

GBSC 724 Advanced Special Topics in Metabolomics

Population Scale Metabolomics: Newborn Screening

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1

Lecture Overview

- Introduction and historical perspective
- Disorders
- Methods
- Logistics, ethical issues, and future considerations

Prologue: the Impact of Newborn Screening

- JS was born in1955 with phenylketonuria (PKU).
 Undiagnosed, he developed severe intellectual disability and was institutionalized at the age of 20.
- JD was born in1965 with PKU. NBS was now available and led to a diagnosis at 2 weeks of age. He was placed on a special diet, and grew to be normal adult.
- ES was born in a state without medium chain acyl-CoA dehydrogenase (MCAD) deficiency screening in 1999. Undiagnosed, she died in her sleep at 15 months of age.
- RD was born on the same day, but 20 miles away, just across the border in a state where MCAD screening was offered.
 She was placed on dietary therapy and grew to be a normal adult.

3

Newborn Screening: One of the Ten Great Public Health Achievements Worldwide, 2001–2010

"Improvements in technology and endorsement of a uniform newborn-screening panel of diseases have led to earlier life-saving treatment and intervention for at least 4000 additional newborns each year with selected genetic and endocrine disorders."

> Morbidity & Mortality Weekly Report. 2011; 60(24):814-818 © 2011 Centers for Disease Control and Prevention (CDC)

What is Newborn Screening (NBS)?

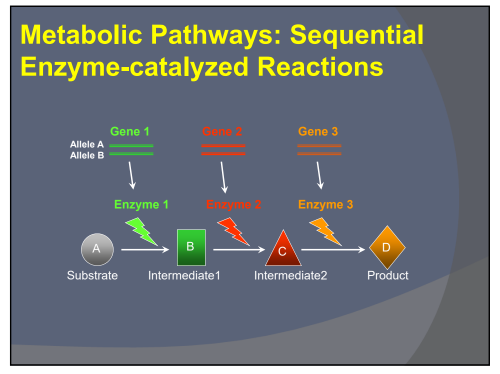


Approximately 1 in 300 newborns has a condition detectable by modern NBS

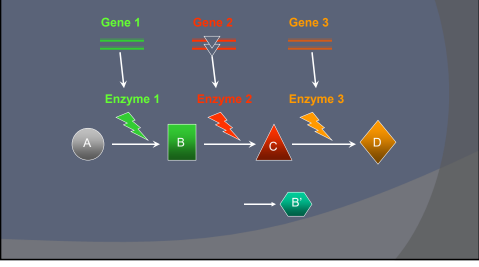
*USA: 4 million births/year

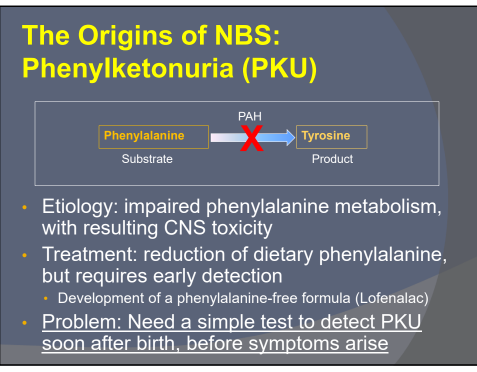
- Population scale screening of all newborns* for the presence of *treatable* conditions that are not otherwise evident at birth
 - screening vs. diagnostic testing
- State specific programs (no federal mandate) with significant variability
 - disorders detected
 - follow-up procedures



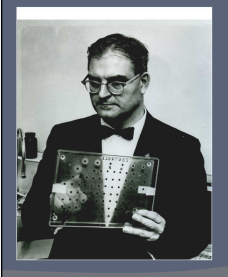








Robert Guthrie Pioneered the First NBS Test for PKU in 1961



- Filter paper containing blood from newborns applied to a seeded agar plate
- Bacteria only grow in the presence of phenylalanine
 - Large colonies = PKU
- Paradigm: one test for one disorder

9

A Brief History of Newborn Screening: the Early Years

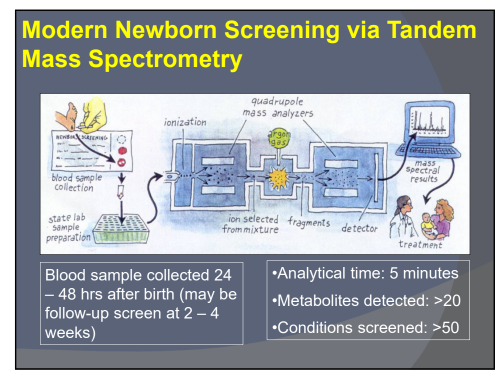
- 1961: Robert Guthrie develops screening test for PKU
- 1962: Massachusetts pilots state-wide PKU screening
- 1965: Over 50% of states have mandated PKU screening
- 1968: WHO publishes Principles and Practices of Screening for Disease
 - Wilson-Jungner principles (early screening criteria)
 - 1970s 1990s: most states screen for ~6 conditions

