

# Acetaminophen overdose

*N*-acetyl-*p*-aminophenol  
(APAP)

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# Objectives

- By the end of this lecture, participants should be able to:
- Know the potential toxic dose of APAP according to age
- Know the symptoms and signs of APAP OD
- Know the indications of NAC therapy

# APAP

- Acetaminophen has been approved for OTC use since 1960
- 1<sup>st</sup> cases of hepatic damage after APAP OD 1966
- Therapeutic dose of acetaminophen is 10-15 mg/kg/dose in children and 325-1000 mg/dose every 4-6 hours in adults, with a maximum of 4g/day

# Toxic dose of APAP

## ■ Children:

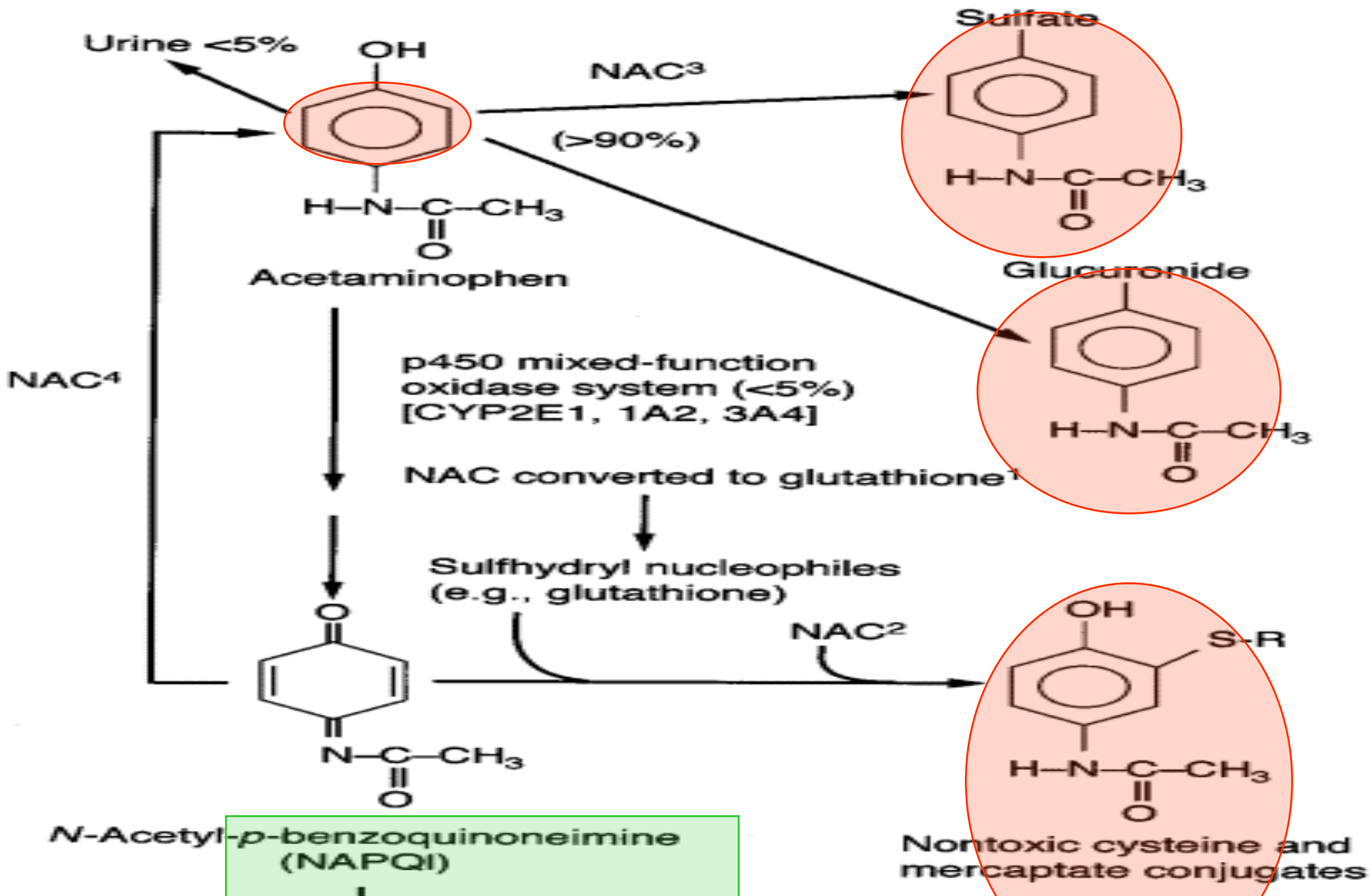
- < 12 months 150 mg/kg
- 1 – 6 y 200 mg/kg
- 1 – 6 y with risk factors 150 mg/kg
- 7 – 12 y 150 mg/kg

