

# Package ‘adsorptionMCMC’

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**Type** Package

**Title** Bayesian Estimation of Adsorption Isotherms via MCMC

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**Author** Paul Angelo C. Manlapaz [aut, cre] (ORCID:  
<<https://orcid.org/0000-0002-1203-2064>>)

**Maintainer** Paul Angelo C. Manlapaz <[pacmanlapaz@gmail.com](mailto:pacmanlapaz@gmail.com)>

**Description** Provides tools for Bayesian parameter estimation of adsorption isotherm models using Markov Chain Monte Carlo (MCMC) methods. This package enables users to fit non-linear and linear adsorption isotherm models—Freundlich, Langmuir, and Temkin—within a probabilistic framework, capturing uncertainty and parameter correlations. It provides posterior summaries, 95% credible intervals, convergence diagnostics (Gelman-Rubin), and visualizations through trace and density plots. With this R package, researchers can rigorously analyze adsorption behavior in environmental and chemical systems using robust Bayesian inference. For more details, see Gilks et al. (1995) <[doi:10.1201/b14835](https://doi.org/10.1201/b14835)>, and Gamerman & Lopes (2006) <[doi:10.1201/9781482296426](https://doi.org/10.1201/9781482296426)>.

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mcmc_freundlichLM	<i>MCMC Analysis for Freundlich Isotherm Linear Model</i>
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### Description

Performs Bayesian parameter estimation using Markov Chain Monte Carlo (MCMC) to estimate the parameters of the Freundlich isotherm based on its linearized form:  $\log(Q_e) = \log(K_f) + (1/n)\log(C_e)$ . This method provides a probabilistic interpretation of the model parameters and accounts for their uncertainties. It supports multiple MCMC chains and computes convergence diagnostics (Gelman-Rubin).

### Arguments

Ce	Numeric vector of equilibrium concentrations.
Qe	Numeric vector of adsorbed amounts.
burnin	Integer specifying the number of burn-in iterations (default is 1000).
mcmc	Integer specifying the total number of MCMC iterations (default is 5000).
thin	Integer specifying the thinning interval (default is 10).
verbose	Integer controlling the frequency of progress updates (default is 100).
plot	Logical; if TRUE, trace and density plots of the MCMC chains are shown (default is FALSE).
n_chains	Number of independent MCMC chains (default = 2).
seed	Optional integer for reproducibility.

### Value

A list containing:

**mcmc\_results** An object of class `mcmc.list` containing posterior samples from all MCMC chains. Each chain includes samples of the intercept and slope.

**Kf\_mean** Posterior mean estimate of the Freundlich constant (Kf).

**n\_mean** Posterior mean estimate of the Freundlich exponent (n).

**logKf\_mean** Posterior mean of  $\log(K_f)$  from the first MCMC chain.

**inv\_n\_mean** Posterior mean of  $(1/n)$  (the slope) from the first MCMC chain.

**logKf\_sd** Posterior standard deviation of  $(\log(Kf))$ , quantifying uncertainty in the intercept estimate.

**inv\_n\_sd** Posterior standard deviation of  $(1/n)$ , quantifying uncertainty in the slope estimate.

**logKf\_ci** 95% credible interval for  $(\log(Kf))$  from the first MCMC chain, representing the posterior uncertainty in the intercept.

**inv\_n\_ci** 95% credible interval for  $(1/n)$  from the first MCMC chain, representing the posterior uncertainty in the slope.

**gelman\_diag** Gelman-Rubin diagnostic output from `coda::gelman.diag()`, used to assess convergence of the multiple MCMC chains. A potential scale reduction factor (PSRF) close to 1 indicates good convergence.

**mcmc\_summary** Summary statistics of the first MCMC chain, including means, standard deviations, quantiles, and sample sizes for each parameter.

### Author(s)

Paul Angelo C. Manlapaz

### References

Gilks, W. R., Richardson, S., & Spiegelhalter, D. J. (1995). *Markov Chain Monte Carlo in Practice*. Chapman and Hall/CRC.

