# Continuous monitoring of A/B tests without pain: Optional stopping in Bayesian testing

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Abstract—A/B testing is one of the most successful applications of statistical theory in the Internet age. A crucial problem of Null Hypothesis Statistical Testing (NHST), the backbone of A/B testing methodology, is that experimenters are not allowed to continuously monitor the results and make decisions in real time. Many people see this restriction as a setback against the trend in the technology toward real time data analytics. Recently, Bayesian Hypothesis Testing, which intuitively is more suitable for real time decision making, attracted growing interest as a viable alternative to NHST. While corrections of NHST for the continuous monitoring setting are well established in the existing literature and known in A/B testing community, the debate over the issue of whether continuous monitoring is a proper practice in Bayesian testing exists among both academic researchers and general practitioners. In this paper, we formally prove the validity of Bayesian testing under proper stopping rules, and illustrate the theoretical results with concrete simulation illustrations. We point out common bad practices where stopping rules are not proper, and discuss how priors can be learned objectively. General guidelines for researchers and practitioners are also provided.

Category and Subject Descriptors: G.3 [Probability and Statistics]: Statistical Computing

**Keywords:** A/B testing, controlled experiments, Bayes factor, optional stopping, continuous monitoring

#### I. Introduction

Many online service companies nowadays have been using online controlled experiments, a.k.a. A/B Testing, as a scientifically grounded way to evaluate changes and comparing different alternatives. A/B testing plays a leading role in establishing the mantra of data driven decision making, and is one of the basic pillars in Data Science.

Most of A/B tests are conducted using the statistical theory of frequentist Null Hypothesis Statistical Testing (NHST), namely t-test or z-test. Experimenters using NHST summarize the test result in a p-value and reject the null hypothesis  $H_0$  when the p-value is less than a prescribed significance level  $\alpha$  (often 0.05). The interpretation is that assuming all model assumptions are correct, doing so we can control the Type-I error, i.e. the probability of making a false rejection when  $H_0$  is true, to be no greater than  $\alpha$ .

Recently, interests in using Bayesian model comparison for two sample hypothesis testing are growing [8; 21; 15]. The type of statistical interpretations we make from Bayesian tests is fundamentally different from NHST. Under the Bayesian framework, we assume there is a prior probability  $P(H_1)$  for  $H_1$  (the alternative) to be true, and similarly  $P(H_0)$  for  $H_0$  to be true. The ratio between the two is called the prior odds. After collecting data from an experiment, we update prior odds into posterior odds (abbreviated as PostOdds) using the Bayes Rule:

$$PostOdds := \frac{P(H_1|Data)}{P(H_0|Data)} = \frac{P(H_1)}{P(H_0)} \frac{P(Data|H_1)}{P(Data|H_0)}, \quad (1)$$

which is commonly referred as

Posterior Odds = Prior Odds  $\times$  Bayes Factor.

Note that the Bayes Factor (BF) is the likelihood ratio of observing the data between  $H_1$  and  $H_0$ . From the posterior odds, it is straightforward to calculate the posterior probabilities  $P(H_1|Data)$  and  $P(H_0|Data)$ . Moreover we have

$$\frac{P(H_1|PostOdds)}{P(H_0|PostOdds)} = PostOdds, \tag{2}$$

and therefore

$$P(H_1|PostOdds) = PostOdds/(PostOdds + 1).$$
 (3)

Equation (2) (or its equivalent form (3)) has the following interpretation: when observing a posterior odds K, rejecting  $H_1$  will expose us to a risk of a false rejection/discovery with probability  $P(H_0|Data) = 1/(K+1)$ . In other words, the whole experiment result can be summarized by posterior odds.<sup>2</sup>

In this paper we are interested in a common practice called continuous monitoring or optional stopping. This practice is best described as the following Example.

Example 1 (Optional Stopping):

We observe data sequentially and at any time we can conduct statistical analysis on the data already observed. Let  $t=1,\ldots,N$  be all the interim check-points that we can take a peek at our A/B test results.<sup>3</sup> For any given metric M, let  $R_t$  be its test result at check-point t. We define an event  $S_t$  that is observed at time t and stop the experiment at the first t such that the event happens  $(R_t \in S_t)$  and return result  $R_t$ , e.g. when we deem the result is "significant" or "conclusive". Typically, the event  $S_t$  is defined as p-value  $< \alpha$  or  $P(H_0|Data) < r$ . If this event didn't happen at t = N

<sup>&</sup>lt;sup>1</sup>For this paper, we assume readers are already familiar with the concepts of Null Hypothesis Statistical Testing (NHST) in controlled experiments. Readers new to these concepts and A/B testing should refer to references such as Kohavi et al. [17].

<sup>&</sup>lt;sup>2</sup>(2) and (3) is a special case of Theorem 1 proven later.

<sup>&</sup>lt;sup>3</sup>A test result could be something like a test statistic, p-value or Bayes Factor/posterior. We use this vague notion when the detail is not important.

we return test result  $R_N$ . A general version involves infinite horizon.

Pitfalls of continuous monitoring under NHST framework have been documented in various publications[1; 24; 13]. We say the interpretation of the result is unchanged with continuous monitoring if the validity of the interpretation holds regardless of whether continuous monitoring is used. NHST is valid for fixed horizon test. But it is known to underestimate Type-I error when continuous monitoring is used. To quickly see why, if experimenters are allowed to stop the first time pvalue is less than 5%, we will only reject more often, but no less comparing to a fixed horizon design, because the event of rejection in a fixed horizon design, i.e. only reject at time N, is strictly a subset of the event of rejection in the continuously monitoring design. As a result, if the Type-I error in the fixed horizon design is 5%, the Type-I error with continuous monitoring will in general exceed 5%. An application of the law of iterated logarithm shows when incoming data are i.i.d. continuous monitoring will inflate Type-I error to 100% when the horizon N goes to infinity, see Siegmund [24]. Johari et al. [13] provided simulation results showing that the inflation of Type-I error is significant and could be typically above 50% or more.

In the Bayesian framework with continuous monitoring, posterior is still defined as in (1). However, the key problem is that, when a data-adaptive stopping rule is applied, it is unclear how to compute the Bayes Factor  $\frac{P(Data|H_1)}{P(Data|H_0)}$  because the probability space is on stochastic processes without a fixed length. This BF is easy to calculate if we are allowed to ignore the random stopping time and treat it as fixed. The main result of this paper justifies this practice.

