

Package ‘isogeochem’

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Type Package

Title Tools For Carbonate Isotope Geochemistry

Version 0.0.0.9300

Description This collection of functions makes working with stable oxygen and clumped isotope data simpler and more reproducible.

License GPL (>= 3)

URL <https://github.com/davidbajnai/isogeochem>

Encoding UTF-8

LazyData true

Roxygen list(markdown = TRUE)

RoxygenNote 7.1.1

Suggests shades,
knitr,
rmarkdown

VignetteBuilder knitr

Imports graphics,
reconPlots (>= 0.1.0),
grDevices

Remotes andrewheiss/reconPlots

Depends R (>= 2.10)

BugReports <https://github.com/davidbajnai/isogeochem/issues>

R topics documented:

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| | |
|--------|---|
| a18c_w | <i>Calculate the 18O/16O fractionation factor between carbonate and water</i> |
|--------|---|

Description

a18c_w() calculates the equilibrium 18O/16O oxygen isotope fractionation factor between carbonate and water for a given temperature.

Usage

```
a18c_w(temp, min = "calcite", eq = "Daeron19")
```

Arguments

| | |
|------|---|
| temp | Crystallization temperature in degrees Celsius. |
| min | Mineralogy. Options are "calcite" (default), "aragonite", and "dolomite". |
| eq | Equation used to calculate the equilibrium 18O/16O oxygen isotope fractionation factor between calcium carbonate and water. For calcite choose from "Daeron19" (default), "Watkins13", "Coplen07", "K097", and "F077". For aragonite and dolomite eq need not be specified. |

Value

Returns the equilibrium 18O/16O oxygen isotope fractionation factor "alpha"

References

Calcite:

O'Neil, J. R., Clayton, R. N., & Mayeda, T. K. (1969). Oxygen isotope fractionation in divalent metal carbonates. *The Journal of Chemical Physics*, 51(12), 5547-5558. <https://doi.org/10.1063/1.1671982>

Kim, S.-T., & O'Neil, J. R. (1997). Equilibrium and nonequilibrium oxygen isotope effects in synthetic carbonates. *Geochimica et Cosmochimica Acta*, 61(16), 3461-3475. [https://doi.org/10.1016/S0016-7037\(97\)00169-5](https://doi.org/10.1016/S0016-7037(97)00169-5)

Coplen, T. B. (2007). Calibration of the calcite–water oxygen-isotope geothermometer at Devils Hole, Nevada, a natural laboratory. *Geochimica et Cosmochimica Acta*, 71(16), 3948-3957. <https://doi.org/10.1016/j.gca.2007.05.028>

Watkins, J. M., Nielsen, L. C., Ryerson, F. J., & DePaolo, D. J. (2013). The influence of kinetics on the oxygen isotope composition of calcium carbonate. *Earth and Planetary Science Letters*, 375, 349-360. <https://doi.org/10.1016/j.epsl.2013.05.054>

Daëron, M., Drysdale, R. N., Peral, M., Huyghe, D., Blamart, D., Coplen, T. B., et al. (2019). Most Earth-surface calcites precipitate out of isotopic equilibrium. *Nature Communications*, 10, 429. <https://doi.org/10.1038/s41467-019-08336-5>

Aragonite:

Dettman, D. L., Reische, A. K., & Lohmann, K. C. (1999). Controls on the stable isotope composition of seasonal growth bands in aragonitic fresh-water bivalves (unionidae). *Geochimica et Cosmochimica Acta*, 63(7-8), 1049-1057. [https://doi.org/10.1016/s0016-7037\(99\)00020-4](https://doi.org/10.1016/s0016-7037(99)00020-4)

Dolomite:

Vasconcelos, C., McKenzie, J. A., Warthmann, R., & Bernasconi, S. M. (2005). Calibration of the d18O paleothermometer for dolomite precipitated in microbial cultures and natural environments. *Geology*, 33(4), 317-320. <https://www.doi.org/10.1130/g20992.1>

Examples

```
a18c_w(25, "calcite", "Coplen07") # Returns 1.030249 and 30.25
a18c_w(25, "aragonite") # Returns 1.000913 and 0.91
a18c_w(25, "dolomite") # Returns 1.031456 and 31.46
```

| | |
|-------|-------------------------------------|
| d17Oc | <i>Triple oxygen isotope values</i> |
|-------|-------------------------------------|

Description

d17Oc() calculates equilibrium calcite d18O, d17O, and D17O values for a given temperature.