A Brief History of Newborn Screening: the Era of Mass Spectrometry



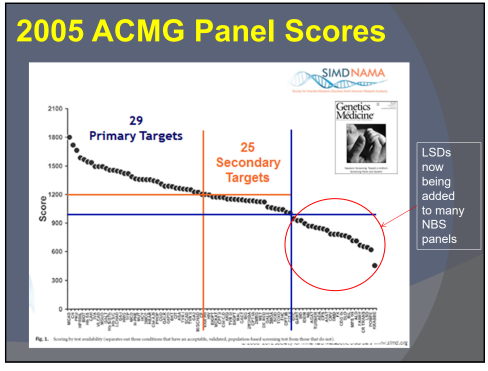
- 1990s early 2000s: Development and implementation of MSMS for newborn screening
 - New paradigm: one test for multiple disorders
- 2002: Maternal and Child Health Bureau commissions ACMG to recommend a uniform panel of conditions for NBS
 - 2005: ACMG ENS report identifies 29 core conditions and 25 secondary conditions (designated by HHS as the national standard for NBS – but not federally mandated)
 - 2009: All states screen for at least 29 disorders; approximately 20 states screen for 40+ disorders





Criteria for Inclusion in the ACMG Core Screening Panel (2006)

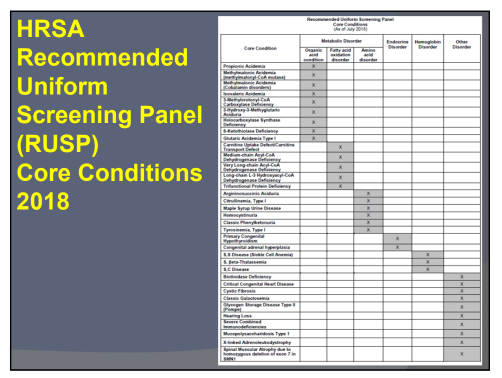
- An effective treatment is available
- Demonstrated benefits of early detection and treatment (clinical utility)
- The condition does not usually produce symptoms within 24 – 48 hrs after birth
- A sensitive, specific, and cost-effective test is available that can detect the condition within this time frame
- See http://mchb.hrsa.gov/screening/ for more about the ENS task force



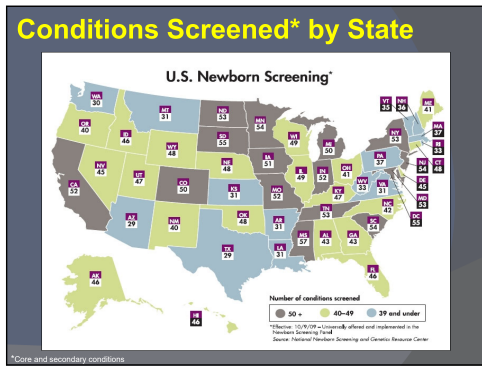
Screened Disorders in the United States

- Currently, 35 core conditions are on the Recommended Uniform Screening Panel (RUSP)
 - 20 classified as metabolic disorders (eg, PKU)
 - 2 endocrine disorders (eg, CAH)
 - 3 hemoglobin disorders (eg, sickle cell anemia)
 - 10 other conditions (eg, hearing loss, cystic fibrosis)
- Also 26 secondary conditions (may lack an effective therapy or have an unclear natural hx) that can be detected when screening for core disorders
 - 22 metabolic
 - 1 hemoglobinopathy
 - 3 other

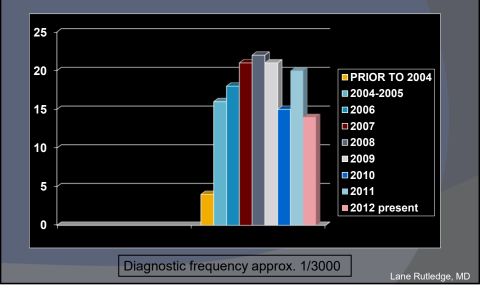
National Newborn Screening & Global Resource Center (NNSGRC)



