PARACETAMOL



Paracetamol IS A HIGH-RISK MEDICINE USE WITH CAUTION AND ENSURE THE DIRECTIONS WITHIN THIS PROTOCOL ARE FOLLOWED CAREFULLY Areas where **Protocol/Guideline SESLHD** Facilities applicable Medical Officers, Nurse Practitioners, Nurses/Midwives (under Authorised approved SESLHD Nurse/Midwife medicine protocols) **Prescribers:** Adverse events associated with paracetamol toxicity have been **Important Safety** associated with: Considerations The concurrent use of multiple paracetamol-containing products and confusion between different strengths, formulations, and route of administration Incorrect dosing including using Actual Body Weight instead of Ideal Body Weight (IBW) in obese paediatric patients and prescribing standard dosing regimens in patients with identified risk factors for toxicity Inadequate review of treatment Accidental overdose through ongoing administration of regular and PRN paracetamol Electronic Medication Management systems provide safeguard (e.g., prescribing and administration alerts) to assist with safe prescribing and administration of paracetamol where eMM is implemented. Hybrid (electronic and paper) medication management systems require caution to avoid duplication of doses or dose administration within 4 to 6-hour dosing interval. Information about paracetamol administration must be included in handover. Prior to prescribing or administering paracetamol clinicians MUST: 1. ascertain if paracetamol has been recently ingested, 2. check that no other formulations of paracetamol are concurrently prescribed or administered, 3. ensure the time interval between doses are appropriate and that the administration of the dose will not exceed the safe maximum daily dose of paracetamol (from all sources including combinations containing paracetamol). This protocol sets out measures to reduce the likelihood of adverse events with use of paracetamol in SESLHD facilities.

Medicine Guideline for the Safe Use of

PARACETAMOL



SAFETY ALERT	UPDATED: Critical disruption to supply – Intravenous (IV) paracetamol solution for injection/infusion
	Stock of IV paracetamol solution for injection/infusion is reserved for patients in whom oral , enteral , or rectal routes of administration are not appropriate (for example, patients who are nil by mouth). Regular review of IV paracetamol orders is to occur every 6 to 8 hours , with the view of de-escalating to an alternative dose form as soon as possible.
	Clinicians are to consider the use of alternative routes of administration or alternative analgesic agents for the treatment of mild-moderate pain where clinically appropriate. Alternative agents may include non-steroidal anti-inflammatory drugs (NSAIDs). Advice may need to be sought from the local Acute Pain Specialist team.
Indication for use	Consider non-pharmacological intervention prior to paracetamol use.
	 Paracetamol is an effective analgesic and antipyretic agent and may be used as first line therapy for: mild to moderate pain the symptoms of fever, when temperature is above 37.5°C
	The <u>SESLHD Medicines Formulary</u> outlines the paracetamol preparations available for inpatient initiation or continuation and the associated prescribing restrictions.
Clinical condition	Before prescribing paracetamol a full medical history and medication history (to include over the counter products and complementary medicines) should be obtained to determine the potential for paracetamol toxicity and/or any possible adverse drug reaction and to identify paracetamol intake from all sources.
	According to emerging evidence paracetamol use during pregnancy may influence premature closure of the foetal ductus arteriosus; it is recommended that use in pregnancy is limited to the minimum dose and duration clinically necessary.

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Proposed Place in Therapy	Oral route Oral paracetamol is recommended for first line use wherever possible.
	Rectal route Rectal paracetamol is effective and indicated when oral dosing is not possible. Absorption of paracetamol given rectally is erratic and the time taken to achieve maximum concentration is unpredictable. Oral dosing should be resumed as soon as possible.
	Intravenous route Intravenous (IV) paracetamol is reserved for short-term management of mild to moderate pain or fever when oral/NG or rectal administration is not feasible, limited to patients who are nil by mouth or under specialist supervision in anaesthesia, intensive care, or pain management. Due to the critical disruption to supply its use should be reviewed prior to every dose (i.e., every 6 or 8 hours), with a prompt transition to oral, enteral, or rectal administration when possible.
Adjunctive Therapy	Paracetamol is first-line analgesic in adults and children for acute nociceptive pain because of its favourable adverse effect profile. It can be used alone, or as a component of multimodal analgesia. Multimodal analgesia combines analgesic agents with different mechanisms of action which can result in synergistic effects while reducing individual drug dosages. Most commonly, paracetamol is combined with a non-steroidal anti-inflammatory drug (NSAID) or an opioid regimen.
Contra-indications	 Patients with a previous history of hypersensitivity to paracetamol or to any of the product's excipients. IV Paracetamol: Severe liver disease Rectal Paracetamol: Avoid use in neutropenic patients



