

Package ‘tidywater’

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Description Provides multiple water chemistry-based models and published empirical models in one standard format. As many models have been included as possible, however, users should be aware that models have varying degrees of accuracy and applicability. To learn more, read the references provided below for the models implemented. Functions can be chained together to model a complete treatment process and are designed to work in a 'tidyverse' workflow. Models are primarily based on these sources:
Benjamin, M. M. (2002, ISBN:147862308X),
Crittenden, J. C., Trussell, R., Hand, D., Howe, J. K., & Tchobanoglous, G., Borchardt, J. H. (2012, ISBN:9781118131473),
USEPA. (2001) <https://www.epa.gov/sites/default/files/2017-03/documents/wtp_model_v._2.0_manual_508.pdf>.

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balance_ions	<i>Add an ion to balance overall charge in a water</i>
--------------	--

Description

This function takes a water defined by [define_water](#) and balances charge. For a single water use `balance_ions`; for a dataframe use `balance_ions_chain`. Use [pluck_water](#) to get values from the output water as new dataframe columns.

Usage

```
balance_ions(water, anion = "cl", cation = "na")

balance_ions_chain(
  df,
  input_water = "defined_water",
  output_water = "balanced_water",
  anion = "cl",
  cation = "na"
)
```

Arguments

water	Water created with define_water , which may have some ions set to 0 when unknown
anion	Selected anion to use to for ion balance when more cations are present. Defaults to "cl". Choose one of c("cl", "so4").
cation	Selected cation to use to for ion balance when more anions are present. Defaults to "na". Choose one of c("na", "k", "ca", or "mg").
df	a data frame containing a water class column, which has already been computed using define_water_chain
input_water	name of the column of water class data to be used as the input for this function. Default is "defined_water".
output_water	name of the output column storing updated water classes. Default is "balanced_water".

Details

If more cations are needed, sodium will be added. User may specify which cation ("na", "k", "ca", or "mg") to use for balancing. If calcium and magnesium are not specified when defining a water with [define_water](#), they will default to 0 and not be changed by this function unless specified in the cation argument. Anions are added by default with chloride. User may specify which anion ("cl", "so4") to use for balancing. This function is purely mathematical. User should always check the outputs to make sure values are reasonable for the input source water.

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

`balance_ions` returns a single water class object with updated ions to balance water charge.

`balance_ions_chain` returns a dataframe with a new column with the ion balanced water

Examples

```
water_defined <- define_water(7, 20, 50, 100, 80, 10, 10, 10, 10) %>%
  balance_ions()

water_defined <- define_water(7, 20, 50, tot_hard = 150) %>%
  balance_ions(anion = "so4")

example_df <- water_df %>%
  define_water_chain() %>%
  balance_ions_chain(anion = "so4", cation = "ca")
```

```
# Initialize parallel processing
library(furrr)
# plan(multisession)
example_df <- water_df %>%
  define_water_chain() %>%
  balance_ions_chain()

# Optional: explicitly close multisession processing
# plan(sequential)
```

biofilter_toc	<i>Determine TOC removal from biofiltration using Terry & Summers BDOC model</i>
---------------	--

Description

This function applies the Terry model to a water created by [define_water](#) to determine biofiltered DOC (mg/L). All particulate TOC is assumed to be removed so TOC = DOC. For a single water use `biofilter_toc`; for a dataframe use `biofilter_toc_chain`. Use [pluck_water](#) to get values from the output water as new dataframe columns. For most arguments in the `_chain` helper "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```
biofilter_toc(water, ebct, ozonated = TRUE)
```

```
biofilter_toc_chain(
  df,
  input_water = "defined_water",
  output_water = "biofiltered_water",
  ebct = "use_col",
  ozonated = "use_col"
)
```

Arguments

<code>water</code>	Source water object of class "water" created by define_water .
<code>ebct</code>	The empty bed contact time (min) used for the biofilter.
<code>ozonated</code>	Logical; TRUE if the water is ozonated (default), FALSE otherwise.
<code>df</code>	a data frame containing a water class column, which has already been computed using define_water_chain . The df may include a column indicating the EBCT or whether the water is ozonated.
<code>input_water</code>	name of the column of Water class data to be used as the input for this function. Default is "defined_water".
<code>output_water</code>	name of the output column storing updated parameters with the class, Water. Default is "biofiltered_water".

Details

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

`biofilter_toc` returns water class object with modeled DOC removal from biofiltration.

`biofilter_toc_chain` returns a data frame containing a water class column with updated DOC, TOC, and UV254 water slots.

Source

Terry and Summers 2018

Examples

```
library(tidywater)
water <- define_water(ph = 7, temp = 25, alk = 100, toc = 5.0, doc = 4.0, uv254 = .1) %>%
  biofilter_toc(ebct = 10, ozonated = FALSE)

library(purrr)
library(tidyr)
library(dplyr)

example_df <- water_df %>%
  define_water_chain() %>%
  biofilter_toc_chain(input_water = "defined_water", ebct = 10, ozonated = FALSE)

example_df <- water_df %>%
  define_water_chain() %>%
  mutate(
    BiofEBCT = c(10, 10, 10, 15, 15, 15, 20, 20, 20, 25, 25, 25),
    ozonated = c(rep(TRUE, 6), rep(FALSE, 6))
  ) %>%
  biofilter_toc_chain(input_water = "defined_water", ebct = BiofEBCT)

# Initialize parallel processing
library(furrr)
# plan(multisession)
example_df <- water_df %>%
  define_water_chain() %>%
  biofilter_toc_chain(input_water = "defined_water", ebct = c(10, 20))

# Optional: explicitly close multisession processing
# plan(sequential)
```

blend_waters	<i>Determine blended water quality from multiple waters based on mass balance and acid/base equilibrium</i>
--------------	---

Description

This function takes a vector of waters defined by [define_water](#) and a vector of ratios and outputs a new water object with updated ions and pH. For a single blend use `blend_waters`; for a dataframe use `blend_waters_chain`. Use [pluck_water](#) to get values from the output water as new dataframe columns.

Usage

```
blend_waters(waters, ratios)
```

```
blend_waters_chain(df, waters, ratios, output_water = "blended_water")
```

Arguments

waters	Vector of source waters created by define_water . For chain function, this can include quoted column names and/or existing single water objects unquoted.
ratios	Vector of ratios in the same order as waters. (Blend ratios must sum to 1). For chain function, this can also be a list of quoted column names.
df	a data frame containing a water class column, which has already been computed using define_water_chain
output_water	name of output column storing updated parameters with the class, water. Default is "blended_water".

Details

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.[#]

Value

`blend_waters` returns a water class object with blended water quality parameters.

`blend_waters_chain` returns a data frame with a water class column containing blended water quality

See Also[define_water](#)**Examples**

```

water1 <- define_water(7, 20, 50)
water2 <- define_water(7.5, 20, 100, tot_nh3 = 2)
blend_waters(c(water1, water2), c(.4, .6))

library(dplyr)

example_df <- water_df %>%
  slice_head(n = 3) %>%
  define_water_chain() %>%
  chemdose_ph_chain(naoh = 22) %>%
  mutate(
    ratios1 = .4,
    ratios2 = .6
  ) %>%
  blend_waters_chain(
    waters = c("defined_water", "dosed_chem_water"),
    ratios = c("ratios1", "ratios2"), output_water = "Blending_after_chemicals"
  )

waterA <- define_water(7, 20, 100, tds = 100)
example_df <- water_df %>%
  slice_head(n = 3) %>%
  define_water_chain() %>%
  blend_waters_chain(waters = c("defined_water", waterA), ratios = c(.8, .2))

# Initialize parallel processing
library(furrr)
# plan(multisession)
example_df <- water_df %>%
  define_water_chain() %>%
  balance_ions_chain() %>%
  chemdose_ph_chain(naoh = 22, output_water = "dosed") %>%
  blend_waters_chain(waters = c("defined_water", "dosed", "balanced_water"), ratios = c(.2, .3, .5))

# Optional: explicitly close multisession processing
# plan(sequential)

```

bromatecoeffs

Data frame of bromate coefficients for predicting bromate formation during ozonation

Description

A dataset containing coefficients for calculating ozone formation

Usage

```
bromatecoeffs
```

Format

A dataframe with 30 rows and 10 columns

model First author of source model

ammonia Either T or F, depending on whether the model applies to waters with ammonia present.

A First coefficient in bromate model

a Exponent in bromate model, associated with Br-

b Exponent in bromate model, associated with DOC

c Exponent in bromate model, associated with UVA

d Exponent in bromate model, associated with pH

e Exponent in bromate model, associated with Alkalinity

f Exponent in bromate model, associated with ozone dose

g Exponent in bromate model, associated with reaction time

h Exponent in bromate model, associated with ammonia (NH₄⁺)

i Exponent in bromate model, associated with temperature

I Coefficient in bromate model, associated with temperature in the exponent. Either i or I are used, not both.

Source

Ozekin (1994), Sohn et al (2004), Song et al (1996), Galey et al (1997), Siddiqui et al (1994)

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

calculate_activity	<i>Calculate activity coefficients</i>
--------------------	--

Description

This function calculates activity coefficients at a given temperature based on equation 5-43 from Davies (1967), Crittenden et al. (2012)

Usage

```
calculate_activity(z, is, temp)
```

Arguments

<code>z</code>	Charge of ions in the solution
<code>is</code>	Ionic strength of the solution
<code>temp</code>	Temperature of the solution in Celsius

Value

A numeric value for the activity coefficient.

Examples

```
calculate_activity(2, 0.1, 25)
```

<code>calculate_corrosion</code>	<i>Calculate six corrosion and scaling indices (AI, RI, LSI, LI, CSMR, CCPP)</i>
----------------------------------	--

Description

This function takes an object created by [define_water](#) and calculates corrosion and scaling indices. For a single water, use `calculate_corrosion`; to apply the calculations to a dataframe, use `calculate_corrosion_once`.

Usage

```
calculate_corrosion(
  water,
  index = c("aggressive", "ryznar", "langelier", "ccpp", "laronskold", "csmr"),
  form = "calcite"
)

calculate_corrosion_once(
  df,
  input_water = "defined_water",
  index = c("aggressive", "ryznar", "langelier", "ccpp", "laronskold", "csmr"),
  form = "calcite"
)
```

Arguments

<code>water</code>	Source water of class "water" created by define_water
<code>index</code>	The indices to be calculated. Default calculates all six indices: "aggressive", "ryznar", "langelier", "ccpp", "laronskold", "csmr" CCPP may not be able to be calculated sometimes, so it may be advantageous to leave this out of the function to avoid errors

form	Form of calcium carbonate mineral to use for modelling solubility: "calcite" (default), "aragonite", or "vaterite"
df	a data frame containing a water class column, created using define_water
input_water	name of the column of water class data to be used as the input. Default is "defined_water".

Details

Aggressiveness Index (AI), unitless - the corrosive tendency of water and its effect on asbestos cement pipe.

Ryznar Index (RI), unitless - a measure of scaling potential.

Langelier Saturation Index (LSI), unitless - describes the potential for calcium carbonate scale formation. Equations use empirical calcium carbonate solubilities from Plummer and Busenberg (1982) and Crittenden et al. (2012) rather than calculated from the concentrations of calcium and carbonate in the water.

Larson-skold Index (LI), unitless - describes the corrosivity towards mild steel.

Chloride-to-sulfate mass ratio (CSMR), mg Cl/mg SO₄ - indicator of galvanic corrosion for lead solder pipe joints.

Calcium carbonate precipitation potential (CCPP), mg/L as CaCO₃ - a prediction of the mass of calcium carbonate that will precipitate at equilibrium. A positive CCPP value indicates the amount of CaCO₃ (mg/L as CaCO₃) that will precipitate. A negative CCPP indicates how much CaCO₃ can be dissolved in the water.

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

`calculate_corrosion` returns a data frame with corrosion and scaling indices as individual columns.

`calculate_corrosion_once` returns the a data frame containing specified corrosion and scaling indices as columns.

Source

AWWA (1977)

Crittenden et al. (2012)

Langelier (1936)

Larson and Skold (1958)

Merrill and Sanks (1977a)

Merrill and Sanks (1977b)

Merrill and Sanks (1978)

Nguyen et al. (2011)

Plummer and Busenberg (1982)

Ryznar (1944)

Schock (1984)

Trussell (1998)

U.S. EPA (1980)

See reference list at <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

Examples

