# Package 'RLumCarlo'

November 27, 2019

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Type Package
Title Monte-Carlo Methods for Simulating Luminescence Phenomena
Version 0.1.0.9000-133
Date 2019-11-27
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Description A collection of functions to simulate luminescence production in dosimetric materials us-
       ing Monte-Carlo methods.
       Implemented are models for delocalised, localised and tunnelling transitions.
       Supported stimulation methods are TL, CW-OSL, LM-OSL, LM-IRSL, and ITL (ISO-TL).
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License GPL-3
BugReports https://github.com/R-Lum/RLumCarlo/issues
Depends R (>= 3.3.0),
       utils,
       magrittr
URL https://CRAN.R-project.org/package=RLumCarlo
LinkingTo Rcpp (>= 1.0.2),
       RcppArmadillo (>= 0.9.700.2.0)
Imports abind (>= 1.4-5),
       doParallel (>= 1.0.15),
       foreach (>= 1.4.7),
       khroma (>= 1.2.0),
       parallel,
       methods,
       Rcpp (>= 1.0.2)
Suggests R.rsp (>= 0.43.1),
       testthat (>= 2.0.0)
Encoding UTF-8
```

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## VignetteBuilder R.rsp RoxygenNote 7.0.1

## **R** topics documented:

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### **Description**

A collection of functions to simulate luminescence production in dosimetric materials using Monte-Carlo methods. Implemented are models for delocalised, localised and tunnelling transitions. Supported stimulation modes are TL, CW-OSL, LM-OSL, LM-IRSL, and ITL (ISO-TL).

## **Details**

## **Funding**

The development of RLumCarlo benefitted from the support by various funding bodies:

- The initial work by Johannes Friedrich, Sebastian Kreutzer and Christoph Schmidt was supported by the Deutsche Forschungsgemeinschaft (DFG, 2015–2018, SCHM 3051/4-1, "Modelling quartz luminescence signal dynamics relevant for dating and dosimetry", SCHM 3051/4-1).
- Later work (2018-2019) was secured through the project "ULTIMO: Unifying Luminescence Models of quartz and feldspar DAAD: Deutscher Akademischer Austauschdienst (German Academic Exchange Service). Framework: DAAD PPP USA 2018, ID: 57387041.
- The work of Sebastian Kreutzer as maintainer of the package was supported by LabEx LaS-cArBx (ANR n. ANR-10-LABX-52) between 2017 and 2019.

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#### Author(s)

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Christoph Schmidt, University of Bayreuth (Germany),
Ena Rajovic, University of Bayreuth (Germany),
Alex Roy Duncan, University of Bayreuth (Germany),
Christian Laag, University of Bayreuth (Germany)

#### References

Pagonis, V., Friedrich, J., Discher, M., Müller-Kirschbaum, A., Schlosser, V., Kreutzer, S., Chen, R., 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

plot\_RLumCarlo

Plot RLumCarlo Monte-Carlo Simulation Results

### **Description**

Visualise 'RLumCarlo' modelling results without extracting the values manually. Typically visualised values are the averaged signal or the number of remaining electrons, with a polygon indicating modelling uncertainties.

#### Usage

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

## **Arguments**

object

list of class RLumCarlo\_Model\_Output (**required**): input object to be plotted, usually the required input object is generated by one of the functions starting with run\_. Alternatively a list of such objects can be provided.

plot\_uncertainty

logical (with default): type of the displayed uncertainty. Allowed values are range, sd (standard deviation) and var (variance). NULL disables the uncertainty visualisation.

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

add logical (with default): allows overplotting of results by adding curves to an ex-

isting plot. This argument is handled automatically if object is of type list

further arguments that can be passed to control the plot output largely following the argument names in graphics::plot.default. Currently supported are: xlab, ylab, xlim, ylim, main, lwd, type, pch, lty,col, grid, legend. The arguments lwd, type, pch, lty, col can be provided as a vector if object is a list

#### **Details**

For colouring the curves, the package khroma::khroma-package is used to provide colours that can be best distinguished, in particular by colour-blind users.

### Value

This function returns a graphical output

#### **Function version**

0.1.0

#### How to cite

Friedrich, J., Kreutzer, S., 2019. plot\_RLumCarlo(): Plot RLumCarlo Monte-Carlo Simulation Results. Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

#### Author(s)

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

run\_MC\_CW\_IRSL\_LOC

Monte-Carlo Simulation for CW-IRSL (localized transitions)

#### **Description**

Runs a Monte-Carlo (MC) simulation of continuous wave infrared stimulated luminesence (CW-IRSL) using the generalized one trap (GOT) model. Localized transitions refer to transitions which do not involve the conduction or valence band. These transitions take place between the ground state and an excited state of the trapped charge, and also involve an excited state of the recombination center.

## Usage

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

#### **Arguments**

A	numeric ( <b>required</b> ): The optical excitation rate from the ground state of the trap to the excited state ( $s^{-1}$ )
times	numeric (with default): The sequence of time steps within the simulation (s)
clusters	numeric (with default): The number of MC runs (unitless)
n_filled	integer (with default): The number of filled electron traps at the beginning of the simulation (unitless)
r	numeric (with default): The retrapping ratio for localized transitions
method	character ( <i>with default</i> ): Sequential 'seq' or parallel 'par' processing. In the parallel mode the function tries to run the simulation on multiple CPU cores (if available) with a positive effect on the computation time.
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**

## The model

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

where in the function:

A := optical excitation rate from the ground state into the excited state of the trapped charge ( $s^{-1}$ ) r := retrapping ratio for localized transitions

t := time (s)

i .= time (s)

n := number of filled electron traps

#### Value

This function returns an object of class RLumCarlo\_Model\_Output which is a list consisting of an array with dimension length(times) x clusters and a numeric time vector.

## **Function version**

0.1.0

#### How to cite

Kreutzer, S., 2019. run\_MC\_CW\_IRSL\_LOC(): Monte-Carlo Simulation for CW-IRSL (localized transitions). Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

## 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

#### **Further reading**

Chen, R., McKeever, S.W.S., 1997. Theory of Thermoluminescence and Related Phenomena. WORLD SCIENTIFIC. doi: 10.1142/2781

## **Examples**