## ■ Youth & Adult

- 7.5 - 10 g

# Metabolic Pathways

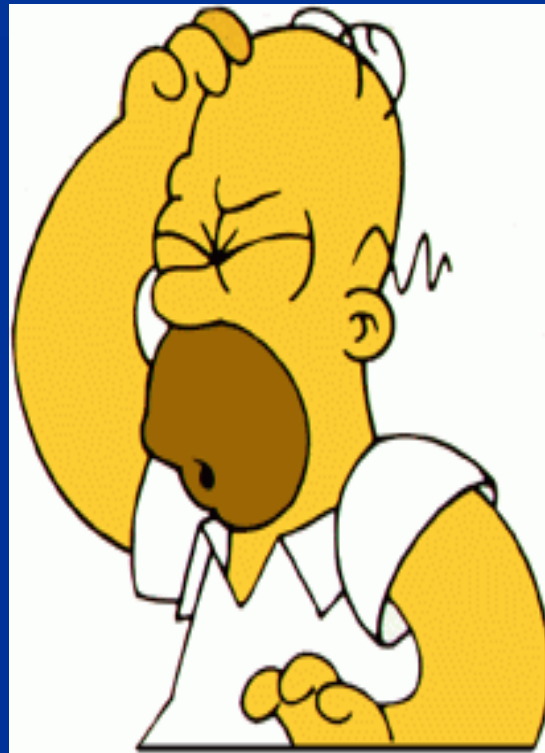
- Hepatic glucuronide conjugation(40-65%)
  - Hepatic sulfate conjugation(20-45%)
- inactive metabolites excreted in the urine.
- 90%
- Excretion of unchanged APAP in the urine (5%).
  - Oxidation by P450 cytochromes (CYP 2E1, 1A2, and 3A4) to **NAPQI** (5-15%)
    - GSH combines with NAPQI
    - nontoxic cysteine/mercaptate conjugates
    - excreted in urine.



Toxicity due to covalent bonding to nucleophilic tissue macromolecules:

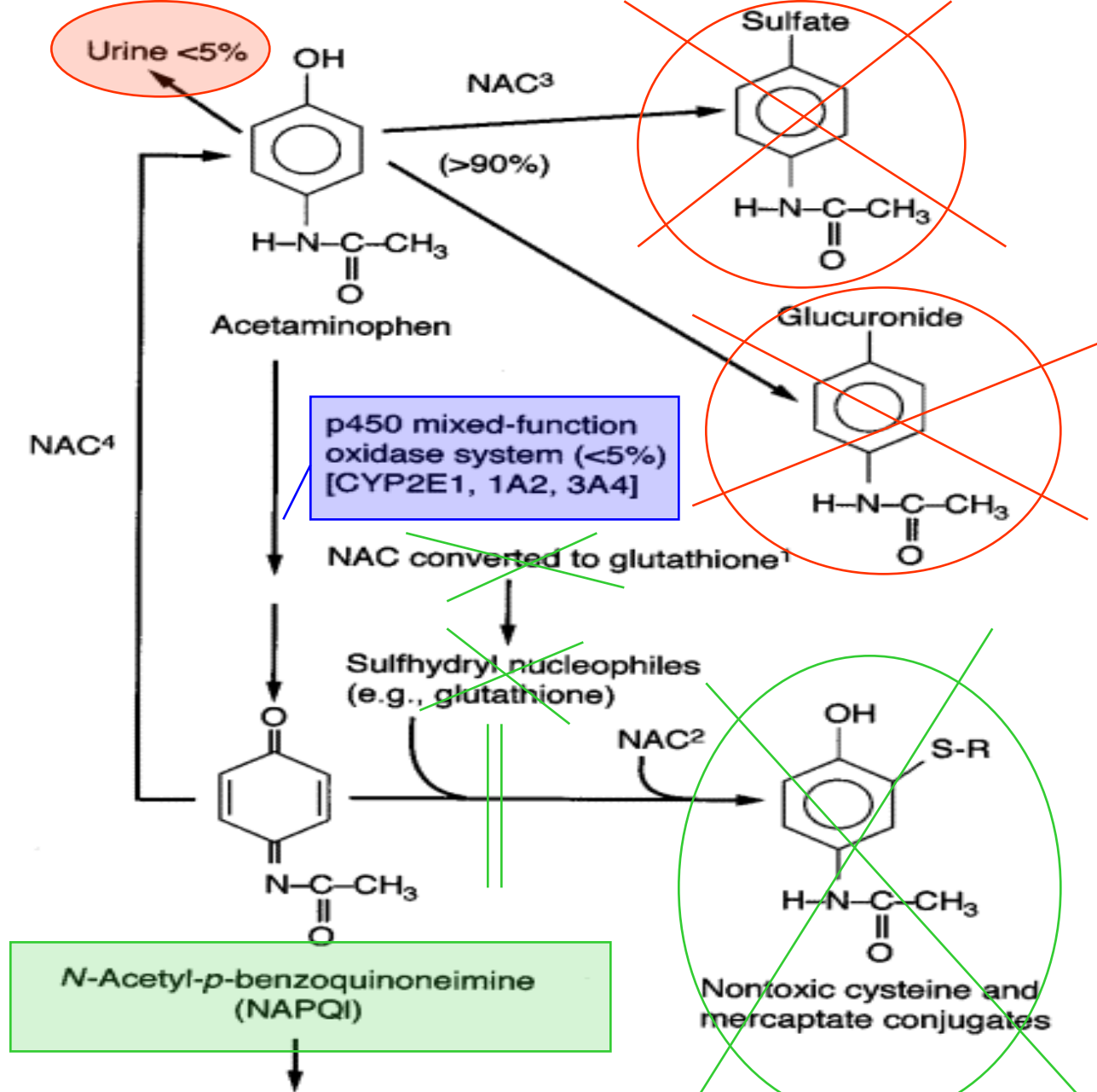
NAC may improve outcome by a variety of mechanisms<sup>1-4</sup>

