship
The PDSA’s Audi Gerstein Scholarship Program provides financial assistance to senior high school students and college students who are suffering from ITP or a similar platelet disorder. In 2014 PDSA awarded its sixth $1,000 Audi Gerstein Scholarship to a college student with ITP. Several other students received $100 book awards.

“I just want to say that we LOVE the Buzzy and my son was so excited to get one of his own so that no matter where we have to go for draws, we always know we have a Buzzy available. The conference calls for parents of ITP kids are fabulous, too, both for my son AND the rest of our family. It means so much to us to have others to talk to who understand what we are going through. We are grateful for the help and support we have received from the PDSA as we navigate this journey with Chronic ITP. Thank you!” – Jana R.

“My ITP turned out to be a blessing in disguise. It has taught me some important lessons and has helped me become more comfortable with myself. Not only that but it gave me the chance to seize opportunities I probably would have never have gotten the chance to have.” – Julia Gabor, $1,000 Audi Gerstein Scholarship winner 2014, New York, NY, St. Louis University

“I wanted to thank you so much for letting me take part of the conference call yesterday evening. I am new to ITP as my 9-year-old son was just diagnosed 3 weeks ago, partaking in the conference call gave me a better understanding of what we are facing. Although I have researched it extensively and asked his pediatric Hematologist one million questions, there is always something else to learn. The parent to parent part of the conference call is priceless, this is a wonderful service that you are providing and I thank you from the bottom of my heart. Thanks again!” – Gloria A.

Attendees of the 2014 Facilitators’ Workshop in Manhattan Beach, CA
During the month of September we celebrated National ITP Awareness Month. ITP patients, family members, friends and our industry partners joined PDSA to raise public awareness of ITP. A number of activities moved us closer to our goal of increasing the public’s understanding of ITP and other platelet disorders and communicating the important resources and support PDSA offers for ITP patients, families, and health care providers.

**Sport Purple for Platelets Day**
The last Friday in September found people young and old, from schools to business offices, sporting shades of purple.

“Just found out about this organization and proud to see what you guys are doing! I had ITP until I had a splenectomy (sadly enough) few years ago. I surely would’ve loved to see all the support and help you all are giving to each other! Keep that up!” – Sarah B.

**Awareness Materials**
PDSA staff distributed more than 2,500 awareness bands, 2,500 ITP awareness pins, and more than 2,500 educational brochures, pamphlets and school information packets.

**International ITP Alliance**
PDSA collaborated with European ITP patient advocacy groups to share ideas and strategies to foster greater awareness and understanding of ITP, a little-known disease around the world.

“I’m so excited that I found this! I have been dealing with ITP for so long and as of the past year and half have had so many issues. So nice to see what has been done for others. I feel hopeful again.” – Cathy S.

**PDSA Packets**
More than 500 educational packets were distributed to hematologists and other medical professionals.
During 2014 there were many outstanding fundraising events held that supported PDSA and helped raise awareness of ITP and PDSA, including:

**Pump it Up for Platelets**

It was another great year for PDSA fundraisers and our signature event *Pump It Up For Platelets!* ~ for a world free of ITP thanks to the continued hard work of our organizers who dedicate their time and resources to ensure successful events. These events continue to grow, inspire, raise public awareness, and connect those affected by ITP with others in the community.

Many friends of PDSA held a variety of fundraisers, raising close to $23,000 for programs and research: Archer & Greiner Law Firm Fundraiser, Conestoga-Rovers & Associates Fundraiser, Triathlon for Ayla Charness, Hope for Haley Benefit Ride, Lewis University/Tri-Beta Biological Honor Society, Montreal Half Marathon, Party for Platelets, Paula Lagree Dinner Party and BBQ, Pedal for Platelets: El Tour de Tucson, Showdown at Sundown, and Giving Tuesday.

“Just spent almost 2 hours on a conference call sponsored by PDSA for kids with ITP and their parents. This is why we are trying to raise funds for PDSA. I got to talk to a Pediatric Hematologist from Boston who is an ITP expert and has experience with Nplate.” – Tammi C.
Contributions in 2014

Each year thousands of individuals, organizations, corporations and foundations help support PDSA through donations of gifts, tributes in memory or honor of family and friends, sponsorships of new programs and special events, and contributions to the annual fund. PDSA receives NO federal funding. Since 1998, it has been the caring members of the ITP community who have given life to PDSA’s mission of helping children, adults and families affected by ITP and other platelet disorders. Every gift is important and deeply appreciated. The following list includes gifts received between January 1, 2014 and December 31, 2014.

