

# Neonatal Syndrome

## Multisystem Maladaptation

Hypoxic Ischemic Syndrome

Perinatal Asphyxia

Hypoxic Ischemic Asphyxial Syndrome

Neonatal Maladjustment Syndrome

Dummy Foals



# Changes in Behavior

















# Neonatal Intensive Care



# Hypoxic-Ischemic Syndrome

- Human Neonates - cerebral palsy
  - Prolonged Stage II
  - Lawsuits
  - Clinical studies on onset
    - Intranatal
    - Prenatal
    - Postnatal
- Experimental Studies
  - Hypoxic ischemic insults
  - Hypoxic ischemic encephalopathy (HIE)





# Neonatal Problems

## Hypoxic Ischemic Asphyxial Disease

- Selective neuronal pathology
- Renal pathology
- Gastrointestinal pathology
- Metabolic failure
- Cardiovascular pathology
- Endocrine abnormalities
- Pulmonary pathology



# Neonatal Problems

- Hypoxic ischemic asphyxial disease?
  - Often no evidence
- Inflammatory placental disease
  - Strong correlation
- Role of inflammatory mediators?
  - Cytokines, local vasoactive mediators
  - Primary effect?
  - Secondary hypoxic ischemic insult?

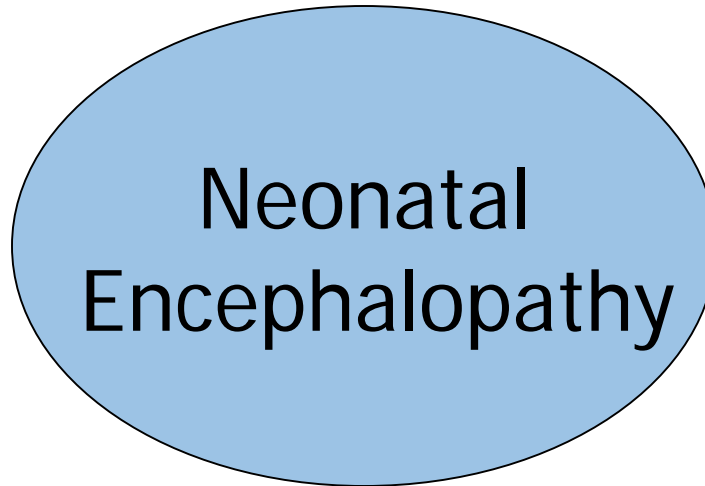
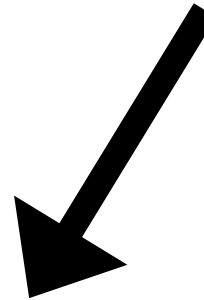
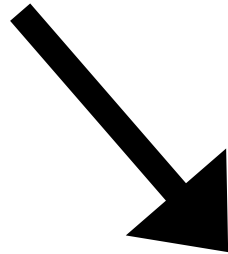




Hypoxic  
Ischemic  
Insults



Inflammatory  
Insults



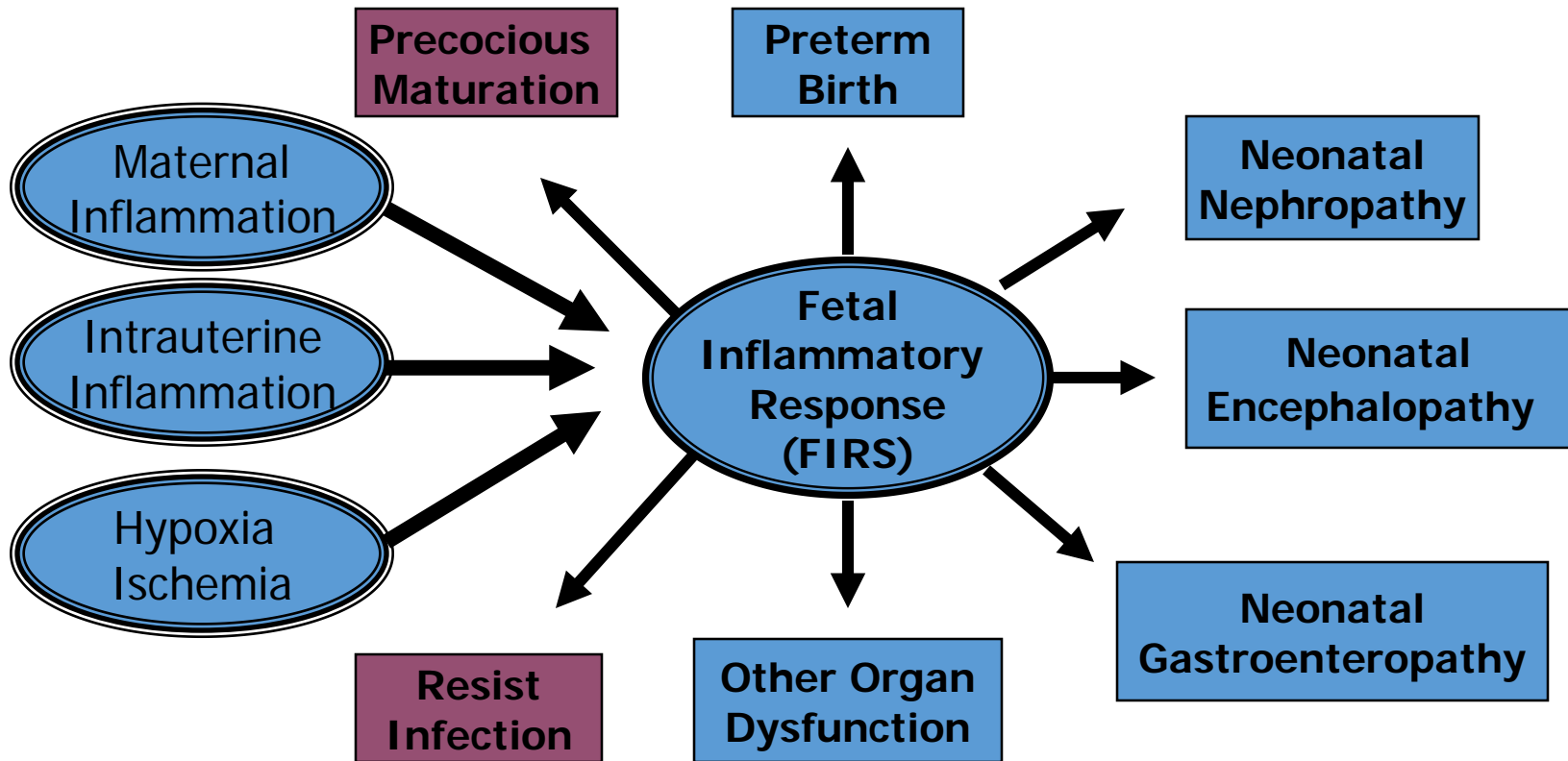
Neonatal  
Encephalopathy

# Role of Placentitis

- Many neonatal diseases
  - Multiple etiologies
  - Disruption of fetal life
    - Predispose to neonatal disease
    - Origin of the neonatal disease
- Placentitis - untreated
  - Neonatal diseases
    - CNS, Renal, GI
- Placentitis - treated
  - Protects against neonatal diseases





