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Author Marie-Karelle Riviere and Jacques-Henri Jourdan
Maintainer Jacques-Henri Jourdan <jacques-henri.jourdan@normalesup.org>
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**dfcomb-package**

**Phase I/II Adaptive Dose-Finding Design for Combination Studies**

**Description**

Phase I/II adaptive dose-finding design for combination studies where toxicity rates are supposed to increase with both agents.

**Details**

The DESCRIPTION file:

- **Package:** dfcomb
- **Type:** Package
- **Title:** Phase I/II Adaptive Dose-Finding Design for Combination Studies
- **Version:** 3.1-1
- **Date:** 2022-12-26
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- **Description:** Phase I/II adaptive dose-finding design for combination studies where toxicity rates are supposed to increase.
- **License:** GPL-3
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- **LinkingTo:** BH (>= 1.55), Rcpp, RcppProgress (>= 0.2.1)
- **NeedsCompilation:** yes

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- CombIncrease_sim: Combination design Simulator using Logistic model
- dfcomb-package: Phase I/II Adaptive Dose-Finding Design for Combination Studies

**Author(s)**

Marie-Karelle Riviere and Jacques-Henri Jourdan

Maintainer: Jacques-Henri Jourdan <jacques-henri.jourdan@normalesup.org>

**References**

Combination determination with logistic model

Description

CombIncrease_next is used to determine the next or recommended combination in a phase I combination clinical trial using the design proposed by Riviere et al. entitled "A Bayesian dose-finding design for drug combination clinical trials based on the logistic model".

Usage

CombIncrease_next(ndose_a1, ndose_a2, target, target_min, target_max, prior_tox_a1, prior_tox_a2, cohort, final, pat_incl, dose_adm1, dose_adm2, tite=FALSE, toxicity, time_full=0, time_tox=0, time_follow=0, c_e=0.85, c_d=0.45, c_stop=0.95, c_t=0.5, c_over=0.25, cmin_overunder=2, cmin_mtd=3, cmin_recom=1, early_stop=1, alloc_rule=1, nburn=2000, niter=5000)

Arguments

ndose_a1           Number of dose levels for agent 1.
ndose_a2           Number of dose levels for agent 2.
target             Toxicity (probability) target.
target_min         Minimum of the targeted toxicity interval.
target_max         Maximum of the targeted toxicity interval.
prior_tox_a1       A vector of initial guesses of toxicity probabilities associated with the doses of agent 1. Must be of length ndose_a1.
prior_tox_a2       A vector of initial guesses of toxicity probabilities associated with the doses of agent 2. Must be of length ndose_a2.
cohort             Cohort size.
final              A boolean with value TRUE if the trial is finished and the recommended combination for further phases should be given, or FALSE (default value) if the combination determination is performed for the next cohort of patients.
pat_incl           Current number of patients included.
dose_adm1          A vector indicating the dose levels of agents 1 administered to each patient included in the trial. Must be of length pat_incl.
dose_adm2          A vector indicating the dose levels of agents 2 administered to each patient included in the trial. Must be of length pat_incl.
tite               A boolean indicating if the toxicity is considered as a time-to-event outcome (TRUE), or as a binary outcome (default value FALSE).
toxicity           A vector of observed toxicities (DLTs) for each patient included in the trial. Must be of length pat_incl. This argument is used/required only if tite=FALSE.
time_full          Full follow-up time window. This argument is used only if tite=TRUE.
time_tox  A vector of times-to-toxicity for each patient included in the trial. If no toxicity was observed for a patient, must be filled with +Inf. Must be of length pat_incl. This argument is used/required only if tite=TRUE.

time_follow  A vector of follow-up times for each patient included in the trial. Must be of length pat_incl. This argument is used/required only if tite=TRUE.

c_e  Probability threshold for dose-escalation. The default value is set at 0.85.

c_d  Probability threshold for dose-deescalation. The default value is set at 0.45.

c_stop  Probability threshold for early trial termination. The default value is set at 0.95.

c_t  Probability threshold for early trial termination for finding the MTD (see details). The default value is set at 0.5.

c_over  Probability threshold to control over-dosing (see details).

cmin_overunder  Minimum number of cohorts to be included at the lowest/highest combination before possible early trial termination for over-toxicity or under-toxicity (see details). The default value is set at 2.

cmin_mtd  Minimum number of cohorts to be included at the recommended combination before possible early trial termination for finding the MTD (see details). The default value is set at 3.

cmin_recom  Minimum number of cohorts to be included at the recommended combination at the end of the trial. The default value is set at 1.

alloc_rule  Integer (1, 2, or 3) indicating which allocation rule is used (see details). The default value is set at 1.

early_stop  Integer (1, 2, or 3) indicating which early stopping rule is used (see details). The default value is set at 1.

nburn  Number of burn-in for HMC. The default value is set at 2000.

niter  Number of iterations for HMC. The default value is set at 5000.