Usage

```
d17Oc(temp, d18ow_VSMOW, eq18 = "Daeron19", lambda = 0.528)
```

Arguments

| | |
|-------------|--|
| temp | Calcite growth temperature (°C). |
| d18ow_VSMOW | Water d18O value expressed on the VSMOW scale (‰). |
| eq18 | Equation used to calculate the equilibrium 18O/16O oxygen isotope fractionation factor between calcite and water. Options are "Daeron19" (default), "Watkins13", "Coplen07", "K097", and "F077". |
| lambda | Triple oxygen isotope reference slope. Default is 0.528. |

Details

$$\theta_{A/B} = \frac{\alpha_{A/B}^{17}}{\alpha_{A/B}^{18}}$$

$$\delta'^{17}O_{w,VSMOW} = \beta \times \delta'^{18}O_{w,VSMOW} + \gamma \text{ where } \beta = 0.528 \text{ and } \gamma = 0$$

$$\Delta^{17}O = \delta'^{17}O_{c,VSMOW} - \lambda \times \delta'^{18}O_{c,VSMOW}$$

Value

Returns a data frame:

- d18O value of the carbonate expressed on the VSMOW scale (‰).
- d18O value of the carbonate expressed on the VSMOW scale (‰).
- D17O value of the carbonate expressed on the VSMOW scale (‰).

References

Guo, W., & Zhou, C. (2019). Triple oxygen isotope fractionation in the DIC-H₂O-CO₂ system: A numerical framework and its implications. *Geochimica et Cosmochimica Acta*, 246, 541-564. <https://www.doi.org/10.1016/j.gca.2018.11.018>

See Also

Other equilibrium_carbonate: [D47c\(\)](#), [D48c\(\)](#), [d18Oc\(\)](#)

Examples

```
d17Oc(10,-1) # Returns d18Oc = 32.44, d17Oc = 16.91, D17O = -0.084
d17Oc(10,-1)[,3] # Returns D17O = -0.084
prime(d17Oc(10,-1)[,2]) # Returns d'17O = 16.77
```

| | |
|-------|---|
| d18Oc | <i>Equilibrium carbonate d18O value</i> |
|-------|---|

Description

d18Oc() calculates the equilibrium d18O value of a carbonate grown at a given temperature.

Usage

```
d18Oc(temp, d18Ow_VSMOW, min = "calcite", eq = "Daeron19")
```

Arguments

| | |
|-------------|---|
| temp | Crystallization temperature (°C). |
| d18Ow_VSMOW | Water d18O value expressed on the VSMOW scale (‰). |
| min | Mineralogy. Options are "calcite" (default), "aragonite", and "dolomite". |
| eq | Equation used to calculate the equilibrium 18O/16O oxygen isotope fractionation factor between calcium carbonate and water. Options depend on mineralogy. For calcite choose from "Daeron19" (default), "Watkins13", "Coplen07", "K097", and "F077". For aragonite and dolomite the eq need not be specified. |

Value

Returns the equilibrium carbonate d18O value expressed on the VSMOW scale (‰).

Note

Use [to_VSMOW\(\)](#) and [to_VPDB\(\)](#) to convert between the VSMOW and VPDB scales.

References

References are listed in the description of [a18c_w\(\)](#).

See Also

[d180w\(\)](#) calculates the d18O value of the ambient water from the d18O value of a carbonate and its growth temperature.