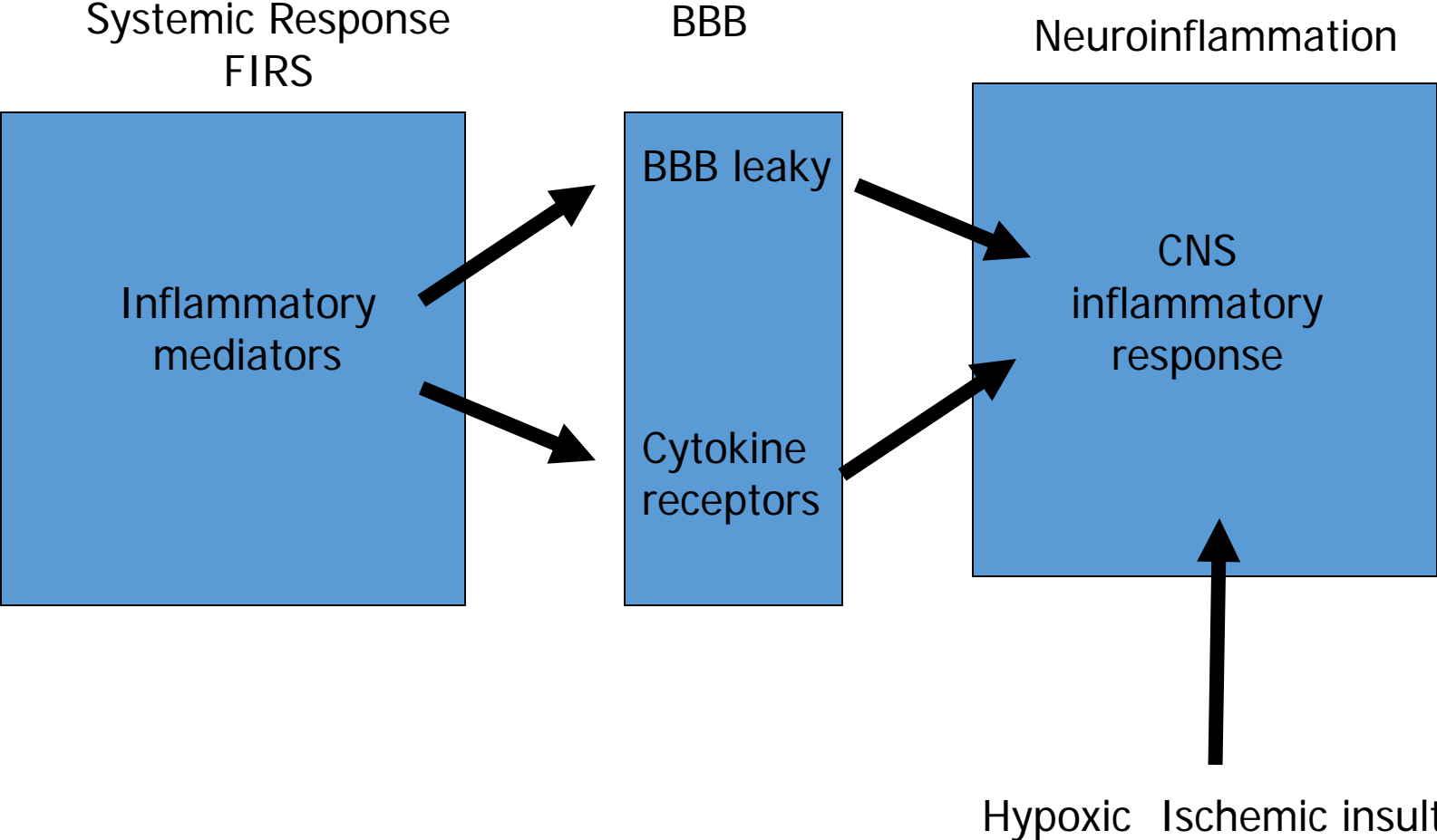


# Septic Encephalopathy

- Fetal
  - Neuroinflammation
  - FIRS (Fetal Inflammatory Response Syndrome)
    - Fetal placentitis
- Maternal
  - Maternal placentitis
  - SIRS
  - Focal maternal infections



# Septic Encephalopathy



# Neuroinflammation

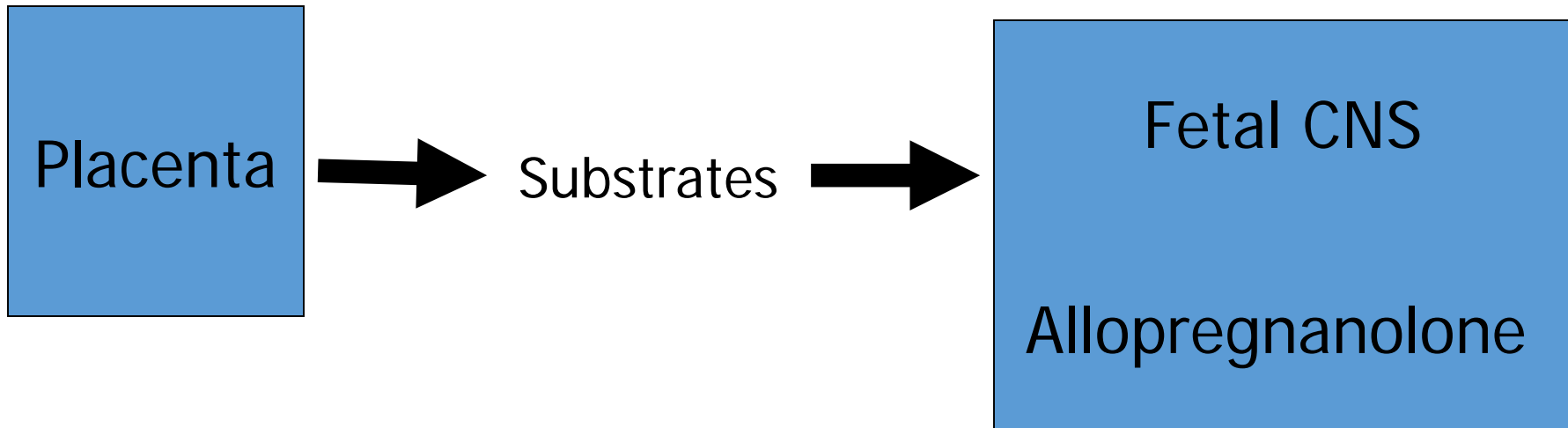
- Important in the pathogenesis of
  - Septic encephalopathy
  - Hypoxic ischemic encephalopathy
- Microglia cells are key
  - Up-regulation of proinflammatory cytokines
  - Up-regulation of trophic factors
- Can result in
  - Morphological alterations
  - Biochemical alterations
  - Functional alterations

# Neuroinflammation

- Response depends on mix
  - Proinflammatory
  - Anti-inflammatory
  - Specific mediators
- Mild disease – often no morphologic changes
  - Motor
  - Perceptual, visual
  - Behavioral
  - Cognition
  - Excitatory responses
- Excitotoxicity



# Neurosteroids



- Protect the brain during fetal life
- Responsible for the somnolence
- At birth
  - Removal of the placental
  - Levels drop rapidly
  - Fetus to "awake up"

# Neurosteroids

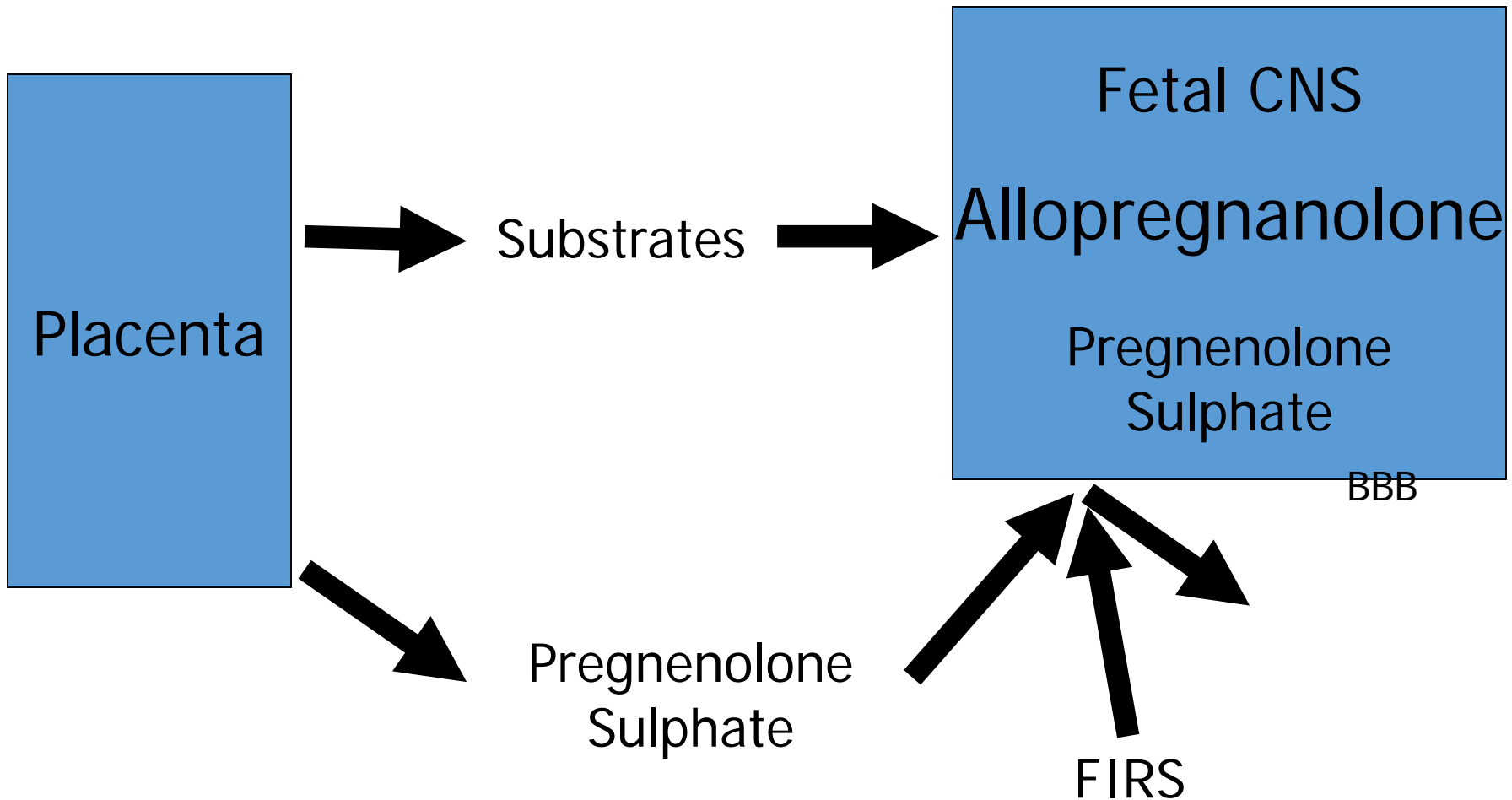
- Allopregnanolone
  - Brain levels induced by
    - Inflammatory mediators
    - Hypoxic ischemic insults
  - Protect against neuroexcitatory toxicity
  - Marked anti-seizure actions
  - Raise seizure threshold
  - Induces somnolence

