ATTACHMENT 5: PRACTITIONER INFORMED CONSENT

Screening, Brief Intervention, Referral and Treatment (SBIRT) Practitioner Survey Consent to Participate in Research

RTI International is currently conducting an evaluation of the Screening, Brief Intervention, Referral and Treatment (SBIRT) program designed to help expand care for substance use and misuse. The SBIRT program includes screening, brief intervention, brief treatment, and referrals for persons at risk for dependence on alcohol or drugs. This survey is sponsored by the Department of Health and Human Services and is conducted in collaboration with this facility. We are asking you as a staff member of this facility to participate in this important anonymous survey.

You will be asked questions about your background, work environment, and your experiences with the SBIRT program. Completing the survey will only take about <u>18</u> minutes, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. There is a very important benefit to the study in that it will help us to gain knowledge about substance use care and prevention programs.

All the information you provide in this survey will be kept strictly private and will not be shared with anyone from this facility. RTI will not be documenting your name and will not be able to connect your survey responses to your name. At no time will your employer or anyone else not connected with the research see your responses. If the results of this study are presented at scientific meetings or published in scientific journals, no information will be included that could identify you or your answers personally.

Your decision to take part in this evaluation study and your participation is completely voluntary. You can refuse any part of the study and you can stop participating at any time. You can refuse to answer any question.

If you have any questions about the study, you may call (project team member's name and toll-free telephone number). If you have any questions about your rights as a study participant, you may call (IRB contact person).

Completion and return of the survey implies your consent to participate in this research. Please keep this form for your records.