Other equilibrium_carbonate: [D47c\(\)](#), [D48c\(\)](#), [d170c\(\)](#)

Examples

```
d180c(33.7, -13.54, eq="Coplen07") # Returns 14.58
to_VPDB(d180c(12, -6.94, min="aragonite")) # Returns -5.21
d180c(25, -10.96, min="dolomite") # Returns 20.15
```

| | |
|-------|-----------------------------------|
| d180w | <i>Calculate water d18O value</i> |
|-------|-----------------------------------|

Description

[d180w\(\)](#) calculates the d18O value of the ambient water. from the d18O value of a carbonate and its growth temperature.

Usage

```
d180w(temp, d180c_VSMOW, min = "calcite", eq = "Daeron19")
```

Arguments

| | |
|-------------|---|
| temp | Crystallization temperature (°C). |
| d180c_VSMOW | Carbonate d18O value expressed on the VSMOW scale (‰). |
| min | Mineralogy. Options are "calcite" (default), "aragonite", and "dolomite". |
| eq | Equation used to calculate the equilibrium 18O/16O oxygen isotope fractionation factor between calcium carbonate and water. Options depend on mineralogy. For calcite choose from "Daeron19" (default), "Watkins13", "Coplen07", "K097", and "F077". For aragonite and dolomite the eq need not be specified. |

Value

Returns the water d18O value expressed on the VSMOW scale (‰).

Note

Use [to_VSMOW\(\)](#) and [to_VPDB\(\)](#) to convert between the VSMOW and VPDB scales.

References

References are listed in the description of `a18c_w()`.

See Also

`d180c()` calculates the equilibrium d18O value of a carbonate grown at a given temperature.

`temp_d180()` calculates growth temperatures from oxygen isotope data.

Examples

```
d180w(33.7, 14.58, "calcite", "Coplen07") # Returns -13.54
d180w(25, to_VSMOW(-7.47), "aragonite") # Returns -6.53
d180w(25, 20.43, "dolomite") # Returns -10.69
```

D47c

Carbonate D47 for a given temperature

Description

`D47c()` calculates the equilibrium carbonate D47 value for a given temperature.

Usage

```
D47c(temp, eq = "Fiebig21")
```

Arguments

| | |
|-------------------|--|
| <code>temp</code> | Carbonate growth temperature (°C). |
| <code>eq</code> | Equation used for the calculation. Options are "Fiebig21" (default) and "Petersen19". "Fiebig21" refers to the CDES90 calibration in Fiebig et al. (2021). "Petersen19" refers to the synthetic-only D47-RFACBr,WD "UNICAL" calibration of Petersen et al. (2019). |

Value

Returns the carbonate D47 value expressed on the CDES90 scale.

References

Petersen, S. V., Defliese, W. F., Saenger, C., Daëron, M., Huntington, K. W., John, C. M., et al. (2019). Effects of improved ^{17}O correction on interlaboratory agreement in clumped isotope calibrations, estimates of mineral-specific offsets, and temperature dependence of acid digestion fractionation. *Geochemistry, Geophysics, Geosystems*, 20(7), 3495-3519. <https://www.doi.org/10.1029/2018GC008127>

Fiebig, J., Daëron, M., Bernecker, M., Guo, W., Schneider, G., Boch, R., et al. (2021). Calibration of the dual clumped isotope thermometer for carbonates. *Geochimica et Cosmochimica Acta*. <https://www.doi.org/10.1016/j.gca.2021.07.012>

See Also

[temp_D47\(\)](#)

Other equilibrium_carbonate: [D48c\(\)](#), [d170c\(\)](#), [d180c\(\)](#)

Examples

```
D47c(33.7) # Returns 0.5713
```

| | |
|------|--|
| D48c | <i>Carbonate D48 for a given temperature</i> |
|------|--|

Description

D48c() calculates the equilibrium carbonate D48 value for a given temperature.

