RAPID SEQUENCE INTUBATION (RSI)

I. Overview

Rapid Sequence Intubation (also known as Rapid Sequence Induction, or RSI) is a method of intubating patients who present with issues that make intubation difficult (e.g. gag reflex, clenched jaw, patient combativeness, etc.). Intubation is accomplished by sedating and paralyzing the patient, allowing for easier intubation. Although the physical endotracheal intubation procedure itself remains the same, the serious nature of using paralytic medications requires excellent critical thinking skills, advanced pharmacology knowledge, and continuous training.

RSI utilizes a sedative, an analgesic, a short term paralytic, and a long term paralytic when necessary. In addition, atropine may be utilized for bradycardic patients, while lidocaine may be used in cases of increased intracranial pressure (ICP) per service protocols.

Because of the potential complications involved with RSI, not all paramedics are eligible to perform the procedure. A training program approved by the ADH Section of EMS is required for every service applying to practice RSI procedures, as well as a thorough documentation process for each procedure attempted. The performance of this skill by paramedics will be limited to patients 8 years of age and older. Air medical services may use RSI for pediatric patients under 8 years of age only when performed by a flight nurse. The EMS medical director must play a key role in the selection of RSI approved paramedic providers, their training, and evaluation of the effectiveness of the program.

Training places a heavy emphasis on quality of skills, enhanced pharmacological knowledge, and critical thinking/decision making (who should and should not receive RSI). Quality Assurance/Quality Improvement (QA/QI) is critical to the success of an RSI protocol. Cases should be reviewed as soon as possible following an RSI procedure with timely feedback given to the paramedic. Paramedics making questionable decisions or having poor intubation rates should be identified and remediated immediately. If improvement criteria are not met, the paramedic must be removed from the approved practicing RSI provider list.

II. Qualifications to Participate in RSI:

The service medical director and training coordinator will select individuals to participate in program. All participants must meet training requirements as outlined in Section VI.

III. Responsibilities of Ambulance Service Medical Director:

A. Approve an RSI protocol and training program to be submitted for review/approval by the Section of EMS

B. Select/approve paramedic RSI providers
C. Monitor RSI education and skills training- all paramedics should receive RSI skills validation annually

D. Review all RSI attempts through submitted patient care reports (PCR) in an expeditious manner

E. Evaluate reports to ensure skill competency- paramedics must have at a minimum three (3) successful intubations every six (6) months

F. Initiate retraining/remediation as needed- paramedics who have not performed an RSI procedure within six (6) months must be checked off by the EMS agency training department and medical director

IV. Implementation of an RSI Protocol

A. Utilization in the pre-hospital emergency setting:

Ambulance services electing to perform RSI procedures will be required to submit the following to the Section for **review and approval prior to initiation of the protocol**:

1. The service medical director’s proposed RSI procedure protocol

2. A statement must be attached to the protocol stating that appropriate education and training will be provided to approved paramedics prior to utilization of skill/procedures

3. Paramedics who wish to be approved by the Section of EMS to perform RSI must complete the mandatory training courses (outlined in Section VI). All paramedics providing RSI must be signed off by an approved list of instructors and by their service medical director before practicing RSI in the field

B. Paramedics performing RSI procedures in a hospital setting:

Pursuant to ACT 293 of 1981, if a hospital wishes to permit an Arkansas licensed Emergency Medical Services Provider (EMSP) to perform specified procedures within the emergency department or as a member of an emergency code team functioning elsewhere in the hospital, the following action must be taken:

1. The medical staff must approve the privileges granted to the individual functioning as an EMSP with the concurrence of the hospital's governing body. Specific policies governing the supervision and the procedures to be performed by the EMSP must be developed by the medical staff and also approved by the hospital's governing body. In no event, however, may an EMSP perform a procedure on a patient in a hospital that he or she is not certified to do by the Section of EMS and Trauma Systems, Arkansas Department of Health(ADH).
2. Approved EMSPs in a hospital setting must function in accordance with physician's orders and under the direct supervision of either the physician or the registered nurse responsible for emergency services within a hospital.

3. A roster with the delineation of privileges will be maintained in the files of the supervisor for the respective department of employment and in the files of the administrator.

4. Participants must complete training program as outlined below (Section VI, Outline of Initial Training) and documentation of training will be kept on file in employees training file at the facility. A verification form regarding the monitoring of two RSI procedures must also be kept on file by the hospital. Participant must complete training and monitoring program prior to utilization of skill set.

Each paramedic participant must complete the specified training program for the ambulance service/hospital where he/she is employed. If a paramedic works at multiple ambulance services or hospitals, he/she must complete the training program for each employer and obtain approval from each service medical director or hospital governing body by which they are employed.

V. RSI Requirements

Once an approval of an RSI protocol has been obtained from the Section of EMS, the following must be conducted:

A. A copy of all RSI ePCRs will be sent to the ambulance service medical director and training coordinator immediately following the call. The ambulance service medical director will review each call for appropriate and inappropriate treatment decisions to use RSI.

B. Ambulance services will utilize the Section of EMS RSI Tracking Form (may be found on ADH website) for each patient on whom the RSI procedure is performed. The electronic form shall be submitted on the first day of the month listing all RSI procedures performed for the previous month.

C. The participating ambulance service will be required to provide a quarterly report to the Section of EMS regarding utilization of procedure and a QA/QI program update related to RSI procedures, including remediation and/or removal of RSI approved providers.

