## Package 'RLumCarlo'

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```
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Description A Collection of Functions to Simulate Luminescence Production in Minerals using
      Monte-Carlo methods.
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      magrittr
URL https://CRAN.R-project.org/package=RLumCarlo
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```

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plot\_RLumCarlo

Plot results from Monte-Carlo simulations with RLumCarlo

## Description

Plot results from Monte-Carlo simulations with RLumCarlo

## Usage

```
plot_RLumCarlo(object, times = NULL, plot_uncertainty = "range",
    norm = FALSE, add = FALSE, ...)
```

## **Arguments**

object data.frame (required)

times numeric (optinal): Optional vector for the x-axis

plot\_uncertainty

logical (with default): Enable/disable uncertainty polygon plot

norm logical (with default): Normalise curve to the highest intensity

add logical (with default): allow overplotting of results

further arguments that can be passed to control the plot output. Currently sup-

ported are: xlab, xlim, ylim, main, lwd, type, pch, lty,col, grid, legend

## Value

This function returns a graphical output

## **Function version**

0.1.0

#### How to cite

Friedrich, J., Kreutzer, S., 2019. plot\_RLumCarlo(): Plot results from Monte-Carlo simulations with RLumCarlo. Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

#### Author(s)

Johannes Friedrich, University of Bayreuth (Germany), Sebastian Kreutzer, IRAMAT-CRP2A, Université Bordeaux Montaigne (France)

run\_MC\_CW\_IRSL\_LOC

Run Monte-Carlo simulation for CW-IRSL for localised transition

## **Description**

Runs a Monte-Carlo (MC) simulation of constant wave infrared stimulated luminesence (CW-IRSL) using the generalized one trap (GOT) model.

## Usage

```
run_MC_CW_IRSL_LOC(A, times, clusters = 10, n_filled = 100, r,
  method = "par", output = "signal", ...)
```

## **Arguments**

Α	numeric ( <b>required</b> ): The transition probability (cm <sup>3</sup> /s).
times	numeric (with default): The number of MC runs.
clusters	numeric (with default): The number of clusters.
n_filled	integer (with default): The number of electron traps that are filled at the beginning of the simulation.
r	numeric (with default): The retrapping ratio.
method	character (with default): sequential 'seq' or parallel processing 'par'
output	<pre>character (with default): output is either the 'signal' (the default) or 'remaining_e' (the remaining charges, electrons, in the trap)</pre>
	further arguments

#### **Details**

$$I_{LOC}(t) = -dn/dt = A * (n^2/(r+n))$$

## Value

This function returns an array with dimension length(times) x length(r) x clusters

#### **Function version**

0.1.0

#### How to cite

Kreutzer, S., 2019. run\_MC\_CW\_IRSL\_LOC(): Run Monte-Carlo simulation for CW-IRSL for localised transition. Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

#### Author(s)

Sebastian Kreutzer, IRAMAT-CRP2A, UMR 5060, CNRS - Université Bordeaux Montaigne (France)

#### References

Pagonis, V. and Kulp, C., 2017. Monte Carlo simulations of tunneling phenomena and nearest neighbor hopping mechanism in feldspars. Journal of Luminescence 181, 114–120. doi: 10.1016/j.jlumin.2016.09.014.

#### **Examples**

run\_MC\_CW\_IRSL\_TUN

Run Monte-Carlo simulation for CW-IRSL

## Description

Runs a Monte-Carlo (MC) simulation of constant wave infrared stimulated luminesence (CW-IRSL) using the model.

### Usage

```
run_MC_CW_IRSL_TUN(A, rho, times, clusters = 10, r = NULL, N_e = 200,
method = "seq", output = "signal", ...)
```

## **Arguments**

```
A numeric (required): The transition probability (cm^3/s).

rho numeric (required): The calculated dimesionless Charge density (normally written Rho').

times numeric (with default): The number of MC runs.

clusters numeric (with default): The number of clusters.
```

```
r numeric (with default): The retrapping ratio.

N_e numeric (with default): The number of electrons

method character (with default): sequential 'seq' or parallel processing 'par'

output character (with default): output is either the 'signal' (the default) or 'remaining_e' (the remaining charges, electrons, in the trap)

... further arguments
```

#### **Details**

####equation here please####

#### Value

This function returns a list.