```
run_MC_CW_IRSL_LOC(
    A = 0.12,
    times = 0:100,
    clusters = 50,
    n_filled = 100,
    r = 1e-7,
    method = "seq",
    output = "signal"
) %>%
plot_RLumCarlo(legend = TRUE)
```

run\_MC\_CW\_IRSL\_TUN

Run Monte-Carlo Simulation for CW-IRSL (tunneling transitions)

## Description

Runs a Monte-Carlo (MC) simulation of continuous wave infrared stimulated luminesence (CW-IRSL) using the model for tunneling translations. Tunneling refers to quantum mechanical tunneling processes from the excited state of the trapped charge, into a recombination center.

### Usage

```
run_MC_CW_IRSL_TUN(
    A,
    rho,
    times,
    clusters = 10,
    r_c = 0,
    delta.r = 0.1,
    N_e = 200,
    method = "seq",
    output = "signal",
    ...
)
```

#### **Arguments**

A	numeric ( <b>required</b> ): The optical excitation rate from the ground state of trap to the excited state of trap ( $s^{-1}$ ).
rho	numeric ( <b>required</b> ): The density of recombination centers (defined as $\rho$ ' in Huntley 2006) (unitless).
times	numeric (with default): The sequence of time steps within the simulation (s).
clusters	numeric (with default): The number of MC runs (unitless).
r_c	numeric (with default): Critical distance (>0) that must be provided if the sample has been thermally and/or optically pretreated. This parameter expresses the fact that electron-hole pairs within a critical radius $r_c$ have already recombined.
delta.r	numeric (with default): Increments of the unitless distance parameter r
N_e	numeric (width default): The total number of electron traps available (unitless).
method	character ( <i>with default</i> ): Sequential 'seq' or parallel 'par' processing. In the parallel mode the function tries to run the simulation on multiple CPU cores (if available) with a positive effect on the computation time.
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**

#### The model

$$I_{TUN}(r',t) = -dn/dt = A*exp(-(\rho')^{-1/3}*r')*n(r',t)$$

Where in the function:

A := excitation rate from ground state of the trap to the excited state (1/s)

r' := the unitless tunneling radius

 $\rho'$  := rho' the unitless density of recombination centres (see Huntley (2006))

t := time(s)

n := the instantaneous number of electrons

### Value

This function returns an object of class  $RLumCarlo\_Model\_Output$  which is a list consisting of an array with dimension length(times) x length(r) x clusters and a numeric time vector.

## **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 (tunneling transitions). Function version 0.2.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

#### Author(s)

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

#### References

Huntley, D.J., 2006. An explanation of the power-law decay of luminescence. Journal of Physics: Condensed Matter, 18(4), 1359. doi: 10.1088/09538984/18/4/020

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

### **Further reading**

Aitken, M.J., 1985. Thermoluminescence dating. Academic Press.

Jain, M., Guralnik, B., Andersen, M.T., 2012. Stimulated luminescence emission from localized recombination in randomly distributed defects. J. Phys.: Condens. Matter 24, 385402. doi: 10.1088/09538984/24/38/385402

Chen, R., McKeever, S.W.S., 1997. Theory of Thermoluminescence and Related Phenomena. WORLD SCIENTIFIC. doi: 10.1142/2781

#### **Examples**