Theorem 1: Let  $\mathbf{X}_t$  be all the observed data up to time t and  $BF_t$  be the Bayes Factor defined as  $\frac{P(\mathbf{X}_t|H_1)}{P(\mathbf{X}_t|H_0)}$  and posterior odds  $PostOdds_t$  defined as in (1) with a pre-defined, known prior odds  $P(H_1)/P(H_0)$ . Let  $\tau$  be any stopping time defined by a proper stopping rule. That is, a mechanism for deciding whether to continue or stop on the basis of *only the present and past events* and  $\tau$  is finite almost surely. Then the interpretation of posterior odds remains unchanged with optional stopping at  $\tau$ . Specifically, we have

$$P(H_1|PostOdds_{\tau}) = PostOdds_{\tau}/(PostOdds_{\tau} + 1).$$
 (4)

Theorem 1 says (3) is correct even when the posterior is calculated under a fixed horizon model by treating the random stopping time  $\tau$  simply as a fixed time. In other words, Equation (4) guarantees that the fixed horizon Bayesian test result remains the same interpretation even with continuous monitoring, provided that the Bayes Factor (and hence the posterior odds) are calculated using all available observations up to the stopping time  $\tau$ , and the stopping rule is properly defined to be based on only the present and past events. In particular, the theorem does not hold if Bayes Factor is calculated on a selected subset of the observations available at time t, or if the stopping rule peeked ahead into the future. These requirements are met in all common practices of continuous monitoring as in Example 1 where the stopping

time is called a "hitting time". In conclusion, Theorem 1 formally endorsed the practice of continuous monitoring in the framework of Bayesian Hypothesis Testing. This is in stark contrast to NHST, where special adjustment has to be done. Still, practices like "re-analyze the same data using continuous monitoring after failed to reject using all data" is not supported by Theorem 1. More bad practices are discussed later in Section V.

At the time of writing, to authors' best knowledge there is still a lack of general agreement on whether continuous monitoring is a proper practice when Bayes test is used because there is no concrete theorem or proof like Theorem 1, more details in Section II. The purpose of this paper is to provide arguments assessable by practitioners and engineers, while at the same time provide rigorous proofs for researchers in A/B testing community as well as related fields. With this main goal, the contributions of this paper are

- 1) We formally prove Theorem 1 in Section IV.
- 2) We adopt a simulation approach as in Rouder [20], to help readers understand the result and gain intuitions. We also emphasize what Theorem 1 does not guarantee.
- 3) For practitioners, we make recommendations on when and when not to use continuous monitoring. We put emphases on cases where Theorem 1 does not apply.
- 4) We discuss the importance of a known prior odds in Theorem 1 and learning this odds objectively from historical experiment data.

All model assumptions required in our models are taken as granted. Although both NHST and Bayesian tests make extra model assumptions, the latter requires more such as prior and distribution under  $H_1$ . In practice many people use subjective priors or so called non-informative priors. These practices have been criticized a lot since there are no agreement among researchers and practitioners on which prior is appropriate. However, with the existence of rich historical A/B tests data, we can learn prior objectively from the empirical data. We discuss this in more detail in Section VII.

The rest of the paper is structured as follows. We review related work in the next Section. Section III illustrates Theorem 1 using a simple simulation setup. Proof of the main theorem is in Section IV, with intuitive explanations. We emphasize bad practices where Theorem 1 does not apply in Section V. More simulation study and discussions about pros and cons of continuous monitoring are presented in Section VI. Section VII studies the important practical issue of objective prior learning. Section VIII concludes the paper with practical recommendations.

#### II. RELATED WORK

The need of a different theory to allow continuous monitoring in NHST framework has long been known as the subject of sequential hypothesis testing [25]. Sequential hypothesis testing and later on group sequential testing have been widely used in Clinical Trials [1]. The idea of sequential test was only recently popularized by Johari et al. [13] in A/B testing community, by including it as part of the offering of the commercial A/B testing platform Optimizely. Despite it being

newly introduced to A/B testing community, the theories behind sequential tests under NHST frameworks are well known by statisticians and practitioners in related areas such as clinical trials, psychology, econometrics and other social sciences.

Bayesian hypothesis testing, on the other hand, is much less accepted and established than its frequentist counterpart. This was largely due to the need of prior knowledge that commonly requires a subjective choice or so called "noninformative" priors which also lacks justification. Putting the issue of choosing a prior aside, many Bayesians have argued that Bayesian reasoning should be immune to stopping rule. For example, Dawid [6] brought up the notion of conditional independence and argued that posterior based on stopping time shouldn't alter likelihood ratios. This issue is also discussed in Berger and Berry [3] and later Berger and Bayarri [2] referred to the idea as the "stopping rule principle" and said "once the data have been obtained, the reasons for stopping experimentation should have no bearing on the evidence reported about unknown model parameters." Although the idea is well received by leading Bayesians, there are still a lot of debates going on among researchers and practitioners on whether Bayesian testing and more generally Bayesian analysis is adjustment-free when optional stopping is applied. John K. Kruschke made the point that Bayesian testing can be biased under optional stopping in 2013 [18], in which he used simulation to show Type-I error could be higher when using stopping rule based on Bayes Factor. Andrew Gelman claimed that optional stopping "is Kosher" in Bayesian analysis in 2014 [12]. This debate is also still heated in Psychon. Bull. Rev., a journal where Bayesian hypothesis testing is relatively well received. In a 2014 paper Rouder [20] used simulation to support the case of Bayesian test with optional stopping, and to counter criticisms from Erica et al. [11] and Sanborn and Hills [22], both published also in 2014. Recently Schönbrodt [23] also supports optional stopping for Bayesian test with discussions on the issue of parameter estimation bias. However, no proof in a concrete setting is provided in the literature beyond simulation studies. A lot of dispute mentioned above are due to misunderstanding of what kind of promises are Bayesian tests making and should still keep with optional stopping. This paper set out to settle this issue with a rigorous theorem. We believe the lack of a general agreement on this issue even in mid 2010s is a clear sign that this is still a big problem for researchers and practitioners in various areas. This is especially the case for A/B testing community because 1) data are always received in near real time in a sequential fashion, 2) the technology enables and even encourages experimenters to frequently check out the test results.