2018 RUSP mmended Uniform Screening Panel SECONDARY² CONDITIONS ³ (As of July 2018) Secondary Metabolic Disorder Other Disorder Hemoglobin Disorder Secondary Condition Fatty acid Amino oxidation acid disorders disorder Organic acid Conditions Methylmalonic acidemia with homocystinuria Malonic acidemia isobutyrylglycinuria 2-Methylbutyrylglycinuria 2-Methyloutysygigycnuna 3-Methylglutaconic aciduria 2-Methyl-3-hydroxybutyric aciduria Short-chain acyl-CoA dehydrogenase deficiency Medium/short-chain L-3-hydroxyacyl-CoA dehydrogenase deficiency Charlie acidurais hans u CoA dehydrogenase deficiency Glutaria acidemia type II Medium-chain ketoacyl-CoA thiolase deficiency 2.4 Diencyl-CoA reductase deficiency Camitine pathiofyttansferase type II deficiency Camitine pathiofyttansferase type II deficiency Camitine pathiofyttansferase type II deficiency Camitine pathiofittansferase type II deficiency Argininemia X rgininemia Citrullinemia, type II Benign hyperphenylalaninemia Biopterin defect in cofactor biosynthesis Biopterin defect in cofactor regeneration Tyrosinemia, type II Tyrosinemia, type III Various other hemoglobinopathies Galactoepimerase deficiency ctokinase deficiency T-cell related lymphocyte defici Selection of conditions based upon "Weatoom Screening: Towards a Uniform Screening Panel and System." Ge 2027 as automoted by the American College of Medical Genetics (ACMG) and commissioned by the Health Res Discretes that can be detected in the attentional disposals of a core discrete. Nomendature for Conditions based upon "Naming and Country Disorders (Conditions) included in Newsom 50 117 (§) Super Scho-314. etic Med. 2006; 8(5) Suppl: S12-



Alabama NBS: New Diagnoses Since Initiation of Expanded Newborn Screening



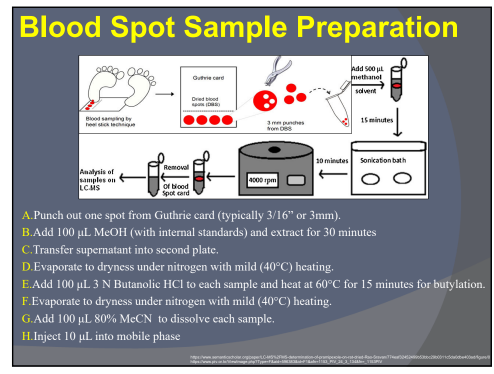
19

Overview of a Modern Newborn Screening Workflow Follow-up testing Required to confirm or Sample collection by refute screening results heel stick at 24 – 48 hrs · Vary significantly by state · Most infants (90%) with Transport to NBS program abnormal NBS results have normal follow-up Prematurity Invalid sample Screen positive Screen negative TPN or certain •unsat •unsat •sample <24 hrs •delivered >14 days •TPN or transfusion •prematurity formulas · If disease is confirmed then treatment is Results sent to Physician initiated immediately immediately referring contacted by phone physician ↓ Repeat sample Referral for follow-up to confirm diagnosis and begin treatment requested

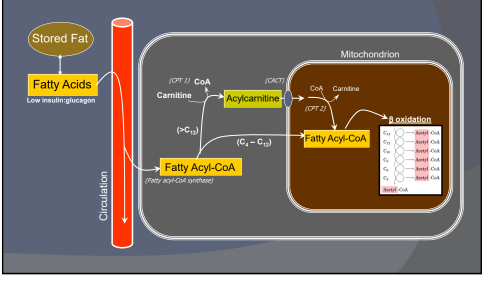
Analysis of Metabolites



- Small molecule substrates or products of enzyme-catalyzed reactions
 - Targeted metabolomics
 - Biomarkers
 - · Precise instrumental analysis techniques
 - Accurate and appropriate reference ranges
 - · Caution: overreliance on ref ranges
 - Quality control extremely important



