TO BE RESCINDED

3701-45-02 **Required screening: facility requirements.**

| (A) All newborn children shall be screened in accordance with the procedures set forth in Chapter 3701-45 of the Administrative Code for the presence of the following |
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| genetic, endocrine, or metabolic disorders: |
| (1) Phenylketonuria; |
| (2) Homocystinuria; |
| (3) Galactosemia; |
| (4) Hypothyroidism; |
| (5) Sickle cell and other hemoglobinopathies; |
| (6) Medium chain acyl-coA dehydrogenase deficiency; |
| (7) Maple syrup urine disease; |
| (8) Propionic Aciduria; |
| (9) Isovaleric Aciduria; |
| (10) Methylmalonic Aciduria; |
| (11) Citrullinemia; and |
| (12) Argininosuccinic Aciduria. |
| (B) All hospitals and freestanding birthing centers that are required by this chapter to cause specimens to be collected for newborn screening for genetic, endocrine, or metabolic disorders shall: |
| (1) Designate a newborn screening coordinator and physician responsible for the |

coordination of the facility's newborn screening;

(2) Notify the chief of the Ohio department of health bureau of public laboratories of the name of the individual designated as the newborn screening

coordinator on a yearly basis and whenever the designated individual changes; and

(3) Develop a written protocol for tracking newborn screening. The protocol must include a requirement that the name of the physician attending the child after birth or a designee be placed on the specimen slip sent with the initial specimen to the Ohio department of health public health laboratory.

Effective: 04/14/2003

R.C. 119.032 review dates: 01/17/2003

CERTIFIED ELECTRONICALLY

Certification

04/04/2003

Date

Promulgated Under: 119.03 Statutory Authority: 3701.501 Rule Amplifies: 3701.501

Prior Effective Dates: 1/31/02, 1/13/01, 2/15/99