SCREENING OF BIOLOGICAL SPECIMENS BY ENZYME MULTIPLIED IMMUNOASSAY TECHNIQUE (EMIT)

18.1 METHOD

This test method may be used to presumptively identify several drugs or drug metabolites and/or drug classes as being present in biological specimens.

18.2 PRINCIPLE

Immunoassays are rapid scientific tests (competitive or noncompetitive assays) that use antibodies to detect and identify chemical substances. In forensic toxicology, these are typically used to screen biological samples for the presence of a specific drug, drug metabolite or class of drugs in urine or extracted biological specimens.

The EMIT assay is based on competition between drug in the specimen and drug tagged to the enzyme glucose-6-phosphate dehydrogenase for antibody binding sites. Enzyme activity decreases upon binding of the drug to the antibody. The enzyme converts glucose-6-phosphate into glucose-6-phosphate gluconate. The reaction is coupled to the conversion of NAD to NADH. Enzyme activity is determined by spectrophotometric measurement of the absorbance of the NADH. An increase in absorbance indicates the presence of the drug in the analyzed sample.

Specificity (the degree to which the assay correctly identifies only the compound(s) of interest) is a critical component of immunoassays. Cross-reactivity to structurally similar compounds is inherent. Understanding the compounds that exhibit cross-reactivity is important to data interpretation.

18.3 SPECIMENS

Specimens include whole blood, serum, plasma, urine, vitreous humor and tissue homogenate. Standard volume for extracted specimens is 1 mL. Urine specimens do not require any sample preparation; approximately 8 drops is sufficient for un-extracted analysis. If serum/plasma specimen volume is limited, these matrices may be tested without extraction; however, caution should be used when interpreting the results.

NOTE: Analysis of tissue homogenate specimens requires matrix-matching of calibrators and controls.

18.4 REAGENTS, MATERIALS AND EQUIPMENT

18.4.1 REAGENTS

- Acetonitrile (ACN), reagent grade
- Certified blank blood and/or other biological matrices
- Certified blank urine
- Deionized water (DI H2O), laboratory general-use
- Dilute aqueous Emit® Drug Assay Buffer

*Emit® Drug Assay Buffer Concentrate (from TCA reagent kit) is diluted 1:14 with DI H2O. (For example: Add 5 mL concentrated EMIT buffer to 70 mL of DI H2O and mix, for total volume of 75 mL dilute aqueous EMIT buffer). This solution is used to prepare the working blood EMIT buffer.*
18.4.2 MATERIALS

- Disposable extraction tubes (16 x 125 mm recommended) with closures
- Disposable 12 x 75 mm tubes with closures
- Fisherbrand conical sample cups (or equivalent)
- Disposable centrifuge tubes with closures
- Laboratory glassware (graduated cylinders, flasks)

18.4.3 EQUIPMENT

- Olympus AU400e
- Calibrated, adjustable piston pipettes and verified, adjustable repeater-pipette with disposable pipette tips
- General-use equipment (centrifuge, vortex mixer, evaporator)

18.5 STANDARDS, CALIBRATORS AND CONTROLS

18.5.1 STANDARDS

- Working standard
- Working control standard
- Urine working control standard

See Appendix A for target compounds, concentrations and preparation information.

18.5.2 CALIBRATORS – All calibrations are single-point, qualitative only.

18.5.2.1 Blood, Serum, Plasma, Vitreous or Tissue Homogenate:
a. Calibrators are prepared in certified blank blood at the time of analysis using the working standard. The preparation of the calibrators is detailed in 18.6 SAMPLE PREPARATION. If necessary, calibrators may be prepared in alternate matrices (e.g., tissue homogenate).

b. Calibrations are generated from calibrator 1 for barbiturates, cannabinoids and cocaine metabolite, and from calibrator 2 for amphetamines, benzodiazepines and opiates. Single-point calibration levels represent the cutoff concentrations.

18.5.2.2 Urine calibration is performed as necessary, when testing of urine specimens is performed. The Emit® II Plus calibrators (level 1, 2, and 3) are supplied as liquids, ready to use, and kept refrigerated. The calibrators are stable until the expiration date printed on the vials.