```
run_MC_CW_IRSL_TUN(
    A = 0.8,
    rho = 1e-4,
    times = 0:50,
    r_c = 0.05,
    delta.r = 0.1,
    method = "seq",
    clusters = 10,
    output = "signal") %>%
    plot_RLumCarlo(norm = TRUE, legend = TRUE)
```

 $run\_MC\_CW\_OSL\_DELOC$ 

Run Monte-Carlo Simulation for CW-OSL (delocalized transitions)

## Description

Runs a Monte-Carlo (MC) simulation of continuous wave optically stimulated luminesence (CW-OSL) using the one trap one recombination center (OTOR) model. The term delocalized here refers to the involvement of the conduction band.

## Usage

```
run_MC_CW_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 optical excitation rate from trap to conduction band $(s^{-1})$
times	numeric (with default): The sequence of temperature steps within the simulation (s)
clusters	numeric (with default): The number of MC runs (unitless)
N_e	integer (with default): The total number of electron traps available (unitless)
n_filled	integer (with default): The number of filled electron traps at the beginning of the simulation (unitless)
R	numeric (with default): The retrapping ratio for delocalized transitions (unitless)
method	character (with default): Sequential 'seq' or parallel 'par' processing. In the parallel mode the function tries to run the simulation on multiple CPU cores (if available) with a positive effect on the computation time.
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**

#### The model

```
I_{DELOC}(t) = -dn/dt = A * (n^2/(N * R + n(1 - R)))
```

Where in the function:

t := time

A := the optical excitation rate from trap to conduction band

 $n := n_{filled}$ , the instantaneous number of electrons

 $N := N_e$  the available number of electron traps available

R := retrapping ratio for delocalized transitions

#### Value

This function returns an object of class RLumCarlo\_Model\_Output which is a list consisting of an array with dimension length(times) x clusters and a numeric time vector.

#### **Function version**

0.1.0

#### How to cite

Kreutzer, S., 2019. run\_MC\_CW\_OSL\_DELOC(): Run Monte-Carlo Simulation for CW-OSL (delocalized transitions). Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

#### 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

#### **Further reading**

Chen, R., McKeever, S.W.S., 1997. Theory of Thermoluminescence and Related Phenomena. WORLD SCIENTIFIC. doi: 10.1142/2781

### **Examples**

```
## brief example
run_MC_CW_OSL_DELOC(
A = 0.12,
R = 0.1,
 times = 0:10,
 clusters = 10,
 method = "seq") %>%
plot_RLumCarlo(legend = TRUE)
## A long example
## Not run:
A \leftarrow c(0.1, 0.3, 0.5, 1)
times <- seq(0, 60, 1)
s <- 1e12
E <- 1
R \leftarrow 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} \leftarrow c(200, 500, 100, 70) # number of filled traps
method <-"par"
output <- "signal"
col \leftarrow c(1,2,3,4) # ifferent colours for the individual curves
plot_uncertainty <- c(TRUE,FALSE,TRUE,FALSE) # do you want to see the uncertainty?</pre>
add_TF <- c(FALSE,rep(TRUE, (length(R)-1)))</pre>
#loop to plot different curves into one plotfor
for (u in 1:length(R)){
 results <- run_MC_CW_OSL_DELOC(</pre>
  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 = FALSE,
col = col[u],
main = "Delocalised Transition")
# add your legend with your parameters
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)),
  bty = "n",
   text.col = col)
## End(Not run)
```

run\_MC\_ISO\_DELOC

Run Monte-Carlo Simulation for ISO-TL (delocalized transitions)

### **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 (required): The frequency factor of the trap (s^-1)

E numeric (required): Thermal activation energy of the trap (eV)

T numeric (with default): Constant stimulation temperature (degress C)

times numeric (with default): The sequence of time steps within the simulation (s)
```

clusters numeric (with default): The number of MC runs (unitless)

N\_e integer (with default): The total number of electron traps available (unitless)

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

simulation (unitless)

R numeric (with default): The delocalized retrapping ratio (unitless)

method character (with default): Sequential 'seq' or parallel 'par' processing. In the

parallel mode the function tries to run the simulation on multiple CPU cores (if

available) with a positive effect on the computation time.

output character (with default): Output is either the 'signal' (the default) or 'remaining\_e'

(the remaining charges, electrons, in the trap)

... further arguments

#### **Details**

#### The model

$$I_{DELOC}(t) = -dn/dt = (s * exp(-E/(k_B * T_{ISO}))) * (n^2/(N * R + n(1 - R)))$$

Where in the function:

t := time

 $k_B := Boltzmann constant$ 

 $T_{ISO}$  = temperature of the isothermal experiment

 $n := n_{filled}$ , the number of filled electron traps at the beginning of the simulation

E := the trap depth in eV

s :=the frequency factor in  $(s^{-1})$ 

 $N := N_e$ , the total number of electron traps available (unitless)

R := the retrapping ratio for delocalized transitions

#### Value

This function returns an object of class RLumCarlo\_Model\_Output which is a list consisting of an array with dimension length(times) x clusters and a numeric time vector.

## **Function version**

0.1.0

## How to cite

Kreutzer, S., 2019. run\_MC\_ISO\_DELOC(): Run Monte-Carlo Simulation for ISO-TL (delocalized transitions). Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

#### Author(s)

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

run\_MC\_ISO\_LOC

#### 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

#### **Further reading**

Chen, R., McKeever, S.W.S., 1997. Theory of Thermoluminescence and Related Phenomena. WORLD SCIENTIFIC. doi: 10.1142/2781

#### **Examples**