### Examples

```
Ce <- c(0.01353, 0.04648, 0.13239, 0.27714, 0.41600, 0.63607, 0.80435, 1.10327, 1.58223)
Qe <- c(0.03409, 0.06025, 0.10622, 0.12842, 0.15299, 0.15379, 0.15735, 0.15735, 0.16607)
mcmc_freundlichLM(Ce, Qe, burnin = 1000, mcmc = 5000, thin = 10,
                  verbose = 100, plot = TRUE, n_chains = 2, seed = 123)
```

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mcmc\_freundlichNLM      *MCMC Analysis for Freundlich Isotherm Non-linear Model*

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### Description

Performs Bayesian parameter estimation using Markov Chain Monte Carlo (MCMC) for the non-linear Freundlich isotherm model:  $Q_e = K_f * C_e^{1/n}$ . This approach is applied to obtain a probabilistic distribution of the model parameters, capturing uncertainties and potential correlations between them.

### Arguments

Ce	Numeric vector of equilibrium concentrations.
Qe	Numeric vector of adsorbed amounts.
burnin	Integer specifying the number of burn-in iterations (default is 1000).
mcmc	Integer specifying the total number of MCMC iterations (default is 5000).

thin	Integer specifying the thinning interval (default is 10).
verbose	Integer controlling the frequency of progress updates (default is 100).
plot	Logical; if TRUE, trace and density plots of the MCMC chains are shown (default is FALSE).
n_chains	Number of independent MCMC chains (default = 2).
seed	Optional integer for reproducibility.

### Value

A list containing:

- Kf\_mean** Posterior mean estimate of Freundlich constant (Kf).
- n\_mean** Posterior mean estimate of Freundlich exponent (n).
- logKf\_mean** Posterior mean of (log(K\_f)).
- inv\_n\_mean** Posterior mean of (1/n).
- logKf\_sd** Posterior standard deviation for (log(Kf)).
- inv\_n\_sd** Posterior standard deviation for (1/n).
- logKf\_ci** 95% credible interval for (log(Kf)).
- inv\_n\_ci** 95% credible interval for (1/n).
- gelman\_diag** Gelman-Rubin diagnostics (only if multiple chains).
- mcmc\_summary** Summary statistics for each parameter.

### Author(s)

Paul Angelo C. Manlapaz

### References

Gilks, W. R., Richardson, S., & Spiegelhalter, D. J. (1995). *Markov Chain Monte Carlo in Practice*. Chapman and Hall/CRC.

### Examples

```
Ce <- c(0.01353, 0.04648, 0.13239, 0.27714, 0.41600, 0.63607, 0.80435, 1.10327, 1.58223)
Qe <- c(0.03409, 0.06025, 0.10622, 0.12842, 0.15299, 0.15379, 0.15735, 0.15735, 0.16607)
mcmc_freundlichNLM(Ce, Qe, burnin = 1000, mcmc = 5000, thin = 10,
                   verbose = 100, plot = TRUE, n_chains = 2, seed = 123)
```

**Description**

Performs Bayesian parameter estimation using Markov Chain Monte Carlo (MCMC) to estimate the parameters of the Langmuir isotherm based on its first linear form:  $C_e / Q_e = 1 / (Q_{\max} * b) + C_e / Q_{\max}$ . This method provides a probabilistic interpretation of the model parameters and accounts for their uncertainties. It supports multiple MCMC chains and computes convergence diagnostics (Gelman-Rubin).

**Arguments**

<code>Ce</code>	Numeric vector of equilibrium concentrations.
<code>Qe</code>	Numeric vector of adsorbed amounts.
<code>burnin</code>	Integer specifying the number of burn-in iterations (default is 1000).
<code>mcmc</code>	Integer specifying the total number of MCMC iterations (default is 5000).
<code>thin</code>	Integer specifying the thinning interval (default is 10).
<code>verbose</code>	Integer controlling the frequency of progress updates (default is 100).
<code>plot</code>	Logical; if TRUE, trace and density plots of the MCMC chains are shown (default is FALSE).
<code>n_chains</code>	Number of independent MCMC chains (default = 2).
<code>seed</code>	Optional integer for reproducibility.