#### III. BAYESIAN PROMISES IN FIXED HORIZON TEST

Instead of explaining and proving Theorem 1 right away, we explain what are we going to prove in Theorem 1 for the basic fixed horizon test before we go into optional stopping. This section serves two purposes. First, a big part of the

debates about whether Bayesian test is biased with continuous monitoring is due to wrong interpretations of the Bayesian test result itself, and using frequentist measurements such as Type-I error to evaluate a Bayesian test result. This is largely due to most researchers, especially statisticians, are trained with frequentist statistics and methodologies. A correct interpretation of Bayesian posterior odds and Bayes Factor is a prerequisite for readers to understand and appreciate Theorem 1 and rest of the paper. To this end, for most data scientists and engineers, we found replicable simulation results is more tangible and concrete than probability formulas. Secondly, through the simulation results, we hope readers will glimpse some intuitions on why Theorem 1 is indeed expected even with optional stopping. Simulation setup in this section follows prior work of Rouder [20].

Recall Posterior Odds = Prior Odds × Bayes Factor. Prior odds is considered known and is independent of the observations collected for the test. Prior odds is easy to interpret and interpretation of Bayes Factor, hence the posterior odds is the essence of a Bayesian test. Without loss of generality, from now on we will assume a prior odds of 1:1 and leave the problem of how to objectively pick prior odds in Section VII. In this case *Posterior Odds and Bayes Factor are the same*. Readers can treat them as interchangeable in this paper.

We consider a simple problem of testing a normal mean. We observe N i.i.d. observations  $X_i, i=1,\ldots,N$  from a normal distribution  $N(\mu,1)$  with unknown mean  $\mu.$   $\mu=0$  under the null hypothesis  $H_0$ , and  $\mu=\delta$  under the alternative hypothesis  $H_1$ . Equivalently, sample mean  $\overline{X}$  is the sufficient statistics and it has distribution N(0,1/N) under  $H_0$  and  $N(\delta,1/N)$  under  $H_1$ . Note that this is even simpler than a A/B test because there is only one group. For two sample A/B test we replace  $\overline{X}$  by  $\Delta=\overline{X}_T-\overline{X}_C$ , e.g. difference of two sample means, and the test is essentially the same as one sample test. For details see Section VII.

The Bayes Factor is

$$\frac{P(\overline{X}|H_1)}{P(\overline{X}|H_0)} = \frac{\exp(-(\overline{X} - \delta)^2/(2/N))}{\exp(-(\overline{X})^2/(2/N))} = \exp\left(\frac{N}{2}\delta(2\overline{X} - \delta)\right) \quad (5)$$

Conditioning on observing a  $\overline{X}$ , if we plug it into Equation 5 and get a number K, what does it mean? To illustrate this, we simulate 100,000 runs and each run we simulate N=100 observations  $X_i, i=1,\ldots,N$ . Since we assume prior odds 1:1, we simulate 50,000 runs under  $H_1$ , where  $X_i \sim N(\delta,1)$  and the other 50,000 runs under  $H_0$  where  $X_i \sim N(0,1)$ . At the end of each run, we calculate Bayes Factor based on Equation 5. The end result of this simulation is 100,000 Bayes Factors, half of them are from  $H_1$  and half of them from  $H_0$ .

We did this simulation for  $\delta=0.2$  and N=100. Figure 1 shows histograms of those Bayes Factors in log scale, grouped by the ground truth models  $H_0$  and  $H_1$ . Bayes Factors from  $H_1$  are shown on top and those from  $H_0$  are shown at the bottom. What does it mean if we observe a Bayes Factor of 2.1? On Figure 1, we first group Bayes Factors close to 2.1 together in to the same bin. There are about 4,000 runs from  $H_1$  (height of the red bar) that produced a Bayes Factor close

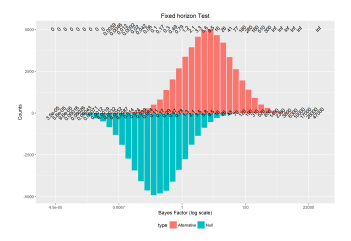


Fig. 1. Histograms of Bayes Factor simulated from both models. For each bin, the number on top are the ratio of simulated Bayes Factors from alternative to those from null.

to 2.1 (among 50,000 simulation runs), while around 2,000 (height of the blue bar) are from  $H_0$  (among 50,000 simulation runs). The actual ratio of those from  $H_1$  to  $H_0$  is shown on top of the plot and is 2.1, which is the same as the Bayes Factor 2.1 we started with. In fact, if you go through each bin carefully in Figure 1, you will find the number on the x-axis, which represents Bayes Factor value calculated from Equation 5 and grouped into each bin, are very close to the actual observed count ratio of those from  $H_1$  (top red) to those from  $H_0$ (bottom green), except those at the far tail on both sides. Is this a coincidence? Of course not. When observed a Bayes Factor of K, we know both model  $H_0$  and  $H_1$  could result in such a Bayes Factor. This simulation we did let us replay the data generation process and observe how likely it is for  $H_1$  to generate such a Bayes Factor and how likely for  $H_0$  respectively, which are represented, after binning similar Bayes Factors together, by the height of the top and bottom histograms. Our interest is the odds of this Bayes Factor being from  $H_1$  to  $H_0$ , which is the ratio of heights between the red bar and the green bar. We will expect the observed ratio to be close to the true underlying odds, within some small expected error due to 1) simulation randomness and 2) discretization used in binning similar Bayes Factor together. The error from simulation randomness is smaller for those center bins, i.e. bins where more Bayes Factors are observed from 100,000 simulation runs, and are larger for those at the tails.<sup>4</sup> What we observed so far can be summarized as: For each posterior odds bin

> True underlying odds = Observed ratio = Bayes Factor calculated from Equation 5.

This simulation illustrated two things:

1) Bayes Factor can be conceptually "materialized" as the ratio of the bar heights from the  $H_1$  histogram and  $H_0$ 

- histogram. An observed Bayes Factor of K means it is K times more likely to be generated from  $H_1$  than  $H_0$ .
- 2) For the fixed horizon case, Equation 5 is the same as the true odds (at least they must be very close).

**Bayesian Promise** The above simulation illustrates the Bayesian Promises: Equation (2) (and (3) as its direct consequence). After we've collected all the simulation results, we summarize each run into a Bayes Factor/posterior odds. Conditioned on any posterior odds, the observed ratio is an estimate for LHS of (2), and the posterior odds calculated from 5 is the RHS of (2). In other words, we have seen from simulation that without continuous monitoring, a fixed horizon Bayesian test of  $H_1$  vs.  $H_0$  keeps the Bayesian Promise (2).

