Study Information Sheet for ED Clinicians

Study Title: A pilot study to test the acceptability and feasibility of brief motivational interview intervention to help patients formulate their goals for medical care in the emergency department

Version: 5/24/2017

Principal Investigator: Kei Ouchi, MD, MPH
Purpose of the Research: This is a research study to test the acceptability and feasibility of our brief motivational interview intervention to facilitate advance care planning (ACP) conversation on older adults with serious co-morbid illness being discharged from the emergency department (ED).

Sponsor of the Research: Emergency Medicine Foundation.

How we obtained your name and contact information: The participants are identified based on their clinical roles as attending physicians and physician assistants belonging to the Department of Emergency Medicine at Brigham and Women's Hospital. We asked your directors of service to identify you.

Why are we asking you to participate?
We are asking you to participate because we want to test the acceptability of our intervention when performed in the ED.

How many people are anticipated to participate?
We hope to recruit 5 to 10 attending physicians and 5 to 10 physician assistants to participate in this study. We hope to recruit 125 patients to participate in this study with you.

How long it will take to complete the study?
The study will take place in the ED observation unit. We anticipate that each patient encounter/intervention administration and answering of acceptability survey will add 8 minutes total to your routine clinical practice. We will plan to continue enrollment until we complete 125 patient encounters total. With your permission, we will video-record your patient encounter to ensure the accuracy of our intervention administration. Deciding not to participate will have no effect on your employment or professional standing in the hospital.

Remuneration: There is no remuneration for this study.

Confidentiality and Data Security
The data will be collected in the form of qualitative (e.g. your comments) and quantitative (e.g. how many minutes does it take to read off the scripted text in a patient encounter) formats. All data will be de-identified (we will not be able to match specific data to specific participant) and stored in a secure location in PI's office, which will be destroyed upon completion of this study.

What are the risks associated with participation?
There is no clear foreseeable risk associated with participation in this study other than 8 extra minutes of your time during your routine clinical practice in the ED observation unit. Your participation is voluntary and you may stop at any time.

**Questions:**

The PI can be contacted at the following 24 hours / day and 7 days per week:

Cell – 857-205-4947  
Email – kouchi@partners.org

If you’d like to speak to someone not involved in this research about your rights as a research subject, or any concerns or complaints you may have about the research, contact the Partners Human Research Committee at 857-282-1900.