**Value**

A list containing:

**mcmc\_results** An object of class `mcmc.list` containing posterior samples from all MCMC chains.

**Qmax\_mean** Posterior mean estimate of ( $Q_{\max}$ ).

**b\_mean** Posterior mean estimate of Langmuir constant ( $b$ ).

**slope\_mean** Posterior mean of the slope ( $(1/Q_{\max})$ ).

**intercept\_mean** Posterior mean of the intercept ( $(1/(Q_{\max}*b))$ ).

**slope\_sd** Posterior standard deviation of the slope.

**intercept\_sd** Posterior standard deviation of the intercept.

**slope\_ci** 95% credible interval for the slope.

**intercept\_ci** 95% credible interval for the intercept.

**gelman\_diag** Gelman-Rubin convergence diagnostic.

**mcmc\_summary** Summary statistics of the first MCMC chain.

**Author(s)**

Paul Angelo C. Manlapaz

## References

Gilks, W. R., Richardson, S., & Spiegelhalter, D. J. (1995). *Markov Chain Monte Carlo in Practice*. Chapman and Hall/CRC.

## Examples

```
Ce <- c(0.01353, 0.04648, 0.13239, 0.27714, 0.41600, 0.63607, 0.80435, 1.10327, 1.58223)
Qe <- c(0.03409, 0.06025, 0.10622, 0.12842, 0.15299, 0.15379, 0.15735, 0.15735, 0.16607)
mcmc_langmuirLM1(Ce, Qe, burnin = 1000, mcmc = 5000, thin = 10,
                 verbose = 100, plot = TRUE, n_chains = 2, seed = 123)
```

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mcmc\_langmuirLM2

*MCMC Analysis for Langmuir Isotherm Linear (Form 2) Model*


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## Description

Performs Bayesian parameter estimation using Markov Chain Monte Carlo (MCMC) to estimate the parameters of the Langmuir isotherm using its second linear form:  $1 / Q_e = 1 / Q_{max} + (1 / (Q_{max} * b)) * (1 / C_e)$  This method provides a probabilistic interpretation of the model parameters and accounts for their uncertainties. It supports multiple MCMC chains and computes convergence diagnostics (Gelman-Rubin).

## Arguments

Ce	Numeric vector of equilibrium concentrations.
Qe	Numeric vector of adsorbed amounts.
burnin	Integer specifying the number of burn-in iterations (default is 1000).
mcmc	Integer specifying the total number of MCMC iterations (default is 5000).
thin	Integer specifying the thinning interval (default is 10).
verbose	Integer controlling the frequency of progress updates (default is 100).
plot	Logical; if TRUE, trace and density plots of the MCMC chains are shown (default is FALSE).
n_chains	Number of independent MCMC chains (default = 2).
seed	Optional integer for reproducibility.

## Value

A list containing:

**mcmc\_results** An object of class `mcmc.list` with posterior samples from all chains.

**Qmax\_mean** Posterior mean estimate of ( $Q_{max}$ ).

**b\_mean** Posterior mean estimate of ( $b$ ).

**slope\_mean** Posterior mean of slope ( $1 / (Q_{max} * b)$ ).

**intercept\_mean** Posterior mean of intercept ( $(1/Q_{max})$ ).

**slope\_sd** Posterior standard deviation of the slope.  
**intercept\_sd** Posterior standard deviation of the intercept.  
**slope\_ci** 95% credible interval for the slope.  
**intercept\_ci** 95% credible interval for the intercept.  
**gelman\_diag** Gelman-Rubin convergence diagnostic.  
**mcmc\_summary** Summary statistics of the first MCMC chain.

### Author(s)

Paul Angelo C. Manlapaz

### References

Gilks, W. R., Richardson, S., & Spiegelhalter, D. J. (1995). *Markov Chain Monte Carlo in Practice*. Chapman and Hall/CRC.