What Theorem 1 tells us is that the same Bayesian Promise is kept when optional stopping is used with a proper stopping rule. The fixed horizon case is a special case of proper stopping rule where the stopping time is fixed and independent of data.

#### IV. PROOF OF MAIN THEOREM

We now prove Theorem 1. Readers who only need intuition are recommended to skip the proof and jump to Section IV-A. Both sides of the (4) are random variables depending on  $PostOdds_{\tau}$ . It is equivalent to the following:

$$\begin{split} &\frac{P(H_1|PostOdds_{\tau} = K)}{P(H_0|PostOdds_{\tau} = K)} = \frac{P(H_1 \text{ and } PostOdds_{\tau} = K)}{P(H_0 \text{ and } PostOdds_{\tau} = K)} \\ &= \frac{P(PostOdds_{\tau} = K|H_1)}{P(PostOdds_{\tau} = K|H_0)} \times \frac{P(H_1)}{P(H_0)} = K, \end{split}$$

for any K where  $P(PostOdds_{\tau} = K) > 0$ . Let  $K' = K \times P(H_0)/P(H_1)$ . The event  $\{PostOdds_{\tau} = K\}$  is equivalent to  $\{BF_{\tau} = K'\}$ . The last equality above after rearranging the prior odds  $P(H_1)/P(H_0)$  to the right side becomes

$$\frac{P(BF_{\tau} = K'|H_1)}{P(BF_{\tau} = K'|H_0)} = K'.$$
 (6)

Without loss of generality, we only need to prove (6).

We first prove for the fixed horizon case, which is a direct result of *likelihood ratio identity*, or *change of measure* [9]. For any fixed t, let  $\mathbb{Q}_t = P(\cdot|H_1)$  and  $\mathbb{P}_t = P(\cdot|H_0)$  be the probability measure under  $H_1$  and  $H_0$  respectively for observations up to t, both have a density function with respect to Lebesgue measure on the real line. Let A be any event observable at time t.<sup>5</sup> The likelihood ratio identity<sup>6</sup> ensures

$$\mathbb{Q}_t(A) = E_{\mathbb{P}_t} \left( \mathbb{1}_A \times \frac{d\mathbb{Q}_t}{d\mathbb{P}_t} \right),$$

where  $d\mathbb{Q}_t/d\mathbb{P}_t$  is the likelihood ratio and  $\mathbb{1}_A$  is the binary indicator function for event A. Recall Bayes Factor is defined as the likelihood ratio. Replace  $d\mathbb{Q}_t/d\mathbb{P}_t$  by  $BF_t$ , set  $A=\{BF_t=K'\}$  in the above to get

$$\mathbb{Q}_t(BF_t = K') = E_{\mathbb{P}_t}(\mathbb{1}_A) \cdot K' = K' \cdot \mathbb{P}_t(BF_t = K'), \tag{7}$$
 which is (6).

<sup>&</sup>lt;sup>4</sup>Some bins on the two tails are either showing an observed ratio of 0 or Inf, for the obvious reason.

 $<sup>{}^5</sup>A \in \mathcal{F}_t$  where  $\mathcal{F}_t$  represents the set of measurable events at time t.  $\mathcal{F}_t$  is called a filtration because  $\mathcal{F}_s \subseteq \mathcal{F}_t$  for any  $s \le t$ .

 $<sup>^6</sup>$ It is also called change of measure identity because the equation transforms an expectation under a measure  $\mathbb P$  into an expectation under another measure  $\mathbb Q$ . This is a special case of the Radon-Nykodym Theorem in measure theory.

We can generalize this argument for random time  $\tau$ . Theorem 1 requires  $\tau$  to be a stopping time t so that the event  $\{\tau=t\}$  is observable at time t. This is a necessary requirement to ensure that we can apply the likelihood ratio identity for the event  $\{BF_t=K' \text{ and } \tau=t\}$  (observable at t) to get

$$\mathbb{Q}_t(BF_t = K', \tau = t) = K' \times \mathbb{P}_t(BF_t = K', \tau = t). \tag{8}$$

If  $\tau$  can only take value from 1 to a maximum horizon N (experiment stop at N no matter what, which covers all practical cases), summing up (8) over all t entails

$$P(BF_{\tau} = K'|H_{1}) = \sum_{t=1}^{N} P(BF_{\tau} = K', \tau = t|H_{1})$$

$$= \sum_{t=1}^{N} \mathbb{Q}_{t}(BF_{t} = K', \tau = t) = \sum_{t=1}^{N} K'\mathbb{P}_{t}(BF_{t} = K', \tau = t)$$

$$= \sum_{t=1}^{N} K'P(BF_{\tau} = K', \tau = t|H_{0}) = K'P(BF_{\tau} = K'|H_{0})$$

which is (6) and the proof is completed. Notice how we changed  $BF_{\tau}=K'$  to  $BF_{t}=K'$  once we restrict ourselves to the set  $\tau=t$  in the second and fifth equality. The essence of the proof is to show

$$\frac{P(BF_{\tau} = K', \tau = t|H_1)}{P(BF_{\tau} = K', \tau = t|H_0)} = K'$$
(9)

for every  $t \leq N$  by applying likelihood ratio identity (8) and then sum up both numerator and denominator of (9) over t to recover (6). For potentially unbounded  $\tau$ , we just need to sum up to infinity and the result still holds because the sum series of both numerator and denominator of (9) are finite.

**Important Remark**. We make the remark that Theorem 1 does not require observation X to be sequential i.i.d. observations as in earlier simulation examples. All we used in the proof is the likelihood ratio identity for the whole path of observations  $X_t$  up to t for any t. In a typical A/B test with user level tracking, users who visit the site multiple times will provide multiple observations sequentially. Since there is a strong between-user correlation, when we look at  $\mathbf{X}_t$ , which includes all sequential observations (page-views) from different users, they are not independent. However, at any given time t, we can always first aggregate  $X_t$  to the randomization unit, which is user. Take the metric Revenue per user as example,  $X_t$  is the sequence of revenues for each page-view up to time t. For each user, we can sum up revenues as  $Y_{it}$ ,  $i = 1, ..., N_t$  where  $N_t$  is number of unique users. We can treat  $Y_{it}$  as i.i.d. when computing likelihoods under both  $H_1$  and  $H_0$ . For ratio metrics such as Click-Through-Rate(CTR),  $Y_{it}$  can either be CTR for each user and average of  $Y_{it}$  is the average CTR over all users — a double average metric, or  $Y_{it}$  can be a pair  $(Clicks_{it}, PageViews_{it})$  and the metric is the sum of clicks over all users divided by sum of page-views. Delta Method is required to compute the likelihood for the latter case. The main point is that the original sequential observations might not be *i.i.d.* but after aggregated to randomization unit level, likelihood ratio can be easily calculated using those aggregated values which can be assumed *i.i.d.* because of the randomization design.

