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## **A sample size calculator for SMART pilot studies\***

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#### **Abstract**

In clinical practice, as well as in other areas where interventions are provided, a sequential individualized approach to treatment is often necessary, whereby each treatment is adapted based on the object's response. An adaptive intervention is a sequence of decision rules which formalizes the provision of treatment at critical decision points in the care of an individual. In order to inform the development of an *adaptive intervention*, scientists are increasingly interested in the use of *sequential multiple assignment randomized trials (SMART)*, which is a type of multi-stage randomized trial where individuals are randomized repeatedly at critical decision points to a set treatment options. While there is great interest in the use of SMART and in the development of adaptive interventions, both are relatively new to the medical and behavioral sciences. As a result, many clinical researchers will first implement a **SMART** pilot study (i.e., a small-scale version of a SMART) to examine feasibility and acceptability considerations prior to conducting a full-scale SMART study. A primary aim of this paper is to introduce a new methodology to calculate minimal sample size necessary for conducting a SMART pilot.

## **1 Introduction**

In the medical and behavioral health sciences, researchers have successfully established evidence-based treatments for a variety of health disorders. However, even with such treatments, there is heterogeneity in the type of individuals who respond and do not respond to treatment. Treatment effects may also vary over time (within the same individual): a treatment that improves outcomes in the short-run for an individual may not improve outcomes longer-term. Further, certain evidence based treatments may be too expensive to provide to all individuals; in such cases, health care providers may reserve these treatments for individuals who do not respond to less costly alternatives. The converse is also true: certain treatments are more ideally suited as maintenance treatments, and may be reserved for individuals who respond to earlier treatments in order to sustain improvements in outcomes. As a result, clinical researchers have recently shown great interest in developing sequences of treatments that are adapted over time in response to each individual's needs. This approach is promising because it allows clinicians to capitalize on the heterogeneity of

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treatment effect. An adaptive intervention offers a way to guide the provision of treatments over time, leading to such individualized sequence of treatments.

An *adaptive intervention*  $[1, 2, 3]$  is a sequence of decision rules that formalizes the provision of treatment at critical decision points in the course of care. In other words, an adaptive intervention is a guideline that can aid clinicians in deciding which treatment to use, for which individuals to use them, and when to use them. Figure 1 depicts a concrete example of an adaptive intervention for young children who are initially diagnosed with pediatric anxiety disorder.

In this example of *adaptive intervention*, first, clinicians offer the medication *sertraline* [4] for initial 12 weeks. If the child does not show an adequate response to it at the end of week 12, the clinician offers to augment the treatment with a combination of the medication sertraline and *cognitive behavioral therapy* [5] for additional 12 weeks. Otherwise the clinician would continue the *sertraline* medication for another 12 weeks. In this *adaptive* intervention, response is defined based on a measure of improvement, for example, based on a cut-off of five or less on the Clinical Global Impression-Improvement Scale [6]. Change in the *Pediatric Anxiety Rating Scale* could also be used to define response/non-response [7]. An *adaptive intervention* is also known as an *adaptive treatment strategy* [8] or a *dynamic* treatment regime [9].

Recently, methodologists introduced a specific type of randomized trial design known as a Sequential Multiple Assignment Randomized Trials [1, 10, 11] to inform the development of high-quality, empirically-supported adaptive interventions. A SMART is a type of multi-stage trial where each subject is randomly (re)assigned to one of various treatment options at each stage. Each stage corresponds to a critical treatment decision point. Each randomization is intended to address a critical scientific question concerning the provision of treatment at that stage; together, these help to inform the development of a high-quality adaptive intervention. Lei et al. [1] reviews a number of SMART studies in behavioral interventions science. Also, see work by Almirall et al. [10]

An example *SMART* is provided in Figure 2. This example could be used to develop an adaptive intervention for children who are diagnosed with pediatric anxiety disorder. At the first stage, there are two treatment options, *sertraline* medication or *cognitive behavioral*  $thereasy(CBT)$ . Each subject is randomly assigned to one of the initial treatment options and the assigned treatment is conducted for the first 12 weeks. At the end of week 12, each subject's response to the treatment is assessed based on *Clinical Global Impression*-Improvement Scale [4] and categorized as a responder or as a non-responder. Based on this, those who do not respond to the initial treatment are again randomly assigned to one of two secondary treatment options: One is a switch strategy whereby the child is switched to the stage 1 treatment option they were not offered at first. The second option is the combination of both *sertraline* medication and *cognitive behavioral therapy(CBT)*. For those who responded by the end of 12 weeks, continually initial intervention will be used. As with stage 1, both stage 2 treatments are provided for 12 weeks.

In a SMART such as the one shown in Figure 2, research outcomes may be collected at the end of week 24 or throughout, from baseline to week 24. Research outcomes may be continuous, e.g., Pediatric Anxiety Rating Scale [4] or discrete, e.g., Clinical Global Impression-Improvement Scale [4]. Note that the measure of response versus nonresponse at week 12 is not necessarily a research outcome. It is purely a criterion to categorize participants of the first stage intervention into a group of responders or that of non-responders.

The SMART study described above can be used to address three key scientific questions in the development of an *adaptive intervention* for *pediatric anxiety disorder:* (1) 'Which treatment to use in stage 1, medication or CBT?' and (2) 'Which tactic is best for nonresponders to stage 1 treatment, switch or augment?' Both of these questions involve randomized comparisons. Question (1) is addressed by comparing outcome measures of subgroup A, B and C in the Figure 2 with outcome measures of subgroup D,E and F in the Figure 2. Question (2) is addressed by comparing subgroup B and C when medication sertraline is stage 1 treatment and subgroup E and F when *CBT* is stage 1 treatment. Lastly, (3) This *SMART* design could also be used to compare the following four *adaptive* interventions contained within it.

- **1.** First, offer sertraline medication for 12 weeks. If the patient does not respond well to initial medication at the end of week 12, augment by initiating a combination therapy (sertraline and CBT) for next 12 weeks. Otherwise continue with the medication *sertraline* for another 12 weeks. (Children in subgroups A and B provided data for this adaptive intervention.)
- **2.** First, offer sertraline medication for 12 weeks. If the patient does not respond well to initial medication at the end of week 12, switch the treatment to CBT for next 12 weeks. Otherwise continue with the medication sertraline for another 12 weeks. (Children in subgroups A and C provided data for this adaptive intervention.)
- **3.** First, offer CBT medication for 12 weeks. If the patient does not respond well to initial medication at the end of week 12, augment by initiating a combination therapy (sertraline and CBT) for next 12 weeks. Otherwise continue with the medication CBT for another 12 weeks. (Children in subgroups D and E provided data for this *adaptive intervention*.)
- **4.** First, offer CBT medication for 12 weeks. If the patient does not respond well to initial medication at the end of week 12, switch the treatment to *sertraline* medication for next 12 weeks. Otherwise continue with the medication CBT for another 12 weeks. (Children in subgroups D and F provided data for this adaptive intervention.)

