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## The IFOPA Mission

*To eliminate FOP as a health concern through education, advocacy, research and support.*

## The IFOPA Vision

*To instill hope worldwide while searching for a cure.*
What is Fibrodysplasia Ossificans Progressiva (FOP)?

One of the rarest, most disabling genetic conditions known to medicine, FOP causes bone to form in muscles, tendons, ligaments and other connective tissues. Bridges of extra bone develop across the joints, progressively restricting movement and creating a second skeleton that encases the body in a prison of bone.

This immobilizing and catastrophic genetic condition strikes approximately one in 2 million people. It is estimated that 2,500 people worldwide have FOP, although there are only 700 confirmed cases and some 285 cases in the United States.

FOP, known by many names throughout its history, first appeared on the medical landscape more than 250 years ago when a London physician, John Freke wrote in a letter to The Royal Society of Medicine in which he described a 14-year old boy “with many large swellings on his back…they arise from all the vertebrae of the neck and reach down to the os sacrum; they likewise arise from every rib of his body, and joining together in all parts of his back, as the ramifications of coral do, they make, as it were, a fixed bony pair of bodice.”

The disease eventually became known as myositis ossificans progressiva, which means “muscle turns progressively to bone.” The condition was officially modified to its current name in the 1970’s by Dr. Victor McKusick of Johns Hopkins University School of Medicine, to acknowledge that other soft (fibrous) tissues in addition to muscle are replaced by bone.

Symptoms of FOP almost always begin in the first two decades of life, and the majority of patients learn they are affected before the age of ten. Inflamed and painful swellings, usually in the shoulder and back areas, are the first signs of the condition. These swellings eventually clear up, but leave behind an area of mature bone.

FOP patients experience different rates of bone formation, and the exact rate of progression is unpredictable. The bone growth can occur spontaneously, or can be triggered as a result of trauma, an injury, bump or fall. The extra bones grow in the same order that they are first formed in the fetus – from the head, shoulders, arms and progressing to the hips and knees. As the condition worsens, patients eventually have joints locked in fixed positions.
The International FOP Association

The International FOP Association is a 501(c) (3) non-profit organization formed to support those afflicted by the rare genetic condition Fibrodysplasia Ossificans Progressiva. The IFOPA's mission is to eliminate FOP as a health concern through support, research, advocacy and education. The vision of the IFOPA is to search for a cure while instilling hope worldwide.

The IFOPA was established in 1988 by Jeannie L. Peeper, a woman with FOP. The focus of the IFOPA has been, and will always be, to find a treatment and cure for FOP. Until that day, the organization strives to improve the lives of people around the world who suffer from the condition while continuing to support research.

The IFOPA's programs and services are focused in four primary categories – research, education, support and advocacy.

Since its founding in 1988, the IFOPA has directed over five million dollars to the Center for FOP Research at the University of Pennsylvania School of Medicine. Nearly $500,000 is restricted each year from IFOPA operations for research.

Awareness is among the top objectives of the IFOPA. Nearly 90 percent of patients are misdiagnosed, with the correct diagnosis taking an average of four years. Potentially harmful tests, procedures and biopsies are often conducted on patients before a correct diagnosis is made. The organization provides education to the medical community on early signs of FOP and distributes information about the condition via the IFOPA website.

The Betty Ann Laue FOP Library is a central repository for FOP information that offers scientific and medical articles, interesting stories on FOP members, information materials, videos, special interest articles and copies of medical publications for families, physicians and those interested in FOP.
Support to patients and families ranks as a top programmatic objective.

Periodic symposiums bring together FOP patients with health professionals that specialize in FOP research and care. These gatherings provide a wonderful opportunity for patients to gather in a collaborative forum to discuss FOP research, exchange information and experiences, and share coping strategies.

The Traveling Resource Center (TRC) provides the global FOP community with a traveling showcase of tools and devices available for use in daily living. The TRC has had a presence in numerous FOP gatherings across the world.

The IFOPA provides L.I.F.E. Awards (Living Independently with Full Equality) to its members and is designed to promote health, welfare and independence. Examples of the awards range from equipment, walkers, wheelchairs, adaptive beds, vehicle adaptation, tuition/registration for educational purposes, specialized bicycles, computer hardware or assistive technology and reading aids.

The FOP Connection is a newsletter distributed to the FOP community and features stories about members, fundraising events, research updates, helpful suggestions and resources for living with FOP. An online member forum provides a vital medium to connect members for discussion, support and research news. Periodic web-based E-Learning events spotlight issues that are of relevance to members.

The IFOPA sponsors a mentoring program that provides members with the support they need to better manage the daily challenges of living with FOP. Additionally, new members receive a comprehensive resource package which includes the following items: What is FOP? A Guidebook for Families, a medical binder which includes a flash drive containing important medical information, a portable pill holder which holds one dose of emergency medication, and medical forms to customize with their medical information.

IFOPA’s financial support comes almost exclusively from families, friends and others who join forces by holding barbecues, golf events, auctions and other events to raise the funds to support the organization’s annual operating budget. Nearly half of the one million dollars generated each year is directed toward research efforts at the FOP Research Lab at the University of Pennsylvania.
Our Founder
President, Jeannie L. Peeper

Jeannie Peeper’s pioneering spirit, vision and unwavering devotion to the cause of finding a treatment and cure for FOP has been a source of strength and beacon of hope to the many patients and families within the FOP community.

Founded as a pen pal group in 1988, she led the organization through a period of unprecedented growth into a world class non-profit that has emerged as the number one global resource on FOP. Today, the IFOPA serves over 700 patients and families in 52 countries through education, public awareness, program-based initiatives, networking opportunities, and support to families while instilling hope through research while searching for a cure for FOP.

A native of Ypsilanti, Michigan, Jeannie’s life began to change at an early age with the onset of FOP. Instead of allowing the condition to overwhelm her, she began a journey that allowed her to earn a college degree, and create an organization that has ended the isolation formerly associated with the disease to connect and support persons with FOP and their families.

Jeannie has served as President of the IFOPA since its founding. In 1989, she began working with Dr. Frederick Kaplan, who was established as the IFOPA medical advisor. In 1992, Dr. Kaplan and Dr. Eileen Shore founded the FOP research program at the University of Pennsylvania, and it remains the preeminent center for FOP research in the world. Since then, the IFOPA has provided more than five million dollars to the quest for treatments and a cure. A team of over 15 researchers led by Drs. Kaplan and Shore are working and progressing steadfastly to solve the mystery of FOP.