# Neurosteroids

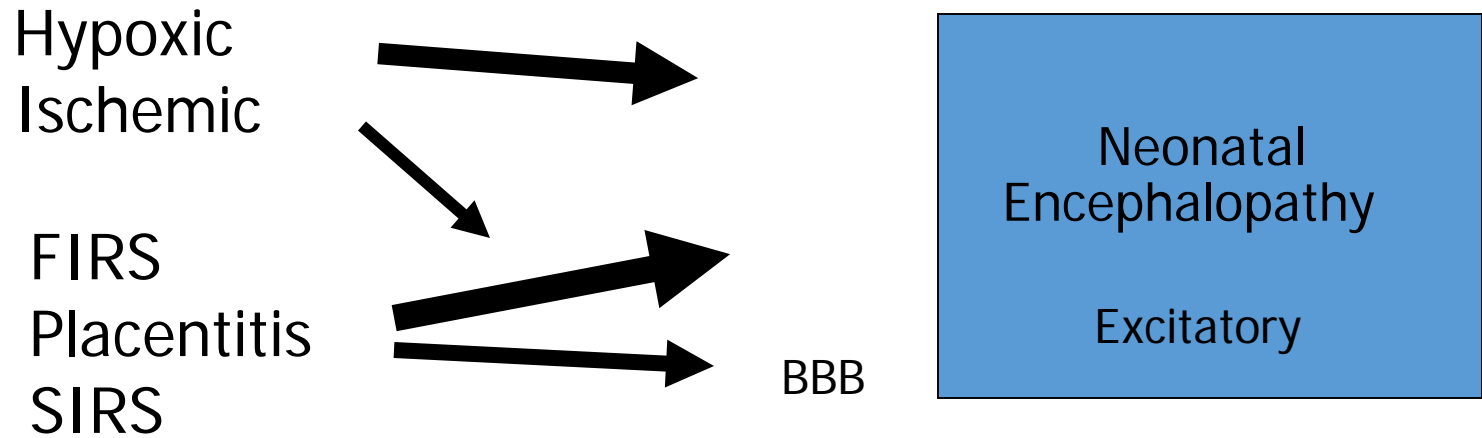
- Pregnenolone and pregnenolone sulphate
  - Placenta also secretes
  - Excitatory action in the brain
  - Cross the blood brain barrier
    - Normal – slow
    - Abnormal BBB – rapid transfer
    - Inflammation
    - Hypoxic ischemic insult



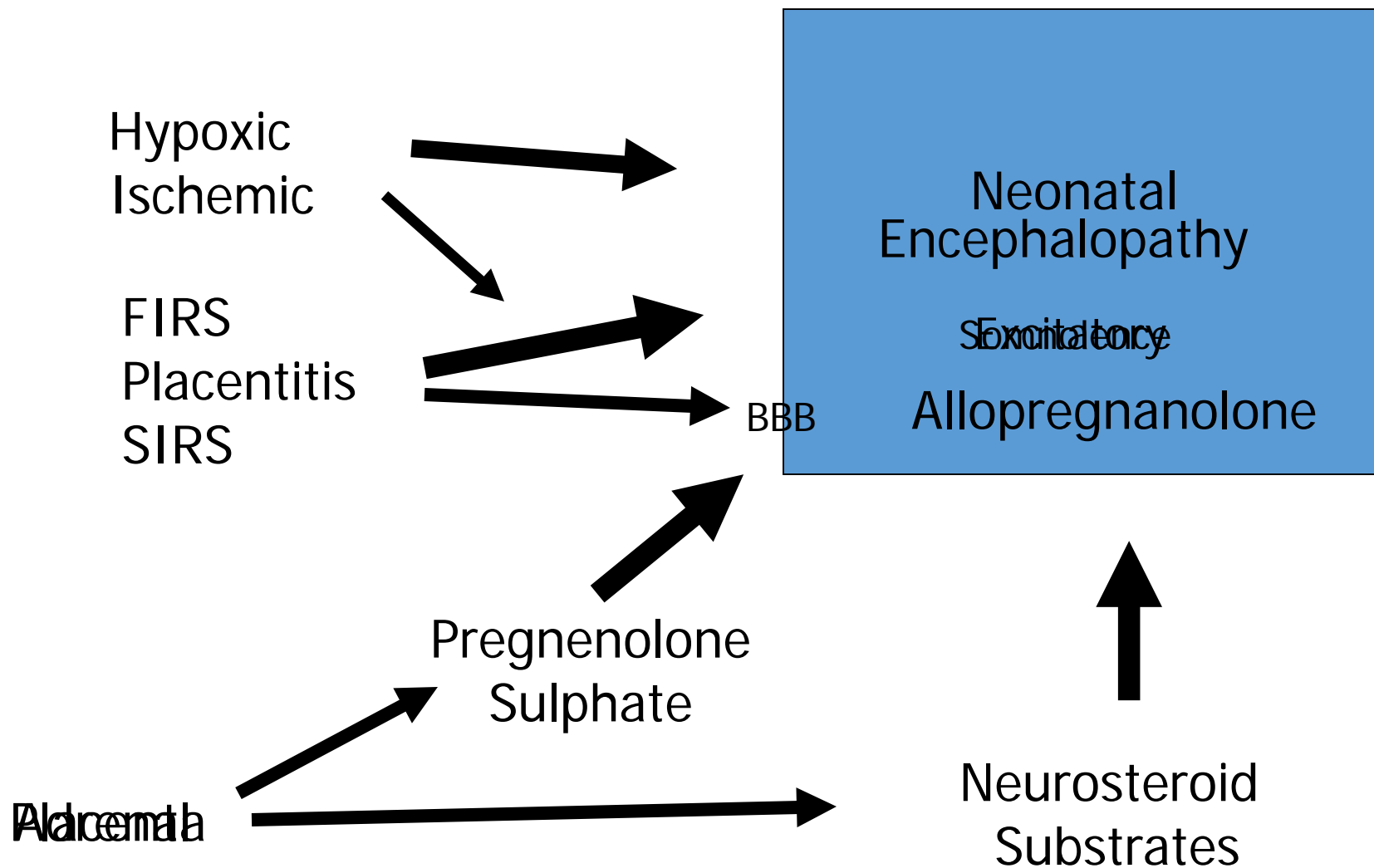
# Neurosteroids



# Neonatal Encephalopathy



# Neonatal Encephalopathy





# Typical Clinical Course

- Born near normal behavior
- Initial signs – excitatory
  - Constant activity – wandering, not lie down
  - Hyper-responsiveness
  - Hypertonus
  - Culminating in tonic-clonic seizure-like behavior
- Onset of somnolent phase
  - Stress induced adrenal steroidogenesis
  - Neuroinflammation induces neurosteroids
  - Healing period
- Recovery

# Typical Clinical Course

- **Born seizure-like behavior**
  - Less placental steroidogenesis
    - Lower levels protective neurosteroids
  - Inflammatory mediators
    - Induced blood brain barrier deficits
    - Allow sulfated neurosteroids into CNS
- **With neonatal stress onset of somnolent phase**
  - Stress induced adrenal steroidogenesis
  - Neuroinflammation induced CNS neurosteroids
  - Healing period