The Circle of HOPE

CIRCLE OF HOPE ($5,000+) – INDIVIDUALS
Louise Kittel
Barbara & Peter Pruitt, Jr.
Jacqueline & Bruce Prescott

CIRCLE OF HOPE ($3,000 – $5,000) – INDIVIDUALS
Matthew Linde

CIRCLE OF HOPE ($2,000 – $2,999) – INDIVIDUALS
Cindy & Philip Ayliff
Michael Tarantino, MD
Kimberly & Rodney Hall
Daisie Wong
Joan & Richard Jordan
Cheri & Derek Zimmerman
Karen Rosenbaum

CIRCLE OF HOPE ($1,000 – $1,999) – INDIVIDUALS
Anonymous
Janice Ippolito
Karen & Adam Avrick
Eric Johnson
Kenda & Brian Bamesberger
Ann & William Jones
Bonnie & Mark Barnes
Diane & Robert Joseph
Mary & James Benvenuto
Caroline & Ken Kruse
Amita & Ranjive Bhalla
Irene & David Kuter, MD, DPhil
Brendan Cameron
David Lihani
Audrey & Jay Charness
Mary Lou & James Lyons
Melanie & Will Deaver
Eleonore & Russell McCabe
Robert Feiner
Daniel Miles
Joseph Fitzgerald
Rae Ellen Pistone
Allison & Keith Flowers
Brian Pomianowski
Jan & Steve Gardner
Nancy & Michael Potthast
Terry Gernsheimer, MD
Neil Prescott
Barry Gesser
Shirley & Peter Pruitt, Sr.
Terri & Michael Goldberg
Barbara & Mark Rice
Robert Halonen
Steven Rodgers
Kala Harvey
Lois Umhoefer
Kristin Henrikson & Jon Brandt
Elizabeth & Robert Welch
Matthew Heyman
Jess Weiner
Madeline & John Hromyak
Neil Weiner

Matching Gifts
AT&T United Way Employee Giving Campaign
The Boeing Company
IBM Employee Services Center
JP Morgan Chase & Co.
MGM Resorts Foundation
Nestle
Pfizer Foundations Matching Gifts Programs
Progressive Casualty Insurance Company
Raytheon
Schwab Charitable Fund
Wells Fargo Community Support Campaign

Income Funding

- Contributions 17%
- Sales & Other Revenue 3%
- Advertising 5%
- Membership Dues 13%
- Foundation Grants 49%
- Conference 13%

How the Funds Were Used

- Patient Education 37%
- Programs 11%
- Patient Support 23%
- General and Administrative 22%
- Fundraising 7%
CIRCLE OF HOPE – EVENT ORGANIZERS

Pump It Up for Platelets!
Chicago – $9,211.31
Kim & Jim Everett – Organizer
Susan Frank – Organizer
Trish & Paul Santaromana – Organizer
Emily Auterson – Team Organizer

Pump It Up for Platelets!
Everett, WA – $8,922.50
Jenn & Andy Lindal – Organizers

Pump It Up for Platelets!
Hannibal, NY – $1,700
Arlene Laut – Organizer

Pump It Up for Platelets!
Iowa – $1,069.42
Sherry Frazier – Organizer

Pump It Up for Platelets!
Irvine, CA – $3,455
Melissa Hilsabeck – Organizer

Pedal for Platelets!
El Tour de Tucson – $15,074
Peter Pruitt – Organizer
Michael Tarantino, MD – Organizer

100 Mile Challenge – $1,777
Kristin Hunt – Organizer

The McGuirl Family – Organizer
Meredith Prescott – Organizer

Pump It Up for Platelets!
Livingston, NJ – $36,740

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Pump It Up for Platelets!
Livingston, NJ – $36,740
The McGuirl Family – Organizer
Meredith Prescott – Organizer