18.5.3 CONTROLS

NOTE: Controls (positive or negative) must make up at least 10% of the extracted batch (based on number of case specimen samples), with case specimens bracketed by positive controls. Positive controls in both whole blood and/or alternate matrices may be used to bracket case specimens. If a batch contains more than 20 case specimens, an additional positive control (low or high) must be extracted and analyzed mid-run.

Additional negative or positive controls are extracted/analyzed, as needed, to bring the total percentage of controls to 10% of the batch. The additional controls are spaced as evenly as possible throughout the test batch. When analyzing compounds in multiple matrices, both whole blood and alternate matrix controls apply towards 10% of the batch.

18.5.3.1 Blood, Serum, Plasma, Vitreous, or Tissue Homogenate

18.5.3.1.1 Negative Controls

At least two negative whole blood controls are tested in each batch. One is to function as the matrix blank for the spectrophotometer, and the remaining functions as a negative control. All controls are prepared using certified blank blood. When analyzing tissue homogenate, two negative controls must be prepared in that matrix. If analyzing only tissue homogenate specimens, blood negative controls are not required.

18.5.3.1.2 Positive Controls

At least two whole blood positive controls are tested with every batch. The preparation of the positive whole blood controls is detailed in 18.6 SAMPLE PREPARATION. When analyzing tissue homogenate, at least two positive controls must be prepared in that matrix. If analyzing only tissue homogenate specimens, blood positive controls are not required.

18.5.3.2 Urine

18.5.3.2.1 Quality Control

The Green QC rack (see 18.7.3.1) is designated for running urine controls, on days urine testing is performed, and consists of:

a. The negative control, made up of certified blank urine.
b. The urine positive control; prepare by adding 30 µL of the urine EMIT working control standard (using a calibrated pipette) to 1.0 mL blank urine in a 12 x 75 mm tube. Cap the tube and vortex-mix. Transfer contents to a sample cup.

NOTE: Multiple positive control sample cups may be filled from this control. Where the green QC rack is run concurrently with the test batch, a sample of the spiked control serves as the positive control at the start of the batch, to bracket case specimens.

18.5.3.2.2 Within-batch Quality Controls

At least one negative control (certified negative urine) and one positive control are tested at the start of every batch. Note that if the green QC rack is run concurrently with the test batch, this fulfills the requirement.

18.6 SAMPLE PREPARATION (Blood, Vitreous, Tissue Homogenate, Urine)

18.6.1 Blood, Vitreous, or Tissue Homogenate

NOTE: Serum or plasma will be run with contemporary blood calibrators and controls. Serum or plasma specimens may be extracted or un-extracted if volume is limited. Skip to step 18.6.1.11 for un-extracted serum or plasma samples (see also 18.3).

18.6.1.1 Label a clean extraction tube for each member of the test batch. (i.e., calibrators, controls, or case samples).

18.6.1.2 Using a calibrated pipette, add 1 mL of certified blank blood into each of the two calibrator tubes (low and high), the matrix blank, and the positive and negative control(s).

18.6.1.3 Using a calibrated pipette and the working standard, spike the calibrators as follows: Use a calibrated pipette to add 20 µL of working standard to the calibrator 1 tube and 40 µL to the calibrator 2 tube.

18.6.1.4 Using a calibrated pipette and the working control standard, spike the low and high positive controls as follows: Use a calibrated pipette to add 10 µL of working control standard to the low positive control tube and 20 µL to the high positive control tube.

18.6.1.5 Using a calibrated pipette, sample 1 mL of each case specimen into its respective tube.

18.6.1.6 To each tube, add 1 mL of methanol, followed immediately by 3 mL of acetonitrile. Vortex mix approximately 30 seconds.

18.6.1.7 Centrifuge the tubes for 5 minutes at 2000rpm (recommended for 16 x 125 mm tubes) to achieve precipitation.

18.6.1.8 Decant the supernatant into a conical centrifuge tube, and evaporate under air at 50°C to approximately 100 µL.

18.6.1.9 Remove the tubes from the evaporator and add 350 µL of working blood EMIT buffer and vortex mix.

18.6.1.10 Centrifuge the tubes for 5 minutes at 2000 rpm.
18.6.1.11 Transfer the clear supernatant to labeled conical sample cups for analysis on the Olympus AU400e. See section 18.7 INSTRUMENTAL PARAMETERS for guidance.

