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# Local Anesthesia Toxicity

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# **Local Anesthesia Toxicity**

Nicole McCleery, RN, BSN, CCRN

#### Signs and symptoms Pathophysiology **Nursing Implication** • The classic description is Systemic toxicity from local progressive "biphasic" effect on the anesthetic (LA) occurs due to CNS then to the CVS accidental intravascular injection, CNS excitation progresses to absorption from the tissues or seizure or CNS depression. This is repeated doses without balanced followed by CVS excitation elimination (Neal et al., 2012). (tachycardia or ventricular The pathophysiology of LAs are 0 0 Stage 1: Get help arrhythmia) then depression thought to be an extension of their 0 Stage 2: Initial Focus (bradycardia or asystole) uses (Ciechanowicz & Patil, 2012). Blocking cardiac voltage-gated Cardiovascular system toxicity is sodium channels, prevention myocyte classically three phases depolarization, blocking Initial phase: hypertension and repolarization via potassium tachycardia channels and blocking sarcoplasmic Intermediate: myocardial reticulum voltage-dependent calcium depression and hypotension channels to limit the increase of Terminal: peripheral vasodilation, calcium available for contraction instability severe hypotension and (Ciechanowicz & Patil, 2012). arrhythmias (bradycardia, 0 Myocyte ATP is reduced which limits Management conduction blocks or asystole) energy available for connecting actin-0 (Christie, Picard & Weinberg, myosin cycle and ion channel 2015). involvement is extensive Early: metallic taste, auditory (Ciechanowicz & Patil, 2012). changes, visual disturbance (mainly Bupivacaine, Levobupivacaine and focusing), lightheadedness, Ropivacaine are long acting amideapprehension, drowsiness and based LA most commonly used in numbness of tongue and lips; clinical practice (Ciechanowicz & Restlessness, agitation, myoclonus, Patil, 2012). nystagmus and slurred speech 0 LAs rapidly cross cell membranes occur at high doses and toxicity can act in many sites The early signs are caused by including ionotropic and blocking inhibitory pathways in metabotropic. cerebral cortex- which allows for In the brain, LA affects inhibitory and disinhibition of facilitator neurons excitatory pathways resulting in excitatory cell In the heart, LAs can cause dominance causing dizziness or conduction blocks through sodium, lightheadedness (Dewaele & Santos, of Local Anesthetic Systemic Toxicity potassium and calcium channelsresulting in dysrhythmias and Tachycardia and hypertension can reduced contractility (Christie, Picard & Weinberg, 2015). Get Help LA can disrupt intracellular signal Initial Focus originating at metabotropic receptors- leading to reduced monophosphate concentrations and to asystole), chest pain, shortness decreased contractility. 0 In addition, the heart has preference for fats and ketone bodies. To be oxidized, the fats must be carried across the mitochondrial membranes by a translocase system which is inhibited by concentrations of LA

leading to enhanced potency (Christie, Picard & Weinberg, 2015). LAST can be fatal if it is not

 A checklist was developed by the American Society of Regional Anesthesia and Pain Medicine (ASRA) for the management of LAST

 Airway management: ventilate with 100% oxygen Seizure management: benzodiazepines preferred. AVOID Propofol in patients with cardiovascular

Stage 3: Cardiac Arrhythmia

- Initiate ACLS if needed Avoid vasopressin
  - calcium channel blockers, beta blockers or LA Reduce epinephrine
- does to < 1mcg/kg

2016). <u>Stage 5</u>: Stabilization

Detection

See Table 1

molecules from the plasma phase decreasing toxicity and potentially

Stage 4: Lipid Emulsion Management

 Bolus 1.5 ml/kg IV over 2 minutes Continuous infusion

LAST.

LAST.

outcomes

Pain Management (ASRA)

help influence a positive patient outcome.

0

0

 May repeat bolus The lipid emulsion creates a lipid phase that extracts the lipid soluble

reversing LAST (Nicholas & Thornton,

ASRA concludes:

0

Be prepared Risk reduction



Treatment (Neal et al., 2012).

a adequate oxygenation and ventilation throughoutprevent hypercarbia, a Assess-cardiovascular status throughout Start intralipid RX in case traditional therapy is not investigately effective.