# What happens in OD ?



- Saturation of glucuronidation and sulfation pathways
- Amount of APAP metabolized by p450 cytochromes to **NAPQI** increases
- Normally NAPQI is detoxified by reduced **GSH** (glutathione) and **thiol-containing substances**
- In OD: rate and quantity of NAPQI formation overwhelms GSH supply and regeneration:
  - elimination of NAPQI prolonged
  - free NAPQI binds critical cell proteins with sulfhydryl groups
  - cellular dysfunction and cell death.
- Animal models: hepatotoxicity when GSH stores fall <30% of baseline





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# Factors which adversely affect APAP metabolism

- Up regulation (i.e. induction) of CYP 2E1 enzyme activity
- Decreased glutathione stores
  - Eating
  - NAC
- Frequent dosing interval of APAP
- Prolonged duration of excessive dosing

(Kuffner et al. 2001)

# Clinical manifestation

- I 0.5-24h n/v, anorexia, asymptomatic
- II 24-48 h resolution of stage I sxs  
RUQ pain, elevation of PTT, INR,  
bili + enzymes (at the latest by 36h)
- III 48-96h coagulopathy, peaking of enzymes,  
acidosis, hypoglycemia, bleeding  
diathesis, jaundice, anuria, cerebral  
edema, coma. ARF in 25% of pts  
with hepatotoxicity
- IV 4-14d resolution

# Diagnosis

- In the patient with a history of APAP overdose, a serum APAP level should be measured between 4 and 24 hours after ingestion
- The value obtained should be evaluated according to the Rumack-Matthew nomogram for determining the risk of hepatotoxicity and the need for NAC therapy

# Toxicological History

- Often incomplete, unreliable or unobtainable
- Sources – Patient, friends, family, EMS, or pill containers
- PMHx, liver/renal disease, concurrent medications, previous overdoses, PSHx, substance abuse

# The **5W's** of toxicology

- **Who** – pt's age, weight, relation to others
- **What** – name and dose of medication, coingestants and amount ingested
- **When** – time of ingestion, single vs. multiple ingestions
- **Where** – route of ingestion, geographical location
- **Why** – intentional vs. unintentional

# Management Guidelines

- **A**irway
- **B**reathing
- **C**irculation
- **D**econtamination
  - AC
- **F**ind antidote
  - NAC