Acylcarnitines: Intermediates of Fatty/Organic Acid Oxidation



23

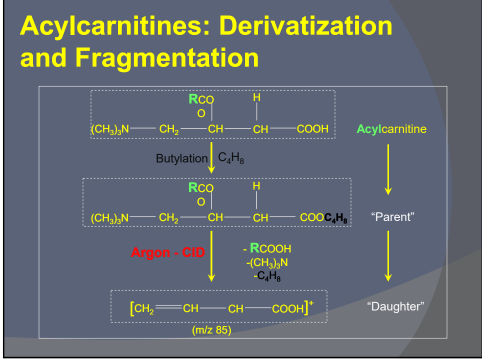
Acylcarnitines as Biomarkers

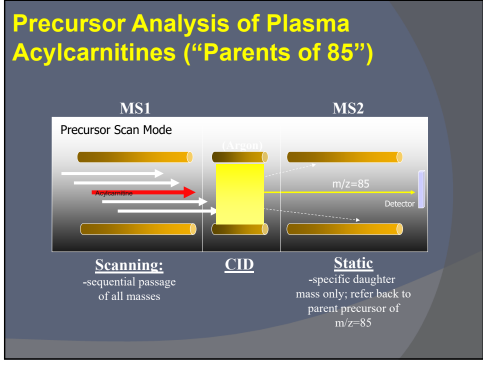


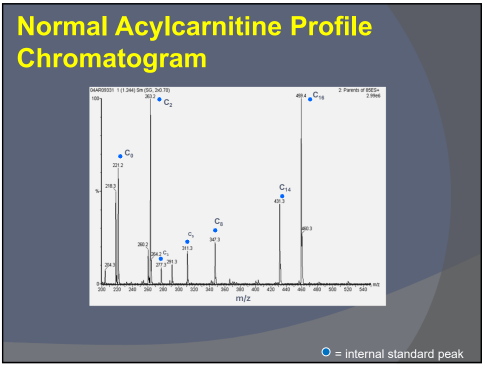
- Deficient fatty/organic acid oxidation results in accumulation of one or more <u>size-specific</u> acylcarnitines in blood
 - Effectively measured via MSMS
- Initial basis for expanded newborn screening
- Disorders detected
 - Fatty acid oxidation disorders
 - Organic acid disorders
 - Other conditions identified
 - Ketosis, acidosis, catabolism, liver disease, renal disease, MCT feeding, etc
- Methodology
 - MSMS analysis of butylated acylcarnitines
 - Quantification of >30 acylcarnitines
 - Analytical time: ~2 hrs

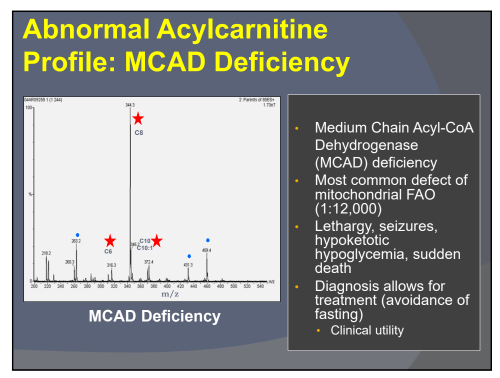
Acylcarnitine Analysis Sample requirements • Plasma (>1 mL) 20 ul used in assay Limitations Interfering substances Results generally not considered to be diagnostic (enzyme activity and/or sequence analysis) Confounders Liver/kidney disease (AC-DCs) • Ketosis (C2, C4-OH, C12:1, C14:1) • MCT oil (C8, C10) Valproate (C0, C8, C10) • Carnitine supplements (short chain ACs) • Cefotaxime (C14:1, C16:1-OH) Cheese (C3)