Precautions	Hypovolemia: Use the IV formulation with caution in patients with severe hypovolemia (e.g., due to dehydration or blood loss).
	Clinical judgement should be used to adjust dose/frequency of paracetamol for patients with any of the following potential risk factors that may increase the risk of acute liver injury: systemic sepsis/febrile illness (particularly in children) prolonged fasting, vomiting or dehydration chronic malnutrition hepatic impairment renal impairment sustained administration of high doses chronic, excessive alcohol use frail elderly patients underweight patients prolonged use of IV preparations
	Chronic use of interacting medicines Warfarin:
Important Drug Interactions	The anticoagulant effect of warfarin may be enhanced by prolonged regular use of paracetamol with increased risk of bleeding; occasional doses have no significant effect. Warfarin dosage may require reduction if paracetamol and warfarin are taken concurrently for a prolonged period of time.
	Other hepatotoxic medications:
	The risk of paracetamol toxicity may be increased in patients receiving other potentially hepatotoxic drugs or drugs that induce liver microsomal enzymes such as carbamazepine, phenytoin, rifampicin, and isoniazid.
	Probenecid: Probenecid may increase the serum concentration of paracetamol by inhibiting their hepatic glucuronidation.
	Cholestyramine: Cholestyramine reduces the absorption of paracetamol if given within 1 hour of paracetamol.



Formulati	on	Dose and freq	uency	Max dose in 24
	Weight			hours
Oral /	> 50kg	1 g every four hours	to six	Maximum of 4 doses in 24 hours
Rectal	≤ 50 kg and > 33 kg	15 mg/kg/dose four to six hou	•	60 mg/kg/day (not exceeding 3g)
	≤ 33 kg and > 10 kg	15 mg/kg/dose four to six hou	rs	60 mg/kg/day (not exceeding 2g)
Oral contr (665 mg)	rolled release	2 tablets every 8 hours swallo whole.		Maximum 6 tablets (3990 mg) daily
	Weight > 50kg	1 g every four	to six	Maximum of 4 doses in 24
IV	≤ 50 kg and > 33 kg	15 mg/kg/dose four to six hou	•	hours 60 mg/kg/day (not exceeding 3g)
	≤ 33 kg and > 10 kg	15 mg/kg/dose four to six hou		60 mg/kg/day (not exceeding 2g)
Children	3 months to 1'	1 voars *		
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Route	Dose and 15 mg/kg/ four to six exceed 1 In acute in	frequency dose [#] every hours (do not g per dose) njury or surgery bading dose of may be	Maximu 24 hour of 48 ho Subseq maximu	im of 90 mg/kg in rs for a maximum ours uent days: im of 60 mg/kg in
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Route Oral Rectal IV # for obese	Dose and 15 mg/kg/ four to six exceed 1 In acute in an initial lo 20mg/kg [#] considered Loading d 40mg/kg [#] every 6 to Round dos nearest su strength 15mg/kg [#] (do not ex- dose)	frequency dose [#] every hours (do not g per dose) njury or surgery bading dose of may be d. ose 30- then 20mg/kg [#] 8 hours. se down to uppository every 6 hours xceed 1 g per	Maximu 24 hour of 48 hour Subseq maximu 24 hour After 8 o review r Do not total in Maximu 24 hour	im of 90 mg/kg in rs for a maximum ours uent days: im of 60 mg/kg in rs days: specialist recommended exceed 4 grams 24 hours im of 60 mg/kg in



Duration of therapy	Paracetamol requires regular medical review to ensure treatment continues to be appropriate
	Oral and rectal paracetamol for acute pain or symptomatic high fever: Review no later than 24 hours after commencement and at least every 48 hours thereafter.
	IV paracetamol: Review prior to every dose (i.e., every 6 or 8 hours) and replace with enteral paracetamol at the earliest opportunity. If dosing for longer than 48 hours, monitoring of liver function tests, including International Normalised Ratio (INR) should be carried out.
	Patients receiving paracetamol for chronic pain: Review no later than 48 hours after commencement, then as required. Care should be taken when prescribing paracetamol for extended periods of time in children, frail elderly and/or patients with risk factors for toxicity.