```
run_MC_ISO_DELOC(
    s = 3.5e12,
    E = 1.45,
    T = 200,
    R = 1,
    method = 'seq',
    times = 0:100) %>%
plot_RLumCarlo(legend = TRUE)
```

run\_MC\_ISO\_LOC

Run Monte-Carlo simulation for ISO-TL (localized transitions)

### **Description**

Runs a Monte-Carlo (MC) simulation of isothermally stimulated luminesence (ISO-TL or ITL) using the generalized one trap (GOT) model. Localized transitions refer to transitions which do no involve the conduction or valence band. These transitions take place between the ground state and an excited state of the trapped charge, and also involve an excited state of the recombination center.

#### Usage

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

#### **Arguments**

```
s numeric (required): The frequency factor of the trap (s^-1)

E numeric (required): Thermal activation energy of the trap (eV)
```

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Т numeric (with default): Constant stimulation temperature (degrees C) times numeric (with default): The sequence of time steps within the simulation (s) numeric (with default): The number of MC runs (unitless) clusters n filled integer (with default): The number of filled electron traps at the beginning of the simulation (unitless) numeric (with default): The retrapping ratio for localized transitions. r character (with default): Sequential 'seq' or parallel 'par' processing. In the method parallel mode the function tries to run the simulation on multiple CPU cores (if available) with a positive effect on the computation time. character (with default): output is either the 'signal' (the default) or 'remaining\_e' output (the remaining charges/electrons in the trap) further arguments

#### **Details**

#### The model

$$I_{LOC}(t) = -dn/dt = (s * exp(-E/(k_B * T_{ISO}))) * (n^2/(r+n)))$$

Where in the function:

t := time(s)

 $k_B := Boltzmann constant$ 

 $T_{ISO}$  "= isothermal temperature

 $n := n_filled$ 

s := frequency factor of the trap (1/s)

E := activation energy of the trap (eV)

r := retrapping ratio for localized transitions

#### Value

This function returns an object of class RLumCarlo\_Model\_Output which is a list consisting of an array with dimension length(times) x clusters and a numeric time vector.

#### **Function version**

0.1.0

#### How to cite

Kreutzer, S., 2019. run\_MC\_ISO\_LOC(): Run Monte-Carlo simulation for ISO-TL (localized transitions). Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

## Author(s)

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

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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

### **Examples**

```
run_MC_ISO_LOC(
    E = 1.45,
    s = 3.5e12,
    T = 200,
    times = 0:100,
    method = 'seq',
    r = 1) %>%
plot_RLumCarlo(legend = TRUE)
```

run\_MC\_ISO\_TUN

Monte-Carlo Simulation for ISO-TL (tunneling transitions)

## **Description**

Runs a Monte-Carlo (MC) simulation of isothermally stimulated luminesence (ISO-TL or ITL) using the tunneling (TUN) model. Tunneling refers to quantum mechanical tunneling processes from the excited state of the trapped charge, into the recombination center.

## Usage

```
run_MC_ISO_TUN(
    E,
    s,
    T = 200,
    rho,
    times,
    clusters = 10,
    r_c = 0,
    delta.r = 0.1,
    N_e = 200L,
    method = "par",
    output = "signal",
    ...
)
```

#### **Arguments**

```
E numeric (required): Thermal activation energy of the trap (eV).

s numeric (required): The effective frequency factor for the tunneling process (s^-1).

T numeric (with default): Constant stimulation temperature (degrees C).
```

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rho	numeric ( <b>required</b> ): The dimensionless density of recombination centres (defined as $\rho$ ' in Huntley 2006) (unitless).
times	numeric (with default): The sequence of time steps within the simulation (s).
clusters	numeric (with default): The number of MC runs (unitless).
r_c	numeric ( <i>with default</i> ): Critical distance (>0) that must be provided if the sample has been thermally and/or optically pretreated. This parameter expresses the fact that electron-hole pairs within a critical radius r_c have already recombined.
delta.r	numeric (with default): Fractional change of the dimensionless distance of nearest recombination centres (r')
N_e	numeric (width default): The total number of electron traps available (unitless).
method	character ( <i>with default</i> ): Sequential 'seq' or parallel 'par' processing. In the parallel mode the function tries to run the simulation on multiple CPU cores (if available) with a positive effect on the computation time.
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**

#### The model

