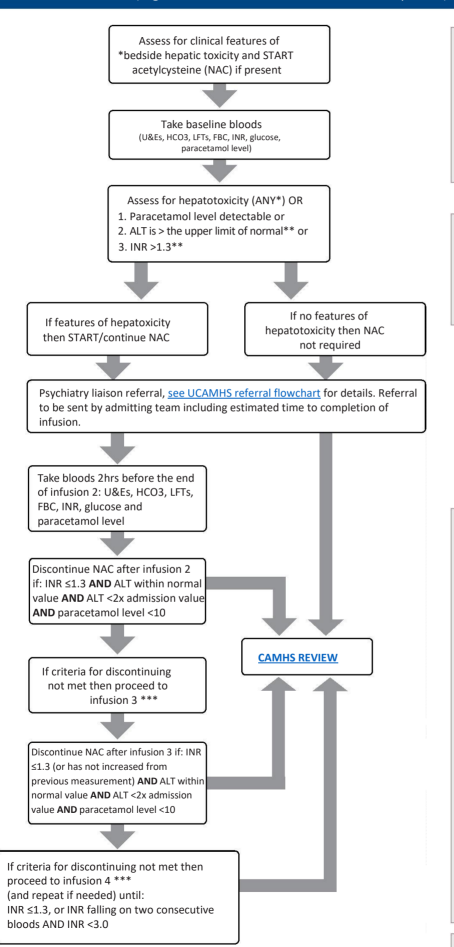
### Paracetamol overdose presenting >24hrs

(Ingested total overdose in ≤1 hour time period)



- \*Clinical judgement required
- Bedside hepatic toxicity: Jaundice, tender liver, hypoglycaemia, encephalopathy, unexplained lactic acidosis.
- Ensure no doubt about time of ingestion or type.
- If uncertainty then treat and review with bloods.
  - \*\*Clinical judgement required
- Some patients have a chronically raised ALT/INR.
- Review old LFTs/INRs and if chronic derangement discuss with a senior clinician before proceeding to NAC.

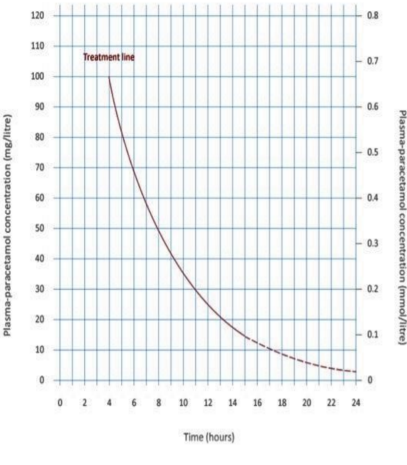
#### **Blood monitoring**

- Checking a paracetamol level 2hrs before the end of bag 2 is NEW for this protocol.
- U&E, HCO3, glu, LFTs, FBC and INR should be done 2hrs before the end of each infusion 2.
   Ensure results are READY for the end of the infusion.
- If unable to achieve blood sampling at the correct time and a delay of >90 minutes is predicted then proceed to the next infusion to avoid prolonged omission of NAC. Bloods should be checked at the earliest opportunity and discontinuation criteria referred to.
- Capillary Blood Glucose (CBG) 6 hourly while on NAC.
- If rapid or progressive biochemical deterioration then discuss with senior and consider referral to regional transplant centre.
- IV NAC can be associated with minor rise in INR without an acute liver injury. (see guideline for details)

\*\*\* Bloods should be done 2 hours before the end of infusion 3 and 4.

# Paracetamol treatment nomogram and 12-hr shortened N-acetylcysteine dosing schedule (SNAP protocol)





Reproduced courtesy of MHRA

If unclear which of the five paracetamol overdose protocols to follow, discuss with ED / paeds Reg / Cons.

In situations where paracetamol levels will be used to determine need for acetylcysteine (refer to appropriate protocol), plot the measured plasma concentration (in mg/L) against the time since ingestion. If plasma level falls above the line then give acetylcysteine as detailed below.

The nomogram is less accurate between 15-24 hours and accurate ingestion time is even more vital.

Actual weight should be used for calculating both the toxic dose and the acetylcysteine dose - up to a maximum of 110 kg

Reactions to acetylcysteine include flushing, nausea & vomiting.

Please use 'Acetylcysteine Antidote Adverse Effects – Features & Management' guidance to document any adverse events and guide further management.

Hypersensitivity and anaphylactoid reactions with acetylcysteine are not contraindications as the benefit of treatment still outweighs the risk of not treating.

True anaphylaxis is rare with acetylcysteine but can be managed by stopping the infusion and then restarting at a slower rate.

Table 1. 12-hr shortened N-acetylcysteine dosing schedule (SNAP protocol).

