PRF By The Numbers







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Produced by Leslie B. Gordon, MD, PhD; Medical Director

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PRF By The Numbers: A Data Sharing Tool

PRF By The Numbers is a data sharing tool originating from The Progeria Research Foundation's programs and services.

We translate information collected within our programs and services, and develop charts and graphs which track our progress from year to year.

This allows you to assess where we've been, and the improvements we've made for children with Progeria.



Why Sharing Data Is Essential

According to the National Institutes of Health: "data sharing is essential for expedited translation of research results into knowledge, products, and procedures to improve human health."

http://grants.nih.gov/grants/guide/notice-files/NOT-OD-03-032.html

In other words, everyone benefits by knowing and learning as much as possible about Progeria - the scientific and medical communities, the public, and the children.





PRF By The Numbers...Here's How It Works

- We take raw data collected through our programs and services, remove any personal information to protect the participant, and present it to you in a format that is engaging and informative.
- PRF programs and services include:

The PRF International Registry

The PRF Diagnostics Program

The PRF Cell & Tissue Bank

The PRF Medical & Research Database

PRF Research Grants

🧗 Scientific Workshops

Clinical Trial Funding and Participation



Our Target Audience

> PRF By The Numbers is intended for a broad array of users



Families and children with Progeria



The general public and nonscientists of all ages



Scientists



Physicians



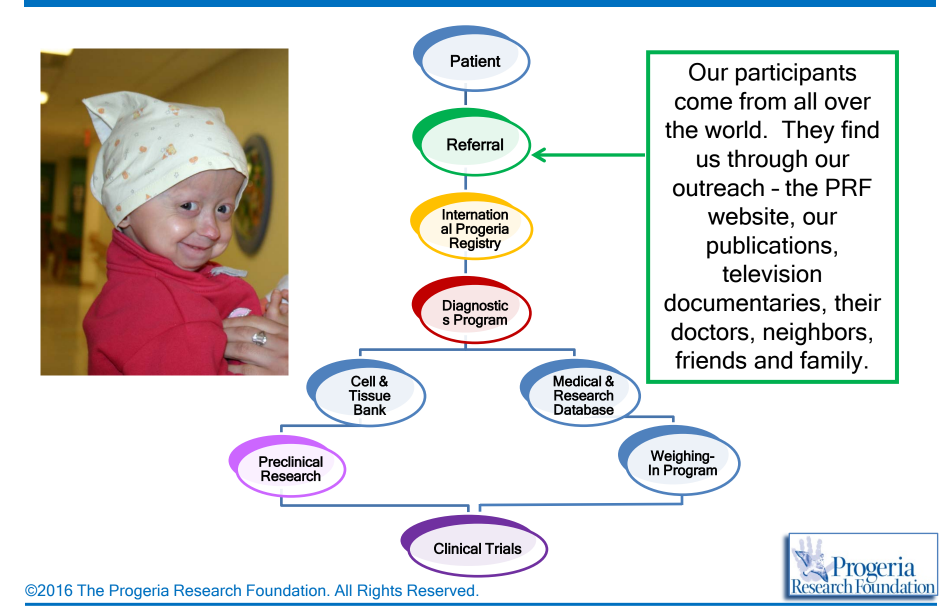
The media

- ➤ This means that different types of slides will be of interest depending on who is looking at the information. We have designed this slide set so that you can pull out what is most important to you.
- We love suggestions if you don't see some facts and figures here that you think would be informative, please let us know at

info@progeriaresearch.org



PRF Programs: It All Starts With The Children



Program Collaborations For Success



PRF Cell & Tissue Bank Core Laboratory



PRF Medical & Research Database PRF Cell & Tissue Bank PRF Diagnostics Program



PRF Cell & Tissue Bank : Lymphoblast Cell Line Generation

PRF Diagnostics Program Sequencing Laboratory

PREVENTION GENETICS

DISEASE PREVENTION THROUGH GENETIC TESTING

Progeria Research Foundation

PRF Clinical Trials





PRF Cell Bank Submission: Immortalized Fibroblast Cell Lines



Non-HGPS Progeroid Patient Diagnosis



PRF Cell & Tissue Bank : iPS Cell Line Generation



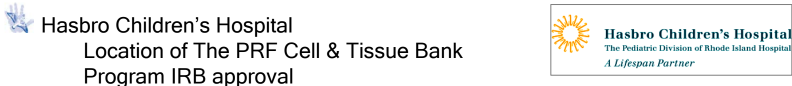
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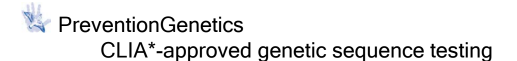
Our Program Collaborators

Our collaborating institutions are crucial to our ability to help children with Progeria. We are extremely grateful for these ongoing partnerships:



Brown University Location of The PRF Medical & Research Database Program IRB approval







BROWN

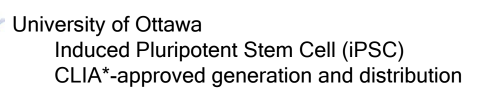
BROWN

Alpert Medical School

School of Public Health

Rutgers University Cell and DNA Repository
CLIA*-approved lymphoblast generation and distribution







Our Clinical Trial Collaborators

Our collaborating institutions are crucial to our ability to help children with Progeria



Harvard University - Associated Hospitals:

Boston Children's Hospital
Brigham and Women's Hospital
Dana Farber Cancer Institute









NIH - funded Clinical and Translational Study Unit at Boston Children's Hospital





Number of Living PRF-Identified Cases

As of July 1, 2016:

Total Number of Children with Progeria Worldwide:

134

HGPS* worldwide:



HGPS* in the United States:



Progeroid Laminopathies** worldwide:



Progeroid Laminopathies** in the United States:



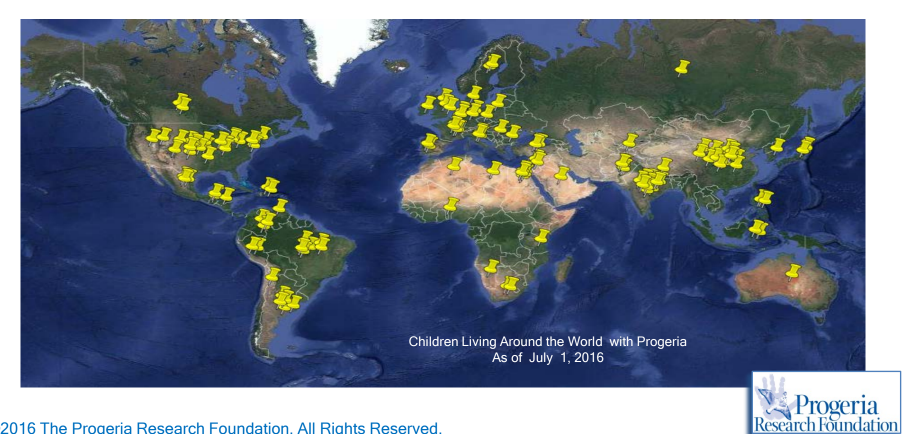
*Children in the HGPS category have a progerin-producing mutation in the LMNA gene