#### **Function version**

0.2.0

#### How to cite

Friedrich, J., Kreutzer, S., 2019. run\_MC\_CW\_IRSL\_TUN(): Run Monte-Carlo simulation for CW-IRSL. Function version 0.2.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

## Author(s)

Johannes Friedrich, University of Bayreuth (Germany), Sebastian Kreutzer, IRAMAT-CRP2A, UMR 5060, Université Bordeaux Montaigne (France)

#### References

Pagonis, V. and Kulp, C. (2017) 'Monte Carlo simulations of tunneling phenomena and nearest neighbor hopping mechanism in feldspars', Journal of Luminescence. Elsevier, 181, pp. 114–120. doi: 10.1016/j.jlumin.2016.09.014.

run\_MC\_CW\_OSL\_DELOC

Run Monte-Carlo simulation for CW-OSL for delocalized transition

## **Description**

Runs a Monte-Carlo (MC) simulation of constant wave infrared stimulated luminesence (CW-OSL) using the one trap one recombination center (OTOR) model.

## Usage

```
run_MC_CW_OSL_DELOC(A, times, clusters = 10, N_e = 200,
    n_filled = N_e, R, method = "par", output = "signal", ...)
```

#### **Arguments**

Α	numeric (required): The transition probability (cm^3/s).
times	numeric (with default): The number of MC runs.
clusters	numeric (with default): The number of clusters.
N_e	integer (with default): The number of electrons.
n_filled	integer (with default): The number of electron traps that are filled at the beginning of the simulation.
R	numeric (with default): The retrapping ratio.
method	character (with default): sequential 'seq' or parallel processing 'par'
output	<pre>character (with default): output is either the 'signal' (the default) or 'remaining_e' (the remaining charges, electrons, in the trap)</pre>
	further arguments

## **Details**

$$I_{DELOC}(t) = -dn/dt = p(t) * (n^2/(NR + n(1-R)))$$

#### Value

This function returns an array with dimension length(times) x length(r) x clusters

## **Function version**

0.1.0

#### How to cite

Kreutzer, S., 2019. run\_MC\_CW\_OSL\_DELOC(): Run Monte-Carlo simulation for CW-OSL for delocalized transition. Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

#### Author(s)

Sebastian Kreutzer, IRAMAT-CRP2A, UMR 5060, CNRS - Université Bordeaux Montaigne (France)

## End(Not run)

#### References

Pagonis, V. and Kulp, C., 2017. Monte Carlo simulations of tunneling phenomena and nearest neighbor hopping mechanism in feldspars, Journal of Luminescence 181, 114–120. doi: 10.1016/j.jlumin.2016.09.014

```
## Example 1: Simulate CW-OSL
## Not run:
run_MC_CW_OSL_DELOC(
A = 0.12,
R = 1,
times = 0:100) %>%
  plot_RLumCarlo(legend = T)
## End(Not run)
#' @examples
## Example 2: Simulate CW-OSL DELOC with several parameter changes
## Not run:
# define your parameters
A=c(0.1,0.3,0.5,1)
times=seq(0,60,1)
s=1e12
E=1
R<-c(1e-7,1e-6,0.01,0.1) # sequence of different R values
clusters=1000 # number of Monte Carlo simulations
N_e = c(200, 500, 700, 400) # number of free electrons
n_{filled} = c(200, 500, 100, 70) # number of filled traps
method="par"
output ="signal"
col=c(1,2,3,4) # ifferent colours for the individual curves
plot_uncertainty <- c(T,F,T,F) # do you want to see the uncertainty?
add_{TF} \leftarrow c(F,rep(T, (length(R)-1)))
for (u in 1:length(R)){
results <-run_MC_CW_OSL_DELOC(A=A[u], times, clusters =clusters, N_e = N_e[u],
                       n_filled = n_filled[u], R=R[u], method = method, output = output)
plot_RLumCarlo(results,add=add_TF[u],legend = F, col=col[u], main=" your plot")
legend("topright",ncol=4,cex=0.55,title = "parameters" ,legend=c(paste0("A = ", A),
                                                    paste0("n_filled = ", n_filled),
                                                           paste0("N_e = ", N_e),
                                                  paste0("R = ", R)), text.col=col)
```

run\_MC\_ISO\_DELOC

Run Monte-Carlo simulation for ISO for delocalized transition

## **Description**

Runs a Monte-Carlo (MC) simulation of isothermally stimulated luminesence (ISO-TL or ITL) using the one trap one recombination center (OTOR) model. Delocalized refers to involvement of the conduction band.