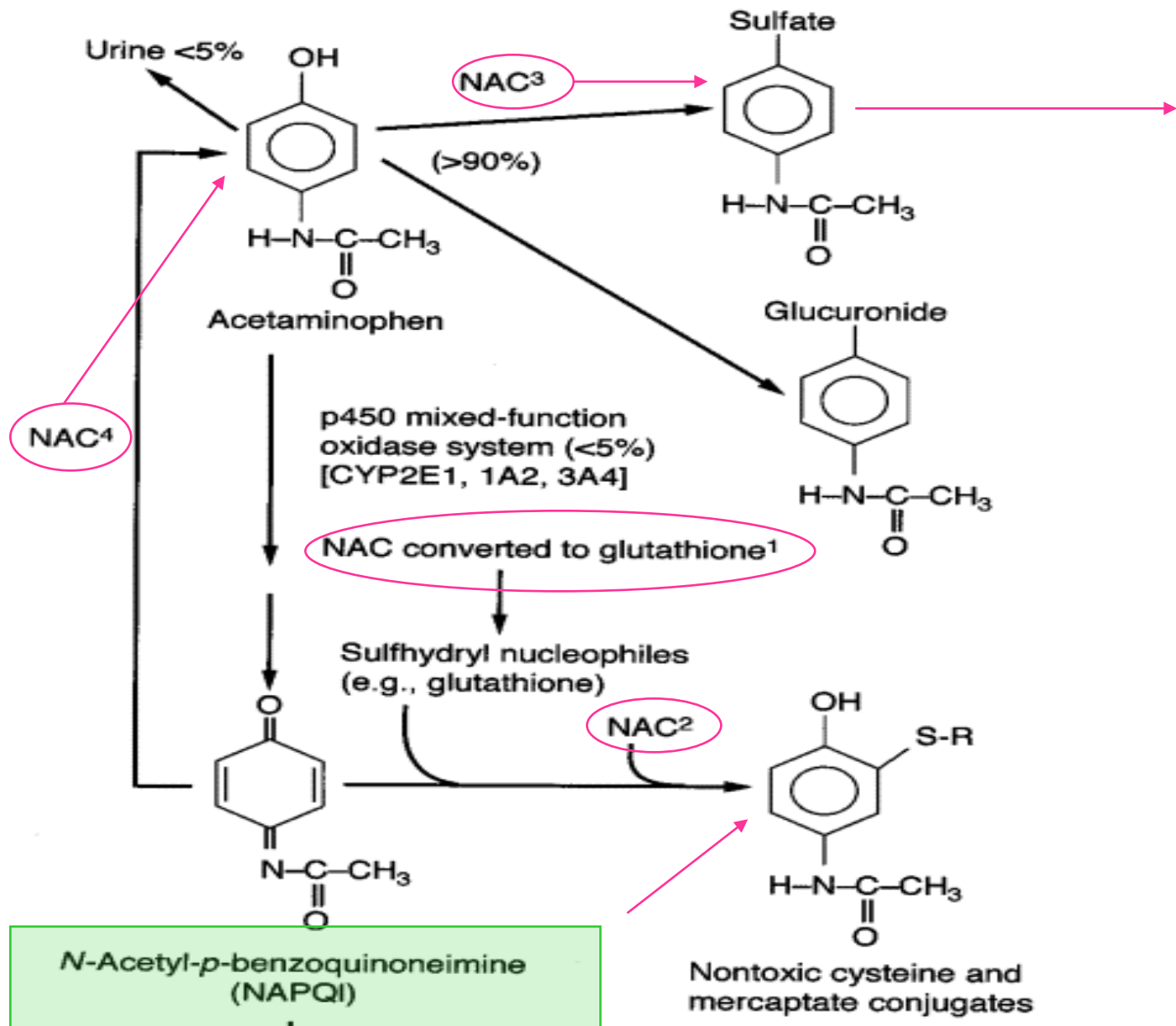
# NAC

- Early → Prevents binding of NAPQI to hepatocytes
- GSH precursor → increases GSH stores
- Increases sulfation metabolism of APAP → less NAPQI formed
- Reduces NAPQI back to APAP (at least in animal models)
- Sulfur group of NAC binds and detoxifies NAPQI to cysteine and mercaptate conjugate (= GSH substitute)



# NAC

- Late (12-24h) → Modulates the inflammatory response
- Antioxidant, free radical scavenger
- Reservoir for thiol groups (i.e. GSH)
- Impairs WBC migration and function → antiinflammatory
- Positive inotropic and vasodilating effects (NO) → improves microcirculatory blood flow and O<sub>2</sub> delivery to tissues
- Decreases cerebral edema formation, prevents progression of hepatic encephalopathy and improves survival