# Changes in responsiveness

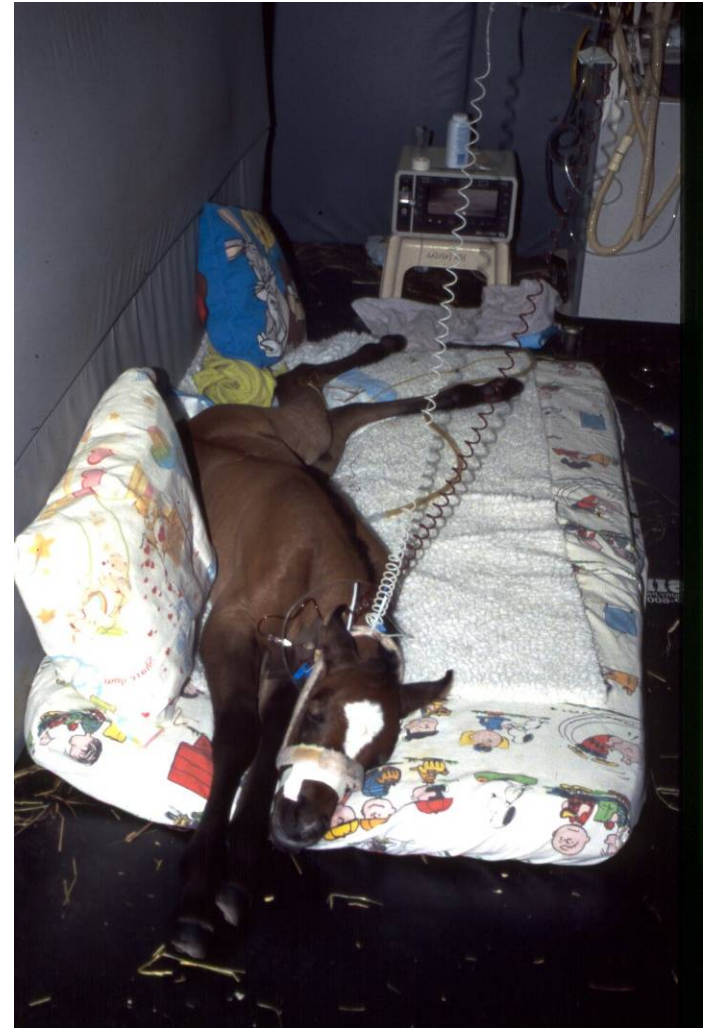




# Changes in muscle tone



# Changes in muscle tone





# Changes in behavior



# Brain stem damage





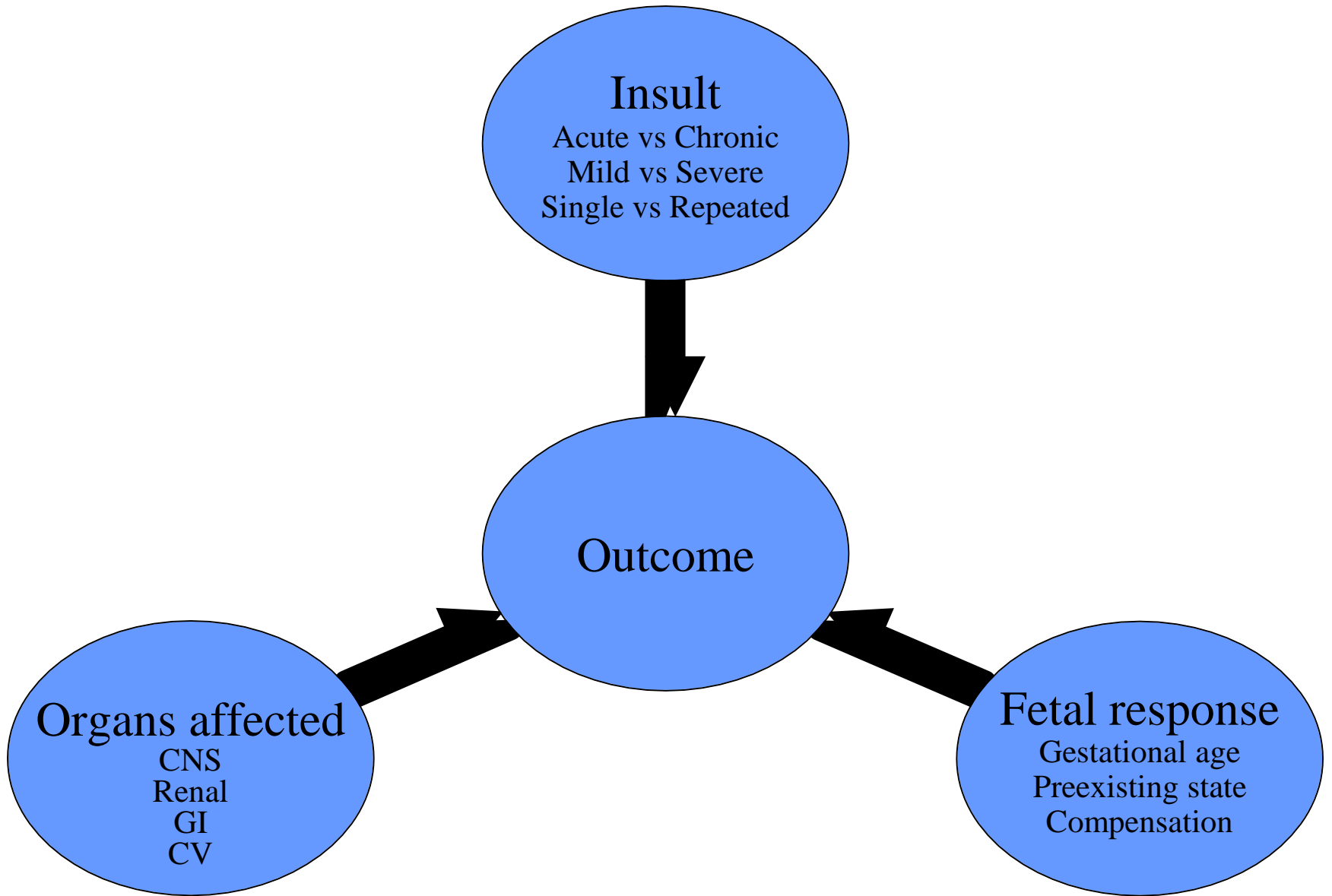
# Seizure-like behavior



# Terms

## Generic Description of Signs

- Neonatal Encephalopathy (NE)
- Neonatal Gastroenteropathy (NG)
- Neonatal Nephropathy (NN)
- Neonatal Metabolic Maladaptation
- Neonatal Cardiovascular Maladaptation



## Insult

Acute vs Chronic  
Mild vs Severe  
Single vs Repeated

## Outcome

## Organs affected

CNS  
Renal  
GI  
CV

## Fetal response

Gestational age  
Preexisting state  
Compensation

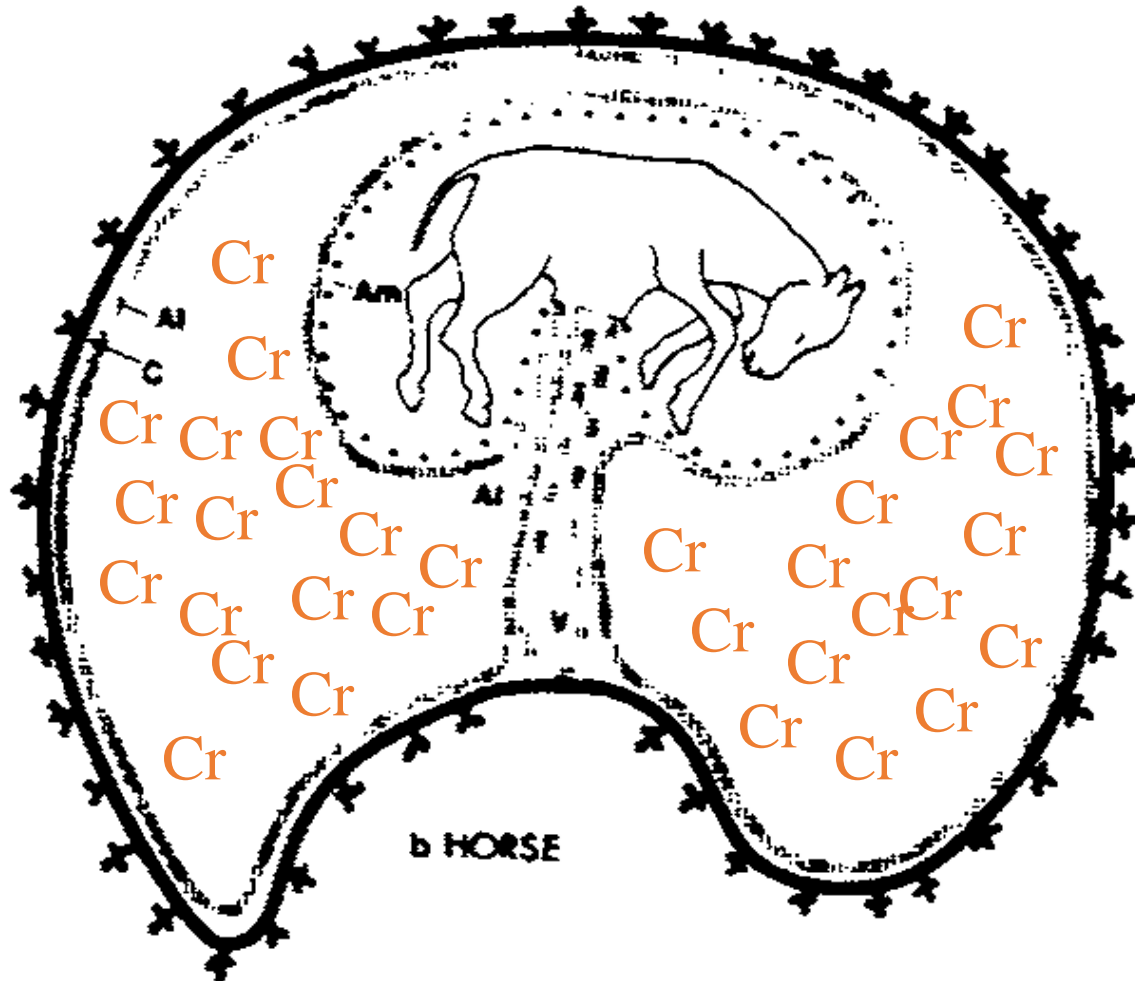
# Intrauterine Challenge

- Indications at birth of intrauterine

- Cr level
- Hypochlo
- High PCV
- High birth
- Persistent
- Ca levels
- Fibrinoge
- WBC
- Low corti
- Lactate le