The list of IFOPA achievements under Jeannie’s leadership are multiple, and include hosting four international symposiums for families and medical researchers, establishing a website, participating in many media projects, hosting organizational/strategic planning meetings for the IFOPA, supporting many family gatherings, visiting with donors and providing valuable counsel to many FOP families.

Jeannie is the symbol of the FOP cause, emblematic of the close-knit community that is unrivaled in its linkage of patients, doctors, scientists, researchers, families and the laboratory all united in the search for treatments and a cure. She has provided the passion and the dream to help the next generation of FOP patients.
A Message from Jeannie . . .

As I reflect on 2011 and all of the amazing accomplishments of the IFOPA, I would like to personally thank each of you for your extraordinary dedication and generosity.
Let’s celebrate our successes and YOU!

1. To our families who have fundraised this year: You have done a phenomenal job. We could not reach our goals and achieve what we have done without the extraordinary work of your grass-roots efforts. We love you and thank you.

2. To our donors who generously continue to give to the IFOPA each year: We are extremely grateful for your support of the mission of the IFOPA. Thank you for keeping us in your heart, for your warm thoughts and for your generous gifting.

3. To the many volunteers who give unselfishly of their time and talents at fundraising to help make each event a huge success. Our dedicated volunteer members who serve on committees and provide support on our projects and our volunteers who have worked diligently at the IFOPA office this year: We thank you and appreciate you!

4. To the FOP Community worldwide for your support to members and their families: Your knowledge, strength, inspiration and wisdom provide encouragement and hope. There is presently an amazing connection of people on Facebook sharing their lives with each other and I always look forward to reading what our friends around the globe are doing. As well as, the International President’s Council members who translate a variety of important documents for our membership. You are all so incredible!

5. To Dr. Fred Kaplan, Dr. Eileen Shore, Dr. Bob Pignolo and their research team: Your extraordinary love and dedication for us will enable us to achieve the ultimate goal of a treatment and a cure for FOP. That is a blessing and a miracle. The extraordinary Strategies for the Treatment of FOP meeting held at The University of Pennsylvania earlier this year was a remarkable milestone in our quest for a cure, and has engaged and energized researchers in an outstanding way.

6. To the IFOPA Board of Directors who has given graciously of their time, talents and effort to reorganize the office priorities and staffing needs to keep the organization moving forward: It has been a great labor of love: Your exceptional diligence demonstrates the wonderful leadership and unity of our organization. A special thanks to Mark Gambaiana, Christine Ford and Gail Weakland for traveling to the IFOPA office to conduct potential employee interviews. Mission accomplished!

The excitement of the coming year.......2012.....what will it bring us? We don’t know for sure, but with the recent new research developments we have HOPE that clinical trials may be a possibility in the not too distant future. We continue on our journey with each other, with love, courage and HOPE.

With Love and Gratitude,

Jeannie L. Peeper
Founder and President
Thank You from the IFOPA Board Chair . . .

The IFOPA enjoyed an exceptionally strong 2011 year on multiple fronts. A budget dotted with black ink, the ability to send significant resources to the FOP research lab at the University of Pennsylvania, and a renewed interest in conducting family fundraisers across the globe were all notable highlights.

The IFOPA enjoyed a solid fiscal year in 2011, with revenues of $850,999 and expenses of $694,641, strengthening the organization’s net asset position by $156,358. At a time of continued economic distress and numerous challenges for non-profit organizations, our performance was noteworthy.

I extend deep personal thanks to all those families and friends who have conducted various fundraising events as our organization is funded exclusively from private contributions. Sometime in 2012, we will top $15 million in revenue since Jeannie Peeper founded the IFOPA in 1988, a monumental milestone. Without our family fundraisers, we would not be able to fund the FOP research lab, conduct programs and provide support and assistance to members and families.

The crown jewel of the 2011 year was the August scientific workshop where 27 researchers representing 11 universities and one pharmaceutical company assembled at the University of Pennsylvania. Vigorous discussion and the establishment of priorities for developing effective treatments and an eventual cure were primary takeaways from this gathering of FOP “all-star” researchers. We look forward to capitalizing on the momentum generated from the workshop, and the continued devotion of scientists and researchers at the FOP lab, many of whom have dedicated their life’s work to our cause.

Since 2009, I have met several FOP families across the country. An incredible dedication to the cause, perseverance, resiliency and optimism for the future are all common denominators of members and families. I have tremendous respect for those pioneering FOP families who have been championing this cause for many years. Having the opportunity to work with our founder, Jeannie Peeper, is a remarkable privilege. Jeanne’s pioneering spirit, vision and unwavering devotion to the cause of finding a treatment and cure for FOP provides inspiration and propels us all to keep moving forward.

Looking ahead to 2012, our 24th year of operation, the IFOPA will again focus on its programs and services in the areas of research, education, support and advocacy. On behalf of the IFOPA Board of Directors, I encourage your continued involvement and support!

Mark Gambaiana
IFOPA Board Chair

Vigorous discussion and the establishment of priorities for developing effective treatments and an eventual cure were primary takeaways from this gathering of FOP “all-star” researchers.
2011 in Review
Scientific Workshop for a Cure
Strategies for the Treatment of FOP

By Frederick S. Kaplan, MD
Robert J. Pignolo, MD, PhD
Eileen M. Shore, PhD

In mid-August, 2011, 27 scientists (13 from the University of Pennsylvania and 14 from other institutions) met in Philadelphia for an informal, but intense two-day workshop on: “Strategies for the Treatment of FOP.” The workshop was co-organized by Frederick S. Kaplan, MD, Robert J. Pignolo, MD, PhD, and Eileen M. Shore, PhD, with the tireless and gracious support of Mrs. Amanda Cali. The workshop was sponsored by generous contributions from Mrs. Diane Weiss, the IFOPA, the University of Pennsylvania, and an anonymous donor. The primary goal of the workshop was to establish priorities for developing effective treatments and an eventual cure for FOP. An important secondary goal of the workshop was to stimulate and strengthen basic scientific and translational collaborations that will lead to the development of a cure.

Participants were provided with an extensive reading list of the most up-to-date scientific articles in the field of FOP research a month prior to the meeting, and were encouraged to scour the details before arriving in Philadelphia. All clearly did.