## Usage

```
run_MC_ISO_DELOC(s, E, T = 20, times, clusters = 10, N_e = 200,
    n_filled = N_e, R, method = "par", output = "signal", ...)
```

## **Arguments**

S	numeric ( <b>required</b> ): Escape frequency of the trap (s^-1).
E	numeric (required): Thermal activation energy of the trap (eV).
Т	numeric (with default): Temperature (deg. C).
times	numeric (with default): The specified time within the simulation with the same syntax as the function seq().
clusters	numeric (with default): The number of MC runs.
N_e	integer (with default): The number of electrons.
n_filled	integer (with default): The number of electron traps that are filled at the beginning of the simulation.
R	numeric (with default): The retrapping ratio.
method	character (with default): sequential 'seq' or parallel processing 'par'
output	<pre>character (with default): output is either the 'signal' (the default) or 'remaining_e' (the remaining charges, electrons, in the trap)</pre>
	further arguments

## **Details**

$$I_{DELOC}(t) = -dn/dt = p(t)*(n^2/(NR + n(1-R)))$$

Where in the function  $n := n_filled := N := N_e$ 

## Value

This function returns an array with dimension length(times) x length(r) x clusters

## **Function version**

0.0.1

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#### How to cite

Kreutzer, S., 2019. run\_MC\_ISO\_DELOC(): Run Monte-Carlo simulation for ISO for delocalized transition. Function version 0.0.1. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

#### Author(s)

Sebastian Kreutzer, IRAMAT-CRP2A, UMR 5060, CNRS - Université Bordeaux Montaigne (France)

#### References

Pagonis, V., Friedrich, J., Discher, M., Müller-Kirschbaum, A., Schlosser, V., Kreutzer, S., Chen, R. and Schmidt, C., 2019. Excited state luminescence signals from a random distribution of defects: A new Monte Carlo simulation approach for feldspar. Journal of Luminescence 207, 266–272. doi: 10.1016/j.jlumin.2018.11.024

Reuven, C. and S. Mckeever, 1997. Theory of thermoluminescence and related phenomena.

## **Examples**

```
##=========##
## Example 1: Simulate ITL
##========##

## Not run:
run_MC_ISO_DELOC(
    s = 3.5e12,
    E = 1.45,
    T = 200,
    R = 1,
    times = 0:10000) %>%
    plot_RLumCarlo(legend = T)

## End(Not run)
```

run\_MC\_ISO\_LOC

Run Monte-Carlo simulation for ITL for localised transition

## Description

Runs a Monte-Carlo (MC) simulation of isothermally stimulated luminesence (ISO-TL or ITL) using the genralized one trap (GOT) model. Localized refers to excitation of an electron before it recombines, but without the involvement of the conduction band.

## Usage

```
run_MC_ISO_LOC(s, E, T = 20, times, clusters = 10, n_filled = 100, r,
  method = "par", output = "signal", ...)
```

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## **Arguments**

S	numeric ( <b>required</b> ): Escape frequency of the trap (s^-1).
Е	numeric (required): Thermal activation energy of the trap (eV).
Т	numeric (with default): Temperature (deg. C).
times	numeric (with default): The specified time within the simulation with the same syntax as the function seq().
clusters	numeric (with default): The number of MC runs.
n_filled	integer (with default): The number of electron traps that are filled at the beginning of the simulation.
r	numeric (with default): The retrapping ratio for localized models (dimensionless).
method	character (with default): sequential 'seq' or parallel processing 'par'
output	<pre>character (with default): output is either the 'signal' (the default) or 'remaining_e' (the remaining charges, electrons, in the trap)</pre>
	further arguments

#### **Details**

$$I_{LOC}(t) = -dn/dt = p(t) * (n^2/(r+n))$$

Where in the function  $n := n_filled := N := N_e$ 

#### Value

This function returns an array with dimension length(times) x length(r) x clusters

## **Function version**

0.0.1

## How to cite

Kreutzer, S., 2019. run\_MC\_ISO\_LOC(): Run Monte-Carlo simulation for ITL for localised transition. Function version 0.0.1. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