### Examples

```
Ce <- c(0.01353, 0.04648, 0.13239, 0.27714, 0.41600, 0.63607, 0.80435, 1.10327, 1.58223)
Qe <- c(0.03409, 0.06025, 0.10622, 0.12842, 0.15299, 0.15379, 0.15735, 0.15735, 0.16607)
mcmc_langmuirLM2(Ce, Qe, burnin = 1000, mcmc = 5000, thin = 10,
                 verbose = 100, plot = TRUE, n_chains = 2, seed = 123)
```

---

mcmc\_langmuirLM3

*MCMC Analysis for Langmuir Isotherm Linear (Form 3) Model*

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### Description

Performs Bayesian parameter estimation using Markov Chain Monte Carlo (MCMC) to estimate the parameters of the Langmuir isotherm using its third linear form:  $Q_e = Q_{max} - (1 / b) * (Q_e / C_e)$  This method provides a probabilistic interpretation of the model parameters and accounts for their uncertainties. It supports multiple MCMC chains and computes convergence diagnostics (Gelman-Rubin).

### Arguments

Ce	Numeric vector of equilibrium concentrations.
Qe	Numeric vector of adsorbed amounts.
burnin	Integer specifying the number of burn-in iterations (default is 1000).
mcmc	Integer specifying the total number of MCMC iterations (default is 5000).
thin	Integer specifying the thinning interval (default is 10).
verbose	Integer controlling the frequency of progress updates (default is 100).
plot	Logical; if TRUE, trace and density plots of the MCMC chains are shown (default is FALSE).
n_chains	Number of independent MCMC chains (default = 2).
seed	Optional integer for reproducibility.

**Value**

A list containing:

**mcmc\_results** Combined posterior samples (mcmc.list).

**Qmax\_mean** Posterior mean of Qmax.

**b\_mean** Posterior mean of b.

**slope\_mean** Posterior mean of slope (-1/b).

**intercept\_mean** Posterior mean of Qmax.

**slope\_sd** Standard deviation of slope.

**intercept\_sd** Standard deviation of intercept.

**slope\_ci** 95% credible interval for slope.

**intercept\_ci** 95% credible interval for intercept.

**gelman\_diag** Gelman-Rubin convergence diagnostic.

**mcmc\_summary** Summary of main chain.

**Author(s)**

Paul Angelo C. Manlapaz

**References**

Gilks, W. R., Richardson, S., & Spiegelhalter, D. J. (1995). *Markov Chain Monte Carlo in Practice*. Chapman and Hall/CRC.

**Examples**

```
Ce <- c(0.01353, 0.04648, 0.13239, 0.27714, 0.41600, 0.63607, 0.80435, 1.10327, 1.58223)
Qe <- c(0.03409, 0.06025, 0.10622, 0.12842, 0.15299, 0.15379, 0.15735, 0.15735, 0.16607)
mcmc_langmuirLM3(Ce, Qe, burnin = 1000, mcmc = 10000, thin = 10,
                 verbose = 100, plot = TRUE, n_chains = 2, seed = 123)
```

---

mcmc\_langmuirLM4

*MCMC Analysis for Langmuir Isotherm Linear (Form 4) Model*

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**Description**

Performs Bayesian parameter estimation using Markov Chain Monte Carlo (MCMC) to estimate the parameters of the Langmuir isotherm using its fourth linear form:  $Q_e / C_e = b * Q_{max} - b * Q_e$ . This method provides a probabilistic interpretation of the model parameters and accounts for their uncertainties. It supports multiple MCMC chains and computes convergence diagnostics (Gelman-Rubin).



**Arguments**

Ce	Numeric vector of equilibrium concentrations.
Qe	Numeric vector of adsorbed amounts.
burnin	Integer specifying the number of burn-in iterations (default is 1000).
mcmc	Integer specifying the total number of MCMC iterations (default is 5000).
thin	Integer specifying the thinning interval (default is 10).
verbose	Integer controlling the frequency of progress updates (default is 100).
plot	Logical; if TRUE, trace and density plots of the MCMC chains are shown (default is FALSE).
n_chains	Number of independent MCMC chains (default = 2).
seed	Optional integer for reproducibility.

