



IN PARTNERSHIP WITH



WHO WE SERVE

PC Project provides comprehensive patient support and diagnostics while uniting patients, researchers, physicians, and industry partners on a global level to advance research and drug development for meaningful treatments, and ultimately a cure, for PC and similar rare skin disorders.



OUR VISION

A day when PC sufferers will live without excruciating pain, isolation, and embarrassment.



OUR MISSION

PC Project is dedicated to finding effective treatments and a cure for Pachyonychia Congenita, a painful, debilitating skin disorder, through helping and connecting patients, empowering research, and partnering with industry.

How skin conditions **IMPACT** patients



Pachyonychia Congenita is a painful and debilitating skin disorder caused by a mutation in one of five keratin genes: *KRT6a*, *KRT6b*, *KRT6c*, *KRT16* and *KRT17*. These mutations can cause painful calluses, blisters, cysts, and thickened nails. The worst symptoms are the painful calluses which make walking difficult or impossible. Many patients crawl and rely on walking aids such as crutches or wheelchairs.

OUR STORY

A 501(c)(3) nonprofit, PC Project was established in 2003 by a family who was not content to watch their loved ones with PC suffer and do nothing. Founded and still motivated by love and genuine care, PC Project now serves patients worldwide and facilitates research and drug development.

HOW WE MAKE AN IMPACT

PC Project achieves its mission through two pivotal and foundational programs: the International Pachyonychia Congenita Consortium (IPCC) and the International Pachyonychia Congenita Research Registry (IPCRR).

These powerful initiatives unite patients, scientists, physicians, and industry partners, establishing PC Project as the global repository for PC support and information.

Through collaborative efforts, the organization is committed to transforming PC comprehension, enhancing patient quality of life, and propelling research and drug development in the pursuit of meaningful treatments, and ultimately a cure, for PC.

**SCAN TO LEARN MORE ABOUT
PACHYONYCHIA CONGENITA PROJECT**



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
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



AT A GLANCE HOW WE PROVIDE SUPPORT


- Patient Registry (IPCRR)
- Diagnostic Genetic Testing
- Patient Support Meetings & Online Forums
- Raising Awareness & Education
- Medical Guidance
- Resources, Information on PC, Individualized Support
- Scientific Symposiums
- Collaboration with Scientists, Clinicians, and Industry

GET SOCIAL WITH US

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