```
water <- define_water(
  ph = 8, temp = 25, alk = 200, tot_hard = 200,
  tds = 576, cl = 150, so4 = 200
)
corrosion_indices <- calculate_corrosion(water)

water <- define_water(ph = 8, temp = 25, alk = 100, tot_hard = 50, tds = 200)
corrosion_indices <- calculate_corrosion(water, index = c("aggressive", "ccpp"))

library(dplyr)

example_df <- water_df %>%
  slice_head(n = 2) %>% # used to make example run faster
  define_water_chain() %>%
  calculate_corrosion_once(index = c("aggressive", "ccpp"))
```

calculate_hardness	<i>Calculate hardness from calcium and magnesium</i>
--------------------	--

Description

This function takes Ca and Mg in mg/L and returns hardness in mg/L as CaCO₃

Usage

```
calculate_hardness(ca, mg, type = "total", startunit = "mg/L")
```

Arguments

ca	Calcium concentration in mg/L as Ca
mg	Magnesium concentration in mg/L as Mg
type	"total" returns total hardness, "ca" returns calcium hardness. Defaults to "total"
startunit	Units of Ca and Mg. Defaults to mg/L

Value

A numeric value for the total hardness in mg/L as CaCO₃.

Examples

```
calculate_hardness(50, 10)

water_defined <- define_water(7, 20, 50, 100, 80, 10, 10, 10, 10, tot_po4 = 1)
calculate_hardness(water_defined@ca, water_defined@mg, "total", "M")
```

chemdose_chloramine	<i>Calculate chlorine and chloramine Concentrations with the breakpoint chlorination approach</i>
---------------------	---

Description

[chemdose_chloramine](#), adopted from the U.S. EPA's Chlorine Breakpoint Curve Simulator, calculates chlorine and chloramine concentrations based on the two papers Jafvert & Valentine (Environ. Sci. Technol., 1992, 26 (3), pp 577-586) and Vikesland et al. (Water Res., 2001, 35 (7), pp 1766-1776). Required arguments include an object of class "water" created by [define_water](#), chlorine dose, and reaction time. The function also requires additional water quality parameters defined in [define_water](#) including temperature, pH, and alkalinity.

Usage

```
chemdose_chloramine(
  water,
  time,
  cl2 = 0,
  nh3 = 0,
  use_free_cl_slot = FALSE,
  use_tot_nh3_slot = FALSE
)

chemdose_chloramine_chain(
  df,
  input_water = "defined_water",
  output_water = "chlorinated_water",
  time = "use_col",
  cl2 = "use_col",
  nh3 = "use_col",
  use_free_cl_slot = "use_col",
  use_tot_nh3_slot = "use_col"
)
```

Arguments

water	Source water object of class "water" created by define_water
time	Reaction time (minutes). Time defined needs to be greater or equal to 1 minute.
cl2	Applied chlorine dose (mg/L as Cl ₂), defaults to 0. If not specified, use free_chlorine slot in water.
nh3	Applied ammonia dose (mg/L as N), defaults to 0. If not specified, use tot_nh3 slot in water.
use_free_cl_slot	Defaults to FALSE. If TRUE, uses free_chlorine slot in water. If TRUE AND there is a cl2 input, both the free_chlorine water slot and chlorine dose will be used.
use_tot_nh3_slot	Defaults to FALSE. If TRUE, uses tot_nh3 slot in water. If TRUE AND there is a nh3 input, both the tot_nh3 water slot and ammonia dose will be used.
df	a data frame containing a water class column, which has already been computed using define_water_chain . The df may include a column named for the applied chlorine dose (cl2_dose), and a column for time in hours.
input_water	name of the column of water class data to be used as the input for this function. Default is "defined_water".
output_water	name of the output column storing updated parameters with the class, water. Default is "chlorinated_water".

Details

The function will calculate the chlorine and chloramine concentrations and update the "water" class object proceed to the next steps of the treatment chain.

Value

chemdose_chloramine returns a water class object with predicted chlorine and chloramine concentrations.

chemdose_chloramine_chain returns a data frame containing water class column with updated chlorine residuals.

Source

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

Examples

```
breakpoint <- define_water(7.5, 20, 65, free_chlorine = 5, tot_nh3 = 1) %>%
  chemdose_chloramine(time = 40, cl2 = 2, nh3 = 1, use_free_cl_slot = TRUE)

library(dplyr)

breakpoint <- water_df %>%
```

```

mutate(free_chlorine = 5, tot_nh3 = 1) %>%
slice_head(n = 3) %>%
define_water_chain() %>%
mutate(
  time = 8,
  cl2dose = c(2, 3, 4)
) %>%
chemdose_chloramine_chain(
  input_water = "defined_water",
  cl2 = cl2dose,
  use_free_cl_slot = TRUE,
  use_tot_nh3_slot = TRUE
)

# Initialize parallel processing
library(furrr)
# plan(multisession)

example_df <- water_df %>%
  define_water_chain() %>%
  chemdose_chloramine_chain(
    input_water = "defined_water", cl2 = c(2, 4), nh3 = 2, time = 8
  )

# Optional: explicitly close multisession processing
# plan(sequential)

```

chemdose_chlordecay *Calculate chlorine decay*

Description

calculates the decay of chlorine or chloramine based on the U.S. EPA's Water Treatment Plant Model (U.S. EPA, 2001). For a single water use `chemdose_chlordecay`; for a dataframe use `chemdose_chlordecay_chain`. For most arguments in the `_chain` helper "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```

chemdose_chlordecay(
  water,
  cl2_dose,
  time,
  treatment = "raw",
  cl_type = "chlorine",

```

```

    use_chlorine_slot = FALSE
  )

  chemdose_chlordecay_chain(
    df,
    input_water = "defined_water",
    output_water = "disinfected_water",
    cl2_dose = "use_col",
    time = "use_col",
    treatment = "use_col",
    cl_type = "use_col",
    use_chlorine_slot = "use_col"
  )

```

Arguments

water	Source water object of class "water" created by define_water
cl2_dose	Applied chlorine or chloramine dose (mg/L as cl2). Model results are valid for doses between 0.995 and 41.7 mg/L for raw water, and for doses between 1.11 and 24.7 mg/L for coagulated water.
time	Reaction time (hours). Chlorine decay model results are valid for reaction times between 0.25 and 120 hours. Chloramine decay model does not have specified boundary conditions.
treatment	Type of treatment applied to the water. Options include "raw" for no treatment (default), "coag" for water that has been coagulated or softened.
cl_type	Type of chlorination applied, either "chlorine" (default) or "chloramine".
use_chlorine_slot	Defaults to FALSE. When TRUE, uses either free_chlorine or combined_chlorine slot in water (depending on cl_type). If 'cl2_dose' argument, not specified, chlorine slot will be used. If 'cl2_dose' specified and use_chlorine_slot is TRUE, all chlorine will be summed.
df	a data frame containing a water class column, which has already been computed using define_water_once . The df may include a column named for the applied chlorine dose (cl2), and a column for time in hours.
input_water	name of the column of water class data to be used as the input for this function. Default is "defined_water".
output_water	name of the output column storing updated parameters with the class, water. Default is "disinfected_water".

Details

Required arguments include an object of class "water" created by [define_water](#), applied chlorine/chloramine dose, type, reaction time, and treatment applied (options include "raw" for no treatment, or "coag" for coagulated water). The function also requires additional water quality parameters defined in [define_water](#) including TOC and UV254. The output is a new "water" class with the calculated total chlorine value stored in the 'free_chlorine' or 'combined_chlorine' slot, depending on what type of chlorine is dosed. When modeling residual concentrations through a

unit process, the U.S. EPA Water Treatment Plant Model applies a correction factor based on the influent and effluent residual concentrations (see U.S. EPA (2001) equation 5-118) that may need to be applied manually by the user based on the output.

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.[#]

Value

`chemdose_chlordecay` returns an updated disinfectant residual in the `free_chlorine` or `combined_chlorine` water slot in units of M. Use [convert_units](#) to convert to mg/L.

`chemdose_chlordecay_chain` returns a data frame containing a water class column with updated chlorine residuals.

Source

U.S. EPA (2001)

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

Examples

```
example_cl2 <- define_water(8, 20, 66, toc = 4, uv254 = 0.2) %>%
  chemdose_chlordecay(cl2_dose = 2, time = 8)

example_cl2 <- define_water(8, 20, 66, toc = 4, uv254 = 0.2, free_chlorine = 3) %>%
  chemdose_chlordecay(cl2_dose = 2, time = 8, use_chlorine_slot = TRUE)

library(dplyr)

example_df <- water_df %>%
  mutate(br = 50) %>%
  define_water_chain() %>%
  chemdose_chlordecay_chain(input_water = "defined_water", cl2_dose = 4, time = 8)

example_df <- water_df %>%
  mutate(
    br = 50,
    free_chlorine = 2
  ) %>%
  define_water_chain() %>%
  mutate(
    cl2_dose = seq(2, 24, 2),
    ClTime = 30
  ) %>%
  chemdose_chlordecay_chain(
    time = ClTime,
```

```

    use_chlorine_slot = TRUE,
    treatment = "coag",
    cl_type = "chloramine"
  )

# Initialize parallel processing
library(furrr)
# plan(multisession)
example_df <- water_df %>%
  mutate(br = 50) %>%
  define_water_chain() %>%
  chemdose_chlordecay_chain(cl2_dose = 4, time = 8)

# Optional: explicitly close multisession processing
# plan(sequential)

```

chemdose_dbp

Calculate DBP formation

Description

Calculates disinfection byproduct (DBP) formation based on the U.S. EPA's Water Treatment Plant Model (U.S. EPA, 2001). Required arguments include an object of class "water" created by [define_water](#) chlorine dose, type, reaction time, and treatment applied (if any). The function also requires additional water quality parameters defined in [define_water](#) including bromide, TOC, UV254, temperature, and pH. For a single water use `chemdose_dbp`; for a dataframe use `chemdose_dbp_chain`. For most arguments in the `_chain` helper "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```

chemdose_dbp(
  water,
  cl2,
  time,
  treatment = "raw",
  cl_type = "chlorine",
  location = "plant",
  correction = TRUE,
  coeff = NULL
)

chemdose_dbp_chain(
  df,
  input_water = "defined_water",

```

```

    output_water = "disinfected_water",
    cl2 = "use_col",
    time = "use_col",
    treatment = "use_col",
    cl_type = "use_col",
    location = "use_col",
    correction = TRUE,
    coeff = NULL
)

chemdose_dbp_once(
  df,
  input_water = "defined_water",
  cl2 = "use_col",
  time = "use_col",
  treatment = "use_col",
  cl_type = "use_col",
  location = "use_col",
  correction = TRUE,
  coeff = NULL,
  water_prefix = TRUE
)

```

Arguments

water	Source water object of class "water" created by define_water
cl2	Applied chlorine dose (mg/L as Cl ₂). Model results are valid for doses between 1.51 and 33.55 mg/L.
time	Reaction time (hours). Model results are valid for reaction times between 2 and 168 hours.
treatment	Type of treatment applied to the water. Options include "raw" for no treatment (default), "coag" for water that has been coagulated or softened, and "gac" for water that has been treated by granular activated carbon (GAC). GAC treatment has also been used for estimating formation after membrane treatment with good results.
cl_type	Type of chlorination applied, either "chlorine" (default) or "chloramine".
location	Location for DBP formation, either in the "plant" (default), or in the distributions system, "ds".
correction	Model calculations are adjusted based on location and cl_type. Default value is TRUE.
coeff	Optional input to specify custom coefficients to the dbp model. Must be a data frame with the following columns: ID, and the corresponding coefficients A, a, b, c, d, e, f, and ph_const for each dbp of interest. Default value is NULL.
df	a data frame containing a water class column, which has already been computed using define_water . The df may include columns for the other function arguments.

input_water	name of the column of water class data to be used as the input for this function. Default is "defined_water".
output_water	name of the output column storing updated parameters with the class, water. Default is "disinfected_water".
water_prefix	name of the input water used for the calculation, appended to the start of output columns. Default is TRUE. Change to FALSE to remove the water prefix from output column names.

Details

The function will calculate haloacetic acids (HAA) as HAA5, and total trihalomethanes (TTHM). Use `summarize_wq(water, params = c("dbps"))` to quickly tabulate the results.

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

`chemdose_dbp` returns a single water class object with predicted DBP concentrations.

`chemdose_dbp_chain` returns a data frame containing a water class column with predicted DBP concentrations.

`chemdose_dbp_once` returns a data frame containing predicted DBP concentrations as columns.

Source

TTHMs, raw: U.S. EPA (2001) equation 5-131

HAAs, raw: U.S. EPA (2001) equation 5-134

TTHMs, treated: U.S. EPA (2001) equation 5-139

HAAs, treated: U.S. EPA (2001) equation 5-142

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

Examples