Details

Allocation rule:

- alloc_rule=1 (Riviere et al 2014): If P(toxicity probability at combination (i,j) < target) > c_e: among combinations in the neighborhood (-1, +1), (0, +1), (+1, 0), (+1, -1), choose the combination with a higher estimated toxicity probability than the current combination and with the estimated toxicity probability closest to target. If P(toxicity probability at combination (i,j) > target) > 1-c_d: among neighborhood (-1, -1), (-1, 0), (0, -1), (+1, -1), choose the combination with a lower estimated toxicity probability than the current combination and with the estimated toxicity probability closest to target. Otherwise, remain on the same combination.

- alloc_rule=2: Among combinations already tested and combinations in the neighborhood (-1, 0), (-1, +1), (0, +1), (+1, 0), (+1, -1), (0, -1), (-1, -1) of a combination tested, choose the combination with the highest posterior probability to be in the targeted interval [target_min, target_max] while controlling overdosing i.e. P(toxicity probability at combination (i,j) > target_max) < c_over.
• alloc_rule=3: Among combinations in the neighborhood (-1, 0), (-1, +1), (0, +1), (+1, 0),
  (+1, -1), (0, -1), (-1, -1) of the current combination, choose the combination with the highest
  posterior probability to be in the targeted interval \([\text{target}_\text{min}, \text{target}_\text{max}]\) while controlling
  overdosing i.e. \(P(\text{toxicity probability at combination (i,j)} > \text{target}_\text{max}) < \text{c}\_\text{over}\).

Early stopping for over-dosing: If the current combination is the lowest (1, 1) and at least \(c\_\text{min}\_\text{overunder}\) cohorts have been included at that combination and \(P(\text{toxicity probability at combination (i,j)} > \text{target}) >= \text{c}\_\text{stop} \) then stop the trial and do not recommend any combination.

Early stopping for under-dosing: If the current combination is the highest and at least \(c\_\text{min}\_\text{overunder}\) cohorts have been included at that combination and \(P(\text{toxicity probability at combination (i,j)} < \text{target}) >= \text{c}\_\text{stop} \) then stop the trial and do not recommend any combination.

Early stopping for identifying the MTD:

• early_stop=1 (Riviere et al 2014): No stopping rule, include patients until maximum sample
  size is reached.
• early_stop=2: If the next recommended combination has been tested on at least \(c\_\text{min}\_\text{mtd}\) cohorts and has a posterior probability to be in the targeted interval \([\text{target}_\text{min}, \text{target}_\text{max}]\) that is \(>= \text{c}\_t\) and also control overdosing i.e. \(P(\text{toxicity probability at current combination } > \text{target}_\text{max}) < \text{c}\_\text{over}\) then stop the trial and recommend this combination.
• early_stop=3: If at least \(c\_\text{min}\_\text{mtd}\) cohorts have been included at the next recommended
  combination then stop the trial and recommend this combination.

Stopping at the maximum sample size: If the maximum sample size is reached and no stopping rule
is met, then the recommended combination is the one that was tested on at least \(c\_\text{min}\_\text{recom}\) cohorts
and with the highest posterior probability to be in the targeted interval \([\text{target}_\text{min}, \text{target}_\text{max}]\).