#### Author(s)

Sebastian Kreutzer, IRAMAT-CRP2A, UMR 5060, CNRS - Université Bordeaux Montaigne (France)

#### References

Pagonis, V., Friedrich, J., Discher, M., Müller-Kirschbaum, A., Schlosser, V., Kreutzer, S., Chen, R. and Schmidt, C., 2019. Excited state luminescence signals from a random distribution of defects: A new Monte Carlo simulation approach for feldspar. Journal of Luminescence 207, 266–272. doi: 10.1016/j.jlumin.2018.11.024

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#### **Examples**

run\_MC\_ISO\_TUN

Run Monte-Carlo Simulation for Isothermal Measurements for Tunneling Transition

## Description

Runs a Monte-Carlo (MC) simulation of isothermally stimulated luminesence (ISO-TL or ITL) using the tunneling (TUN) model. Tunneling refers to the movement of electrons from a trap directly to the recombination center.

#### Usage

```
run_MC_ISO_TUN(E, s, T = 200, rho, times, clusters = 10, r = NULL, N_e = 200, method = "par", output = "signal", ...)
```

## **Arguments**

E	numeric (required): Thermal activation energy of the trap (eV).
S	numeric ( <b>required</b> ): Escape frequency of the trap (s^-1).
T	numeric (required): Temperature (deg. C).
rho	numeric ( <b>required</b> ): The calculated dimesionless charge density (normally written Rho') (also defined as $(4 * pi rho/3)^1/3*r$ ) where rho := the density of recombination centers within the material given in m^3 and r is _).
times	numeric (with default): The specified time within the simulation with the same syntax as the function seq().
clusters	numeric (with default): The number of MC runs.
r	numeric (with default): The radius of tunneling (dimensionless).

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N_e	numeric (with default): The number of electrons.
method	character (with default): sequential 'seq' or parallel processing 'par'
output	<pre>character (with default): output is either the 'signal' (the default) or 'remaining_e'   (the remaining charges, electrons, in the trap)</pre>
	further arguments

#### **Details**

$$I_{TUN}(t) = -dn/dt = A * (n^2/(r+n))$$

Where in the function  $n := n_filled := N := N_e := rho := rho'$  :=  $code\{r_c\} := code\{rho'_c\}$ 

#### Value

This function returns a list.

#### **Function version**

0.1.0

#### How to cite

Friedrich, J., Kreutzer, S., 2019. run\_MC\_ISO\_TUN(): Run Monte-Carlo Simulation for Isothermal Measurements for Tunneling Transition. Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

## Author(s)

Johannes Friedrich, University of Bayreuth (Germany), Sebastian Kreutzer, IRAMAT-CRP2A, UMR 5060, CNRS - Université Bordeaux Montaigne (France)

#### References

Pagonis, V. and Kulp, C., 2017. Monte Carlo simulations of tunneling phenomena and nearest neighbor hopping mechanism in feldspars. Journal of Luminescence 181, 114–120. doi: 10.1016/j.jlumin.2016.09.014

Pagonis, V., Friedrich, J., Discher, M., Müller-Kirschbaum, A., Schlosser, V., Kreutzer, S., Chen, R. and Schmidt, C., 2019. Excited state luminescence signals from a random distribution of defects: A new Monte Carlo simulation approach for feldspar. Journal of Luminescence 207, 266–272. doi: 10.1016/j.jlumin.2018.11.024

For a discussion of tunneling see: Aitken, M.J., 1985. Thermoluminescence dating. 276-280. doi: 10.1002/gea.3340020110

```
s = 1e10,
T = 200,
rho = 0.007,
times = 0:5000) %>%
  plot_RLumCarlo(legend = TRUE)
## End(Not run)
```

run\_MC\_LM\_OSL\_DELOC

Run Monte-Carlo simulation for LM-OSL for delocalized transition

## **Description**

Runs a Monte-Carlo (MC) simulation of linearly modulated optically stimulated luminesence (LM-OSL) using the one trap one recombination center (OTOR) model.

