

PSRC Recommendations for Reporting Events and Interventions

The Pediatric Sedation Research Consortium (PSRC) employs and **intervention-based event reporting paradigm.** This means events are only documented and considered to be clinically significant if associated with an intervention performed in response to said event.

The severity of documented events can be designated based on the nature of the event itself, or based on the intervention associated with the event. We recommend classifying events into one of three categories:

1.	Critical event	Life-threatening, requires immediate attention and intervention.
2.	High risk event	Requires advanced level of skill and/or capacity to manage, and has potential to become a critical event if untreated.
3.	Event	Any other event that does not meet the criteria for a critical or high risk event.

A critical event includes the following:

- 1. Death
- 2. Cardiac arrest
- 3. Clinical/radiologic suspicion for pulmonary aspiration
- 4. Allergic reaction, anaphylaxis
- 5. Any event (other than the aforementioned) requiring:
 - a. ETT/NTT
 - b. Supraglottic/LMA
 - c. Chest compressions
 - d. Epinephrine, IV or IM
 - e. Muscle relaxant/paralytic agent
 - f. Vasopressor, IV (not incl. epi)
 - g. RRT, code team, or emergency anesthesia

An example of how critical events would be reported:

Critical event	n (%) N=30,000
Clinical/radiologic suspicion for pulmonary aspiration	5 (0.02)
Cardiac arrest	1 (0.003)
Death	1 (0.003)
Airway obstruction, complete ¹	50 (0.2)
Apnea ²	100 (0.3)
Hypoxia ³	200 (0.7)

¹Requiring ETT/NTT, muscle relaxant/paralytic agent, and/or chest compressions

²Requiring ETT/NTT, and/or muscle relaxant/paralytic agent

³Requiring ETT/NTT, epinephrine, IV or IM, and/or RRT, code team, or emergency anesthesia called.



A high risk event includes the following:

- 1. Apnea
- 2. Airway obstruction, complete
- 3. Laryngospasm
- 4. Seizure
- 5. Any event (other than the aforementioned) requiring:
 - a. BMV
 - b. CPAP/PEEP
 - c. Oral airway
 - d. Laryngospasm notch
 - e. Additional sedative to relieve laryngospasm
 - f. Atropine
 - g. Flumazenil
 - h. Naloxone
 - i. Unplanned hospital admission or increase in level of care

An example of how high risk events would be reported:

High risk event	n (%) N=30,000
Apnea	240 (0.8)
Airway obstruction, complete	150 (0.5)
Laryngospasm	150 (0.5)
Seizure	30 (0.1)
Hypoxia ¹	300 (1)
Bradycardia ²	30 (0.1)

¹Requiring BMV, oral airway, naloxone

 $^{2}\mbox{Requiring atropine, unplanned hospital admission or increase in level of care$

An event can be attributed to only one of the three categories; each event occurrence is exclusive to a single category. It can only be a critical event, high risk event, or an event. You cannot have an occurrence that is both a critical and high risk event, nor can you have an occurrence that is both a high risk event, nor can you have an occurrence that is both a high risk event.

Investigators may wish to study outcomes that reflect specific systems (e.g. airway/breathing, cardiovascular). Events and interventions are already categorized in the database by system. One way to report these outcomes would be to report only the events specific to the system of interest in the categories of critical events, high risk events, and events (e.g. airway/breathing critical events, airway/breathing high risk events, airway/breathing high risk events, airway/breathing high risk events. For example, airway/breathing high risk events could be reported as follows:

rway/breathing high risk event	n (%) N=30,000
onea	240 (0.8)
rway obstruction, complete	150 (0.5)
aryngospasm	150 (0.5)
ypoxia ¹	300 (1)
ypoxia ¹	3

¹Requiring BMV, oral airway, naloxone



Alternatively, investigators may report each system-based event and the frequency of interventions associated with each type of event, without using the designations of critical or high risk. For example:

Event	n (%) N=30,000	Intervention	n (%) N=30,000
Apnea	100 (0.3)	BMV	100 (0.3)
		Supplemental oxygen	100 (0.3)
Airway obstruction, complete	50 (0.2)	ETT/NTT	5 (0.02)
		Repositioning	50 (0.2)
		Supplemental oxygen	50 (0.2)
Laryngospasm	15 (0.05)	Laryngospasm notch	15 (0.05)
	. ,	Muscle relaxant/paralytic agent	5 (0.02)
Hypoxia	500 (0.7)	ETT/NTT	10 (0.03)
		Supplemental oxygen	500 (0.7)
		Repositioning	500 (0.7)
Wheezing/bronchospasm	100 (0.3)	Supplemental oxygen	10 (0.03)
		Albuterol	90 (0.3)
		Epinephrine, IV or IM	10 (0.03)

Note that the number of interventions listed for one event may not equal the number of events due to the fact that multiple interventions can be performed in response to one single event.

Here is an example of how events (that are neither critical or high risk) could be reported:

Event	n (%) N=30,000	Intervention	n (%) N=30,000
Excessive secretions	600 (2)	Suction	600 (2)
		Repositioning	50 (0.2)
Inadequate depth of sedation	300 (1)	Unanticipated pause in procedure	240 (0.8)
		Discontinued procedure	30 (0.1)
		Aborted procedure	30 (0.1)
		Child Life Specialist	240 (0.8)
		Physical restraints	120 (0.4)
Vomiting	300 (1)	Suction	200 (0.7)
-		Ondansetron	100 (0.3)
		Unanticipated pause in procedure	300 (1)

If you have any questions about this document, please contact the SPS Research Committee:

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