18.6.2 Urine

Urine samples require no sample preparation. A few drops (8-10) of sample are transferred into labeled conical sample cups for analysis on the Olympus AU400e (see 18.7 INSTRUMENTAL PARAMETERS).

18.7 INSTRUMENTAL PARAMETERS

18.7.1 INSTRUMENT MAINTENANCE

Daily, weekly, monthly, 3-month, and 6-month maintenance is required to keep the AU400e in working order. See the Procedure for Instrument Maintenance and Performance Verification (PTmp12504) and refer to the Olympus AU400e Maintenance Manual for instructions on routine maintenance as well as troubleshooting any issues that may arise. Maintenance records are filed in the instrument maintenance/QC binder.

18.7.2 INSTRUMENT CALIBRATION

18.7.2.1 From the main computer screen, under USER, select “Start Condition/New Data Index”. Change the data index to reflect the current time and set the operator name.

18.7.2.2 Under USER, select “Order Calibration from Racks”, select which sample type (serum or urine- select serum if using blood, serum, plasma, vitreous or tissue homogenate). Click on “Start Entry” and enter “1” or “3” under “Profile” if using a serum panel, “2” for the urine panel. Click “Entry” to make the selection, and then click on “Exit.”

NOTE: Serum panel “1” includes all compound classes listed in 18.5.2.1.b. Serum panel “3” includes all compound classes listed, with the exception of cannabinoids; panel “3” is used for specimens that will be screened with a separate method for the presence of cannabinoids.

18.7.2.3 For blood (and other associated matrices), load the extracted matrix blank in position 1 of the blue sample rack, designated for the reagent blank. Load the supernatant from the 20 µL calibrator 1 in position 1, and the supernatant from the 40 µL calibrator 2 in position 2, of the red-striped yellow sample rack, designated for calibration.

18.7.2.4 For the urine reagent blank, use a blue sample rack with DI H₂O in position 10. For urine calibration, fill positions 1, 2 and 3 in a yellow sample rack with calibrators 1, 2 and 3, respectively.

NOTE: The urine calibration is performed weekly, or as needed, with calibration data filed in the instrument maintenance/QC binder.

18.7.3 ORDERING CONTROL SAMPLES

NOTE: Ensure that step 18.7.2.1 (setting the operator name and data index) is performed before ordering any quality controls or sample tests.
18.7.3.1 The negative, low, and high controls for blood and other associated matrices are ordered as samples. See section 18.7.4 ORDERING SAMPLE TESTS.

18.7.3.2 To order urine controls, select “Order QC from Green Rack” under USER. Select urine for the sample type. Fill the labeled sections of a green sample rack with the negative and positive controls.

NOTE: The spiked urine control (as described in 18.5.3.2.1.b), ordered as a “QC” in the green rack and as a “sample” when run at the end of the batch, serve to bracket case specimens.

18.7.4 ORDERING SAMPLE TESTS

18.7.4.1 Under USER, choose “Select Report Format”. Select either the urine report or blood report appropriately (choose the blood report format that corresponds to panel “1” or panel “3”).

18.7.4.2 Under USER, select “Order Sample Tests”. Set the sample type appropriately.

18.7.4.3 Click on “Start Entry” at the bottom of the screen, and enter the sample ID (for example “Negative Control” or an ST# associated with a case).

18.7.4.4 Click on “Profile” and select “1” or “3” for serum and “2” for urine.

18.7.4.5 Click on “Entry” at the bottom of the screen to accept the selections. Continue as above until all sample tests have been ordered.

18.7.4.6 Samples are placed in labeled conical sample cups in gray sample racks for urine, and in red-striped gray racks for blood and other associated matrices. Ensure that the position of each sample cup matches the position ordered as above in 18.7.4.3.

18.7.4.7 Once all calibration, quality control samples, and case samples have been ordered, and the racks have been placed in the rack loading area of the AU400e, the green “PLAY” arrow is pressed at the top of the screen to start the analysis.