```
example_dbp <- define_water(8, 20, 66, toc = 4, uv254 = .2, br = 50) %>%
  chemdose_dbp(cl2 = 2, time = 8)
example_dbp <- define_water(7.5, 20, 66, toc = 4, uv254 = .2, br = 50) %>%
  chemdose_dbp(cl2 = 3, time = 168, treatment = "coag", location = "ds")

library(dplyr)

example_df <- water_df %>%
  mutate(br = 50) %>%
  define_water_chain() %>%
```

```

    chemdose_dbp_chain(input_water = "defined_water", cl2 = 4, time = 8)

example_df <- water_df %>%
  mutate(br = 50) %>%
  slice_sample(n = 3) %>%
  define_water_chain() %>%
  mutate(
    cl2_dose = c(2, 3, 4),
    time = 30
  ) %>%
  chemdose_dbp_chain(cl2 = cl2_dose, treatment = "coag", location = "ds", cl_type = "chloramine")

# Initialize parallel processing
library(furrr)
# plan(multisession)
example_df <- water_df %>%
  mutate(br = 50) %>%
  define_water_chain() %>%
  chemdose_dbp_chain(cl2 = 4, time = 8)

# Optional: explicitly close multisession processing
# plan(sequential)

library(dplyr)

water <- water_df %>%
  slice(1) %>%
  mutate(br = 50) %>%
  define_water_chain() %>%
  chemdose_dbp_once(cl2 = 10, time = 8)

```

chemdose_ph

Calculate new pH and ion balance after chemical addition

Description

Calculates the new pH, alkalinity, and ion balance of a water based on different chemical additions. For a single water use `chemdose_ph`; for a dataframe use `chemdose_ph_chain`. Use [pluck_water](#) to get values from the output water as new dataframe columns. For most arguments in the `_chain` helper "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```

chemdose_ph(
  water,

```

```
hcl = 0,
h2so4 = 0,
h3po4 = 0,
hno3 = 0,
co2 = 0,
naoh = 0,
caoh2 = 0,
mgoh2 = 0,
na2co3 = 0,
nahco3 = 0,
caco3 = 0,
caso4 = 0,
caocl2 = 0,
cac12 = 0,
cl2 = 0,
naocl = 0,
nh4oh = 0,
nh42so4 = 0,
alum = 0,
ferricchloride = 0,
ferricsulfate = 0,
ach = 0,
kmno4 = 0,
naf = 0,
na3po4 = 0,
softening_correction = FALSE
)

chemdose_ph_chain(
  df,
  input_water = "defined_water",
  output_water = "dosed_chem_water",
  hcl = "use_col",
  h2so4 = "use_col",
  h3po4 = "use_col",
  hno3 = "use_col",
  co2 = "use_col",
  naoh = "use_col",
  na2co3 = "use_col",
  nahco3 = "use_col",
  caoh2 = "use_col",
  mgoh2 = "use_col",
  caocl2 = "use_col",
  cac12 = "use_col",
  cl2 = "use_col",
  naocl = "use_col",
  nh4oh = "use_col",
  nh42so4 = "use_col",
```

```

    caco3 = "use_col",
    caso4 = "use_col",
    alum = "use_col",
    ferricchloride = "use_col",
    ferricsulfate = "use_col",
    ach = "use_col",
    kmno4 = "use_col",
    naf = "use_col",
    na3po4 = "use_col",
    softening_correction = "use_col",
    na_to_zero = TRUE
)

```

```

chemdose_ph_once(
  df,
  input_water = "defined_water",
  hcl = "use_col",
  h2so4 = "use_col",
  h3po4 = "use_col",
  hno3 = "use_col",
  co2 = "use_col",
  naoh = "use_col",
  na2co3 = "use_col",
  nahco3 = "use_col",
  caoh2 = "use_col",
  mgoh2 = "use_col",
  caocl2 = "use_col",
  cac12 = "use_col",
  cl2 = "use_col",
  naocl = "use_col",
  nh4oh = "use_col",
  nh42so4 = "use_col",
  caco3 = "use_col",
  caso4 = "use_col",
  alum = "use_col",
  ferricchloride = "use_col",
  ferricsulfate = "use_col",
  ach = "use_col",
  kmno4 = "use_col",
  naf = "use_col",
  na3po4 = "use_col"
)

```

Arguments

water	Source water object of class "water" created by define_water
hcl	Amount of hydrochloric acid added in mg/L: $\text{HCl} \rightarrow \text{H} + \text{Cl}$
h2so4	Amount of sulfuric acid added in mg/L: $\text{H}_2\text{SO}_4 \rightarrow 2\text{H} + \text{SO}_4$

h3po4	Amount of phosphoric acid added in mg/L: $\text{H}_3\text{PO}_4 \rightarrow 3\text{H} + \text{PO}_4$
hno3	Amount of nitric acid added in mg/L: $\text{HNO}_3 \rightarrow \text{H} + \text{NO}_3$
co2	Amount of carbon dioxide added in mg/L: $\text{CO}_2 (\text{gas}) + \text{H}_2\text{O} \rightarrow \text{H}_2\text{CO}_3^*$
naoh	Amount of caustic added in mg/L: $\text{NaOH} \rightarrow \text{Na} + \text{OH}$
caoh2	Amount of lime added in mg/L: $\text{Ca}(\text{OH})_2 \rightarrow \text{Ca} + 2\text{OH}$
mgoh2	Amount of magnesium hydroxide added in mg/L: $\text{Mg}(\text{OH})_2 \rightarrow \text{Mg} + 2\text{OH}$
na2co3	Amount of soda ash added in mg/L: $\text{Na}_2\text{CO}_3 \rightarrow 2\text{Na} + \text{CO}_3$
nahco3	Amount of sodium bicarbonate added in mg/L: $\text{NaHCO}_3 \rightarrow \text{Na} + \text{H} + \text{CO}_3$
caco3	Amount of calcium carbonate added (or removed) in mg/L: $\text{CaCO}_3 \rightarrow \text{Ca} + \text{CO}_3$
caso4	Amount of calcium sulfate added (for post-RO condition) in mg/L: $\text{CaSO}_4 \rightarrow \text{Ca} + \text{SO}_4$
caocl2	Amount of Calcium hypochlorite added in mg/L as Cl_2 : $\text{CaOCl}_2 \rightarrow \text{Ca} + 2\text{OCl}$
cac12	Amount of calcium chloride added in mg/L: $\text{CaCl}_2 \rightarrow \text{Ca}^{2+} + 2\text{Cl}^-$
cl2	Amount of chlorine gas added in mg/L as Cl_2 : $\text{Cl}_2(\text{g}) + \text{H}_2\text{O} \rightarrow \text{HOCl} + \text{H} + \text{Cl}$
naocl	Amount of sodium hypochlorite added in mg/L as Cl_2 : $\text{NaOCl} \rightarrow \text{Na} + \text{OCl}$
nh4oh	Amount of ammonium hydroxide added in mg/L as N: $\text{NH}_4\text{OH} \rightarrow \text{NH}_4 + \text{OH}$
nh42so4	Amount of ammonium sulfate added in mg/L as N: $(\text{NH}_4)_2\text{SO}_4 \rightarrow 2\text{NH}_4 + \text{SO}_4$
alum	Amount of hydrated aluminum sulfate added in mg/L: $\text{Al}_2(\text{SO}_4)_3 \cdot 14\text{H}_2\text{O} + 6\text{HCO}_3 \rightarrow 2\text{Al}(\text{OH})_3(\text{am}) + 3\text{SO}_4 + 14\text{H}_2\text{O} + 6\text{CO}_2$
ferricchloride	Amount of ferric Chloride added in mg/L: $\text{FeCl}_3 + 3\text{HCO}_3 \rightarrow \text{Fe}(\text{OH})_3(\text{am}) + 3\text{Cl} + 3\text{CO}_2$
ferricsulfate	Amount of ferric sulfate added in mg/L: $\text{Fe}_2(\text{SO}_4)_3 \cdot 8.8\text{H}_2\text{O} + 6\text{HCO}_3 \rightarrow 2\text{Fe}(\text{OH})_3(\text{am}) + 3\text{SO}_4 + 8.8\text{H}_2\text{O} + 6\text{CO}_2$
ach	Amount of aluminum chlorohydrate added in mg/L: $\text{Al}_2(\text{OH})_5\text{Cl} \cdot 2\text{H}_2\text{O} + \text{HCO}_3 \rightarrow 2\text{Al}(\text{OH})_3(\text{am}) + \text{Cl} + 2\text{H}_2\text{O} + \text{CO}_2$
kmno4	Amount of potassium permanganate added in mg/L: $\text{KMnO}_4 \rightarrow \text{K} + \text{MnO}_4$
naf	Amount of sodium fluoride added in mg/L: $\text{NaF} \rightarrow \text{Na} + \text{F}$
na3po4	Amount of trisodium phosphate added in mg/L: $\text{Na}_3\text{PO}_4 \rightarrow 3\text{Na} + \text{PO}_4$
softening_correction	Set to TRUE to correct post-softening pH (caco3 must be < 0). Default is FALSE. Based on WTP model equation 5-62
df	a data frame containing a water class column, which has already been computed using define_water_chain The df may include columns named for the chemical(s) being dosed.
input_water	name of the column of water class data to be used as the input for this function. Default is "defined_water".
output_water	name of the output column storing updated parameters with the class, water. Default is "dosed_chem_water".
na_to_zero	option to convert all NA values in the data frame to zeros. Default value is TRUE.

Details

The function takes an object of class "water" created by [define_water](#) and user-specified chemical additions and returns a new object of class "water" with updated water quality. Units of all chemical additions are in mg/L as chemical (not as product).

chemdose_ph works by evaluating all the user-specified chemical additions and solving for what the new pH must be using [uniroot](#) to satisfy the principle of electroneutrality in pure water while correcting for the existing alkalinity of the water that the chemical is added to. Multiple chemicals can be added simultaneously or each addition can be modeled independently through sequential doses.

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

chemdose_ph returns a water class object with updated pH, alkalinity, and ions post-chemical addition.

chemdose_ph_chain returns a data frame containing a water class column with updated pH, alkalinity, and ions post-chemical addition.

chemdose_ph_once returns a data frame with columns for pH and alkalinity post-chemical addition.

