Newborn Screening ACT Sheet
[Elevated C5-DC Acylcarnitine]
Glutaryl-CoA Dehydrogenase Deficiency

**Differential Diagnosis:** Glutaric aciduria (GA-1)

**Condition Description:** GA-I is caused by a defect of glutaryl-CoA dehydrogenase which limits the metabolism of glutaryl-CoA to crotonyl-CoA, resulting in increased glutaric acid (toxic) and its metabolites.

**You Should Take the Following Actions:**

- Contact family **IMMEDIATELY** to inform them of the newborn screening result.
- Consult with pediatric metabolic specialist.
- Evaluate the newborn for macrocephaly and muscle hypotonia, initiate confirmatory/diagnostic testing as recommended by metabolic specialist.
- Refer to metabolic specialist to be seen as soon as possible but not later than three weeks.
- Educate family about diagnostic possibilities, complexity of diagnostic work-up and the possibility of neurodegenerative crisis with an intercurrent infectious illness.
- **IMMEDIATE** treatment with IV glucose is needed for intercurrent infectious illness.
- Report findings to newborn screening program.

**Diagnostic Evaluation:** Urine organic acid analysis will reveal elevated glutaric acid and 3-hydroxyglutaric acid, should be ordered promptly and is often diagnostic. If urine organic acids don’t confirm the diagnosis, the metabolic specialist will consider analyzing glutarylcarnitine in urine and 3-hydroxyglutaric acid in blood and CSF, enzyme assay in fibroblasts, and molecular analysis of the GCDH gene.

**Clinical Considerations:** The neonate with glutaric acidemia type I is usually macrocephalic but otherwise asymptomatic. Later signs include metabolic ketoacidosis, failure to thrive, and sudden onset of dystonia and athetosis due to irreversible striatal damage. With appropriate treatment, 60-70% of patients will not suffer neurodegenerative disease.

**Additional Information:**
(Click on the name to take you to the website. Complete URLs are listed in the Appendix)

- New England Metabolic Consortium—Emergency Protocols
- Gene Tests/Gene Clinics
- Genetics Home Reference

**Referral (local, state, regional and national):**

- Testing
  - Clinical

**Disclaimer:** These standards and guidelines are designed primarily as an educational resource for physicians to help them provide quality medical services. Adherence to these standards and guidelines does not necessarily ensure a successful medical outcome. These standards and guidelines should not be considered inclusive of all proper procedures and tests or exclusive of other procedures and tests that are reasonably directed to obtaining the same results. In determining the propriety of any specific procedure or test, the healthcare provider should apply his or her own professional judgment to the specific clinical circumstances presented by the individual patient or specimen. It may be prudent, however, to document in the patient’s record the rationale for any significant deviation from these standards and guidelines.
APPENDIX: Resources with Full URL Addresses

Additional Information:

New England Metabolic Consortium—Emergency Protocols
http://www.childrenshospital.org/newenglandconsortium/NBS/descriptions/GAI.html

Gene Tests/Gene Clinics


Referral (local, state, regional and national):

Testing http://biochemgen.ucsd.edu/UCSDW3BG/Labchoose.asp

Clinical
http://www.genetests.org/servlet/access?id=8888891&key=RRqZcXXEUiAx9&fcn=y&fw=zLsf&filename=/clinicsearch/searchclinic.html

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