Bayesian approaches in clinical trials

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23/04/2023

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1 Introduction

1.1 General notation and abbreviations

- iid: Independent and identically distributed.
- pdf: Probability density function. Most often denoted as $f(\cdot)$. For bivariate pdf we use the notation $f_2(\cdot)$.
- cdf: Cumulative distribution function. Most often denoted as $F(\cdot)$. For bivariate pdf we use the notation $F_2(\cdot)$.
- N_2 : Bivariate cdf of the Gaussian distribution.
- ϕ : pdf of the standard Gaussian distribution.
- Φ : cdf of the standard Gaussian distribution.
- Φ^{-1} : Quantile function of the standard Gaussian distribution function.

1.2 'Power' vocabulary

In their supplement Kunzmann et al. [1] provide a literature review of the terminology used in articles. We provide here a summary of this terminology:

- Frequentist power: Probability of rejection given that the alternative hypothesis is true.
- Average power: Prior averaged probability of rejection. Often also called 'probability of success', 'assurance', 'Bayesian predictive power'.
- **Prior adjusted power**: Joint probability of rejection and that the treatment effect is effective.

1.3 Some 'Bayesian' concepts

- **Prior**: A random variable Θ with pdf $f(\theta)$ representing that the uncertainty of a parameter of interest.
- **Design prior**: Prior used before data collection as data generating mode [2].
- Analysis prior: Prior used for Bayesian analysis of the collected data [2].
- Prior predictive distribution: Situation before a sample was taken. Let θ be a realisation of a random variable Θ with pdf $f(\theta)$. Then for a future observation \tilde{X} the pdf is

$$f(\tilde{x}) = \int_{\Theta} f(\tilde{x}, \theta) d\theta = \int_{\Theta} \underbrace{f(\tilde{x}|\theta)}_{likelihood} \underbrace{f(\theta)}_{prior} d\theta.$$

• Posterior predictive distribution: Situation after a sample was taken. Let θ be a realisation of a random variable Θ with pdf $f(\theta)$. Then for a future observation \tilde{X} and observed X (since X is independent \tilde{X}) the pdf is

$$f(\tilde{x}|x) = \int_{\Theta} f(\tilde{x}|\theta, x) f(\theta|x) d\theta = \int_{\Theta} \underbrace{f(\tilde{x}|\theta)}_{likelihood} \underbrace{f(\theta|x)}_{prior} d\theta.$$

- Improper prior: A prior with $\int_{\Theta} f(\theta) = \infty$.
- **Jeffrey's prior**: For an unknown parameter θ Jeffrey's (scalar) prior is defined as $f(\theta) \propto \sqrt{I(\theta)}$, where $I(\theta)$ is the expected Fisher information of θ [3]. Jeffrey's prior can be improper [3]. Bayesian point estimates using Jeffrey's prior are often very close to maximum likelihood estimators [3].

Example: Jeffrey's prior for the binomial model

The likelihood of the binomial model is

$$f(x|\theta) = \binom{n}{x} \theta^x (1-\theta)^{n-x}$$

and thus

$$L := log(f(x|\theta)) = x \log(\theta) + (n-x) \log(1-\theta).$$

Simple algebra leads to

$$\frac{dL}{d\theta} = \frac{x}{\theta} - \frac{n-x}{1-\theta}, \quad \frac{d^2L}{d\theta^2} = -\frac{x}{\theta^2} - \frac{n-x}{(1-\theta)^2}.$$

The expected Fisher information is

$$I(\theta) = -E_{\theta}\left(\frac{d^2L}{d\theta^2}\right) = \frac{n\theta}{\theta^2} + \frac{n-n\theta}{(1-\theta)^2} = \frac{n}{\theta(1-\theta)} \propto \frac{1}{\theta(1-\theta)}.$$

Thus, Jeffrey's prior for the binomial model is

$$f(\theta) \propto \frac{1}{\sqrt{\theta(1-\theta)}} = \theta^{-0.5}(1-\theta)^{-0.5} = beta(0.5, 0.5).$$

Jeffrey's prior for the binomial model is a proper prior [3].

1.4 Used R libraries

- # All tidyverse functions
- library(tidyverse)
- # For frequentist sample size calculation
- library(epiR)
- # Multivariate Gaussian and t-distributions
- library(mvtnorm)
- # Beta binomial distribution
- library(extraDistr)
- # Agresti-Caffo confidence intervals for risk difference
- library(PropCIs)

2 Power and sample size calculations

2.1 Background

The terminology of 'power' is often imprecisely used [1]. Kunzmann et al. suggest to use the neutral term 'probability to reject'. The classical (frequentist) 'power' is defined as the probability to reject given that the alternative hypothesis is true. Frequentist power calculations do not include uncertainties of the treatment effect, whereas Bayesian and hybrid approaches include such uncertainties in their calculations. In the following we use different approaches for the calculation of the 'probability to reject' (frequentist, Bayesian and hybrid) for different clinical trial designs.

Definition 'hybrid' ([4], Section 6.5.2)

'[...] we have a prior distribution to use in our study design, but that the conclusions of the study will be entirely classical and will not make use of the prior [...]'

2.2 Two-arm non-inferiority setting

In this section we consider a non-inferiority clinical trial setting with a null hypothesis $H_0: \delta > \delta^*$ and alternative hypothesis $H_a: \delta \leq \delta^*$, where $\delta^* > 0$ is a fixed non-inferiority margin and a treatment effect δ , for example, a continuous difference or a risk difference.



Treatment effect (<--- favors active treatment)

2.2.1 Binomial outcome

Here p_1 and p_0 are event probabilities from an active treatment arm and a control arm, respectively. $\delta = p_1 - p_0$ is the true treatment effect expressed as a risk difference.

In this section we assume that the variances in both groups are known, but might be different.

Working example

We use the SAFE-SSPE trial as a working example [5]. In brief, this non-inferiority randomised placebocontrolled trial compares clinical surveillance versus anticoagulant treatment in low-risk patients with isolated subsegmental pulmonary embolism (SSPE). The primary outcome is the proportion of 3-month recurrence of venous thromboembolism (VTE).

The null hypothesis H_0 is 'clinical surveillance is inferior to anticoagulant treatment' versus the alternative hypothesis H_a 'clinical surveillance is non-inferior to anticoagulant treatment'. Thus, $H_0: p_1 - p_0 > \delta^*$

vs $H_a: p_1 - p_0 \le \delta^*$, where p_1 is the VTE proportion in the clinical surveillance arm and p_0 is the VTE proportion in the control arm.

The non-inferiority margin was set at 3.5% and it was assumed that the proportion of VTE in both groups was 1%.

2.2.1.1 Frequentist approach

Let $Y_i = (Y_{i,1}, Y_{i,2}, \cdots, Y_{i,n_i})^{\top}$ be a sample of size n_i from a binomial distribution $Y_i \sim Binomial(p_i, n_i)$, $i \in \{0,1\}$, where p_i , $i \in \{0,1\}$, are the true event proportions. Denote the estimated event proportions as $\overline{p}_i = \frac{1}{n_i} \sum_{k \leq n_i} Y_{i,k}, \ i \in 0,1$, and the estimated risk difference as $D = \overline{p}_1 - \overline{p}_0$. Then D is asymptotically Gaussian distributed $D \sim N\left(\delta, \frac{\sigma_1^2}{n_1} + \frac{\sigma_0^2}{n_0}\right)$, where $\sigma_i^2 = p_i(1-p_i), \ i \in \{0,1\}$. For notational purposes we denote the variance of the treatment effect as $\sigma_{treat}^2 = \frac{\sigma_1^2}{n_1} + \frac{\sigma_0^2}{n_0}$.

We are interested whether the upper $(1 - \alpha)\%$ -confidence limit of D is smaller than the non-inferiority margin δ^* , that is,

$$D+z_{1-\alpha}\sqrt{\frac{n_0\sigma_1^2+n_1\sigma_0^2}{n_1n_0}}\leq \delta^*\quad \Rightarrow\quad D\leq -z_{1-\alpha}\sqrt{\frac{n_0\sigma_1^2+n_1\sigma_0^2}{n_1n_0}}+\delta^*,$$

where $z_{1-\alpha}=\Phi^{-1}(1-\alpha)$. Note that $D^{\delta^*}_{suc}:=-z_{1-\alpha}\sqrt{\frac{n_0\sigma_1^2+n_1\sigma_0^2}{n_1n_0}}+\delta^*$ is the **required risk difference** for a 'successful' rejection of the null hypothesis. Then

$$\begin{split} P_{\delta}(D \leq D_{suc}^{\delta^*}) &= \Phi\left(-z_{1-\alpha} - \sqrt{\frac{n_1 n_0}{n_0 \sigma_1^2 + n_1 \sigma_0^2}} (\delta - \delta^*)\right) \\ &= \Phi\left(-z_{1-\alpha} - \frac{(\delta - \delta^*)}{\sigma_{treat}}\right), \end{split}$$

since under regularity conditions,

$$Z = \frac{D - \delta^*}{\sigma_{treat}} \to N(0, 1), \quad \min(n_1, n_0) \to \infty.$$

The conditional probability $P_{\delta}(D \leq D_{suc}^{\delta^*})$ is the **probability to reject given** δ . For $\delta > \delta^*$ this is the 'type-I error' and for $\delta \leq \delta^*$ the frequentist 'power'.

For a δ_A it holds that

$$-z_{1-\alpha} - \sqrt{\frac{n_1 n_0}{n_0 \sigma_1^2 + n_1 \sigma_0^2}} (\delta_A - \delta^*) = \Phi^{-1}(1-\beta) = z_{1-\beta}$$

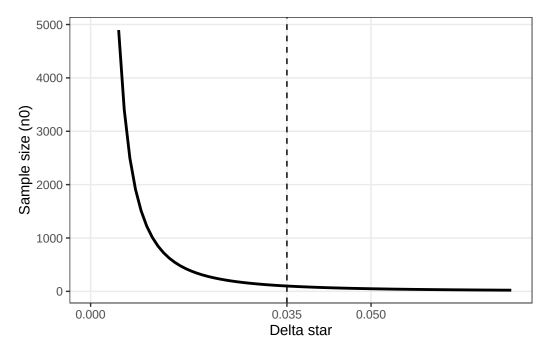
and so the sample size can then be calculated as

$$\frac{(z_{1-\beta}+z_{1-\alpha})^2}{(\delta_A-\delta^*)^2} = \frac{an_0^2}{n_0\sigma_1^2+an_0\sigma_0^2},$$

where $a = n_1/n_0$ is an allocation ratio, such that

$$n_0 = (z_{1-\beta} + z_{1-\alpha})^2 \frac{\sigma_1^2 + a\sigma_0^2}{a(\delta_A - \delta^*)^2}, \quad n_1 = an_0.$$

 δ_0 , δ_A and δ^* are assumed as fixed and known constants in a frequentist approach. Their choices are of high importance, because all trial conclusions are based on those choices and affect the sample size calculation. The plot below shows how the sample size increase as δ^* approaches δ .



Working example (continued)

\$delta

We calculate the required sample size for the SAFE-SSPE trial using a frequentist approach with the following parameters:

```
• p_1 = 0.01, p_0 = 0.01, \delta^* = 0.035, 1 - \beta = 0.8, \alpha = 0.05, a = 1/1
  library(epiR)
  alpha <- 0.05
  beta <- 0.2
  p_0 < 0.01
  p_1 < 0.01
  delta <- p_1 - p_0
  delta_star <- 0.035
  sd_0 \leftarrow sqrt(p_0 * (1 - p_0))
  sd_1 \leftarrow sqrt(p_1 * (1 - p_1))
  a < -1/1
  epi.ssninfb(treat = p_1, control = p_0, delta = delta_star, power = 1 -
       beta, r = a, alpha = alpha, n = NA)
$n.total
[1] 200
$n.treat
[1] 100
$n.control
[1] 100
```

[1] 0.035

\$power

[1] 0.8

[1] 99.93031

```
n_1 <- n_0 * a
n_1
```

[1] 99.93031

Under the specified parameters a sample size of 200 patients (100 per arm) is needed to reject the null hypothesis of inferiority. This is more or less the sample size mentioned in the study protocol of the SAFE-SSPE trial but without dropouts and adjustments for rare events.

2.2.1.2 Hybrid approach: Prior on the risk difference

Suppose that the true treatment effect δ is a realization from a random variable Δ with $f(\delta)$. In this subsection we assume that the design prior comes from a Gaussian distribution function so that $\Delta \sim N\left(d,\frac{\sigma_1^2+\sigma_0^2}{m}\right)$. Note that this prior can be thought as a realisation from m Gaussian 'prior observations' with variance $\sigma_1^2+\sigma_0^2$. Again for notational purposes we denote the variance of the design prior of the treatment effect as $\sigma_{prior}^2=\frac{\sigma_1^2+\sigma_0^2}{m}$.

In the following we will use the following design priors:

- Enthusiastic prior (favors non-inferiority): d = 0, m = 6.6, $P(\Delta > \delta^*) = 0.05$. This prior is centered on the treatment effect such that there is a low probability (here 5%) of inferiority.
- Skeptical prior (favors inferiority): $d = \delta^*$, m = 6.6, $P(\Delta > 0) = 0.05$. This prior is centered on the non-inferiority margin such that there is a low probability (here 5%) of superiority.
- Informative prior (clinical expert knowledge): d = 0 with m = 25.
- Noninformative prior: d = 0 with m = 0.5.



Let

$$AP:=\int_{\Delta}P_{\delta}(D\leq D_{suc}^{\delta^*})f(\delta)d\delta,$$

be the 'average power' [6] (also called 'assurance' [7], 'probability of success' [1], [4] or Bayesian predictive power [8]. [1] provide in their supplemental section a literature review of the used terminology in publications from clinical trials).