Value

An object of class "CombIncrease_next" is returned, consisting of determination of the next combi-
nation and estimations. Objects generated by CombIncrease_next contain at least the following
components:

\begin{itemize}
  \item \text{n\_pat\_comb} Number of patients per combination.
  \item \text{n\_tox\_comb} Number of observed toxicities per combination.
  \item \text{pi} Estimated toxicity probabilities (if the start-up ended).
  \item \text{ptox\_inf} Estimated probabilities that the toxicity probability is inferior to target (if the
  start-up ended).
  \item \text{ptox\_inf\_targ} Estimated probabilities of underdosing, i.e. to be inferior to \text{target}_\text{min} (if the
  start-up ended).
  \item \text{ptox\_targ} Estimated probabilities to be in the targeted interval \([\text{target}_\text{min},\text{target}_\text{max}]\)
  (if the start-up ended).
  \item \text{ptox\_sup\_targ} Estimated probabilities of overdosing, i.e. to be superior to \text{target}_\text{max} (if the
  start-up ended).
  \item \text{(cdose1, cdose2)} NEXT RECOMMENDED COMBINATION.
  \item \text{inconc} Boolean indicating if trial must stop for under/over dosing.
  \item \text{early\_conc} Boolean indicating if trial can be stopped earlier for finding the MTD.
\end{itemize}
CombIncrease_sim

**Author(s)**
Jacques-Henri Jourdan and Marie-Karelle Riviere-Jourdan <eldamjh@gmail.com>

**References**

**See Also**
CombIncrease_sim.

**Examples**
```r
prior_a1 = c(0.12, 0.2, 0.3, 0.4, 0.5)
prior_a2 = c(0.2, 0.3, 0.4)
toxicity1 = c(0,0,0,0,0,0,1,0,1,0,0,0,0,0,1)
dose1 = c(1,1,2,2,3,3,3,3,3,3,3,3,4,4,4)
dose2 = c(1,1,2,2,3,3,3,2,2,2,1,1,1,1,1,1)
t_tox = c(rep(+Inf,8),2.9,+Inf,4.6,+Inf,+Inf,+Inf,+Inf,+Inf,+Inf,5.2)
follow = c(rep(6,15), 4.9, 3.1, 1.3)

next1 = CombIncrease_next(ndose_a1=5, ndose_a2=3, target=0.3,
                         target_min=0.2, target_max=0.4, prior_tox_a1=prior_a1,
prior_tox_a2=prior_a2, cohort=3, final=FALSE, pat_incl=18,
dose_adm1=dose1, dose_adm2=dose2, toxicity=toxicity1, c_over=1,
cmin_overunder=3, cmin_recom=1, early_stop=1, alloc_rule=1)
next1

next2 = CombIncrease_next(ndose_a1=5, ndose_a2=3, target=0.3,
                         target_min=0.2, target_max=0.4, prior_tox_a1=prior_a1,
prior_tox_a2=prior_a2, cohort=3, final=FALSE, pat_incl=18, dose_adm1=dose1,
dose_adm2=dose2, tite=TRUE, time_full=6, time_tox=t_tox,
time_follow=follow, c_over=1, cmin_overunder=3, cmin_recom=1,
early_stop=1, alloc_rule=1)
next2
```

---

CombIncrease_sim  
**Combination design Simulator using Logistic model**

**Description**
CombIncrease_sim is used to generate simulation replicates of phase I clinical trial for combination studies where the toxicity and efficacy of both agents is assumed to increase with the dose using the design proposed by Riviere et al. entitled "A Bayesian dose-finding design for drug combination clinical trials based on the logistic model".
**CombIncrease_sim**

**Usage**

```r
CombIncrease_sim(ndose_a1, ndose_a2, p_tox, target, target_min, target_max,
    prior_tox_a1, prior_tox_a2, n_cohort, cohort, tite=FALSE, time_full=0,
    poisson_rate=0, nsim, c_e=0.85, c_d=0.45, c_stop=0.95, c_t=0.5, c_over=0.25,
    cmin_overunder=2, cmin_mtd=3, cmin_recom=1, startup=1, alloc_rule=1,
    early_stop=1, init_dose_1=1, init_dose_2=1, nburn=2000, niter=5000, seed=14061991)
```