Fetal foal floating  
in a sea of creatinine







# “Pong”

Thoroughbred foal

Born: May 7 at 6 PM

Admitted: May 8 at 8:53 AM

15 hrs old

# “Pog” History

- Term birth to a multiparas mare
- Normal gestation
- Stage 1 - not observed
- Stage 2 - 10 minutes or less
- Stage 3 - 1 hour
- Assisted to stand after 1.5 hours
  - Nursed from the mare

# “Pog” History

- Never vigorous
- Got up once during night
  - Only for short time
  - Did not nurse
- Bottle-fed 8 oz. of colostrum
- Referred for intensive care
  - Weak
  - Inability to stand

# “Pig” Admission Physical

- Marked oral, nasal, scleral, aural icterus
- Oral, nasal, scleral, aural injection
- Multiple oral petechia
- Marked lingual erythema
- Abdomen
  - Meconium in the right dorsal colon
  - Few borborygmi
  - Fetal/neonatal diarrhea

# “Pig” Admission Physical





**“Pong”**

## Admission Laboratory Data

	<b>Admission</b>	<b>Normal</b>
<b>Fibrinogen</b>	<b>461 mg/dl</b>	<b>150 mg/dl</b>
<b>WBC</b>	<b>800 cells/ul</b>	<b>5-10,000</b>
<b>Neutrophil</b>	<b>42% cells/ul</b>	<b>50-80%</b>
<b>Lymphocytes</b>	<b>38% cells/ul</b>	<b>20-50%</b>
<b>Creatinine</b>	<b>6.46 mg/dl</b>	<b>2.5-4.0</b>
<b>Glucose</b>	<b>44 mg/dl</b>	<b>60 – 120</b>
<b>PCV</b>	<b>54%</b>	<b>30 – 45%</b>
<b>TPP</b>	<b>6.1 gm/dl</b>	<b>4.0 – 5.5</b>

**“Pig”**

## Admission Problems

- **Weakness, somnolence**
- **Not nursing**
- **Lingual erythema**
- **Injection**
- **Petechia**
- **Icterus**
- **Poor perfusion**
- **Diarrhea**
- **↓ WBC,**  
**↑ fibrinogen**
- **↑ PCV, ↑ TPP**
- **↑ Creatinine**
- **Hypoxemia**
- **↑ lactate**

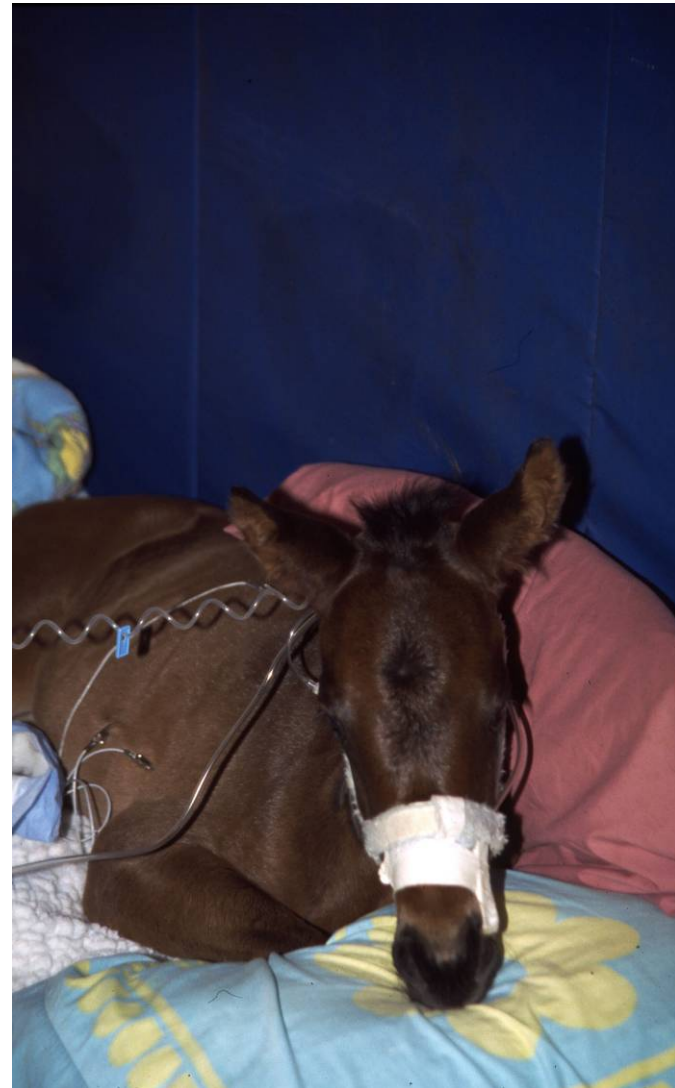
# “Pig” Major Problems



# “Pong”

## Neonatal Encephalopathy

- Periods - bright and active
- Sudden onset of somnolence
  - Somnolence/periods of arousal
- Apparent facial paresis
  - Right ear moves slowly
- Generalized weakness



“Pog”

# Neonatal Encephalopathy

- Periodic apnea
  - Up to 60 sec
  - With clustered breathing
- Inappropriate central tachypnea
- Apneusis (apneustic respiration)
- Hypercapnia
  - Without apnea



“Pog”

# Neonatal Encephalopathy

- Seizure like activity
  - Opisthotonus, tonic/clonic marching activity
  - Minimal nystagmus
- Lingual erythema
- Moderate nasal septum hyperemia
- Hyperresponsive to stimuli
- No suckle or searching

# Neonatal Encephalopathy

## CNS Signs

- Most common and noticeable
  - Signs occur predictably - 90%
- Mild central insult
  - Multifocal lesions
  - Selective neuronal dysfunction
  - Slow maturation of coordination



# Neonatal Encephalopathy

## Signs of CNS disease

- Changes in responsiveness
- Changes in muscle tone
- Changes in behavior
- Signs of brain stem damage
- Seizure-like behavior
- Coma, death



# Neonatal Encephalopathy

## Signs of CNS disease

- Changes in responsiveness
  - Hyperesthesia
  - Hyperresponsiveness
  - Hyperexcitability
  - Hyporesponsiveness
  - Periods of somnolence
  - Unresponsiveness



# Neonatal Encephalopathy

## Signs of CNS disease

- Changes in muscle tone
  - Extensor tonus
  - Hypotonia
  - Neurogenic myotonia
  - Inability to protract legs





# Neonatal Encephalopathy

## Signs of CNS disease



- Changes in behavior
  - Loss of suckle response
  - Loss of tongue curl
  - Loss of tongue coordination
  - Disorientation especially relative to the udder
  - Aimless wandering
  - Blindness
  - Loss of affinity for the dam
  - Abnormal vocalization ("barker")