Remember that in an hybrid approach we are interested in trial conclusions from a frequentist point of view, thus we are interested in

$$D \leq -z_{1-\alpha} \sqrt{\frac{n_0 \sigma_1^2 + n_1 \sigma_0^2}{n_1 n_0}} + \delta^*.$$

By using a design prior we take into account the uncertainty of the treatment effect. The prior pre-

dictive distribution for an estimated risk difference, say \tilde{D} , with a prior $\Delta \sim N\left(d, \sigma_{prior}^2\right)$ includes this uncertainty. For the Gaussian case, the prior predictive distribution of \tilde{D} is given as

$$\tilde{D} \sim N\left(d, \sigma_{treat}^2 + \sigma_{prior}^2\right),$$

since $\tilde{D} \sim N\left(\delta, \sigma_{treat}^2\right)$, see for example [9].

Suppose now that D has a predictive distribution as described above, then

$$AP = \int_{-\infty}^{-z_{1-\alpha}\sqrt{\frac{n_0\sigma_1^2+n_1\sigma_0^2}{n_1n_0}}+\delta^*} f(\tilde{\delta})d\tilde{\delta} = \Phi\left(\frac{1}{\sigma_{prior}}\left[-z_{1-\alpha}\sigma_{treat}-(d-\delta^*)\right]\right),$$

see for example [6]. Note that as $m \to \infty$, then $AP \to \Phi\left(-z_{1-\alpha} - \frac{(d-\delta^*)}{\sigma_{treat}}\right)$, that is, the frequentist power at d.

Working example (continued)

We calculate the AP under the assumed prior distributions and parameters for the SAFE-SSPE trial.

```
m < -6.6
prior_mean <- delta_star</pre>
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
sigma_treat \leftarrow sqrt((sd_0^2/n_0 + sd_1^2/n_1))
AP <- pnorm(1/sigma_prior * (-qnorm(1 - alpha) * sigma_treat -
    (prior_mean - delta_star)))
data_output <- rbind(data_output, data.frame(type = "Skeptical",</pre>
    n_0, n_1, AP = round(AP, 2))
# Informative prior
m <- 25
prior_mean <- 0</pre>
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
sigma_treat \leftarrow sqrt((sd_0^2/n_0 + sd_1^2/n_1))
AP <- pnorm(1/sigma_prior * (-qnorm(1 - alpha) * sigma_treat -
    (prior_mean - delta_star)))
data_output <- rbind(data_output, data.frame(type = "Informative",</pre>
    n_0, n_1, AP = round(AP, 2))
# Noninformative prior
m < -0.5
prior_mean <- 0</pre>
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
sigma_treat \leftarrow sqrt((sd_0^2/n_0 + sd_1^2/n_1))
AP <- pnorm(1/sigma_prior * (-qnorm(1 - alpha) * sigma_treat -
```

```
(prior_mean - delta_star)))

data_output <- rbind(data_output, data.frame(type = "Noninformative",
    n_0, n_1, AP = round(AP, 2)))

data.frame(data_output %>%
    arrange(type))
```

type n_0 n_1 AP

- 1 Enthusiastic 99.93031 99.93031 0.59
- 2 Informative 99.93031 99.93031 0.66
- 3 Noninformative 99.93031 99.93031 0.52
- 4 Skeptical 99.93031 99.93031 0.34

Under an 'enthusiastic prior' we get an average power of 59%. For a 'skeptical prior' the average power decreases to 34%. These values are lower than the frequentist power of 80%, which holds under the assumption that the specified treatment effect (the alternative hypothesis) is true.

Rufibach et al. give a closed a formula for the distribution of $P_{\Delta}(D \leq D_{suc}^{\delta^*})$, where $\Delta \sim N(d, \sigma_{prior}^2)$, and discuss the shape under different prior choices [8]. Kunzmann et al. call this the 'random probability to reject' (RPR), $RPR := P_{\Delta}(D \leq D_{suc}^{\delta^*})$.

For 0 < y < 1, the random variable RPR has a pdf

$$f(y) = \frac{\sigma_{treat}}{\sigma_{prior}} \phi \left(-z_{1-\alpha} \frac{\sigma_{treat}}{\sigma_{prior}} - \frac{(d-\delta^*)}{\sigma_{prior}} - \frac{\sigma_{treat}}{\sigma_{prior}} \Phi^{-1}(y) \right) \left[\phi \left(\Phi^{-1}(y) \right) \right]^{-1},$$

see [8].

Special case

For the case that $n_0 = n_1 = n$ and $\sigma_1^2 = \sigma_0^2 = \sigma^2$, then $\sigma_{treat}^2 = \frac{2\sigma^2}{n}$ and $\sigma_{prior}^2 = \frac{2\sigma^2}{m}$, and the formula above reduces to

$$f(y) = \sqrt{\frac{m}{n}} \phi \left(\sqrt{\frac{m}{n}} \left[-z_{1-\alpha} - \sqrt{\frac{n}{2\sigma^2}} (d - \delta^*) - \Phi^{-1}(y) \right] \right) \left[\phi \left(\Phi^{-1}(y) \right) \right]^{-1}, \quad 0 < y < 1.$$

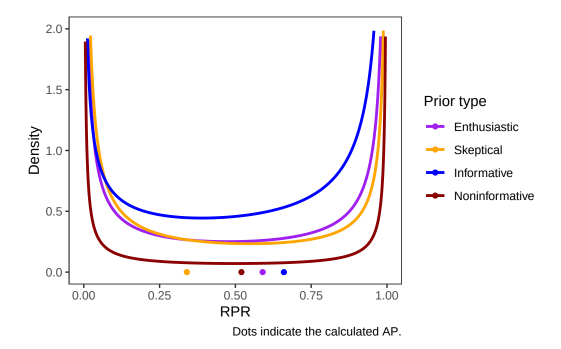
Working example (continued)

We derive the probability densities for the different assumed design priors for the SAFE-SSPE study.

```
# Enthusiastic prior
m <- 6.6
prior_mean <- 0
sigma_prior <- sqrt((sd_0^2 + sd_1^2))/sqrt(m)
sigma_treat <- sqrt((sd_0^2/n_0 + sd_1^2/n_1))

## Rufibach 2016: formula (4)
y <- sqrt(m * n_0 * sd_1^2 + m * n_1 * sd_0^2)/(sqrt(n_1 * n_0 * (sd_0^2 + sd_1^2))) * dnorm(-sqrt(m * n_0 * sd_1^2 + m * n_1 * sd_0^2)/(sqrt(n_1 + n_0 * (sd_0^2 + sd_1^2))) * qnorm(1 - alpha) - sqrt(m)/sqrt((sd_0^2 + sd_1^2)) * (prior_mean - delta_star) - sqrt(m * n_0 * sd_1^2 + m * n_1 * sd_0^2)/(sqrt(n_1 * n_0 * (sd_0^2 + sd_1^2))) * (sqrt(n_1 *
```

```
# Skeptical prior
m < -6.6
prior_mean <- delta_star</pre>
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
sigma_treat \leftarrow sqrt((sd_0^2/n_0 + sd_1^2/n_1))
y <- sigma_treat/sigma_prior * dnorm(-qnorm(1 - alpha) * sigma_treat/sigma_prior -
    (1/sigma_prior) * (prior_mean - delta_star) - (sigma_treat/sigma_prior) *
    qnorm(x)) * (dnorm(qnorm(x)))^(-1)
data_power <- rbind(data_power, data.frame(x, y, type = "Skeptical"))</pre>
# Informative prior
m < -25
prior_mean <- 0</pre>
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
sigma_treat \leftarrow sqrt((sd_0^2/n_0 + sd_1^2/n_1))
y <- sigma_treat/sigma_prior * dnorm(-qnorm(1 - alpha) * sigma_treat/sigma_prior -
    (1/sigma_prior) * (prior_mean - delta_star) - (sigma_treat/sigma_prior) *
    qnorm(x)) * (dnorm(qnorm(x)))^(-1)
data_power <- rbind(data_power, data.frame(x, y, type = "Informative"))</pre>
# Noninformative prior
m < -0.5
prior_mean <- 0</pre>
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
sigma_treat \leftarrow sqrt((sd_0^2/n_0 + sd_1^2/n_1))
```



The cdf of RPR is

$$P(RPR \leq y) = 1 - \Phi\left(\frac{\sigma_{treat}}{\sigma_{prior}} \left[-z_{1-\alpha} - \frac{(d-\delta^*)}{\sigma_{treat}} - \Phi^{-1}(y) \right] \right), \quad 0 < y < 1.$$

Special case

For the case that $n_0 = n_1 = n$ and $\sigma_1^2 = \sigma_0^2 = \sigma^2$, then $\sigma_{treat}^2 = \frac{2\sigma^2}{n}$ and $\sigma_{prior}^2 = \frac{2\sigma^2}{m}$, and the formula above reduces to

$$P(RPR \leq y) = 1 - \Phi\left(\frac{\sigma_{treat}}{\sigma_{prior}} \left[-z_{1-\alpha} - \frac{(d-\delta^*)}{\sigma_{treat}} - \Phi^{-1}(y) \right] \right), \quad 0 < y < 1.$$

Note that if n_0 and n_1 are the planned treatment arm sample sizes from a frequentist power calculation as described in Section 2.1.1 above, then, if $d = \delta_A$,

$$-z_{1-\alpha} - \frac{(d-\delta^*)}{\sigma_{treat}} = \Phi^{-1}(y) = \Phi^{-1}(1-\beta) \quad \Rightarrow \quad P(RPR \leq y) = 0.5, \quad y = 1-\beta.$$

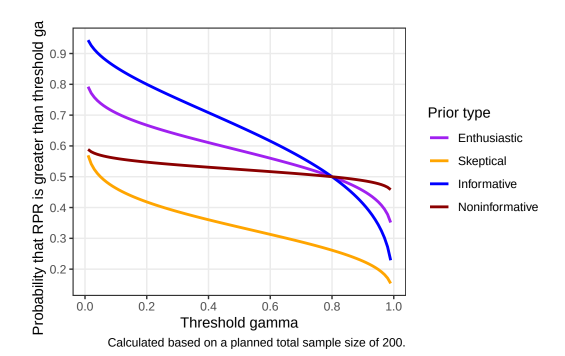
Working example (continued)

With the formula above we can calculate the probability $P(RPR > \gamma)$, $0 < \gamma < 1$, for the SAFE-SSPE study and the above specified design priors.

```
# gamma range
gamma \leftarrow seq(0.01, 0.99, 0.01)
# Centered on p1-p0
prior_mean <- 0</pre>
m_range <- c(0.5, 6.6, 25)
n_0_range <- n_0
data_rpr <- expand.grid(gamma, m_range, n_0_range)</pre>
names(data rpr) <- c("gamma", "m range", "n 0 range")</pre>
data_rpr$n_1_range <- a * data_rpr$n_0_range</pre>
data_rpr$sigma_treat <- sqrt(sd_0^2/data_rpr$n_0_range + sd_1^2/data_rpr$n_1_range)
data rpr$sigma prior <- sqrt(sd 0^2/data rpr$m range + sd 1^2/data rpr$m range)
data_rpr$p_rpr <- pnorm(-qnorm(1 - alpha) * data_rpr$sigma_treat/data_rpr$sigma_prior -
    (1/data_rpr$sigma_prior) * (prior_mean - delta_star) -
      (data_rpr$sigma_treat/data_rpr$sigma_prior) *
    qnorm(data_rpr$gamma))
data rpr$type <- NA
data_rpr$type[data_rpr$m_range == 0.5] <- "Noninformative"</pre>
```

```
data_rpr$type[data_rpr$m_range == 6.6] <- "Enthusiastic"</pre>
data rpr$type[data rpr$m range == 25] <- "Informative"</pre>
# Centered on noninferiority margin
prior_mean <- delta_star</pre>
m_range <- c(6.6)</pre>
n_0_range <- n_0
data_rpr_skep <- expand.grid(gamma, m_range, n_0_range)</pre>
names(data_rpr_skep) <- c("gamma", "m_range", "n_0_range")</pre>
data_rpr_skep$n_1_range <- a * data_rpr_skep$n_0_range</pre>
data_rpr_skep$sigma_treat <- sqrt(sd_0^2/data_rpr_skep$n_0_range +
    sd_1^2/data_rpr_skep$n_1_range)
data_rpr_skep$sigma_prior <- sqrt(sd_0^2/data_rpr_skep$m_range +</pre>
    sd_1^2/data_rpr_skep$m_range)
data_rpr_skep$p_rpr <- pnorm(-qnorm(1 - alpha) *</pre>
                                 data_rpr_skep$sigma_treat/data_rpr_skep$sigma_prior -
    (1/data_rpr_skep$sigma_prior) * (prior_mean - delta_star) -
    (data rpr skep$sigma treat/data rpr skep$sigma prior) * qnorm(data rpr skep$gamma))
data_rpr_skep$type <- "Skeptical"</pre>
data_rpr <- rbind(data_rpr, data_rpr_skep)</pre>
data rpr$type <- factor(data rpr$type, levels = c("Enthusiastic",
    "Skeptical", "Informative", "Noninformative"))
ggplot(data_rpr, aes(x = gamma, y = p_rpr, colour = type)) +
    geom line(linewidth = 1) + theme bw() + theme(panel.grid.minor = element blank()) +
    scale_colour_manual("Prior type", values = c("purple", "orange",
```

```
"blue", "darkred")) + scale_y_continuous(breaks = seq(0,
1, 0.1)) + scale_x_continuous(breaks = seq(0, 1, 0.2)) +
labs(caption = str_glue("Calculated based on a planned total sample size of ",
    round(n_0_range + a * n_0_range, 0), ".")) + xlab("Threshold gamma") +
ylab("Probability that RPR is greater than threshold gamma")
```



For a specific threshold, say $\gamma = 0.6$, and varying 'prior sample sizes' m we obtain

