Fabry disease is an X-linked disorder; males will be symptomatic, carrier females may or may not be symptomatic. Genetic consultation is recommended.

Patients with at least 1 of the following indications:
- Positive or suspected family history
- Angiokeratomas
- Peripheral neuropathy
- Proteinuria
- Cardiovascular disease
- Or any associated feature

*A separate algorithm is available for positive newborn screen results, see Fabry Disease: Newborn Screen-Positive Follow-up algorithm.

**Molecular genetic testing cannot be added to this specimen type and would require a new specimen collection.

**An interpretive report is provided for all tests in this algorithm.