**Arguments**

- `ndose_a1`: Number of dose levels for agent 1.
- `ndose_a2`: Number of dose levels for agent 2.
- `p_tox`: A matrix of the true toxicity probabilities associated with the combinations. True toxicity probabilities should be entered with agent 1 in row and agent 2 in column, with increasing toxicity probabilities with both row and column numbers (see examples).
- `target`: Toxicity (probability) target.
- `target_min`: Minimum of the targeted toxicity interval.
- `target_max`: Maximum of the targeted toxicity interval.
- `prior_tox_a1`: A vector of initial guesses of toxicity probabilities associated with the doses of agent 1. Must be of length `ndose_a1`.
- `prior_tox_a2`: A vector of initial guesses of toxicity probabilities associated with the doses of agent 2. Must be of length `ndose_a2`.
- `n_cohort`: Total number of cohorts to include in the trial.
- `cohort`: Cohort size.
- `tite`: A boolean indicating if the toxicity is considered as a time-to-event outcome (TRUE), or as a binary outcome (default value FALSE).
- `time_full`: Full follow-up time window. This argument is used only if tite=TRUE.
- `poisson_rate`: A value indicating the rate for the Poisson process used to simulate patient arrival, i.e. expected number of arrivals per observation window. This argument is used only if tite=TRUE.
- `nsim`: Number of simulations.
- `c_e`: Probability threshold for dose-escalation. The default value is set at 0.85.
- `c_d`: Probability threshold for dose-deescalation. The default value is set at 0.45.
- `c_stop`: Probability threshold for early trial termination due to over-toxicity or under-toxicity (see details). The default value is set at 0.95.
- `c_t`: Probability threshold for early trial termination for finding the MTD (see details). The default value is set at 0.5.
- `c_over`: Probability threshold to control over-dosing (see details).
- `cmin_overunder`: Minimum number of cohorts to be included at the lowest/highest combination before possible early trial termination for over-toxicity or under-toxicity (see details). The default value is set at 2.
cmin_mtd  Minimum number of cohorts to be included at the recommended combination before possible early trial termination for finding the MTD (see details). The default value is set at 3.

cmin_recom Minimum number of cohorts to be included at the recommended combination at the end of the trial. The default value is set at 1.

startup  Interger (0, 1, 2, or 3) indicating which start-up phase is used (see details). The default value is set at 1.

alloc_rule Interger (1, 2, or 3) indicating which allocation rule is used (see details). The default value is set at 1.

early_stop Interger (1, 2, or 3) indicating which early stopping rule is used (see details). The default value is set at 1.

init_dose_1  Initial dose for agent 1. The default is 1.

init_dose_2  Initial dose for agent 2. The default is 1.

nburn  Number of burn-in for HMC. The default value is set at 2000.

niter  Number of iterations for HMC. The default value is set at 5000.

seed  Seed of the random number generator. Default value is set at 14061991.

Details

Start-up phase:

- startup=0: No startup phase: the first tested combination is forced to be the initial combination. The following ones use the normal allocation rule.
- startup=1 (Riviere et al 2014): Begin at the initial combination and increase both agent (+1, +1) until the first toxicity is observed or maximum combination is reached.
- startup=2: Begin at the initial combination and increase agent 1 (+1, 0) until a toxicity is observed or maximum dose is reached. Then begin at (init_dose1,init_dose2+1) and increase agent 2 (0, +1) until a toxicity is observed or maximum dose is reached.
- startup=3: Begin at the initial combination and increase alternatively each agent (+1, 0) then (0, +1) until the first toxicity is observed or maximum combination is reached.

Allocation rule:

- alloc_rule=1 (Riviere et al 2014): If P(toxicity probability at combination (i,j) < target) > c_e: among combinations in the neighborhood (-1, +1), (0, +1), (+1, 0), (+1, -1), choose the combination with a higher estimated toxicity probability than the current combination and with the estimated toxicity probability closest to target. If P(toxicity probability at combination (i,j) > target) > 1-c_d: among neighborhood (-1, +1), (-1, 0), (0, -1), (+1, -1), choose the combination with a lower estimated toxicity probability than the current combination and with the estimated toxicity probability closest to target. Otherwise, remain on the same combination.
- alloc_rule=2: Among combinations already tested and combinations in the neighborhood (-1, 0), (-1, +1), (0, +1), (+1, 0), (+1, -1), (0, -1), (-1, -1) of a combination tested, choose the combination with the highest posterior probability to be in the targeted interval [target_min, target_max] while controlling overdosing i.e. P(toxicity probability at combination (i,j) > target_max) < c_over.
• alloc_rule=3: Among combinations in the neighborhood (-1, 0), (-1, +1), (0, +1), (+1, 0),
(+1, -1), (0, -1), (-1, -1) of the current combination, choose the combination with the highest
posterior probability to be in the targeted interval \([\text{target}_{\text{min}}, \text{target}_{\text{max}}]\) while controlling
overdosing i.e. \(P(\text{toxicity probability at combination (i,j) > target}_{\text{max}}) < \text{c}_{\text{over}}\).