Usage

```
D48c(temperature, eq = "Fiebig21")
```

Arguments

| | |
|-------------|--|
| temperature | Carbonate growth temperature (°C). |
| eq | Equation used for the calculation. Options are "Fiebig21" (default) and "Swart21". "Fiebig21" refers to the CDES90 calibration in Fiebig et al. (2021). "Swart21" refers to the CDES90 "PBLM1" calibration in Swart et al. (2021). |

Value

Returns the carbonate D48 value expressed on the CDES90 scale (‰).

References

Fiebig, J., Daëron, M., Bernecker, M., Guo, W., Schneider, G., Boch, R., et al. (2021). Calibration of the dual clumped isotope thermometer for carbonates. *Geochimica et Cosmochimica Acta*. <https://www.doi.org/10.1016/j.gca.2021.07.012>

Swart, P. K., Lu, C., Moore, E., Smith, M., Murray, S. T., & Staudigel, P. T. (2021). A calibration equation between D48 values of carbonate and temperature. *Rapid Communications in Mass Spectrometry*, 35(17), e9147. <https://www.doi.org/10.1002/rcm.9147>

See Also

Other equilibrium_carbonate: [D47c\(\)](#), [d170c\(\)](#), [d180c\(\)](#)

Examples

```
D48c(33.7) # Returns 0.237
D48c(33.7,"Swart21") # Returns 0.239
```

devilshole

Devils Hole carbonate d18O time series

Description

A dataset containing the d18O values of the "original" Devils Hole cores.

Usage

devilshole

Format

A data frame with 442 rows and 4 variables:

age Interpolated uranium-series age of the sample expressed as thousands of years before present (ka).

d18O_VSMOW Carbonate d18O value relative to VSMOW (per mille).

d18O_error Standard deviation on the d18O value.

core Name of the core (DHC2-8, DHC2-3, DH-11).

Source

<https://doi.org/10.3133/ofr20111082>

References

Winograd, I. J., Landwehr, J. M., Coplen, T. B., Sharp, W. D., Riggs, A. C., Ludwig, K. R., & Kolesar, P. T. (2006). Devils Hole, Nevada, d18O record extended to the mid-Holocene. *Quaternary Research*, 66(2), 202-212. <https://doi.org/10.1016/j.yqres.2006.06.003>

See Also

Other "datasets": [GTS2020](#), [LR04](#)

GTS2020

Oxygen isotope stratigraphy from the Geologic Time Scale 2020: macrofossils

Description

A dataset containing a compilation of d18O and d13C values of various macrofossils (bivalves, gastropods, belemnites, ammonites) together with information on their age, shell mineralogy, and the climate zone they represent. This dataset is a condensed version of the entire dataset presented in the Geologic Time Scale 2020. Specifically, the full dataset was filtered for those "select" d18O and d13C values that also have age information.

Usage

GTS2020

Format

A data frame with 9676 rows and 8 variables:

age Age of the sample expressed as millions of years before present (Ma).

d18O_VPDB Carbonate d18O value relative to VPDB (per mille).

d13C_VPDB Carbonate d13C value relative to VPDB (per mille).

mineralogy The mineralogy of the carbonate hard part.

group The fossil group (bivalve, gastropod, belemnite, ammonite).

clim_zone The climate zone the fossil represents.

Source

https://download.pangaea.de/dataset/930093/files/GTS2020-App_10.2A.xlsx

References

Grossman, E. L., & Joachimski, M. M. (2020). Oxygen isotope stratigraphy. In F. M. Gradstein, J. G. Ogg, M. D. Schmitz, & G. M. Ogg (Eds.), *Geologic Time Scale 2020: Volume 1* (pp. 279-307): Elsevier. <https://doi.org/10.1016/B978-0-12-824360-2.00010-3>

See Also

Other "datasets": [LR04](#), [devilshole](#)

| | |
|------|---|
| LR04 | <i>A Pliocene-Pleistocene benthic foraminifera d18O stack</i> |
|------|---|

Description

A dataset containing the LR04 benthic d18O stack.