See Also

[define_water](#), [convert_units](#)

Examples

```
water <- define_water(ph = 7, temp = 25, alk = 10)
# Dose 1 mg/L of hydrochloric acid
dosed_water <- chemdose_ph(water, hcl = 1)

# Dose 1 mg/L of hydrochloric acid and 5 mg/L of alum simultaneously
dosed_water <- chemdose_ph(water, hcl = 1, alum = 5)

# Softening:
water2 <- define_water(ph = 7, temp = 25, alk = 100, tot_hard = 350)
dosed_water2 <- chemdose_ph(water2, caco3 = -100, softening_correction = TRUE)

example_df <- water_df %>%
  define_water_chain() %>%
  dplyr::slice_head(n = 3) %>%
  dplyr::mutate(
    hcl = c(2, 4, 6),
    Caustic = 20
  ) %>%
```

```

chemdose_ph_chain(input_water = "defined_water", mgoh2 = c(20, 55), co2 = 4, naoh = Caustic)

# Initialize parallel processing
library(furrr)
# plan(multisession)
example_df <- water_df %>%
  define_water_chain() %>%
  chemdose_ph_chain(naoh = 5)

# Optional: explicitly close multisession processing
# plan(sequential)

example_df <- water_df %>%
  define_water_chain() %>%
  dplyr::slice_head(n = 3) %>%
  chemdose_ph_once(input_water = "defined_water", mgoh2 = 55, co2 = 4)

```

chemdose_toc

Determine TOC removal from coagulation

Description

This function applies the Edwards (1997) model to a water created by [define_water](#) to determine coagulated DOC. Model assumes all particulate TOC is removed; therefore TOC = DOC in output. Coagulated UVA is from U.S. EPA (2001) equation 5-80. Note that the models rely on pH of coagulation. If only raw water pH is known, utilize [chemdose_ph](#) first. For a single water use `chemdose_toc`; for a dataframe use `chemdose_toc_chain`. Use [pluck_water](#) to get values from the output water as new dataframe columns. For most arguments in the `_chain` helper "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```

chemdose_toc(
  water,
  alum = 0,
  ferricchloride = 0,
  ferricsulfate = 0,
  coeff = "Alum"
)

chemdose_toc_chain(
  df,
  input_water = "defined_water",
  output_water = "coagulated_water",

```

```

    alum = "use_col",
    ferricchloride = "use_col",
    ferricsulfate = "use_col",
    coeff = "use_col"
)

chemdose_toc_once(
  df,
  input_water = "defined_water",
  output_water = "coagulated_water",
  alum = "use_col",
  ferricchloride = "use_col",
  ferricsulfate = "use_col",
  coeff = "use_col"
)

```

Arguments

water	Source water object of class "water" created by define_water . Water must include ph, doc, and uv254
alum	Amount of hydrated aluminum sulfate added in mg/L: $\text{Al}_2(\text{SO}_4)_3 \cdot 14\text{H}_2\text{O} + 6\text{HCO}_3 \rightarrow 2\text{Al}(\text{OH})_3(\text{am}) + 3\text{SO}_4 + 14\text{H}_2\text{O} + 6\text{CO}_2$
ferricchloride	Amount of ferric chloride added in mg/L: $\text{FeCl}_3 + 3\text{HCO}_3 \rightarrow \text{Fe}(\text{OH})_3(\text{am}) + 3\text{Cl} + 3\text{CO}_2$
ferricsulfate	Amount of ferric sulfate added in mg/L: $\text{Fe}_2(\text{SO}_4)_3 \cdot 8.8\text{H}_2\text{O} + 6\text{HCO}_3 \rightarrow 2\text{Fe}(\text{OH})_3(\text{am}) + 3\text{SO}_4 + 8.8\text{H}_2\text{O} + 6\text{CO}_2$
coeff	String specifying the Edwards coefficients to be used from "Alum", "Ferric", "General Alum", "General Ferric", or "Low DOC" or data frame of coefficients, which must include: k1, k2, x1, x2, x3, b
df	a data frame containing a water class column, which has already been computed using define_water_chain The df may include columns named for the chemical(s) being dosed.
input_water	name of the column of water class data to be used as the input for this function. Default is "defined_water".
output_water	name of the output column storing updated parameters with the class, Water. Default is "coagulated_water".

Value

chemdose_toc returns a single water class object with an updated DOC, TOC, and UV254 concentration.

chemdose_toc_chain returns a data frame containing a water class column with updated DOC, TOC, and UV254 concentrations.

chemdose_toc_once returns a data frame with columns for updated TOC, DOC, and UV254.

Source

Edwards (1997)

U.S. EPA (2001)

See reference list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

See Also

[chemdose_ph](#)

Examples

```
water <- define_water(ph = 7, temp = 25, alk = 100, toc = 3.7, doc = 3.5, uv254 = .1)
dosed_water <- chemdose_ph(water, alum = 30) %>%
  chemdose_toc(alum = 30, coeff = "Alum")

dosed_water <- chemdose_ph(water, alum = 10, h2so4 = 10) %>%
  chemdose_toc(alum = 10, coeff = data.frame(
    x1 = 280, x2 = -73.9, x3 = 4.96, k1 = -0.028, k2 = 0.23, b = 0.068
  ))

example_df <- water_df %>%
  define_water_chain() %>%
  dplyr::mutate(FerricDose = seq(1, 12, 1)) %>%
  chemdose_toc_chain(ferricchloride = FerricDose, coeff = "Ferric")

# Uncomment below to initialize parallel processing
# library(furrr)
# plan(multisession)
example_df <- water_df %>%
  define_water_chain() %>%
  dplyr::mutate(ferricchloride = seq(1, 12, 1)) %>%
  chemdose_toc_chain(coeff = "Ferric")

# Optional: explicitly close multisession processing
# plan(sequential)

example_df <- water_df %>%
  define_water_chain() %>%
  chemdose_toc_once(input_water = "defined_water", alum = 30)
```

chloramine_conv

Data frame of conversion factors for estimating DBP formation from chloramines

Description

A dataset containing conversion factors for calculating DBP formation

Usage

```
chloramine_conv
```

Format

A dataframe with 17 rows and 3 columns

ID abbreviation of dbp species

alias full name of dbp species

percent specifies the percent of DBP formation predicted from chloramines compared to chlorine, assuming the same chlorine dose applied

Source

U.S. EPA (2001), Table 5-10

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

cl2coeffs

Data frame of Cl2 decay coefficients

Description

A dataset containing coefficients for calculating Cl2 decay

Usage

```
cl2coeffs
```

Format

A dataframe with 3 rows and 4 columns

treatment Specifies the treatment applied to the water

a Coefficient in chlorine decay model, associated with chlorine dose and time

b Coefficient in chlorine decay model, associated with chlorine dose & organics

c Exponent in chlorine decay model, associated with chlorine dose & organics

Source

U.S. EPA (2001)

convert_units	<i>Calculate unit conversions for common compounds</i>
---------------	--

Description

This function takes a value and converts units based on compound name.

Usage

```
convert_units(value, formula, startunit = "mg/L", endunit = "M")
```

Arguments

value	Value to be converted
formula	Chemical formula of compound. Accepts compounds in mweights for conversions between g and mol or eq
startunit	Units of current value, currently accepts g/L; g/L CaCO ₃ ; g/L N; M; eq/L; and the same units with "m", "u", "n" prefixes
endunit	Desired units, currently accepts same as start units

Value

A numeric value for the converted parameter.

Examples

```
convert_units(50, "ca") # converts from mg/L to M by default
convert_units(50, "ca", "mg/L", "mg/L CaCO3")
convert_units(50, "ca", startunit = "mg/L", endunit = "eq/L")
```

convert_water	<i>Convert water class object to a dataframe</i>
---------------	--

Description

This converts a water class to a dataframe with individual columns for each slot (water quality parameter) in the water. This is useful for one-off checks and is applied in all `fn_once` tidywater functions. For typical applications, there may be a `fn_once` tidywater function that provides a more efficient solution.

Use [convert_water](#) to keep all slots in the same units as the water.

Use [convert_watermg](#) to convert to more typical units. Converts the following slots from M to mg/L: na, ca, mg, k, cl, so4, hco3, co3, h2po4, hpo4, po4, ocl, bro3, f, fe, al. Converts these slots to ug/L: br, mn. All other values remain unchanged.

Usage

```
convert_water(water)

convert_watermg(water)
```

Arguments

water A water class object

Value

A data frame containing columns for all non-NA water slots.

A data frame containing columns for all non-NA water slots with ions in mg/L.

Examples

```
library(dplyr)
library(tidyr)

# Generates 1 row dataframe
example_df <- define_water(ph = 7, temp = 20, alk = 100) %>%
  convert_water()

example_df <- water_df %>%
  define_water_chain() %>%
  mutate(to_dataframe = map(defined_water, convert_water)) %>%
  unnest(to_dataframe) %>%
  select(-defined_water)

water_defined <- define_water(7, 20, 50, 100, 80, 10, 10, 10, 10, tot_po4 = 1) %>%
  convert_watermg()
```

correct_k

Correct acid dissociation constants

Description

This function calculates the corrected equilibrium constant for temperature and ionic strength

Usage

```
correct_k(water)
```

Arguments

water Defined water with values for temperature and ion concentrations

Value

A dataframe with equilibrium constants for co3, po4, so4, ocl, and nh4.

Examples

```
water_defined <- define_water(7, 20, 50, 100, 80, 10, 10, 10, 10, tot_po4 = 1)
correct_k(water_defined)
```

dbpcoeffs

Data frame of DBP coefficients for predicting DBP formation

Description

A dataset containing coefficients for calculating DBP formation

Usage

```
dbpcoeffs
```

Format

A dataframe with 30 rows and 10 columns

ID abbreviation of dbp species

alias full name of dbp species

water_type specifies which model the constants apply to, either treated or untreated water

A First coefficient in DBP model

a Second coefficient in DBP model, associated with TOC or DOC

b Third coefficient in DBP model, associated with Cl2

c Fourth coefficient in DBP model, associated with Br-

d Fifth coefficient in DBP model, associated with temperature

e Sixth coefficient in DBP model, associated with pH

f Seventh coefficient in DBP model, associated with reaction time

Source

U.S. EPA (2001)

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

dbp_correction	<i>Data frame of correction factors for estimating DBP formation as a function of location</i>
----------------	--

Description

A dataset containing correction factors for calculating DBP formation

Usage

```
dbp_correction
```

Format

A dataframe with 17 rows and 4 columns

ID abbreviation of dbp species

alias full name of dbp species

plant specifies the correction factor for modelling DBP formation within a treatment plant

ds specifies the correction factor for modelling DBP formation within the distribution system

Source

U.S. EPA (2001), Table 5-7

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

decarbonate_ph	<i>Apply decarbonation to a water</i>
----------------	---------------------------------------

Description

Calculates the new water quality (pH, alkalinity, etc) after a specified amount of CO₂ is removed (removed as bicarbonate). The function takes an object of class "water" and a fraction of CO₂ removed, then returns a water class object with updated water slots. For a single water, use decarbonate_ph; to apply the model to a dataframe, use decarbonate_ph_chain. For most arguments, the _chain helper "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```
decarbonate_ph(water, co2_removed)

decarbonate_ph_chain(
  df,
  input_water = "defined_water",
  output_water = "decarbonated_water",
  co2_removed = "use_col"
)
```

Arguments

water	Source water of class "water" created by define_water
co2_removed	Fraction of CO2 removed
df	a data frame containing a water class column, which has already been computed using define_water_chain . The df may include a column with names for each of the chemicals being dosed.
input_water	name of the column of water class data to be used as the input for this function. Default is "defined_water".
output_water	name of the output column storing updated parameters with the class, water. Default is "dosed_chem_water".

Details

decarbonate_ph uses water@h2co3 to determine the existing CO2 in water, then applies [chemdose_ph](#) to match the CO2 removal.