## A. Intuitive Explanation

There is an intuitive explanation of Theorem 1 using the same simulation procedure in Section III as a thought experiment. Again we assume prior odds is 1:1 so Posterior Odds equals to the Bayes Factor and (4) becomes (6).

We simulate M paths of sequential observations from  $H_1$ and  $H_0$ . M is a very large number, almost infinite. So every path simulated from  $H_1$  which has nonzero probability under  $H_0$  will have the same path simulated under  $H_0$ , and vice versa. For each path we simulate the whole path X up to the fixed final horizon N. For any  $t \leq N$  and any path X, we can calculate a Bayes Factor at time t to be  $BF(\mathbf{X}_t)$  as likelihood ratio  $P(\mathbf{X}_t|H_1)/P(\mathbf{X}_t|H_0)$ . No stopping rule has been introduced yet so everything so far belongs to the fixed horizon case. Define  $Path(\mathbf{X}_t|H_i)$ , i=0,1 to be the set of all paths simulated from  $H_i$ , i = 0, 1 having the same subpath  $\mathbf{X}_t$  up to time t, and let  $|Path(\cdot)|$  denote the total number of paths in a path set, i.e. cardinality. Then for any t and any path X with subpath  $\mathbf{X}_t$ ,  $|Path(\mathbf{X}_t|H_1)|/|Path(\mathbf{X}_t|H_0)| = BF(\mathbf{X}_t)$ . Intuitively, this means for every subpath  $X_t$  simulated from  $H_0$ , there are on average  $BF(\mathbf{X}_t)$  exact subpaths simulated from  $H_1$ . Because this statement is true for any subpath. If we only look at subpaths such that  $BF(\mathbf{X}_t) = K$ , we have  $|Path(\{\mathbf{X}_t : BF(\mathbf{X}_t) = K\}|H_1)|/|Path(\{\mathbf{X}_t : BF(\mathbf{X}_t) = K$  $|K||H_0|| = K$ , for any K.

Now we introduce stopping rule. Pick any path X simulated from  $H_0$ , say the stopping rule will stop at time t and we computed the Bayes Factor to be K. The previous argument shows there will be on average K number of the same exact subpath simulated from  $H_1$ . Here comes the important part! Because the stopping rule does not depend on observations after the stopping time, all subpaths simulated having the exact same subpath  $X_t$  up to t will also have the exact same stopping time at t! (See the next section for examples of bad stopping rules where this property is not true, hence Theorem 1 does not apply.) After we gathered all paths simulated from  $H_0$ with the same Bayes Factor K at time t which also stopped at time t according to the stopping rule, for each one of them we can find K exact same subpaths which also stopped at time t. By one more step of gathering all such set of paths for every possible  $t \leq N$ , it is then intuitively clear that the number of paths gathered together from  $H_1$  and  $H_0$  have a ratio of exactly K. This is exactly what we tried to demonstrate via various simulations in Section III.

# V. BAD PRACTICES

Theorem 1 is a general result with very mild assumptions which are satisfied in most cases. But failure of satisfying those assumptions can result invalid test results. We list three bad practices so readers can be aware of the limitations of the result in this paper. One critical assumption is that the stopping rule is properly defined that only uses information already

 $<sup>^{7}\</sup>tau$  is a stopping time with respect to a filtration  $\mathcal{F}_{t}$  if  $\{\tau \leq t\} \in \mathcal{F}_{t}$ .

observed, without peeking into the future. One example for an improper stopping rule is to reassess all the observations at some time t', and then decide to only use the data up to an earlier time t < t', e.g. stop at t after seeing data at a later time t'. This practice is called *data snooping* and is not supported by Theorem 1. There are two common bad practices related to data snooping:

Example 2 (Re-analysis after Fail to Reject):

Finite horizon test at N failed to reject  $H_0$ . The same data is then reanalyzed using continuous monitoring as in Example 1. Example 3 (Optimal Stopping):

The basic setup is the same as in Example 1. This time we first collect all the data up to finite horizon N. Then, we look at our data and try to find the *best* check-point t so the test result  $R_t$  is the most favorable. The difference between this example and continuous monitoring is that for the latter the decision of stopping the experiment is made without peeking at the data in *future*.

In both examples above, if we collected all the data up to horizon N and did the test, we should always report this test result instead of re-analysing the data or try to cherry pick the optimal stopping time. This is because Bayesian test is consistent: as we observe more data, posterior  $P(H_1|Data)$ converges to 1 if  $H_1$  is true and to 0 otherwise. This means we should always prefer the decision made from more data. However, it is possible that continuous monitoring might have rejected  $H_1$  (if it were used) but the finite horizon test at N does not. Does that mean continuous monitoring makes more error? No! Table I shows continuous monitoring does increase the amount of null rejection, without sacrificing the false discovery rate. In the above case the posterior odds realized in the end of the experiment at horizon N shows  $H_1$  is unlikely to be true. It is fair to say if we had been using continuous monitoring, we would likely be making a false discovery at that time, based on newer observations. However, let K' be the posterior odds reported by continuous monitoring, Theorem 1 guarantees that it is K' to 1 odds that we will see posterior odds increasing to  $\infty$  if we keep getting more data, than decreasing to 0. In other words, it is K'-1 more likely our decision still uphold in the end of the experiment, than reversed as in the hypothetical case.

Another critical assumption in Theorem 1 is that the likelihood ratio has to be correctly calculated with all available observations, *i.e.* the whole subpath  $\mathbf{X}_t$ . In particular, we cannot cherry pick only those observations that favors one hypothesis. Here is another bad example which happens a lot in practice.

Example 4 (Continuous Testing until Win):

With agile development and continuous A/B testing, a team can iteratively modify and test a feature until seeing a successful test result.