## Usage

```
run_MC_LM_OSL_DELOC(A, times, clusters = 10, N_e = 200,
    n_filled = N_e, R, method = "par", output = "signal", ...)
```

## **Arguments**

A	numeric ( <b>required</b> ): The transition probability (cm <sup>3</sup> /s).
times	numeric (with default): The number of MC runs.
clusters	numeric (with default): The number of clusters.
N_e	integer (with default): The number of electrons.
n_filled	integer (with default): The number of electron traps that are filled at the beginning of the simulation.
R	numeric (with default): The retrapping ratio.
method	character (with default): sequential 'seq' or parallel processing 'par'
output	<pre>character (with default): output is either the 'signal' (the default) or 'remaining_e'   (the remaining charges, electrons, in the trap)</pre>
	further arguments

## Details

$$I_{DELOC}(t) = -dn/dt = p(t) * (n^2/(NR + n(1 - R)))$$

## Value

This function returns an array with dimension length(times) x length(r) x clusters

## **Function version**

0.1.0

#### How to cite

Kreutzer, S., 2019. run\_MC\_LM\_OSL\_DELOC(): Run Monte-Carlo simulation for LM-OSL for delocalized transition. Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

#### Author(s)

Sebastian Kreutzer, IRAMAT-CRP2A, UMR 5060, CNRS - Université Bordeaux Montaigne (France)

#### References

Pagonis, V. and Kulp, C. (2017) 'Monte Carlo simulations of tunneling phenomena and nearest neighbor hopping mechanism in feldspars', Journal of Luminescence. Elsevier, 181, pp. 114–120. doi: 10.1016/j.jlumin.2016.09.014.

## **Examples**

run\_MC\_LM\_OSL\_LOC

Run Monte-Carlo simulation for LM-OSL for localized transition

## Description

Runs a Monte-Carlo (MC) simulation of linearly modulated optically stimulated luminesence (LM-OSL) using the generalized one trap (GOT) model.

## Usage

```
run_MC_LM_OSL_LOC(A, times, clusters = 10, n_filled = 100, r,
  method = "par", output = "signal", ...)
```

## **Arguments**

A numeric (**required**): The transition probability (cm^3/s).

times numeric (with default): The number of MC runs.

clusters numeric (with default): The number of clusters.

n\_filled integer (with default): The number of electron traps that are filled at the beginning of the simulation.

```
r numeric (with default):

method character (with default): sequential 'seq' or parallel processing 'par'

output character (with default): output is either the 'signal' (the default) or 'remaining_e'

(the remaining charges, electrons, in the trap)

... further arguments
```

#### **Details**

$$I_{DELOC}(t) = -dn/dt = p(t) * (n^2/(NR + n(1 - R)))$$

#### Value

This function returns an array with dimension length(times) x length(r) x clusters

#### **Function version**

0.0.1

#### How to cite

Kreutzer, S., 2019. run\_MC\_LM\_OSL\_LOC(): Run Monte-Carlo simulation for LM-OSL for localized transition. Function version 0.0.1. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

## Author(s)

Sebastian Kreutzer, IRAMAT-CRP2A, UMR 5060, CNRS - Université Bordeaux Montaigne (France)

#### References

Pagonis, V. and Kulp, C. (2017) 'Monte Carlo simulations of tunneling phenomena and nearest neighbor hopping mechanism in feldspars', Journal of Luminescence. Elsevier, 181, pp. 114–120. doi: 10.1016/j.jlumin.2016.09.014.

```
##===========##
## Example 1: Simulate LM-OSL
##=============##
## Not run:
run_MC_LM_OSL_LOC(
    A = 0.12,
    r = 1,
    times = 0:100) %>%
        plot_RLumCarlo(legend = T)
## End(Not run)
```

run\_MC\_LM\_OSL\_TUN

Run Monte-Carlo simulation for LM-OSL

## Description

Run Monte-Carlo simulation for LM-OSL

#### Usage

```
run_MC_LM_OSL_TUN(A, rho, times, clusters = 10, r = NULL,
  delta.r = 0.1, N_e = 200, method = "par", output = "signal", ...)
```

## **Arguments**

Α	numeric ( <b>required</b> ): The transition probaility (cm^3/s).
rho	numeric (required): The calculated dimesionless Charge density (normally written Rho').
times	vector (with default): The number of MC runs.
clusters	numeric (with default): The number of clusters.
r	numeric (with default): The retrapping ratio.
delta.r	numeric (with default):
N_e	numeric (with default): The number of electrons.
method	character (with default): sequential 'seq' or parallel processing 'par'
output	<pre>character (with default): output is either the 'signal' (the default) or 'remaining_e' (the remaining charges, electrons, in the trap)</pre>
	further arguments

## **Details**

ADD EQUATION

## Value

This function returns a list.