Early stopping for over-dosing: If the current combination is the lowest (1, 1) and at least \(c_{\text{min}}_{\text{over}}\) cohorts have been included at that combination and \(P(\text{toxicity probability at combination (i,j) > target}) \geq c_{\text{stop}}\) then stop the trial and do not recommend any combination.

Early stopping for under-dosing: If the current combination is the highest and at least \(c_{\text{min}}_{\text{under}}\) cohorts have been included at that combination and \(P(\text{toxicity probability at combination (i,j) < target}) \geq c_{\text{stop}}\) then stop the trial and do not recommend any combination.

Early stopping for identifying the MTD:

• early_stop=1 (Riviere et al 2014): No stopping rule, include patients until maximum sample
size is reached.

• early_stop=2: If the next recommended combination has been tested on at least \(c_{\text{min}}_{\text{mtd}}\) cohorts and has a posterior probability to be in the targeted interval \([\text{target}_{\text{min}}, \text{target}_{\text{max}}]\) that is \(\geq c_{\text{t}}\) and also control overdosing i.e. \(P(\text{toxicity probability at current combination > target}_{\text{max}}) < c_{\text{over}}\) then stop the trial and recommend this combination.

• early_stop=3: If at least \(c_{\text{min}}_{\text{mtd}}\) cohorts have been included at the next recommended
combination then stop the trial and recommend this combination.

Stopping at the maximum sample size: If the maximum sample size is reached and no stopping rule
is met, then the recommended combination is the one that was tested on at least \(c_{\text{min}}_{\text{recom}}\) cohorts
and with the highest posterior probability to be in the targeted interval \([\text{target}_{\text{min}}, \text{target}_{\text{max}}]\).

Value

An object of class "CombIncrease_sim" is returned, consisting of the operating characteristics of
the design specified. Objects generated by CombIncrease_sim contain at least the following components:

<table>
<thead>
<tr>
<th>Component</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>rec_dose</td>
<td>Percentage of combination selection.</td>
</tr>
<tr>
<td>n_pat_dose</td>
<td>Mean number of patients at each combination.</td>
</tr>
<tr>
<td>n_tox_dose</td>
<td>Mean number of toxicities at each combination.</td>
</tr>
<tr>
<td>inconc</td>
<td>Percentage of inclusive trials.</td>
</tr>
<tr>
<td>early_conc</td>
<td>Percentage of trials stopping with criterion for finding MTD.</td>
</tr>
<tr>
<td>nsim</td>
<td>Number of simulations (if function stopped while executed, return the current number of simulations performed with associated other outputs).</td>
</tr>
<tr>
<td>pat_tot</td>
<td>Total mean number of patients accrued.</td>
</tr>
<tr>
<td>tab_pat</td>
<td>Vector with the number of patients included for each simulation.</td>
</tr>
</tbody>
</table>

Author(s)

Jacques-Henri Jourdan and Marie-Karelle Riviere-Jourdan <eldamjh@gmail.com>
References

See Also
CombIncrease_next.

Examples

```r
p_tox_sc1 = matrix(c(0.05,0.10,0.15,0.30,0.45,
                      0.10,0.15,0.30,0.45,0.55,
                      0.15,0.30,0.45,0.50,0.60),nrow=5,ncol=3)
prior_a1 = c(0.12, 0.2, 0.3, 0.4, 0.5)
prior_a2 = c(0.2, 0.3, 0.4)

sim1 = CombIncrease_sim(ndose_a1=5, ndose_a2=3, p_tox=p_tox_sc1, target=0.30,
                          target_min=0.20, target_max=0.40, prior_tox_a1=prior_a1,
                          prior_tox_a2=prior_a2, n_cohort=20, cohort=3, tite=FALSE, nsim=2000,
                          c_over=1, cmin_overunder=3, cmin_recom=1, startup=1, alloc_rule=1,
                          early_stop=1, seed=14061991)
sim1

# Dummy example, running quickly
useless = CombIncrease_sim(ndose_a1=3, ndose_a2=2,
                           p_tox=matrix(c(0.05,0.15,0.30,0.15,0.30,0.45),nrow=3), target=0.30,
                           target_min=0.20, target_max=0.40, prior_tox_a1=c(0.2,0.3,0.4),
                           prior_tox_a2=c(0.2,0.3,0.4), n_cohort=2, cohort=2, nsim=1)
```
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