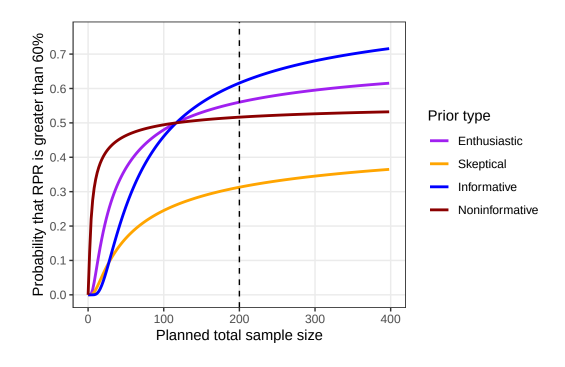
```
gamma <- 0.6

# Centered on p1-p0
prior_mean <- 0

m_range <- c(0.5, 6.6, 25)
n_0_range <- seq(0.01, 300, by = 1)

data_rpr <- expand.grid(m_range, n_0_range)</pre>
```

```
names(data_rpr) <- c("m_range", "n_0_range")</pre>
data_rpr$n_1_range <- a * data_rpr$n_0_range</pre>
data_rpr$sigma_treat <- sqrt(sd_0^2/data_rpr$n_0_range + sd_1^2/data_rpr$n_1_range)
data_rpr$sigma_prior <- sqrt(sd_0^2/data_rpr$m_range + sd_1^2/data_rpr$m_range)</pre>
data rpr$p rpr <- pnorm(data rpr$sigma treat/data rpr$sigma prior *</pre>
    (-qnorm(1 - alpha) - (1/data_rpr$sigma_treat) * (prior_mean -
        delta_star) - qnorm(gamma)))
data_rpr$type <- NA
data_rpr$type[data_rpr$m_range == 0.5] <- "Noninformative"</pre>
data_rpr$type[data_rpr$m_range == 6.6] <- "Enthusiastic"</pre>
data_rpr$type[data_rpr$m_range == 25] <- "Informative"</pre>
# Centered on noninferiority margin
prior_mean <- delta_star</pre>
m_range <- c(6.6)
n_0_{range} \leftarrow seq(0.01, 300, by = 1)
data_rpr_skep <- expand.grid(m_range, n_0_range)</pre>
names(data_rpr_skep) <- c("m_range", "n_0_range")</pre>
data_rpr_skep$n_1_range <- a * data_rpr_skep$n_0_range</pre>
data_rpr_skep$sigma_treat <- sqrt(sd_0^2/data_rpr_skep$n_0_range +
    sd_1^2/data_rpr_skep$n_1_range)
data_rpr_skep$sigma_prior <- sqrt(sd_0^2/data_rpr_skep$m_range +</pre>
    sd_1^2/data_rpr_skep$m_range)
data rpr skep$p rpr <- pnorm(data rpr skep$sigma treat/data rpr skep$sigma prior *
    (-qnorm(1 - alpha) - (1/data_rpr_skep$sigma_treat) * (prior_mean -
```



Note that the average power AP integrates over the whole Δ range. This might include also 'non-favorable' regions. To see that one can decompose AP as follows (see [1], [6]):

$$AP = \overbrace{P(D \leq D_{suc}^{\delta^*}, \Delta > \delta^*)}^{(1)} + \overbrace{P(D \leq D_{suc}^{\delta^*}, 0 < \Delta \leq \delta^*)}^{(2)} + \overbrace{P(D \leq D_{suc}^{\delta^*}, \Delta \leq 0)}^{(3)}$$

where

- (1) Probability of Type-I error,
- (2) 'Non-inferior region, but treatment effect not relevant',
- (3) 'Non-inferior region, treatment effect relevant'.

Spiegelhalter et al. highlight that $AP \approx P(D \leq D_{suc}^{\delta^*}, \Delta \leq 0)$ because the type-I error is often small and one has strong believe for the alternative hypothesis in designing a clinical trial [4]. [1] and [7] discuss the practical relevance of the AP decomposition. For example, pharmaceutical companies might favour (1)+(2)+(3) taking into account shortterm risk, wheras regulators are interested in (3) or (2)+(3), that is non-inferior outcomes (2)+(3) with relevant treatment effects (3).

In the non-inferiority setting we are interested in (2)+(3), that is $\Delta \leq \delta^*$:

$$P(D \leq D_{suc}^{\delta^*}, \Delta \leq \delta^*) = P(D \leq D_{suc}^{\delta^*} | \Delta \leq \delta^*) \\ P(\Delta \leq \delta^*) = \underbrace{E\left[P_{\Delta \leq \delta^*}(D \leq D_{suc}^{\delta^*})\right]}_{EP} P(\Delta \leq \delta^*),$$

Kunzmann et al. denote EP the 'expected power' [1]. Note that

$$\underbrace{P(D \leq D_{suc}^{\delta^*}, \Delta \leq \delta^*)}_{PAP} = \underbrace{E\left[P_{\Delta \leq \delta^*}(D \leq D_{suc}^{\delta^*})\right]}_{EP} \underbrace{P(\Delta \leq \delta^*)}_{constant}.$$

Spiegelhalter calls $P(D \leq D_{suc}^{\delta^*}, \Delta \leq \delta^*)$ the 'prior adjusted power' (PAP).

The AP decomposition can be visualised using the posterior predictive distribution. Let $f(\tilde{\delta})$ be the pdf of the posterior predictive distribution then joint distribution of $\tilde{\delta}$ and δ is

$$f_2(\tilde{\delta}, \delta) = N_2 \left(\begin{pmatrix} d \\ d \end{pmatrix}, \begin{pmatrix} \sigma_{treat}^2 + \sigma_{prior}^2 & \sigma_{prior}^2 \\ \sigma_{prior}^2 & \sigma_{prior}^2 \end{pmatrix} \right),$$

see formula (2.11) in [6].

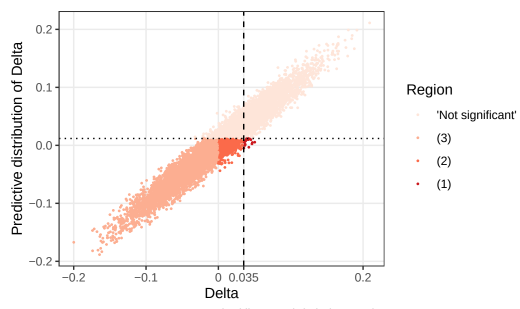
Working example (continued)

For the SAFE-SSPE study the joint distribution $f_2(\tilde{\delta}, \delta)$ under an enthusiastic prior can be plotted as:

```
library(mvtnorm)
library(tidyverse)
library(ggplot2)

delta_star <- 0.035
prior_mean <- 0
m <- 6.6</pre>
sigma_sim <- matrix(c((sd_0^2/n_0 + sd_1^2/n_1 + (sd_1^2 + sd_1^2/n_1))))
```

```
sd_0^2/m, (sd_1^2 + sd_0^2)/m, (sd_1^2 + sd_0^2)/m,
    (sd 1^2 + sd 0^2)/m), nrow = 2, ncol = 2, byrow = T)
set.seed(1)
data_sim <- data.frame(rmvnorm(n = 10000, mean = c(prior_mean,</pre>
    prior_mean), sigma_sim))
names(data_sim) <- c("delta_pred", "delta")</pre>
data_sim$region <- ifelse(data_sim$delta <= 0 & data_sim$delta_pred <=
    -qnorm(1 - alpha) * sqrt(sd_1^2/n_1 + sd_0^2/n_0) -
        (prior_mean - delta_star), 1, 0)
data sim$region <- ifelse(data sim$delta > 0 & data sim$delta <=
    delta_star & data_sim$delta_pred <= -qnorm(1 -</pre>
    alpha) * sqrt(sd_1^2/n_1 + sd_0^2/n_0) - (prior_mean -
    delta_star), 2, data_sim$region)
data_sim$region <- ifelse(data_sim$delta > delta_star &
    data_sim$delta_pred <= -qnorm(1 - alpha) * sqrt(sd_1^2/n_1 +</pre>
        sd_0^2/n_0 - (prior_mean - delta_star), 3,
    data_sim$region)
data_sim$region <- factor(data_sim$region, levels = 0:3,</pre>
    labels = c("'Not significant'", "(3)", "(2)", "(1)"))
ggplot(data_sim, aes(x = delta, y = delta_pred, colour = factor(region))) +
    geom_point(size = 0.3) + theme_bw() + theme(panel.grid.minor = element_blank()) +
    geom_vline(xintercept = 0.035, linetype = "dashed") +
    geom_hline(yintercept = -qnorm(1 - alpha) * sqrt(sd_1^2/n_1 +
        sd_0^2/n_0) - (prior_mean - delta_star), linetype = "dotted") +
    ylab("Predictive distribution of Delta") + xlab("Delta") +
    scale_color_brewer("Region", palette = "Reds") +
```



Dashed line: Noninferiority margin. Dotted line: Upper CI from predicted delta <= non-inferiority margin.

```
data_output <- data.frame(data_sim %>%
    group_by(region) %>%
    summarise(prop = n()/nrow(data_sim)))
data_output
```

```
region prop
1 'Not significant' 0.4148
2 (3) 0.4909
3 (2) 0.0920
4 (1) 0.0023
```

From the above values the average power can be calculated as (1) 0.0023 + (2) 0.092 + (3) 0.4909, which is equal to 0.5852. This is the value we reported above (59%).

The results of EP, PAP and AP are shown in the table below:

```
delta star <- 0.035
prior_mean <- 0</pre>
m < -6.6
set.seed(1)
sd_prior \leftarrow sqrt((sd_1^2 + sd_0^2))/sqrt(m)
draws <- rnorm(10000, mean = prior_mean, sd = sd_prior)</pre>
power_classic <- pnorm(-qnorm(1 - alpha) - sqrt(n_0 * n_1)/sqrt((n_0 *</pre>
    sd_1^2 + n_1 * sd_0^2) * (draws - delta_star)
data_ep <- data.frame(ep = mean(power_classic[draws <= delta_star]),</pre>
    pap = mean(power_classic[draws <= delta_star]) * pnorm(delta_star,</pre>
        mean = prior_mean, sd = sd_prior), ap = mean(power_classic),
    const = pnorm(delta_star, mean = prior_mean, sd = sd_prior),
    type = "Skeptical")
set.seed(1)
prior_mean <- delta_star</pre>
m < -6.6
draws <- rnorm(10000, mean = prior_mean, sd = sd_prior)</pre>
power_classic <- pnorm(-qnorm(1 - alpha) - sqrt(n_0 * n_1)/sqrt((n_0 *</pre>
    sd_1^2 + n_1 * sd_0^2) * (draws - delta_star)
data_ep <- rbind(data_ep, data.frame(ep = mean(power_classic[draws <=</pre>
    delta_star]), pap = mean(power_classic[draws <= delta_star]) *</pre>
```

```
pnorm(delta_star, mean = prior_mean, sd = sd_prior), ap = mean(power_classic),
           const = pnorm(delta_star, mean = prior_mean, sd = sd_prior),
           type = "Enthusiastic"))
prior_mean <- 0</pre>
m < -25
set.seed(1)
draws <- rnorm(10000, mean = prior_mean, sd = sd_prior)</pre>
power\_classic <- pnorm(-qnorm(1 - alpha) - sqrt(n_0 * n_1)/sqrt((n_0 * alpha) - sqrt(n_0 * n_1)/sqrt((n_0 * alpha) - sqrt(n_0 * alpha) - sqrt(n_
            sd_1^2 + n_1 * sd_0^2) * (draws - delta_star)
data_ep <- rbind(data_ep, data.frame(ep = mean(power_classic[draws <=</pre>
           delta_star]), pap = mean(power_classic[draws <= delta_star]) *</pre>
           pnorm(delta star, mean = prior mean, sd = sd prior), ap = mean(power_classic),
           const = pnorm(delta_star, mean = prior_mean, sd = sd_prior),
           type = "Informative"))
prior_mean <- 0</pre>
m < -0.5
set.seed(1)
draws <- rnorm(1e+05, mean = prior_mean, sd = sd_tilde/sqrt(0.5))</pre>
power_classic <- pnorm(-qnorm(1 - alpha) - sqrt(n_0 * n_1)/sqrt((n_0 *</pre>
           sd_1^2 + n_1 * sd_0^2) * (draws - delta_star)
data_ep <- rbind(data_ep, data.frame(ep = mean(power_classic[draws <=</pre>
           delta_star]), pap = mean(power_classic[draws <= delta_star]) *</pre>
           pnorm(delta_star, mean = prior_mean, sd = sd_prior), ap = mean(power_classic),
```

```
const = pnorm(delta_star, mean = prior_mean, sd = sd_prior),
       type = "Noninformative"))
  data_ep$ep <- round(data_ep$ep, 4)</pre>
  data_ep$ap <- round(data_ep$ap, 4)</pre>
  data_ep$pap <- round(data_ep$pap, 4)</pre>
  data_ep$const <- round(data_ep$const, 4)</pre>
  data_ep
      ер
                         const
             pap
                                           type
1 0.7930 0.5857 0.5872 0.7386
                                      Skeptical
2 0.6740 0.3370 0.3437 0.5000
                                  Enthusiastic
3 0.7930 0.5857 0.5872 0.7386
                                    Informative
```

2.2.1.3 Bayesian approach: Prior on the risk difference

4 0.9186 0.6785 0.5233 0.7386 Noninformative

In this section we use a proper Bayesian approach for the calculation of the probability to reject the null hypothesis $H_0: \delta > \delta^*$, where $\delta = p_1 - p_0$ [4] [10]. Suppose that $\Delta \sim N(d, \sigma_{prior}^2)$ is an analysis prior for the treatment effect δ . Remember that $D = \overline{p}_1 - \overline{p}_0$ is the estimated (observed) treatment effect with $D \sim N(\delta, \sigma_{treat}^2)$. A 'trial success' can then be defined as

$$P(\Delta < \delta^*|D) > 1 - \epsilon$$
,

where ϵ is small, say $\epsilon = 0.05$ [4] [10]. Using the above specified prior and likelihood of the data, the posterior distribution is given as

$$\Delta|D \sim N\left(\frac{\frac{d}{\sigma_{prior}^2} + \frac{D}{\sigma_{treat}^2}}{\frac{1}{\sigma_{prior}^2} + \frac{1}{\sigma_{treat}^2}}, \frac{1}{\frac{1}{\sigma_{prior}^2} + \frac{1}{\sigma_{treat}^2}}\right) = N\left(\frac{\sigma_{treat}^2 d + \sigma_{prior}^2 D}{\sigma_{treat}^2 + \sigma_{prior}^2}, \frac{\sigma_{treat}^2 \sigma_{prior}^2}{\sigma_{treat}^2 + \sigma_{prior}^2}\right),$$

see for example [9].