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Prescribing Instructions	 Before prescribing paracetamol (including under a nurse/midwife- initiated protocol), a full medical history, including medication history, should be obtained from the patient or their carer to identify factors with the potential to increase the risk of paracetamol toxicity. Prior to prescribing, clinicians are to ascertain if paracetamol has been recently ingested, check that no other formulations of paracetamol are concurrently prescribed or administered, ensure the time interval between doses are appropriate and that the administration of the dose will not exceed the safe maximum daily dose of paracetamol (from all sources including combinations containing paracetamol).
	The following information is to be documented on all paracetamol orders:
	Accurate weight (for children, frail elderly patients and adults with low body weight)
	 Orders must be expressed in milligrams (mg) or grams (g) per dose
	• For children, frail elderly patients and/or adults less than 50 kg: calculate dose in mg/kg
	 Frequency Route
	 Indication For IV orders: Maximum duration of therapy (or stop date/time)
	For PRN orders: Maximum dose in 24 hours
	 To reduce potential for dosing errors: Paracetamol (and/or paracetamol containing products) should not be prescribed both regularly and 'PRN'. Order in one section of the medication chart ONLY. Ordering in both the regular and as required 'PRN' sections of the chart may potentially lead to overdose. Paracetamol orders should specify a single route of administration (i.e., arcl OR routel OR intraveneus)
	 administration, (i.e., oral OR rectal OR intravenous) particularly relevant for paper-based charts. Do not prescribe multiple formulations concurrently. Different paracetamol-containing products should not be prescribed concurrently.
	 Orders should be written using the active ingredient drug name. Where a brand name is used on the order (e.g., combination products) the active ingredient term 'paracetamol' or 'contains paracetamol' should be documented adjacent to the brand name. Orders must be expressed in milligrams (mg) or grams (g)
	per dose

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Administration Instructions	 When being used to treat fever, patients should have their temperature recorded prior to the first administration of paracetamol to gain accurate baseline of temperature. Prior to administering paracetamol (including nurse/midwife-initiated paracetamol), clinicians are to ascertain: if paracetamol has been recently ingested (by checking with the patient and the medication chart) to ensure sufficient time has lapsed between doses check that no other formulations of paracetamol are concurrently prescribed or administered, and that the administration of the dose will not exceed the safe maximum daily dose of paracetamol (from all sources including combination paracetamol / codeine combinations). if de-escalating to an alternative route is possible (for IV paracetamol ONLY).
	 A second person check is required for administration of: intravenous paracetamol all doses administered to paediatric patients (irrespective of the route of administration) opioids in accordance with NSW health Policy Directive <i>Medication Handling</i> (PD2022_032) requirements. Paracetamol IV Infusion: Administer over 15 minutes. To hang the paracetamol 1 g in 100 mL infusion vial on IV pole, carefully peel the corner of the label on vial in the direction of the arrow (see picture below). Then lift the 2 corners of the vial to reveal perforated area to hang the infusion vial on IV pole (see picture below)
	Peel the corner of the label Lift the 2 corners to reveal perforated area to hang infusion vial on IV pole Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label Image: the corner of the label

Health South Eastern Sydney Local Health District

For patients receiving IV paracetamol, if treatment is to continue
beyond 48 hours, monitoring of liver function tests (LFTs), including
International Normalised Ratio (INR), should be carried out.
It is important for clinicians to promptly identify patients at risk of
developing acute liver injury with paracetamol poisoning. The
prognosis for recovery is good with early recognition and treatment.
Treatment of paracetamol toxicity should be guided by Prescribing
Protocol SESLHDPR/566 - Acetylcysteine IV in Acute Paracetamol
Overdose
Oral liquid Formulations – where possible/appropriate only one
strength of oral liquid paracetamol products should be kept as ward
stock.
Preparations compounded with codeine are to be handled as
accountable drugs (S4D).
Storage of IV paracetamol outside of pharmacy will be limited
and due to the critical disruption to supply.
Patients and/ or their parents or carers being discharged on
paracetamol should be provided with specific information and
education regarding paracetamol administration. They should also
be counselled that many over-the-counter products recommended
for cold, cough, headache etc. may also contain paracetamol and
should not be taken concurrently. Refer to <u>SESLHD Paracetamol</u>
Fact Sheet.
Professional Health Care Interpreters should be utilised for patient
education for patients and/ or carers who are not fluent in English or
who are Deaf.
Safety Alert - UPDATED: Critical disruption to supply – Intravenous
(IV) paracetamol solution for injection/infusion – 01 March 2024
1. NSW Health PD2020 045 - High-Risk Medicines
Management
2. Australian Medicines Handbook – Paracetamol, last
updated January 2020
3. The Children's Hospital at Westmead Pain Management
Guideline, March 2021
4. NSW TAG Inc. PARACETAMOL USE. A Position
Statement of the NSW Therapeutic Advisory Group Inc.
December 2008
5. Pain and analgesia [published 2020 December]. In:
Therapeutic Guidelines. Melbourne: Therapeutic
Guidelines Limited; accessed 25/08/2023.
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Medication Safety Pharmacists Network



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GOVERNANCE		
Enactment date	September 2021	
Reviewed (Version 2)	September 2022	
Reviewed (Version 3)	August 2023	
3.1	March 2024	
Expiry date:	31 March 2025	
Ratification date by SESLHD DTC	7 th March 2024	
Chairperson, SESLHD DTC	Dr John Shephard	
Version Number	3.1	