- 5) Identification of any procedures which are experimental
- 6) A description of any reasonably foreseeable risks or discomforts to the subject
- 7) A description of any benefits to the subject or to others which may reasonably be expected from the research
- 8) A discloaupl an explanation as to whether any medical treatments are available, if injury occurs and, if

so, what they consist of, or where further information may be obtained

- 11) **Research, Rights or Injury:** An explanation of whom to contact for answers to pertinent questions about the research and research subjects' rights, and whom to contact in the event of a research-related injury to the subject
- 12) A statement that participation is voluntary, refusal to participate will involve no penalty or loss of benefits to which the subject may prevaise ensittled, and the subject may prevaise information

a) A statement that identifiers might be removed from the identifiable private information or identifiable biospecimens and that, after such removal, the information or biospecimens could be used for future research studies or distributed to another

14) Add this statement to the bottom of the informed consent: For questions regarding your rights as a participant in this research or IRB approval, contact Dr. Michelle McWhorter, Associate Professor of Biology, IRB Chair, at 937-327-6483, or by email at mmcwhorter@wittenberg.edu.

Drafted 03/14/2016, Approved IRB