Special case

If $n_0=n_1=n$ and $\sigma_1^2=\sigma_0^2=\sigma^2$, then $\sigma_{treat}^2=\frac{2\sigma^2}{n}$ and $\sigma_{prior}^2=\frac{2\sigma^2}{m}$, and the formula above reduces to

$$\Delta | D \sim N\left(\frac{m\delta + nD}{n+m}, \frac{2\sigma^2}{n+m}\right).$$

The trial is successful if

$$\frac{\sigma_{treat}^2 d + \sigma_{prior}^2 D}{\sigma_{treat}^2 + \sigma_{prior}^2} + z_{1-\epsilon} \frac{\sigma_{treat} \sigma_{prior}}{\sqrt{\sigma_{treat}^2 + \sigma_{prior}^2}} \leq \delta^*,$$

where the left side of the above equation is the if the upper $(1 - \epsilon)$ -credible interval. A simple algebraic step gives

$$D \leq -z_{1-\epsilon} \frac{\sigma_{treat}}{\sigma_{prior}} \sqrt{\sigma_{treat}^2 + \sigma_{prior}^2} + \delta^* \left(1 + \frac{\sigma_{treat}^2}{\sigma_{prior}^2} \right) - \frac{\sigma_{treat}^2}{\sigma_{prior}^2} d.$$

Here, $D_{suc}^{d,\delta^*} = -z_{1-\epsilon} \frac{\sigma_{treat}}{\sigma_{prior}} \sqrt{\sigma_{treat}^2 + \sigma_{prior}^2} + \delta^* \left(1 + \frac{\sigma_{treat}^2}{\sigma_{prior}^2}\right) - \frac{\sigma_{treat}^2}{\sigma_{prior}^2} d$ is the **required risk difference** for a successful rejection of the null hypothesis taking into account the uncertainty of δ .

Since $D \sim N(\delta, \sigma_{treat}^2)$,

$$P(D \leq D_{suc}^{d,\delta^*} | \delta) = \Phi\left(-z_{1-\epsilon}\sqrt{1 + \frac{\sigma_{treat}^2}{\sigma_{prior}^2}} - \frac{1}{\sigma_{treat}}\left[\delta - \delta^*\left\{1 + \frac{\sigma_{treat}^2}{\sigma_{prior}^2}\right\}\right] - \frac{\sigma_{treat}}{\sigma_{prior}^2}d\right).$$

This is the **conditional Bayesian probability to reject** (or Bayesian power) under an (assumed) known true treatment effect δ .

Special case

If $n_0 = n_1 = n$ and $\sigma_1^2 = \sigma_0^2 = \sigma^2$, then $\sigma_{treat}^2 = \frac{2\sigma^2}{n}$ and $\sigma_{prior}^2 = \frac{2\sigma^2}{m}$, and the formula above reduces to

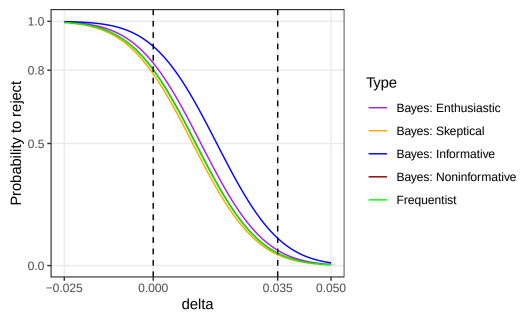
$$P(D \leq D_{suc}^{d,\delta^*}|\delta) = \Phi\left(-z_{1-\epsilon}\sqrt{\frac{n+m}{n}} - \frac{m}{\sqrt{2n\sigma^2}}d - \sqrt{\frac{n}{2n\sigma^2}}\left\{\delta - \delta^*\left(\frac{n+m}{n}\right)\right\}\right).$$

Working example (continued)

We derive the conditional Bayesian probability to reject for the SAFE-SSPE study for different δ values and analysis priors.

```
(sd_1^2 + sd_0^2)) * prior_mean - sqrt((n_1 * n_0)/((n_0 *
    sd_1^2 + n_1 * sd_0^2)) * (delta - delta_star * (1 + (m * 
    (n_0 * sd_1^2 + n_1 * sd_0^2))/(n_1 * n_0 * (sd_1^2 + sd_0^2)))))
data_power <- data.frame(delta, y = ap_bayes, type = "Bayes: Enthusiastic")</pre>
# Skeptical prior
prior_mean <- 0.035</pre>
m < -6.6
ap_bayes <- pnorm(-qnorm(1 - epsilon) * (sqrt(1 + m * (n_0 *
    sd_1^2 + n_1 * sd_0^2/(n_1 * n_0 * (sd_1^2 + sd_0^2)))) -
    (m * sqrt(n_0 * sd_1^2 + n_1 * sd_0^2))/(sqrt(n_1 * n_0) *
        (sd_1^2 + sd_0^2)) * prior_mean - sqrt((n_1 * n_0)/((n_0 *
    sd_1^2 + n_1 * sd_0^2)) * (delta - delta_star * (1 + (m * 
    (n_0 * sd_1^2 + n_1 * sd_0^2))/(n_1 * n_0 * (sd_1^2 + sd_0^2)))))
data_power <- rbind(data_power, data.frame(delta, y = ap_bayes,</pre>
    type = "Bayes: Skeptical"))
# Informative prior
prior_mean <- 0</pre>
m < -25
ap_bayes <- pnorm(-qnorm(1 - epsilon) * (sqrt(1 + m * (n_0 * m_0)))
    sd_1^2 + n_1 * sd_0^2/(n_1 * n_0 * (sd_1^2 + sd_0^2)))) -
    (m * sqrt(n_0 * sd_1^2 + n_1 * sd_0^2))/(sqrt(n_1 * n_0) *
        (sd_1^2 + sd_0^2) * prior_mean - sqrt((n_1 * n_0)/((n_0 *
    sd_1^2 + n_1 * sd_0^2)) * (delta - delta_star * (1 + (m * 
    (n_0 * sd_1^2 + n_1 * sd_0^2))/(n_1 * n_0 * (sd_1^2 + sd_0^2)))))
```

```
data_power <- rbind(data_power, data.frame(delta, y = ap_bayes,</pre>
    type = "Bayes: Informative"))
# Noninformative prior
prior_mean <- 0</pre>
m < -0.5
ap_bayes <- pnorm(-qnorm(1 - epsilon) * (sqrt(1 + m * (n_0 *
    sd_1^2 + n_1 * sd_0^2/(n_1 * n_0 * (sd_1^2 + sd_0^2)))) -
    (m * sqrt(n_0 * sd_1^2 + n_1 * sd_0^2))/(sqrt(n_1 * n_0) *
        (sd_1^2 + sd_0^2) * prior_mean - sqrt((n_1 * n_0)/((n_0 *
    sd_1^2 + n_1 * sd_0^2)) * (delta - delta_star * (1 + (m * 
    (n_0 * sd_1^2 + n_1 * sd_0^2))/(n_1 * n_0 * (sd_1^2 + sd_0^2)))))
data_power <- rbind(data_power, data.frame(delta, y = ap_bayes,</pre>
    type = "Bayes: Noninformative"))
# Frequentist
y_freq <- pnorm(-(delta - delta_star) * sqrt(n_1 * n_0)/sqrt(n_0 *</pre>
    sd_1^2 + n_1 * sd_0^2 - qnorm(1 - alpha)
data_power <- rbind(data_power, data.frame(delta, y = y_freq,</pre>
    type = "Frequentist"))
data_power$type <- factor(data_power$type, levels = c("Bayes: Enthusiastic",
    "Bayes: Skeptical", "Bayes: Informative", "Bayes: Noninformative",
    "Frequentist"))
ggplot(data_power, aes(x = delta, y = y, colour = type)) + geom_line() +
```



For treatment arm sample sizes: n0=100, n1=100

```
data_power$y <- round(data_power$y, 2)
data_power %>%
    filter(delta %in% c("0", "0.035"))
```

delta y type
1 0.000 0.83 Bayes: Enthusiastic
2 0.035 0.06 Bayes: Enthusiastic

0.000 0.78 Bayes: Skeptical 3 0.035 0.04 Bayes: Skeptical 0.000 0.90 Bayes: Informative 5 6 0.035 0.11 Bayes: Informative 0.000 0.80 Bayes: Noninformative 0.035 0.05 Bayes: Noninformative 0.000 0.80 Frequentist 10 0.035 0.05 Frequentist

The Bayesian probability to reject $H_0: \delta > \delta^*$ given $\delta = p_1 - p_0 = 0$ and an informative prior is 90%. This is higher than the frequentist probability to reject of 80% given $\delta = p_1 - p_0 = 0$. In contrast, the probability to reject $H_0: \delta > \delta^*$ for $\delta = 0.035$ under an informative prior is 11%. This is higher than the frequentist 'type-I error' of 5%. The probability to reject under the null hypothesis is higher for the Bayesian approach under an informative prior because the prior has substantial believe in non-inferiority.

Suppose \tilde{D} is calculated from a future sample given we have already observed D. The posterior predictive distribution of \tilde{D} is Gaussian distributed $\tilde{D} \sim N(d, \sigma_{treat}^2 + \sigma_{prior}^2)$ under an analysis prior $\Delta \sim N(d, \sigma_{prior}^2)$. The **Bayesian average (marginal) probability to reject** (BAP) can be calcu-

lated as

$$\begin{split} BAP &= \int_{-\infty}^{-\infty} P(D \leq D_{suc}^{d,\delta^*} | \delta) f(\delta) d\delta = \\ &= \int_{-\infty}^{\infty} \left\{ \int_{-\infty}^{-z_{1-\epsilon} \frac{\sigma_{treat}}{\sigma_{prior}} \sqrt{\sigma_{treat}^2 + \sigma_{prior}^2} + \delta^* \left(1 + \frac{\sigma_{treat}^2}{\sigma_{prior}^2}\right) - \frac{\sigma_{treat}^2}{\sigma_{prior}^2} d}{\sigma_{prior}^2} \int_{-\sigma_{treat}^2}^{\sigma_{treat}^2} d f(\delta) d\delta \right\} f(\delta) d\delta \\ &= \int_{-\infty}^{-z_{1-\epsilon} \frac{\sigma_{treat}}{\sigma_{prior}} \sqrt{\sigma_{treat}^2 + \sigma_{prior}^2}} + \delta^* \left(1 + \frac{\sigma_{treat}^2}{\sigma_{prior}^2}\right) - \frac{\sigma_{treat}^2}{\sigma_{prior}^2} d}{\sigma_{prior}^2} \left\{ \int_{-\infty}^{-\infty} f(\delta) \delta f(\delta) d\delta \right\} d\delta \\ &= \int_{-\infty}^{-z_{1-\epsilon} \frac{\sigma_{treat}}{\sigma_{prior}} \sqrt{\sigma_{treat}^2 + \sigma_{prior}^2}} + \delta^* \left(1 + \frac{\sigma_{treat}^2}{\sigma_{prior}^2}\right) - \frac{\sigma_{treat}^2}{\sigma_{prior}^2} d}{\sigma_{prior}^2} d\delta d\delta \\ &= \Phi \left(-z_{1-\epsilon} \frac{\sigma_{treat}}{\sigma_{prior}} \sqrt{\sigma_{treat}^2 + \sigma_{prior}^2} + \frac{\delta^*}{\sqrt{\sigma_{treat}^2 + \sigma_{prior}^2}} + \frac{\delta^*}{\sqrt{\sigma_{treat}^2 + \sigma_{prior}^2}} \left\{ 1 + \frac{\sigma_{treat}^2}{\sigma_{prior}^2} \right\} \\ &- \frac{d}{\sqrt{\sigma_{treat}^2 + \sigma_{prior}^2}} \left\{ 1 + \frac{\sigma_{treat}^2}{\sigma_{prior}^2} \right\} \right) \\ &= \Phi \left(-z_{1-\epsilon} \frac{\sigma_{treat}}{\sigma_{prior}} - \frac{\sqrt{\sigma_{treat}^2 + \sigma_{prior}^2}}{\sigma_{prior}^2} (d - \delta^*) \right), \end{split}$$

since.