Discussions ensued in a series of small workshop sessions focused on four highly plausible areas of therapeutics, all bolstered by recent advances in FOP research:

1. Down-regulating ACVR1/ALK2 signaling (from the FOP gene and its downstream targets)
2. Inhibiting neuro-inflammatory triggers of heterotopic ossification
3. Blocking responding progenitor cells
4. Altering the molecular microenvironment of the incipient FOP lesion

The meeting was interactive and vibrant, highlighting an agenda that limited formal lectures and promoted brainstorming. The discussions and collaboration-building continued seamlessly well beyond the designated sessions. Participants discussed unpublished data and novel ideas in a dynamic interdisciplinary forum that began in a conference room environment and evolved into informal gatherings.
After brief introductory greetings and scientific updates, the scientists self-assigned themselves to one of three small working groups (BMP pathways; Lesional stem cells; or the Microenvironment of early FOP lesions) for “shop talk” and a discussion of scientific and therapeutic priorities. Each workshop self-selected a group discussion leader who focused the conversation on the broad general question: “What is the biggest gap between what we know and what we need to know in order to achieve successful treatment and a cure for FOP? How do we get there in the short term (1-3 years) and in the long term (3-5 years)?”

Following the introductory workshops, the small working groups reconvened to summarize their proceedings for all in attendance. The remainder of the meeting was spent in several larger workshops where all of the scientists participated in the agenda of each of the smaller working groups.

The workshop participants represented diverse and far-ranging scientific fields relevant to the understanding and treatment of FOP including genetics, cell and molecular biology, immunology, neurology, vascular medicine, structural biology, biomechanics, animal model development, medicinal chemistry, pharmaceutical development, microenvironmental biology, and clinical and translational medicine. Manuel Robert, a 14 year-old young man with FOP from Buenos Aires, Argentina was commissioned to depict the theme of the workshop and the goal of all FOP research in a specially designed, computer-generated poster. Selected from among six renditions, the illustration joins the pantheon of work from historic FOP meetings:

1991: Save the FOP Children (Ashley Kurpiel)
1995: Accentuate the Positive (Sarah Steele)
2001: We are the World (Tiffany Linker)
2007: Together We Can Move Mountains (Hugo Fahlberg)
2011: Cure FOP (Manuel Robert)

The illustrated theme, a mandate in two words “Cure FOP,” says all that needs to be said as the scientists moved the jigsaw pieces of the FOP puzzle into place. The meeting focused on critical pieces that still might be missing as well as currently available knowledge that could be translated most quickly into more effective treatments.
Recalling an earlier time, two decades ago (September, 1991), at the first FOP Symposium in Philadelphia when war was declared on FOP, the scientists recreated a war room atmosphere as they sat face-to-face around a large conference table and sketched the outlines for a final assault on a long-term enemy – FOP itself. The workshop culminated in an intense working session where treatment options, as presently conceived, were prioritized for further basic research and pre-clinical development.

The Workshop think tank atmosphere was palpably exciting with a verve and tone unparalleled in the history of FOP research. “The tools and enthusiasm were never more available to do so much,” said one scientist commenting on the enlightening exchange of ideas. “One could smell the ozone in the room.”

The workshop was peppered with a series of brief inspirational messages from organizing members of the core FOP Laboratory and from The Center for Research in FOP and Related Disorders as well as from the Dean of the Medical School, the Chairman of the Department of Orthopaedic Surgery, members of the IFOPA Board, and most memorably from FOP patients Patrick Doerr and Ian Cali whose messages were moving and inspirational.

The workshop, held at the Inn at Penn on the University of Pennsylvania campus, began with an informal reception at which the attendees were greeted by members of the Penn academic community and representatives of the IFOPA Board of Directors and culminated with a deep appreciation by all of the mysteries and miseries of FOP and the mandate and possibility for a cure. As Ian Cali said, we have taken the step “from hopeless to hopeful.”

Seated (from left to right): Mrs. Amanda Cali, Mr. John Cali, Dr. Robert Pignolo, Dr. Fred Kaplan, Dr. Eileen Shore, Dr. Aris Economides, Dr. Michael Zasloff; Standing (from left to right): Mr. Ian Cali, Mrs. Robin Gambaiana, Mrs. Jennifer Snow, Dr. Viet Le, Dr. Kristi Wharton, Dr. Bjorn Olsen, Dr. Jay Groppe, Dr. Vicki Rosen, Mr. Mark Gambaiana, Dr. Ernestina Schipani, Dr. Corey Hopkins, Dr. Karen Lyons, Mrs. Diane Weiss, Dr. Petra Seemann, Dr. Masahiro Iwamoto, D. Bettina Mucha-Le Ny, Mr. Robert Snow, Dr. Joseph Kitterman, Dr. Emile Mohler, Dr. Malek Kamoun, Dr. Mairizio Pacifici, Dr. Damian Medici, Dr. Lixin Kan, Dr. Ed Hsiao, Dr. Charles Hong, Mrs. Moira Liljesthrom.
IFOPA Awards

Jeannie Peeper International FOP Association (IFOPA) Awards recognize those who make exceptional contributions to the worldwide FOP community through their leadership, service, fundraising and accomplishments. This year’s Awards Selection Committee—Lori Henrotay (Chairperson), Patrick Doerr, Karen Munro, Jeannie Peeper (Honorary Chairperson), and Nancy Sando—would like to thank everyone for your participation. It is very difficult to choose just four winners because all of you are deserving of an award. We hope to continue each year to recognize and highlight the extraordinary accomplishments within our community. Below are brief summaries of each winner’s extraordinary contributions.

President’s Lifetime Leadership Award

Amanda Laue Cali

Upon Ian’s FOP diagnosis in 1994, Amanda embarked as a mother on a tireless journey to fund FOP research, promote awareness and assist the IFOPA in many facets. With her family she established the Ian Cali FOP Research Fund with its primary focus on supporting collaborative ventures between the FOP Lab and other centers of research excellence. She created many first time events including the FOP Parent Meeting in Cranford NJ, the Adult FOP Meeting in Parsippany NJ, Planning for Today and the Pathway to the Future IFOPA Organizational Meeting, and the wonderful Mother’s Retreats. Amanda provided outstanding service as an IFOPA director and as the first Board Chairman after Jeannie’s retirement. Among her many initiatives, she provided important guidance and support in establishing the International President’s Council. We are deeply grateful for her quiet and amazing generosity in providing much valued financial and organizational support on so many occasions over the years.

Outstanding Community Involvement Award

The Weldon Family

In reviewing the Weldon family’s many contributions and supportive activities for the IFOPA and the FOP community, we would like to recognize them for the scope and variety of their fundraising and awareness activities. As a start, of course, there is the establishment of the Weldon FOP Research Fund with its focus on the genetic causes of FOP and the development of safe treatment options. In addition, their extraordinary efforts at organizing events such as the annual golf tournaments and luncheons, dinner dances and celebrity boxing bouts, cocktail parties and comedy shows,
and Friendraisers and recruiting Boston Marathon runners (to name only some!) are truly impressive. The extraordinary outreach of the Weldon family each year to bring FOP awareness through local media coverage via television and radio is amazing and we are deeply grateful for all that they have done for the FOP Community.