Usage

LR04

Format

A data frame with 2115 rows and 3 variables:

age Age of the sample expressed as thousands of years before present (ka).

d18O_VPDB Carbonate d18O value relative to VPDB (per mille).

d18O_error Standard error on the d18O value.

Source

<https://lorraine-lisiecki.com/stack.html>

References

Lisiecki, L. E., & Raymo, M. E. (2005). A Pliocene-Pleistocene stack of 57 globally distributed benthic d18O records. *Paleoceanography*, 20(1), PA1003. <https://doi.org/10.1029/2004pa001071>

See Also

Other "datasets": [GTS2020](#), [devilshole](#)

| | |
|----------|---|
| mix_d17O | <i>Mixing curves in triple oxygen isotope space</i> |
|----------|---|

Description

mix_d17O() produces mixing curves between two endmembers (A and B) in triple oxygen isotope space (d18O vs. D17O).

Usage

```
mix_d17O(d18O_A, d17O_A, d18O_B, d17O_B, lambda = 0.528)
```

Arguments

| | |
|--------|---|
| d18O_A | d18O value of component A (‰). |
| d17O_A | d17O value of component A (‰). |
| d18O_B | d18O value of component B (‰). |
| d17O_B | d17O value of component B (‰). |
| lambda | Triple oxygen isotope reference slope. Default 0.528. |

Value

Returns a data frame:

- d18O value of the mixture at x% mixing (‰).
- d17O value of the mixture x% mixing (‰).
- relative amount of component B in the mixture (%): from 100% A and 0% B to 0% A and 100% B.

See Also

[d17Oc\(\)](#) calculates equilibrium calcite d18O, d17O, and D17O values for a given temperature.

Examples

```
# Mixing between a Mesozoic marine carbonate and a diagenetic carbonate
mix_d17O(d17Oc(10,-1)[1],d17Oc(10,-1)[2],d17Oc(100,0)[1],d17Oc(100,0)[2])
```

| | |
|-------|--|
| prime | <i>Converting delta to delta prime</i> |
|-------|--|

Description

prime() converts "classical delta" values to "delta prime" values.

Usage

```
prime(classical)
```

Arguments

classical "Classical delta" values to be converted (parts per mille).

Details

$$\delta'^{17}O = 1000 \times \ln\left(\frac{\delta^{17}O}{1000} + 1\right)$$

Value

Returns the "delta prime" value (parts per mille).

See Also

[unprime\(\)](#) converts "delta prime" values to "classical delta" values.

Examples

```
prime(10)
```

| | |
|-----------|-----------------------------------|
| temp_d180 | <i>Oxygen isotope thermometry</i> |
|-----------|-----------------------------------|

Description

temp_d180() calculates carbonate growth temperature from oxygen isotope data.

Usage

```
temp_d180(d180c_VSMOW, d180w_VSMOW, eq = "Daeron19")
```

Arguments

| | |
|-------------|--|
| d180c_VSMOW | Carbonate d18O value expressed on the VSMOW scale (‰). |
| d180w_VSMOW | Water d18O value expressed on the VSMOW scale (‰). |
| eq | Equation used to calculate the equilibrium 18O/16O oxygen isotope fractionation factor between calcite and water. For calcite choose from "Daeron19" (default), "Watkins13", "Coplen07", "K097", and "F077". |

Value

Returns the carbonate growth temperature (°C).

Note

Use [to_VSMOW\(\)](#) and [to_VPDB\(\)](#) to convert between the VSMOW and VPDB scales.

References

References are listed in the description of [a18c_w\(\)](#).

See Also

[d180c\(\)](#) calculates the equilibrium d18O value of a carbonate grown at a given temperature.

[d180w\(\)](#) calculates the d18O value of the ambient water from the d18O value of a carbonate and its growth temperature.