In NHST, even if the feature has no effect, there is still  $\alpha$ (typically 0.05) chance that the result could be statistically significant. This means for every 20 iterations, we might just declare a success without really having any true effect. This is like continuous monitoring, but the difference is that here

each new test only uses its own data. Using Bayesian test, if we need to calculate the likelihood ratio up to the t-th test, we have to aggregate all the evidence together, not just looking at the last one. If all tests are independent replications of the same test, aggregating evidence in Bayes test is trivial, we just need to multiply likelihood ratio for each of the replications all together. This way even if we might have a few large likelihood ratio favoring  $H_1$ , but if  $H_0$  is the ground truth there have to be more smaller likelihood ratios so the product is small. In fact it will converge to 0 if we keep doing replication runs. In practice, since iteration runs are not exactly replications, it is still a challenge how we should properly aggregate evidence from multiple experiment runs together. Ignoring the prior runs can still result in more false discovery than nominally controlled. Technically this is the area of multiple testing and selection bias. See Lu and Deng [19] for some preliminary results.

#### VI. BAYESIAN PROMISES WITH OPTIONAL STOPPING

In this section we continue our earlier simulation approach in Section III for Bayesian Promise (2) with optional stopping. We also study the implication of optional stopping on Type-I, Type-II error(power) and point estimation for the effect. In particular, we emphasize two Bayesian Non-Promises.

- 1) Bayesian test does not promise Type-I error control.
- 2) Bayesian optional stopping does not promise unbiased effect estimation using frequentist MLE.

#### A. Stopping Rule Based on Bayes Factor

If rejecting  $H_0$  when observing a posterior odds no less than K exposes us to a risk of false discovery at most 1/(1+K), a natural stopping rule is to prescribe a false discovery rate (FDR) bound and stop the test immediately if observed posterior odds already can guarantee the FDR control. We can set K=9 to guarantee a FDR bound of 10%=1/(9+1).

Similarly, we can early stop for futility and accept  $H_0$  if we believe posterior of null is sufficiently large. A symmetric design is to stop if posterior odds is either no less than K or no greater than 1/K.

Figure 2 illustrated both stopping rules under the same setup in fixed horizon case of Section III, except that we stopped at the first  $t \leq N$  when the above stopping rule is satisfied. In each of the 100,000 simulation runs, regardless of whether this run is early stopped or stopped in the end, we always calculate Bayes Factor based on Equation 5, replacing N by the observed stopping time  $\tau$ . Comparing Figure 2 to Figure 1 shows big differences. The biggest one being the spike at the stopping Bayes Factor boundary K = 9 (and 1/9for futility). However, the interesting observation is, despite the huge histogram shift for both  $H_1(\text{red})$  and  $H_0(\text{green})$ , those numbers on the top margin — ratios of observed Bayes Factors in each bin from  $H_1$  to  $H_0$ , remains very close to the theoretical Bayes Factor value calculated from Equation 5, as in a fixed horizon test. This is exactly what Theorem 1 claims, and this simulation study confirms it!

For many who are used to the frequentist thinking of controlling Type-I error, this result seems odd. If we allow

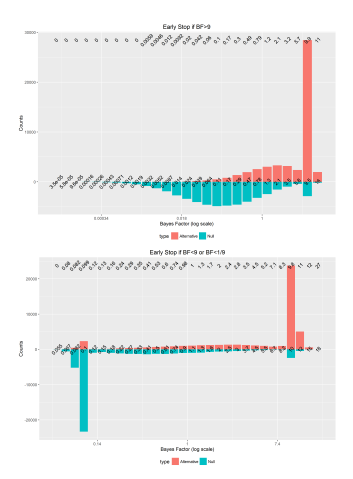


Fig. 2. Histograms of simulated Bayes Factor with optional stopping. Top plot: one-sided stopping. Bottom plot: two-sided stopping.

early stop, and still using the same rejection criteria of BF > 1/K, we will only reject more so we will be inflating the Type-I error. This is correct, but nonetheless does not conflict with the fact that FDR is still controlled below the designed level.

TABLE I IMPACT OF EARLY STOPPING

				Early Stop Rate	
	Type-I	Power	FDR	$H_1$	$H_0$
Fixed Horizon	0.018	0.465	0.037	NA	NA
One-sided Stop	0.060	0.599	0.09	59.5%	NA
Two-sided Stop	0.060	0.598	0.09	64.9%	65.0%

TABLE II POINT ESTIMATE AND STANDARD ERROR WITH EARLY STOPPING: GROUND TRUTH IS 0.2.

	Early Stop	No Early Stop	Unconditioned
Fixed Horizon	NA	NA	0.201(0.0004)
One-sided Stop	0.366(0.0009)	0.112(0.0004)	0.263(0.0007)
Two-sided Stop	0.319(0.001)	0.125(0.0004)	0.251(0.0009)

Table I shows the comparison of three simulation studies we did so far in terms of Type-I error, power and FDR. In the fixed horizon design, when we reject for BF>9, the Type-I error (proportion of false rejection among the null cases) is 0.018. This value increased to 0.06 when continuous monitoring/optional stopping is introduced. Because we are rejecting more, the power of the test is also improved from 0.465 to 0.599. FDR in the finite horizon cases is only 0.037, smaller than the designed bound of 0.1. This is because in the finite horizon cases a lot of rejected cases at the end of the test are actually bearing a BF much larger than the threshold 9, see Figure 1. This suggests that in finite horizon test, using a BF cutoff to calculate FDR might be conservative, also see Efron [10] for the differences of local FDR and FDR. When optional stopping is introduced, FDR become 0.09, very close to the designed level. The small discrepancies here is due to overshoot, i.e. we stop once BF is larger than 9 but not exactly at 9. These over-shoots are reflected in Figure 2 where we found a few bars beyond the spike. In large sample scenario where each individual observation won't make a big change in BF, as in most A/B tests, we can think of the time series of  $BF_t$ as continuous. In this case we can stop the test with a BF almost exactly equal to 9, and the FDR will be also almost exactly 0.1. We saw that FDR control in the fixed horizon setting is conservative because we are wasting sample sizes to collect evidence beyond what we really need, and with early stopping the waste is mitigated. The last two columns in Table I shows the percentage of the simulated experiment with early stopping. We saw majority of the simulated runs stopped earlier. We also calculated that the average length of the simulated runs with early stopping is about 55, much smaller than the fixed horizon of N = 100. Based on Table I, one could argue that early stopping is always superior than the fixed horizon test, and should be recommended if the sole purpose of the study is for hypothesis testing.

