



University
of Manitoba

Research Ethics and Compliance

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HEALTH RESEARCH ETHICS BOARD (HREB)
CERTIFICATE OF FINAL APPROVAL FOR NEW STUDIES
Delegated Review

PRINCIPAL INVESTIGATOR: Dr. Dieter Schonwetter	INSTITUTION/DEPARTMENT: University of Manitoba/College of Dentistry	ETHICS #: HS26761 (H2025:002)
APPROVAL DATE: March 11, 2025	EXPIRY DATE: March 11, 2026	
STUDENT PRINCIPAL INVESTIGATOR SUPERVISOR (If applicable):		

PROTOCOL NUMBER:	PROJECT OR PROTOCOL TITLE: Navigating the AI Landscape: Insights into University Students' and Academics' Use of Artificial Intelligence for Learning, Teaching, and Research
SPONSORING AGENCIES, FUNDING AGENCIES AND/OR COORDINATING GROUPS: No funding	

Submission Date of Investigator Documents: January 28, 2025, March 3, 2025 and March 11, 2025	HREB Receipt Date of Documents: January 28, 2025, March 3, 2025 and March 11, 2025
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THE FOLLOWING ARE APPROVED FOR USE:

Document Name	Version (if applicable)	Date
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Protocol:

Protocol along with proposal as outlined in the revised University of Manitoba Barnabyne Campus Research Ethics Board Submission Form and Letter of Response dated March 3, 2025

January 27, 2025

Consent and Assent Form(s):

Consent Disclosure (as part of survey introduction) for Dental Hygiene and Dentistry Students
Consent Disclosure (as part of survey introduction) for Dental Hygiene and Dental Academics

March 11, 2025
March 11, 2025

Other:

Survey for Dental Hygiene and Dentistry Students
Survey for Dental Hygiene and Dental Academics
Generic Email Recruitment Script (undated)

March 11, 2025
March 11, 2025
Submitted January 28, 2025

CERTIFICATION

The above-named research study/project has been reviewed in a **delegated manner** by the University of Manitoba (UM) Health Research Board (HREB) and was found to be acceptable on ethical grounds for research involving human participants. The study/project and documents listed above was granted final approval by the Chair or Acting Chair, UM HREB.

HREB ATTESTATION

The University of Manitoba (UM) Research Board (HREB) is organized and operates according to Health Canada/ICH Good Clinical Practices, Tri-Council Policy Statement 2, and the applicable laws and regulations of Manitoba. In respect to clinical trials, the HREB complies with the membership requirements for Research Ethics Boards defined in Division 5 of the Food and Drug Regulations of Canada and carries out its functions in a manner consistent with Good Clinical Practices.

QUALITY ASSURANCE

The University of Manitoba Research Quality Management Office may request to review research documentation from this research study/project to demonstrate compliance with this approved protocol and the University of Manitoba Policy on the Ethics of Research Involving Humans.

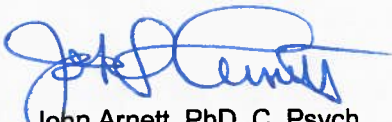
CONFLICT OF INTEREST

Any Principal or Co-Investigators of this study who are members of the UMHREB did not participate in the review or voting of this study.

CONDITIONS OF APPROVAL:

1. The study is acceptable on scientific and ethical grounds for the ethics of human use only. ***For logistics of performing the study, approval must be sought from the relevant institution(s).***
2. This research study/project is to be conducted by the local principal investigator listed on this certificate of approval.
3. The principal investigator has the responsibility for any other administrative or regulatory approvals that may pertain to the research study/project, and for ensuring that the authorized research is carried out according to governing law.
4. **This approval is valid until the expiry date noted on this certificate of approval. A Bannatyne Campus Annual Study Status Report must be submitted to the HREB within 15-30 days of this expiry date.**
5. Any changes of the protocol (including recruitment procedures, etc.), informed consent form(s) or documents must be reported to the HREB for consideration in advance of implementation of such changes on the **Bannatyne Campus Research Amendment Form**.
6. Adverse events and unanticipated problems must be reported to the HREB as per Bannatyne Campus Research Boards Standard Operating procedures.
7. The UM HREB must be notified regarding discontinuation or study/project closure on the **Bannatyne Campus Final Study Status Report**.

Sincerely,



John Arnett, PhD. C. Psych.
Chair, Health Research Ethics Board
Bannatyne Campus