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

A water with updated pH/alk/etc.

`decarbonate_ph_chain` returns a data frame with a column containing a water with updated pH, alk, etc.

See Also

[chemdose_ph](#)

Examples

```
water <- define_water(ph = 4, temp = 25, alk = 5) %>%
  decarbonate_ph(co2_removed = .95)

example_df <- water_df %>%
  define_water_chain() %>%
  decarbonate_ph_chain(input_water = "defined_water", output_water = "decarb", co2_removed = .95)
```

define_water

Create a water class object given water quality parameters

Description

This function takes user-defined water quality parameters and creates an S4 "water" class object that forms the input and output of all tidywater models.

Usage

```
define_water(
  ph,
  temp = 25,
  alk,
  tot_hard,
  ca,
  mg,
  na,
  k,
  cl,
  so4,
  mno4,
  free_chlorine = 0,
  combined_chlorine = 0,
  tot_po4 = 0,
  tot_nh3 = 0,
  tds,
  cond,
  toc,
  doc,
  uv254,
  br,
  f,
  fe,
  al,
  mn,
  no3
)
```

Arguments

ph	water pH
temp	Temperature in degree C
alk	Alkalinity in mg/L as CaCO ₃
tot_hard	Total hardness in mg/L as CaCO ₃
ca	Calcium in mg/L Ca ²⁺
mg	Magnesium in mg/L Mg ²⁺
na	Sodium in mg/L Na ⁺
k	Potassium in mg/L K ⁺
cl	Chloride in mg/L Cl ⁻
so4	Sulfate in mg/L SO ₄ ²⁻
mno4	Permanganate in mg/L MnO ₄ ⁻
free_chlorine	Free chlorine in mg/L as Cl ₂ . Used when a starting water has a free chlorine residual.
combined_chlorine	Combined chlorine (chloramines) in mg/L as Cl ₂ . Used when a starting water has a chloramine residual.
tot_po4	Phosphate in mg/L as PO ₄ ³⁻ . Used when a starting water has a phosphate residual.
tot_nh3	Total ammonia in mg/L as N
tds	Total Dissolved Solids in mg/L (optional if ions are known)
cond	Electrical conductivity in uS/cm (optional if ions are known)
toc	Total organic carbon (TOC) in mg/L
doc	Dissolved organic carbon (DOC) in mg/L
uv254	UV absorbance at 254 nm (cm ⁻¹)
br	Bromide in ug/L Br ⁻
f	Fluoride in mg/L F ⁻
fe	Iron in mg/L Fe ³⁺
al	Aluminum in mg/L Al ³⁺
mn	Manganese in ug/L Mn ²⁺
no3	Nitrate in mg/L as N

Details

Carbonate balance is calculated and units are converted to mol/L. Ionic strength is determined from ions, TDS, or conductivity. Missing values are handled by defaulting to 0 or NA. Calcium defaults to 65 percent of the total hardness when not specified. DOC defaults to 95 percent of TOC.

Value

define_water outputs a water class object where slots are filled or calculated based on input parameters. Water slots have different units than those input into the define_water function, as listed below.

pH pH, numeric, in standard units (SU).
temp temperature, numeric, in °C.
alk alkalinity, numeric, mg/L as CaCO₃.
tds total dissolved solids, numeric, mg/L.
cond electrical conductivity, numeric, uS/cm.
tot_hard total hardness, numeric, mg/L as CaCO₃.
kw dissociation constant for water, numeric, unitless.
alk_eq alkalinity as equivalents, numeric, equivalent (eq).
toc total organic carbon, numeric, mg/L.
doc dissolved organic carbon, numeric, mg/L.
bdoc biodegradable organic carbon, numeric, mg/L.
uv254 light absorption at 254 nm, numeric, cm⁻¹.
dic dissolved inorganic carbon, numeric, mg/L as C.
is ionic strength, numeric, mol/L.
na sodium, numeric, mols/L.
ca calcium, numeric, mols/L.
mg magnesium, numeric, mols/L.
k potassium, numeric, mols/L.
cl chloride, numeric, mols/L.
so4 sulfate, numeric, mols/L.
mno4 permanganate, numeric, mols/L.
no3 nitrate, numeric, mols/L.
hco3 bicarbonate, numeric, mols/L.
co3 carbonate, numeric, mols/L.
h2po4 phosphoric acid, numeric, mols/L.
hpo4 hydrogen phosphate, numeric, mols/L.
po4 phosphate, numeric, mols/L.
nh4 ammonium, numeric, mol/L as N.
h hydrogen ion, numeric, mol/L.
oh hydroxide ion, numeric, mol/L.
tot_po4 total phosphate, numeric, mol/L.
tot_nh3 total ammonia, numeric, mol/L.
tot_co3 total carbonate, numeric, mol/L.

br bromide, numeric, mol/L.
bro3 bromate, numeric, mol/L.
f fluoride, numeric, mol/L.
fe iron, numeric, mol/L.
al aluminum, numeric, mol/L.
mn manganese, numeric, mol/L.
free_chlorine free chlorine, numeric, mol/L.
ocl hypochlorite ion, numeric, mol/L.
combined_chlorine sum of chloramines, numeric, mol/L.
nh2cl monochloramine, numeric, mol/L.
nhcl2 dichloramine, numeric, mol/L.
ncl3 trichloramine, numeric, mol/L.
chcl3 chloroform, numeric, ug/L.
chcl2br bromodichloromethane, numeric, ug/L.
chbr2cl dibromodichloromethane, numeric, ug/L.
chbr3 bromoform, numeric, ug/L.
tthm total trihalomethanes, numeric, ug/L.
mcaa chloroacetic acid, numeric, ug/L.
dmcaa dichloroacetic acid, numeric, ug/L.
tcaa trichloroacetic acid, numeric, ug/L.
mbaa bromoacetic acid, numeric, ug/L.
dbaa dibromoacetic acid, numeric, ug/L.
haa5 sum of haloacetic acids, numeric, ug/L.
bcaa bromochloroacetic acid, numeric, ug/L.
cdbaa chlorodibromoacetic acid, numeric, ug/L.
dcbaa dichlorobromoacetic acid, numeric, ug/L.
tbaa tribromoacetic acid, numeric, ug/L.

Source

Crittenden et al. (2012) equation 5-38 - ionic strength from TDS
 Snoeyink & Jenkins (1980) - ionic strength from conductivity
 Lewis and Randall (1921), Crittenden et al. (2012) equation 5-37 - ionic strength from ion concentrations
 Harned and Owen (1958), Crittenden et al. (2012) equation 5-45 - Temperature correction of dielectric constant (relative permittivity)

Examples

```

water_missingions <- define_water(ph = 7, temp = 15, alk = 100, tds = 10)
water_defined <- define_water(7, 20, 50, 100, 80, 10, 10, 10, 10, tot_po4 = 1)

```

define_water_chain	<i>Apply define_water within a dataframe and output a column of water class to be chained to other tidywater functions</i>
--------------------	--

Description

This function allows `define_water` to be added to a piped data frame. Its output is a water class, and can therefore be chained with "downstream" tidywater functions.

Usage

```
define_water_chain(df, output_water = "defined_water")
```

Arguments

df	a data frame containing columns with all the parameters listed in <code>define_water</code>
output_water	name of the output column storing updated parameters with the class, water. Default is "defined_water".

Details

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

A data frame containing a water class column.

See Also

`define_water`

Examples

```
example_df <- water_df %>%  
  define_water_chain() %>%  
  balance_ions_chain()  
  
example_df <- water_df %>%  
  define_water_chain(output_water = "This is a column of water") %>%  
  balance_ions_chain(input_water = "This is a column of water")  
  
# Initialize parallel processing  
library(furrr)
```

```
# plan(multisession)
example_df <- water_df %>%
  define_water_chain() %>%
  balance_ions_chain()

#' #Optional: explicitly close multisession processing
# plan(sequential)
```

define_water_once	<i>Apply define_water and output a dataframe</i>
-------------------	--

Description

This function allows [define_water](#) to be added to a piped data frame. It outputs all carbonate calculations and other parameters in a data frame. tidywater functions cannot be added after this function because they require a water class input.

Usage

```
define_water_once(df)
```

Arguments

df a data frame containing columns with all the parameters listed in [define_water](#)

Details

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

A data frame containing columns that were filled or calculated based on `define_water`.

See Also

[define_water](#)

Examples

```
example_df <- water_df %>%
  define_water_once()
```

discons*Dissociation constants and standard enthalpy for weak acids/bases*

Description

Equilibrium constants (k) and corresponding standard enthalpy of reaction values (deltah) for significant acids in water influencing pH at equilibrium. Includes carbonate, sulfate, phosphate, and hypochlorite. Standard enthalpy of reaction is calculated by taking the sum of the enthalpy of formation of each individual component minus the enthalpy of formation of the final product. e.g., the standard enthalpy of reaction for water can be calculated as: $\text{deltah_h2o} = \text{deltah_f_oh} + \text{deltah_f_h} - \text{deltah_f_h2o} = -230 + 0 - (-285.83) = 55.83 \text{ kJ/mol}$. See MWH (2012) example 5-5 and Benjamin (2002) eq. 2.96.

Usage

```
discons
```

Format

A dataframe with 8 rows and 3 columns

ID Coefficient type

k Equilibrium constant

deltah Standard enthalpy in J/mol

Source

Benjamin (2015) Appendix A.1 and A.2.

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

dissolve_cu*Calculate Dissolved Copper Concentration*

Description

This function takes a water defined by defined_water and output a column of dissolved copper. It is an empirical model developed based on bench-scale copper solubility testing that can be used to predict copper levels as a function of pH, DIC, and orthophosphate. For a single water, use dissolve_cu; to apply the model to a dataframe use dissolve_cu_chain.

Usage

```
dissolve_cu(water)
```

Arguments

water Source water object of class "water" created by [define_water](#). Water must include ph and dic

Details

Dissolved copper is a function of pH, DIC, and PO4. Output units are in mg/L. For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

`dissolve_cu` returns a column containing dissolved copper concentration in mg/L.

Source

Lytle et al (2018)

Examples

```
example_cu <- define_water(ph = 7.5, alk = 125, tot_po4 = 2) %>%
  dissolve_cu()
```

dissolve_cu_once	<i>Calculate Dissolved Copper Concentration</i>
------------------	---

Description

Calculate Dissolved Copper Concentration

Usage

```
dissolve_cu_once(df, input_water = "defined_water")
```

Arguments

df a data frame containing a water class column, which has already been computed using [define_water_chain](#)

input_water name of the column of Water class data to be used as the input for this function. Default is "defined_water".

Value

dissolve_cu_once returns a data frame containing the original data frame and a column for dissolved copper in mg/L.