```
I_{TUN}(r',t) = -dn/dt = (s * exp(-E/(k_B * T_{ISO}))) * exp(-(\rho')^{-1/3} * r') * n(r',t)
```

Where in the function:

E := thermal activation energy (eV)

s := the effective frequency factor for the tunneling process ( $s^{-1}$ )

 $T_{ISO}$  := the temperature of the isothermal experiment (degrees C)

 $k_B := Boltzmann constant$ 

r' := the unitless tunneling radius

 $\rho$  := rho the unitless density of recombination centres see Huntley (2006)

t := time(s)

n := the instantaneous number of electrons

## Value

This function returns an object of class  $RLumCarlo\_Model\_Output$  which is a list consisting of an array with dimension length(times) x length(r) x clusters and a numeric time vector.

#### **Function version**

0.1.0

#### How to cite

Friedrich, J., Kreutzer, S., 2019. run\_MC\_ISO\_TUN(): Monte-Carlo Simulation for ISO-TL (tunneling transitions). Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

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#### 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

Further reading Aitken, M.J., 1985. Thermoluminescence dating. Academic Press.

Huntley, D.J., 2006. An explanation of the power-law decay of luminescence. Journal of Physics: Condensed Matter, 18(4), 1359.doi: 10.1088/09538984/18/4/020

Jain, M., Guralnik, B., Andersen, M.T., 2012. Stimulated luminescence emission from localized recombination in randomly distributed defects. J. Phys.: Condens. Matter 24, 385402. doi: 10.1088/09538984/24/38/385402

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

### **Examples**

```
## short example
run_MC_ISO_TUN(
E = .8,
s = 1e16,
T = 50,
rho = 1e-4.
 times = 0:100,
clusters = 10,
N_e = 2,
 r_c = 1e-4
 delta.r = 0.5,
method = "seq") \%>\%
plot_RLumCarlo(legend = TRUE)
## Not run:
## long (meaningful) example
results <- run_MC_ISO_TUN(</pre>
E = .8,
s = 1e16,
 T = 50,
 rho = 1e-4,
 times = 0:100,
 clusters = 1000,
N_e = 200,
 r_c = 1e-4
delta.r = 0.5,
method = "par")
plot_RLumCarlo(results, legend = TRUE)
## End(Not run)
```

run\_MC\_LM\_OSL\_DELOC

Run Monte-Carlo Simulation for LM-OSL (delocalized transitions)

### **Description**

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

## Usage

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

# **Arguments** A

A	numeric ( <b>required</b> ): The optical excitation rate from trap to conduction band $(s^{-1})$
times	numeric (with default): The sequence of time steps within the simulation (s)
clusters	numeric (with default): The number of MC runs (unitless)
N_e	integer (with default): The total number of electron traps available (unitless)
n_filled	integer (with default): The number of filled electron traps at the beginning of the simulation (unitless)
R	numeric (with default): The retrapping ratio for delocalized transitions
method	character ( <i>with default</i> ): Sequential 'seq' or parallel 'par' processing. In the parallel mode the function tries to run the simulation on multiple CPU cores (if available) with a positive effect on the computation time.
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**

## The model

```
I_{DELOC}(t) = -dn/dt = A * (n^2/(N * R + n(1 - R)))
```

Where in the function:

```
t := time(s)
```

```
A := the optical excitation rate from trap to conduction band n := n_filled, the tnstantaneous number of electrons R := the retrapping ratio for delocalized transitions N := N_e, the total number of electron traps available (unitless) P := total stimulation time"
```

#### Value

This function returns an object of class RLumCarlo\_Model\_Output which is a list consisting of an array with dimension length(times) x clusters and a numeric time vector.

#### **Function version**

0.1.0

#### How to cite

Kreutzer, S., 2019. run\_MC\_LM\_OSL\_DELOC(): Run Monte-Carlo Simulation for LM-OSL (delocalized transitions). Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

#### 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

#### **Further reading**

Chen, R., McKeever, S.W.S., 1997. Theory of Thermoluminescence and Related Phenomena. WORLD SCIENTIFIC. doi: 10.1142/2781

## Examples

```
run_MC_LM_OSL_DELOC(
    A = 0.12,
    R = 0.1,
    times = 0:50,
    method = "seq",
    clusters = 10) %>%
plot_RLumCarlo(legend = TRUE)
```

 $run\_MC\_LM\_OSL\_LOC$ 

Run Monte-Carlo Simulation for LM-OSL (localized transitions)

### **Description**

Runs a Monte-Carlo (MC) simulation of linearly modulated optically stimulated luminesence (LM-OSL) using the generalized one trap (GOT) model. Localized transitions refer to transitions which do not involve the conduction or valence band. These transitions take place between the ground state and an excited state of the trapped charge, and also involve a an excited state of the recombination center.

## Usage

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

#### **Arguments**

A	numeric ( <b>required</b> ): The optical excitation rate from trap to conduction band $(s^{-1})$
times	numeric (with default): The sequence of time steps within the simulation (s)
clusters	numeric (with default): The number of MC runs (unitless)
n_filled	integer (with default): The number of filled electron traps at the beginning of the simulation (unitless)
r	numeric (with default): The retrapping ratio for localized transitions
method	character ( <i>with default</i> ): Sequential 'seq' or parallel 'par' processing. In the parallel mode the function tries to run the simulation on multiple CPU cores (if available) with a positive effect on the computation time.
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**

#### The model

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

Where in the function:

A := optical excitation rate from the trap to the conduction band

P := total excitation time

