Package 'RLumCarlo'

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Description A Collection of Functions to Simulate Luminescence Production in Minerals using
      Monte-Carlo methods.
Contact Package Developer Team <sebastian.kreutzer@u-bordeaux-montaigne.fr>
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      utils.
      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 0.1.0.9000-77.

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 (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): 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 0.1.0.9000-77.

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 0.1.0.9000-77.

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 0.1.0.9000-77.

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 (required): 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 0.1.0.9000-77.

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

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 (required): 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 0.1.0.9000-77.

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 (required): Escape frequency of the trap (s^-1).
T	numeric (required): Temperature (deg. C).
rho	numeric (required): 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 0.1.0.9000-77.

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 (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_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 0.1.0.9000-77.

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

 ${\it Run\ Monte-Carlo\ simulation\ for\ LM-OSL\ for\ localized\ transition}$

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 0.1.0.9000-77.

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 (required): 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 0.1.0.9000-77.

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 (required): 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 0.1.0.9000-77.

Author(s)

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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 (required): Escape frequency of the trap (s^{Λ} -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_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 0.1.0.9000-77.

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 (required): 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 0.1.0.9000-77.

Author(s)

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