Examples

```
library(dplyr)
cu_calc <- water_df %>%
  define_water_chain() %>%
  dissolve_cu_once()
```

dissolve_pb	<i>Simulate contributions of various lead solids to total soluble lead</i>
-------------	--

Description

This function takes a water data frame defined by [define_water](#) and outputs a dataframe of the controlling lead solid and total lead solubility. Lead solid solubility is calculated based on controlling solid. Total dissolved lead species (tot_dissolved_pb, M) are calculated based on lead complex calculations. For a single water, use dissolve_pb; to apply the model to a dataframe, use dissolve_pb_once. For most arguments, the _chain and _once helpers "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```
dissolve_pb(
  water,
  hydroxypyromorphite = "Schock",
  pyromorphite = "Topolska",
  laurionite = "Nasanen"
)

dissolve_pb_once(
  df,
  input_water = "defined_water",
  output_col_solid = "controlling_solid",
  output_col_result = "pb",
  hydroxypyromorphite = "Schock",
  pyromorphite = "Topolska",
  laurionite = "Nasanen",
  water_prefix = TRUE
)
```

Arguments

water	Source water object of class "water" created by define_water . Water must include alk and is. If po4, cl, and so4 are known, those should also be included.
hydroxypyromorphite	defaults to "Schock", the constant, K, developed by Schock et al (1996). Can also use "Zhu".
pyromorphite	defaults to "Topolska", the constant, K, developed by Topolska et al (2016). Can also use "Xie".
laurionite	defaults to "Nasanen", the constant, K, developed by Nasanen & Lindell (1976). Can also use "Lothenbach".
df	a data frame containing a water class column, which has already been computed using define_water_chain
input_water	name of the column of water class data to be used as the input. Default is "defined_water".
output_col_solid	name of the output column storing the controlling lead solid. Default is "controlling_solid".
output_col_result	name of the output column storing dissolved lead in M. Default is "pb".
water_prefix	name of the input water used for the calculation, appended to the start of output columns. Default is TRUE. Change to FALSE to remove the water prefix from output column names.

Details

The solid with lowest solubility will form the lead scale (controlling lead solid). Some lead solids have two k-constant options. The function will default to the EPA's default constants. The user may change the constants to hydroxypyromorphite = "Zhu" or pyromorphite = "Xie" or laurionite = "Lothenbach"

Make sure that total dissolved solids, conductivity, or ca, na, cl, so4 are used in `define_water` so that an ionic strength is calculated.

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

`dissolve_pb` returns a one row data frame containing only the controlling lead solid and modeled dissolved lead concentration.

`dissolve_pb_once` returns a data frame containing the controlling lead solid and modeled dissolved lead concentration as new columns.

Source

Code is from EPA's TELSS lead solubility dashboard <https://github.com/USEPA/TELSS> which is licensed under MIT License: Permission is hereby granted, free of charge, to any person obtaining a copy of this software and associated documentation files (the "Software"), to deal in the Software without restriction, including without limitation the rights to use, copy, modify, merge, publish, distribute, sublicense, and/or sell copies of the Software, and to permit persons to whom the Software is furnished to do so, subject to the following conditions: The above copyright notice and this permission notice shall be included in all copies or substantial portions of the Software.

Wahman et al. (2021)

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

Examples

```
example_pb <- define_water(
  ph = 7.5, temp = 25, alk = 93, cl = 240,
  tot_po4 = 0, so4 = 150, tds = 200
) %>%
  dissolve_pb()
example_pb <- define_water(
  ph = 7.5, temp = 25, alk = 93, cl = 240,
  tot_po4 = 0, so4 = 150, tds = 200
) %>%
  dissolve_pb(pyromorphite = "Xie")

example_df <- water_df %>%
  define_water_chain() %>%
  dissolve_pb_once(output_col_result = "dissolved_lead", pyromorphite = "Xie")

# Initialize parallel processing
library(furrr)
# plan(multisession)
example_df <- water_df %>%
  define_water_chain() %>%
  dissolve_pb_once(output_col_result = "dissolved_lead", laurionite = "Lothenbach")

# Optional: explicitly close multisession processing
# plan(sequential)
```

edwardscoeff

Data frame of Edwards model coefficients

Description

A dataset containing coefficients from the Edwards (1997) model for coagulation TOC removal.

Usage

```
edwardscoeff
```

Format

A dataframe with 5 rows and 7 columns:

ID Coefficient type

x3 x3 parameter

x2 x2 parameter

x1 x1 parameter

k1 k1 parameter

k2 k2 parameter

b b parameter

Source

Edwards (1997) Table 2.

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

leadsol_constants	<i>Data frame of equilibrium constants for lead and copper solubility</i>
-------------------	---

Description

A dataset containing equilibrium constants for lead solubility

Usage

```
leadsol_constants
```

Format

A dataframe with 38 rows and 3 columns

Solids:

species_name Name of lead solid or complex with possible _letter to cite different references

constant_name Reference ID for constants

log_value Equilibrium constant log value

source Source for equilibrium constant value

Source

Benjamin (2010)

Lothenbach et al. (1999)

Nasanen & Lindell (1976)

Powell et al. (2009)

Powell et al. (2005)

Schock et al. (1996)

Topolska et al. (2016)

Xie & Giammar (2007)

Zhu et al. (2015)

Wahman et al. (2021)

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

modify_water

Modify a single slot in a water class object

Description

This function a single slot of a water class object without impacting the other parameters. For example, you can manually update "tthm" and the new speciation will not be calculated. This function is designed to make sure all parameters are stored in the correct units when manually updating a water. Some slots cannot be modified with this function because they are interconnected with too many others (usually pH dependent, eg, hco3). For those parameters, update [define_water](#).

Usage

```
modify_water(water, slot, value, units)
```

```
modify_water_chain(  
  df,  
  input_water = "defined_water",  
  output_water = "modified_water",  
  slot = "use_col",  
  value = "use_col",  
  units = "use_col"  
)
```

Arguments

water	A water class object
slot	A character string of the slot in the water to modify, eg, "tthm"
value	New value for the modified slot

units	Units of the value being entered, typically one of c("mg/L", "ug/L", "M", "cm-1"). For ions any units supported by convert_units are allowed. For organic carbon, one of "mg/L", "ug/L". For uv254 one of "cm-1", "m-1". For DBPs, one of "ug/L" or "mg/L".
df	a data frame containing a water class column, which has already been computed using define_water_chain
input_water	name of the column of water class data to be used as the input for this function. Default is "defined_water".
output_water	name of the output column storing updated parameters with the class, water. Default is "modified_water".

Value

A data frame containing columns of selected parameters from a list of water class objects.

`modify_water_chain` returns a data frame containing a water class column with updated slot

Examples

```
water1 <- define_water(ph = 7, alk = 100, tds = 100, toc = 5) %>%
  modify_water(slot = "toc", value = 4, units = "mg/L")
```

```
library(dplyr)
```

```
example_df <- water_df %>%
  define_water_chain() %>%
  modify_water_chain(slot = "br", value = 50, units = "ug/L")
```

```
# Un-comment below to initialize parallel processing
# library(furrr)
# plan(multisession)
example_df <- water_df %>%
  define_water_chain() %>%
  mutate(bromide = rep(c(20, 30, 50), 4)) %>%
  modify_water_chain(slot = "br", value = bromide, units = "ug/L")

# Optional: explicitly close multisession processing
# plan(sequential)
```

mweights

Molar weights of relevant compounds

Description

A dataset containing the molar weights of several compounds in g/mol. Column names are lower-case chemical formulas (with no charge), with the exception of the following coagulants: `alum` = $\text{Al}_2(\text{SO}_4)_3 \cdot 14\text{H}_2\text{O}$, `ferricchloride` = FeCl_3 , `ferricsulfate` = $\text{Fe}_2(\text{SO}_4)_3 \cdot 8\text{H}_2\text{O}$,

Usage

```
mweights
```

Format

A dataframe with one row and one column per compound

ozonate_bromate	<i>Calculate bromate formation</i>
-----------------	------------------------------------

Description

Calculates bromate (BrO₃⁻, ug/L) formation based on selected model. Required arguments include an object of class "water" created by [define_water](#) ozone dose, reaction time, and desired model. The function also requires additional water quality parameters defined in [define_water](#) including bromide, DOC or UV254 (depending on the model), pH, alkalinity (depending on the model), and optionally, ammonia (added when defining water using the tot_nh3 argument.) For a single water use `ozonate_bromate`; for a dataframe use `ozonate_bromate_chain`. Use [pluck_water](#) to get values from the output water as new dataframe columns. For most arguments in the `_chain` helper "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```
ozonate_bromate(water, dose, time, model = "Ozekin")
```

```
ozonate_bromate_chain(
  df,
  input_water = "defined_water",
  output_water = "ozonated_water",
  dose = "use_col",
  time = "use_col",
  model = "use_col"
)
```

Arguments

water	Source water object of class "water" created by define_water
dose	Applied ozone dose (mg/L as O ₃). Results typically valid for 1-10 mg/L, but varies depending on model.
time	Reaction time (minutes). Results typically valid for 1-120 minutes, but varies depending on model.
model	Model to apply. One of c("Ozekin", "Sohn", "Song", "Galey", "Siddiqui")
df	a data frame containing a water class column, which has already been computed using define_water_once . The df may include a column named for the applied chlorine dose (cl2), and a column for time in minutes.

input_water	name of the column of water class data to be used as the input for this function. Default is "defined_water".
output_water	name of the output column storing updated parameters with the class, water. Default is "ozonated_water".

Details

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

`ozonate_bromate` returns a single water class object with calculated bromate (ug/L).

`ozonate_bromate_chain` returns a data frame containing a water class column with updated bro3.

Source

Ozekin (1994), Sohn et al (2004), Song et al (1996), Galey et al (1997), Siddiqui et al (1994)

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

Examples

```
example_dbp <- define_water(8, 20, 66, toc = 4, uv254 = .2, br = 50) %>%
  ozonate_bromate(dose = 1.5, time = 5, model = "Ozekin")
example_dbp <- define_water(7.5, 20, 66, toc = 4, uv254 = .2, br = 50) %>%
  ozonate_bromate(dose = 3, time = 15, model = "Sohn")
```

```
library(dplyr)
```

```
example_df <- water_df %>%
  slice_head(n = 6) %>%
  mutate(br = 50) %>%
  define_water_chain() %>%
  mutate(
    dose = c(seq(.5, 3, .5)),
    OzoneTime = 30
  ) %>%
  ozonate_bromate_chain(time = OzoneTime, model = "Sohn")
```

```
# Initialize parallel processing
library(furrr)
# plan(multisession)
example_df <- water_df %>%
  mutate(br = 50) %>%
```

```

define_water_chain() %>%
  ozonate_bromate_chain(dose = 4, time = 8)

# Optional: explicitly close multiseession processing
# plan(sequential)

```

pactocoeffs	<i>Data frame of PAC TOC model coefficients</i>
-------------	---

Description

A dataset containing coefficients for calculating PAC TOC removal

Usage

```
pactocoeffs
```

Format

A dataframe with 4 rows and 3 columns

pactype Specifies PAC type

A Constant in the PAC model

a Coefficient in PAC model, associated with DOC0

b Coefficient in PAC model, associated with dose

c Coefficient in PAC model, associated with time

Source

Cho (2007)

pac_toc	<i>Calculate DOC Concentration in PAC system</i>
---------	--

Description

Calculates DOC concentration multiple linear regression model found in 2-METHYLISOBORNEOL AND NATURAL ORGANIC MATTER ADSORPTION BY POWDERED ACTIVATED CARBON by HYUKJIN CHO (2007). Assumes all particulate TOC is removed when PAC is removed; therefore TOC = DOC in output. For a single water use pac_toc; for a dataframe use pac_toc_chain. Use [pluck_water](#) to get values from the output water as new dataframe columns. For most arguments in the _chain helper "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

water must contain DOC or TOC value.

Usage

```
pac_toc(water, dose, time, type = "bituminous")

pac_toc_chain(
  df,
  input_water = "defined_water",
  output_water = "pac_water",
  dose = "use_col",
  time = "use_col",
  type = "use_col"
)
```

Arguments

water	Source water object of class "water" created by define_water
dose	Applied PAC dose (mg/L). Model results are valid for doses concentrations between 5 and 30 mg/L.
time	Contact time (minutes). Model results are valid for reaction times between 10 and 1440 minutes
type	Type of PAC applied, either "bituminous", "lignite", "wood".
df	a data frame containing a water class column, which has already been computed using define_water_chain . The df may include columns named for the dose, time, and type
input_water	name of the column of water class data to be used as the input for this function. Default is "defined_water".
output_water	name of the output column storing updated parameters with the class, water. Default is "pac_water".

Details

The function will calculate DOC concentration by PAC adsorption in drinking water treatment. UV254 concentrations are predicted based on a linear relationship with DOC.

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

`pac_toc` returns a water class object with updated DOC, TOC, and UV254 slots.