Special case

If $n_0 = n_1 = n$ and $\sigma_1^2 = \sigma_0^2 = \sigma^2$, then $\sigma_{treat}^2 = \frac{2\sigma^2}{n}$ and $\sigma_{prior}^2 = \frac{2\sigma^2}{m}$, and the formula above reduces to

$$BAP = -z_{1-\epsilon}\sqrt{\frac{m}{n}} - \sqrt{\frac{(m+n)m}{2n\sigma^2}}(d-\delta^*)$$

Working example (continued)

We derive the Bayesian average probability to reject for the SAFE-SSPE study.

```
### Hybrid AP
sigma_treat \leftarrow sqrt((sd_0^2/n_0 + sd_1^2/n_1))
digit_round <- 4</pre>
# Enthusiastic prior
m < -6.6
prior_mean <- 0</pre>
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
AP <- pnorm(1/sigma_prior * (-qnorm(1 - alpha) * sigma_treat -
    (prior_mean - delta_star)))
data_output <- data.frame(type = "Enthusiastic", n_0, n_1, AP = round(AP,</pre>
    digit_round))
# Skeptical prior
m < -6.6
prior_mean <- delta_star</pre>
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
AP <- pnorm(1/sigma_prior * (-qnorm(1 - alpha) * sigma_treat -
    (prior_mean - delta_star)))
data_output <- rbind(data_output, data.frame(type = "Skeptical",</pre>
    n_0, n_1, AP = round(AP, digit_round)))
# Informative prior
m < -25
prior_mean <- 0</pre>
```

```
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
AP <- pnorm(1/sigma_prior * (-qnorm(1 - alpha) * sigma_treat -
    (prior_mean - delta_star)))
data_output <- rbind(data_output, data.frame(type = "Informative",</pre>
    n_0, n_1, AP = round(AP, digit_round)))
# Noninformative prior
m < -0.5
prior_mean <- 0</pre>
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
AP <- pnorm(1/sigma_prior * (-qnorm(1 - alpha) * sigma_treat -
    (prior_mean - delta_star)))
data_output <- rbind(data_output, data.frame(type = "Noninformative",</pre>
    n_0, n_1, AP = round(AP, digit_round)))
data_output <- data_output %>%
    select(type, n_0, n_1, AP)
### Bayesian AP
sigma_treat \leftarrow sqrt((sd_0^2/n_0 + sd_1^2/n_1))
# Enthusiastic prior
prior_mean <- 0</pre>
m < -6.6
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
```

```
ap_bayes <- pnorm(-qnorm(1 - alpha) * sigma_treat/sigma_prior -</pre>
    sqrt(sigma_treat^2 + sigma_prior^2)/sigma_prior^2 * (prior_mean -
        delta_star))
data_output2 <- data.frame(type = "Enthusiastic", n_0, n_1, AP_bayes = round(ap_bayes,</pre>
    digit_round))
# Skeptical prior
prior_mean <- delta_star</pre>
m < -6.6
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
ap_bayes <- pnorm(-qnorm(1 - alpha) * sigma_treat/sigma_prior -</pre>
    sqrt(sigma_treat^2 + sigma_prior^2)/sigma_prior^2 * (prior_mean -
        delta_star))
data_output2 <- rbind(data_output2, data.frame(type = "Skeptical",</pre>
    n_0, n_1, AP_bayes = round(ap_bayes, digit_round)))
# Informative prior
prior_mean <- 0</pre>
m < -25
sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
ap_bayes <- pnorm(-qnorm(1 - alpha) * sigma_treat/sigma_prior -</pre>
    sqrt(sigma_treat^2 + sigma_prior^2)/sigma_prior^2 * (prior_mean -
        delta_star))
data_output2 <- rbind(data_output2, data.frame(type = "Informative",</pre>
```

```
n_0, n_1, AP_bayes = round(ap_bayes, digit_round)))
  # Noninformative prior
  prior_mean <- 0</pre>
  m < -0.5
  sigma_prior \leftarrow sqrt((sd_0^2 + sd_1^2))/sqrt(m)
  ap_bayes <- pnorm(-qnorm(1 - alpha) * sigma_treat/sigma_prior -</pre>
      sqrt(sigma_treat^2 + sigma_prior^2)/sigma_prior^2 * (prior_mean -
          delta_star))
  data_output2 <- rbind(data_output2, data.frame(type = "Noninformative",</pre>
      n_0, n_1, AP_bayes = round(ap_bayes, digit_round)))
  data_output <- left_join(data_output, data_output2 %>%
      select(type, AP_bayes), by = "type")
  data_output %>%
      select(type, n_0, n_1, AP_hybrid = AP, AP_bayes)
                               n_1 AP_hybrid AP_bayes
            type
                      n_0
    Enthusiastic 99.93031 99.93031
                                       0.5856
                                                0.5937
2
       Skeptical 99.93031 99.93031
                                     0.3363
                                                0.3363
3
     Informative 99.93031 99.93031
                                       0.6631
                                                0.7149
4 Noninformative 99.93031 99.93031
                                       0.5237
                                                0.5239
```

The average Bayesian power under an informative prior is 72% compared to 66% from a hybrid approach.

2.2.1.4 Hybrid approach: Prior on event probabilities

In the former subsections we assumed that the prior was directly specified on the treatment effect δ . In this subsection design priors are specified on the event probabilities p_i , $i = \{0, 1\}$. Suppose that these design priors are beta distributed $\pi_i \sim Beta(a_i, b_i)$, $i = \{0, 1\}$. It is well known (see for example [9]) that the prior predictive distribution of the number of events R_i , $i = \{0, 1\}$, from a future binomial sample of size n_i , $i = \{0, 1\}$, is beta-binomial distributed $R_i \sim betabinom(r_i|n_i, a_i, b_i)$ with

$$E(R_i) = n_i \frac{a_i}{a_i + b_i}, \quad Var(R_i) = \frac{n_i a_i b_i (n_i + a_i + b_i)}{(a_i + b_i)^2 (a_i + b_i + 1)}.$$

Working example (continued)

For the SAFE-SSPE trial we specify the following design beta priors for the expected event probabilites $p_1 = p_0 = 0.01$.

Enthusiastic prior: The beta prior for the active treatment arm is centered at 0.01 (the expected event proportion) with $P(\pi_1 > 0.05) = 0.025$, that is, the probability that the prior is greater than the safety margin is 2.5%. The mean of the beta prior for the control arm is centered at 0.03 (the safety margin) with $P(\pi_0 > 0.05) = 0.05$.

- Parameters of beta prior for active treatment arm: $a_1 = 0.5, b_1 = 49.5$.
- Parameters of beta prior for control treatment arm: $a_0=7.2,\,b_0=232.8.$

```
library(extraDistr)

x <- seq(0.001, 0.999, 0.001)

## Enthusiastic prior

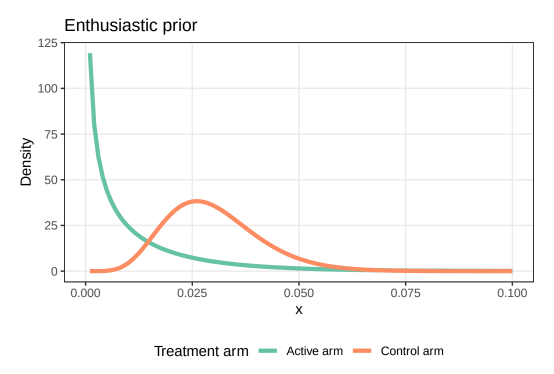
a_1 <- 0.5

b_1 <- (1 - p_1)/p_1 * a_1

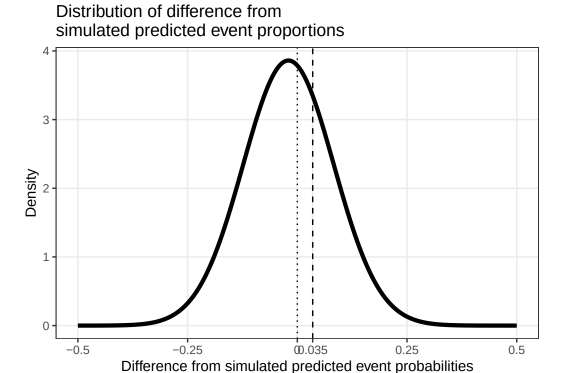
a_0 <- 7.2

b_0 <- (1 - 0.03)/0.03 * a_0
```

```
prior_0 <- dbeta(x, a_0, b_0)</pre>
prior_1 <- dbeta(x, a_1, b_1)</pre>
prior plot <- data.frame(x, prior 0, prior 1)</pre>
prior_plot_long <- pivot_longer(prior_plot, cols = c(prior_0,</pre>
    prior_1))
prior_plot_long$group <- ifelse(str_detect(prior_plot_long$name,</pre>
    "_0") == T, "Control arm", "Active arm")
fig1 <- ggplot(prior_plot_long, aes(x = x, y = value,
    colour = group)) + geom_line(linewidth = 1.5) +
    theme_bw() + theme(panel.grid.minor = element_blank(),
    legend.position = "bottom", legend.direction = "horizontal") +
    scale_x_continuous(limits = c(0, 0.1)) + ylab("Density") +
    labs(caption = str_glue("Assumed event probabilities: Control arm (p_0=",
        p_0, "), ", "Active arm (p_1=", p_1, ")")) +
    scale_color_brewer("Treatment arm", palette = "Set2") +
    scale_linetype("Type") + ggtitle("Enthusiastic prior")
diff_prior <- data.frame(x = rbbinom(1e+06, size = ceiling(n_1),
    a_1, b_1)/ceiling(n_1) - rbbinom(1e+06, size = ceiling(n_0),
    a_0, b_0)/ceiling(n_0))
fig2 <- ggplot(diff_prior, aes(x), group = x) + geom_density(linewidth = 1.5,
    bw = 0.1) + theme_bw() + theme(panel.grid.minor = element_blank()) +
    xlab("Difference from simulated predicted event probabilities") +
    ggtitle("Distribution of difference from\nsimulated predicted event proportions") +
    ylab("Density") + geom_vline(xintercept = delta_star,
    linetype = "dashed") + geom_vline(xintercept = 0,
```



Assumed event probabilities: Control arm (p_0=0.01), Active arm (p_1=0.01)



Skeptical prior: The beta prior for the active treatment arm is centered at 0.05 (the safety margin) with

 $P(\pi_1 \le 0.01) = 0.025$, that is the probability that the prior is smaller or equal than the expected event proportion is 2.5%. The mean of the beta prior for the control arm is is centered at 0.01 (the expected event proportion) with $P(\pi_0 > 0.05) = 0.025$.

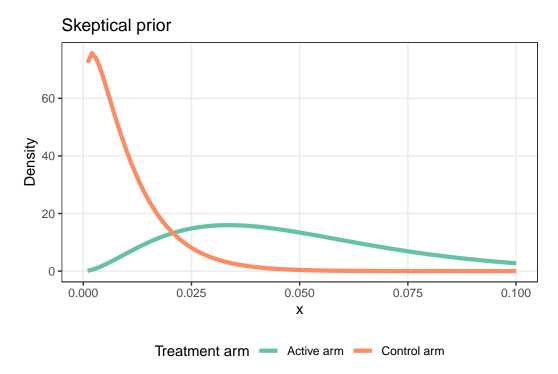
- Parameters of beta prior for active treatment arm: $a_1 = 2.84$, $b_1 = 53.96$.
- Parameters of beta prior for control treatment arm: $a_i = 1.24, b_i = 122.76$.

```
x \leftarrow seq(0.001, 0.999, 0.001)
# Skeptical prior
a 1 <- 2.84
b_1 \leftarrow (1 - 0.05)/0.05 * a_1
a_0 < 1.24
b \ 0 \ \leftarrow \ (1 - 0.01)/0.01 * a \ 0
prior_0 <- dbeta(x, a_0, b_0)</pre>
prior_1 <- dbeta(x, a_1, b_1)</pre>
prior_plot <- data.frame(x, prior_0, prior_1)</pre>
prior_plot_long <- pivot_longer(prior_plot, cols = c(prior_0,</pre>
    prior 1))
prior_plot_long$group <- ifelse(str_detect(prior_plot_long$name,</pre>
    "_0") == T, "Control arm", "Active arm")
fig1 <- ggplot(prior_plot_long, aes(x = x, y = value,
    colour = group)) + geom_line(linewidth = 1.5) +
    theme_bw() + theme(panel.grid.minor = element_blank(),
    legend.position = "bottom", legend.direction = "horizontal") +
    scale_x_continuous(limits = c(0, 0.1)) + ylab("Density") +
    labs(caption = str_glue("Assumed event probabilities: Control arm (p_0=",
        p_0, "), ", "Active arm (p_1=", p_1, ")")) +
```

```
scale_color_brewer("Treatment arm", palette = "Set2") +
    scale_linetype("Type") + ggtitle("Skeptical prior")

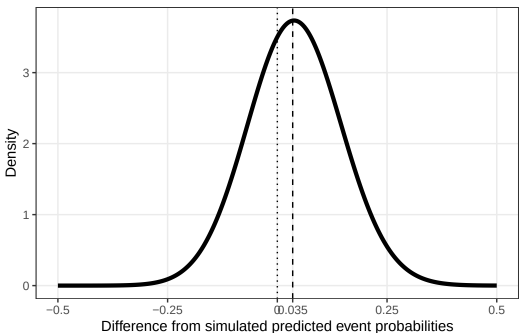
diff_prior <- data.frame(x = rbbinom(1e+06, size = ceiling(n_1),
    a_1, b_1)/ceiling(n_1) - rbbinom(1e+06, size = ceiling(n_0),
    a_0, b_0)/ceiling(n_0))

fig2 <- ggplot(diff_prior, aes(x), group = x) + geom_density(linewidth = 1.5,
    bw = 0.1) + theme_bw() + theme(panel.grid.minor = element_blank()) +
    xlab("Difference from simulated predicted event probabilities") +
    ggtitle("Distribution of difference from\nsimulated predicted event proportions") +
    ylab("Density") + geom_vline(xintercept = delta_star,
    linetype = "dashed") + geom_vline(xintercept = 0,
    linetype = "dotted") + scale_x_continuous(breaks = c(-0.5,
    -0.25, 0, 0.035, 0.25, 0.5), labels = c(-0.5, -0.25,
    0, 0.035, 0.25, 0.5), limits = c(-0.5, 0.5))</pre>
```



Assumed event probabilities: Control arm (p_0=0.01), Active arm (p_1=0.01)

Distribution of difference from simulated predicted event proportions



Informative prior: The beta prior for the active treatment arm and control arm is centered at 0.01 (the

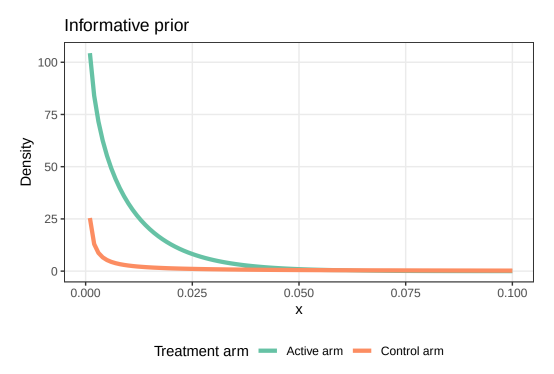
expected event proportion) with $P(\pi_1 \le 0.05) = 0.01$ and $P(\pi_1 \le 0.05) = 0.05$.