^{**} Those in the Progeroid Laminopathy category have a mutation in the lamin pathway but don't produce progerin



PRF-Identified Cases Reside In 45 Countries

Algeria	Canada	Denmark	Germany	Ireland	Mexico	Philippines	Serbia	Tanzania
Argentina	Chile	Dominican Republic	Guatemala	Israel	Namibia	Poland	South Africa	Togo
Australia	China	Egypt	Honduras	Italy	Nepal	Portugal	South Korea	Turkey
Belgium	Colombia	England	India	Japan	Pakistan	Russia	Sweden	USA
Brazil	Czech Republic	France	Indonesia	Libya	Peru	Saudi Arabia	Tajikistan	Venezuela



...and Speak 30 Languages

Arabic	English	Indonesian	Marathi	Spanish	Tamil
Chinese	French	Italian	Polish	Swahili	Telugu
Czech	German	Japanese	Portuguese	Swedish	Turkish
Danish	Hebrew	Kannada	Russian	Tagalog	Urdu
Dutch	Hindi	Korean	Serbian	Tajik	Uzbek

прогерии исследовательский фонд

مؤسسة أبحاث الشياخ

早衰症研究基金會

Progeria रिसरच फाउंडेशन



조로증 연구 재단

Progeria Araştırma Vakfı

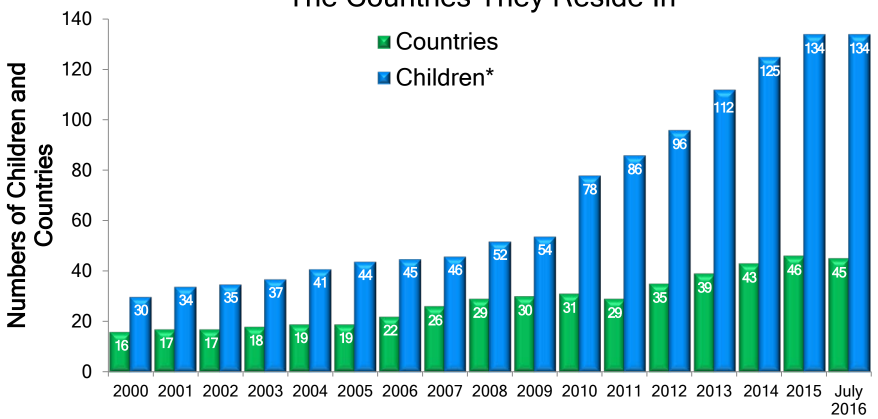
早老症研究財団

బాలుడ బాలిక వయస్స ముదరుకండానే వృద్ధాప్యరూపంలోనికి వచ్చుట రీసెర్చ్ ఫ్రాండేషన్



Every Year Our Numbers Grow

Living Children PRF Has Identified with Progeria and The Countries They Reside In*



Year

*When a child passes away, numbers are decreased.

Numbers include those with HGPS and genetically confirmed Progeroid Laminopathies



Tracking Children with Progeria Through Prevalence

➤ How does PRF estimate how many children we are searching for, and in what countries? We use *population prevalence*.

Prevalence is the proportion of children with Progeria per total population.



How Prevalence Is Estimated

- ➤ At PRF, we use a formula based on the number of children we've identified in the US. We then expand that out to the world population.
- ➤ We do this because we have the most complete reporting for the US and since Progeria has no gender, ethnic, or other biases, we assume that the prevalence in the US is the same prevalence in other countries.
- ➤ PRF estimates prevalence for years when the official US census provides a reliable population number.



USA Prevalence of Progeria

January 2016 population statistics:



👺 The US population was:

322,761,807 people



Number of PRF-identified children with Progeria in the US:





Prevalence of HGPS in the US: 16 in 323 million is about

1 in 20 million people





Source: 2016 US population: http://www.census.gov/#

Prevalence and World Population of Progeria

Given the world population as of January 2016

there are between 350 and 400 children living with Progeria worldwide.



PRF strives to find every child with Progeria because in order to help every child, we must find every child



Using Prevalence To Find Children In A Certain Country

We can now use the total population estimates for any given country, in order to understand whether we have found most or all children in a particular country.

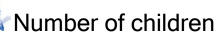
For example, as of January 2016:



👺 Brazil's population was estimated as



205,809,000 people



👺 Number of children living with Progeria in Brazil is







International Progeria Registry*

Program Goals:

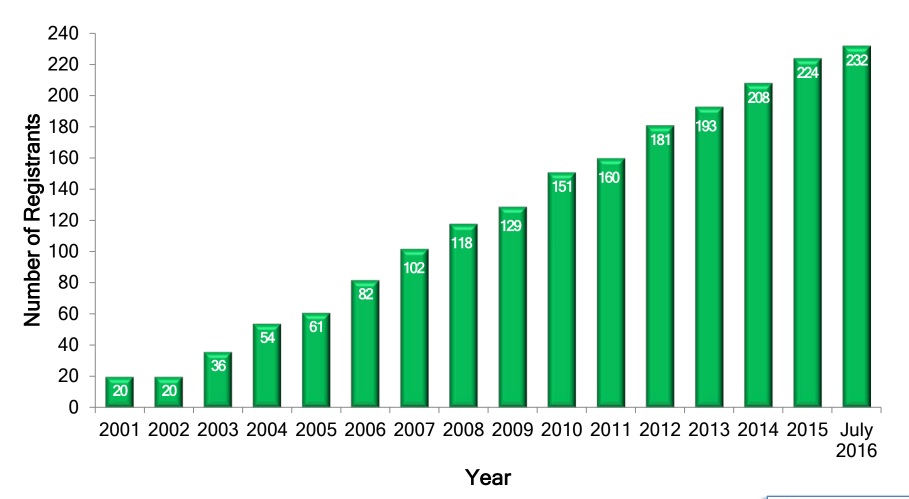
- > Patient identification
- > Outreach to patient families and their physicians
- > A springboard for program enrollment