18.8 CRITERIA FOR BATCH ACCEPTANCE

If the analysis of the batch meets the criteria listed below, the results for the specimens are accepted.

18.8.1 Controls

18.8.1.1 Negative control – blood and associated matrices

The negative control must read negative for all target compounds/classes. If the negative control is positive for any target compound/class (response of >100) all unknowns must be reanalyzed for that target compound/class.

18.8.1.2 Negative control – urine

The negative control must read negative (<25) for all target compounds/classes. If any target compound/class reads 25 or
higher, all unknowns must be reanalyzed for that target compound/class.

18.8.1.3 Positive controls – blood and associated matrices

a. The low control is prepared to illicit responses slightly above the cutoff concentration (i.e., response just over 100). In the event that one or more target compound/class does not read >100, that target compound/class response is multiplied by 0.75, and truncated to a whole number, to determine the new cutoff response. Any unknown samples reading greater than or equal to the new cutoff response shall be confirmed for that target compound/class. For example, if the opiate control reads 94, the resulting cutoff is 70. All cases with an opiate response ≥70 will be confirmed for opiates.

b. The high control must read positive for all target compounds/classes (>100). In addition, the response for the high control should be greater than the response for each analyte in the low control. If a target compound/class response is negative (or less than the low control response for that target compound/class), all unknowns must be reanalyzed for that target compound/class.

18.8.1.4 Positive controls – urine

a. Positive quality controls (spiked using working control standard - see 18.5.3.2.1.b) must read >80 for target compounds/classes.

b. If any of the above criteria are not met for a target compound/class, all unknowns must be reanalyzed for that target compound/class.

18.8.2 Case Specimens

If the target compound/class response printed on the report is >100, (or greater than or equal to the new cutoff as described in 18.8.1.3.a), it is reported as presumptive positive.

18.9 REPORTS AND DOCUMENTATION

18.9.1 Batch documentation includes instrument report printouts for all members of the test batch, including case samples, quality control, and calibration data (for blood and other associated matrices only) and the Excel worklist.

18.9.2 The analyst performing the test reviews the instrument reports, adds initials to each quality control and calibration printout, adds initials to each specimen report, and signs and dates the worklist prior to submission for peer review.

18.9.3 The peer review process includes verification that the calibration and all quality controls are acceptable (or that positive low blood control responses <100 are appropriately documented and/or new cutoff values are determined), and a report is included for all samples listed on the worklist. Batch review is documented by the reviewer adding initials/date to the quality control and calibration reports and signing/dating the worklist.
18.9.4 For blood and other associated matrices, the original sample report and copies of the calibration, negative controls, positive controls, and worklist are included in the respective case file. The original calibration data, control results, and worklist are filed in the batch file.

For urine, the original sample report and copies of the control results and the worklist are included in each respective case file. The original control results and worklist are filed in the batch file.

18.10 REPORTING

18.10.1 Any positive results reported from this assay are indicated as “presumptive positive” in the LIMS panel.

18.10.2 All presumptive positive results that are chosen to be confirmed must be confirmed by a separate method, in the same matrix or from a different matrix from the same case; if the confirmation method indicates results are not reportable, the presumptive positive EMIT result(s) is removed from the LIMS panel.

18.11 REFERENCES

- Behring Diagnostics. EMIT II Package Inserts (for reagent kits).
APPENDIX A

NOTE: Refer to the *Standard Solution Preparation procedure (PQ12702)* for additional information.

Stock standards (1.0 mg/mL) are purchased from an approved reference material supplier and include the following:

- Benzoylecgonine
- Morphine
- Oxazepam
- Secobarbital
- 11-nor-9-COOH-Δ9-THC
- d-Methamphetamine

**Preparation**

Adjustments to the final volume of prepared standards are permitted, provided that the proportions and final concentration is maintained.

