### 1. PURPOSE

The purpose of this standard operating procedure (SOP) is to specify who has the authority to sign documents on behalf of the Health Sciences Research Ethics Board (HSREB), and to describe the responsibilities of such individuals, and the circumstances under which signing authority may be delegated.

### 2. GENERAL POLICY STATEMENT

Research Ethics Boards (REBs) are accountable for their activities and decisions, and appropriate controls must be applied to ensure that documents related to REB review and approvals of research are signed by a person or persons having the appropriate authority to do so.

### 3. RESPONSIBILITY

This SOP applies to the HSREB Chair, Vice-Chair(s), REB members and OHRE staff.

The HSREB Chair is responsible for signing documents related to the HSREB review and approval of research. If the task of signing is delegated, the responsibility for oversight remains with the HSREB Chair.

### 4. DEFINITIONS

See Glossary of Terms

### 5. SPECIFIC POLICIES AND PROCEDURES.

#### 5.1. Delegation of Signing Authority

**5.1.1.** The HSREB Chair may delegate signing authority for documents related to HSREB review and approval;

**5.1.2.** The HSREB Chair may only delegate signing authority to Vice-Chair(s) and OHRE staff with the skill and knowledge necessary to effectively exercise the authority;

**5.1.3.** The HSREB Chair may not delegate his/her signing authority to consultants;

**5.1.4.** The HSREB Chair may delegate signing authority indefinitely, or for defined periods of time (e.g., for absences);

**5.1.5.** Delegation of signing authority must be made in writing and kept on file.
5.2. **REB Reviews, Approvals and Other Correspondence with the Investigator**

5.2.1. The results of reviews and decisions made by the HSREB, either the Full Board or delegated review, including investigation of ongoing research, that grant or may appear to grant investigators with initial or continuing approval of research involving human participants, or suspends or terminates such research, must be signed by the HSREB Chair, Vice-Chair(s), or as otherwise designated in writing by the HSREB Chair;

5.2.2. Any letters, memos, or e-mails between the HSREB and or OHRE and investigators that provide written information concerning the review of research (e.g. requests for consent form changes, requests for additional information, renewal reminder notices) and that do not grant or appear to grant approval of the research, may be signed and sent by the appropriate OHRE staff as delegated in writing by the HSREB chair;

5.2.3. Individuals must sign their own name. The individual’s title must be documented.

5.3. **Correspondence with External Agencies**

5.3.1. Unless authorizing signature by the VP Research is required, the HSREB Chair, Vice-Chair(s) or designee signs all correspondence to federal government agencies (Health Canada, OHRP, FDA) and funding agencies or sponsors as applicable.

6. **REFERENCES**


6.2. The International Conference on Harmonization Guidelines for Good Clinical Practice, Section 3;


7. **SOP HISTORY**

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