Registry forms available at www.progeriaresearch.org/patient-registry

*PRF International Registry includes those with genetically confirmed or clinically suspected Progeria, as well as those with other possible progeroid syndromes



226 Children Have Registered With PRF





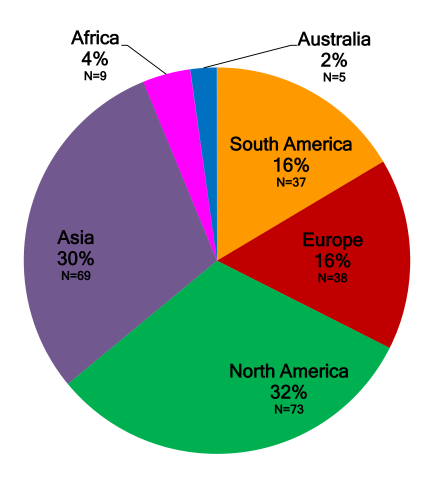
...From 55 Countries

Algeria	Canada	Dominican Republic	Honduras	Ireland	Morocco	Poland	Senegal	Switzerland
Argentina	Chile	England	Hong Kong	Israel	Netherlands	Portugal	South Africa	Tanzania
Australia	China	Finland	India	Italy	Pakistan	Puerto Rico	South Korea	Togo
Belgium	Colombia	France	Indonesia	Japan	Panama	Romania	Spain	Turkey
Brazil	Czech Republic	Germany	Iran	Libya	Peru	Russia	Sri Lanka	USA
Bulgaria	Denmark	Guatemala	Iraq	Mexico	Philippines	Saudi Arabia	Sweden	Venezuela
								Vietnam



...And All Continents

Participation (%) By Continent





PRF Diagnostics Program

Program Goal:

➤ Genetic Sequence Testing for Progeria-causing mutations

Pre-requisites for Testing:

- ➤ Registration with PRF International Registry
- One or more of the following



👺 Family history - proband, prenatal



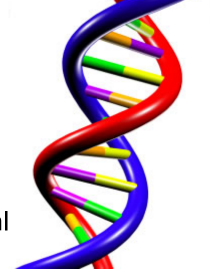
👺 Phenotypic presentation - proband, postnatal



Relative of positive proband



www.progeriaresearch.org/diagnostic testing





Diagnostics Testing Summary

As of July 1, 2016:

Total Number of Proband Tests Performed:



Exon 11 (HGPS) Mutations:



Other Progeroid Laminopathies (Exons 1 - 12):



Zmpste24 Mutations:



Average Number of Patients Tested Per Year:



All tests are performed in a Clinical Laboratory Improvement Amendments (CLIA) certified facility.



Mutations Identified Through PRF Diagnostics Program

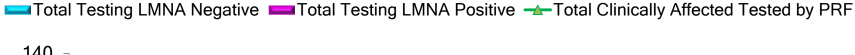
DNA Mutation	Amino Acid Effect	Zygosity	Progerin Producing?	Number Diagnosed				
	Classic HGPS - LMNA Mutation							
1824 C>T, exon 11	G608G	heterozygous	Yes	72				
	Non Classic HG	PS-LMNA Mutation						
1822 G>A, exon 11	G608S	heterozygous	Yes	4				
1821 G>A, exon 11	V607V	heterozygous	Yes	2				
1868 C>G, exon 11	T623S	heterozygous	Yes	1				
1968+5 G>C, intron 11		heterozygous	Yes	2				
1968+1 G>C, intron 11		heterozygous	Yes	2				
1968+2 T>A, intron 11		heterozygous	Yes	1				
1968+1 G>A, intron 11		heterozygous	Yes	1				
	Progeroid Laminopathy- LMNA Mutation							
1579 C>T, exon 9	A527C	heterozygous	No	1				
1579 C>T, exon 9	A527C	homozygous	No	6				
1580G>T, exon9	A527L	Homozygous	No	1				
1619 T>C, exon 10	M540T	homozygous	No	1				
331 G>A, exon 1	G111L	heterozygous	No	1				
	Progeroid Laminopa	athy-Zmpste24 Mutatio	n					
1274T>C, exon 10	L425P	homozygous	No	2				

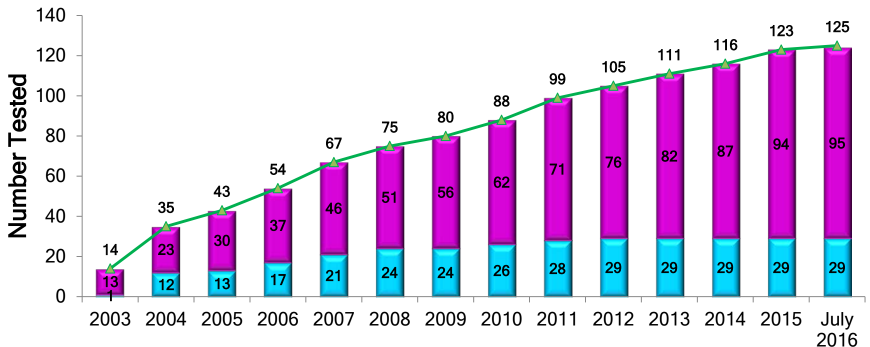
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As of July 1, 2016

Longitudinal Testing Data for PRF Diagnostics Program

Number of Affected Children/Adults Tested and the Number Testing Positive for *LMNA* Gene Mutation*





*Graph does not include Parents/Siblings tested

Year



PRF Cell & Tissue Bank

Program Goals:

- Provide a resource for researchers worldwide
- ➤ Ensure the sufficient availability of genetic and biological materials essential for research aimed at understanding the pathophysiology of disease and the links between Progeria, aging and heart disease
- Obtain long-term clinical data



Resource information available at: www.progeriaresearch.org/cell tissue bank



PRF Cell & Tissue Bank Holdings

As of July 1, 2016:

Total Number of Cell Lines:



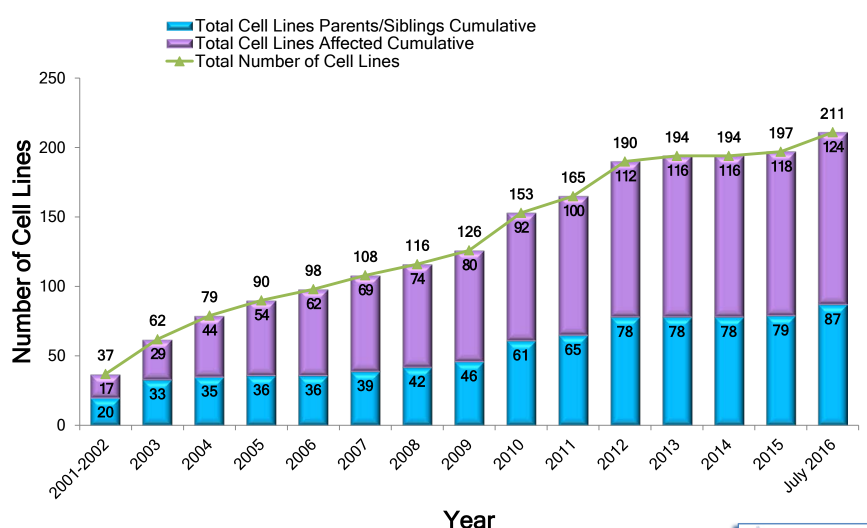
- Dermal Fibroblast Lines from 44 affected, 21 parents and 0 siblings
- 13 Immortalized Fibroblast Cell Lines from 2 affected and 8 parents
- Lymphoblast Lines from 67 affected, 45 parents and 8 siblings
- 10 Induced Pluripotent Stem Cell Lines from 2 affected and 2 parents

Mutations Available in PRF Cell & Tissue Bank

DNA Mutation	Amino Acid Effect			Cell Type DFN=Dermal Fibroblast LBV= Lymphoblast		
	Classic HGPS	S - LMNA Mutation				
c.1824 C>T, exon 11	p.G608G	heterozygous	Yes	DFN, LBV, iPSC		
	Non Classic HG	PS- LMNA Mutation				
c.1822 G>A, exon 11	p.G608S	heterozygous	Yes	DFN, LBV		
c.1821 G>A, exon 11	p.V607V	heterozygous	Yes	DFN		
c.1868 C>G, exon 11	p.T623S	heterozygous	Yes	LBV		
c.1762 T>C, exon 11	p.C588R	heterozygous	No	DFN		
c.1968+5 G>C, intron 11		heterozygous	Yes	DFN		
c.1968+1 G>A, intron 11		heterozygous	Yes	LBV		
c.1968+2 T>C		heterozygous	Yes	DFN		
c.973 G>A, exon 6	p.A325A	heterozygous	No	DFN		
	Progeroid Laminopathy- LMNA Mutation					
c.1579 C>T, exon 9	p.A527C	heterozygous	No	LBV		
c.1579 C>T, exon 9	p.A527C	homozygous	No	LBV		
c.1580 C>T, exon 9	p.A527L	Homozygous	No	LBV		
c.1619 T>C, exon 10	p.M540T	homozygous	No	DFN		
c.331 G>A, exon 1	p.G111L	heterozygous	No	DFN, LBV		
Progeroid Laminopathy- Zmpste24 Mutation						
c.1274 T>C, exon 10	p.L425P	homozygous	No	DFN, LBV		
c.743 C>T, exon 6 &	p.P248L	heterozygous	No	DFN		
c.1349 G>A, exon 10	p.T450S	, , , , , , , , , , , , , , , , , , , ,				



Number Of Cell Lines By Year





PRF Cell & Tissue Bank Distribution

As of July 1, 2016:



Research Teams From



Countries Have Received

678 Cell Lines

103 DNA Samples

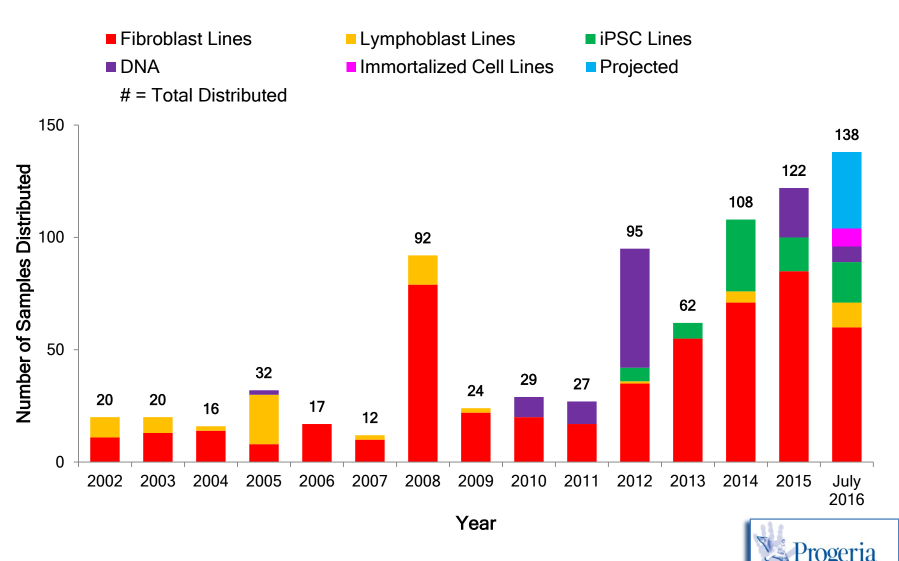
34 Types of Tissues



Senescent Progeria Fibroblasts in Culture



Biological Sample Distribution Over Time



USA Cell & Tissue Bank Recipients



Recipient	Institution
Angelika Amon	Massachusetts Institute of Technology
Stelios Andreadis	U. of Buffalo
Shelley Berger	U of Pennsylvania
Bruce Blazer	U. of Minnesota
Ted Brown	Institute for Basic Research (IBR)
Judy Campisi	Buck Institute
Kan Cao	U. of Maryland
Francis Collins	National Genome Research Institute
Lucio Comai	U. of Southern California
John Cooke	Houston Methodist Research Institute
Mauro Costa-Mattioli	Baylor College of Medicine
Adrienne Cox	U. of North Carolina at Chapel Hill
Greg Crawford	Duke University Medical Center
Antonei Csoka	Howard University
Kris Dahl	Carnegie Mellon University
George Daley	Boston Children's Hospital
Channing Der	U. of North Carolina at Chapel Hill
Mohanish Deshmukh	U. of North Carolina at Chapel Hill
Dennis Discher	U. of Pennsylvania

