## Instructions for Using 'seqDesign' and Generating Output Tables and Figures Summarizing Operating Characteristics of the Trial Design

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**Step 1.** Specify the per-arm sample sizes in the placebo and vaccine arm, and the total (maximum) number of vaccine arms:

```
N.pla <- 1900
N.vax <- 1100
N.vax.arms <- 4
```

**Step 2.** Simulate data-sets (for each component of aveVElist), apply the monitoring procedures, and extract results needed for generating output tables and figures:

```
aveVElist <- list(-2, -1.5, -1, 0, 0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8)
aveVElist[12:19] <- lapply(aveVElist[-(1:3)], function(aveVE) { rep(aveVE, 4) })</pre>
aveVElist[[20]] \leftarrow rep(0.5, 2)
aveVElist[[21]] <- rep(0.5, 3)</pre>
aveVElist[[22]] \leftarrow c(0, 0,
                                  0,
                                         0.4)
aveVElist[[23]] \leftarrow c(0, 0, 0.3, 0.4)
aveVElist[[23]] \leftarrow c(0.2, 0.2, 0.3, 0.4)
aveVElist[[24]] \leftarrow c(0, 0,
aveVElist[[25]] \leftarrow c(0, 0,
                                  0.3.0.6
aveVElist[[26]] \leftarrow c(0, 0,
                                  0.45, 0.6)
aveVElist[[27]] \leftarrow c(0.3, 0.3, 0.45, 0.6)
aveVElist[[28]] \leftarrow c(0.3, 0.45, 0.45, 0.6)
aveVElist[[29]] \leftarrow rep(0, 2)
aveVElist[[30]] <- c(0.4, 0)
aveVElist[[31]] \leftarrow c(0.4, 0.4)
aveVElist[[32]] \leftarrow rep(0, 3)
aveVElist[[33]] <- c(0.4, 0,
aveVElist[[34]] \leftarrow c(0.4, 0.4, 0)
aveVElist[[35]] \leftarrow c(0.4, 0.4, 0.4)
aveVElist[[36]] <- c(0.4, 0,
aveVElist[[37]] \leftarrow c(0.4, 0.4, 0,
aveVElist[[38]] \leftarrow c(0.4, 0.4, 0.4, 0)
aveVElist[[39]] \leftarrow rep(0.4, 4)
for (i in 1:length(aveVElist)){
  simTrial(N=c(N.pla, rep(N.vax, length(aveVElist[[i]]))), aveVE=c(0, aveVElist[[i]]),
            VEmodel="half", vePeriods=c(1, 27, 79), enrollPeriod=78, enrollPartial=13,
            enrollPartialRelRate=0.5, dropoutRate=0.05, infecRate=0.04, fuTime=156,
            visitSchedule=c(0, (13/3)*(1:4), seq(13*6/3, 156, by=13*2/3)),
            missVaccProb=c(0,0.05,0.1,0.15), VEcutoffWeek=26, nTrials=30,
            stage1=78, saveDir="./", randomSeed=300)
  monitorTrial(dataFile=
                pasteO("simTrial_nPlac=", N.pla, "_nVacc=",
                       paste(rep(N.vax, length(aveVElist[[i]])), collapse="_"),
                        "_aveVE=", paste(aveVElist[[i]], collapse="_"), "_infRate=0.04.RData"),
```

```
stage1=78, stage2=156, harmMonitorRange=c(10,100), alphaPerTest=0.0106,
               minCnt=50, minPct=0.33, week1=26, minCnt2=2, week2=52, nonEffInterval=20,
               nullVE=0, altVE=0.4, highVE=0.6, alpha=0.025, estimand="combined",
               VEcutoffWeek=26, saveDir="./")
  censTrial(dataFile=
            pasteO("simTrial_nPlac=", N.pla, "_nVacc=",
                   paste(rep(N.vax, length(aveVElist[[i]])), collapse="_"),
                   "_aveVE=", paste(aveVElist[[i]], collapse="_"), "_infRate=0.04.RData"),
            monitorFile=
            pasteO("monitorTrial_nPlac=", N.pla, "_nVacc=",
                   paste(rep(N.vax, length(aveVElist[[i]])), collapse="_"),
                   "_aveVE=", paste(aveVElist[[i]], collapse="_"), "_infRate=0.04_combined.RData"),
            stage1=78, stage2=156, saveDir="./")
  if (i %in% 22:28){
      rankTrial(censFile=
                pasteO("trialDataCens_nPlac=", N.pla, "_nVacc=",
                       paste(rep(N.vax, length(aveVElist[[i]])), collapse="_"),
                       "_aveVE=", paste(aveVElist[[i]], collapse="_"), "_infRate=0.04_combined.RData"),
                idxHighestVE=2, headHead=matrix(c(4,3), nrow=1, ncol=2),
                poolHead=matrix(c(3,4,1,2), nrow=1, ncol=4), stage1=78, stage2=156,
                alpha=0.025, saveDir="./")
VEpowerPP(dataList=
          as.list(paste0("simTrial_nPlac=", N.pla, "_nVacc=", N.vax, "_aveVE=",
                        do.call("c", aveVElist[4:11]), "_infRate=0.04.RData")),
          VEcutoffWeek=26, stage1=78, alpha=0.025,
          outName=paste0("VEpwPP_nPlac=", N.pla, "_nVacc=", N.vax, "_infRate=0.04.RData"),
          saveDir="./")
```

Step 3. Update the variables N.pla, N.vax, N.vax.arms, and srcDir in the first R chunk of seqDesignReport.Rnw in the extdata subdirectory and compile the PDF report. The full path to seqDesignReport.Rnw can be obtained by:

```
system.file("extdata/seqDesignReport.Rnw", package="seqDesign")
```

Some changes in table/figure captions and figure labels might be needed.

Note: The sample PDF report generated by seqDesignReport.Rnw can be found in seqDesignSampleReport.pdf.