# Changes in behavior



# “Pog” Neonatal Encephalopathy



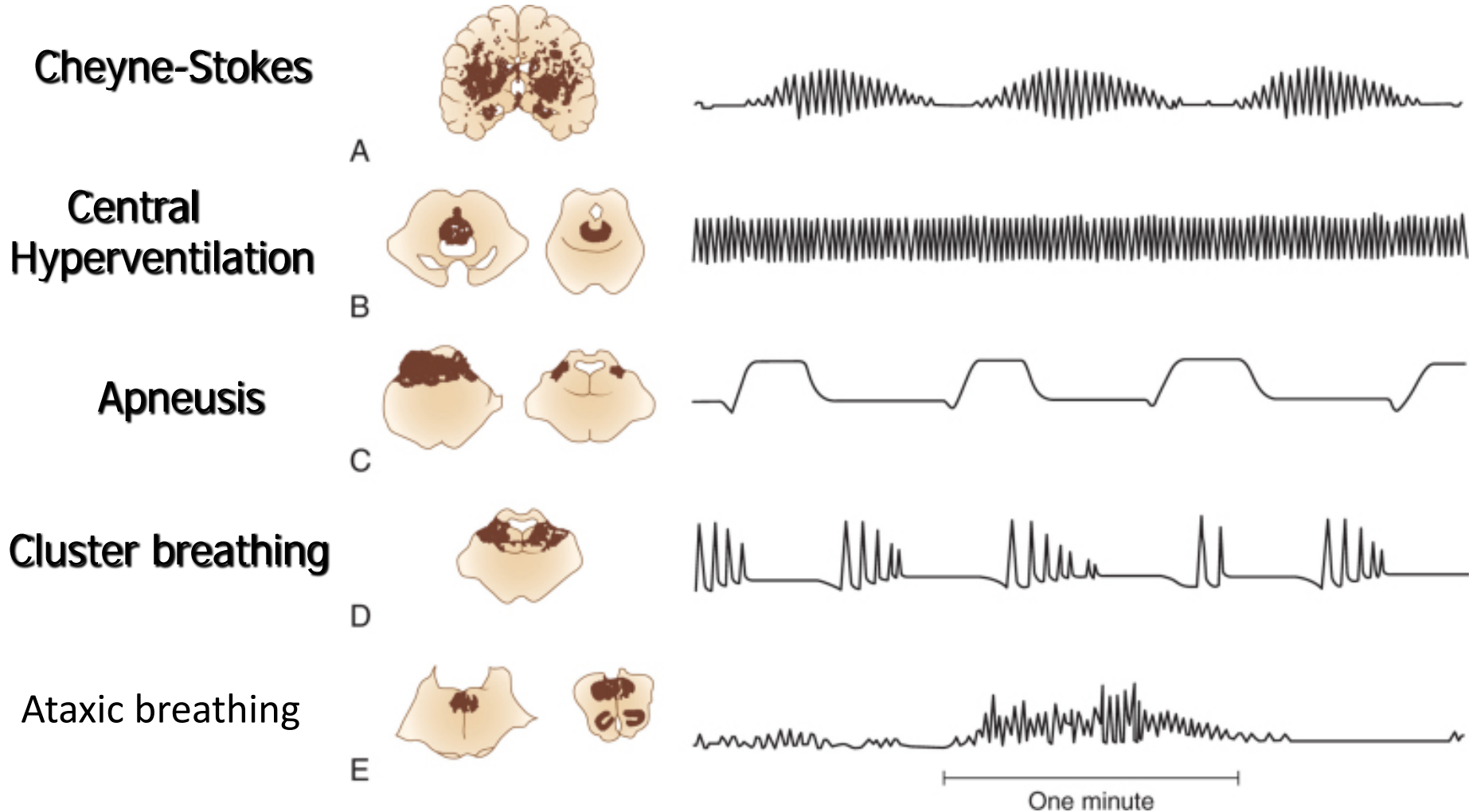
# Neonatal Encephalopathy

## Signs of CNS disease

- Changes in respiratory patterns
  - Central tachypnea (midbrain)
  - Apneusis (pontine)
  - Apnea (> 20 seconds, midbrain)
  - Cluster breathing (high medullary)
  - Ataxic breathing (medulla)
  - Cheyne-Stokes breathing - very rare
- Central hypercapnia



# Central Respiratory Patterns





# Neonatal Encephalopathy

## Signs of CNS disease

- Signs of brain stem damage

- Loss of thermoregulatory control
- Weakness
- Anisocoria (3rd nerve, one side)
- Pupillary dilation (midbrain)
- Pinpoint pupils (pontine)
- Hypotension
- Loss of consciousness (reticular formation)
- Vestibular signs - circling, head tilt
- Facial nerve paresis



# Neonatal Encephalopathy

## Signs of CNS disease

- Seizure-like behavior (tonic/clonic generalized)
  - Marching type behavior (clonic, partial or gen)
  - Abnormal extensor tone (tonic, partial or gen)
  - Seizures
- Coma, death



“Dong”

# Neonatal Encephalopathy Treatment

- Nutrition
  - Not nursing
  - Trophic feeding
  - Parenteral Nutrition
- Respiratory
  - Intranasal oxygen
  - Caffeine
  - Positive Pressure Ventilation
- Seizures
  - Phenobarbital

# “Pong”

## Neonatal Encephalopathy

- Hospital day 2
  - Seizures – resolved with phenobarbital therapy
  - Began ventilation
- Hospital day 3 – standing
- Hospital day 5 – nursing from bottle, more aware
- Hospital day 6 – off intranasal oxygen
- Hospital day 9 – nursing from mare

# “Pong”

## Neonatal Nephropathy

- Creatinine level slow to drop
  - Above normal until hospital day 11
- High fractional excretion of Na
  - As high as 2.18% - normal for neonatal foal <0.3%
  - Still > 1% at discharge (day 20)
- Development of significant edema
  - Persisted until day 6



# Neonatal Nephropathy

- Second most common target - 45%
- Common disease states
  - Mild decrease GFR
  - Mild acute tubular necrosis
  - Mild tubular dysfunction
  - Maldistribution of renal blood flow
- Less common disease states
  - Severe acute tubular necrosis
  - Irreversible acute damage
  - Chronic renal disease



# Neonatal Nephropathy

- Oliguria
- Anuria
- Edema formation
- Fluid overload
- Weight gain
- Persistently elevated Cr
- Birth Cr slow to drop
- Abnormal fraction excretions
- High amikacin trough levels
- Slow response to fluid challenges





# “Pong” Neonatal Gastroenteropathy

- Fetal/neonatal diarrhea
- Retained meconium
- Too much abdominal fill for not being fed
- Abnormal abdominal palpation
  - One loop of bowel thickened wall
- Day 7 began passing feces
  - Frequency > 24 hours
  - Enema dependent
- Day 17 resolved



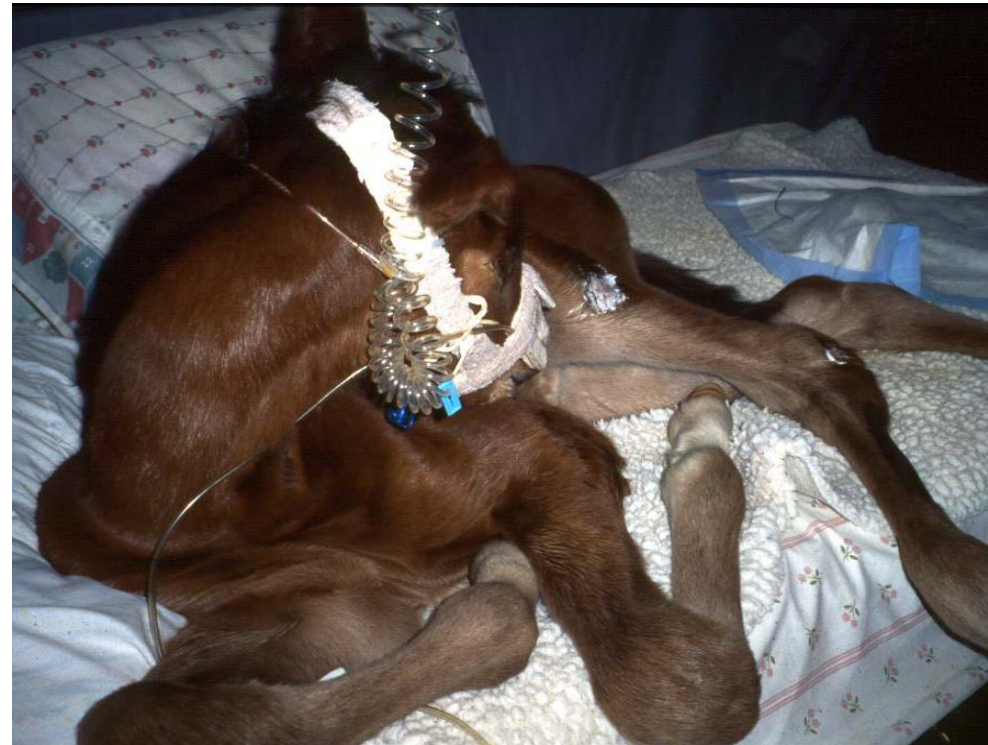
# Neonatal Gastroenteropathy

- Third most common target - 40%
  - Especially when metabolic demands (digestion) are superimposed on cardiopulmonary instability
- Predisposition to sepsis and SIRS
  - Translocation of bacteria through the GI tract



# Neonatal Gastroenteropathy

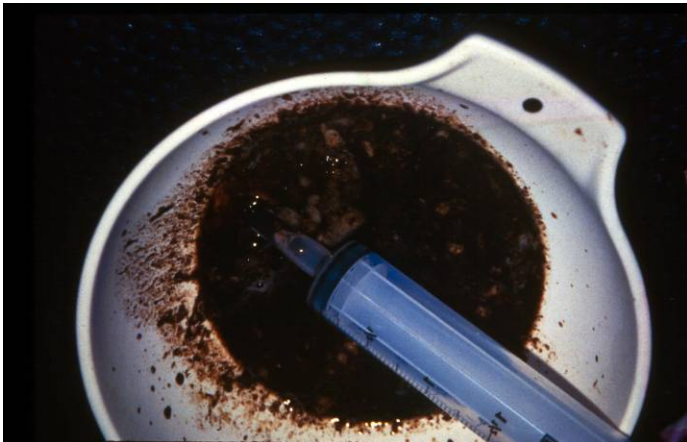
- Dysphagia
- Colic
- Abdominal distension
- Gastric reflux
- Diarrhea
- Constipation
- Dietary intolerance
  - Milk replacer
  - Other specie's milk
  - Frozen mare's milk
  - Fresh mare's milk