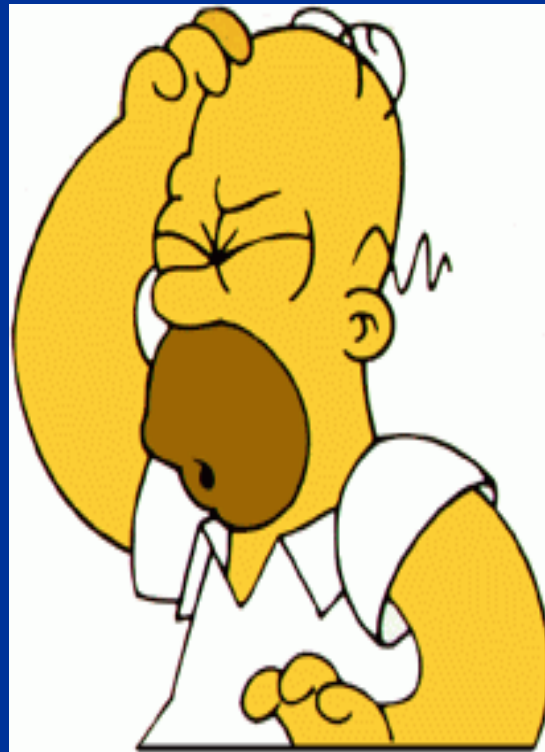


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 NAC may improve outcome by a variety of mechanisms<sup>1-4</sup>

