

5	SOP: Emergency Use Post-Review				
Ā	NUMBER	DATE	AUTHOR	APPROVED BY	PAGE
	HRP-027	08/15/2014	L. Smith	C. Kiel	1 of 1

# 1 PURPOSE

- 1.1 This procedure establishes the process to communicate the review of:
  - 1.1.1 Emergency use of a drug, biologic, or device in a life-threatening situation.
  - 1.1.2 Compassionate use of an unapproved device without an IDE for a serious condition.
- 1.2 The process begins when the <u>Designated Reviewer</u> has notified IRB staff of whether an actual or proposed use has followed or will follow FDA regulations and guidance.
- 1.3 The process ends when the IRB staff has communicated the results to the physician and if necessary initiated the non-compliance process.

## 2 REVISIONS FROM PREVIOUS VERSION

2.1 None

## 3 POLICY

- 3.1 Whenever possible physicians are to notify the IRB of a proposed emergency use of a drug, biologic, or device in a life-threatening situation in advance of the use.
- 3.2 Physicians are to notify the IRB of a proposed compassionate use of an unapproved device without an IDE for a serious condition.
- 3.3 Data obtained from uses covered by this procedure cannot be used in a non-exempt systematic investigation designed to develop or contribute to generalizable knowledge.

## 4 **RESPONSIBILITIES**

4.1 IRB staff carry out these procedures.

## 5 PROCEDURE

- 5.1 If the <u>Designated Reviewer</u> has indicated that the proposed use will follow FDA regulations:
  - 5.1.1 Complete a "TEMPLATE LETTER: Pre-Review of Emergency Use Criteria Met (HRP-570)" or equivalent, and send to the physician.
  - 5.1.2 Set a 5 day deadline for receipt of the 5 day report.
- 5.2 If the <u>Designated Reviewer</u> has indicated that the proposed use will NOT follow FDA regulations, complete a "TEMPLATE LETTER: Pre-Review of Emergency Use Criteria Not Met (HRP-571)" or equivalent, and send to the physician.
- 5.3 If the <u>Designated Reviewer</u> has indicated that the actual use followed FDA regulations
  - 5.3.1 Complete a "TEMPLATE LETTER: Review of Emergency Use Criteria Met (HRP-572)" or equivalent, and send to the physician.
  - 5.3.2 For uses of drugs and biologics, set a 30 day deadline for receipt of a protocol.
- 5.4 If the <u>Designated Reviewer</u> has indicated that the proposed use did NOT follow FDA regulations:
  - 5.4.1 Complete a "TEMPLATE LETTER: Review of Emergency Use Criteria Not Met (HRP-573)" or equivalent and send to the physician.
  - 5.4.2 Manage under "SOP: New Information (HRP-024)" as Non-Compliance.

### 6 MATERIALS

- 6.1 SOP: New Information (HRP-024)
- 6.2 TEMPLATE LETTER: Pre-Review of Emergency Use Criteria Not Met (HRP-571)
- 6.3 TEMPLATE LETTER: Review of Emergency Use Criteria Met (HRP-572)
- 6.4 TEMPLATE LETTER: Review of Emergency Use Criteria Not Met (HRP-573)
- 6.5 TEMPLATE LETTER: Pre-Review of Emergency Use Criteria Met (HRP-570)
- 6.6 WORKSHEET: Emergency Use (HRP-322)

### 7 REFERENCES

7.1 21 CFR §50.23; 21 CFR §56.104(c).