Regimen	First in	fusion	Second (& extended) infusion			
Infusion fluid	200mL sodium chloride	0.9% or 5% glucose	1000mL sodium chloride 0.9% or 5% glucose			
Preparation	Use 250mL infusion bag and add required volur		Add required volume of acetylcysteine to 1000mL infusion bag			
Duration of infusion	2 ho	ours	10 hours			
Drug dose	100mg/kg acc	etylcysteine	200mg/kg acetylcysteine			
Weight (kg)	Weight (kg) Ampoule volume (mL)		Ampoule volume (mL)	Infusion rate (mL/h)		
30-39	18	109	35	104		
40-49	23	112	45	105		
50-59	28	114	55	106		
60-69	33 117		65	107		
70-79	38	119	75	108		
80-89	43	122	85	109		
90-99	48 124		95	110		
100-109	53	127	105	111		
≥ 110	55	128	110	111		

Each ampoule = 200mg/mL acetylcysteine. Dose calculation based on weight in middle of band. Ampoule rounded up to nearest whole number.

Acetylcysteine Prescribing and Administration Chart for 12-hr shortened N-acetylcysteine dosing schedule (SNAP protocol) – **RHC Glasgow** 

### Infusion 1 & 2 only

Please ensure that acetylcysteine is also prescribed on the patient's HEPMA Kardex.

Name:
Address:
DoB:
CHI:
Affix patient data label

Weight:....kgs (DO NOT USE If <30kg or patient <6 years of age)

Infusion 1 Acetylcysteine			steine 100n	ng/kg over 2	2 hours					
Prescription					Preparation	Administration checks				
Date	Time	Dose (mL)	Diluent (200mL)	Infusion rate (mL/hr)	Prescriber's signature	Prepared/ Checked by	Date Time	Volume remaining (mL)	Volume infused (mL)	Checked by
				75						
Comme	Comments: Stopped by:									
			Date:	Time	Signature					

Infusion 2 Acetylcysteine 200mg/kg over 10 hours										
Prescription						Preparation	Administration checks			
Date	Time	Dose (mL)	Diluent (1000mL)	Infusion rate (mL/hr)	Prescriber's signature	Prepared/ Checked by	Date Time	Volume remaining (mL)	Volume infused (mL)	Checked by
Comments: Stop			Stopped b	Stopped by:						
				Date:	Time	Signature				

Extended treatment

If extended treatment with acetylcysteine is required (see clinical guideline), continue at the dose and infusion rate used for the second infusion and prescribe.

Recheck U&Es, bicarbonate, LFTs, FBC and INR 2 hours before the end of infusions 3 and 4 to assess the need to continue.

Refer to appropriate protocol regarding discontinuation of extended treatment





## <u>Acetylcysteine Antidote Adverse Effects – Features & Management</u>

REACTION to acetylo			COIVIPLICATIONS of paracetamol ingestion							
None		Wheeze			Abnormal liver function		Encephalopathy			
Flushing		Hypotens	sion		Acute kidney injury		Haemorrhage			
Vomiting		Other:			Hypoglycaemia		Other:			
Rash		Specify.			Acidosis		Specify			
Date and time of reaction Initial					Date and time of reaction Initial					
MANAGEMENT OF SI	DE EFFEC	TS								
<ul> <li>N-acetylcysteine may cause anaphylactoid reactions in 2% of cases with this protocol. Flushing, pruritus, rash, hypotension, angioedema, bronchospasm and vomiting are most common.</li> <li>Reactions can be managed by stopping the infusion. Consider chlorphenamine for flushing/itch, nebulised salbutamol if there is bronchospasm and ondansetron if there are GI side effects.</li> <li>Restart the infusion once the reaction has resolved at half the rate to completion of infusion.</li> <li>Previous reaction is NOT a contra-indication to N-acetylcysteine and cases should receive treatment if indicated. Reactions are now considerably less common with the 12-hour SNAP protocol compared to standard regimes.</li> </ul>										
Ondansetron oral or Body weight	· IV slow	(over2mi	ins) injection	a and vomiting) - Age 6 mo	nths-1	6 years				
Up to 10kg			Dos	ng three times daily						
10 - 40kg			1	ng three times daily						
41kg and above			1	8mg three times daily						
Chlorphenamine ORAL (Rash and itch)										
Age				Dos	se					
1-23 months				1m	g twice perday					
2-5 years				1m	g 4-6 hourly (maximum 6	mg pe	erday)			
6-11 years				2m	g 4-6 hourly (maximum 1	2mg p	perday)			
12-16 years			4mg 4-6 hourly (maximum 24mg perday)							
Chlorphenamine IVI	NJECTIC	<u>)N</u> (Rash a	nd itch)							
Age				Dos	se					
1-5 months 2					250 micrograms/kg (maximum four times daily)					
6 months - 5 years			.5mg (maximum four times daily)							
6 - 11 years				5m	g (maximum four times da	aily)				
12 - 16 years 10n					ng (maximum four times daily)					