Other thermometry: [temp_D47\(\)](#), [temp_D48\(\)](#)

Examples

```
temp_d180(14.58, -13.54, "Coplen07") # Returns 33.7
```

| | |
|----------|------------------------------------|
| temp_D47 | <i>Clumped isotope thermometry</i> |
|----------|------------------------------------|

Description

[temp_D47\(\)](#) calculates carbonate growth temperature from D47 value.

Usage

```
temp_D47(D47_CDES90, D47_error, eq = "Petersen19")
```

Arguments

| | |
|------------|--|
| D47_CDES90 | Carbonate D47 values expressed on the CDES90 scale: referenced to 90°C acid digestion (‰). |
| D47_error | Error on the D47 value. Optional. |
| eq | Equation used for the calculation. Options are "Petersen19" (default) and "Kele15". "Petersen19" refers to the synthetic-only IUPAC-reprocessed "Br UNICAL" calibration of Petersen et al. (2019). "Kele14" refers to the Kele et al. (2015) calibration reprocessed by Bernasconi et al. (2020) using the IUPAC parameters. |

Value

Returns the carbonate growth temperature (°C), and — if D47_error is specified — also the error.

References

References are listed at [D47c\(\)](#).

See Also

[D47c\(\)](#) calculates the equilibrium carbonate D47 value for a given temperature.

Other thermometry: [temp_D48\(\)](#), [temp_d180\(\)](#)

Examples

```
temp_D47(0.580) # Returns 33.7
temp_D47(0.580, 0.004) # Returns 33.7 and 1.9
```

| | |
|----------|---|
| temp_D48 | <i>Dual clumped isotope thermometry</i> |
|----------|---|

Description

temp_D48() calculates carbonate growth temperature from D47 and D48 values.

Usage

```
temp_D48(
  D47_CDES90,
  D48_CDES90,
  D47_error,
  D48_error,
  ks,
  add = FALSE,
  col,
  pch
)
```

Arguments

| | |
|------------|--|
| D47_CDES90 | Carbonate D47 values expressed on the CDES90 scale (‰). |
| D48_CDES90 | Carbonate D48 values expressed on the CDES90 scale (‰). |
| D47_error | Error on the D47 value. Optional. |
| D48_error | Error on the D48 value. Optional. |
| ks | Kinetic slope. Has to be negative! |
| add | Add graphics to an already existing plot? Default FALSE. |
| col | Graphical parameter. Optional. |
| pch | Graphical parameter. Optional. |

Details

The function calculates a D47 value as an intersect of two curves:

- the equilibrium D47 vs D48 curve from Fiebig et al. (2021)
- the kinetic slope

The resulting D47 value is then converted to temperature using the [D47c\(\)](#) function, i.e., by default the equation of Petersen et al. (2019). This is not consistent and I will fix it in a later version. In any case, the resulting discrepancy is smaller than the temperature error.

Value

Returns the carbonate growth temperature (‰).

References

References are listed at [D48c\(\)](#) and [D47c\(\)](#).

See Also

[D47c\(\)](#) calculates the equilibrium carbonate D47 value for a given temperature.

[D48c\(\)](#) calculates the equilibrium carbonate D48 value for a given temperature.

Other thermometry: [temp_D47\(\)](#), [temp_d180\(\)](#)

Examples

```
temp_D48(0.617, 0.139, ks = -0.6) # Returns 44
temp_D48(0.546, 0.277, ks = -1)  # Returns 33
```

to_VPDB

Converting between VSMOW and VPDB scales

Description

to_VPDB() converts d18O values expressed on the VSMOW scale to the VPDB scale.

Usage

```
to_VPDB(d180_VSMOW, eq = "IUPAC")
```

Arguments

| | |
|------------|--|
| d180_VSMOW | d18O values expressed on the VSMOW scale (‰). |
| eq | Equation used for the conversion. Options are "IUPAC", and "Coplen83". The default is "IUPAC": the IUPAC recommended equation listed in Brand et al. (2014) and Kim et al. (2015). To use the equation listed in Coplen et al. (1983) and the Hoefs book, set the parameter to "Coplen83". |

Details

The IUPAC recommended equation to convert between the scales is:

$$\delta^{18}O_{VPDB} = 0.97001 \times \delta^{18}O_{VSMOW} - 29.99$$

Value

Returns the d18O value expressed on the VPDB scale (‰).