Emerging Leader Award

**Ashley Kurpiel**

Ashley is an incredible role model and an amazing spokesperson for the FOP Community and the mission of the IFOPA. Ashley’s many speaking engagements have included traveling the country speaking at various FOP family fundraisers, local rotary clubs and her most recent speech at Starr’s Mill High School Baccalaureate. Over the years, Ashley has been instrumental in promoting FOP awareness by the many press and media outreach she has done including: ABC News 20/20, Mystery Diagnosis, BBC The Skeleton Key, YouTube, Tyra Banks Show and others. Ashley also played an important role in representing FOP as a spokesperson at events, including Capital Hills Awareness Days. She has provided support in any way possible; assisting with the Teen/Young Adult Meeting, Symposia, hosting fundraising events, including her most recent American 5K Run/Walk event, supporting International IFOPA Awareness Day and many other IFOPA sponsored events. Ashley, we have seen you grow up from a child to an amazing talented woman. We are very proud of you and want thank you for all that you do for the FOP Community.

Outstanding International Leadership Award

**Roger zum Felde**

Roger’s extraordinary leadership as the FOP e.V. – CEO of FOP Germany, the first international FOP organization, and his guidance as the IFOPA International President’s Council representative for Germany exemplifies the amazing community network that we all represent. His steadfast commitment to promote FOP awareness through personal interviews with the Press and other media such as YouTube videos and presentations at service clubs, such as Rotary and Mercedes-Benz Auto, is a remarkable testimony of his pledge to make a difference. Roger’s wonderful personality shines brightly and provides an inspiration to others. Roger’s tremendous support in joining our team effort in fundraising for FOP research and the IFOPA brings us one step closer to our dream of a treatment and a cure for FOP.
2011 was a very active year in terms of international FOP meetings. We are pleased to report that meetings were held in Argentina, Germany, Italy and the Netherlands. A brief summary of each meeting is provided below in chronological order; for more detail, including pictures for each event, please see the June 2012 online issue of FOP Connection.

Italy

The 5th Italian FOP conference was held in Rome on March 24-26 2011, an event held annually since FOP Italia was founded in 2006. The conference is organized under the guidance of Mr. Enrico Cristoforetti, President of the Association, and with the medical/scientific support of Genova University. Held over three days, the conference provided both a research and clinical component. Regarding research, the conference provided a general update on research results, treatment guidelines, and open discussions among the doctors, experts and audience. On the clinical side, Italian patients (of whom there are 25 in Italy) benefited from the attendance of a pool of American, German and Italian doctors and other experts in clinical research.

Argentina

The Second FOP Latin America meeting was held May 12th-15th in Buenos Aires, Argentina and organized by Fundación FOP with the support of the International FOP Association. With a focus on FOP families, the meeting was attended by 21 families from Brazil, Chile, Venezuela, Colombia, Argentina and the United States. The meeting was organized around family workshops which enabled families to increase their knowledge of FOP, exchange experiences, and meet with specialists and doctors. Physicians attending included Drs. Fred Kaplan and Robert Pignolo from the United States, Drs. Carmen de Cunto and Patricia Delai from Sao Paulo, Brazil, and Dr. Gabriela Scagnet and her team of dental specialists from the Universidad de Buenos Aires. Also attending were doctors with FOP patients and those interested in the pathology. FOP patients had the opportunity for clinical consultations with Drs. Kaplan, Pignolo, De Cunto and Delai, and odontological consultations with Dr Scagnet. Each
family in attendance also received the Family Guide printed in Spanish, the Medical Binder, an Emergency Card and a catalog of tools developed by Fundación FOP.

**Germany**

The German FOP group met on July 29-31, 2011 in the town of Meinerzhagen in the North-Rhine-Westphalia district of Germany. With 59 members attending, the meeting began socially with the introduction of new members, followed by reports from Petra Sommer, the group’s chairperson, and Roger zum Felde. Events reported on included fund-raising activities, TV and newspaper appearances, recognition by the Bavarian ministry of Petra Sommer’s extraordinary work, and an announcement of Roger’s Outstanding International Leadership Award from the IFOPA. On the following day, Petra Seeman of the Berlin-Brandenburg Center for Regenerative Therapies spoke about research on FOP. This was followed by a presentation from Professor Lutz Schomburg on personalized medicine and a new blood test. The afternoon of the first full day focused on FOP Germany’s plans for the coming year, as well as computer lessons for attendees. On the final day, Sunday, the group received information about various forms of personal insurance available, and as in the previous two years, a person with FOP was given the opportunity to talk about his/her life. This year, Meike Mrugallas from Switzerland shared his experiences. The meeting concluded with a review and discussion of the association’s finances.

**The Netherlands**

The Third Netherlands FOP Symposium was held on November 12-13, 2011 at the Marriott Hotel in Amsterdam. Through the generosity of donors, including the Marriott Hotel, the organizers were able to obtain subsidies from the government to organize the symposium. Research presentations were provided by scientists from Leiden University Medical Center), VU University Medical Center) and the Berlin Brandenburg Center for Regenerative Therapies. At the end of the first day, workshops regarding stress management and breathing techniques were conducted by hypnotherapist Dick Wernars. On Sunday, consultation sessions were provided and the weekend concluded with a canal cruise!
International President’s Council Activities in 2011

Our International President’s Council (IPC) is a network of volunteer representatives from 16 countries around the world. IPC members help the FOP community in their country or region by keeping FOP families informed about treatments and research, and providing family support. FOP volunteers establish local FOP associations, create local FOP family networks, and raise funds for FOP research.

Led by Moira Liljesthrom, IPC chairperson from Buenos Aires, and Chris Bedford-Gay of the UK, IPC focused in 2011 on translating many of our IFOPA documents into major world languages such as French, German, Spanish, Italian, Dutch, Swedish, Hindi and several others. This is a very large effort involving many volunteer translators, a number of whom are IPC representatives. Recruiting translators is challenging as translating can be very time-consuming. Documents may be as long as 50 or more pages which means that translating even one of these into another language is a large effort. Once translated, documents are then made available on our IFOPA website. We owe a great vote of thanks to Moira, Chris and all volunteers who invest so much time in making sure these important documents and papers about FOP can be read by as many people as possible around the world.

