

SICKLE CELL DISEASE COALITION

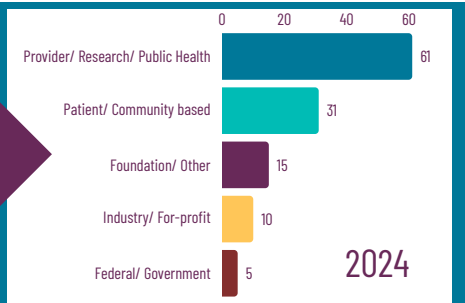


OCTOBER 2024

SCDC SNAPSHOT

Established in 2016 by the American Society of Hematology, the Sickle Cell Disease Coalition (SCDC) is an alliance of diverse stakeholders and multidisciplinary experts from around the world united to raise awareness, broaden education, expand research, advance access to care, and improve health outcomes and quality of life for those impacted by sickle cell disease globally.

SCDC Membership 100+ Organizations



SCDC Goals

- Broaden, amplify, and harmonize voices within the sickle cell stakeholder community
- Unite diverse stakeholders to lead coordinated and collaborative SCD initiatives
- Promote evidence-based interventions and shared decision-making related to sickle cell care

SCDC Subcommittees & Current Priorities



Access to Care

Facilitating networking & knowledge-sharing to improve SCD care



Blood Donor Diversity

Spreading awareness about SCD & need for more diverse blood donors



Global Perspectives

Indexing of SCD stakeholders in regions most impacted by sickle cell



Research & Clinical Trials

Promoting the development & implementation of patient-centered approaches to SCD research



Sickle Cell Trait (SCT)

Promoting best practices for SCT awareness, screening, documentation & high-quality genetic counseling

To learn more about SCDC and opportunities to collaborate:



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[@ConquerSCD](https://www.instagram.com/ConquerSCD)



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