**Value**

A list with:

**mcmc\_results** Combined posterior samples (mcmc.list).

**Qmax\_mean** Posterior mean of Qmax.

**b\_mean** Posterior mean of b.

**intercept\_mean** Posterior mean of intercept (b \* Qmax).

**intercept\_sd** Posterior standard deviation of intercept.

**intercept\_ci** 95% credible interval for intercept.

**slope\_mean** Posterior mean of slope (-b).

**slope\_sd** Posterior standard deviation of slope.

**slope\_ci** 95% credible interval for slope.

**gelman\_diag** Gelman-Rubin convergence diagnostics.

**mcmc\_summary** Summary of the first MCMC chain.

**Author(s)**

Paul Angelo C. Manlapaz

**References**

Gilks, W. R., Richardson, S., & Spiegelhalter, D. J. (1995). *Markov Chain Monte Carlo in Practice*. Chapman and Hall/CRC.

**Examples**

```
Ce <- c(0.01353, 0.04648, 0.13239, 0.27714, 0.41600, 0.63607, 0.80435, 1.10327, 1.58223)
Qe <- c(0.03409, 0.06025, 0.10622, 0.12842, 0.15299, 0.15379, 0.15735, 0.15735, 0.16607)
mcmc_langmuirLM4(Ce, Qe, burnin = 1000, mcmc = 10000, thin = 10,
  verbose = 100, plot = TRUE, n_chains = 2, seed = 123)
```

mcmc\_langmuirNLM

*MCMC Analysis for Langmuir Isotherm Non-linear Model***Description**

This function performs Bayesian parameter estimation using Markov Chain Monte Carlo (MCMC) simulation to estimate the parameters of the Langmuir isotherm using the non-linear model:  $Q_e = (Q_{max} * K_L * C_e) / (1 + K_L * C_e)$ . This approach is applied to obtain a probabilistic distribution of the model parameters, capturing uncertainties and potential correlations between them.

**Arguments**

Ce	Numeric vector of equilibrium concentrations.
Qe	Numeric vector of adsorbed amounts.
burnin	Integer specifying the number of burn-in iterations (default is 1000).
mcmc	Integer specifying the total number of MCMC iterations (default is 5000).
thin	Integer specifying the thinning interval (default is 10).
verbose	Integer controlling the frequency of progress updates (default is 100).
plot	Logical; if TRUE, trace and density plots of the MCMC chains are shown (default is FALSE).
n_chains	Number of independent MCMC chains (default = 2).
seed	Optional integer for reproducibility.

**Value**

A list containing:

**Qmax\_mean** Posterior mean estimate of the Langmuir maximum adsorption capacity (Qmax).

**Kl\_mean** Posterior mean estimate of the Langmuir constant (Kl).

**Qmax\_sd** Posterior standard deviation for (Qmax).

**Kl\_sd** Posterior standard deviation for (Kl).

**Qmax\_ci** 95% credible interval for (Qmax).

**Kl\_ci** 95% credible interval for (Kl).

**gelman\_diag** Gelman-Rubin diagnostics (only if multiple chains).

**mcmc\_summary** Summary statistics for each parameter.

**Author(s)**

Paul Angelo C. Manlapaz

**Examples**

```
Ce <- c(0.01353, 0.04648, 0.13239, 0.27714, 0.41600, 0.63607, 0.80435, 1.10327, 1.58223)
Qe <- c(0.03409, 0.06025, 0.10622, 0.12842, 0.15299, 0.15379, 0.15735, 0.15735, 0.16607)
mcmc_langmuirNLM(Ce, Qe, burnin = 1000, mcmc = 5000, thin = 10,
  verbose = 100, plot = TRUE, n_chains = 2, seed = 123)
```

**Description**

Performs Bayesian parameter estimation using Markov Chain Monte Carlo (MCMC) to estimate the parameters of the Temkin isotherm based on its linearized form:  $Q_e = aT + bT * \log(C_e)$ . This method provides a probabilistic interpretation of the model parameters and accounts for their uncertainties. It supports multiple MCMC chains and computes convergence diagnostics (Gelman-Rubin).

**Arguments**

Ce	Numeric vector of equilibrium concentrations.
Qe	Numeric vector of adsorbed amounts.
Temp	Numeric value of temperature in Kelvin.
burnin	Integer specifying the number of burn-in iterations (default is 1000).
mcmc	Integer specifying the total number of MCMC iterations (default is 5000).
thin	Integer specifying the thinning interval (default is 10).
verbose	Integer controlling the frequency of progress updates (default is 100).
plot	Logical; if TRUE, trace and density plots of the MCMC chains are shown (default is FALSE).
n_chains	Number of independent MCMC chains (default = 2).
seed	Optional integer for reproducibility.