Note that the type of design used for SMART study described in Figure 2 is one of the most frequently used design. It has been used in *SMART* study of *adolescent marijuana use* [12], cocaine dependence [13, 14] and youth with conduct disorders [15]. For more recent ongoing SMART studies, visit the website: <http://methodology.psu.edu/ra/smart/projects>.

For detailed data analysis method regarding SMART, see the work of Nahum-Shani et al. [11,16].

Despite the advantages of *SMARTs*, it is fairly new to clinical research. Therefore, researchers may have concerns over the feasibility or acceptability of conducting a SMART. Feasibility refers to the capability of the investigators to perform the SMART and the ability of clinical staff (i.e., staff providing treatment) to treat subjects with the *adaptive* interventions in the *SMART*. For example, psychologists or psychiatrists delivering the stage 1 treatments may have concerns about the way non-response is defined; it is important to work out these concerns prior to a full-scale SMART study. Acceptability refers to the tolerability of the *adaptive interventions* being studied from the perspectives of study participants, as well as the appropriateness of the decision rules from the perspective of the clinical staff. For instance, some parents may object to a switch strategy (they may, instead, prefer an augmentation or an intensification strategy). If this happens often, investigators may re-consider the acceptability of the switch strategy prior to conducting a full-scale SMART. In such cases, researchers may conduct SMART pilot study to resolve feasibility and acceptability concerns prior to performing the full-scale SMART study.

The design of any study (pilot or full-scale randomized trial) requires researchers to select an appropriate sample size in order to conduct the study. In full scale randomized trials (including SMARTs), the sample size is typically determined to ensure sufficient statistical power to detect a minimally clinically significant treatment effect. For example, in a full scale SMART study, such as the one shown in Figure 2, the sample size could be determined to provide sufficient power (e.g., 80%) to detect a minimaly clinically significant treatment effect between any two of the four embedded adaptive interventions [17].

However, because the primary aim of pilot studies centers on acceptability and feasibility considerations, the sample size for pilot studies is not based on statistical power considerations [18, 19, 20, 21]. For the  $SMARKT$  pilot study, the goal is to examine feasibility and acceptability of conducting a full-scale trial. One approach for selecting a sample size achieving this is to observe sufficient number of participants for each subgroup from A to E in Figure 2. This is because each subgroup corresponds to a particular sequence of treatments and if the investigator does not have an ample amount of participants in each group, they cannot detect potential problems regarding feasibility or acceptability of certain sequence of treatments prior to conducting full-scale *SMART*. The primary aim of this paper is to introduce a new method which calculates a minimal sample size of SMART pilot study.

In Section 2, we develop a methodology for calculating a minimal sample size for SMART pilot studies that are like the pediatric anxiety disorder SMART presented above. In Section 3, we verify the result using simulations. We also compare our proposed methodology with an pre-existing method [22]. In Section 4, we extend the method in Section 2 to other types of SMART designs (the pediatric anxiety SMART described above represents just one type of SMART design). In Section 5, we provide a summary and discussions including areas for future work.

#### **2 A method for calculating the sample size for a SMART pilot study**

#### **2.1 The Proposed Approach**

In this section, we develop a sample size calculator for SMART pilot studies. We first develop an approach for the *SMART* study shown in Figure 2. Note that the method we will provide can be used in any area of SMART study whose design is identical to the one in Figure 2. Later, in Section 4, we generalize the method for other types of SMART designs. The approach provides investigators planning a SMART pilot study a principled way to choose a sample size for the pilot study, such that a minimal number of participants are observed in subgroups A-F in Figure 2. This is important because if the investigators do not observe sufficient number of participants of one particular sequence of treatments, the investigator cannot judge whether the sequence of treatments is actually feasible or can be accepted. For example, suppose that to examine feasibility and acceptability concerns, an investigator wishes to observe at least three participants in each of the subgroups A-F in Figure 2: in this case, how many participants should the investigators recruit in the study? Because the exact number of non-responders is unknown ahead of the pilot study, a probabilistic argument is necessary to answer such a question.

To formalize this idea, we first define some notations. Let N denote the total sample size of the  $SMRAT$  pilot study. For simplicity, we assume N is always a multiple of two; later we discuss the implications of this. Let  $m$  denote the minimum number of participants that an investigator would like to observe in subgroups A-F. Let  $q_j$  denote the anticipated rate of non-response to stage 1 treatment where  $j = SERT$  or CBT and let  $q = min(q_{SERT}, q_{CBT})$ , which will be used as a common non-response rate; the implications of using the minimum will also be discussed later. Lastly, a lower bound for the probability of the event that each subgroup will have at least m number of participants is denoted as  $k$ . Note that m, q and  $k$ are all provided by the investigator planning the SMART pilot. Hence, our goal is to provide a formulae for N as a function of m, q and k. More formally, the goal is to find a smallest N which satisfies

 $P$ ( all subgroups A-F have at leastm participants ) > k.

Using our notation, the above is equivalent to

$$
\mathbb{P}(M_A \ge m, M_B \ge m, M_C \ge m, M_D \ge m, M_E \ge m \text{ and } M_F \ge m) > k \tag{1}
$$

where  $M_A$  stands for the number of participants who fall into subgroup A.  $M_B$ ,  $M_C$ ,  $M_D$ ,  $M_E$ and  $M_F$  are defined in a similar way, respectively. Note that  $M_A$ ,  $M_B$ ,  $M_C$ ,  $M_D$ ,  $M_E$  and  $M_F$ are all discrete random variables. Next, we re-express (1) as

$$
\mathbb{P}(M_A \ge m, M_B \ge m \text{ and } M_C \ge m) \cdot \mathbb{P}(M_D \ge m, M_E \ge m \text{ and } M_F \ge m) > k. \tag{2}
$$

This is because, by the design of  $SMRAT$  study in Figure 2, any event of  $M_A$ ,  $M_B$  and  $M_C$  is independent to that of  $M_D$ ,  $M_E$  and  $M_F$ . Next let  $M_{NS}$  denote the number of non-responders out of N who were initially assigned to *sertraline* medication; and, similarly, let  $M_{\text{NC}}$  denote

the number of non-responders initially assigned to *CBT*. We first consider the leftmost probability term involving  $M_A$ ,  $M_B$  and  $M_C$ . Notice the event that  $M_B \ge m$  and  $M_C \ge m$  is equivalent to the event that  $M_{\text{NS}} \geq 2m$ . This is because, once the number of non-responders of sertraline medication is greater than or equal to  $2m$ , regardless of whether  $M_{\text{NS}}$  is odd or even, both  $M<sub>B</sub>$  and  $M<sub>C</sub>$  would be at least m due to a block randomization[23] with equal probabilities. For the case, when  $M_{\text{NS}}$  is odd, we exclude a participant and proceed the second stage randomization. Therefore, we get

$$
\mathbb{P}(M_A \ge m, M_B \ge m \text{ and } M_C \ge m) = \mathbb{P}(M_A \ge m, M_{NS} \ge 2m) = \mathbb{P}\left(\frac{N}{2} - M_{NS} \ge m, M_{NS} \ge 2m\right).
$$