`pac_toc_chain` returns a data frame containing a water class column with updated DOC, TOC, and UV254 slots

Source

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>
CHO(2007)

Examples

```
water <- define_water(toc = 2.5, uv254 = .05, doc = 1.5) %>%
  pac_toc(dose = 15, time = 50, type = "wood")

library(dplyr)

example_df <- water_df %>%
  define_water_chain("raw") %>%
  mutate(dose = seq(11, 22, 1), PACTime = 30) %>%
  pac_toc_chain(input_water = "raw", time = PACTime, type = "wood")

# Initialize parallel processing
library(furrr)
# plan(multisession)
example_df <- water_df %>%
  define_water_chain("raw") %>%
  pac_toc_chain(input_water = "raw", dose = 4, time = 8)

# Optional: explicitly close multisession processing
# plan(sequential)
```

plot_ions

Create summary plot of ions from water class

Description

This function takes a water data frame defined by `define_water` and outputs an ion balance plot.

Usage

```
plot_ions(water)
```

Arguments

water Source water vector created by link function here

Value

A ggplot object displaying the water's ion balance.

Examples

```
water <- define_water(7, 20, 50, 100, 20, 10, 10, 10, tot_po4 = 1)
plot_ions(water)
```

pluck_water

Pluck out a single parameter from a water class object

Description

This function plucks one or more selected parameters from selected columns of water class objects. The names of the output columns will follow the form water_parameter

Usage

```
pluck_water(df, input_waters = c("defined_water"), parameter)
```

Arguments

df	a data frame containing a water class column, which has already been computed using define_water
input_waters	vector of names of the columns of water class data to be used as the input for this function.
parameter	vector of water class parameters to view outside the water column. Can also specify "all" to get all non-NA water slots.

Value

A data frame containing columns of selected parameters from a list of water class objects.

See Also

[convert_water](#)

Examples

```
pluck_example <- water_df %>%
  define_water_chain("raw") %>%
  pluck_water(input_waters = c("raw"), parameter = c("hco3", "doc"))
```

```
library(furrr)
# plan(multisession)
pluck_example <- water_df %>%
  define_water_chain() %>%
  pluck_water(parameter = c("ph", "alk"))
```

```
# Optional: explicitly close multisession processing
```

```
# plan(sequential)
```

solvecost_chem	<i>Determine chemical cost</i>
----------------	--------------------------------

Description

This function takes a chemical dose in mg/L, plant flow, chemical strength, and \$/lb and calculates cost.

Usage

```
solvecost_chem(dose, flow, strength = 100, cost, time = "day")
```

Arguments

dose	Chemical dose in mg/L as chemical
flow	Plant flow in MGD
strength	Chemical product strength in percent. Defaults to 100 percent.
cost	Chemical product cost in \$/lb
time	Desired output units, one of c("day", "month", "year"). Defaults to "day".

Value

A numeric value for chemical cost, \$/time.

Examples

```
alum_cost <- solvecost_chem(dose = 20, flow = 10, strength = 49, cost = .22)

library(dplyr)
cost_data <- tibble(
  dose = seq(10, 50, 10),
  flow = 10
) %>%
  mutate(costs = solvecost_chem(dose = dose, flow = flow, strength = 49, cost = .22))
```

solvecost_labor	<i>Determine labor cost</i>
-----------------	-----------------------------

Description

This function takes number of FTE and annual \$/FTE and determines labor cost

Usage

```
solvecost_labor(fte, cost, time = "day")
```

Arguments

fte	Number of FTEs. Can be decimal.
cost	\$/year per FTE
time	Desired output units, one of c("day", "month", "year"). Defaults to "day".

Value

A numeric value for labor \$/time.

Examples

```
laborcost <- solvecost_labor(1.5, 50000)

library(dplyr)
cost_data <- tibble(
  fte = seq(1, 10, 1)
) %>%
  mutate(costs = solvecost_labor(fte = fte, cost = .08))
```

solvecost_power	<i>Determine power cost</i>
-----------------	-----------------------------

Description

This function takes kW, % utilization, \$/kWhr and determines power cost.

Usage

```
solvecost_power(power, utilization = 100, cost, time = "day")
```

Arguments

power	Power consumed in kW
utilization	Amount of time equipment is running in percent. Defaults to continuous.
cost	Power cost in \$/kWhr
time	Desired output units, one of c("day", "month", "year"). Defaults to "day".

Value

A numeric value for power, \$/time.

Examples

```
powercost <- solvecost_power(50, 100, .08)

library(dplyr)
cost_data <- tibble(
  power = seq(10, 50, 10),
  utilization = 80
) %>%
  mutate(costs = solvecost_power(power = power, utilization = utilization, cost = .08))
```

solvecost_solids	<i>Determine solids disposal cost</i>
------------------	---------------------------------------

Description

This function takes coagulant doses in mg/L as chemical, removed turbidity, and cost (\$/lb) to determine disposal cost.

Usage

```
solvecost_solids(
  alum = 0,
  ferricchloride = 0,
  ferricsulfate = 0,
  flow,
  turb,
  b = 1.5,
  cost,
  time = "day"
)
```

Arguments

alum	Hydrated aluminum sulfate $\text{Al}_2(\text{SO}_4)_3 \cdot 14\text{H}_2\text{O} + 6\text{HCO}_3 \rightarrow 2\text{Al}(\text{OH})_3(\text{am}) + 3\text{SO}_4 + 14\text{H}_2\text{O} + 6\text{CO}_2$
ferricchloride	Ferric Chloride $\text{FeCl}_3 + 3\text{HCO}_3 \rightarrow \text{Fe}(\text{OH})_3(\text{am}) + 3\text{Cl} + 3\text{CO}_2$
ferricsulfate	Amount of ferric sulfate added in mg/L: $\text{Fe}_2(\text{SO}_4)_3 \cdot 8.8\text{H}_2\text{O} + 6\text{HCO}_3 \rightarrow 2\text{Fe}(\text{OH})_3(\text{am}) + 3\text{SO}_4 + 8.8\text{H}_2\text{O} + 6\text{CO}_2$
flow	Plant flow in MGD
turb	Turbidity removed in NTU
b	Correlation factor from turbidity to suspended solids. Defaults to 1.5.
cost	Disposal cost in \$/lb
time	Desired output units, one of c("day", "month", "year"). Defaults to "day".

Value

A numeric value for disposal costs, \$/time.

Source

<https://water.mecc.edu/courses/ENV295Residuals/lesson3b.htm#:~:text=From%20the%20diagram%2C%20for%20example>

Examples

```
alum_solidscost <- solvecost_solids(alum = 50, flow = 10, turb = 2, cost = 0.05)

library(dplyr)
cost_data <- tibble(
  alum = seq(10, 50, 10),
  flow = 10
) %>%
  mutate(costs = solvecost_solids(alum = alum, flow = flow, turb = 2, cost = 0.05))
```

solvect_chlorine

Determine disinfection credit from chlorine.

Description

This function takes a water defined by [define_water](#) and other disinfection parameters and outputs a data frame of the required CT (ct_required), actual CT (ct_actual), and giardia log removal (glog_removal). For a single water, use solvect_chlorine; to apply the model to a dataframe, use solvect_chlorine_once. For most arguments, the _chain and _once helpers "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```

solvect_chlorine(water, time, residual, baffle, free_cl_slot = "residual_only")

solvect_chlorine_once(
  df,
  input_water = "defined_water",
  time = "use_col",
  residual = "use_col",
  baffle = "use_col",
  free_cl_slot = "residual_only",
  water_prefix = TRUE
)

```

Arguments

water	Source water object of class "water" created by define_water . Water must include ph and temp
time	Retention time of disinfection segment in minutes.
residual	Minimum chlorine residual in disinfection segment in mg/L as Cl ₂ .
baffle	Baffle factor - unitless value between 0 and 1.
free_cl_slot	Defaults to "residual_only", which uses the residual argument. If "slot_only", the model will use the free_chlorine slot in the input water. "sum_with_residual", will use the sum of the residual argument and the free_chlorine slot.
df	a data frame containing a water class column, which has already been computed using define_water_chain
input_water	name of the column of Water class data to be used as the input for this function. Default is "defined_water".
water_prefix	name of the input water used for the calculation will be appended to the start of output columns. Default is TRUE.

Details

CT actual is a function of time, chlorine residual, and baffle factor, whereas CT required is a function of pH, temperature, chlorine residual, and the standard 0.5 log removal of giardia requirement. CT required is an empirical regression equation developed by Smith et al. (1995) to provide conservative estimates for CT tables in USEPA Disinfection Profiling Guidance. Log removal is a rearrangement of the CT equations.

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

`solvect_chlorine` returns a data frame containing required CT (mg/Lmin), *actual CT* (mg/Lmin), and giardia log removal.

`solvect_chlorine_once` returns a data frame containing the original data frame and columns for required CT, actual CT, and giardia log removal.

Source

Smith et al. (1995)

USEPA (2020)

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

Examples

```
example_ct <- define_water(ph = 7.5, temp = 25) %>%
  solvect_chlorine(time = 30, residual = 1, baffle = 0.7)
library(dplyr)
ct_calc <- water_df %>%
  define_water_chain() %>%
  solvect_chlorine_once(residual = 2, time = 10, baffle = .5)

chlor_resid <- water_df %>%
  mutate(br = 50) %>%
  define_water_chain() %>%
  mutate(
    residual = seq(1, 12, 1),
    time = seq(2, 24, 2),
    baffle = 0.7
  ) %>%
  solvect_chlorine_once()
```

solvect_o3

Determine disinfection credit from ozone.

Description

This function takes a water defined by `define_water()` and the first order decay curve parameters from an ozone dose and outputs a dataframe of actual CT, and log removal for giardia, virus, and crypto. For a single water, use `solvect_o3`; to apply the model to a dataframe, use `solvect_o3_once`. For most arguments, the `_once` helper "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```
solvect_o3(water, time, dose, kd, baffle)
```

```
solvect_o3_once(
  df,
  input_water = "defined_water",
  time = "use_col",
  dose = "use_col",
  kd = "use_col",
  baffle = "use_col",
  water_prefix = TRUE
)
```

Arguments

water	Source water object of class "water" created by define_water() . Water must include ph and temp
time	Retention time of disinfection segment in minutes.
dose	Ozone dose in mg/L. This value can also be the y intercept of the decay curve (often slightly lower than ozone dose.)
kd	First order decay constant. This parameter is optional. If not specified, the default ozone decay equations will be used.
baffle	Baffle factor - unitless value between 0 and 1.
df	a data frame containing a water class column, which has already been computed using define_water_chain() .
input_water	name of the column of Water class data to be used as the input for this function. Default is "defined_water".
water_prefix	name of the input water used for the calculation will be appended to the start of output columns. Default is TRUE.

Details

First order decay curve for ozone has the form: $\text{residual} = \text{dose} * \exp(\text{kd} * \text{time})$. kd should be a negative number. Actual CT is an integration of the first order curve. The first 30 seconds are removed from the integral to account for instantaneous demand.

When kd is not specified, a default decay curve is used from the Water Treatment Plant Model (2002). This model does not perform well for ozone decay, so specifying the decay curve is recommended.

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

`solvect_o3` returns a data frame containing actual CT (mg/L*min), giardia log removal, virus log removal, and crypto log removal.

`solvect_o3_once` returns a data frame containing the original data frame and columns for required CT, actual CT, and giardia log removal.

