UNIVERSITY OF MICHIGAN
CONSENT TO BE SCREENED FOR ELIGIBILITY IN A RESEARCH STUDY

INFORMATION ABOUT THIS DOCUMENT:
You are being invited to take part in a research study conducted at several different locations (multi-site research). The University of Michigan is providing IRB oversight for all sites in this study. This consent form includes information that applies to all study sites.

Study Title: Persist Study

GENERAL INFORMATION

We’re doing a study to learn about new programs aimed at helping people manage chronic pain and medications. The program sessions focus on educational information and strategies for managing pain and medications. Before you can join the study, we’ll need to make sure you qualify. To find out whether you qualify, we’ll ask you some questions about your treatment, substance use, and pain experience. These questions will take about 5 minutes to answer.

If you qualify to be in the study and you are interested in joining, we’ll give you another consent form to read and sign. That form will explain the rest of the study.

Taking this survey to find out whether you qualify for our study is voluntary. You don’t have to take part if you don’t want to. Choosing not to take part won’t affect your medical care in any way. It’s possible that some of the questions may make you feel uncomfortable. You can skip any questions that you don’t want to answer, whatever the reason, and you don’t have to tell us why. Even if you do qualify for the study and decide to join, you can change your mind later and leave the study.

Determining whether you qualify for the study won’t benefit you directly.

Like the information in your medical record, the records we create in this study will remain confidential and protected. The computerized surveys are designed and administered using the REDCap database (https://www.project-redcap.org/). REDCap is dedicated to protect all customer data using industry best standards. For more information, REDCap security and privacy statements can be found at https://www.iths.org/wp-content/uploads/About-REDCap-Vanderbilt.pdf All paper surveys will be entered into the REDCap database and destroyed or kept in a locked file cabinet. Your answers to the survey will not be connected to your name unless you qualify and sign up for the next part of the study. If you sign up for the next part of the study, we will include your screening survey answers with the rest of your study data.

CONTACT INFORMATION
To find out more about the study, to ask a question or express a concern about the study, or to talk about any problems you may have as a study subject, you may contact one of the following:

<table>
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<tr>
<th>Site Principal Investigators:</th>
<th>Mark Ilgen, PhD and Lewei (Allison) Lin, MD</th>
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</table>
| Site Principal Investigator Contact: | 2800 Plymouth Road  
Ann Arbor, MI 48109  
Telephone: (734) 845-3646 (Ilgen)  
(734) 845-3637 (Lin) |
| Site Study Coordinator: | Mandy Lewis, MS |
| Site Study Coordinator Contact: | 2800 Plymouth Road  
Ann Arbor, MI 48109  
Telephone: (734) 936-1386 |

You may also express a concern about a study by contacting the Institutional Review Board:

University of Michigan Medical School Institutional Review Board (IRBMED)  
2800 Plymouth Road  
Building 520, Room 3214  
Ann Arbor, MI 48109-2800  
734-763-4768  
E-mail: irbmed@umich.edu

If you are concerned about a possible violation of your privacy or concerned about a study, you may contact the University of Michigan Health System Compliance Help Line at 1-866-990-0111.

**VERBAL CONSENT**

Do you agree to participate in this research study?

- [ ] Yes  
- [ ] No

I confirm that this individual has verbally agreed to participate in this research study.  
Screening ID#_______  
Staff Signature: ________________________ RA#: ____________________  
Date: ________________________________