References

References are listed at [to_VSMOW\(\)](#).

See Also

[to_VSMOW\(\)](#) converts d18O values expressed on the VPDB scale to the VSMOW scale.

Examples

```
to_VPDB(0) # Returns -29.99
to_VPDB(0, eq = "Coplen83") # Returns -29.98
```

to_VSMOW

*Converting between VPDB and VSMOW scales***Description**

to_VSMOW() converts d18O values expressed on the VPDB scale to the VSMOW scale.

Usage

```
to_VSMOW(d18O_VPDB, eq = "IUPAC")
```

Arguments

| | |
|-----------|--|
| d18O_VPDB | d18O values expressed on the VPDB scale (‰). |
| eq | Equation used for the conversion. Options are "IUPAC", and "Coplen83". The default is "IUPAC": the IUPAC recommended equation listed in Brand et al. (2014) and Kim et al. (2015). To use the equation listed in Coplen et al. (1983) and the Hoefs book, set the parameter to "Coplen83". |

Details

The IUPAC recommended equation to convert between the scales is:

$$\delta^{18}O_{VSMOW} = 1.03092 \times \delta^{18}O_{VPDB} + 30.92$$

Value

Returns the d18O value expressed on the VSMOW scale (‰).

References

- Coplen, T. B., Kendall, C., & Hopple, J. (1983). Comparison of stable isotope reference samples. *Nature*, 302, 236-238. <https://doi.org/10.1038/302236a0>
- Brand, W. A., Coplen, T. B., Vogl, J., Rosner, M., & Prohaska, T. (2014). Assessment of international reference materials for isotope-ratio analysis (IUPAC Technical Report). *Pure and Applied Chemistry*, 86(3), 425-467. <https://doi.org/10.1515/pac-2013-1023>
- Kim, S.-T., Coplen, T. B., & Horita, J. (2015). Normalization of stable isotope data for carbonate minerals: Implementation of IUPAC guidelines. *Geochimica et Cosmochimica Acta*, 158, 276-289. <https://doi.org/10.1016/j.gca.2015.02.011>

See Also

[to_VPDB\(\)](#) converts d18O values expressed on the VSMOW scale to the VPDB scale.

Examples

```
to_VSMOW(0) # Returns 30.92
to_VSMOW(0, eq = "Coplen83") # Returns 30.91
```

unprime

Converting delta prime to delta

Description

unprime() converts "delta prime" values to "classical delta" values.

Usage

```
unprime(prime)
```

Arguments

prime "Delta prime" values to be converted (parts per mille).

Details

$$\delta^{17}O = 1000 \times e^{(\frac{\delta^{17}O}{1000} + 1)}$$

Value

Returns the "classical delta" value (parts per mille).

See Also

[prime\(\)](#) converts "classical delta" values to "delta prime" values.

Examples

```
unprime(9.950331)
```

| | |
|------|---|
| xDIC | <i>Dissolved inorganic carbon species</i> |
|------|---|

Description

xDIC() calculates the relative abundance of the DIC species as a function of solution temperature, pH, and salinity.

Usage

```
xDIC(temp, pH, S)
```

Arguments

| | |
|------|---|
| temp | The temperature of the solution (°C). |
| pH | The pH of the solution. |
| S | The salinity of the solution (g/kg, ‰). |

Value

Returns a data frame with the relative abundance of the DIC species:

- Relative abundance of dissolved CO₂ (%).
- Relative abundance of bicarbonate ion (%).
- Relative abundance of carbonate ion (%).

Examples

```
xDIC(25, 7, 30)
```

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