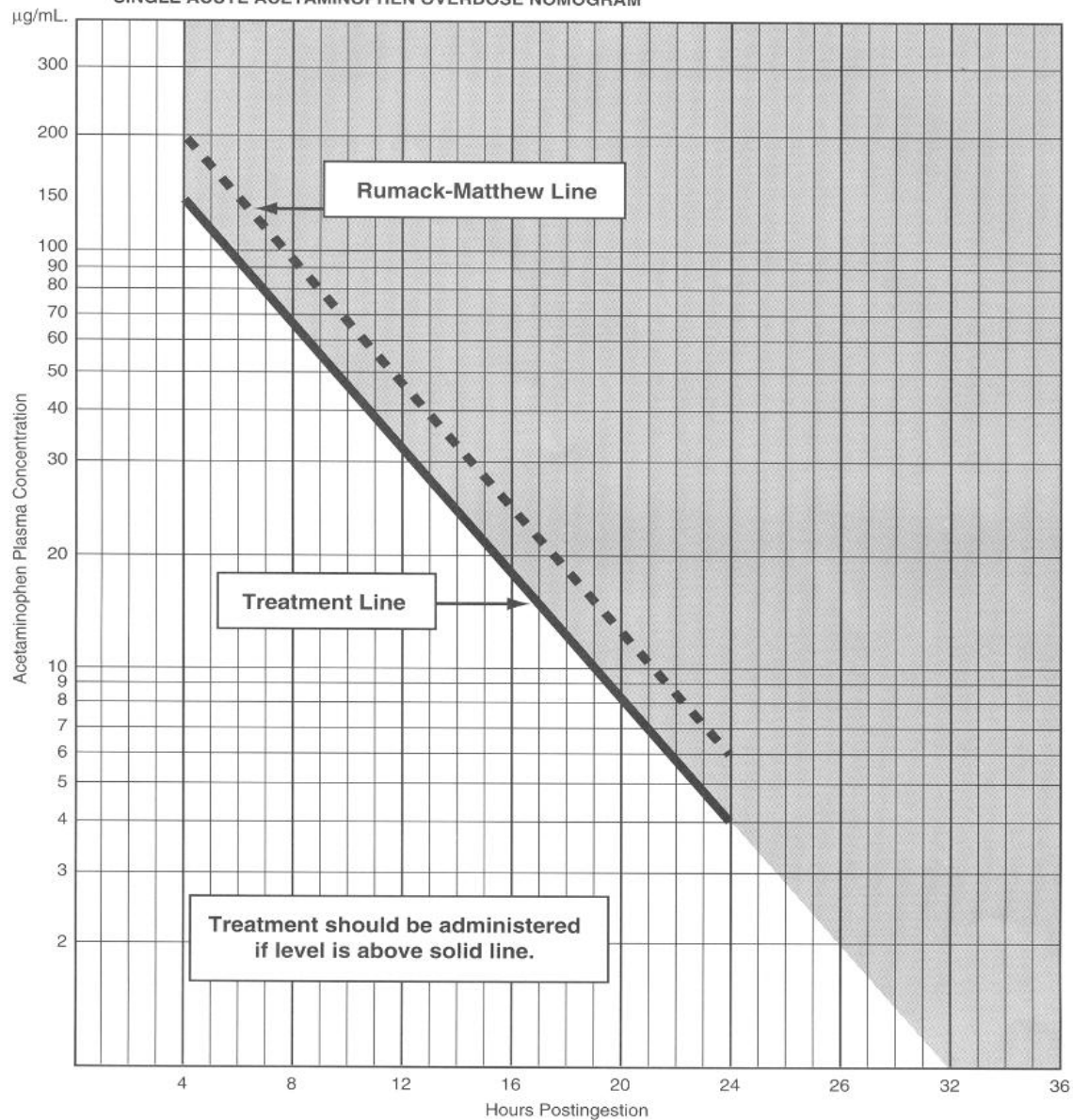
# NAC

- NAC should optimally be given within 8 to 10 hours after ingestion
- More delayed therapy is associated with a progressive increase in hepatic toxicity
- some benefit may still be seen 24 hours or later after ingestion

# What is the Rumack-Matthew nomogram?



### SINGLE ACUTE ACETAMINOPHEN OVERDOSE NOMOGRAM



The nomogram has been developed to estimate the probability of whether a plasma acetaminophen concentration in relation to the interval postingestion will result in hepatotoxicity and, therefore, whether acetylcysteine therapy should be administered.

#### CAUTIONS FOR USE OF THIS CHART:

1. Time coordinates refer to time postingestion.
2. Graph relates only to plasma concentrations following a single, acute overdose ingestion.
3. The Treatment Line is plotted 25% below the Rumack-Matthew Line to allow for potential errors in plasma acetaminophen assays and estimated time from ingestion of an overdose.

# Rumack-Matthew nomogram

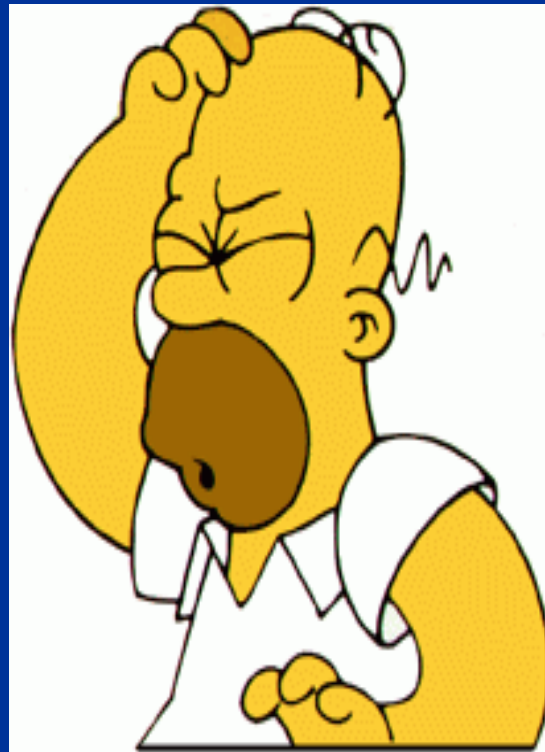
- APAP level to predict which patients will develop an AST elevation  $>1000$  IU/L without antidotal treatment
- Derived from acute ingestion of immediate release acetaminophen
- Begins at 4 h post ingestion
- Recommended line of treatment has been lowered by 25% to increase its sensitivity

**What percent of pts whose APAP level falls above the upper line of the Rumack-Matthew nomogram will develop hepatotoxicity?**

**(defined as elevation of the plasma transaminases above 1,000 U/L)**

**60%**

# When to give NAC?





# Indication for NAC

- APAP level above the treatment line
- Hx of significant APAP ingestion presenting close to 8h (give while waiting for level)
- All APAP ingestions who present late >24h with either detectable APAP or elevated transaminases
- Chronic ingestions (>4g/day in adult, >120mg/d in child) with elevated transaminases
- Hx of exposure and FHF