A similar argument can be applied to  $M_D$ ,  $M_E$  and  $M_F$  and we have

$$
\mathbb{P}(M_{\textnormal{D}} \geqslant m, M_{\textnormal{E}} \geqslant m \textnormal{and} M_{\textnormal{F}} \geqslant m) = \mathbb{P}(M_{\textnormal{D}} \geqslant m, M_{\textnormal{NC}} \geqslant 2m) = \mathbb{P}\Big(\frac{N}{2} - M_{\textnormal{NC}} \geqslant m, M_{\textnormal{NC}} \geqslant 2m\Big).
$$

Re-expressing  $(2)$ , our goal is to find the smallest N such that

$$
\mathbb{P}\left(\frac{N}{2} - M_{\rm NS} \ge m, M_{\rm NS} \ge 2m\right) \cdot \mathbb{P}\left(\frac{N}{2} - M_{\rm NC} \ge m, M_{\rm NC} \ge 2m\right) > k \tag{3}
$$

which is equivalent to,

$$
\mathbb{P}\left(\frac{N}{2} - m \ge M_{\rm NS} \ge 2m\right) \cdot \mathbb{P}\left(\frac{N}{2} - m \ge M_{\rm NC} \ge 2m\right) > k.
$$
 (4)

Now note that

$$
M_{NS} = \sum_{i=1}^{N/2} X_i, M_{NC} = \sum_{i=1}^{N/2} Y_i,
$$

where  $X_i = 1$  if the ith participant assigned to *sertraline* medication did not respond well or  $X_i = 0$  otherwise. Since the probability of non-response to *sertraline* is assumed to be  $q$ , we have that  $X_n$  has a Bernoulli distribution with success probability  $q$  [24]. Similarly,  $Y_n$  has a Bernoulli distribution with success probability  $q$  (recall the assumption that the probability of non-response is assumed to be q for both sertraline and CBT). Therefore,  $M_{\text{NS}}$  and  $M_{\text{NC}}$ have identical distributions, which we denote by the random variable Mq. Further, given the result that the sum of independent identically distributed Bernoulli random variables has a Binomial distribution [24], we have that

$$
\mathbb{P}\left(\frac{N}{2} - m \ge M_q \ge 2m\right)^2 > k,\tag{5}
$$

where  $M_{q}$  Binomial  $\left(\frac{N}{2}, q\right)$  or, equivalently,

$$
\left(\mathbb{P}\left(\frac{N}{2} - m \ge M_q\right) - \mathbb{P}(2m - 1 \ge M_q)\right)^2 > k \tag{6}
$$

#### holds as well.

As a side note, if we have an odd number of participants, it is impossible to assign an equal number of participants to each initial intervention. Therefore we set  $N$  to be a multiple of 2, this is because, by the design of our SMART study in Figure 2, there is a block randomization in stage  $1$  [23]. Setting N as a multiple of 2 allows us to assign equal number of participants to two treatment options provided at the first stage. Additionally we use a minimum value of the two non-response rate  $(q_{\text{SERT}}, q_{\text{CBT}})$  as a common non-response rate $(q)$ . This is because, by using a minimum value of two non-response rates, we will get a robust sample size which satisfies (6).

**2.1.1 Implementation—**For fixed values for  $m$ ,  $k$  and  $q$ (i.e., provided by the scientists designing a  $SMRAT$  pilot), a suitable value of N can be found by searching for the smallest N such that (6) holds true. This is possible because (6) is an inequality with respect to N assuming that  $m$ ,  $k$  and  $q$  are given. This can be easily accomplished using any computer program capable of calculating upper tail probabilities for random variables with Binomial distributions(e.g., the *pbinom* function in R [25]).

Using the implementation outlined above, Table 1 provides values of N for a range of inputs of m, k, and q. For example, suppose an investigator wishes that at least  $3(m)$  participants are observed in each subgroup with probability greater than  $0.8(k)$ , and assumes that the common non-response rate is  $0.30(q)$ . Based on the Table 1 below, the investigator needs to recruit at least 58 participants for the SMART pilot study. Note that in this paper, we provide sample sizes for the q values in a range between  $0.2$  to  $0.8$  because for the non-response rate values below 0.2 or above 0.8, as it may not be feasible to conduct a SMART studies.

**2.1.2 A Pre-existing Approach—**A similar approach to calculate the sample size for SMART pilot studies was first proposed by Almirall et al. [22]. Their proposal centered on finding the smallest sample size  $N$  which satisfies

 $\mathbb{P}(M_{\rm B} > m-1, M_{\rm C} > m-1, M_{\rm E} > m-1 \text{ and } M_{\rm F} > m-1) > k$ .

This differs from our proposed approach, which requires that all six subgroups A-F have at least m participants with probability greater than  $k$ (see  $(1)$ ). The use of this objective function was based on the argument that in typical *SMART* studies, the rate of non-response is often not very large(i.e less than equal to  $0.60$ ). Therefore, in such settings it is highly likely that if the condition that the number of participants in subgroups B, C, E and F are respectively greater than  $m-1$  was required, the number of responders in subgroups A and D would also be greater than  $m-1$ , respectively.

On top of that the pre-existing method had an assumption that the event:  $M_B > m - 1$  &  $M_C$  $> m - 1$  is equivalent to the event:  $M_{\text{NS}} > 2m - 2$ , which may not be true. Consider the

case when  $M_{\text{NS}} = 2m - 1$ . If  $M_{\text{NS}} = 2m - 1$ , either  $M_{\text{B}}$  or  $M_{\text{C}}$  should be  $m - 1$ , which violates the condition given:  $M_B > m - 1$  &  $M_C > m - 1$ . In other words, the condition:  $M_B > m - 1$  &  $M_C > m - 1$  implies that  $M_{NS} > 2m - 2$ , but not necessarily in the other way around.

Therefore the sample size we get from the pre-existing method will not guarantee that the investigator would observe at least m number of people for each subgroup with probability greater than  $k$ . In next section, we will conduct a simulation study to check validity of the pre-existing method by comparing the simulation result of the pre-existing method with that of the new method introduced in Section 2.1.

#### **3 Simulation**

A simulation experiment is conducted (i) to verify that sample sizes obtained under the proposed approach satisfy equation (6) under a variety of realistic values for  $m$ ,  $k$  and  $q$ , and (ii) to compare the performance of the proposed method with the pre-existing method by Almirall et al. [22], described above.

The simulation experiment is conducted in the following way for each combination of values of  $m$ ,  $k$  and  $q$ .

- **1.** Firstly, the values m, k and q are used to calculate the minimum suggested sample size N based on the proposed methodology.
- **2.** Secondly, using this sample size N, we simulate the flow of participants through one realization of the SMART shown in Figure 2. Specifically, we divide the total sample size( $N$ ) by 2. Then by *rbinom* function [25] in R, we obtain the number of responders and non-responders for each pilot SMART simulation, which allows us to get the number of participants in each of the subgroups A-E.
- **3.** Thirdly, we check if the number of participants in each subgroup is greater than pre-specified  $m$  or not. If the condition is met, we count it as a successful SMART pilot study. This process is repeated for 10,000 times. In the end, after 10,000 simulations, we obtain the proportion of successes out of 10,000. This represents an estimate of the left-hand side of expression (1), which we take it as a true proportion, denoted as  $\rho$ , since we are conducting 10,000 times of *Monte* Carlo simulation.
- **4.** Lastly, the proportion(ρ) obtained in previous step is compared with a prespecified lower bound for the proportion(k). If this proportion( $\rho$ ) is greater than k value, we conclude the sample size obtained from the proposed method is valid. Otherwise, the proposed sample size is invalid. The Table 2 provides the results of this experiment.