**Value**

A list containing:

**mcmc\_results** An object of class `mcmc.list` containing posterior samples from all MCMC chains.

**aT\_mean** Posterior mean estimate of Temkin constant (aT).

**bT\_mean** Posterior mean estimate of Temkin constant (bT).

**aT\_raw\_mean** Posterior mean of the intercept (aT) from the linear model.

**bT\_raw\_mean** Posterior mean of the slope (b\_T) from the linear model.

**aT\_sd** Posterior standard deviation of (aT).

**bT\_sd** Posterior standard deviation of (bT).

**aT\_ci** 95% credible interval for (aT).

**bT\_ci** 95% credible interval for (bT).

**gelman\_diag** Gelman-Rubin convergence diagnostic.

**mcmc\_summary** Summary statistics from the first chain.

**Author(s)**

Paul Angelo C. Manlapaz

**References**

Gilks, W. R., Richardson, S., & Spiegelhalter, D. J. (1995). *Markov Chain Monte Carlo in Practice*. Chapman and Hall/CRC.

**Examples**

```
Ce <- c(0.01353, 0.04648, 0.13239, 0.27714, 0.41600, 0.63607, 0.80435, 1.10327, 1.58223)
Qe <- c(0.03409, 0.06025, 0.10622, 0.12842, 0.15299, 0.15379, 0.15735, 0.15735, 0.16607)
mcmc_temkinLM(Ce, Qe, 298, burnin = 1000, mcmc = 5000, thin = 10,
              verbose = 100, plot = TRUE, n_chains = 2, seed = 123)
```

---

mcmc\_temkinNLM

*MCMC Analysis for Temkin Isotherm Non-linear Model*

---

**Description**

Performs Bayesian parameter estimation using Markov Chain Monte Carlo (MCMC) for the non-linear Temkin isotherm model:  $Q_e = (R * T / b_T) * \ln(A_T * C_e)$  This approach is applied to obtain a probabilistic distribution of the model parameters, capturing uncertainties and potential correlations between them.

**Arguments**

Ce	Numeric vector of equilibrium concentrations.
Qe	Numeric vector of adsorbed amounts.
Temp	Numeric value of temperature in Kelvin.
burnin	Integer specifying the number of burn-in iterations (default is 1000).
mcmc	Integer specifying the total number of MCMC iterations (default is 5000).
thin	Integer specifying the thinning interval (default is 10).
verbose	Integer controlling the frequency of progress updates (default is 100).
plot	Logical; if TRUE, trace and density plots of the MCMC chains are shown (default is FALSE).
n_chains	Number of independent MCMC chains (default = 2).
seed	Optional integer for reproducibility.

**Value**

A list containing:

**AT\_mean** Posterior mean estimate of Temkin constant (AT).

**bT\_mean** Posterior mean estimate of Temkin constant (bT).

**logAT\_mean** Posterior mean of (log(AT)).

**bT\_sd** Posterior standard deviation for (bT).

**logAT\_sd** Posterior standard deviation for (log(AT)).

**logAT\_ci** 95% credible interval for (log(AT)).

**bT\_ci** 95% credible interval for (bT).

**gelman\_diag** Gelman-Rubin diagnostics (only if multiple chains).

**mcmc\_summary** Summary statistics for each parameter.

**Author(s)**

Paul Angelo C. Manlapaz

**References**

Gilks, W. R., Richardson, S., & Spiegelhalter, D. J. (1995). *Markov Chain Monte Carlo in Practice*. Chapman and Hall/CRC.

**Examples**

```
Ce <- c(0.01353, 0.04648, 0.13239, 0.27714, 0.41600, 0.63607, 0.80435, 1.10327, 1.58223)
Qe <- c(0.03409, 0.06025, 0.10622, 0.12842, 0.15299, 0.15379, 0.15735, 0.15735, 0.16607)
mcmc_temkinNLM(Ce, Qe, 298, burnin = 1000, mcmc = 5000, thin = 10,
               verbose = 100, plot = TRUE, n_chains = 2, seed = 123)
```

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