Give Myself Away Project – $1,235.91
Poonam Kharel – Organizer

Hope for Hayley – $1,251
Kimberly & Ron Sewald – Organizers

Lagree Fundraisers - $2,074.18
Paula Lagree – Organizer

Lyons Fundraiser – $1,883
Kristie Lyons – Organizer

Molino Fundraisers – $4,159
Yvonne Molino – Organizer

Parents Partner for PDSA – $1,160
Kim Barber – Organizer

St. Paul’s Elementary Fundraiser – $1,180.41
Terra-Lee Gratton – Organizer

Tri Beta Bio Honor Society/
Lewis University – $1,052
Jennifer Roberts, PhD – Organizer

SUPPORTERS ($1,000+)

Barnstormers Aerial Advertising
Bleeding & Clotting Institute
The Boeing Company

Carchia Financial Management Group
of Wells Fargo Financial
Design Distributors

Greg Norman’s Australian Grill
Lake County Press, Inc.

MEMBERS AND SUPPORTERS ($500 – $999)

Anchor Associates Group
Archer & Greiner Foundation
Barbara & Jim Brady
Brock Bridge Elementary
School/Angie Como
Karen & Michael Bukiet
CP4, Inc.
Salvatore DiFazio
Joyce Fleming
Ann George
Randy Gilman

Rebecca Goldberg
Michael Grunwell
Chris Gunder
Douglas Hovanec
Sherrill Hudson
Paula & Thomas Huntzinger
Mary & Edison Jean
Tom Joplin
Dana Matthews
Laurie & Brian McAleenan
Janet & William McAllister

Katharine & Edward McCleary
The McGuirl Family
Mike Molnar
Terry O’Connor
OCS, LLC
Susan & Dale Paynter
Keith Pulley
Jennifer & Sanjay Puri
William Resnick
Howard Rosenthal
Beth Siegelbaum

Bunnie Stevenson
Sachiko Terrible
Raju Tuladhar
Sonia & Jeremy Vandama
Walmart
Sharon Webb
Becky & Timothy Wirtz
Laura Weiner

MEMBERS AND SUPPORTERS ($250 – $499)

9 Dots Technology
American Vending Sales
Susan Anderson
Anonymous
Rebecca Antas
Roya Ara
Lois & David Bacon
Wendy Bauer, MD
Mona Bobrow
Sangeeta & Richard Bolesla
Southwest Dog Obedience Club/
Julie Bremer & Jennifer Necker
Timothy Byun
Peter Cabrera
Donna Carter
Kenneth Christiansen
Maureen & Gregory Church
Dru & Todd Davies
Michael De Michele
Rita Effron
Magda El Nokaly
Lainey Fallek
Sarabeth Fields
Sherry Frazier
Terri & Mark Friedman
Michelle Gass
Angela & David Girgus
Denise & John Gordon
Lori Gottsegen
Bruce Gross
Hanson Chiropractic
Deborah Harlan
Fred Hasler
Eva & Istvan Hegedus
Jonathan Hilsabeck
Teri Howe
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Frank Laezza
June Lee
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Rachel & Hope Lomita
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Linda Meisel
Mercedes Benz of Coral Gables/
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Colleen Miller
Cheryl Miller
Beverly Miller
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Nora Thornber
Ronald Virgin
Risa & Rob Weinstock
Laura & Kevin White
Mike Wicks
Benjamin Willig
Gayle Wills
Margaret Zivelonghi

*Denotes deceased donor
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<td>Charles Rosner</td>
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<td>Corney Urps</td>
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<td>Wells Fargo Community Support Campaign</td>
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<td>Dagus Zuchowski</td>
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WE HAVE RECEIVED MANY ANONYMOUS DONATIONS FROM THE FOLLOWING UNITED WAY/CFC CHAPTERS:

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United Way of Delaware CFC
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What our members are saying:

“Hello, I want to thank you from the heart for keeping me on your email list. God bless you for the information you send to help all who have ITP. With having so many doctor bills and hospital bills it’s difficult to have any extra funds. The information helps to ask the doctors what we need to do in ways to make life better. I am now watching what I eat and exercise more. That does help. Take care and GOD bless all of you for caring the way you do.” – Brenda A.

“PDSA is a wonderful organization with lots of information on ITP. Have been a member for 12 years. It has been a great source for me.” – Joann B.

“Thank you so much for doing what you do and being here for us. I receive so much valuable information from you.” – Pat H.