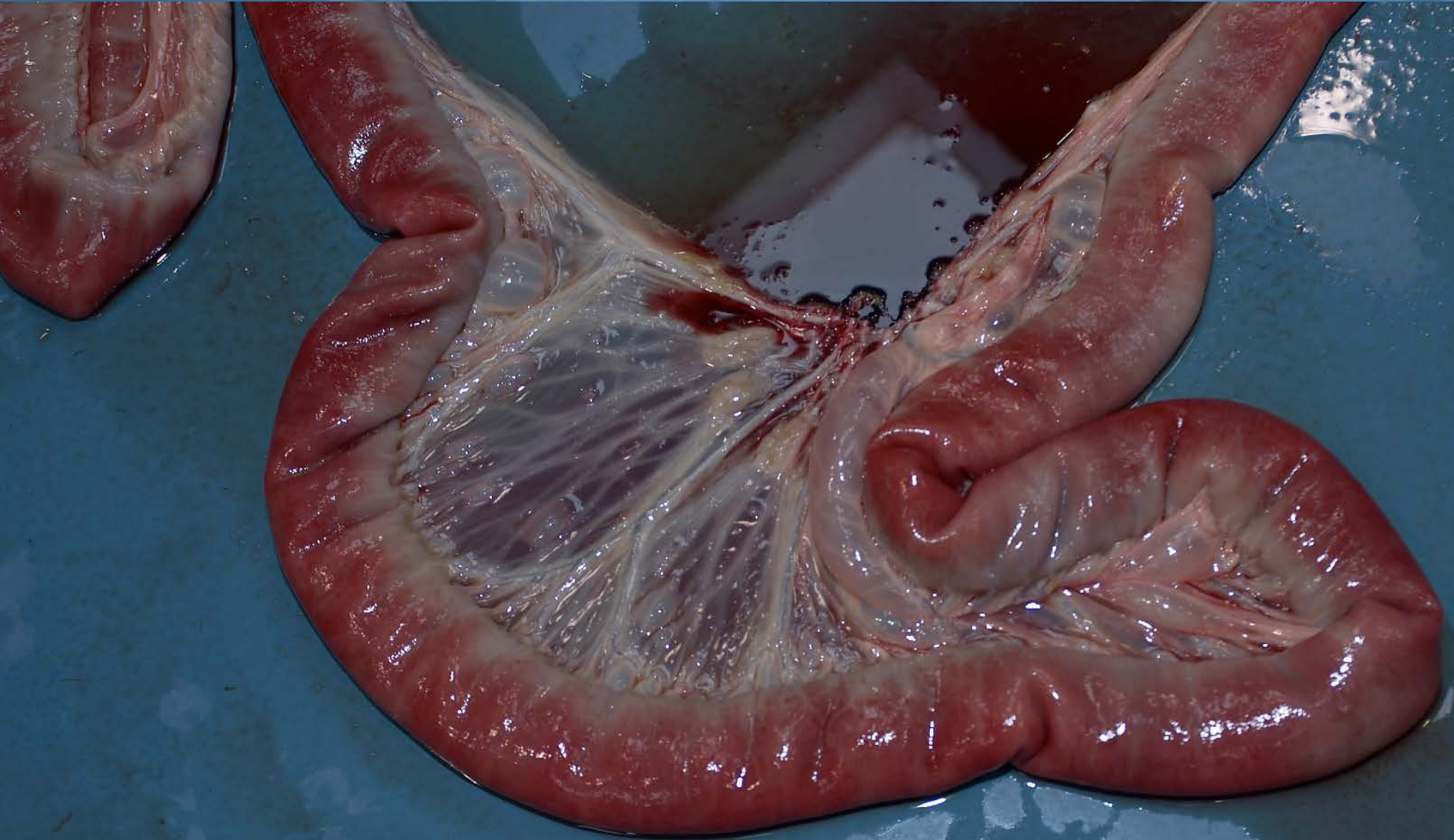
# Neonatal Gastroenteropathy

- Mild indigestion
- Dysmotility
- Ileus
- Diapedesis of blood into the lumen
- Mucosal edema
- Epithelial necrosis
- Development of intussusceptions or structures
- Hemorrhagic gastritis or enteritis/colitis
- Pneumatosis intestinalis

# Neonatal Gastroenteropathy







# Neonatal Syndrome

## Cardiovascular tract

- Less commonly affected – 10 %
- Poorly responsive peripheral vasculature
  - To hypovolemic challenges
  - To endogenous/exogenous adrenergic agents
- Cardiac disease
  - Inappropriate bradycardia
  - Premature ventricular contractions
  - Supraventricular tachycardia
  - Ventricular tachycardia
- Persistent fetal circulation/PPH
- Cardiovascular collapse
  - Refractory hypotension
  - Cardiovascular shock
  - Septic shock



# “Pong”

## Metabolic Maladaptation

- Hypoglycemia at admission – 44 mg/dl
- Hyperglycemic on glucose infusion – 243 mg/dl
  - Glucose diuresis
  - Hyponatremia, hypochloremia, hypokalemia
    - Diuresis, plasma osmotic effects
- Insulin therapy
  - Constant infusion regular insulin IV
  - Begun hospital day 2, weaned day 4

# Neonatal Metabolic Maladaptation

## Signs of Metabolic Disease

- Hypoglycemia
- Hyperglycemia
- Hypocalcemia
- Hypercalcemia
- Hyperlipemia/hyperlipidemia
- Slow response
  - To changing metabolic demands

# Neonatal Syndrome



• NE - *Neonatal Encephalopathy*



• NN - *Neonatal Nephropathy*



• NG - *Neonatal Gastroenteropathy*



• NMM - *Neonatal Metabolic Maladaptation*

• NCM - *Neonatal Cardiovascular Maladaptation*

• NAM - *Neonatal Autonomic Maladaptation*



• NEM - *Neonatal Endocrine Maladaptation*



# “Pog” Problems

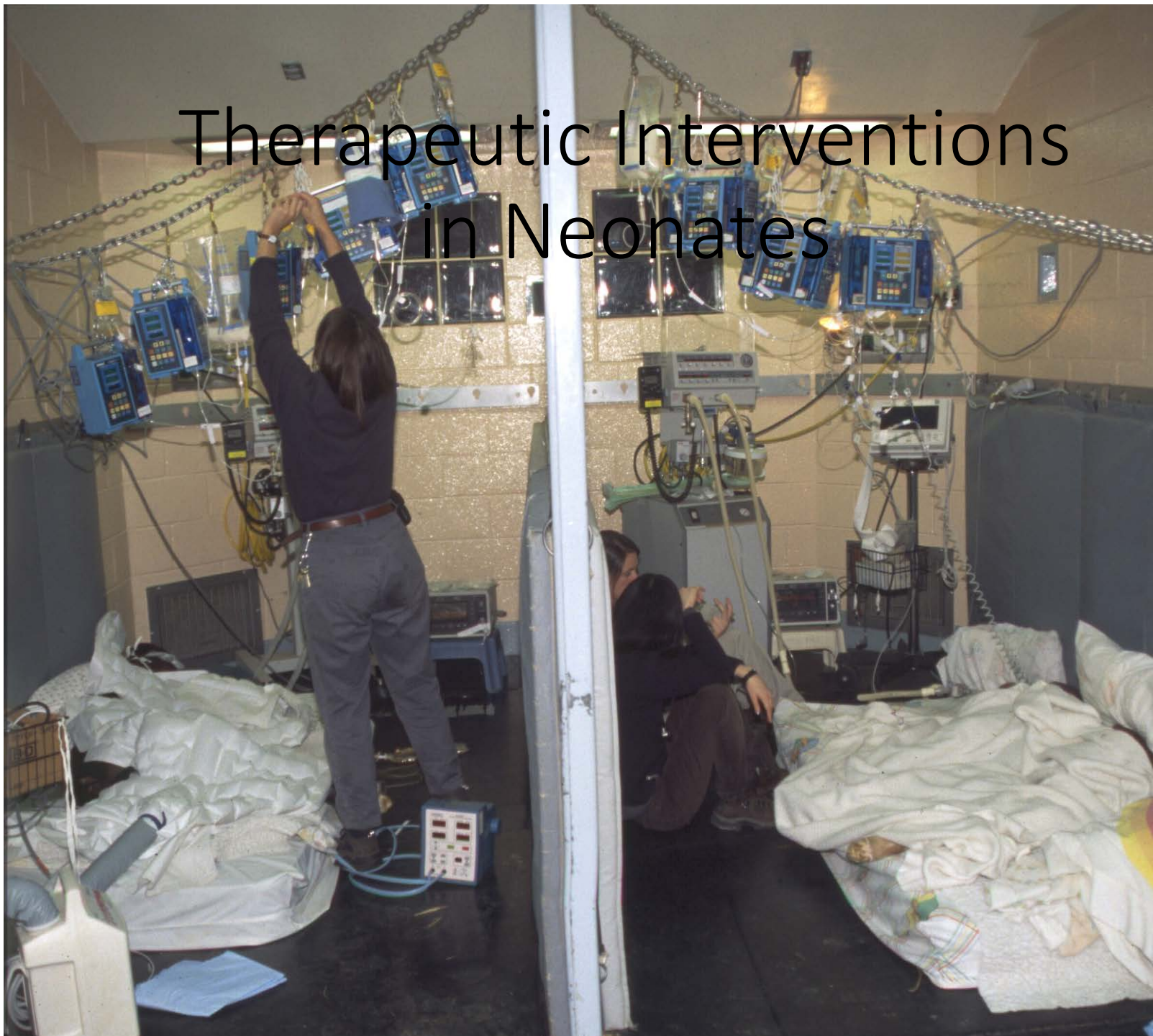
- Sepsis
  - Bacteremia - *Pantoea agglomerans*
- Septic shock
- Neonatal Encephalopathy
  - Central Respiratory failure – ventilation therapy
- Neonatal Nephropathy
- Neonatal Gastroenteropathy