Conclusion

• When the effectiveness of the checklist was researched by Neal et al. (2012),

Prevention measures (in a checklist form) have been put in place to help

results showed improved outcomes and excellent medical management of

reduce the risk of LAST by The American Society of Regional Anesthesia and

This is a helpful tool utilized by anesthesia staff to help focus on the immediate

needs of the patient and manage cardiac events for someone diagnosed with

By educating all staff, effective management of this emergent situation will

Comprehensive training on LAST guidelines and treatment, along with best

practice measures for patients receiving LA are essential for optimistic patient

DIAGNOSIS AND TREATMENT OF LOCAL ANESTHETIC TOXICITY



Table 2. Image retrieved from http://www.nysora.com/regionalanesthesia/foundations-of-ra/3075-toxicity-of-local-anesthetics.html

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Dest LAST events at www.lipidrescue.org and report use of lipid to www.lipidregistry.org

Table 1. Checklist created by ASRA. Image provided by Neal, Mulroy & Weinberg, 2012.

administration. It can result from the patient's risk factors, current medications or inadvertent injection directly into the vascular systemresulting in immediate absorption of the anesthetic agent into an exceptionally vascular area (Fencl. 0

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- 2015). According to Stannard (2015), 20 out of 10,000 peripheral nerve blocks (PNBs) and 4 per 10,000 epidurals result in LAST. PNBs improve short-term pain
- control, lowers pain scores from0-72 hours post procedure and reduces hospital length of stay (Joshi, Gandhi, Shah, Gadsden, & Corman, 2016). Early response at the first sign of

Introduction

Local anesthesia (LA) provides a

used for over 100 years (Fencl,

stimuli into the central nervous

system (CNS) making a surgical

According to Noble, "altering the

passage of stimuli from smaller

diameter neurons in a confined area

with lower drug dosages is called

around a surgical incision" (Noble,

administration one must be aware

of the risk of local anesthetic

systemic toxicity (LAST) (Fencl,

It is a very serious and sometimes

LA, such as the injection of LA

procedure less painful for the

patient (Noble, 2015).

2015, p.325).

2015).

When performing LA

fatal complication to

2015).

way to relieve temporary pain in a

LA prevents the passage of surgical

small part of the body and has been

toxicity is pertinent and improves chances of successful treatment. Once the reaction is noticed. immediate supportive care needs to be initiated due to the chance of severe cardiac depression. Advanced Cardiac Life Support (ACLS) should be started immediately and is considered the first-line treatment for this

complication (Noble, 2015).



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2013). 0 occur after injection without 0

epinephrine (Weinburg, 2002). Cardiovascular manifestations: Dysrhythmias and conduction delays (from prolonger PR interval

of breath, palpitations, lightheadedness, diaphoresis,







0

recognized early enough!

is Different from Other Cardiac Arrest Scenarios □ Airway management: ventilate with 100% oxygen Seizure suppression: benzodiazepines are preferred; AVOID propofel in patients having signs of cardiovascular instability

- Management of Cardiac Arrhythmia

- REDUCE individual epinephrine doses to <1 mcg/kg</p>

- Repeat bolus once or twice for persistent cardiovascular collarse
- Double the infusion rate to 0.5 mL/kg/min if blood pressure remains low
- Continue infusion for at least 10 minutes after attaining circulatory stability
- C Recommended upper limit: Approximately 10 mL/kg lipid emulsion

over the first 30 minutes

The Pharmacologic Treatment of Local Anesthetic Systemic Toxicity (LAST)

**Checklist for Treatment** 

AMERICAN SOCIETY OF

REGIONAL ANESTHESIA AND PAIN MEDICINE

- Alert the nearest facility having cardiopulmonary bypass capability
- - D Basic and Advanced Cardiac Life Support (ACLS) will require
  - justment of medications and perhaps prolonged effor
  - AVOID vasopressin, calcium channel blockers, beta blockers, or local anesthetic
- □ Lipid Emulsion (20%) Therapy (values in parenthesis are for 70kg patient)
- Bolus 1.5 mL/kg (lean body mass) intravenously over 1 minute (~100mL)
- Continuous infusion 0.25 mL/kg/min (~18 mL/min; adjust by roller clamp)