Notice that the number of non-responsers could be an odd number. In this case, we subtract one from the number of non-responsers and divide by two. Then we use this value to check if it is greater than  $m$  or not. This is to (i) to get a conservative sample size and (ii) to avoid having non integer value of participants in each subgroup. If both values are greater than  $m$ , then we count this trial SMART pilot as a successful SMART pilot study.

From the simulation result, we can assess if the sample sizes we get from the proposed method, which are in Table 1, are valid or not. For instance, when  $m = 3$ ,  $k = 0.80$  and  $q = 0.30$ , we need to have at least 58 participants to conduct a *SMART* pilot study based on the Table 1. Then, from Table 2, we can see that out of 10,000 simulations, roughly in 8,160 (10,000·0.816) times, the condition that there are 3 or more people in each subgroup is satisfied. Since all the values we get from the simulation are greater than corresponding  $k$ value which is in the left end column, we can say that our new method developed in previous section is valid.

To see whether the method discussed in Section 2.1 is an improved version, a simulation study is also conducted, in a same manner, for the pre-existing method [22]. As one can see in Table 3, pre-existing method failed to prove its validity. Again, this is because of the assumptions discussed in the Section 2.1.2.

### **4 Extensions to other SMART pilot studies**

Not all *SMART* studies will be like the type shown in Figure 2. In the *SMART* in Figure 2, all non-responders were re-randomized at the second stage regardless of the initial treatment assignment; i.e., re-randomization to second-stage treatment depended only on response/ non-response status. In a second type of commonly-used SMART design, re-randomization at the second stage depends on both initial treatment and response/non-response status. In a third type of commonly-used SMART design, both responders and non-responders are re-randomized at the second stage. In this section, we extend the methods of Section 2.1 to these two types of SMART designs.

#### **4.1 Re-randomization depends on initial treatment and response status**

In this section, we consider SMART studies where re-randomization to second-stage treatment depends on the choice of initial treatment as well as response/non-response status.

As an example, consider the SMART shown in Figure 3. This SMART study was designed to develop adaptive interventions for improving linguistic and social communication outcomes among children with autism spectrum disorders who are minimally verbal [26]. Specifically, this SMART examined the effects of three adaptive interventions involving different provisions of a speech generating device (SGD; a type of Augmentative and Alternative Communication Interventions). This SMART was facilitated to answer two scientific questions in the context of a behavioral language intervention (BLI) for children with autism [27, 28]. Initially, all children were randomized at stage 1 to BLI versus BLI+SGD for 12 weeks to answer question (1): Is providing SGD more effective at initial stage? At the end of week 12, each participant is categorized as a responder or a nonresponder to stage 1 treatment based on 14 measures including: 7 communication variables from natural language sample with blinded assessor and 7 communication variables from intervention transcripts [26]. All responders continued on stage 1 treatment for an additional 12 weeks. All non-responders to BLI+SGD received intensified BLI+SGD. Non-responders to BLI were re-randomized to intensified BLI versus BLI+SGD to answer question (2): For non-responders to BLI, is providing SGD with BLI as a rescue intervention more efficacious than intensifying the initial intervention? Total number of spontaneous communicative

utterances, primary outcome of the study, was collected at week 24 with a follow-up collection at week 36.

The derivation of the sample size formulae for a pilot study of a *SMART* of this type is similar to the derivation in Section 2.1. One difference in the notation is that in this SMART design, there are 5 subgroups, labeled A to E. Our goal is to determine the smallest  $N$  which guarantees that

 $P$ ( all subgroups A-E have at least *m* participants ) > k

Using arguments similar to those used in section 2.1 (see Appendix A), one can show that this inequality is identical to

$$
\left[\mathbb{P}\left(\frac{N}{2} - m \ge M_q\right) - \mathbb{P}(2m - 1 \ge M_q)\right] \cdot \left[\mathbb{P}\left(\frac{N}{2} - m \ge M_q\right) - \mathbb{P}(m - 1 \ge M_q)\right] \tag{7}
$$

Notice that, unlike with the inequality given in equation (6), the left-hand-side of inequality (7) does not reduce to the square of a probability. This is due to the imbalance in the SMART design shown in Figure 3(only non-responders to one of the initial treatments are re-randomized) relative to the design shown in Figure 2(where all non-responders are re-randomized). Given k, q, and m, a solution for N in expression (7) can be found using an approach that is similar to the one described earlier to solve expression (6). Table 4 provides a minimal sample size for the type of SMART designs in Figure 3.

See the work of Kilbourne et al. [29], which employs a SMART of this type to enhance outcomes of a mental disorders program.

#### **4.2 Both responders and non-responders are re-randomized**

In this section, we consider a third type of SMART design where both responders and non-responders are re-randomized. As an example of this type of design, we present a study of individuals with alcoholic use disorder. The example SMART design is shown in Figure 4. The goal of this SMART study, which is reviewed in greater detail in Lei et al. [1], was to develop adaptive interventions for individuals with alcoholic use disorders. This SMART was used to answer three scientific questions regarding the use of naltrexone medication (NTX) [30], an opioid receptor antagonist, for the management and prevention of relapse among individuals with alcohol use disorder. All participants were provided  $NTX$  medication as a stage 1 treatment. Non-response to  $NTX$  was measured on a weekly basis. Participants were randomized initially to two different definitions for non-response to NTX–a lenient versus a more stringent definition–to answer the question (1): What extent of weekly drinking activity is best regarded as non-response? The lenient definition of non-response was defined as having five or more heavy drinking days per week, whereas the stringent definition of non-response was defined as having two or more heavy drinking days per week. Participants identified as non-responders to NTX were re-randomized to the combination of combined behavioral intervention (CBI) [31, 32], medical management  $(MM)$  [33] versus to the combination of *NTX, CBI* and *MM*. This randomization answers

the question (2): What type of treatments would be useful for subjects who do not respond well to *NTX*? If participants had not been identified as non-responders by week 8, they were said to be responders to stage 1 intervention. Responders were re-randomized to the NTX versus to the combination of NTX and telephone disease management (TDM) to answer the question (3): What type of treatments would be effective for reducing the chance of relapse among people who responded well to NTX? Primary outcomes included the percentage of heavy drinking days and percentage of drinking days of the last two months of the study.