Another special feature of IPC involves providing our medical binders to international IFOPA members with FOP. The medical binder enables keeping good records of your medical history readily available, a key to receiving quality care for regular doctor visits and hospital stays, but especially in an emergency situation. Medical binder contents typically include emergency contact numbers, current and past prescriptions, medical history, history of flare ups, medications and supplements, as well as basic information regarding FOP for healthcare professionals who may be uninformed regarding this rare disease. In 2011, medical binders, along with flash drives and pill pods, were sent to members with FOP in 22 countries in four continents. This program is ongoing and many others will receive medical binders in 2012.
Quality of LIFE Awards 2011

The Quality of LIFE (Living Independently with Full Equality) Award assists IFOPA members who have FOP to live more independently in their daily lives. IFOPA provides a lifetime maximum of $1500 for items that improve the health, welfare and independence of eligible recipients. Qualifying items include aids for daily living such as adaptive equipment, walkers and other mobility aids, wheelchairs, specialized bicycles, reading aids, computer equipment, as well as tuition or registration for vocational training, to name a few examples. For more information on eligibility, qualifying items and how to apply, please see the Member Center on the IFOPA website. Funding for LIFE awards is provided by donors and family fundraisers who direct their donations to the LIFE Award Fund.

In 2011, nearly to $4000 was awarded to six IFOPA members with FOP. The following items were purchased to help them live independently: camera bag; two iPads; power lift chair; Mac Book computer, two cases of a prescribed nutritional drink; and repairs to a van to enable easier and safer access and egress.

Mentoring Program

The Mentoring Program was created to enable new families/individuals with FOP to have contact with someone who can acquaint them with the IFOPA and its resources and education. We are also happy to speak with existing IFOPA members to help them cope with their ongoing challenges (flare-ups, school issues, care, etc.). Each of our mentors either has FOP or is a parent or caregiver of someone with FOP.

Our activities during 2011 were varied. First, we made contact with at least three new families who came aboard as members of the IFOPA during the year. Next, with help from Teresa Caruso, the IFOPA established a Facebook page for women’s issues. Women who wish to join must be over 18 because subjects being discussed may sometimes be age sensitive. We have also provided international mentoring, however as each country and culture is unique this is an ongoing challenge. Last, as 2011 drew to a close, our mentoring program assisted in choreographing communications between Dr. Kaplan’s team at the FOP Lab and a team of doctors regarding the impending oral surgery of an FOP patient.

As the IFOPA continues to grow and we promote our Mentoring Program, we hope to identify new members who might need a Mentor. We are also seeking established members to take our training and join this program to mentor others. If you are interested, or know of a family that might appreciate contact with a mentor, please contact Nancy Sando (nasando@chartermi.net).
IFOPA Revenue Rebounds in 2011

Bolstered by an increase in the number of family and friends conducting fundraisers, the IFOPA enjoyed a solid revenue year in 2011.

A total of $893,213 was generated from all revenue sources, the largest total since 2008 and more than $200,000 above last year’s figure.

“I extend deep personal thanks to the donors who supported our cause so generously, and to those families and friends who organized and implemented fundraising events in 2011,” said Mark Gambaiana, Board Chair. “We are completely dependent upon private gift support to fund the IFOPA. We are able to fund research and provide programs and services because of those families and friends who are willing to fundraise.”

More than 30 families and friends conducted events during 2011, with half of those drawing support from over 50 donors and $10,000 in revenue.

Two family fundraisers set the pace of activity for 2011. Lincoln’s Legacy, a series of special events in honor of five-year-old Lincoln Wheelock of Des Moines, IA, generated over $95,000 last year from a winter carnival, a trike-a-thon, dinner and auction, and off road race. A “Bingo for a Cure” event in honor of five-year-old Joshua Scoble of Emmaus, PA and related events raised over $80,000 in support for the IFOPA. In July, the IFOPA received a generous gift of $104,100 from the estate of the late Kevin James and made in honor of FOP member Margherita Russo of Adelaide, South Australia. The legacy gift was the third largest gift in the history of the IFOPA.

The major revenue categories included $383,407 from IFOPA member events, $311,512 from outright and major contributions, $51,722 from direct mail appeals and $26,711 from third-party events, office fundraisers and grants.

All events and gifts are important to the success of the IFOPA.

Throughout its 23-year history, the IFOPA is truly unique among the nation’s universe of not-for-profit organizations. Since a modest beginning in 1988, when a total of $1,450 was generated during the inaugural year, family fundraising has become a major focus of the IFOPA’s annual operations. From 1988 through 2011, a remarkable $14,690,091 has been generated to support the IFOPA cause. The majority of this financial support has come from the deeply devoted families who rally together and form family fundraisers to advance the quest for treatments and a cure while providing necessary services and support to FOP patients and families.

Examples of family fundraisers over the years include bake sales and suppers, barn dances and bingo, chicken barbecues and spaghetti dinners, garage sales and silent auctions, country fairs and benefit concerts, drawings and rodeos, sales of holiday cards and embroidered quilts, 5K runs and road races, ice fishing contests and winter carnivals, golf tournaments and bowling parties, and wine tasting events and lemonade stands.

Meanwhile, the IFOPA is grateful to the many families who conduct various fundraisers within their communities and circles of influence. Year after year, these special events provide the majority of revenue to propel the IFOPA mission forward.
Focus on Fundraising

Hand Crafted Jewelry

Severine Letartre and her mother Martine founded an association in 2001 to have their Jewelry Fundraising go towards helping Dr. Kaplan's FOP research team. Their efforts not only help the IFOPA but also inform people about FOP, research and how they can give private donations. Each piece of jewelry is designed and hand crafted by Severine and her mom. Together they invite friends to come to their home in December before Christmas and May before Mother's Day to purchase the beautiful jewelry. If someone can’t attend they have a blog http://sephyr29.skyrock.com where people can see the jewelry and phone in their request. Their friends have also organized private jewelry sales and people are happy to give money for FOP research in exchange for hand crafted jewelry. Thank you Severine and Martine for using your creativity to help the FOP community!

Morgan Lawler Spaghetti Dinner Fundraiser

After learning that the younger sibling of two students attending the Gordon W. Mitchell Middle School in East Bridgewater, MA, was suffering from FOP, Stan Piltch, the Principal decided to hold a fundraiser to benefit The Morgan Fund.