Recipient	Institution			
Martin Dorf	Harvard Medical School			
Stephen Doxsey	U. of Massachusetts Medical School			
Jack Elias	Brown University School of Medicine			
Mike Erdos	National Institutes of Health			
Jed Fahey	Johns Hopkins University			
Shridar Ganesan	Cancer Institute of New Jersey			
Abhimanyu Garg	U. of Texas Southwestern Medical Center			
Thomas Glover	U.of Michigan Medical School			
David Gilbert	Florida State University			
Robert Goldman	Northwestern University			
Susana Gonzalo	St. Louis School of Medicine			
Lilian Grigorian	Cedars Sinai Medical Center			
Curtis Harris	National Institutes of Health			
Martin Hetzer	Salk Institute			
Steve Horvath	UCLA			
Vishwanath lyer	U. of Texas Austin			
Jose Jalife	University of Michigan			
David Kaplan	Tufts University			
Karen Katula	UNC - Greensboro Progeria			
As of Jul				

USA Cell & Tissue Bank Recipients



Recipient	Institution
Timothy Kowalik	U. of Massachusetts Medical School
Dmitri Krainc	Massachusetts General Hospital
Jan Lammerding	Harvard University
Dudley Lamming	U of Wisconsin-Madison
Jeanne Lawrence	U. of Massachusetts Medical School
Joan Lemire	Tufts University School of Medicine
Kam Leong	Columbia University
Jason Lieb	U. of North Carolina at Chapel Hill
Shigemi Matsuyama	Case Western Reserve University
Andrew Mendelsohn	Regenerative Sciences Institute
Jeffrey Miner	Washington University
Tom Misteli	National Cancer Institute
Marsha Moses	Boston Children's Hospital
Elizabeth Nabel	National Heart, Lung & Blood Institute
Timothy Osborne	Sanford Burnham Medical Research Institute
Junko Oshima	U. of Washington
Bryce Paschal	U. of Virginia
Mary Patti	Joslin Diabetes Center

Recipient	Institution
Joseph Rabinowitz	Temple University
Ana Robles	National Cancer Institute
David Sabatini	Whitehead Institute
John Sedivy	Brown University
Christian Sell	Drexel University College of Medicine
Andrew Sonis	Boston Children's Hospital
Earl Stadtman	National Heart, Lung & Blood Institute
Dylan Taatjes	U. of Colorado
Marc Tatar	Brown University
George Truskey	Duke University
Alan Waldman	University of South Carolina
Steve Warren	Emory University School of Medicine
Howard Worman	Columbia University
Tom Wight	Hope Heart Institute
Yue Zou	East Tennessee State University



International Cell & Tissue Bank Recipients



University of Pecs, Hungary

University of Cambridge, England

University of Milan, Italy

CNR Institute of Cellular Biology & Neurobiology

Kirsztian Kvell

Chiara Lanzuolo

Caterina La Porta

Delphine Larrieu

International Cell & Tissue Bank Recipients



















Recipient	Institution
Lucia Latella	National Research Council (CNR) Rome, Italy
Giovanna Lattanzi	ITOI-CNR Unit of Bologna, Italy
Jean-Marc Lemaitre	Institute of Functional Genomics, Montpellier, France
Nicolas Levy	Génétique Médicale et Développement, Faculté de Médecine de la Timone, France
Frank Lyko	German Cancer Research Institute Heidelberg, Germany
Thorston Marquart	University of Münster, Germany
Scott Maynard	Danish Cancer Society Research Institute, Denmark
Ohad Medalia	University of Zurich, Switzerland
Denis Mottet	University of Liege, Belgium
Luis Pereira de Almeida	Center for Neuroscience and Cell Biology (CNC), Portugal
Neale Ridgway	University of Halifax, Canada
Kanda Sangthongpitag	Experimental therapeutics Centre, Singapore
Ok Sarah Shin	Korea University Guro Hospital, Korea
Michael Speicher	Medical University of Graz, Austria
William Stanford	University of Toronto, Canada
Michael Walter	University of Münster, Germany
Herbert Waldman	Max Planck Institute, Germany
Miguel Weil	Tel Aviv university, Israel
Jesus Vazquez Cobos	Centro Nacional de Investigaciones Cardiovasculares, Spain
Alex Zhavoronkov	Federal Clinical Research Centre, Russia
Zhongjun Zhou	University of Hong Kong, China Proceria

PRF Medical & Research Database

Program Goals:

Collect the patient health records for living and deceased children with Progeria

- Obtain long-term clinical data
- Abstract data for longitudinal and crosssectional analyses
- Better understand the clinical disease process in Progeria and aging related diseases
- Develop treatment strategies and recommendations for health care professionals and families



How The PRF Medical & Research Database Works

- Project staff obtain the patient's medical records and film studies from birth throughout the participant's lifespan.
- Medical records include visits to: primary care physicians, specialty physicians, hospital emergency rooms, hospital admissions, dentists, physical therapy, occupational therapy and school health records.
- Retrospective data abstraction protocol allows for specifically targeted or broad spectrum of data.

Enrollment information available at: www.progeriaresearch.org/medical_database



Medical & Research Database Participation

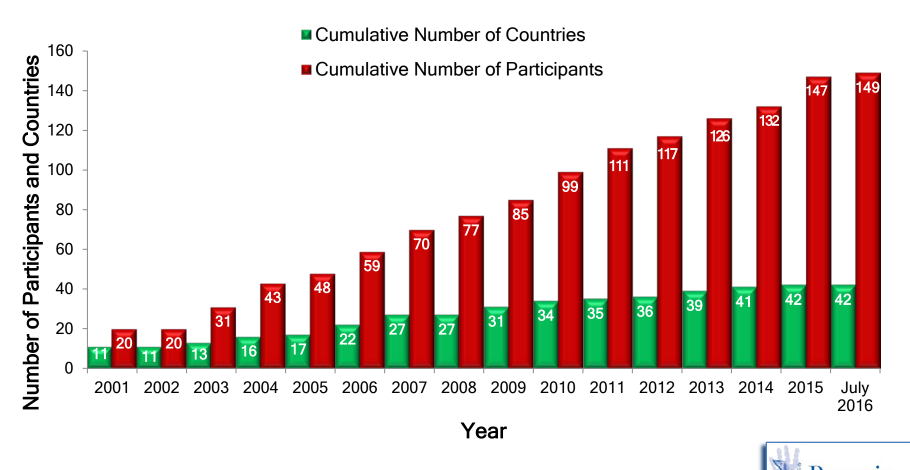
Participants are enrolled from 41 countries and 1 US territory