The variables  $m$ ,  $k$ , and  $q$  are defined as in Section 2.1 (see the Appendix B). In this type of SMART, there are 8 subgroups, labeled A through H. In addition, randomization occurs both for responders and non-responders. Our goal is to find a smallest  $N$  which satisfies

 $P$ ( all subgroups A-H have at least *m* participants ) > k

In Appendix B we show that the above equation is true if and only if

$$
\left[\mathbb{P}\left(\frac{N}{2} - 2m \ge M_q\right) - \mathbb{P}(2m - 1 \ge M_q)\right]^2 > k,\tag{8}
$$

As you can see in expression (8), unlike in expression (7), since the design is perfectly symmetric we have two identical probability terms multiplied each other. In addition, unlike expression (6), instead of m, 2m was subtracted from  $\frac{N}{2}$ . This is because, for the type of SMART designs described in Figure 4, responders were also randomized. For more detailed explanation on how this influences the method, see Appendix B. Again, given  $m$ ,  $k$  and  $q$ , a solution for N in expression  $(8)$  can be found using an approach that is similar to the one described earlier to solve expression (6). Table 5 provides a minimal sample size for the type of SMART design in Figure 4.

A number of other SMART studies are similar to the type shown in Figure 4. These studies include a SMART for developing an adaptive reinforcement-based behavioral intervention for woman who are pregnant and abusing drugs [34]; a SMART study aimed at developing an adaptive intervention involving individual and family-delivered cognitive behavioral therapy among children with depression; and a SMART designed to develop an adaptive intervention for children with autism spectrum disorders who are minimally verbal. All three of these studies are currently in the field.

#### **5 Discussion**

This manuscript presents pilot sample size calculators for three of the most common types of sequential multiple assignment randomized trial (SMART) designs. As stated in the introduction, researchers use *SMARTs* to inform the development of *adaptive interventions*. More specifically, *SMART* designs can be used to address critical scientific questions that need to be answered in order to construct high-quality *adaptive interventions*. Over the last 15 years, SMART designs have become more popular among clinical and health service researchers. However, some researchers may have concerns regarding the feasibility

of conducting a full scale SMART or the acceptability of the treatments or adaptive interventions embedded in a SMART design. Such researchers may choose to conduct a smaller-scale pilot *SMART* prior to conducting a full-scale *SMART*. Specifically, a *SMART* pilot study is a small scale version of a full scale SMART study, where the primary purpose is to examine the acceptability and feasibility issues. See the following papers for more detailed explanations and concrete examples of *SMART* pilot studies [15, 22, 35].

This paper develops an approach for determining the minimum sample size necessary for conducting a pilot SMART. The number of participants for SMART pilot study should be enough to address concerns in feasibility and acceptability of full-scale SMART study. The paper introduces one way to operationalize this, which is to ensure that each subgroup corresponding to sequence of treatments to observe some minimum number $(m)$ of participants. This approach was used to select the sample sizes for two recent SMART pilot studies: 1) SMART for developing an adaptive intervention for adolescent depression [35], 2) SMART for adolescent conduct problems [15]. Further, the methods are developed for three of the most commonly used types of *SMART* designs. Finally, we compare our proposed method with the pre-existing, related method to calculate a sample size for a SMART pilot [22] and explain how the proposed methodology is an improvement on the pre-existing one. In addition, the characteristics of the methodologies developed in this paper were examined thoroughly via Monte Carlo simulation. Specifically, for each type of SMART design, 10,000 simulation SMART pilot studies were conducted with different combinations of values of  $m, k$  and  $q$  via statistical software R. In all possible combinations of  $m, k$  and  $q$ , the simulation study supported that the condition imposed on the sample  $size(M)$  was met.

The method may be conservative in that, based on the way the rate of non-response is elicited from the scientist, the method may suggest a sample size that is as large or larger than the sample size actually needed to meet the constraint. Specifically, our proposed approach elicits the minimum value of the non-response rates to first-stage treatments. This was done to minimize the burden on the investigator of having to guess/provide two non-response rate. In settings where the two non-response rates differ, using the minimum for both may lead to conservative sample size requirements, relative to a method which uses both of two different non-response rates. For the future work, one can possibly develop a new methodology to calculate minimum sample size for a SMART pilot using two non-response rates. Also one can further investigate, in which circumstances (i.e. which combinations of  $m$ ,  $k$  and  $q$ ), a method that uses two non-response rate values results in substantially small sample size than the method introduced in this paper.

Some suggestions on choosing values for  $m$ ,  $k$  and  $q$  are provided in this paragraph. Concerning  $q$ : Existing data from previous studies (not necessarily a previous  $SMARK$ study) are often used to obtain estimates of q. Typical values of non-response rates for SMART ranges from 0.3 to 0.7. Concerning m: In many cases, we have found that investigators are interested in observing between 3 and 5 participants for each subgroup of a pilot SMART. Note that for typical pilot studies, resources, including the maximum number of participants that could be afforded in a pilot study, are often limited. And observing between 3 to 5 people for each subgroup is typically enough to assess feasibility

and acceptability issues regarding *adaptive interventions*. Concerning  $k$ : typical values range from 0.8 to 0.95.

This manuscript provides a way to choose a sample size for a pilot *SMART*, to examine feasibility and acceptability concerns before conducting a full scale SMART study. Another possible approach is to choose a sample size so that investigators may observe an estimate of response/non-response rate with pre-specified amount of precision. Researchers may want to adopt this approach to estimate non-response rate. By using the estimate of non-response rate, researchers can implement the methodologies described in the paper. For instance, suppose we want a sample size N which allows us to estimate non-response rate( $q$ ) within the margin of error of 0.1 with significance level 0.05. Then, we estimate  $q$  by the proportion of non-responders among total sample  $(\hat{q})$ . Since the number of non-responders follows a Binomial distribution with parameters q and  $N[24, 36]$ , after some calculation, we get  $N=$ 100. Note that the above example is just to illustrate another way to calculate sample size for pilot study. In this way, one can come up with an alternative way to develop a sample size calculator for SMART pilot study. For more detailed technical explanation, see Appendix C.

#### **6 Acknowledgements**

There are numerous people who I would like to thank. Dr. Daniel Almirall, from the Institute of Social Research at the University of Michigan, introduced this whole new research area regarding SMART. He also gave numerous suggestions on possible directions of the work. In addition, Dr. Edward Ionides, from the Department of Statistics at the University of Michigan, provided invaluable advice in communications in academia. His insights helped me to sharpen the whole arguments in the paper. I was also lucky to talk with  $Dr$ . Lu Xi, from the Institute of Social Research at the University of Michigan, who always patiently answered my questions regarding SMART and related fields. Lastly, I also want to show my gratitude to all the members of the statistical reinforcement learning lab in the Department of Statistics at the University of Michigan for providing a wonderful environment to get involved in research.

## **Appendix**

#### **A Appendix A**

Here, we provide a mathematical derivation of equation (7). All the variables used here are defined in a similar way as in Section 2.1. Recall that what we want to calculate is the smallest N such that

 $P$ ( all subgroups A-E have at least *m* participants ) > k

holds. Using mathematical expression, one can write it as

 $\mathbb{P}(M_A \ge m, M_B \ge m, M_C \ge m, M_D \ge m \text{ and } M_E \ge m) > k$ .