```
\begin{split} t &:= time \ (s) \\ n &:= n\_filled, \ the \ instantaneous \ number \ of \ electrons \\ r &:= the \ retrapping \ ratio \ for \ localized \ transitions \\ P &:= the \ total \ stimulation \ period \ (s) \end{split}
```

#### Value

This function returns an object of class RLumCarlo\_Model\_Output which is a list consisting of an array with dimension length(times) x clusters and a numeric time vector.

#### **Function version**

0.1.0

#### How to cite

Kreutzer, S., 2019. run\_MC\_LM\_OSL\_LOC(): Run Monte-Carlo Simulation for LM-OSL (localized transitions). Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

#### 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

### **Examples**

```
## short example
run_MC_LM_OSL_LOC(
A = 1,
s = 1e8,
E = 0.5,
 times = 0:40,
clusters = 10,
n_filled = 10,
 r = 1e-7,
method = "seq",
output = "signal") %>%
plot_RLumCarlo(legend = TRUE)
## Not run:
## the long (meaningful) example
results <- run_MC_LM_OSL_LOC(</pre>
A = 1,
s = 1e8,
E = 0.6,
 times = 0:100,
 clusters = 100,
n_filled = 100,
```

```
r = 1e-7,
method = "par",
output = "signal")

## plot
plot_RLumCarlo(results, legend = TRUE)

## End(Not run)
```

run\_MC\_LM\_OSL\_TUN

Run Monte-Carlo Simulation for LM-OSL (tunneling transitions)

#### **Description**

Runs a Monte-Carlo (MC) simulation of linearly modulated optically stimulated luminesence (LM-OSL) using the tunneling (TUN) model. Tunneling refers to quantum mechanical tunneling processes from the excited state of the trapped charge, into a recombination center.

## Usage

```
run_MC_LM_OSL_TUN(
    A,
    rho,
    times,
    clusters = 10,
    r_c = 0,
    delta.r = 0.1,
    N_e = 200,
    method = "par",
    output = "signal",
    ...
)
```

#### **Arguments**

numeric (required): The effective optical excitation rate for the tunneling pro-Α rho numeric (required): The dimensionless density of recombination centers (defined as  $\rho$ ' in Huntley 2006) (unitless) times numeric (with default): The sequence of time steps within the simulation (s) clusters numeric (with default): The number of MC runs numeric (with default): Critical distance (>0) that is to be used if the sample has r\_c 1 been thermally and/or optically pretreated. This parameter expresses the fact that electron-hole pairs within a critical radius r\_c have already been recombined. delta.r numeric (with default): Increments of r\_c (unitless) numeric (width default): The total number of electron traps available (unitless) N\_e

method character (with default): Sequential 'seq' or parallel 'par' processing. In the

parallel mode the function tries to run the simulation on multiple CPU cores (if

available) with a positive effect on the computation time.

output character (with default): output is either the 'signal' (the default) or 'remaining\_e'

(the remaining charges, electrons, in the trap)

... further arguments

#### **Details**

#### The model

$$I_{TUN}(r',t) = -dn/dt = (A * t/P) * exp(-(\rho')^{-1/3} * r') * n(r',t)$$

Where in the function:

A := the optical excitation rate for the tunneling process

t := time(s)

P := maximum stimulation time

r' := the unitless tunneling radius

 $\rho$  := rho the unitless density of recombination centres see Huntley (2006)

n := the instantaneous number of electrons

#### Value

This function returns an object of class  $RLumCarlo\_Model\_Output$  which is a list consisting of an array with dimension length(times) x length(r) x clusters and a numeric time vector.

#### **Function version**

0.1.0

## How to cite

Friedrich, J., Kreutzer, S., 2019. run\_MC\_LM\_OSL\_TUN(): Run Monte-Carlo Simulation for LM-OSL (tunneling transitions). Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

#### Author(s)

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

#### References

Huntley, D.J., 2006. An explanation of the power-law decay of luminescence. Journal of Physics: Condensed Matter, 18(4), 1359.doi: 10.1088/09538984/18/4/020

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

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Further reading Aitken, M.J., 1985. Thermoluminescence dating. Academic Press.

Jain, M., Guralnik, B., Andersen, M.T., 2012. Stimulated luminescence emission from localized recombination in randomly distributed defects. J. Phys.: Condens. Matter 24, 385402. doi: 10.1088/09538984/24/38/385402

#### **Examples**