Chile **England** Indonesia Libya Peru Russia Tanzania Argentina Australia China France Ireland Mexico **Philippines** Senegal Togo Poland South Africa Turkey Belgium Colombia Germany Israel Morocco Denmark Honduras South Korea USA Italy Netherlands Portugal Brazil Dominican Republic Pakistan Venezuela Canada India Puerto Rico Spain Japan Romania Sweden Vietnam



Database Longitudinal Enrollment

Children Enrolled in The PRF Medical & Research Database and the Countries of Residence



Types Of Data Collected

➤ Participants with Medical Records Reports:



Participants with Radiology Studies:









PRF Weighing-In Program

- ➤ A sub-program of The PRF Medical & Research Database
- Collects weight-for-age data prospectively:



Home scale provided by PRF



Parents weigh child weekly or monthly



Report weights electronically





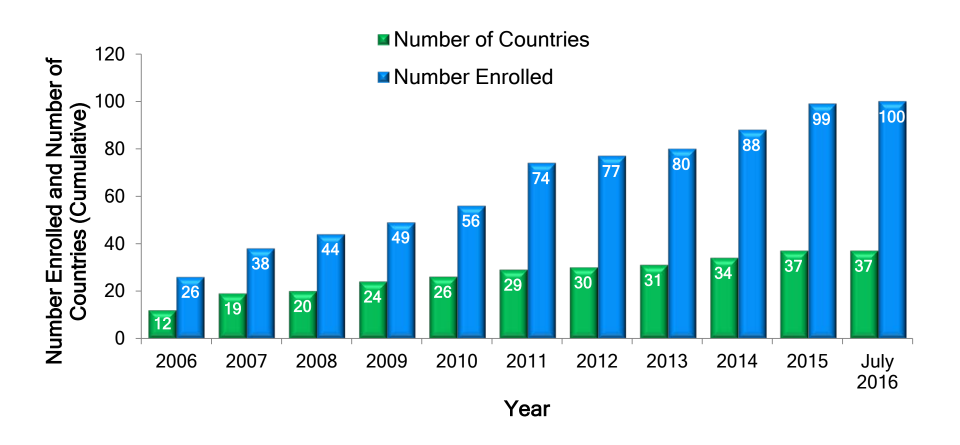
Weighing-In Program Participation

Participants are enrolled from 36 countries and 1 US territory

Puerto Rico South Korea **USA** Argentina China England Pakistan Israel Venezuela Australia Colombia Germany Italy Peru Romania Spain Tanzania Belgium Denmark Honduras Japan **Philippines** Vietnam Russia Brazil **Dominion** Mexico Poland Senegal Togo India Republic



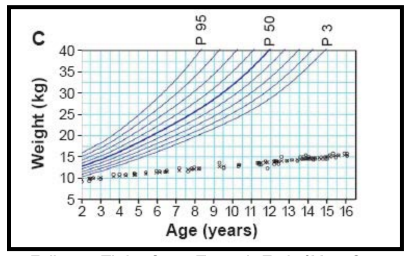
Participants Enrolled In The PRF Weighing-In Program and Countries of Residence





Clinical Trials And The Weighing-In Program

- Data from this program were key in the development of primary outcome measure for the first drug treatment trial for Progeria.
- ➤ As of July 1, 2016, **75** children from The PRF Weighing-In Program have entered clinical treatment trials using this data.







PRF-Funded Clinical Treatment Trials





Clinical Drug Treatment Trials

Goals:

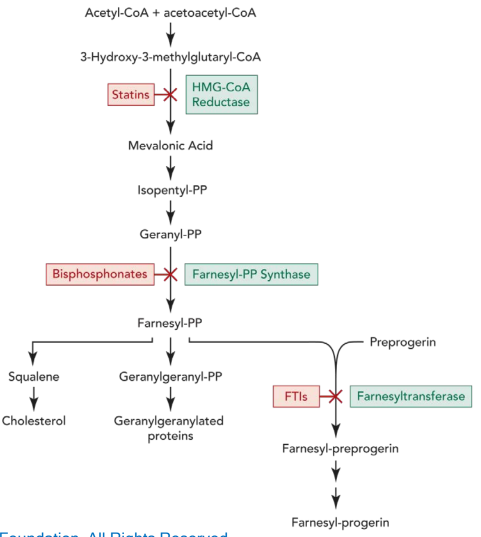
- ➤ To define the natural history of HGPS in quantifiable terms that will expand our ability to measure treatment outcome
- ➤ To assess the safety of new treatments for HGPS
- ➤ To measure effects of treatments for children with HGPS on disease status, changes in health, and survival





Current Therapeutic Intervention Strategies

Medications That Inhibit Farnesylation of Progerin





PRF Funds Clinical Treatment Trials

	Year	Drug(s)	Phase	Location	#	Countries	
	2007- 2010	Lonafarnib	2	Boston	28	17	
	2009	Lonafarnib Pravastatin Zoledronate	Feasibility	Boston	5	1	
	2009- 2013	Lonafarnib Pravastatin Zoledronate	2	Boston	45	24	
	2014- present	Lonafarnib	2	Boston		33 countries led to date	13
	O						
©2016 The Pr	ogeria Rese	earch Foundation. All	Rights Reserve	d.		K	Progeria

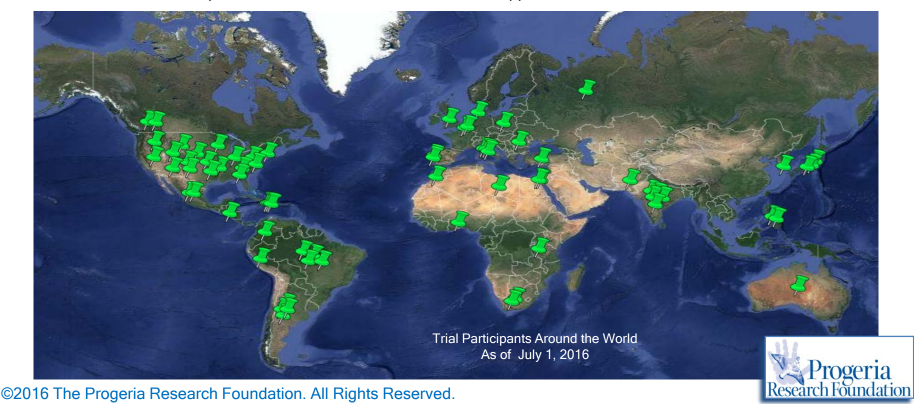
Research Foundation

Participation in PRF Clinical Trials

78

Children have participated in PRF Clinical Trials from 32 countries:

Poland Argentina Canada **England** Italy Morocco South Africa Togo South Korea Australia Colombia Honduras Japan Pakistan Portugal Turkey **USA** Belgium Denmark India Libya Peru Romania Sweden Mexico Tanzania Venezuela Brazil Dominican Republic **Philippines** Russia Israel



Treatment Trial Collaborations For Success

> The children are seen by physicians from:













Data were also generated by scientists from:

Alpert Medical School at Brown University

Brown University School of Public Health

University of California Los Angeles

National Human Genome Research Institute

Schering-Plough Research Institute

Lonafarnib generously provided by Merck













Clinical Treatment Trial Efficacy Results

Lonafarnib, a type of farnesyltransferase inhibitor (FTI) is our first treatment for Progeria.

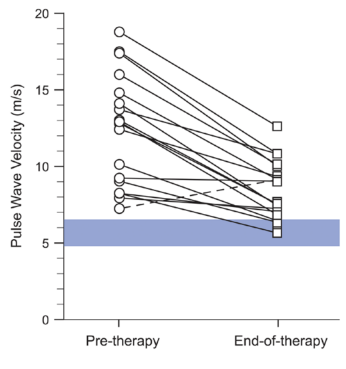
> Results showed improvement in:

Rate of weight gain

Increased vascular distensibility

Improved bone structure

Better neurosensory hearing



Gordon et al, PNAS, 2011



Clinical Treatment Trial Publications



Drug Effect:

Clinical Trial of the Protein Farnesylation Inhibitors Lonafarnib, Pravastatin, and Zoledronic Acid in Children With Hutchinson-Gilford Progeria Syndrome. Gordon et al, *Circulation*, 2016 Jul 12;134(2):114-25.

Seeking a Cure for One of the Rarest Diseases: Progeria. Collins FS. Circulation, 2016 Jul 12;134(2):126-9.

Clinical Trial of a Farnesyltransferase Inhibitor in Children with Hutchinson-Gilford Progeria Syndrome, Gordon et al, *Proceedings of the National Academy of Sciences*, 2012 Sep 24.

Neurologic Features of Hutchinson-Gilford Progeria Syndrome after Lonafarnib Treatment - *Neurology*, 2013, 81:427-430.



Dermatology:

Initial Cutaneous Manifestations of Hutchinson-Gilford Progeria Syndrome - *Pediatric Dermatology*, 2014,1-7.



X-ray:

A Prospective Study of Radiographic Manifestations in Hutchinson-Gilford Progeria Syndrome, Cleveland et al., *Pediatric Radiology*, 2012 Sep;42(9):1089-98. Epub 2012 Jul 1.



Cardiology:

Mechanisms of Premature Vascular Aging in Children with Hutchinson-Gilford Progeria Syndrome. Gerhard-Herman M, et al., *Hypertension*. 2012 Jan;59(1):92-97; Epub 2011 Nov 14.



Skeleton:

Hutchinson-Gilford progeria is a skeletal dysplasia. Gordon, et al., J Bone Miner Res. 2011 Jul;26(7):1670-9.



PRF Grants Program

Program Goals:

- Attract high level researchers to the field of Progeria
- Foster high quality publications
- Stimulate novel research that will lead to larger grants from other resources such as NIH, Ellison Foundation, and others
- Provide ability for researcher to thrive in the field
- Foster researchers of interest to PRF's mission

Grants program information available at www.progeriaresearch.org/research funding opportunities



PRF Medical Research Committee

Volunteer MRC Reviews Grant Applications Semi-annually



Back Row (L to R): Tom Misteli, PhD; Judy Campisi, PhD; Christine Harling-Berg, PhD;

Leslie Gordon, PhD, MD; Ted Brown, MD, PhD

Front Row (L to R): Frank Rothman, PhD; Tom Glover, PhD; Bryan Toole, PhD (chair)

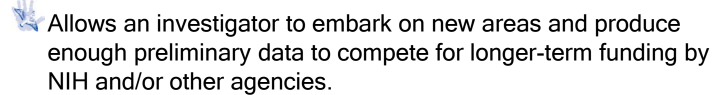
Not Pictured: Monica Kleinman, MD

PRF Granting Structure

Innovator Awards:



🌄 2 years, up to \$75,000 per year



Established Investigator Awards:



Up to 3 years, up to \$100,000 per year.



For senior investigators established either in the field of Progeria or a field that can be directly applied to Progeria

Specialty Awards:



Funding amounts and lengths flexible



👺 For smaller, technology-driven projects, e.g., sequencing, drug screening, obtaining cell lines, antibody preparation, animal models, other

Grant Funding Rates And Topics

As of April 1, 2016, The PRF funding rate is 35%

- Since inception, 175 grant applications received and 62 funded
- PRF has funded 54 researchers from 44 institutions in 13 countries
 - Lamina A, progerin, Lamin B in HGPS and aging
 - Genetics and nuclear function
 - Preclinical Drug Therapy
 - Molecular Abnormalities and Therapies
 - Vascular Pathology
 - Mouse Models
 - Stem Cell Investigations and Therapy
 - Clinical Trials



USA PRF Grantees



GRANTEE NAME	INSTITUTION
Jemima Barrowman	Johns Hopkins University
Juan Carlos Belmonte	Salk Institute for Biological Studies
Ted Brown	The Institute for Basic Research in Developmental Disabilities
Kan Cao	National Institutes of Health; University of Maryland
Christopher Carroll	Yale University
Francis Collins	National Institute of Health
Lucio Comai	University of Southern California
John P. Cooke	Houston Methodist Research Institute
Kris Dahl	Carnegie Mellon University
Jed W. Fahey	Johns Hopkins School of Medicine
Loren Fong	UCLA
Michael Gimbrone	Brigham & Women's Hospital
Thomas W. Glover	University of Michigan
Robert Goldman	Northwestern University
Leslie B. Gordon	Tufts University School of Medicine; Brown University
John Graziotto	Massachusetts General Hospital
Brian Kennedy	Buck Institute for Research on Aging
Jan Lammerding	Cornell University
Dudley Lamming	University of Wisconsin Madison
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USA PRF Grantees