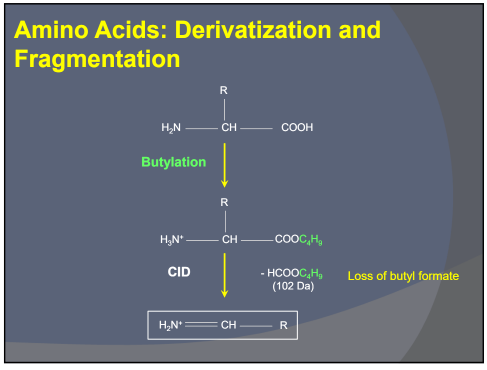


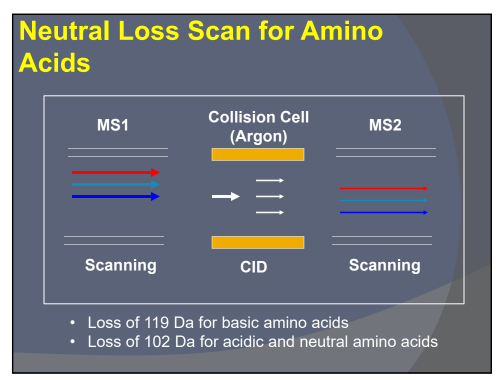


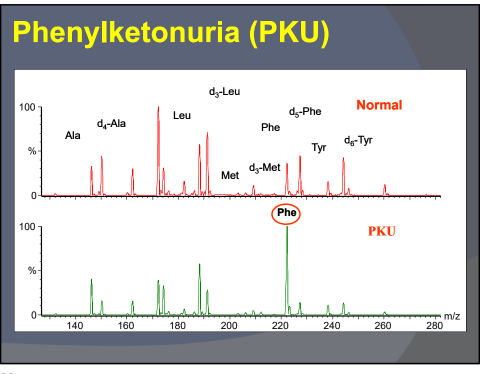












Benefits of Newborn Screening



- 4000 5000 newborns/yr experience significantly improved health outcomes¹
- prevents diagnostic odysseys
- Cost-effectiveness (congenital hypothyroidism):
 - Annual economic cost of screening and early treatment for CH is 20-fold less than treating severely affected patients who were not screened
 - (\$400 M vs. \$20 M)²

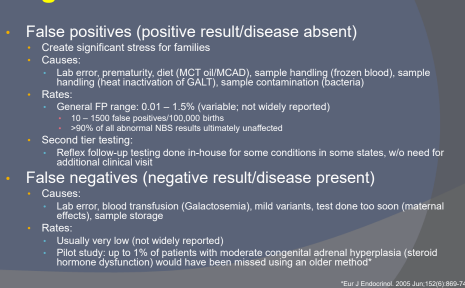
Limitations of NBS



- False positives
- False negatives
- Many types of metabolic disorders are not screened
- Questionable clinical utility for some screened disorders
- Lack of clinical and laboratory expertise
- Significant financial constraints

35

False Positives and False Negatives



Newborn Screening: Ethical Issues



- Privacy
 - Sample retention and security of stored data
- Clinical utility is questionable for some screened disorders
 - Severe forms of certain disorders that may present before NBS results are available
 - Very rare disorders with small numbers of affected patients, making outcomes uncertain
 - Very mild, ill-defined phenotypes
 - Lack of treatment options

37

The Future of Newborn Screening



Where Does NBS Go From Here?



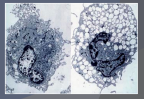
- The existing NBS model continues to evolve
 - More conditions being added or considered for screening (eg, LSDs)
 - Changes to current screening criteria proposed
- Next generation DNA sequencing: the new screening paradigm?
 - Potential for massive expansion of genetic screening

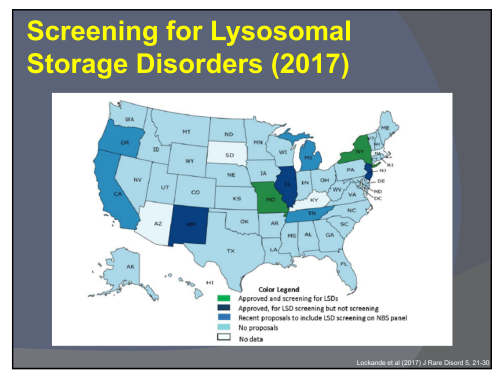
39

Newborn Screening for Lysosomal Storage Disorders (LSDs)

- LSDs: disorders of lysosomal enzymes that degrade/recycle cellular waste products.
- Accumulating materials cause progressive damage to multiple organs, incl CNS
 - Often early mortality w/o treatment
- Estimated incidence: 1:5000 10,000
- LSDs as candidates for NBS:
 - Usually not apparent at birth
 - Diagnosis is often delayed
 - Growing number of therapeutic options and demonstrated benefits of early treatment
- Multiplex screening methods now available
 Several programs now offering or piloting limited LSD
- screening (Alabama 2021: Pompe)







Should We Screen for Diseases Without an Effective Therapy?

- Cornerstone of traditional screening: must be an effective treatment available
- However, it has been suggested that future screening should consider other benefits:
 - avoiding diagnostic odysseys
 - making preparations for disease
 - reproductive decisions
 - early access to promising new therapies

The Next Big Thing: Next Generation Sequencing (NGS)?



- DNA sequencing-based methods may represent the future of genetic screening
- Will initially take the form of small scale, targeted panels
 - The National Institute of Child Health and Human Development (NICHD) is currently funding efforts to develop DNA-based screening.
- Ultimately, the entire genome of all newborns may be routinely sequenced at birth
- Paradigm shift? Functional (biochemical) testing to confirm molecular screening (see below)

43

Obstacles to NGS Screening Cost Must be cost effective: current NBS testing costs ~\$2.00/disorder. Current genome sequencing costs about \$1000 (w/o interpretation) Costs are falling rapidly; may become cost-effective in the next 5 - 10 years Infrastructure Bioinformatics: data storage and analysis Expansion of follow-up programs? Genetic counseling Ethical considerations Security/privacy Variants of unknown significance Incidental findings 44