Table II shows the sample mean, which is the frequentest MLE for the effect in this one sample test scenario. Those are average of sample means taken from those 50,000 runs simulated under  $H_1$ . For fixed horizon test, it is centered around the true effect 0.2. However, when we use optional stopping, we get biased estimation when averaged across all of the 50,000 runs (numbers in parentheses are standard error of the average MLE). We further separate the cases where early stopping happened or not. We can see that in both one-sided and two-sided stopping, when early stopping happened, the average MLE are much higher than the true effect. When early stopping didn't happen, the average is lower. This reflect a common criticism of Bayesian optional stopping that it produces biased effect estimation, especially when early stopping is triggered. This is actually a misunderstanding of the Bayesian Promise. To be clear, our Theorem 1 didn't make any promises regarding effect estimation. We assume the alternative model, i.e. distribution of the effect under alternative is known and how to objectively learn this distribution is another orthogonal task we will mention later in Section VII.<sup>8</sup> Also, if we were to make some promises about effect estimation, under Bayesian framework we should be looking at posterior mean instead of frequentist MLE. Just like we never make promises for Type-I error, a lot of misunderstanding of Bayesian promises is due to mixing Bayesian test with frequentist methods [12]. Note that even using frequentist NHST, conditioned on a result that  $H_0$  is rejected, the MLE will still over-estimate the true effect due to post selection bias.

We do not want to underplay the importance of effect estimation. An extended discussion of Bayesian effect size estimation with optional stopping is beyond the scope of this paper and recently has been nicely discussed in Schönbrodt [23]. If the purpose of the study is for effect estimation, then a more proper stopping rule should be based on accuracy requirement, e.g. stop when required posterior standard deviation is achieved.

#### B. General Stopping Rule

Theorem 1 holds for general stopping rules, not only those based on BF cutoff values. For experimenters who want to "hack" p-values, they could choose to stop once p-value is less than  $\alpha$ . Here we did the simulation study with the stopping rule with both criteria: 1) p-value less than 0.1, and 2) the sample sizes is at least 10.

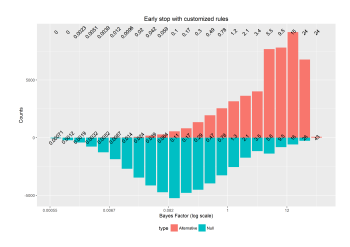


Fig. 3. Using a stopping rule based on p-value and minimum required sample size

Figure 3 shows the simulation results, this time with a rather bizarre histogram for  $H_1$  runs. The important part is the observed actual ratio on the top margin still closely tracks the theoretical Bayes Factor values on the x-axis.

#### C. Composite Alternative

So far in this section we have been using a overly simple alternative model  $H_1$  where the treatment effect is assumed to be known. This is not very realistic since we never know the effect so that alternative is always a composite alternative where  $\delta$  can be anything nonzero. In Bayesian model comparison we need to put a prior distribution for  $\delta$  under  $H_1$ , in

addition to the prior odds. Following [13] and [8], we put a normal prior  $N(0,\sigma_0^2)$ ). Under this  $H_1, X_i \sim N(0,\sigma_0^2+1)$  and the formula for Bayes Factor assuming a fixed sample size N changes to

$$\frac{N(\overline{X};0,\sigma_0^2+1/N)}{N(\overline{X};0,1/N)} \tag{10}$$

A similar simulation to those above in this section is run by setting N=1,000. We also set  $\sigma_0=0.1$  to generate 50,000 independent  $\delta$  first for each of the simulation runs from  $H_1$ . At the end of each runs(or at the stopping time) we compute Bayes Factor based on (10) with N for fixed horizon setting or  $\tau$  in its place when optional stopping is introduced. Figure 4 shows the results for both fixed horizon setting and optional stopping with BF cutoff at 9. In the fixed horizon setting, the histogram is much more dispersed than the previous precise alternative case. Some BF is as large as several thousands so we only show those no greater than 100. Early stopping effectively eliminated those extremely large BF, creating spikes around 9. We hope readers at this point already noticed that the top margin numbers are very close to the theoretical BF values on the x-axis.

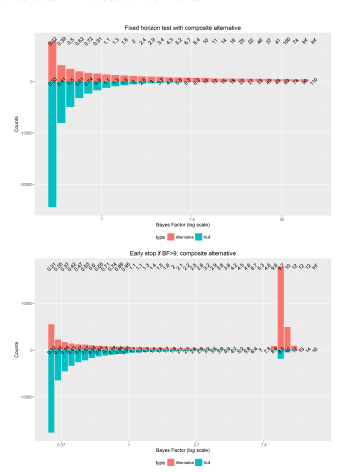


Fig. 4. Composite  $H_1$  with Normal prior. Top: fixed horizon(x-axis limit to 100). Bottom: early stopping when BF<sub>i</sub>.9.

 $<sup>^{8}</sup>$ In this particular simulation study we fix the alternative to be 0.2 under  $H_{1}$ , a known quantity so estimate the effect makes no sense at all.

#### VII. OBJECTIVE PRIOR LEARNING

Theorem 1 and all the discussion of this paper so far assumes two things are known:

- 1) model under  $H_1$ , i.e. the effect distribution under  $H_1$ ,
- 2) prior probability  $P(H_1)$  and  $P(H_0)$ , or equivalently the prior odds  $P(H_1)/P(H_0)$ .

With this crucial assumption we argued that the Bayesian Promises in Equations (2) and (3) do not change when optional stopping is used. However, in reality we almost always don't know both the model under  $H_1$  and the prior probabilities. Some people rely on subjective belief, and of course are hard to be accepted by scientific community. A lot of practical application of Bayesian Model use "non-informative" prior. These are less subjective but ironically still contains strong prior information. For A/B testing platform at scale (tens or hundreds of experiments per month), rich historical data exist and prior can be learned objectively from the empirical data by making the assumption that the current is like the past using hierarchical model [8]. This idea is similar to [14] which applied empirical Bayes techniques to signal processing.

Hierarchical model prior learning involves two steps. First, two sample t-test are transformed into a one sample problem with proper rescale so that the prior are defined in scaleless effect size. Then we use a hierarchical model where both the model under  $H_1$  and prior probabilities are unknown parameters. Fitting the hierarchical model can be done using either MLE (Empirical Bayes) as in [8] or Hierarchical Bayes (Full Bayes).

