Policy Manual



Adverse Event Reporting Procedure

Approval Date: 11Nov2013 Revision Date: 28May2015

Revision Date: 9OCT2017 To accommodate eSOURCE implementation.

Revision Date: 310CT2017 Cosmetic Only

Revision Date: 11DEC2021 Revision Date: 23JAN2024

1. Enter or update AE during visit while in room with subject.

- 2. Progress Note at procedure level or in main progress note any special circumstances, discussion regarding AE, meds taken, etc.
- 3. If AE is serious or unexpected, the sponsor will be notified within 24 hours.
- 4. Verbally discuss details of AE with Investigator for triage. Investigator determines severity, causality, and any treatment (if deemed necessary by Investigator).
- 5. Progress Note discussion with Investigator as he/she determined severity, causality, and any treatment given. This is documented in source by study staff for the Investigator.
- 6. Enter AE into CRF per sponsor's requirements.
- 7. Update AE as necessary. Follow up with subject until resolved or end of study as determined by sponsor.