**Working standard (used for all matrices except urine)**

Using a calibrated pipette, add the following volumes of each compound to a 50 mL Class A volumetric flask and bring to nominal volume with methanol.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Volume (µL)</th>
<th>Final Standard Concentration (mg/L)</th>
<th>Cal Level 1 Concentration (mg/L)</th>
<th>Cal Level 2 Concentration (mg/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>benzoylecgonine</td>
<td>250</td>
<td>5</td>
<td>0.10 mg/L</td>
<td>-</td>
</tr>
<tr>
<td>morphine</td>
<td>62.5</td>
<td>1.25</td>
<td>-</td>
<td>0.05 mg/L</td>
</tr>
<tr>
<td>oxazepam</td>
<td>125</td>
<td>2.5</td>
<td>-</td>
<td>0.10 mg/L</td>
</tr>
<tr>
<td>secobarbital</td>
<td>250</td>
<td>5</td>
<td>0.10 mg/L</td>
<td>-</td>
</tr>
<tr>
<td>(-)-11-nor-9-COOH-Δ9-THC</td>
<td>25</td>
<td>0.5</td>
<td>0.01 mg/L</td>
<td>-</td>
</tr>
<tr>
<td>d-methamphetamine</td>
<td>250</td>
<td>5</td>
<td>-</td>
<td>0.20 mg/L</td>
</tr>
</tbody>
</table>

**Working control standard (used for all matrices except urine)**

Using a calibrated pipette, add the following volumes of each compound to a 10 mL Class A volumetric flask and bring to nominal volume with methanol.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Volume (µL)</th>
<th>Final Standard Concentration (mg/L)</th>
<th>Low Pos Ctl Concentration (mg/L)</th>
<th>High Pos Ctl Concentration (mg/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>benzoylecgonine</td>
<td>125</td>
<td>12.5</td>
<td>0.125 mg/L</td>
<td>0.25 mg/L</td>
</tr>
<tr>
<td>morphine</td>
<td>60</td>
<td>6</td>
<td>0.06 mg/L</td>
<td>0.12 mg/L</td>
</tr>
<tr>
<td>oxazepam</td>
<td>200</td>
<td>20</td>
<td>0.20 mg/L</td>
<td>0.40 mg/L</td>
</tr>
<tr>
<td>secobarbital</td>
<td>150</td>
<td>15</td>
<td>0.15 mg/L</td>
<td>0.30 mg/L</td>
</tr>
<tr>
<td>(-)-11-nor-9-COOH-Δ9-THC</td>
<td>12.5</td>
<td>1.25</td>
<td>0.0125 mg/L</td>
<td>0.025 mg/L</td>
</tr>
<tr>
<td>d-methamphetamine</td>
<td>200</td>
<td>20</td>
<td>0.20 mg/L</td>
<td>0.40 mg/L</td>
</tr>
</tbody>
</table>
Urine working control standard

Using a calibrated pipette, add the following volumes of each compound to a 10 mL Class A volumetric flask and bring to nominal volume with methanol.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Volume (µL)</th>
<th>Final Standard Concentration (mg/L)</th>
<th>Pos Ctl Concentration (mg/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>benzoylecgonine</td>
<td>125</td>
<td>12.5</td>
<td>0.375 mg/L</td>
</tr>
<tr>
<td>morphine</td>
<td>125</td>
<td>12.5</td>
<td>0.375 mg/L</td>
</tr>
<tr>
<td>oxazepam</td>
<td>80</td>
<td>8.0</td>
<td>0.24 mg/L</td>
</tr>
<tr>
<td>secobarbital</td>
<td>80</td>
<td>8.0</td>
<td>0.24 mg/L</td>
</tr>
</tbody>
</table>
### LIST OF CHANGES

