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Get Involved! Take the Survey at stopsarcoidosis.org/ACTeNOW

THE TIME TO ACTE SNOW!

Current medications prescribed for sarcoidosis were determined without adequate representation of Black Americans in clinical trials, despite their being **2.5x more likely** to have sarcoidosis than White Americans.

ADVANCE CLINICAL TRIALS FOR EQUITY

IN SARCOIDOSIS

TAKE THE FIRST STEP

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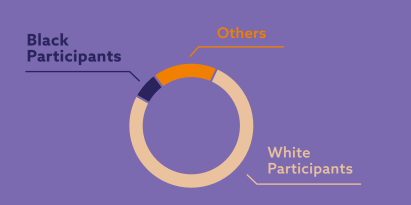
Complete FSR's Patient Survey for Black Americans with sarcoidosis (Incentives provided) by visiting:

STOPSARCOIDOSIS.ORG/ACTENOW

Increasing representation of Black Americans in clinical trials and research is the pathway to developing and identifying the most effective treatments and better health outcomes for Black patients with sarcoidosis.

According to the FDA, Only 7% of clinical trial participants are Black

Without proper representation of Black Americans in clinical trials, therapies may not account for greater health risks, comorbidities, and higher hospitalization and mortality rates.



THERE ARE MORE SARCOIDOSIS CLINICAL TRIALS NOW THAN EVER, SO THE TIME TO ACTe IS NOW!

CLINICAL TRIALS & RESEARCH CAN BE AS SIMPLE AS...

BENEFITS INCLUDE...

- more frequent communication with providers
- increased access to medication, expert clinicians, and advanced technologies
- monitoring regardless of insurance status

- surveys, patient registries, or interviews
- health education programs
- trying medications that have been FDA approved for safety in other diseases
- a simple blood draw
- trying new potential therapies with close oversight

Help improve care for yourself, family members impacted by sarcoidosis, and future generations! Be part of the solution. ACTe Now. Learn How. Visit stopsarcoidosis.org/ACTeNOW



FOUNDATION FOR SARCOIDOSIS RESEARCH

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