

**Institutional Review Board
Informed Consent Document for Research**

Principal Investigator: Sandra F. Simmons, PhD and Eduard E. Vasilevskis, MD, MPH **Revision Date:** August 20, 2018
Study Title: A Randomized Controlled Trial to Deprescribe for Older Patients with Polypharmacy Transferred from the Hospital to Skilled Nursing Facilities
Institution/Hospital: Vanderbilt University Medical Center

Statement by Surrogate Agreeing to Patient's Participation

I, _____ [name of decision-maker/surrogate],
am the _____ [state relationship to participant]
of _____ [state participant's name]. I have read the informed
consent document or it has been explained to me. I have had the opportunity to ask any questions and all of my
questions have been answered. I have been informed that an investigational treatment may be administered to
_____ [participant's name]. I believe receiving such treatment
would be in the interests of _____ [participant's name] and is consistent with
what he/she would have decided had he/she been able to do so.

Your decision to allow your family member/friend to participate in this research study is voluntary. You may choose not to allow his/her participation and he/she will receive alternative treatments without affecting his/her healthcare/services or other rights. You are also free to withdraw him/her from this study at any time. In the event new information becomes available that may affect the risks or benefits associated with this research study or your willingness to allow continued participation in this research study, you will be notified so that you can make an informed decision whether or not to continue your family member/friend's participation in this study.

Your family member/friend will periodically be re-evaluated for the capacity to give consent. If he/she is found to be capable, continued participation in this study would only occur with his/her consent.

_____/_____/_____
Signature of Health Care Decision-Maker/Surrogate Date

_____/_____/_____
Signature of Witness Date

_____/_____/_____
Name and Signature of person obtaining consent Date

Date of IRB Approval: 09/10/2019
Date of Expiration: 09/09/2020

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