



IN PARTNERSHIP WITH



WHO WE SERVE

We strive to serve the health care providers who care for people with HS, the patients who live with HS, and the caregivers who support people with HS.



OUR VISION

The vision of the HS Foundation is a world where no one suffers from HS.



OUR MISSION

The mission of the HS Foundation is to improve the lives of people affected by HS through advocacy, education, and research.

How skin conditions **IMPACT** patients



Hidradenitis Suppurativa is an inflammatory disease that impacts roughly 1% of the US population. The hallmark of the condition is lumps that appear in the folds of the skin, which can cause significant pain, often draining an odorous fluid, creating an embarrassing situation for the person with HS. There is a lengthy delay in proper diagnosis and treatment, leading to poor quality of life outcomes.

OUR STORY

The HS Foundation was created in 2005 to disseminate much-needed information about this relatively unknown disease. Led by the top HS experts in the US, along with patient advocates, the organization seeks to be the trusted resource for patients and health care providers searching for the latest information on HS.

HOW WE MAKE AN IMPACT

Today, the HSF still exists to be a trusted resource, but has expanded its scope to fund clinical research to find better treatments for HS, and ultimately find a cure for this debilitating disease.

It is estimated that HS affects approximately 1% of the population, yet the average time to get a correct diagnosis is seven-ten years.

This disease also disproportionately affects women, as well as people with richly pigmented skin. Over the past few years, the HSF has added several programs and resources with the hope of improving the lives of those with HS.

The Foundation works closely with industry to promote education around the globe, as well as funding research opportunities with the goal of projects leading to larger grant opportunities.

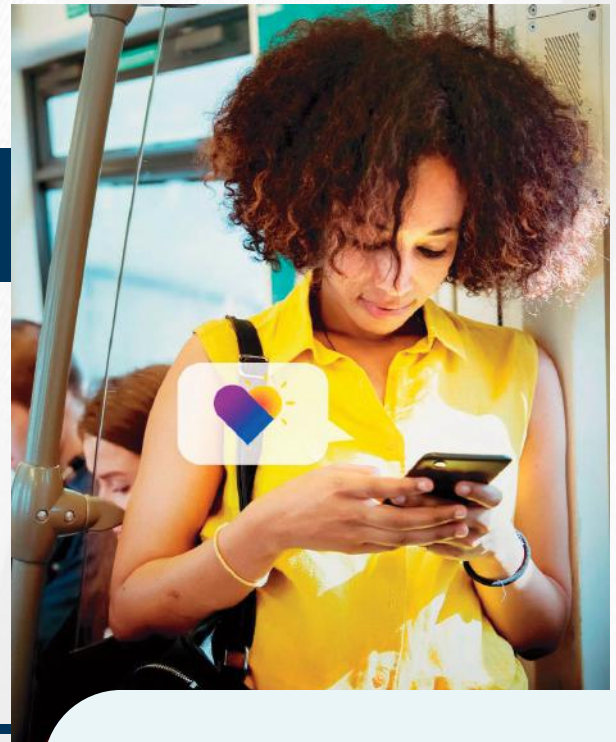
SCAN TO LEARN MORE ABOUT HS



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
info@hs-foundation.org
hs-foundation.org




AT A GLANCE HOW WE PROVIDE SUPPORT


- Research Funding
- Treatment Guidelines
- Health Care Provider Education
- Patient Education
- HS Expert Mentoring
- Career Development Awards
- Education Outside of Dermatology
- Plain Language Research Summaries
- Connection to Support Groups
- Prior Authorization Templates

GET SOCIAL WITH US

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