- Parameters of beta prior for active treatment arm: $a_1 = 0.8, b_1 = 79.2$.
- Parameters of beta prior for control treatment arm: $a_0 = 0.03, b_0 = 2.97.$

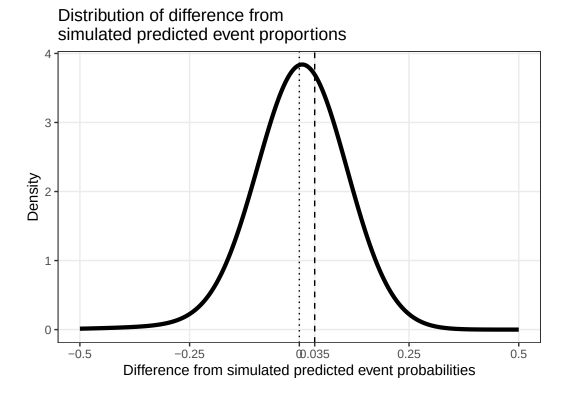
```
x \leftarrow seq(0.001, 0.999, 0.001)
## Informative prior
a_1 <- 0.8
b_1 \leftarrow (1 - p_1)/p_1 * a_1
a 0 < -0.03
b_0 < (1 - p_0)/p_0 * a_0
prior 0 <- dbeta(x, a 0, b 0)</pre>
prior_1 <- dbeta(x, a_1, b_1)</pre>
prior_plot <- data.frame(x, prior_0, prior_1)</pre>
prior_plot_long <- pivot_longer(prior_plot, cols = c(prior_0,</pre>
    prior_1))
prior_plot_long$group <- ifelse(str_detect(prior_plot_long$name,</pre>
    " 0") == T, "Control arm", "Active arm")
fig1 <- ggplot(prior_plot_long, aes(x = x, y = value,
    colour = group)) + geom_line(linewidth = 1.5) +
    theme_bw() + theme(panel.grid.minor = element_blank(),
    legend.position = "bottom", legend.direction = "horizontal") +
    scale_x = c(0, 0.1) + ylab("Density") +
    labs(caption = str_glue("Assumed event probabilities: Control arm (p_0=",
        p_0, "), ", "Active arm <math>(p_1=", p_1, ")")) +
    scale_color_brewer("Treatment arm", palette = "Set2") +
    scale linetype("Type") + ggtitle("Informative prior")
```

```
diff_prior <- data.frame(x = rbbinom(1e+06, size = ceiling(n_1),
    a_1, b_1)/ceiling(n_1) - rbbinom(1e+06, size = ceiling(n_0),
    a_0, b_0)/ceiling(n_0))

fig2 <- ggplot(diff_prior, aes(x), group = x) + geom_density(linewidth = 1.5,
    bw = 0.1) + theme_bw() + theme(panel.grid.minor = element_blank()) +
    xlab("Difference from simulated predicted event probabilities") +
    ggtitle("Distribution of difference from\nsimulated predicted event proportions") +
    ylab("Density") + geom_vline(xintercept = delta_star,
    linetype = "dashed") + geom_vline(xintercept = 0,
    linetype = "dotted") + scale_x_continuous(breaks = c(-0.5,
    -0.25, 0, 0.035, 0.25, 0.5), labels = c(-0.5, -0.25,
    0, 0.035, 0.25, 0.5), limits = c(-0.5, 0.5))</pre>
```



Assumed event probabilities: Control arm (p_0=0.01), Active arm (p_1=0.01)



Noninformative prior: Agresti and Min suggest to use a 'diffuse' prior and recommend to use Jeffrey's

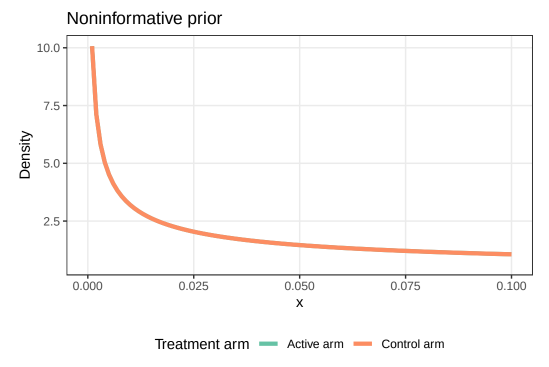
prior [11]. Jeffrey's prior for a binomial likelihood is beta(0.5, 0.5)-distributed.

- Parameters of beta prior for active treatment arm: $a_1 = 0.5, b_1 = 0.5$.
- Parameters of beta prior for control treatment arm: $a_0 = 0.5, b_0 = 0.5$.

```
## Noninformative prior
a_0 < -0.5
b_0 < -0.5
a_1 < 0.5
b_1 < 0.5
prior_0 <- dbeta(x, a_0, b_0)</pre>
prior_1 <- dbeta(x, a_1, b_1)</pre>
prior_plot <- data.frame(x, prior_0, prior_1)</pre>
prior_plot_long <- pivot_longer(prior_plot, cols = c(prior_0,</pre>
    prior_1))
prior_plot_long$group <- ifelse(str_detect(prior_plot_long$name,</pre>
    "_0") == T, "Control arm", "Active arm")
fig1 <- ggplot(prior_plot_long, aes(x = x, y = value,
    colour = group)) + geom_line(linewidth = 1.5) +
    theme_bw() + theme(panel.grid.minor = element_blank(),
    legend.position = "bottom", legend.direction = "horizontal") +
    scale_x_continuous(limits = c(0, 0.1)) + ylab("Density") +
    labs(caption = str_glue("Assumed event probabilities: Control arm (p_0=",
        p_0, "), ", "Active arm <math>(p_1=", p_1, ")") +
    scale_color_brewer("Treatment arm", palette = "Set2") +
    scale_linetype("Type") + ggtitle("Noninformative prior")
diff_prior <- data.frame(x = rbbinom(1e+06, size = ceiling(n_1),
```

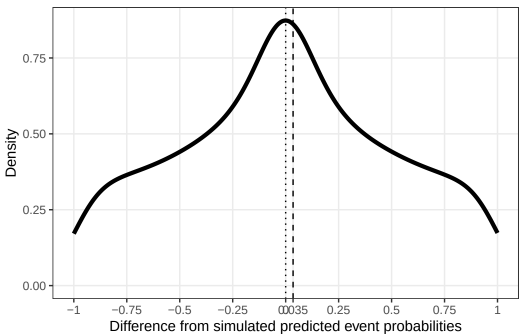
```
a_1, b_1)/ceiling(n_1) - rbbinom(1e+06, size = ceiling(n_0),
a_0, b_0)/ceiling(n_0))

fig2 <- ggplot(diff_prior, aes(x), group = x) + geom_density(linewidth = 1.5,
bw = 0.1) + theme_bw() + theme(panel.grid.minor = element_blank()) +
    xlab("Difference from simulated predicted event probabilities") +
    ggtitle("Distribution of difference from\nsimulated predicted event proportions") +
    ylab("Density") + geom_vline(xintercept = delta_star,
    linetype = "dashed") + geom_vline(xintercept = 0,
    linetype = "dotted") + scale_x_continuous(breaks = c(-1, -0.75, -0.5,
    -0.25, 0, 0.035, 0.25, 0.5, 0.75, 1), labels = c(-1, -0.75, -0.5, -0.25,
    0, 0.035, 0.25, 0.5, 0.75, 1), limits = c(-1, 1))</pre>
```



Assumed event probabilities: Control arm (p_0=0.01), Active arm (p_1=0.01)

Distribution of difference from simulated predicted event proportions



One can calculate the average power as follows:

2.2.1.4.1 Approach 1: Joint prior predictive probability

The joint probability approach has been described in [6] and includes the following steps:

- Calculate the joint pdf for all pairs $r_i \in \{0,1,\cdots,n_i\}, i \in \{0,1\}.$
- The sum of the joint predictive probabilites over r_i pairs where the (1α) -upper confidence limit of the risk difference is smaller than the non-inferiority margin is the average power.

Here we use Agresti-Caffo confidence intervals for the risk difference [12].

```
x <- c()
library(extraDistr)
library(PropCIs)
res_ap <- c()
## Enthusiastic prior
a_1 < 0.5
b_1 \leftarrow (1 - p_1)/p_1 * a_1
a_0 <- 7.2
b_0 < (1 - 0.03)/0.03 * a_0
r_0 \leftarrow 0:(ceiling(n_0)-1)
r_1 < 0: (ceiling(n_1) - 1)
data_points <- expand.grid(r_1 = r_1, r_0 = r_0)
data_points$pdf_r_1 <- dbbinom(data_points$r_1, size = ceiling(n_1), a_1,
```

```
b_1)
data_points$pdf_r_0 <- dbbinom(data_points$r_0, size = ceiling(n_0), a_0,
    b 0)
data_points$joint_pdf <- data_points$pdf_r_1 * data_points$pdf_r_0</pre>
data_points$sig <- NA
for (i in 1:nrow(data_points)) {
    data_points$sig[i] <- ifelse(wald2ci(data_points$r_1[i], n_1, data_points$r_0[i],</pre>
        n_0, conf.level = 0.95, adjust = "AC")$conf.int[2] <= delta_star,</pre>
        1, 0)
}
res_ap <- data.frame(type = "Enthusiastic",</pre>
                              ap = sum(data_points$joint_pdf[data_points$sig == 1]))
## Skeptical prior
a_1 <- 2.84
b_1 \leftarrow (1 - 0.05)/0.05 * a_1
a_0 < 1.24
b_0 < (1 - 0.01)/0.01 * a_0
r_0 \leftarrow 0:(ceiling(n_0)-1)
r_1 \leftarrow 0:(ceiling(n_1)-1)
data_points <- expand.grid(r_1 = r_1, r_0 = r_0)
data_points$pdf_r_1 <- dbbinom(data_points$r_1, size = ceiling(n_1), a_1,</pre>
    b_1)
```

```
data_points$pdf_r_0 <- dbbinom(data_points$r_0, size = ceiling(n_0), a_0,</pre>
    b 0)
data_points$joint_pdf <- data_points$pdf_r_1 * data_points$pdf_r_0</pre>
data_points$sig <- NA
for (i in 1:nrow(data_points)) {
    data_points$sig[i] <- ifelse(wald2ci(data_points$r_1[i], n_1, data_points$r_0[i],
        n_0, conf.level = 0.95, adjust = "AC")$conf.int[2] <= delta_star,</pre>
        1, 0)
}
res_ap <- rbind(res_ap, data.frame(type = "Skeptical",</pre>
                            ap = sum(data_points$joint_pdf[data_points$sig == 1])))
## Informative prior
a_1 <- 0.8
b_1 \leftarrow (1 - p_1)/p_1 * a_1
a_0 < 0.03
b_0 < (1 - p_0)/p_0 * a_0
r_0 \leftarrow 0:(ceiling(n_0)-1)
r_1 \leftarrow 0:(ceiling(n_1)-1)
data_points <- expand.grid(r_1 = r_1, r_0 = r_0)
data_points$pdf_r_1 <- dbbinom(data_points$r_1, size = ceiling(n_1), a_1,</pre>
    b_1)
data_points$pdf_r_0 <- dbbinom(data_points$r_0, size = ceiling(n_0), a_0,
```

```
b_0)
data_points$joint_pdf <- data_points$pdf_r_1 * data_points$pdf_r_0</pre>
data_points$sig <- NA</pre>
for (i in 1:nrow(data_points)) {
    data_points$sig[i] <- ifelse(wald2ci(data_points$r_1[i], n_1, data_points$r_0[i],</pre>
        n_0, conf.level = 0.95, adjust = "AC")$conf.int[2] <= delta_star,</pre>
        1, 0)
}
res_ap <- rbind(res_ap, data.frame(type = "Informative",</pre>
                            ap = sum(data_points$joint_pdf[data_points$sig == 1])))
## Noninformative prior
a_1 < 0.5
b_1 < 0.5
a_0 < -0.5
b_0 < 0.5
r_0 \leftarrow 0:(ceiling(n_0)-1)
r_1 \leftarrow 0:(ceiling(n_1)-1)
data_points <- expand.grid(r_1 = r_1, r_0 = r_0)
data_points$pdf_r_1 <- dbbinom(data_points$r_1, size = ceiling(n_1), a_1,</pre>
data_points$pdf_r_0 <- dbbinom(data_points$r_0, size = ceiling(n_0), a_0,</pre>
    b_0)
```

```
data_points$joint_pdf <- data_points$pdf_r_1 * data_points$pdf_r_0
  data_points$sig <- NA
  for (i in 1:nrow(data_points)) {
      data_points$sig[i] <- ifelse(wald2ci(data_points$r_1[i], n_1, data_points$r_0[i],
          n_0, conf.level = 0.95, adjust = "AC")$conf.int[2] <= delta_star,</pre>
          1, 0)
  }
  res_ap <- rbind(res_ap, data.frame(type = "Noninformative prior",</pre>
                              ap = sum(data_points$joint_pdf[data_points$sig == 1])))
  res_ap$ap <- round(res_ap$ap, 2)</pre>
  res_ap
                  type
1
          Enthusiastic 0.76
2
             Skeptical 0.09
3
           Informative 0.55
```