## **Function version**

0.1.0

## How to cite

Friedrich, J., 2019. run\_MC\_LM\_OSL\_TUN(): Run Monte-Carlo simulation for LM-OSL. Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

## Author(s)

Johannes Friedrich, University of Bayreuth (Germany)

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#### References

Pagonis, V. and Kulp, C. (2017) 'Monte Carlo simulations of tunneling phenomena and nearest neighbor hopping mechanism in feldspars', Journal of Luminescence. Elsevier, 181, pp. 114-120. doi: 10.1016/j.jlumin.2016.09.014.

## **Examples**

```
## Not run:
##TODO: Primary example, should be verified
run_MC_LM_OSL_TUN(A = 10000, rho = 0.0001, times = 1:100, clusters = 10, r = NULL,
delta.r = 0.1,
N_e = 200, method = "par", output = "signal") %>%
plot_RLumCarlo(norm = T)
## End(Not run)
```

run\_MC\_TL\_DELOC

Run Monte-Carlo simulation for TL for delocalized transition

## **Description**

Runs a Monte-Carlo (MC) simulation of thermo-luminesence (TL) using the one trap one recombination center (OTOR) model. Delocalized refers to involvement of the conduction band.

## Usage

```
run_MC_TL_DELOC(s, E, times, clusters = 10, N_e = 200,
 n_filled = N_e, R, method = "par", output = "signal", ...)
```

## **Arguments**

S	numeric ( <b>required</b> ): Escape frequency of the trap (s^-1).
E	numeric (required): Thermal activation energy of the trap (eV).
times	numeric (with default): The specified time within the simulation with the same syntax as the function seq().
clusters	numeric (with default): The number of MC runs.
N_e	integer (with default): The number of electrons.
n_filled	integer (with default): The number of electron traps that are filled at the beginning of the simulation.
R	numeric (with default): The retrapping ratio.
method	character (with default): sequential 'seq' or parallel processing 'par'
output	<pre>character (with default): output is either the 'signal' (the default) or 'remaining_e'   (the remaining charges, electrons, in the trap)</pre>
	further arguments

run\_MC\_TL\_DELOC

#### **Details**

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$$I_{DELOC}(t) = -dn/dt = p(t) * (n^2/(NR + n(1 - R)))$$

where in the function N := N e := n := n filled

#### Value

This function returns an array with dimension length(times) x length(r) x clusters

#### **Function version**

0.0.1

#### How to cite

Kreutzer, S., 2019. run\_MC\_TL\_DELOC(): Run Monte-Carlo simulation for TL for delocalized transition. Function version 0.0.1. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

#### Author(s)

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#### References

Pagonis, V., Friedrich, J., Discher, M., Müller-Kirschbaum, A., Schlosser, V., Kreutzer, S., Chen, R. and Schmidt, C., 2019. Excited state luminescence signals from a random distribution of defects: A new Monte Carlo simulation approach for feldspar. Journal of Luminescence 207, 266–272. doi: 10.1016/j.jlumin.2018.11.024

Reuven, C. and S. Mckeever, 1997. Theory of thermoluminescence and related phenomena.

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```
## Not run:
# define your parameters
times=seq(100,450,1)
s=rep(3.5e12,4)
E=rep(1.45,4)
R<-c(0.7e-6,1e-6,0.01,0.1)
clusters=1000
N_e = c(400, 500, 700, 400)
n_filled =c(400, 500, 300, 70)
method="par"
output ="signal"
col=c(1,2,3,4) # different colours for the individual curves
plot_uncertainty <- c(TRUE,TRUE,TRUE,TRUE) # do you want to see the uncertainty?</pre>
add_TF <- c(FALSE,rep(TRUE, (length(R)-1)))</pre>
for (u in 1:length(R)){
results <-run_MC_TL_DELOC(times=times, s=s[u], E=E[u], clusters =clusters, N_e = N_e[u],
                         n_filled = n_filled[u], R=R[u], method = method, output = output)
plot_RLumCarlo(results,add=add_TF[u],legend = FALSE, col=col[u], main=" your plot", ylim=c(0,20))
legend("topright",ncol=5,cex=0.55,title = "parameters" ,legend=c(paste0("E = ", E),
                                                           paste0("s = ", s),
paste0("n_filled = ", n_filled),
                                                                   paste0("N_e = ", N_e),
                                                         paste0("R = ", R)), text.col=col)
## End(Not run)
```

run\_MC\_TL\_LOC

Run Monte-Carlo simulation for TL for localised transition

## Description

Runs a Monte-Carlo (MC) simulation of thermo-luminesence (TL) using the generalized one trap (GOT) model. Localized refers to excitation of an electron before it recombines, but without the involvement of the conduction band.

## Usage