By the independence of the group with *behavioral language intervention* and the group with both

behavioral language intervention and speech generating device, the above is same as

 $\mathbb{P}(M_A \geq m, M_B \geq m \text{ and } M_C \geq m) \cdot \mathbb{P}(M_D \geq m, M_E \geq m) > k$ .

Let  $M_{\text{NB}}$  denote the number of non-responders of *behavioral language intervention*. Note that the event:  $M_B \ge m$  and  $M_C \ge m$  is same as the event:  $M_{NB} \ge 2m$  due to a block randomization with equal probabilities [23]. Therefore we get

$$
\mathbb{P}(M_A \ge m, M_B \ge m \text{ and } M_C \ge m) = \mathbb{P}\left(\frac{N}{2} - M_{\text{NB}} \ge m, M_{\text{NB}} \ge 2m\right).
$$

Let  $M<sub>NBS</sub>$  denote the number of non-responders of the initial intervention which involves both behavioral language intervention and speech generating device. Then,

$$
\mathbb{P}(M_{\text{D}} \geq m, M_{\text{E}} \geq m) = \mathbb{P}\Big(\frac{N}{2} - M_{\text{NBS}} \geq m, M_{\text{NBS}} \geq m\Big)
$$

Therefore we find the smallest  $N$ , which satisfies

$$
\mathbb{P}\left(\frac{N}{2}-M_{\text{NB}} \geqslant m, M_{\text{NB}} \geqslant 2m\right) \cdot \mathbb{P}\left(\frac{N}{2}-M_{\text{NBS}} \geqslant m, M_{\text{NBS}} \geqslant m\right) > k,
$$

One can re-write above as

$$
\mathbb{P}\left(\frac{N}{2}-M_{\rm q}\geqslant m, M_{\rm q}\geqslant 2m\right)\cdot\mathbb{P}\left(\frac{N}{2}-M_{\rm q}\geqslant m, M_{\rm q}\geqslant m\right)>k,
$$

where  $M_q$  follows a Binomial distribution with size parameter  $\frac{N}{2}$  and probability parameter q. Re-arranging  $M_q$ , we have

$$
\mathbb{P}\left(\frac{N}{2} - m \ge M_q \ge 2m\right) \cdot \mathbb{P}\left(\frac{N}{2} - m \ge M_q \ge m\right) > k,
$$

which is analogous to

$$
\left[\mathbb{P}\left(\frac{N}{2}-m\geq M_{\rm q}\right)-\mathbb{P}\big(2m-1\geqslant M_{\rm q}\big)\right]\cdot\left[\mathbb{P}\Big(\frac{N}{2}-m\geqslant M_{\rm q}\Big)-\mathbb{P}\big(m-1\geqslant M_{\rm q}\big)\right]>k\,.
$$

## **Appendix**

#### **B Appendix B**

Here, we provide a mathematical derivation of equation (8). All the variables used here are defined in a similar way as in Section 2.1. Recall that what we want to calculate is the smallest N such that

 $P$ ( all subgroups A – H have at least *m* participants ) > k

holds. Using mathematical expression, one can write it as

$$
\mathbb{P}(M_A \geqslant m, M_{\small \textrm{B}} \geqslant m, M_{\small \textrm{C}} \geqslant m, M_{\small \textrm{D}} \geqslant m, M_{\small \textrm{E}} \geqslant m, M_{\small \textrm{F}} \geqslant m, M_{\small \textrm{G}} \geqslant m \textrm{ and } M_{\small \textrm{H}} \geqslant m) > k \,.
$$

By design, we know that the group with the lenient definition of non-response and the group with the stringent definition of non-response are independent. Therefore, the above expression is same as,

$$
\mathbb{P}(M_A \ge m, M_B \ge m, M_C \ge m \text{ and } M_D \ge m) \cdot \mathbb{P}(M_E \ge m, M_F \ge m, M_G \ge m \text{ and } M_H \ge m) > k
$$

Let  $M<sub>NL</sub>$  denote the number of non-responders for the initial intervention with lenient definition of non-response. Similarly we define  $M_{\text{NS}}$  as the number of non-responders for the initial intervention with stringent definition of non-response. Our next step is to re-express above expression in terms of  $M_{NL}$ ,  $M_{NS}$  and N. Note that the event:  $M_C \ge m$  and  $M_D$  is same as the event:  $M_{NL} \ge 2m$ . In addition, the event:  $M_A \ge m$  and  $M_B \ge m$  is same as the event:  $\frac{N}{2} - M_{NL} \ge 2m$ . Analogous arguments can be applied to the event involving subgroup E through H. Therefore we get,

$$
\mathbb{P}(M_\text{A}\geqslant m, M_\text{B}\geqslant m, M_\text{C}\geqslant m \text{ and } M_\text{D}\geqslant m)=\mathbb{P}\Big(\frac{N}{2}-M_\text{NL}\geqslant 2m \text{ and } M_\text{NL}\geqslant 2m\Big)
$$

and

$$
\mathbb{P}(M_{\text{E}} \ge m, M_{\text{F}} \ge m, M_{\text{G}} \ge m \text{ and } M_{\text{H}} \ge m) = \mathbb{P}\Big(\frac{N}{2} - M_{\text{NS}} \ge 2m \text{ and } M_{\text{NS}} \ge 2m\Big).
$$

Therefore our goal is to find a sample size  $N$ , which satisfies

$$
\mathbb{P}\left(\frac{N}{2} - M_{\rm NL} \ge 2m \text{ and } M_{\rm NL} \ge 2m\right) \cdot \mathbb{P}\left(\frac{N}{2} - M_{\rm NS} \ge 2m \text{ and } M_{\rm NS} \ge 2m\right) > k
$$

One can re-write above as

$$
\mathbb{P}\left(\frac{N}{2}-M_{\mathrm{q}}\geq 2m, M_{\mathrm{q}}\geq 2m\right)\cdot\mathbb{P}\left(\frac{N}{2}-M_{\mathrm{q}}\geq 2m, M_{\mathrm{q}}\geq 2m\right) > k,
$$

where  $M_q$  follows a Binomial distribution with size parameter N and probability parameter <sup>q</sup>. Then one can further simplify as

$$
\mathbb{P}\Big(\frac{N}{2}-2m\geq M_{\rm q}\geq 2m\Big)\cdot\mathbb{P}\Big(\frac{N}{2}-2m\geq M_{\rm q}\geq 2m\Big]>k,
$$

where the above is equivalent to

$$
\left[\mathbb{P}\left(\frac{N}{2}-2m\geq M_{\rm q}\right)-\mathbb{P}\left(2m-1\geq M_{\rm q}\right)\right]^2>k\,.
$$

### **Appendix**

## **C Appendix C**

In this section, we provide technical explanation of finding sample size using margin of error. Recall that the number of non-responders follows a Binomial distribution with parameters q and  $N[24, 36]$ . One can show that  $\hat{q}$ , a proportion of non-responders among total sample, is an unbiased estimate of  $q$  and its variance and standard deviation are below [37]:

$$
Var(\hat{q}) = \frac{q(1-q)}{N} \Rightarrow sd(\hat{q}) = \sqrt{\frac{q(1-q)}{N}}.
$$

