The Problem

Sickle Cell Disease (SCD) is one of the most common genetic diseases globally, but Africans are especially burdened by the disease: nearly 2% of births in sub-Saharan Africa are affected by SCD. While solutions based largely on infection control have reduced the effects of the disease globally particularly among children, it is still difficult to predict, prevent and effectively manage the end-stage organ damage typical of this genetic disorder.

Project Strategy

1. To develop prognostic markers of organ damage through the discovery of key genetic factors and mechanisms that influence the pathogenesis of SCD.
2. To build capacity and career pipelines in Africa to support translational and patient-centred research to advance the development of innovative therapy for SCD.

Outcomes to Date

SickleGenAfrica’s preliminary results have identified that quantitative differences in key hemolysis defense proteins influence the development of end-organ damage in SCD. The network is working to validate these results studying the largest global cohort of SCD patients, and animal models to learn more about their affects in the disease process. As this large-scale research project is in its infancy, the project is currently working to establish their core investigator networks and building infrastructure to support crucial laboratory and clinical work, which will enable it to make massive strides in understanding SCD in the coming years.

Project Leads

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Project Sites

A: Ghana  
West African Genetic Medicine Centre (WAGMC), University of Ghana, Accra. Kwame Nkrumah University of Science and Technology, Kumasi.

B: Nigeria  

C: Tanzania  
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