GRANTEE NAME	INSTITUTION
Joan Lemire	Tufts University of Medicine
Jason Lieb	University of North Carolina
Monica Mallampalli	The Johns Hopkins School of Medicine
Susan Michaelis	The Johns Hopkins School of Medicine
Thomas Misteli	National Cancer Institute
Marsha Moses	Harvard Medical School; Boston Children's Hospital
Junko Oshima	University of Washington
Bryce Paschal	University of Virginia
Joseph Rabinowitz	Temple Medical School
John M. Sedivy	Brown University
Dale Shumaker	Northwestern University
Michael Sinensky	East Tennessee State University
Brian Snyder	Beth Israel Hospital
Dylan Taatjes	University of Colorado
Jakub Tolar	University of Minnesota
Katherine Ullman	University of Utah
Thomas Wight	Benaroya Research Institute
Katherine Wilson	Johns Hopkins University
Stephen Young	UCLA
Yue Zou	EastTennessee State University

As of April 1, 2016

International PRF Grantees

















GRANTEE NAME	INSTITUTION
Vincente Andres Garcia	Centro Nacional de Investigaciones Cardiovasculares, Spain
Samuel Benchimol	York University, Toronto, Canada
Bum-Joon Park	Pusan National University, Korea
Claudia Cavadas	University of Coimbra, Portugal
Jesus Vazquez Cobos	Centro Nacional de Investigaciones Cardiovasculares, Spain
Thomas Dechat	Medical University of Vienna, Austria
Karima Djabali	Technical University of Munich, Germany
Maria Eriksson	Karolinska Institute, Sweden
Gerardo Ferbeyre	Université de Montreal, Canada
Celia Ferrera de Oliviera Aveleira	University of Coimbra, Portugal
Evgeny Makarov	Brunel University, England
Charlotte Sorenson	Aarhus University, Denmark
William Stanford	University of Toronto, Canada
Colin Stewart	Institute of Medical Biology, Singapore
Anthony Weiss	University of Sydney, Australia
Zhongjun Zhou	University of Hong Kong, China

PRF Scientific Meetings

Meeting Goals:

➤ To promote collaboration between basic and clinical scientists toward progress in Progeria, cardiovascular, and aging research

PRF has held



international scientific meetings







7 Workshops Promoting Global Interest In Progeria, Cardiovascular Disease And Aging

These are large multi-day workshops open to all scientists. Clinical and basic researchers spend intense days sharing data and planning new collaborations for progress towards treatments and cure.

Various NIH Institutes have funded all 7 workshops through R13 and other granting mechanisms

Other organizations have also generously sponsored workshops





american federation for aging research











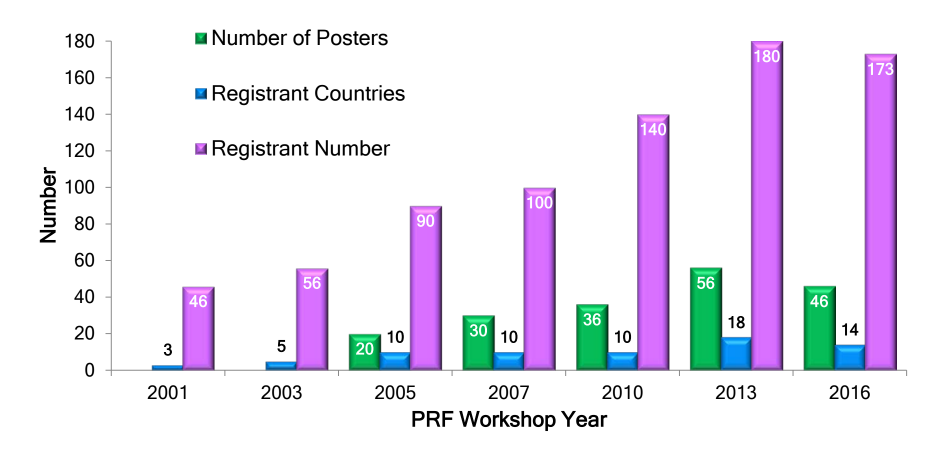




THE MAX AND VICTORIA DREYFUS FOUNDATION



Growth of Global Interest In PRF Workshops





4 Subspecialty Scientific Meetings

Small, focused meetings designed to promote and support work in areas of high interest for Progeria

👺 First Genetics Consortium Meeting - "Searching for the Progeria Gene", August 23, 2002, Brown University, Providence, RI



Second Genetics Consortium Meeting - "Postgene Discovery", July 30, 2003, Bethesda, MD



👺 Bone Marrow Transplant Meeting - "Forging Ahead by Exploring Potential Treatments", April 25-26, 2004, National Institutes of Health, Bethesda, MD



👺 New Frontiers in Progeria Research (2012), Boston, MA









Scientific Publications

As of July 1, 2016:

Scientific articles have been published citing PRF Cell & Tissue Bank resources:



Publication list at www.progeriaresearch.org/cell tissue bank

Scientific articles have been published citing The PRF Medical & Research Database:



Publication list at www.progeriaresearch.org/medical_database

Scientific articles have been published from clinical trial data



See slide #69



Progeria Clinical Care Handbook

The Progeria Handbook. A Guide for Families & Health Care Providers of Children with Progeria. *The Progeria Research Foundation.* Leslie B. Gordon (editor) 2010.



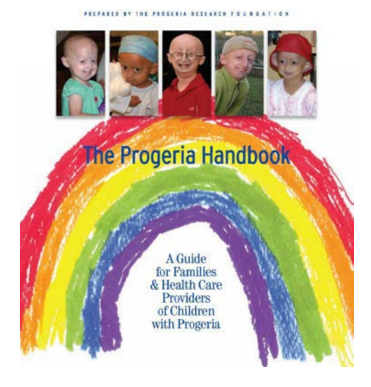
Provided in English, Spanish and Portuguese



Expert contributors from Boston Children's Hospital



Number of Progeria Care Handbooks distributed to families of those with Progeria and their care givers:







NIH Natural History Study

➤ From 2005-2006, PRF participated in an NIH/NHGRI sponsored natural history study that included **15** children with Progeria conducted at the NIH Clinical Research Center.

Goal: to understand the disease processes that drive Progeria.



Phenotype and Course of Hutchinson-Gilford Progeria Syndrome Merideth et al, NEJM, 2008, vol 358, 592-604







The Progeria Research Foundation

Finding...





Together We WILL Find The Cure!

www.progeriaresearch.org

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