```
##the short example
run_MC_LM_OSL_TUN(
 A = 1,
 rho = 1e-7,
 times = 0:10,
 clusters = 10,
N_e = 100,
 r_c = 0.001,
 delta.r = 1e-1,
method = "seq",
output = "signal") %>%
plot_RLumCarlo(norm = TRUE)
## Not run:
## the long (meaningful) example
results <- run_MC_LM_OSL_TUN(
 A = 1,
 rho = 1e-3,
 times = 0:1000,
 clusters = 30,
 N_e = 100,
 r_c = 0.1
 delta.r = 1e-1,
 method = "par",
 output = "signal")
plot_RLumCarlo(results, norm = TRUE)
## End(Not run)
```

run\_MC\_TL\_DELOC

Run Monte-Carlo Simulation for TL (delocalized transitions)

## **Description**

Runs a Monte-Carlo (MC) simulation of thermoluminesence (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,
```

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```
N_e = 200,
n_filled = N_e,
R,
method = "par",
output = "signal",
...
)
```

#### **Arguments**

S	numeric ( <b>required</b> ): The frequency factor of the trap (s^-1)
E	numeric (required): Thermal activation energy of the trap (eV)
times	numeric (with default): The sequence of temperature steps within the simulation (s)
clusters	numeric (with default): The number of MC runs (unitless)
N_e	integer (with default): The total number of electron traps available (unitless)
n_filled	integer (with default): The number of filled electron traps at the beginning of the simulation (unitless)
R	numeric (with default): Retrapping ratio for delocalized transitions
method	character ( <i>with default</i> ): Sequential 'seq' or parallel 'par'processing. In the parallel mode the function tries to run the simulation on multiple CPU cores (if available) with a positive effect on the computation time.
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**

#### The model

```
I_{DELOC}(t) = -dn/dt = (s*exp(-E/(k_B*T)))*(n^2/(N*R+n(1-R))))
```

```
Where in the function:
```

```
E := the thermal activation enery (eV)

s := the frequency factor in (s^-1)

t := time

k_B := \text{Boltzmann constant}

T := temperature

n := n_filled, the instantaneous number of electrons

N := N_e, the total number of electron traps available (unitless)
```

### Value

This function returns an object of class RLumCarlo\_Model\_Output which is a list consisting of an array with dimension length(times) x clusters and a numeric time vector.

## **Function version**

0.1.0

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#### How to cite

Kreutzer, S., 2019. run\_MC\_TL\_DELOC(): Run Monte-Carlo Simulation for TL (delocalized transitions). Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

#### 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

#### **Further reading**

Chen, R., McKeever, S.W.S., 1997. Theory of Thermoluminescence and Related Phenomena. WORLD SCIENTIFIC. doi: 10.1142/2781

### **Examples**

```
## the short example
run_MC_TL_DELOC(
s = 3.5e12,
E = 1.45,
R = 0.1,
method = 'seq',
clusters = 3,
 times = 150:350) %>%
plot_RLumCarlo(legend = TRUE)
## Not run:
## the long (meaningful) example
# define your parameters
times <- seq(100, 450, 1)
s \leftarrow rep(3.5e12, 4)
E < - rep(1.45, 4)
R \leftarrow c(0.7e-6, 1e-6, 0.01, 0.1)
clusters <- 300
N_e < -c(400, 500, 700, 400)
n_{filled} \leftarrow 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?
add_TF <- c(FALSE, rep(TRUE, (length(R) - 1)))</pre>
# loop to plot different curves into one plot
for (u in 1:length(R)){
 results <- run_MC_TL_DELOC(
  times=times,
  s = s[u],
  E = E[u],
  clusters = clusters,
```

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```
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))
#add your legend with your parameters
legend("topright",
  ncol = 5,
  cex = 0.55,
  bty = "n",
  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 (localized transitions)

## **Description**

Runs a Monte-Carlo (MC) simulation of thermoluminesence (TL) using the generalized one trap (GOT) model. Localized transitions refer to transitions which do not involve the conduction or valence band. These transitions take place between the ground state and an excited state of the trapped charge, and also involve an excited state of the recombination center.

## Usage

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

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#### **Arguments**

numeric (required): The frequency factor of the trap (s^-1) s Ε numeric (required): Thermal activation energy of the trap (eV) times numeric (with default): The sequence of temperature steps within the simulation clusters numeric (with default): The number of MC run (unitless) integer (with default): The number of filled electron traps at the beginning of the n\_filled simulation (unitless) numeric (with default): The localized retrapping ratio (unitless) r character (with default): Sequential 'seq' or parallel 'par' processing. In the method parallel mode the function tries to run the simulation on multiple CPU cores (if available) with a positive effect on the computation time. character (with default): output is either the 'signal' (the default) or 'remaining\_e' output (the remaining charges/electrons in the trap)

... further arguments

#### **Details**

#### The model

$$I_{LOC}(t) = -dn/dt = (s * exp(-E/(k_B * T))) * (n^2/(r+n))$$

Where in the function:

E :=the thermal activation energy (eV)

s :=the frequency factor for the trap (s^-1)

t := time(s)

 $k_B := Boltzmann constant$ 

T := temperature (degrees C)

n := the instantaneous number of electrons

r := the retrapping ratio for localized transitions

## Value

This function returns an object of class RLumCarlo\_Model\_Output which is a list consisting of an array with dimension length(times) x clusters and a numeric time vector.

#### **Function version**

0.1.0

#### How to cite

Kreutzer, S., 2019. run\_MC\_TL\_LOC(): Run Monte-Carlo Simulation for TL (localized transitions). Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

#### 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., reutzer, 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

#### **Examples**