The event was a Spaghetti Dinner with a silent auction and DJ for entertainment held on June 17th. In order to make the Spaghetti Dinner more eventful and attractive to community members, Principal Piltch and Mrs. Shel Stephen (a 5th grade teacher at the school) vowed to shave and cut their hair if the event was successful at reaching its goal. The goal was set high, at raising $20,000. The community reacted very positively and the Dinner had an overwhelming attendance which contributed to the fundraiser exceeding the goal! Good news for everyone, except perhaps Principal Piltch who had his head shaved and Mrs. Stephen who had her hair cut by more than 15 inches with all their students cheering them on. The smiles on Principal Piltch and Mrs. Stephen told everyone they enjoyed getting new hair styles!

Just prior to the Spaghetti Dinner an article was placed in the Boston Globe, in which hundreds of donations were sent in from all over the state of Massachusetts. On June 16th the school held a bake sale and a Hat Day, in which 1,200 FOP awareness bracelets were sold. Morgan’s brother and sister’s school had a lot of fun fundraising with the community – Thank you!
Justin Time Cure FOP Fundraiser

From July 22nd to the 24th, the “Justin time, Cure FOP” fundraiser ‘pooled’ in some cool cash for the IFOPA, in honor of Justin Henke, 11, of Middletown, DE. Teammates of Justin’s from his winter swim team (Greater Newark Boys & Girls Club), as well as from his summer team (Delaware Aquatics Riptide), kicked it into high gear swimming loads of laps for FOP, while land-loving friends and adults took many positive steps walking for FOP. Knowing that the greatest prize of all is a cure for FOP, youth participants were also motivated by marvelously donated prizes from ‘bone’afide businesses in the area. Stuffed among some of the prizes is one happy bird, Justin.

Ride for Joshua’s Future of Promises

The first annual Ride for Joshua’s Future of Promises was held on August 6th. It was a successful day and the heavy rain held off until the bikers completed their ride. A little over one hundred people registered for the ride. It was quite a sight to see the bikes lined up and getting ready to depart from Indian Trail Park in Northampton.

Despite the heavy downpour people stayed for the festivities. We had a ton of food, a basket raffle and numerous drawings; one of the prizes donated was two airline tickets to anywhere in the continental United States! Everyone had a great time and we were able to educate more people about FOP. At one point Joshua got a hold of the microphone and was thanking everyone for coming. A couple of people commented on how great it was to see Joshua and interact with him. His presence seemed to help them better understand what FOP is and how it affects the person afflicted and their family.

We hope to make future rides even more successful! A huge Thank You to both Kelly Beers and Lori Schaller for organizing this event!

Golf for a Cure in Honor of Shane Terry

Golf for a Cure in Honor of Shane Terry took place in Watertown, New York on July 23rd and the weather was beautiful in the mid-80s. This was Shane’s third golf tournament and he started the festivities with his famous saying, “Send It!” A local candy store helped golfers begin the day with sweet energy by providing goody bags for each golf cart which included not only candy but peanuts and golf tees too. 119 golfers participated in three divisions (Men, Women and Co-Ed) with first and second place winners of each division receiving a $25.00 restaurant gift card. Shane enjoyed participating in the tournament by getting around the course in a special street golf cart to play and visit with each.
team. After golfing, 150 people participated in a special dinner and Chinese Auction. It was a full day of fun and $6,000 was raised. Thank you to everyone who organized and participated in this fundraising event.

Lincoln’s Legacy Event Generates Over $50,000

A special fundraising event in Sioux City, Iowa held in honor of Lincoln Wheelock has added over $50,000 to FOP research. The event, conducted on August 19, attracted over 350 people to a dinner that featured a silent auction, drawings for special items and outright contributions. Over 150 items were contributed.

The event was organized by Lincoln’s grandmother, Marilyn Wheelock, and included a special appearance by Dr. Fred Kaplan from the University of Pennsylvania’s FOP research laboratory. Dr. Kaplan updated the audience on the progress of FOP research, including the recent scientific workshop. Dr. Kaplan also met with the medical staff at Mercy Medical Center during his visit to Sioux City.

Lincoln’s Legacy is an annual fundraising program designed to raise awareness and financial support for FOP. The major fundraiser is held every other year in Sioux City. Other events/activities held included a winter carnival, fun run, and a trike-a-thon. Since 2009, over $180,000 has been raised to support FOP research in honor of Lincoln, 5, who lives in Des Moines, IA with his parents, Lee and Trisha, and a brother, Hudson.

Erin’s Day

On Friday, September 23, 2011, the Danzer family, along with supportive friends and family, held the 3rd Annual Erin’s Day Golf Tournament, Dinner, Silent Auction, and Opportunity Drawing, at The Crossings Golf Resort in Carlsbad, California. There were 39 eager golfers, with eight being sponsored by Echo Pacific Construction, a wonderful and supportive company for the last three years. We are so thankful for them. The weather was perfect for a golf outing, 75 degrees, under partly cloudy skies. Also perfect weather for Erin to drive around in the golf cart with her Aunt Terri to take photos. Ken Danzer’s claim to fame this evening was to say, with a chuckle, that he broke his record by only losing 17 balls.

After the tournament, approximately 60 more people joined in the fun, eating hors d’oeuvres and listening to 50s music and even a bit of bluegrass, all played by a band called Too Much Fun! Dinner was wonderful. MC Don Brister, from Santa Maria, who is a former IFOPA board member and has been a dedicated friend of the FOP community for many years, started off the evening by talking about his being on the IFOPA board and about the important scientific meeting that had just recently taken place in August with Dr. Kaplan, Dr. Shore, Dr. Pignolo, and many others, in Philadelphia. Glen Eckart,
father of Cassie Eckart, a member with FOP, was also in attendance and helped in all areas, including presenting the Mission of Hope video and a really sweet power point presentation of various pictures of Erin’s Day events throughout the years. Denise Vietti from the IFOPA office worked tirelessly also to help make Erin’s Day successful.

There were approximately 150 items donated for the auction and drawing, including an afghan donated by Mildred Riewald of Santa Maria in Honor of Stephanie Snow, and an oil painting painted by the talented Cassie Eckart. One of the highlights of the evening was of Don Brister presenting a check to Erin for $1,000 from Santa Maria Kiwanis for Kids. She had a big smile on her face. At the end of the evening, Erin received a standing ovation for just being Erin. That really put a smile on her face! What a great way to end the evening!

Major Gift Received in Honor of FOP Member

The IFOPA has received a gift valued at $104,100 from the estate of the late Kevin R. James of South Australia.

The gift is the third largest single gift ever received in the 23-year history of the IFOPA and was made in honor of FOP member Margherita Russo. Ms. Russo is a Senior Paralegal for the Webster Lawyers law firm in Adelaide, South Australia.