Then the goal is to find a sample size  $N$  which satisfies

$$
2 \cdot \sqrt{\frac{q(1-q)}{N}} = 0.1.
$$

However, since we do not know the true value of q, we instead use  $\frac{1}{2}$  as a value of q to find conservative sample size of N[36, 38]. By solving above formula after plugging in  $\frac{1}{2}$  to q, we have

$$
\frac{1}{\sqrt{N}} = 0.1 \Rightarrow N = 100.
$$

#### **References**

- [1]. Lei H, Nahum-Shani I, Lynch K, Oslin D, and Murphy SA. A "smart" design for building individualized treatment sequences. Annual review of clinical psychology, 8, 2012.
- [2]. Murphy Susan A and Collins Linda M. Customizing treatment to the patient: Adaptive treatment strategies. Drug and alcohol dependence, 88(Suppl 2):S1, 2007.
- [3]. Murphy Susan A, Lynch Kevin G, Oslin David, McKay James R, and TenHave Tom. Developing adaptive treatment strategies in substance abuse research. Drug and alcohol dependence, 88: S24–S30, 2007. [PubMed: 17056207]
- [4]. Compton Scott N, Walkup John T, Albano Anne Marie, Piacentini John C, Birmaher Boris, Sherrill Joel T, Ginsburg Golda S, Rynn Moira A, McCracken James T, Waslick Bruce D, et al. Child/adolescent anxiety multimodal study (cams): rationale, design, and methods. Child and Adolescent Psychiatry and Mental Health, 4(1):1, 2010. [PubMed: 20051130]
- [5]. Compton Scott N, March John S, Brent David, Albano Anne Marie, Weersing V Robin, and Curry John. Cognitive-behavioral psychotherapy for anxiety and depressive disorders in children and adolescents: an evidence-based medicine review. Journal of the American Academy of Child & Adolescent Psychiatry, 43(8):930–959,2004. [PubMed: 15266189]
- [6]. Busner Joan and Targum Steven D. The clinical global impressions scale: applying a research tool in clinical practice. Psychiatry (Edgmont), 4(7):28, 2007.
- [7]. Caporino Nicole E, Brodman Douglas M, Kendall Philip C, Albano Anne Marie, Sherrill Joel, Piacentini John, Sakolsky Dara, Birmaher Boris, Compton Scott N, Ginsburg Golda, et al. Defining treatment response and remission in child anxiety: signal detection analysis using the pediatric anxiety rating scale. Journal of the American Academy of Child & Adolescent Psychiatry, 52(1):57–67, 2013. [PubMed: 23265634]

- [8]. Murphy Susan A and McKay James R. Adaptive treatment strategies: An emerging approach for improving treatment effectiveness. Clinical Science, 12:7–13, 2004.
- [9]. Murphy Susan A. Optimal dynamic treatment regimes. Journal of the Royal Statistical Society: Series B (Statistical Methodology), 65(2):331–355, 2003.
- [10]. Almirall Daniel, Nahum-Shani Inbal, Sherwood Nancy E, and Murphy Susan A. Introduction to smart designs for the development of adaptive interventions: with application to weight loss research. Translational behavioral medicine, 4(3):260–274, 2014. [PubMed: 25264466]
- [11]. Nahum-Shani Inbal, Qian Min, Almirall Daniel, Pelham William E, Gnagy Beth, Fabiano Gregory A, Waxmonsky James G, Yu Jihnhee, and Murphy Susan A. Experimental design and primary data analysis methods for comparing adaptive interventions. Psychological methods, 17(4):457, 2012. [PubMed: 23025433]
- [12]. Brown Pamela C, Budney Alan J, Thostenson Jeff D, and Stanger Catherine. Initiation of abstinence in adolescents treated for marijuana use disorders. Journal of substance abuse treatment, 44(4):384–390, 2013. [PubMed: 23085041]
- [13]. McKay James R. Treating substance use disorders with adaptive continuing care. American Psychological Association, 2009.
- [14]. McKay James R, Van Horn Deborah HA, Lynch Kevin G, Ivey Megan, Cary Mark S, Drapkin Michelle L, Coviello Donna M, and Plebani Jennifer G. An adaptive approach for identifying cocaine dependent patients who benefit from extended continuing care. Journal of consulting and clinical psychology, 81(6):1063, 2013. [PubMed: 24041231]
- [15]. August Gerald J, Piehler Timothy F, and Bloomquist Michael L. Being "smart" about adolescent conduct problems prevention: Executing a smart pilot study in a juvenile diversion agency. Journal of Clinical Child & Adolescent Psychology, (ahead-of-print):1–15,2014.
- [16]. Nahum-Shani Inbal, Qian Min, Almirall Daniel, Pelham William E, Gnagy Beth, Fabiano Gregory A, Waxmonsky James G, Yu Jihnhee, and Murphy Susan A. Q-learning: A data analysis method for constructing adaptive interventions. Psychological methods, 17(4):478, 2012. [PubMed: 23025434]
- [17]. Scott Alena I, Levy Janet A, and Murphy Susan A. Evaluation of sample size formulae for developing adaptive treatment strategies using a smart design. 2007.
- [18]. Kraemer Helena Chmura, Mintz Jim, Noda Art, Tinklenberg Jared, and Yesavage Jerome A. Caution regarding the use of pilot studies to guide power calculations for study proposals. Archives of General Psychiatry, 63(5):484–489, 2006. [PubMed: 16651505]
- [19]. Thabane Lehana, Ma Jinhui, Chu Rong, Cheng Ji, Ismaila Afisi, Rios Lorena P, Robson Reid, Thabane Marroon, Giangregorio Lora, and Goldsmith Charles H. A tutorial on pilot studies: the what, why and how. BMC medical research methodology, 10(1):1, 2010. [PubMed: 20053272]
- [20]. Lancaster Gillian A, Dodd Susanna, and Williamson Paula R. Design and analysis of pilot studies: recommendations for good practice. Journal of evaluation in clinical practice, 10(2):307– 312, 2004. [PubMed: 15189396]
- [21]. Leon Andrew C, Davis Lori L, and Kraemer Helena C. The role and interpretation of pilot studies in clinical research. Journal of psychiatric research, 45(5):626–629, 2011. [PubMed: 21035130]
- [22]. Almirall Daniel, Compton Scott N, Gunlicks-Stoessel Meredith, Duan Naihua, and Murphy Susan A. Designing a pilot sequential multiple assignment randomized trial for developing an adaptive treatment strategy. Statistics in medicine, 31(17):1887–1902, 2012. [PubMed: 22438190]
- [23]. Suresh KP. An overview of randomization techniques: An unbiased assessment of outcome in clinical research. Journal of human reproductive sciences, 4(1):8,2011. [PubMed: 21772732]
- [24]. Ross Sheldon M. Introduction to probability models. Academic press, 2014.
- [25]. R Core Team. R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria, 2014. URL [http://www.R-project.org/.](http://www.R-project.org/)
- [26]. Kasari Connie, Kaiser Ann, Goods Kelly, Nietfeld Jennifer, Mathy Pamela, Landa Rebecca, Murphy Susan, and Almirall Daniel. Communication interventions for minimally verbal children with autism: a sequential multiple assignment randomized trial. Journal of the American Academy of Child & Adolescent Psychiatry, 53(6):635–646, 2014. [PubMed: 24839882]