2.2.1.4.2 Approach 2: Sampling from prior predictive distribution

4 Noninformative prior 0.40

The second approach samples q event outcomes from the predictive distribution and calculates for each draw whether the $(1-\alpha)$ -upper confidence limit of the risk difference is smaller than the non-inferiority margin. The sum of positive outcomes divided by the number of samples q gives the average power. Also here we use Agresti-Caffo confidence intervals for the risk difference [12].

```
# Number of draws
q <- 10000
res_ap <- c()
## Enthusiastic prior
a_1 < 0.5
b_1 \leftarrow (1 - p_1)/p_1 * a_1
a_0 < 7.2
b_0 < (1 - 0.03)/0.03 * a_0
set.seed(1)
r_1 \leftarrow rbbinom(q, size = ceiling(n_1), a_1, b_1)
r_0 \leftarrow rbbinom(q, size = ceiling(n_0), a_0, b_0)
res <- c()
for (i in 1:length(r_1)) {
    res <- c(res, ifelse(wald2ci(r_1[i], n_1, r_0[i], n_0, conf.level = 0.95,
        adjust = "AC")$conf.int[2] <= delta_star, 1, 0))</pre>
}
res_ap <- data.frame(type = "Enthusiastic", ap = sum(res)/q)</pre>
## Skeptical prior
a_1 <- 2.84
b_1 < (1 - 0.05)/0.05 * a_1
a_0 <- 1.24
```

```
b_0 < (1 - p_0)/p_0 * a_0
set.seed(1)
r_1 \leftarrow rbbinom(q, size = ceiling(n_1), a_1, b_1)
r_0 \leftarrow rbbinom(q, size = ceiling(n_0), a_0, b_0)
res <- c()
for (i in 1:length(r_1)) {
    res \leftarrow c(res, ifelse(wald2ci(r_1[i], n_1, r_0[i], n_0, conf.level = 0.95,
        adjust = "AC")$conf.int[2] <= delta_star, 1, 0))</pre>
}
res_ap <- rbind(res_ap, data.frame(type = "Skeptical", ap = sum(res)/q))</pre>
## Informative prior
a_1 <- 0.8
b_1 \leftarrow (1 - p_1)/p_1 * a_1
a_0 < 0.03
b_0 < (1 - p_0)/p_0 * a_0
set.seed(1)
r_1 \leftarrow rbbinom(q, size = ceiling(n_1), a_1, b_1)
r_0 \leftarrow rbbinom(q, size = ceiling(n_0), a_0, b_0)
res <- c()
for (i in 1:length(r_1)) {
    res <- c(res, ifelse(wald2ci(r_1[i], n_1, r_0[i], n_0, conf.level = 0.95,
```

```
adjust = "AC")$conf.int[2] <= delta_star, 1, 0))</pre>
}
res_ap <- rbind(res_ap, data.frame(type = "Informative", ap = sum(res)/q))</pre>
## Noninformative prior
a_0 < -0.5
b_0 < -0.5
a_1 < 0.5
b_1 < 0.5
set.seed(1)
r_1 \leftarrow rbbinom(q, size = ceiling(n_1), a_1, b_1)
r_0 \leftarrow rbbinom(q, size = ceiling(n_0), a_0, b_0)
res <- c()
for (i in 1:length(r_1)) {
    res <- c(res, ifelse(wald2ci(r_1[i], n_1, r_0[i], n_0, conf.level = 0.95,
        adjust = "AC")$conf.int[2] <= delta_star, 1, 0))</pre>
}
res_ap <- rbind(res_ap, data.frame(type = "Noninformative", ap = sum(res)/q))</pre>
res_ap$ap <- round(res_ap$ap, 2)</pre>
res_ap
          type
 Enthusiastic 0.76
```

- 2 Skeptical 0.09
- 3 Informative 0.55
- 4 Noninformative 0.45

2.2.1.5 Bayesian approach: Prior on event probabilities

In the former subsection we assumed that the prior was directly specified on the treatment effect. In this subsection we assume that that design prior is specified on the event probabilities p_i , $i = \{0, 1\}$. Suppose that the priors on the event probability are beta distributed $\pi_i \sim Beta(a_i, b_i)$ and that we have observed a binomial distributed random sample of size n_i and number of events r_i . Then the posterior distribution is

$$\pi_i | r_i \sim Beta\left(a_i + r_i, b_i + n_i - r_i\right) = \frac{\pi_i^{a_i + r_i - 1} (1 - \pi_i)^{b_i + n_i - r_i - 1}}{B(a_i + r_i, b_i + n_i - r_i)}$$

with $B(\cdot)$ the beta function and

$$\mu_{i,r_i} := E(\pi_i|r_i) = \frac{a_i + r_i}{a_i + b_i + n_i}, \quad \sigma^2_{i,r_i} := Var(\pi_i|r_i) = \frac{(a_i + r_i)\,(b_i + n_i - r_i)}{(a_i + b_i + n_i)^2(a_i + b_i + n_i + 1)}.$$

Working example (continued)

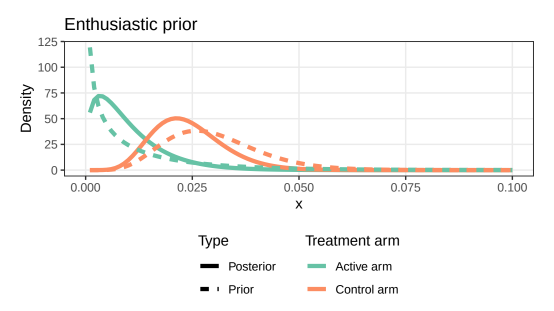
We use the same prior specifications as from the former subsection for the analysis priors.

Enthusiastic prior: The beta prior for the active treatment arm is centered at 0.01 (the expected event proportion) with $P(\pi_1 > 0.05) = 0.025$, that is, the probability that the prior is greater than the safety margin is 2.5%. The mean of the beta prior for the control arm is centered at 0.03 (the safety margin) with $P(\pi_0 > 0.05) = 0.05$.

- Parameters of beta prior for active treatment arm: $a_1 = 0.5, b_1 = 49.5$.
- Parameters of beta prior for control treatment arm: $a_0 = 7.2, b_0 = 232.8$.

```
x <- seq(0.001, 0.999, 0.001)
## Enthusiastic prior</pre>
```

```
a_1 < 0.5
b_1 \leftarrow (1 - p_1)/p_1 * a_1
a_0 < 7.2
b_0 < (1 - 0.03)/0.03 * a_0
prior_0 <- dbeta(x, a_0, b_0)</pre>
prior_1 <- dbeta(x, a_1, b_1)</pre>
posterior_0 <- dbeta(x, a_0 + n_0 * p_0, b_0 + n_0 - n_0 * p_0)
posterior_1 <- dbeta(x, a_1 + n_1 * p_1, b_1 + n_1 - n_1 * p_1)
var_posterior_0 \leftarrow ((a_0 + n_0 * p_0) * (b_0 + n_0 - n_0 * p_0))/((a_0 + n_0) + n_0)
    b_0 + n_0^2 * (a_0 + b_0 + n_0 + 1)
var_posterior_1 \leftarrow ((a_1 + n_1 * p_1) * (b_1 + n_1 - n_1 * p_1))/((a_1 + n_1) + n_2)
    b_1 + n_1^2 * (a_1 + b_1 + n_1 + 1)
prior_plot <- data.frame(x, prior_0, prior_1, posterior_0, posterior_1)</pre>
prior_plot_long <- pivot_longer(prior_plot, cols = c(prior_0, prior_1,</pre>
    posterior_0, posterior_1))
prior_plot_long$type <- ifelse(str_detect(prior_plot_long$name, "posterior") ==</pre>
    T, "Posterior", "Prior")
prior_plot_long$group <- ifelse(str_detect(prior_plot_long$name, "_0") ==</pre>
    T, "Control arm", "Active arm")
ggplot(prior_plot_long, aes(x = x, y = value, linetype = type, colour = group)) +
    geom_line(linewidth = 1.5) + theme_bw() + theme(panel.grid.minor = element_blank(),
    legend.position = "bottom", legend.direction = "vertical") + scale_x_continuous(limits = c
    0.1)) + ylab("Density") + labs(caption = str_glue("Assumed event probabilities: Control arm
    p_0, "), ", "Active arm (p_1=", p_1, ")\nPosterior standard deviation: Control arm=",
    round(sqrt(var_posterior_0), 3), ", ", "Active arm=", round(sqrt(var_posterior_1),
        3))) + scale_color_brewer("Treatment arm", palette = "Set2") +
    scale_linetype("Type") + ggtitle("Enthusiastic prior")
```



Assumed event probabilities: Control arm (p_0=0.01), Active arm (p_1=0.01) Posterior standard deviation: Control arm=0.008, Active arm=0.008

Skeptical prior: The beta prior for the active treatment arm is centered at 0.05 (the safety margin) with $P(\pi_1 \le 0.01) = 0.025$, that is the probability that the prior is smaller or equal than the expected event proportion is 2.5%. The mean of the beta prior for the control arm is is centered at 0.01 (the expected event proportion) with $P(\pi_0 > 0.05) = 0.025$.

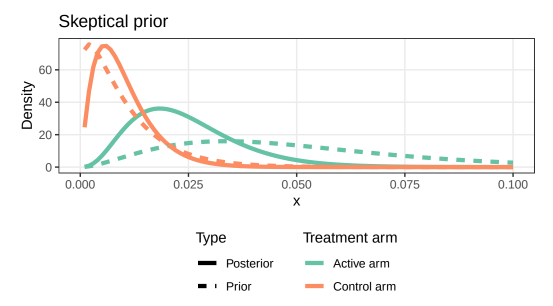
- Parameters of beta prior for active treatment arm: $a_1 = 2.84$, $b_1 = 53.96$.
- Parameters of beta prior for control treatment arm: $a_i = 1.24, b_i = 122.76.$

```
x <- seq(0.001, 0.999, 0.001)

# Skeptical prior
a_1 <- 2.84
b_1 <- (1 - 0.05)/0.05 * a_1
a_0 <- 1.24
b_0 <- (1 - p_0)/p_0 * a_0

prior_0 <- dbeta(x, a_0, b_0)</pre>
```

```
prior_1 <- dbeta(x, a_1, b_1)</pre>
posterior_0 <- dbeta(x, a_0 + n_0 * p_0, b_0 + n_0 - n_0 * p_0)
posterior_1 <- dbeta(x, a_1 + n_1 * p_1, b_1 + n_1 - n_1 * p_1)
var_posterior_0 \leftarrow ((a_0 + n_0 * p_0) * (b_0 + n_0 - n_0 * p_0))/((a_0 + n_0) + n_0)
    b_0 + n_0^2 * (a_0 + b_0 + n_0 + 1)
var_posterior_1 \leftarrow ((a_1 + n_1 * p_1) * (b_1 + n_1 - n_1 * p_1))/((a_1 + n_1) + n_1)
    b_1 + n_1^2 * (a_1 + b_1 + n_1 + 1)
prior_plot <- data.frame(x, prior_0, prior_1, posterior_0, posterior_1)</pre>
prior_plot_long <- pivot_longer(prior_plot, cols = c(prior_0, prior_1,</pre>
    posterior_0, posterior_1))
prior_plot_long$type <- ifelse(str_detect(prior_plot_long$name, "posterior") ==</pre>
    T, "Posterior", "Prior")
prior_plot_long$group <- ifelse(str_detect(prior_plot_long$name, "_0") ==</pre>
    T, "Control arm", "Active arm")
ggplot(prior_plot_long, aes(x = x, y = value, linetype = type, colour = group)) +
    geom_line(linewidth = 1.5) + theme_bw() + theme(panel.grid.minor = element_blank(),
    legend.position = "bottom", legend.direction = "vertical") + scale_x_continuous(limits = c
    0.1)) + ylab("Density") + labs(caption = str_glue("Expected event probabilities: Control as
    p_0, "), ", "Active arm (p_1=", p_1, ")\nPosterior standard deviation: Control arm=",
    round(sqrt(var_posterior_0), 3), ", ", "Active arm=", round(sqrt(var_posterior_1),
        3))) + scale_color_brewer("Treatment arm", palette = "Set2") +
    scale_linetype("Type") + ggtitle("Skeptical prior")
```



Expected event probabilities: Control arm (p_0=0.01), Active arm (p_1=0.01) Posterior standard deviation: Control arm=0.007, Active arm=0.012

Informative prior: The beta prior for the active treatment arm and control arm is centered at 0.01 (the expected event proportion) with $P(\pi_1 \le 0.05) = 0.01$ and $P(\pi_1 \le 0.05) = 0.05$.

- Parameters of beta prior for active treatment arm: $a_1 = 0.8, b_1 = 79.2$.
- Parameters of beta prior for control treatment arm: $a_0 = 0.03, b_0 = 2.97.$

```
x <- seq(0.001, 0.999, 0.001)

## Informative prior
a_1 <- 0.8

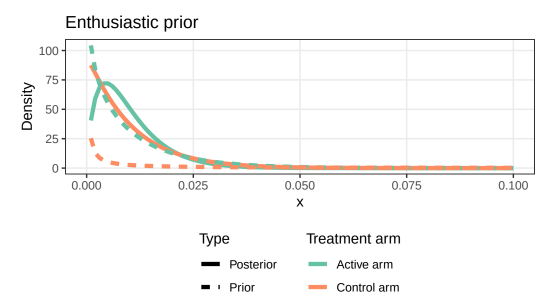
b_1 <- (1 - p_1)/p_1 * a_1
a_0 <- 0.03

b_0 <- (1 - p_0)/p_0 * a_0

prior_0 <- dbeta(x, a_0, b_0)
prior_1 <- dbeta(x, a_1, b_1)

posterior_0 <- dbeta(x, a_0 + n_0 * p_0, b_0 + n_0 - n_0 * p_0)</pre>
```

```
posterior_1 <- dbeta(x, a_1 + n_1 * p_1, b_1 + n_1 - n_1 * p_1)
var_posterior_0 <- ((a_0 + n_0 * p_0) * (b_0 + n_0 - n_0 * p_0))/((a_0 + n_0) + n_0) <- (a_0 + a_0) + a_0 <- (a_0 + a_0) <- 
          b_0 + n_0^2 * (a_0 + b_0 + n_0 + 1)
var_posterior_1 \leftarrow ((a_1 + n_1 * p_1) * (b_1 + n_1 - n_1 * p_1))/((a_1 + n_1 + n_1))
          b_1 + n_1^2 * (a_1 + b_1 + n_1 + 1)
prior_plot <- data.frame(x, prior_0, prior_1, posterior_0, posterior_1)</pre>
prior_plot_long <- pivot_longer(prior_plot, cols = c(prior_0, prior_1,</pre>
          posterior_0, posterior_1))
prior_plot_long$type <- ifelse(str_detect(prior_plot_long$name, "posterior") ==</pre>
          T, "Posterior", "Prior")
prior_plot_long$group <- ifelse(str_detect(prior_plot_long$name, "_0") ==</pre>
          T, "Control arm", "Active arm")
ggplot(prior_plot_long, aes(x = x, y = value, linetype = type, colour = group)) +
          geom_line(linewidth = 1.5) + theme_bw() + theme(panel.grid.minor = element_blank(),
          legend.position = "bottom", legend.direction = "vertical") + scale_x_continuous(limits = c
          0.1)) + ylab("Density") + labs(caption = str_glue("Assumed event probabilities: Control arm
          p_0, "), ", "Active arm (p_1=", p_1, ")\nPosterior standard deviation: Control arm=",
          round(sqrt(var_posterior_0), 3), ", ", "Active arm=", round(sqrt(var_posterior_1),
                     3))) + scale_color_brewer("Treatment arm", palette = "Set2") +
           scale_linetype("Type") + ggtitle("Enthusiastic prior")
```



Assumed event probabilities: Control arm (p_0=0.01), Active arm (p_1=0.01) Posterior standard deviation: Control arm=0.01, Active arm=0.007

Noninformative prior: Agresti and Min suggest to use a 'diffuse' prior and recommend to use Jeffrey's prior [11]. Jeffrey's prior for a binomial likelihood is beta(0.5, 0.5)-distributed.