```
run_MC_TL_LOC(s, E, times, clusters = 10, n_filled = 100, r,
  method = "par", output = "signal", ...)
```

#### **Arguments**

S	numeric ( <b>required</b> ): Escape frequency of the trap ( $s^{-1}$ ).
Е	numeric (required): Thermal activation energy of the trap (eV).
times	numeric (with default): The specified time within the simulation with the same syntax as the function seq().
clusters	numeric (with default): The number of MC runs.
n_filled	integer (with default): The number of electron traps that are filled at the beginning of the simulation.

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```
r numeric (with default): The retrapping ratio.

method character (with default): sequential 'seq' or parallel processing 'par'

output character (with default): output is either the 'signal' (the default) or 'remaining_e'

(the remaining charges, electrons, in the trap)

... further arguments
```

#### **Details**

$$I_{LOC}(t) = -dn/dt = p(t) * (n^2/(r+n))$$

where in the function  $n := n_filled$ 

#### Value

This function returns an array with dimension length(times) x length(r) x clusters

#### **Function version**

0.1.0

#### How to cite

Kreutzer, S., 2019. run\_MC\_TL\_LOC(): Run Monte-Carlo simulation for TL for localised transition. Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

## Author(s)

Sebastian Kreutzer, IRAMAT-CRP2A, UMR 5060, CNRS - Université Bordeaux Montaigne (France)

#### References

Pagonis, V., Friedrich, J., Discher, M., Müller-Kirschbaum, A., Schlosser, V., Kreutzer, S., Chen, R. and Schmidt, C., 2019. Excited state luminescence signals from a random distribution of defects: A new Monte Carlo simulation approach for feldspar. Journal of Luminescence 207, 266–272. doi: 10.1016/j.jlumin.2018.11.024

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## End(Not run)

run\_MC\_TL\_TUN

Run Monte-Carlo Simulation for TL using Tunnelling Transition

## Description

Runs a Monte-Carlo (MC) simulation of thermo-luminesence (TL) using the tunneling (TUN) model. Tunneling transitions refers to the direct movement of electrons from a trap directly to the recombination centre.

## Usage

```
run_MC_TL_TUN(s, E, rho, r_c, times, clusters = 10, N_e = 200,
  delta.r = 0.1, method = "par", output = "signal", ...)
```

## Arguments

S	list ( <b>required</b> ): Escape frequency of the trap (s^-1).
E	numeric (required): Thermal activation energy of the trap (eV).
rho	numeric (required): The calculated dimesionless Charge density.
r_c	numeric (with default): The dimensionless minimal critical radius.
times	vector (with default): The number of MC runs.
clusters	numeric (with default): The number of clusters.
N_e	numeric (with default): The number of electrons
delta.r	numeric (with default): The approriate distance interval along the r axis (dimensionless).
method	character (with default): sequential 'seq' or parallel processing 'par'
output	<pre>character (with default): output is either the 'signal' (the default) or 'remaining_e' (the remaining charges, electrons, in the trap)</pre>
	further arguments

## **Details**

$$I_{TUN}(t) = -dn/dt = A * (n^2/(r+n))$$

where in the function  $N := N_e := rho' \} := \code{r_c} := \code{rho'_c}$ 

## Value

This function returns an array with dimension length(times) x length(r) x clusters

## **Function version**

0.1.0

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#### How to cite

Friedrich, J., Kreutzer, S., 2019. run\_MC\_TL\_TUN(): Run Monte-Carlo Simulation for TL using Tunnelling Transition. Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence PhenomenaR package version 0.1.0.9000-79.

#### Author(s)

Johannes Friedrich, University of Bayreuth (Germany), Sebastian Kreutzer, IRAMAT-CRP2A, UMR 5060, Université Bordeaux Montaigne (France)

#### References

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