- [27]. Kaiser Ann Pand Goetz Lori. Enhancing communication with persons labeled severely disabled. Research and Practice for Persons with Severe Disabilities, 18(3):137–142, 1993.
- [28]. Kaiser Ann P, Hancock Terry B, and Nietfeld Jennifer P. The effects of parent-implemented enhanced milieu teaching on the social communication of children who have autism. Early Education and Development, 11(4):423–446, 2000.
- [29]. Kilbourne Amy M, Almirall Daniel, Eisenberg Daniel, Waxmonsky Jeanette, Goodrich David E, Fortney John C, Kirchner JoAnn E, Solberg Leif I, Main Deborah, Bauer Mark S, et al. Protocol: Adaptive implementation of effective programs trial (adept): cluster randomized smart trial comparing a standard versus enhanced implementation strategy to improve outcomes of a mood disorders program. Implementation Science, 9(1):132, 2014. [PubMed: 25267385]
- [30]. Volpicelli Joseph R, Alterman Arthur I, Hayashida Motoi, and O'Brien Charles P. Naltrexone in the treatment of alcohol dependence. Archives of general psychiatry, 49(11):876–880,1992. [PubMed: 1345133]
- [31]. Longabaugh Richard, Zweben Allen, LoCastro Joseph S, and Miller William R. Origins, issues and options in the development of the combined behavioral intervention. Journal of Studies on Alcohol and Drugs, (15):179,2005.
- [32]. Anton Raymond F, O'Malley Stephanie S, Ciraulo Domenic A, Cisler Ron A, Couper David, Donovan Dennis M, Gastfriend David R, Hosking James D, Johnson Bankole A, LoCastro Joseph S, et al. Combined pharmacotherapies and behavioral interventions for alcohol dependence: the combine study: a randomized controlled trial. Jama, 295(17):2003–2017, 2006. [PubMed: 16670409]
- [33]. Pettinati Helen M, Weiss Roger D, Miller William R, Donovan Dennis, Ernst Denise B, Rounsaville Bruce J, COMBINE Monograph Series, and Mattson Margaret E. Medical management treatment manual. A Clinical Research Guide for Medically Trained Clinicians Providing Pharmacotherapy as Part of the Treatment for Alcohol Dependence. COMBINE Monograph Series, 2, 2004.
- [34]. Jones Hendree E, O'Grady Kevin E, and Tuten Michelle. Reinforcement-based treatment improves the maternal treatment and neonatal outcomes of pregnant patients enrolled in comprehensive care treatment. The American Journal on Addictions, 20(3):196–204, 2011. [PubMed: 21477047]
- [35]. Gunlicks-Stoessel Meredith, Mufson Laura, Westervelt Ana, Almirall Daniel, and Murphy Susan. A pilot smart for developing an adaptive treatment strategy for adolescent depression. Journal of Clinical Child & Adolescent Psychology, (ahead-of-print):1–15, 2015.
- [36]. Blyth Colin Rand Still Harold A. Binomial confidence intervals. Journal of the American Statistical Association, 78(381):108–116,1983.
- [37]. Casella George and Berger Roger L. Statistical inference, volume 2. Duxbury Pacific Grove, CA, 2002.
- [38]. Morisette Jeffrey Tand Khorram Siamak. Exact binomial confidence interval for proportions. Photogrammetric engineering and remote sensing, 64(4):281–282,1998.

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**SERT -> Sertraline Medication** CBT -> Cognitive Behavioral Therapy





**SERT <-> Sertraline Medication CBT** <-> Cognitive Behavioral Therapy  $R \leftarrow$ > Randomization

#### **Figure 2:**  An example SMART for pediatric anxiety disorder patients



**BLI** <-> Behavioral Language Intervention **SGD <-> Speech Generating Device**  $R \leftarrow$  Randomization

#### **Figure 3:**

An example SMART for Children with Autism





An example SMART for alcoholic patients

#### **Table 1:**

Minimal sample size of SMART pilot study based on the proposed method

	Range of <i>q</i> :	0.20	0.30	0.40	0.50	0.60	0.70	0.80
$k = 0.80$	$m = 3$	88	58	42	34	28	32	50
$k = 0.80$	$m = 4$	112	74	54	42	36	42	64
$k = 0.80$	$m = 5$	136	90	66	52	44	50	76
$k = 0.90$	$m = 3$	100	64	48	36	32	38	60
$k = 0.90$	$m = 4$	126	82	60	46	40	48	74
$k = 0.90$	$m = 5$	150	98	72	56	48	56	86

#### **Table 2:**

Simulation table of the sample sizes based on the proposed method

	Range of $q$ :	0.20	0.30	0.40	0.50	0.60	0.70	0.80
$k = 0.80$	$m = 3$	0.807	0.816	0.821	0.860	0.809	0.810	0.815
$k = 0.80$	$m = 4$	0.810	0.828	0.814	0.820	0.834	0.844	0.821
$k = 0.80$	$m = 5$	0.811	0.825	0.820	0.835	0.838	0.830	0.813
$k = 0.90$	$m = 3$	0.911	0.902	0.921	0.903	0.931	0.912	0.910
$k = 0.90$	$m = 4$	0.906	0.911	0.921	0.913	0.925	0.920	0.912
$k = 0.90$	$m = 5$	0.903	0.906	0.915	0.918	0.926	0.902	0.901

#### **Table 3:**

Simulation table of the sample sizes based on the pre-existing method

	Range of q:	0.20	0.30	0.40	0.50	0.60	0.70	0.80
$k = 0.80$	$m = 3$	0.633	0.662	0.623	0.616	0.475	0.194	0.000
$k = 0.80$	$m = 4$	0.650	0.664	0.650	0.672	0.549	0.353	0.000
$k = 0.80$	$m = 5$	0.675	0.691	0.678	0.641	0.592	0.000	0.000
$k = 0.90$	$m = 3$	0.790	0.780	0.766	0.803	0.769	0.401	0.000
$k = 0.90$	$m = 4$	0.796	0.822	0.818	0.813	0.786	0.000	0.000
$k = 0.90$	$m = 5$	0.821	0.820	0.831	0.841	0.806	0.459	0.000

#### **Table 4:**

Minimal sample size of SMART pilot study for nonverbal children with autism



#### **Table 5:**

Minimal sample size of SMART pilot study for alcoholic patient

	Range of $q$ :	0.20	0.30	0.40	0.50	0.60	0.70	0.80
$k = 0.80$	$m = 3$	88	58	42	36	42	58	88
$k = 0.80$	$m=4$	112	74	54	46	54	74	112
$k = 0.80$	$m = 5$	136	90	66	56	66	90	136
$k = 0.90$	$m = 3$	100	64	48	40	48	64	100
$k = 0.90$	$m = 4$	126	82	60	50	60	82	126
$k = 0.90$	$m = 5$	150	98	72	60	72	98	150