“The IFOPA is deeply grateful to Mr. James for his foresight and generosity in providing for this extraordinary gift,” said Mark Gambaiana, board member and fund development chair. “This estate provision provides a substantial boost to our FOP research efforts at a critical moment in our quest for treatments and a cure.”

The estate gift is included in the Jeannie L. Peeper Heritage Roll of Honor, a legacy society established to recognize and honor those individuals and families who have included the IFOPA in their estate plans.
2011 Fundraising

The IFOPA would like to extend a special thank you to the many individuals who supported and were involved in the recent fundraising events. We appreciate the following fundraisers for their contributions:

- All American 5K: In Honor of Ashley Kurpiel
- Bingo for a Cure! in Honor of Joshua Scoble
- Charleston Marathon in Honor of Ashley Kurpiel
- Christmas Card Campaign in Honor of Yorick Blonk
- Coins for a Cure in Honor of Matthew Wadd
- Disney Princess Half Marathon in Honor of Wayne Gopshes
- Emily Collins Fundraiser at Nicholas School
- Erin’s Day in Honor of Erin Danzer
- Francis Marion Dirt Dash Half Marathon & 5K in Honor of Ashley Kurpiel
- “Friends of Oliver” in Honor of Oliver Bedford-Gay
- FOP Awareness Week at Woodstock Academy in Honor of Jasmin Ariel Floyd
- Golf for a Cure in Honor of Shane Terry
- Hand Crafted Jewelry Fundraiser in Honor of Severine Letartre
- Hat Days in Honor of Morgan Lawler
- Harford Chapter No. 83, Order of the Eastern Star in Honor of Nathaniel Padilla
- Hofstra University Students Fundraiser to Fight FOP in Honor of Laura Rossano
- IFOPA Bid to a Cure
- Jeans and Green Day: In Honor of Morgan Lawler
- Justin Time - Cure FOP in Honor of Justin Henke
- Keystone Candle Fundraiser in Honor of Joshua Scoble
- Kros Trivia Night in Honor of Lucas Whitmore
- “Leap for Lincoln” Winter Carnival in Honor of Lincoln Wheelock
- Lincoln’s Legacy in Honor of Lincoln Wheelock
- Luciana’s Walk in Honor of Luciana Wulkan
- Racing for a Cure in Honor of Joshua Scoble
- Ride for Joshua’s Future of Promises in Honor of Joshua Scoble
- Ring Toss & Hat Booth, JNES Spring Carnival in Honor of Dilyn Martin
- Scandinavian FOP Association Collection Project to Fund FOP Research
- Spaghetti Dinner & Auction in Honor of Morgan Lawler
- Strike Out FOP in Honor of Lucas Whitmore
- Tav-on-the-Ave: In Honor of Lincoln Wheelock
- Teamwork Makes the Dreamwork in Honor of Damian Fredericks
- Tenth Annual Comedy Show in Memory of BJ Giannone
- Trike-4-Lincoln: In Honor of Lincoln Wheelock
- Unmasking the Mystery of FOP II in Honor of Cody Dennings
- Valentine Card Fundraiser
- 3rd Annual Morgan’s Ride in Honor of Morgan Lawler
- 5th Annual FOP Garage Sale & BBQ in Honor of Dilyn Martin
The IFOPA gratefully acknowledges the generous contributions from friends, families, fundraisers and special events, corporations and foundations that make our programs possible.

The IFOPA enjoyed a solid fiscal year in 2011, with revenues of $850,999 and expenses of $694,641, strengthening the organization’s net asset position by $156,358. As of December 31, 2011, the IFOPA’s net assets stood at $1,578,868.

In 2011, 54 percent of the organization’s revenue came from outright contributions, and 44 percent from net proceeds of special fundraising events. Revenues increased by $246,525 from the previous year. On the expense side, program services accounted for 76 percent of expenditures, including 67 percent for medical research, the highest priority of the organization.

**REVENUE AND PUBLIC SUPPORT**

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<td>Contributions</td>
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<td>Special events, net</td>
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<td>Other</td>
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<td><strong>Total Revenue and Public Support</strong></td>
<td><strong>$850,999</strong></td>
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**EXPENSES**

**Program Services:**

- Medical research: $462,299
- Public awareness: $20,333
- Education and support: $44,963

**Total Program Services:** $527,595

**Support Services:**

- Management and general: $115,538
- Fundraising: $51,508

**Total Support Services:** $167,046

**TOTAL EXPENSES:** $694,641

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<tr>
<td>Net Assets at Dec. 31, 2011</td>
<td><strong>$1,578,868</strong></td>
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</table>
Thank you to the many wonderful families, individuals, organizations, corporations and foundations that supported the IFOPA with generous contributions between January 1, 2011 through December 31, 2011.

**Diamond**

$25,000 and above

- Estate of Kevin Ronald James
- Joshua’s Future of Promise, Inc.
- Ms. Severine Letartre
- Ramex AS

**Platinum**

$10,000-$24,999

- Bridgewater Associates, LP
- Gretchen & John Emmerich
- FOP e.V.
- The Morgan Fund
- Diane N Weiss

**Gold**

$5,000-$9,999

- Erling & Saskia Blonk
- John & Amanda Cali
- Echo Pacific Construction, Inc.
- Mark & Robin Gambaiana
- Herman Goldman Foundation
- Johnson & Johnson Family of Companies
- Lincoln’s Legacy
- Larry & Peggy Nilsen
- Peachtree City Running Club, Inc.
- Skelmir, LLC
- Kristina Sligh & Charles (Skip) Sligh, V
- Strike Out FOP - Lucas Whitmore Benefit
- The Law Offices of Walter W. Whelan
- The Robert J Kahn Foundation
- Ed & Gail Weakland

**Bronze**

$1,000-$2,499

- Adcraft Printwear Company
- AirTran Airways, Inc.
- Anonymous
- Archer Daniels Midland Company
- Ayco Charitable Foundation
- Banco Del Litoral S.A.
- John & Margaret Bannon
- John Borsa
- C.H. Patrick & Co., Inc.
- Michael Calderaro
- Craig & Stephanie Clay
- Clear Creek Independent School District
- CNOS
- Jerome & Sherry Cohen
- Douglas & Sharon Coleman
George Powell  
Pat & Justine Powell  
Power Pool Plus, Inc.  
Deree Reagan  
Recreation Equipment, Inc. (REI)  
Shailaja Reddy & Mike Oswald  
Barry & Gayle Reitman  
Cameron Remmer  
Robin & Gloria Riley  
Ramona & Rex Rock, Sr.  
Roll Giving & Paramount Community Giving  
Santa Maria Kiwanis for Kids  
Robert & Connie Scheigert  
Scientific Workshop Registration  
Elizabeth Sibray & Rebecca Anderson  
Sioux City Bandits  
David Sipple  
Greg & Kathy Smith  
Stan Lawrence Portrait Art Photography  
Tav on the Ave  
Templin Photography  
The Cove Point Foundation - Susan F. Morris  
Michael Thornton  
TMAG  
Todd Construction Management, LLC  
Tulsa Community Foundation  
United Crane Rentals, Inc.  
Unmasking the Mystery FOP  
Greg & Lynn VanBallegooyen  
Wagner, Kuntz & Grabouski, PC  
Jim & Beverly Wharton  
Trisha & Lee Wheelock  
Scott & Lisa Williams  
Dick & Judy Yamane  