<table>
<thead>
<tr>
<th>Revision Date</th>
<th>Description</th>
<th>Page Number</th>
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<tr>
<td>5/30/13</td>
<td>Method approved by the State Toxicologist. See DRA dated 5/28/13. Method released for evidentiary use as of 5/30/13.</td>
<td>All</td>
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<tr>
<td>6/13/14</td>
<td>Added target concentrations to calibrator description table and added table for preparation of working control standard to spike positive controls, in lieu of prepared whole blood controls, to section 18.6.1. Changed opiates cutoff concentration from 20 ng/mL to 50 ng/mL in 18.6.1. Noted calibrator levels (cutoff concentrations) for compound classes in 18.6.2. Changed control description to reflect spiked positive controls in 18.6.3 and 18.7.1.4. Corrected reconstitution expiration for UTAK 0 to 30 days in 18.6.3. Added Keto-Diastix® to materials list in 18.5.2 and amended wording in 18.7.2. Removed 18.11.3, which described presumptive reporting.</td>
<td>3-8</td>
</tr>
<tr>
<td>6/1/15</td>
<td>Added “Technique” to title. Minor changes throughout document needed for clarification or grammatical reasons. Changes made to ‘18.6.3 Controls’ to reflect the need for 10% controls with every batch, that controls must bracket casework, and all batches must end with a positive control.</td>
<td>All</td>
</tr>
<tr>
<td>6/30/15</td>
<td>Preparation instructions for the within-run spiked urine positive control in 18.6.3.2.2 (c) were modified for use of 0.25 mL blank urine (previously 0.5 mL), to provide an opiates concentration above the target cutoff (calibrator level).</td>
<td>7</td>
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<tr>
<td>10/7/15</td>
<td>Replaced “UTAK 0” in 18.9.1.2 with “negative” to reflect use of either the UTAK 0 negative serum control or blank urine.</td>
<td>10</td>
</tr>
<tr>
<td>3/16/16</td>
<td>Edited 18.6.1.4 to indicate use of the working controls standard to prepare positive controls for all matrices. Added wording to 18.9.1.4 to describe acceptance criteria for the within-batch (spiked) positive controls and a note to clarify that the UTAK 5, TCA and acetaminophen/salicylate positive controls (not spiked with working control standard) are positioned in the green quality control sample rack on the instrument. Other minor edits throughout. See DRA dated 1/28/16.</td>
<td>4, 7, 9, 10-11</td>
</tr>
<tr>
<td>11/7/16</td>
<td>Removed “...and Enzymatic Assay” from title, as acetaminophen/salicylates testing is no longer performed. Edited 18.2 to specify compounds/classes for urine versus other specimen types. Removed listed reagents in 18.5.1 for acetaminophen/salicylates, TCA, methadone and PCP and preparation instructions for acetaminophen/salicylates and TCA reagent kits. Removed PCP, methadone and nortriptyline from 18.6.1.2 list of stock standards and from 18.6.1.3 and 18.6.1.4 for preparation of working standard and working control standard. Updated 18.6.2.2 and 18.6.3.2 to reflect that urine calibration and QC check are only performed as needed and to remove acetaminophen/salicylates and TCA references. Removed description of ketones/glucose testing from 18.7.2 and updated 18.9.1.4 to remove acetaminophen/salicylates, TCA, methadone and PCP references and removed NOTE in 18.10.1 for acetaminophen/salicylates criteria. Removed reporting description for ketones/glucose testing in 18.11. Other minor edits throughout.</td>
<td>All</td>
</tr>
<tr>
<td>7/9/19</td>
<td>Removed policy and purpose sections and replaced with 18.1 METHOD. Added specific wording regarding matrix-matching for tissue homogenate in 18.3 SPECIMENS. Edited STANDARDS section – moved list of compounds and preparation to APPENDIX A. Updated 18.6 SAMPLE PREPARATION for use of spiked urine control. Other minor edits throughout. See DRA dated 7/2/19.</td>
<td>All</td>
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<tr>
<td>Date</td>
<td>Changes</td>
<td>References</td>
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<tr>
<td>------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------------</td>
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<tr>
<td>6/15/20</td>
<td>Edited wording in 18.2 and 18.3. In 18.5.3 CONTROLS, edited wording to specify that a positive control is required mid-run if there are more than 20 case specimens. In 18.7.2.3 and 18.7.2.4, clarified loading of blanks and calibrators in analysis racks. In section 18.9.2 REPORTS AND DOCUMENTATION, removed requirement for testing analyst to add the date to each quality control and calibration printout, to align with current practices. Added section 18.11 REFERENCES.</td>
<td>1, 3-6, 8</td>
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<tr>
<td>10/23/20</td>
<td>Corrected typo in title of procedure. Administrative change only. Documented in DRA record for 6/15/20 revision 8.</td>
<td>1</td>
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