- Parameters of beta prior for active treatment arm: $a_1 = 0.5, b_1 = 0.5$.
- Parameters of beta prior for control treatment arm: $a_0 = 0.5, b_0 = 0.5$.

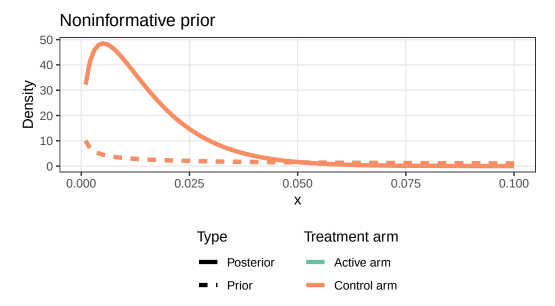
```
## Noninformative prior
a_0 <- 0.5
b_0 <- 0.5
a_1 <- 0.5
b_1 <- 0.5

prior_0 <- dbeta(x, a_0, b_0)
prior_1 <- dbeta(x, a_1, b_1)

posterior_0 <- dbeta(x, a_0 + n_0 * p_0, b_0 + n_0 - n_0 * p_0)

posterior_1 <- dbeta(x, a_1 + n_1 * p_1, b_1 + n_1 - n_0 * p_0)</pre>
```

```
n_1 * p_1
var_posterior_0 \leftarrow ((a_0 + n_0 * p_0) * (b_0 + n_0 - p_0)) * (b_0 + p_0)
          n_0 * p_0))/((a_0 + b_0 + n_0)^2 * (a_0 + b_0 +
          n_0 + 1)
var_posterior_1 \leftarrow ((a_1 + n_1 * p_1) * (b_1 + n_1 - a_1) * (b_1 + a_2) + a_2 + a_3 + a_4 + a_2 + a_4 + a_2 + a_3 + a_4 + a_4
           n_1 * p_1)/((a_1 + b_1 + n_1)^2 * (a_1 + b_1 +
          n_1 + 1)
prior_plot <- data.frame(x, prior_0, prior_1, posterior_0,</pre>
           posterior_1)
prior_plot_long <- pivot_longer(prior_plot, cols = c(prior_0,</pre>
           prior_1, posterior_0, posterior_1))
prior_plot_long$type <- ifelse(str_detect(prior_plot_long$name,</pre>
           "posterior") == T, "Posterior", "Prior")
prior_plot_long$group <- ifelse(str_detect(prior_plot_long$name,</pre>
           "_0") == T, "Control arm", "Active arm")
ggplot(prior_plot_long, aes(x = x, y = value, linetype = type,
           colour = group)) + geom_line(linewidth = 1.5) +
           theme_bw() + theme(panel.grid.minor = element_blank(),
           legend.position = "bottom", legend.direction = "vertical") +
           scale_x_continuous(limits = c(0, 0.1)) + ylab("Density") +
           labs(caption = str_glue("Assumed event probabilities: Control arm (p_0=",
                     p_0, "), ", "Active arm (p_1=", p_1, ")\nPosterior standard deviation: Control arm=",
                     round(sqrt(var_posterior_0), 3), ", ", "Active arm=",
                     round(sqrt(var_posterior_1), 3))) + scale_color_brewer("Treatment arm",
           palette = "Set2") + scale_linetype("Type") + ggtitle("Noninformative prior")
```



Assumed event probabilities: Control arm (p_0=0.01), Active arm (p_1=0.01) Posterior standard deviation: Control arm=0.012, Active arm=0.012

We denote the joint posterior distribution as

$$f_2(\pi_1,\pi_0|r_0,r_1,n_0,n_1) = \prod_{i \in \{0,1\}} \frac{\pi_i^{a_i+r_i-1}(1-\pi_i)^{b_i+n_i-r_i-1}}{B(a_i+r_i,b_i+n_i-r_i)}.$$

and thus, by using Fubini's theorem and the guidance of Patricia Altham [13],

$$\begin{split} P\left(\pi_{1}-\pi_{0} \leq \delta^{*}|\,r_{0},r_{1},n_{0},n_{1}\right) &= \int_{0}^{1} \int_{0}^{\pi_{0}+\delta^{*}} f_{2}(\pi_{1},\pi_{0}|r_{0},r_{1},n_{0},n_{1}) d\pi_{1} d\pi_{0} \\ &= \int_{0}^{1} \frac{\pi_{0}^{a_{0}+r_{0}-1}(1-\pi_{0})^{b_{0}+n_{0}-r_{0}-1}}{B(a_{0}+r_{0},b_{0}+n_{0}-r_{0})} \left(\int_{0}^{\pi_{0}+\delta^{*}} \frac{\pi_{1}^{a_{1}+r_{1}-1}(1-\pi_{1})^{b_{1}+n_{1}-r_{1}-1}}{B(a_{1}+r_{1},b_{1}+n_{1}-r_{1})} d\pi_{1} \right) d\pi_{0} \\ &\stackrel{u=\pi_{0}-\delta^{*}}{=} \int_{-\delta^{*}}^{1-\delta^{*}} \frac{u^{a_{0}+r_{0}-1}(1-u)^{b_{0}+n_{0}-r_{0}-1}}{B(a_{0}+r_{0},b_{0}+n_{0}-r_{0})} \left(\int_{0}^{u} \frac{\pi_{1}^{a_{1}+r_{1}-1}(1-\pi_{1})^{b_{1}+n_{1}-r_{1}-1}}{B(a_{1}+r_{1},b_{1}+n_{1}-r_{1})} d\pi_{1} \right) du \\ &= \cdots \end{split}$$

is the Bayesian probability to reject given assumed r_0, r_1, n_0, n_1 . This integral can be solved using

the integrate function in R, or using the continuation of the formula

$$\begin{split} \cdots &= \int_{-\delta^*}^{1-\delta^*} \frac{u^{a_0+r_0-1}(1-u)^{b_0+n_0-r_0-1}}{B(a_0+r_0,b_0+n_0-r_0)} \left(\sum_{s=a_1+r_1}^{a_1+b_1+n_1-1} \left(a_1+b_1+n_1-1 \right) u^s (1-u)^{a_1+b_1+n_1-1-s} \right) du \\ &= \sum_{s=a_1+r_1}^{a_1+b_1+n_1-1} \left(a_1+b_1+n_1-1 \right) \frac{1}{B(a_0+r_0,b_0+n_0-r_0)} \left(\int_{-\delta^*}^{1-\delta^*} u^{a_0+r_0+s-1} (1-u)^{a_1+b_1+n_1-1+b_0+n_0-r_0-s-1} du \right) \\ &= \sum_{s=a_1+r_1}^{a_1+b_1+n_1-1} \left(a_1+b_1+n_1-1 \right) \frac{1}{B(a_0+r_0,b_0+n_0-r_0)} \left(\int_0^1 z^{a_0+r_0+s-1} (1-z)^{a_1+b_1+n_1-1-s+b_0+n_0-r_0-1} dz \right) \\ &= \sum_{s=a_1+r_1}^{a_1+b_1+n_1-1} \left(a_1+b_1+n_1-1 \right) \frac{B(a_0+r_0+s,a_1+b_1+n_1-1-s+b_0+n_0-r_0)}{B(a_0+r_0,b_0+n_0-r_0)}, \end{split}$$

where the last term in the above formula is the upper tail of a beta-binomial distribution. Altham [13] showed that this could also be written as a hypergeometric distribution (see also [6], [14]).

Working example (continued)

We calculate the Bayesian probability to reject for the SAFE-SSPE study. We show the integral solution and the calculation from the upper tail of a beta-binomial distribution.

Note that for the integral calculation the lower integrand $-\delta^*$ in

$$\int_{-\delta^*}^{1-\delta^*} \frac{u^{a_0+r_0-1}(1-u)^{b_0+n_0-r_0-1}}{B(a_0+r_0,b_0+n_0-r_0)} \left(\int_0^u \frac{\pi_1^{a_1+r_1-1}(1-\pi_1)^{b_1+n_1-r_1-1}}{B(a_1+r_1,b_1+n_1-r_1)} d\pi_1 \right) du$$

needs to be zero, otherwise the function is not defined.

```
# Integral function
inner_func <- function(x) {
    x^(a_1 + ceiling(n_1) * p_1 - 1) * (1 - x)^(b_1 +
        ceiling(n_1) - ceiling(n_1) * p_1 - 1)
}</pre>
```

```
inner_int <- Vectorize(function(u) {</pre>
              u^(a_0 + ceiling(n_0) * p_0 - 1) * (1 - u)^(b_0 +
                            ceiling(n_0) - ceiling(n_0) * p_0 - 1) * integrate(inner_func,
                            0, u)$value
})
## Enthusiastic prior
a_1 < 0.5
b_1 < (1 - p_1)/p_1 * a_1
a_0 < 7.2
b_0 < (1 - 0.03)/(0.03) * a_0
bap_integral <- integrate(inner_int, 0, 1-delta_star)$value/(beta(a_1 +</pre>
             n_1 * p_1, b_1 + n_1 - n_1 * p_1) * beta(a_0 + n_0 * p_0, b_0 + n_0 - n_1) * beta(a_0 + n_0) * p_1 *
            n_0 * p_0)
s \leftarrow ceiling(seq(a_1 + n_1 * p_1, b_1 + n_1 - n_1 *
             p_1 - 1))
bap_tail_betabinomial <- sum(dbbinom(s, size = ceiling(a_1 +</pre>
             b_1 + n_1 - 1, alpha = a_0 + n_0 * p_0, beta = b_0 + 1
             n_0 - n_0 * p_0)
res_bap <- data.frame(type="Enthusiastic", bap_integral, bap_tail_betabinomial)</pre>
# Skeptical prior
a_1 <- 2.84
b_1 < (1 - 0.05)/0.05 * a_1
a_0 <- 1.24
b_0 < (1 - p_0)/p_0 * a_0
```

```
bap_integral <- integrate(inner_int, 0, 1 - delta_star)$value/(beta(a_1 +</pre>
                     n_1 * p_1, b_1 + n_1 - n_1 * p_1) * beta(a_0 + n_0 * p_0, b_0 + n_0 - n_1) * beta(a_0 + n_0) * p_1 *
                    n_0 * p_0)
s \leftarrow ceiling(seq(a_1 + n_1 * p_1, b_1 + n_1 - n_1 *
                     p_1 - 1))
bap_tail_betabinomial <- sum(dbbinom(s, size = ceiling(a_1 +</pre>
                      b_1 + n_1 - 1, alpha = a_0 + n_0 * p_0, beta = b_0 + 1
                     n_0 - n_0 * p_0)
res_bap <- rbind(res_bap, data.frame(type="Skeptical", bap_integral,</pre>
                                                                                                                                                                                                             bap_tail_betabinomial))
# Informative prior
a_1 <- 0.8
b_1 \leftarrow (1 - p_1)/p_1 * a_1
a_0 < 0.03
b_0 < (1 - p_0)/p_0 * a_0
bap_integral <- integrate(inner_int, 0, 1 - delta_star)$value/(beta(a_1 +</pre>
                     n_1 * p_1, b_1 + n_1 - n_1 * p_1) * beta(a_0 + n_0 * p_0, b_0 + n_0 - n_1) * beta(a_0 + n_0) * p_1 *
                     n_0 * p_0)
s \leftarrow ceiling(seq(a_1 + n_1 * p_1, b_1 + n_1 - n_1 *
                      p_1 - 1)
bap_tail_betabinomial <- sum(dbbinom(s, size = ceiling(a_1 +</pre>
                       b_1 + n_1 - 1, alpha = a_0 + n_0 * p_0, beta = b_0 + 1
                     n_0 - n_0 * p_0)
```

```
res_bap <- rbind(res_bap, data.frame(type="Informative", bap_integral,</pre>
                                                                                                                                          bap_tail_betabinomial))
        ## Noninformative prior
       a_0 < -0.5
       b_0 < 0.5
        a_1 < 0.5
       b_1 < 0.5
       bap_integral <- integrate(inner_int, 0, 1 - delta_star)$value/(beta(a_1 +</pre>
                     n_1 * p_1, b_1 + n_1 - n_1 * p_1) * beta(a_0 + n_0 * p_0, b_0 + n_0 - n_1) * p_1
                    n_0 * p_0)
        s \leftarrow ceiling(seq(a_1 + n_1 * p_1, b_1 + n_1 - n_1 *
                      p_1 - 1))
        bap_tail_betabinomial <- sum(dbbinom(s, size = ceiling(a_1 +</pre>
                      b_1 + n_1 - 1, alpha = a_0 + n_0 * p_0, beta = b_0 + n_1 + n_2 + n_3 + n_4 + n_4 + n_5 + n
                     n_0 - n_0 * p_0)
       res_bap <- rbind(res_bap, data.frame(type="Noninformative", bap_integral,</pre>
                                                                                                                                           bap_tail_betabinomial))
        res_bap$bap_integral <- round(res_bap$bap_integral, 2)</pre>
        res_bap$bap_tail_betabinomial <- round(res_bap$bap_tail_betabinomial, 2)</pre>
       res_bap
                                          type bap_integral bap_tail_betabinomial
            Enthusiastic
                                                                                       0.88
                                                                                                                                                                     0.83
1
```

2	Skeptical	0.13	0.12
3	Informative	0.45	0.42
4 Noninformative		0.50	0.38

Small differences between both approaches occur because the 'beta-binomial distribution' approach is a discretization of the integral approach and requires integer steps for the calculation.

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