Source

USEPA (2020) Equation 4-4 through 4-7 https://www.epa.gov/system/files/documents/2022-02/disprof_bench_3rules_final_

See references list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

Examples

```
# Use kd from experimental data (recommended):
define_water(ph = 7.5, temp = 25) %>%
  solvect_o3(time = 10, dose = 2, kd = -0.5, baffle = 0.9)
# Use modeled decay curve:
define_water(ph = 7.5, alk = 100, doc = 2, uv254 = .02, br = 50) %>%
  solvect_o3(time = 10, dose = 2, baffle = 0.5)

library(dplyr)
ct_calc <- water_df %>%
  mutate(br = 50) %>%
  define_water_chain() %>%
  mutate(
    dose = 2,
    O3time = 10,
  ) %>%
  solvect_o3_once(time = O3time, baffle = .7)
```

solvedose_alk

Calculate a desired chemical dose for a target alkalinity

Description

This function calculates the required amount of a chemical to dose based on a target alkalinity and existing water quality. Returns numeric value for dose in mg/L. Uses `uniroot` on the `chemdose_ph` function. For a single water, use `solvedose_alk`; to apply the model to a dataframe, use `solvedose_alk_once`. For most arguments, the `_once` helper "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```
solvedose_alk(water, target_alk, chemical)

solvedose_alk_once(
  df,
  input_water = "defined_water",
  output_column = "dose_required",
  target_alk = "use_col",
  chemical = "use_col"
)
```

Arguments

water	Source water of class "water" created by define_water
target_alk	The final alkalinity in mg/L as CaCO ₃ to be achieved after the specified chemical is added.
chemical	The chemical to be added. Current supported chemicals include: acids: "hcl", "h2so4", "h3po4", "co2", bases: "naoh", "na2co3", "nahco3", "caoh2", "mgoh2"
df	a data frame containing a water class column, which has already been computed using define_water_chain . The df may include a column with names for each of the chemicals being dosed.
input_water	name of the column of water class data to be used as the input. Default is "defined_water".
output_column	name of the output column storing doses in mg/L. Default is "dose_required".

Details

solvedose_alk uses `stats::uniroot()` on [chemdose_ph](#) to match the required dose for the requested alkalinity target.

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

solvedose_alk returns a numeric value for the required chemical dose.

solvedose_alk_once returns a data frame containing the original data frame and columns for target alkalinity, chemical dosed, and required chemical dose.

See Also

[solvedose_ph](#)

Examples

```
dose_required <- define_water(ph = 7.9, temp = 22, alk = 100, 80, 50) %>%
  solvedose_alk(target_alk = 150, "naoh")

library(dplyr)

example_df <- water_df %>%
  define_water_chain() %>%
  mutate(finAlk = seq(100, 210, 10)) %>%
  solvedose_alk_once(chemical = "na2co3", target_alk = finAlk)

# Initialize parallel processing
library(furrr)
# plan(multisession)
example_df <- water_df %>%
  define_water_chain() %>%
  mutate(target_alk = seq(100, 210, 10)) %>%
  solvedose_alk_once(chemical = "na2co3")

# Optional: explicitly close multisession processing
# plan(sequential)
```

solvedose_ph

Calculate a desired chemical dose for a target pH

Description

Calculates the required amount of a chemical to dose based on a target pH and existing water quality. The function takes an object of class "water", and user-specified chemical and target pH and returns a numeric value for the required dose in mg/L. For a single water, use `solvedose_ph`; to apply the model to a dataframe, use `solvedose_ph_once`. For most arguments, the `_once` helper "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```
solvedose_ph(water, target_ph, chemical)

solvedose_ph_once(
  df,
  input_water = "defined_water",
  output_column = "dose_required",
  target_ph = "use_col",
  chemical = "use_col"
)
```

Arguments

water	Source water of class "water" created by define_water
target_ph	The final pH to be achieved after the specified chemical is added.
chemical	The chemical to be added. Current supported chemicals include: acids: "hcl", "h2so4", "h3po4", "co2"; bases: "naoh", "na2co3", "nahco3", "caoh2", "mgoh2"
df	a data frame containing a water class column, which has already been computed using define_water_chain . The df may include a column with names for each of the chemicals being dosed.
input_water	name of the column of water class data to be used as the input. Default is "defined_water".
output_column	name of the output column storing doses in mg/L. Default is "dose_required".

Details

solvedose_ph uses `stats::uniroot()` on [chemdose_ph](#) to match the required dose for the requested pH target.

For large datasets, using `fn_once` or `fn_chain` may take many minutes to run. These types of functions use the `furrr` package for the option to use parallel processing and speed things up. To initialize parallel processing, use `plan(multisession)` or `plan(multicore)` (depending on your operating system) prior to your piped code with the `fn_once` or `fn_chain` functions. Note, parallel processing is best used when your code block takes more than a minute to run, shorter run times will not benefit from parallel processing.

Value

A numeric value for the required chemical dose.

`solvedose_ph_once` returns a data frame containing the original data frame and columns for target pH, chemical dosed, and required chemical dose.

See Also

[chemdose_ph](#), [solvedose_alk](#)

Examples

```
water <- define_water(ph = 7, temp = 25, alk = 10)

# Calculate required dose of lime to reach pH 8
solvedose_ph(water, target_ph = 8, chemical = "caoh2")

example_df <- water_df %>%
  define_water_chain() %>%
  solvedose_ph_once(input_water = "defined_water", target_ph = 8.8, chemical = "naoh")

# Initialize parallel processing
library(dplyr)
```

```
library(furrr)
# plan(multisession, workers = 2) # Remove the workers argument to use all available compute
example_df <- water_df %>%
  define_water_chain() %>%
  mutate(finph = seq(9, 10.1, .1)) %>%
  solvedose_ph_once(chemical = "naoh", target_ph = finph)

# Optional: explicitly close multisession processing
# plan(sequential)
```

solvemass_chem

Convert mg/L of chemical to lb/day

Description

This function takes a chemical dose in mg/L, plant flow in MGD, and chemical strength and calculates lb/day of product

Usage

```
solvemass_chem(dose, flow, strength = 100)
```

Arguments

dose	Chemical dose in mg/L as chemical
flow	Plant flow in MGD
strength	Chemical product strength in percent. Defaults to 100 percent.

Value

A numeric value for the chemical mass in lb/day.

Examples

```
alum_mass <- solvemass_chem(dose = 20, flow = 10, strength = 49)

library(dplyr)
mass_data <- tibble(
  dose = seq(10, 50, 10),
  flow = 10
) %>%
  mutate(mass = solvemass_chem(dose = dose, flow = flow, strength = 49))
```

solvemass_solids	<i>Determine solids lb/day</i>
------------------	--------------------------------

Description

This function takes coagulant doses in mg/L as chemical, removed turbidity, and plant flow as MGD to determine solids production.

Usage

```
solvemass_solids(
  alum = 0,
  ferricchloride = 0,
  ferricsulfate = 0,
  flow,
  turb,
  b = 1.5
)
```

Arguments

alum	Amount of hydrated aluminum sulfate added in mg/L as chemical: $\text{Al}_2(\text{SO}_4)_3 \cdot 14\text{H}_2\text{O} + 6\text{HCO}_3 \rightarrow 2\text{Al}(\text{OH})_3(\text{am}) + 3\text{SO}_4 + 14\text{H}_2\text{O} + 6\text{CO}_2$
ferricchloride	Amount of ferric chloride added in mg/L as chemical: $\text{FeCl}_3 + 3\text{HCO}_3 \rightarrow \text{Fe}(\text{OH})_3(\text{am}) + 3\text{Cl} + 3\text{CO}_2$
ferricsulfate	Amount of ferric sulfate added in mg/L as chemical: $\text{Fe}_2(\text{SO}_4)_3 \cdot 8.8\text{H}_2\text{O} + 6\text{HCO}_3 \rightarrow 2\text{Fe}(\text{OH})_3(\text{am}) + 3\text{SO}_4 + 8.8\text{H}_2\text{O} + 6\text{CO}_2$
flow	Plant flow in MGD
turb	Turbidity removed in NTU
b	Correlation factor from turbidity to suspended solids. Defaults to 1.5.

Value

A numeric value for solids mass in lb/day.

Source

<https://water.mecc.edu/courses/ENV295Residuals/lesson3b.htm#:~:text=From%20the%20diagram%2C%20for%20example>

Examples

```
solids_mass <- solvemass_solids(alum = 50, flow = 10, turb = 20)

library(dplyr)
mass_data <- tibble(
  alum = seq(10, 50, 10),
```

```
    flow = 10
  ) %>%
    mutate(mass = solvemass_solids(alum = alum, flow = flow, turb = 20))
#'
```

<i>solveresid_o3</i>	<i>Determine ozone decay</i>
----------------------	------------------------------

Description

This function applies the ozone decay model to a water from U.S. EPA (2001) equation 5-128. For a single water, use *solveresid_o3*; to apply the model to a dataframe, use *solveresid_o3_once*. For most arguments, the *_once* helper "use_col" default looks for a column of the same name in the dataframe. The argument can be specified directly in the function instead or an unquoted column name can be provided.

Usage

```
solveresid_o3(water, dose, time)

solveresid_o3_once(
  df,
  input_water = "defined_water",
  output_column = "o3resid",
  dose = "use_col",
  time = "use_col"
)
```

Arguments

<i>water</i>	Source water object of class water created by define_water
<i>dose</i>	Applied ozone dose in mg/L
<i>time</i>	Ozone contact time in minutes
<i>df</i>	a data frame containing a water class column, which has already been computed using define_water_chain
<i>input_water</i>	name of the column of Water class data to be used as the input for this function. Default is "defined_water".
<i>output_column</i>	name of the output column storing doses in mg/L. Default is "dose_required".

Value

solveresid_o3 returns a numeric value for the residual ozone.

solveresid_o3_once returns a data frame containing the original data frame and columns for ozone dosed, time, and ozone residual.

Source

U.S. EPA (2001)

See reference list at: <https://github.com/BrownandCaldwell-Public/tidywater/wiki/References>

Examples

```
ozone_resid <- define_water(7, 20, 100, doc = 2, toc = 2.2, uv254 = .02, br = 50) %>%
  solveresid_o3(dose = 2, time = 10)
```

```
library(dplyr)
ozone_resid <- water_df %>%
  mutate(br = 50) %>%
  define_water_chain() %>%
  solveresid_o3_once(dose = 2, time = 10)
```

```
ozone_resid <- water_df %>%
  mutate(br = 50) %>%
  define_water_chain() %>%
  mutate(
    dose = seq(1, 12, 1),
    time = seq(2, 24, 2)
  ) %>%
  solveresid_o3_once()
```

summarize_wq

Create summary table from water class

Description

This function takes a water data frame defined by `define_water` and outputs a formatted summary table of specified water quality parameters.

`summarise_wq()` and `summarize_wq()` are synonyms.

Usage

```
summarize_wq(water, params = c("general"))
```

```
summarise_wq(water, params = c("general"))
```

Arguments

water	Source water vector created by <code>define_water</code> .
params	List of water quality parameters to be summarized. Options include "general", "ions", and "dbps". Defaults to "general" only.

Details

Use `chemdose_dbp` for modeled DBP concentrations.

Value

A knitr_kable table of specified water quality parameters.

Examples

```
# Summarize general parameters
water_defined <- define_water(7, 20, 50, 100, 80, 10, 10, 10, 10, tot_po4 = 1)
summarize_wq(water_defined)

# Summarize major cations and anions
summarize_wq(water_defined, params = list("ions"))
```

water_df	<i>Data frame of water quality parameters</i>
----------	---

Description

A dataset containing fabricated water quality to use as tidywater inputs. Parameters are set to reasonable water quality ranges. Parameters are as follows:

Usage

```
water_df
```

Format

A dataframe with 12 rows and 11 columns:

ph pH in standard units (SU)
temp Temperature in degree C
alk Alkalinity in mg/L as CaCO₃
tot_hard Total hardness in mg/L as CaCO₃
ca_hard Calcium hardness in mg/L as CaCO₃
na Sodium in mg/L Na+
k Potassium in mg/L K+
cl Chloride in mg/L Cl-
so4 Sulfate in mg/L SO₄²⁻
totocl Total chlorine in mg/L as Cl₂
totpo4 Total phosphate in mg/L as PO₄³⁻

Source

Fabricated for use in examples.

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