# Poor prognostic indicators

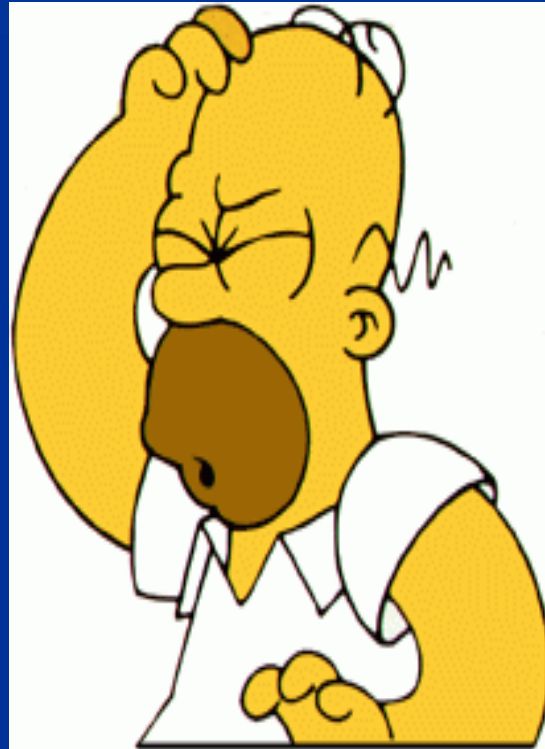
- pH <7.3 (2 days after OD, after fluids)
- Hepatic encephalopathy
- PT >1.8 times normal.
- Serum creatinine >300mmol/L
- Coagulation factor VIII/V ratio of >30

# XR tablets

- Several studies show that elimination of extended and immediate-release acetaminophen are nearly identical after 4 hours.
- some case reports APAP levels falling above the treatment normogram line as late as 11-14 hours post ingestion of the extended-release preparation



# Short cases



- 15 month old child (wt. 10 kg) accidentally took full bottle of Tylenol 60cc(120mg/5cc) 30 mint ago. Clinically looked well. What will be your treatment plan:
  - a) Give Ipecac STAT
  - b) Give 1g/kg activated charcoal
  - c) Insert OGT and perform gastric lavage
  - d) Should be observed for 4h then to do drug level
  - e) None of the above

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- A mother brought her 4 M (5 kg) old son who was febrile for the last 3 days . She was giving him Tylenol (120mg/5 ml) 7ml every 4 h for the last 3 days, she found him today more lethargic, vomiting occasionally, clinically, ill looking slightly jaundiced, afebrile, no meningeal signs, mild injected throat, CSF was obtained & was not suggestive of meningitis. What will be your next step:
  - a) Obtain CBG, LFT, PT, PTT, INR, drug level if abnormal start NAC
  - b) Give activated charcoal immediately
  - c) Admit for observation
  - d) D/C home, most likely it is related to current URTI



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- 19 y old girl brought to ED with GCS 8 following drug ingestion (empty bottle of Tylenol was found in her room). What will be your first response
  - a) 1g/kg activated charcoal STAT
  - b) Orotracheal intubation
  - c) Observation for 4 h
  - d) Do CBC, CBG, PT, PTT, INR, Drug level
  - e) NAC loading dose followed by infusion over 24 h