```
## the short example
run_MC_TL_LOC(
 s = 1e14,
E = 0.9,
 times = 50:100,
 method = "seq",
 clusters = 2,
 r = 1e4) %>%
plot_RLumCarlo()
## Not run:
## the long (meaningful) example
results <- run_MC_TL_LOC(</pre>
 s = 1e14,
E = 0.9,
 times = 50:100,
 method = "par"
 clusters = 100,
 r = 1e4)
## plot
plot_RLumCarlo(results)
## End(Not run)
```

run\_MC\_TL\_TUN

Run Monte-Carlo Simulation for TL (tunneling transitions)

### **Description**

Runs a Monte-Carlo (MC) simulation of thermoluminesence (TL) caused by tunnelling (TUN) transitions. Tunneling refers to quantum mechanical tunneling processes from the excited state of the trapped charge, into a recombination center.

### Usage

```
run_MC_TL_TUN(
    s,
    E,
    rho,
    r_c = 0,
    times,
    clusters = 10,
```

```
N_e = 200,
  delta.r = 0.1,
  method = "par",
  output = "signal",
   ...
)
```

## **Arguments**

S	list ( <b>required</b> ): The frequency factor for the tunneling process (s^-1)
E	numeric (required): Thermal activation energy of the trap (eV)
	• • • • • • • • • • • • • • • • • • • •
rho	numeric ( <b>required</b> ): The dimensionless density of recombination centers (defined as $\rho$ ' in Huntley 2006)
r_c	numeric ( <i>with default</i> ): Critical distance (>0) that is to be used if the sample has been thermally and/or optically pretreated. This parameter expresses the fact that electron-hole pairs within a critical radius r_c have already recombined.
times	vector (wih default): The sequence of time steps within the simulation (s)
clusters	numeric (with default): The number of MC runs (unitless)
N_e	numeric (with default): The total number of electron traps available (unitless)
delta.r	numeric (with default): The increments of r_c (unitless)
method	character ( <i>with default</i> ): Sequential 'seq' or parallel 'par' processing. In the parallel mode the function tries to run the simulation on multiple CPU cores (if available) with a positive effect on the computation time.
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**

#### The model

```
Where in the function:

s := \text{frequency for the tunneling process (s^-1)}

E := \text{thermal activation energy (eV)}

k_B := \text{Boltzmann constant}

T := \text{temperature}

r' := \text{the unitless tunneling radius}

\rho' := \text{rho'}, the unitless density of recombination centres (see Huntley (2006))

t := \text{time (s)}

t := \text{the instantaneous number of electrons}
```

 $I_{TUN}(r',t) = -dn/dt = (s * exp(-E/(k_B * T))) * exp(-(\rho')^{-1/3} * r') * n(r',t)$ 

## Value

This function returns an object of class  $RLumCarlo\_Model\_Output$  which is a list consisting of an array with dimension length(times) x length(r) x clusters and a numeric time vector.

## **Function version**

0.1.0

#### How to cite

Friedrich, J., Kreutzer, S., 2019. run\_MC\_TL\_TUN(): Run Monte-Carlo Simulation for TL (tunneling transitions). Function version 0.1.0. In: Friedrich, J., Kreutzer, S., Pagonis, V., Schmidt, C., 2019. RLumCarlo: Monte-Carlo Methods for Simulating Luminescence Phenomena. R package version 0.1.0.9000-133.

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## **Further reading**

Aitken, M.J., 1985. Thermoluminescence dating. Academic Press.

Jain, M., Guralnik, B., Andersen, M.T., 2012. Stimulated luminescence emission from localized recombination in randomly distributed defects. J. Phys.: Condens. Matter 24, 385402. doi: 10.1088/09538984/24/38/385402

#### **Examples**

```
## the short example
run_MC_TL_TUN(
s = 1e12,
E = 0.9,
rho = 1,
 r_c = 1,
 times = 80:120,
clusters = 2,
method = 'seq',
delta.r = 1e-1) %>%
plot_RLumCarlo()
## Not run:
## the long (meaningful example)
results <- run_MC_TL_TUN(
s = 1e12,
E = 0.9,
rho = 0.01,
 r_c = 0.1,
 times = 80:220,
clusters = 100,
 method = 'par',
 delta.r = 1e-1)
```

```
## plot
plot_RLumCarlo(results)
## End(Not run)
```

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