Suppose observations for treatment and control groups are i.i.d. from two distributions with unknown mean  $\tau_T$  and  $\tau_C$  respectively. Denote our observations by  $Y_i, i=1,\ldots,N_T$  and  $X_i, i=1,\ldots,N_C$ . We test the null hypothesis  $H_0:\tau_T-\tau_C=0$  against the alternative  $H_1:\tau_T\neq\tau_C$ . Without assuming distributions of X and Y, we use the central limit theorem and hence use Wald test which is large sample version of the well-known t-test. The test statistic is

$$\mathbf{Z} := \frac{\overline{X} - \overline{Y}}{\sqrt{\sigma_T^2/N_T + \sigma_C^2/N_C}} = \frac{\Delta}{\sqrt{\sigma_T^2/N_T + \sigma_C^2/N_C}},$$

where  $\sigma_C$  and  $\sigma_T$  are variances of X and Y and  $\Delta$  is the observed metric difference between treatment and control. The variances are also unknown but in large sample scenario we assume they are known and use their estimates. We define  $N_E = 1/(1/N_T + 1/N_C)$  to be the **effective sample size**. And then let  $\sigma^2$  be the **pooled variance** such that  $\sigma^2/N_E = \sigma_T^2/N_T + \sigma_C^2/N_C$ . With the shorthand  $\delta = \Delta/\sigma$ , Z-statistics can be rewritten as

$$Z = \frac{\delta}{\sqrt{1/N_E}}. (11)$$

 $\delta$  is  $\Delta$  scaled by pooled standard deviation and is called the **effect size**. Finally, define

$$\mu := E(\delta) = E(\Delta)/\sigma = (\tau_T - \tau_C)/\sigma \tag{12}$$

is the average treatment effect scaled by  $\sigma$ . When  $\sigma$  is treated as known, inference on  $\tau_T - \tau_C$  and  $\mu$  are equivalent. In

Bayesian analysis it is common to define prior for  $\mu$  as it is scaleless.

Recall  $\mu$  is the average effect size. Under  $H_0$ ,  $\mu=0$ . Under  $H_1$ , we assume a prior  $\pi$  for  $\mu$ . For both cases we observe  $\delta \sim N(\mu, 1/N_E)$ . In addition, we assume a prior probability p for  $H_1$ , and also under  $H_1$ ,  $\pi \sim N(0, V^2)$  for some V. Our challenge is to learn both p and V without the need of subjectively assigning one. Note that the procedure works for different  $\pi$ .

Next we take advantages of historical experiment results and use them to learn the prior parameters p and V. Suppose for a given metric, we have N previously conducted tests with observed effect size and effective sample size  $(\delta_i, N_{Ei}), i = 1, \ldots, N$ . If we know which one of these are from  $H_1$  and  $H_0$ , learning p and V are straightforward. Because those labels are latent, Expectation-Maximization[7] is typically applied for MLE. [8] describes a detailed EM algorithm.

For a full Bayesian inference, we put yet another layer of prior for p and V. Inferences can be done by using MCMC or approximation methods, e.g. variational Bayes. We provide a Stan[4] model in Appendix. Note that in the Stan Model we do not supply prior for p or V and Stan will use uniform prior (in a transformed unconstrained space). The specification of this hyper prior is less critical when the number of historical experiments are at least hundreds, in which case MLE and Bayesian posterior mean are also very close.

In our opinion, learning priors objectively is critical for any study which hopes to use optional stopping and relies on Theorem 1. Once the prior learning has finished, it should be fixed during the experiment stage when continuous monitoring and optional stopping are used.

# VIII. CONCLUSION & RECOMMENDATION

We hope the debate over whether continuous monitoring is a valid practice for Bayesian Hypothesis Testing is settled by Theorem 1 in this paper. The answer is yes and the Bayesian Promises (2) and (3) in fixed horizon case remains *unchanged* when a *proper* stopping rule is used. We emphasize that the correct understanding of (4) and interpretation of Bayesian test result as controlling FDR is critical and we should not mix Bayesian test results with frequentist concepts. Trying to evaluate Type-I error of Bayesian Test under either null or alternative is fallacious because the correct Bayesian interpretation always requires a prior odds weighing the alternative and the null. Our simulation illustrations in Section III and Section VI serve the very goal of helping readers understand what are the promises a Bayesian test tries to make and what not.

Two natural questions are raised by practitioners. 1) Because the fundamental differences in the statistical conclusions we can make from NHST and Bayesian test, which one shall we use in practice? 2) Is the result of this paper suggesting we should always use continuous monitoring for Bayesian tests?

The answer for the first question amounts to the choice between controlling Type-I error and FDR. If false rejection of any single test cost us a lot, and the cost of false rejection is considered higher than false negative, then Type-I error seems to be a better criterion to control, e.g. clinical trial. If our goal is not for each individual test, but our decisions' overall performance on a large set of tests, and the cost of false rejection and false negative are in the same order, then we believe FDR is a better criterion. Large scale A/B testing platform is an example of the latter [16]. In an agile environment where success is built on many small gains, as long as we are shipping more good features than useless ones, we are moving in the right direction.

For the second question, continuous monitoring is not always recommended. In many cases, the goal of the experiment is not only to confirm the existence of the treatment effect, but also to measure it. In A/B tests, it is not uncommon for a feature to have time-varying treatment effect such as weekday and weekend effect. To capture the weekly cycle, running tests for a whole week or multiple of weeks are often necessary. It is also possible that the treatment effect only exists in the weekend and we might early stop the experiment during the first few weekdays result in false negative. Also, conditioned on early stopping happened or not, Bayesian posterior can have a bias for effect estimation [23]. Continuous monitoring should be recommended in many other scenarios. Shutdown a bad experiment is one application in which we want to stop an experiment once we have enough evidence that the treatment is giving user a very bad experience. Another example is comparing a few closely related alternative candidates, e.g. tuning parameters for a backend algorithm, in which case we might assume the ordering of the treatment effects won't be time-varying and hence we can early stop inferior candidates and allocate traffic to outperforming candidates based on Bayesian posterior. The last example is studied in more detail in the literature of multi-armed bandit and Thompson sampling [5] and the result of this paper justifies Thompson sampling for using Bayesian posterior to dynamically change traffic allocation.

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### APPENDIX

# STAN MODEL