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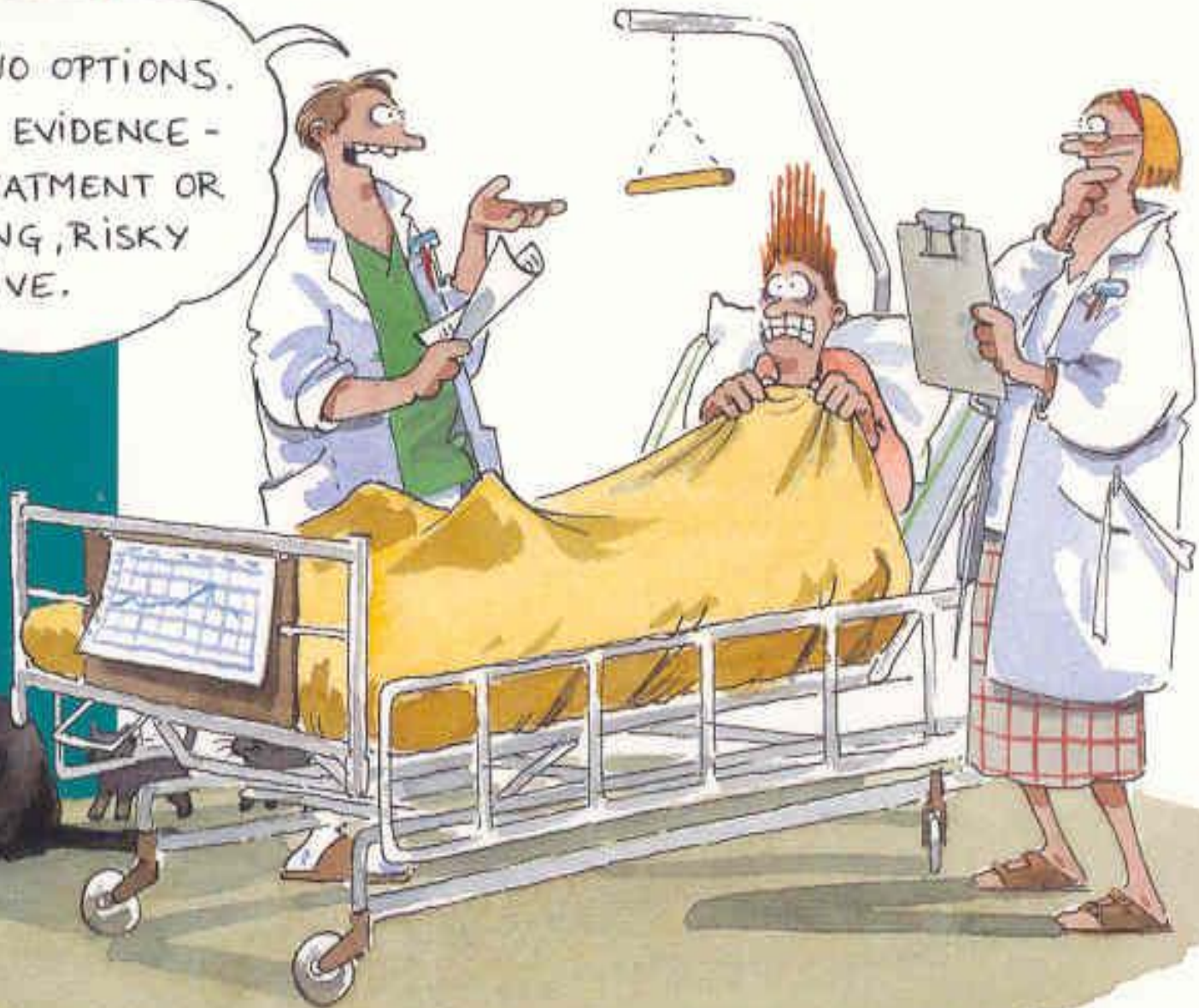
- 3 y old boy with accidental Tylenol ingestion on NAC for high drug level, after 48 h course LFT ,INR are high. What will be your recommendation:
  - a) D/C NAC if drug level undetectable
  - b) D/C NAC and repeat LFT, INR, drug level after 4h
  - c) Continue on NAC until all his labs become normal
  - d) D/C NAC, most likely it is secondary to concurrent viral illness

- 20 yr old pregnant girl ingested 20g of Tylenol in a suicidal gesture 36h ago because she found out it is too late for her to have an abortion. Her APAP is  $<10$  and her AST is 90
- How will you manage her medically?
- She asks you whether her baby will have any defects?

**Questions ??**



WE HAVE TWO OPTIONS.  
EITHER AN EVIDENCE -  
BASED TREATMENT OR  
AN EXCITING, RISKY  
ALTERNATIVE.



Christine Almer



# Paracetamol (acetaminophen)

**poisoning** Vale, JA, Proudfoot, AT. *Lancet* 1995; 346:547

- No deaths in 169 patients with a treatment delay below 10 hours
- In contrast, 200 patients treated at 10 to 24 hours had a 2.0 to 7.4 percent mortality, which was still lower than the 5.3 to 10.7 mortality in 85 patients who received only supportive care.
- There was a 1.6 to 10 percent incidence of liver damage (defined as a plasma ALT or AST level above 1000 IU/L) when the treatment delay was less than 10 hours
- Comparable values were 27 to 63 percent in patients treated at 10 to 24 hours and 58 to 89 percent in those receiving supportive care

# Improved outcome of paracetamol-induced fulminant hepatic failure by late administration of NAC

*Lancet 1990 Jun 30;335(8705):1572-3]*

- The influence of NAC, administered at presentation to hospital, on the subsequent clinical course of 100 patients who developed APAP-induced fulminant hepatic failure was analyzed retrospectively
- Mortality was 37% in patients who received NAC 10-36 h after the overdose, compared with 58% in patients not given the antidote
- In patients given NAC, progression to grade III/IV coma was significantly less common than in those who did not receive the antidote (51% vs 75%)