Lisa & Jerrold Danzer  
Lori & Kenneth Danzer  
Dave Gryp Construction, Inc.  
William & Nancy Davis  
Thomas & Kathy Dawson  
Dishman Sports Group, LLC  
Doctors Express Mgmt of Central Texas LLC  
Mr. Patrick Doerr  
George & Diane Drechsler  
John & Tracy Eck  
Elite Illumination  
Essential Fitness, LLC  
Raoul & Mona Farid  
Christine & James Ford  
Marilee K. Fox-Cichon  
Mark Freitas  
Fulton Elementary  
Funding Factory  
Gary & Deborah Gayton  
GE Foundation  
Andrea Gherini  
Gloucester Terminals/Holt Logistics Corp  
Amy & Raymond Gopshes  
Lisa & Michael Gragnani  
Garret & Deborah Green  
Greenman, Lacy, Klein, O’Harra & Heffron  
Greenstar  
Dason Gwartney  
Nancy Hair  
Greg & Ivy Hamerly  
Lee & Patti Harbin  
Harford Chapter No. 83 OES  
Kevin & Wendy Henke  
Jean-Pierre Henrotay  
Robert Hilongos  
Hogue & Associates  
Keith Houser  
Susan J. Hwang  
I.L.A. Local No 953  
Impact Learning, Inc.  
International Longshoremens’ Assoc Local 1235  
Cindy & Roger Johnson  
JP Morgan Chase Foundation  
Pierre Jraihe  
Daniel Kaler, D.D.S., P.C.  
Dr. Fred Kaplan  
Kelly & Associates Insurance Group, Inc.  
Kathy Kennedy  
James & Fran Kennedy  
Dr. & Mrs. Joseph A. Kitterman  
Linda & Reed Kneebone  
David & Teresa Korol  
Kros Broadcasting, Inc.  
Kysar Chili Cook-Off  
Pamela Laff  

Nate Landendorf  
Katherine Lane  
Julia Lange  
Moira Liljestrom & Frederico Robert  
Rachel Lu  
Geoffrey & Janice McCue  
Margaret & Curtis McWilliams  
McWilliams Farm  
Metro Pilots  
Linda Miller & Timothy Duperron  
Kevin M. Mowry  
Loren & Connie Mulder  
Nakashima  
Lance & Deborah Noe  
Odessa Ofstad  
Doug Olson  
Terry & Karen Olson  
Open Wide Family Dentistry  
P & T Supply & Services, Inc.  
Panera Bread  
Peggy Phillips  
Harry & Beverly Powazek  
PRP Wine International, Inc.  
Charlie & Daphne Purss  
William & Laura Quackenbush  
Lynn E. Refer  
Emily Rocke  
San Diego County Judges Association  
Mark Sargent & Catherine Douglas  
Kathleen & Thomas Schmidt  
Rob & Angelka Scofidio  
Robert & Karen Scott  
Seymour Ford Lincoln  
Bernie & Jerry Shapiro  
Art & Eileen Silva  
Siouxland Golf Charities, Inc.  
Bob & Jennifer Snow  
Philip Spinelli  
Starr’s Mill High School - Class of 2011  
Wendy & Mark Stevenson  
Susquehanna Post 135 American Legion  
The Zartarian Foundation  
Andrea Thrasher  
Dan & Kerry Tomson  
Total Wine & More  
Donald & Donna P. Truby  
Truman Heartland Community Foundation  
Union Bible Study Association, Inc.  
United Way of the Greater Lehigh Valley, Inc.  
W. L. Gore & Assoc., Inc.  
David & Gloria Wadd  
William & Hillary Weldon  
Thomas & Jody Wente  
Doug & Marilyn Wheelock  
Adam & Gretchen Wheelock  
Patrick & Barb Williams

Copper

$500-$999

Addi’s Drawing from the Heart  
Alpeninox UK  
American Maritime Services of NY, Inc.  
BBC Productions Worldwide Productions  
Marvin L Bellin  
Don Brister & Diane Bennett  
David Burack  
Capital One Services, LLC  
Carlsbad High School  
Marie M Carter  
Chippewa Hills School District  
David & Diane Connaughton  
Cut Rate Auto Purchasing Corp.  

Lisa & Jerrold Danzer  
Lori & Kenneth Danzer  
Dave Gryp Construction, Inc.  
William & Nancy Davis  
Thomas & Kathy Dawson  
Dishman Sports Group, LLC  
Doctors Express Mgmt of Central Texas LLC  
Mr. Patrick Doerr  
George & Diane Drechsler  
John & Tracy Eck  
 Elite Illumination  
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Raoul & Mona Farid  
Christine & James Ford  
Marilee K. Fox-Cichon  
Mark Freitas  
Fulton Elementary  
Funding Factory  
Gary & Deborah Gayton  
GE Foundation  
Andrea Gherini  
Gloucester Terminals/Holt Logistics Corp  
Amy & Raymond Gopshes  
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Garret & Deborah Green  
Greenman, Lacy, Klein, O’Harra & Heffron  
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Dason Gwartney  
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Kysar Chili Cook-Off  
Pamela Laff
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Countryside Golf Course
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Jorge & Anna Maria Matarredona
James Maxymillian
RoseMary Mazur
Hope & Dick McBride
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Nichole Weston
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Barry & Sarah Wheelock
Betty Wheelock
Whitewater Challengers Inc.
Calvin & Phyllis Whitmore
Jeff Whitmore
Kris M Whittington
Kathryn Wiegand
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Mary Bartholomew
Deborah & Philip Bastable
Lawrence & Linda Batton
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Christopher Bauman
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Linda & Michael Condrillo
Arthur Connolly, Jr & Joan Connolly
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Kirk & Angela Cox
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Brad & Kelly Crocker
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Jeffrey & Elizabeth Cronston
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