# “Pog” Problems

- Neonatal Metabolic Maladaptation
- Edema
- Urachitis
- Hepatomegaly
- LDN
- Patent Urachus
- Over at knees



# Therapeutic Interventions in Neonates



# Neonatal Syndrome

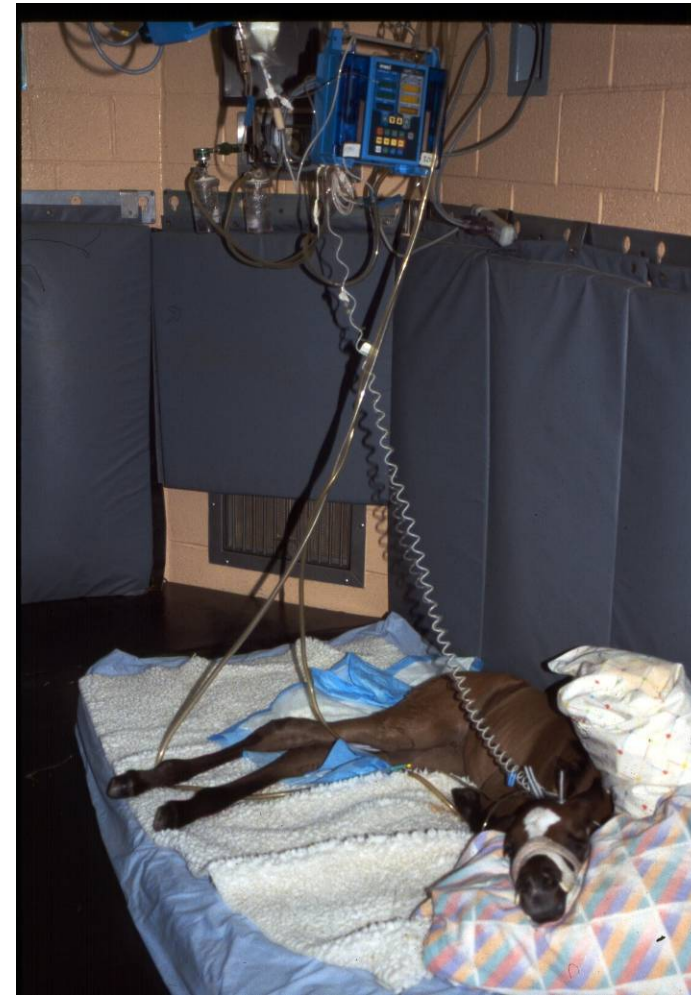
## Clinical Course/Therapeutic Intervention

- As severe organ dysfunction develops
  - Oxygen delivery to the tissues interrupted
  - Progression of more severe disease
- Therapeutic intervention
  - Prevent hypoxic ischemic episodes
  - Support organ system function
    - Allow recovery
  - Prevent secondary sepsis
  - Prevent other complications

# Neonatal Syndrome

## Maintain Tissue Perfusion/Oxygen Delivery

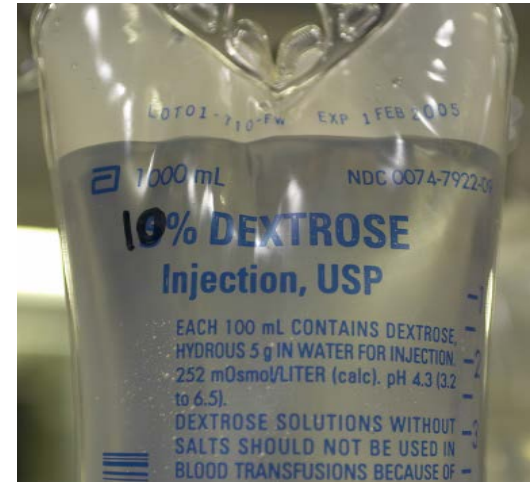
- Adequate cardiac output/perfusion
  - No magic blood pressure value
  - Adequate perfusion reflected by
    - Maintaining urine output
    - Perfusion of the limbs
    - Perfusion of the brain - mental status
    - Perfusion of bowel - GI function
  - Inotrope and pressor therapy



# Neonatal Syndrome

## Maintain Nutrition

- Avoid
  - Catabolic state
  - Hypoglycemia
    - Hypermetabolism
- All compromised neonates
  - Will benefit from glucose therapy
- Hyperglycemia
  - Insulin therapy
- Enteral Nutrition
- Parenteral Nutrition





# NE Therapy

- Support cerebral perfusion
  - Insure volemia
    - Careful fluid replacement
  - Defend perfusion
    - Inopressor therapy
- Insure oxygen delivery
  - Achieve pulmonary O<sub>2</sub> loading
  - Avoid anemia
- Nutritional support
  - Permissive underfeeding





# Therapy

- DMSO
- Mannitol
- Thiamine
- MgSO<sub>4</sub>
- Others



# Seizure Control

Phenobarbital? Midazolam? Others?



# Neonatal Nephropathy

## Therapy for Renal Dysfunction

- Avoid fluid overload
  - Ventral edema
    - Between front legs ("jelly belly")
    - Proximal limbs
    - Back
    - Generalized
  - Monitor body weight at least SID
- Avoid NSAIDs

# Neonatal Nephropathy

## Therapy for Renal Dysfunction

### Fluid restriction

- Most important management tool
- Deliver maintenance fluids or less
  - “Run them dry”
  - Balance nutritional needs/fluid overload
- Watch for onset of diuresis
  - Transition to high output renal failure
  - Initiation of normal renal function



# Neonatal Gastroenteropathy

## Treatment of GI Dysfunction

- Signs of damage lag behind other tissues
- Continued feeding with episodes of hypoxemia
  - May result in further damage
  - Oral feeding undertaken with great care
  - Full nutritional requirements cannot be met enterally
  - Partial parenteral nutrition



# Neonatal Gastroenteropathy

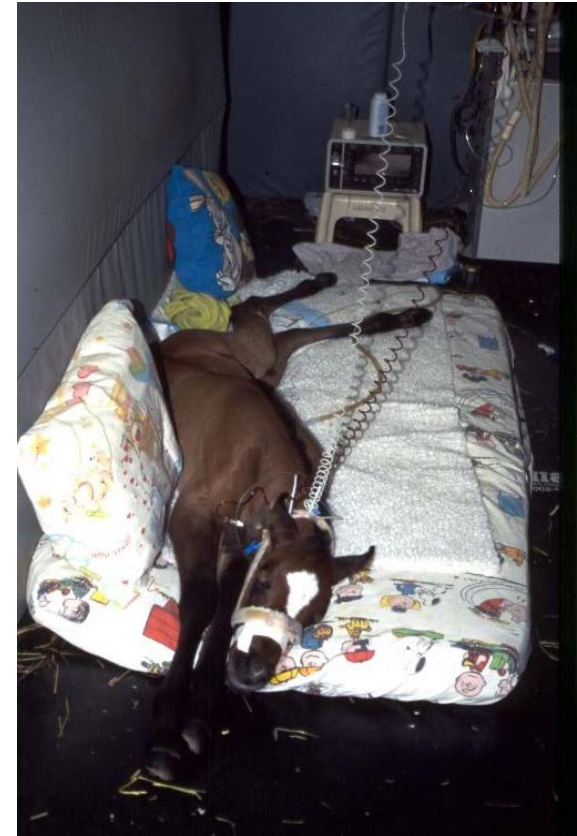
## Treatment of GI Dysfunction

- Important trophic substances in colostrum
  - Only small amounts needed for effect
- Luminal nutrition important to enterocyte health
  - Not feeding increases likelihood of translocation
- Small feedings 1-2 oz QID
  - Fresh colostrum - not refrigerated - best
  - Fresh mare's milk
  - Frozen colostrum or mare's milk
  - Don't use milk replacer

# Neonatal Syndrome

## Recognition/Early Treatment of Secondary Infections

- Very susceptible to infections
- Monitor
  - For localizing signs of infection
  - Repeated blood cultures
- Repeat measurements of IgG
  - Repeated plasma transfusions



“Pog”

## Therapeutic interventions

- INO2
- Fluid boluses
- Dobutamine
- Ticarcillin, clavulanic acid
- Plasma transfusion
- CRI glucose fluids
- Insulin
- Phenobarbital
- Caffeine
- Positive pressure ventilation
- Parenteral Nutrition
- Trophic feedings
- Sucralfate
- Domperidone -- mare
- TMS , Cephalexin
- Bandaging

“Pong”