D. Reports shall consist of the following:

   1. Number of employees trained to perform RSI
   2. Number of patients receiving RSI procedures
   3. Reason patient required RSI procedure
VI. Outline of Initial Training:

Successful completion of an approved “Difficult Airway” course is required for all paramedics prior to performing RSI procedures. Additionally, Department approved instructors are to provide the following educational guidelines for RSI training:

A. Comprehensive ventilation assessment

1. Purpose
2. Procedure
3. Minute volume
4. Alveolar volume
5. Evaluating the effects of artificial ventilation
6. Pulse oximetry
   a. Purpose
   b. Indications
   c. Contraindications
   d. Complications and limitations of the technology
   e. Procedure
7. Blood gas analysis
   a. pH
   b. PaCO2
   c. PaO2
   d. Bicarbonate
   e. Base deficit
8. Capnography review
   a. Purpose
   b. Indications
   c. Contraindications
   d. Complications and limitations
   e. Procedure

B. Review of ventilation devices used by EMTs and AEMTs

1. Manual devices
   a. Purpose
   b. Indications
   c. Contraindications
   d. Complications
   e. Procedures
2. Mechanical devices
   a. Purpose
   b. Indications
   c. Contraindications
   d. Complications
   e. Procedures
C. Assisting patient ventilations

1. Review of techniques used by EMTs and AEMTs
   a. Purpose
   b. Indications
   c. Contraindications
   d. Complications
   e. Procedures
2. Review of the physiologic differences between normal and positive pressure ventilation
3. BiPAP/CPAP
   a. Purpose
   b. Indications
   c. Contraindications
   d. Complications
   e. Procedure
4. Positive End Expiratory Pressure (PEEP)
   a. Purpose
   b. Indications
   c. Contraindications
   d. Complications
   e. Procedure

D. Assessing the airway for successful RSI (look before you leap)

1. LEMON
   a. Look externally
   b. Evaluate 3-3-2
   c. Mallampati score
   d. Obstruction
   e. Neck mobility
2. HEAVEN
   a. Hypoxemia
   b. Extremes of size.
   c. Anatomic abnormalities
   d. Vomit/blood/fluid
   e. Exsanguination
   f. Neck mobility issues.

E. Medications
   Induction agents
   Barbiturates and sedatives
   Thiopental
   Methohexital
   Propofol
   Ketamine
Etomidate

Opiates
Morphine
Fentanyl
Alfentanil

Benzodiazepines
Midazolam
Diazepam
Lorazepam

Premedication agents
Atropine
Lidocaine

Neuromuscular blocking agents
Succinylcholine
Vecuronium
Mivacurium
Rocuronium
Pancuronium
Cisatracurium
Curare

Depolarizing agents
Succinylcholine

Maintenance medications and reversal agents.
Neostigmine
Pyridostigmine
Edrophonium

F. Procedure

G. Age-related variations in pediatric and geriatric patients, to include education from the following resources:

National Emergency Medical Services Education Standards, Paramedic Instructional Guidelines published by NHTSA


VII. Data Collection Requirements

These data points must be captured within the patient care report (PCR). All other required electronic PCR (ePCR) data fields must also be included. All PCR’s where RSI was performed must be reviewed by the EMS agency’s medical director to ensure all data points below are included.

- Date of run: (month/day/year)
- EMS run number
- Name of ambulance service
- Names of all EMS providers providing care - name of paramedic performing RSI procedure
- All patient demographics
- Patient weight
- Vital signs for both pre-RSI procedure and post-RSI procedure (blood pressure, pulse, heart rate/EKG interpretation, respiratory rate, Glasgow Coma Scale (GCS) score, oxygen saturation, ETCO2) **ETCO2 monitoring is required for all RSI procedures.** Vital signs should be recorded every 5-10 minutes.
- Equipment used and size – laryngoscope blade, ET tube size, etc.
- Medications – drug, dose, route, time

The PCR narrative must include the following:

- Indications for RSI procedure vs standard endotracheal intubation
- Type and size of laryngoscope blade used
- The number of attempts* for successful intubation (if not included in the main body of the PCR)
- Total time for the procedure (time from the first drug in RSI procedure until successful intubation)
- ET tube confirmation method (e.g. bilateral breath sounds, equal chest rise, ETCO2 detector, Esophageal Detector Device (EDD), etc.)
- Device used to secure ET tube
- Neck stabilization device utilized (if applicable)
- Complications of RSI procedure, if any (e.g. cervical vertebral injury, aspiration, bradycardia, vomiting, hypertension, etc.)

Additional data is required if RSI was unsuccessful. The following must be documented:

- Unsuccessful attempts* (note paramedic name)
- Rescue airway device attempted and if it was successful
- Suspected reasons for failed intubation (e.g. difficult anatomy, inability to visualize cords, orofacial trauma, inadequate patient paralysis/relaxation, etc.)
* For the purposes of documentation the Section of EMS defines an endotracheal intubation (ETI) attempt as placing a laryngoscope blade into the mouth/oropharynx with an intent to intubate. A nasotracheal intubation (NTI) attempt is defined as when the tube is placed into the nose with the intent to intubate.

Note: Using the laryngoscope blade as an adjunct to use Magill’s forceps is not an ETI. Providers should ensure documentation of the use of Magill’s is noted in the procedure section for each attempt. Providers should ensure documentation of the use of suction is noted in the procedure section for each attempt, and it should be noted if the laryngoscope blade was utilized on the first suction.

VIII. Inappropriate Treatment

In the instance that RSI is performed inappropriately or unnecessarily, the ambulance service medical director will make a written recommendation detailing a plan for the provider’s remediation. If remediation is inadequate, unsuccessful, or refused, the paramedic will be removed from the approved providers list.