

Package ‘eixport’

October 13, 2022

Title Export Emissions to Atmospheric Models

Version 0.5.4

Date 2022-09-26

Description Emissions are the mass of pollutants released into the atmosphere. Air quality models need emissions data, with spatial and temporal distribution, to represent air pollutant concentrations. This package, 'eixport', creates inputs for the air quality models 'WRF-Chem' Grell et al (2005) <[doi:10.1016/j.atmosenv.2005.04.027](https://doi.org/10.1016/j.atmosenv.2005.04.027)>, 'MUNICH' Kim et al (2018) <[doi:10.5194/gmd-11-611-2018](https://doi.org/10.5194/gmd-11-611-2018)>, 'BRAMS-SPM' Freitas et al (2005) <[doi:10.1016/j.atmosenv.2005.07.017](https://doi.org/10.1016/j.atmosenv.2005.07.017)> and 'RLINE' Snyder et al (2013) <[doi:10.1016/j.atmosenv.2013.05.074](https://doi.org/10.1016/j.atmosenv.2013.05.074)>. See the 'eixport' website (<<https://atmoschem.github.io/eixport/>>) for more information, documentation and examples. More details in Ibarra-Espinosa et al (2018) <[doi:10.21105/joss.00607](https://doi.org/10.21105/joss.00607)>.

License MIT + file LICENSE

URL <https://atmoschem.github.io/eixport/>

BugReports <https://github.com/atmoschem/eixport/issues/>

Depends R (>= 3.5.0)

Imports sf, ncdf4, raster, sp, methods, cptcity, utils, parallel, data.table, rgdal

Encoding UTF-8

LazyData no

RoxygenNote 7.2.1

Suggests testthat, covr, lwgeom, units, knitr, rmarkdown, vein

NeedsCompilation no

Author Sergio Ibarra-Espinosa [aut, cre]

(<<https://orcid.org/0000-0002-3162-1905>>),

Daniel Schuch [aut] (<<https://orcid.org/0000-0001-5977-4519>>),

Edmilson Freitas [ths] (<<https://orcid.org/0000-0001-8783-2747>>),

Umur Dinç [ctb] (<<https://orcid.org/0000-0003-2439-8757>>)

Maintainer Sergio Ibarra-Espinosa <zergioibarra@gmail.com>

Repository CRAN

Date/Publication 2022-09-27 08:20:02 UTC

R topics documented:

chem_edgar	2
edgar	4
emisco	5
emis_opt	5
gCO	6
get_edgar	7
Lights	9
rawprofile	10
sfx_explode	11
st_explode	12
to_as4wrf	12
to_brams_spm	14
to_munich	15
to_rline	17
to_wrf	19
wrf_add	21
wrf_create	22
wrf_get	24
wrf_grid	26
wrf_meta	27
wrf_plot	28
wrf_profile	29
wrf_put	31
wrf_raster	32
wrf_summary	33
Index	35

chem_edgar

Agregate EDGAR emissions NetCDF files into a RasterStack by

Description

The Emissions Database for Global Atmospheric Research (EDGAR) is a project from the Joint Research Centre. This function reads the NetCDF and merge/aggregate into diferent chemical mechanisms

Usage

```
chem_edgar(
  path,
  chem,
  merge = FALSE,
  k = rep(1, 34),
  recursive = TRUE,
  verbose = TRUE
)
```

Arguments

path	Character; path to the NetCDF files from EDGAR. The directory must have one file for each of the following pollutants: "voc" from 1 to 25, "co", "nox", "nmvoc", "so2", "nh3", "pm10", "pm2.5", "bc" and "oc"
chem	Character; chemical mechanism: "edgar", "radm", "radmsorg", "cbmz_mosaic", "cptec", "ecb05_opt1", "neu_cb05" (thanks to Daniel Schuch) and "ufpr_cbmz" (thanks to Leila Martins). <ul style="list-style-type: none"> • When chem is "edgar" units are: "g km-2 h-1" • Other mechanisms: gases "mol km-2 h-1" and aerosols: "ug m-2 s-1"
merge	Logical; in the case that there are more than one NetCDF per pollutant, merge = TRUE will merge them with sum. Default is FALSE.
k,	Numeric; Value to factorize each pollutant.
recursive	Logical to search files recursive
verbose	Logical to print more information

Value

RasterStack

Note

Molecular weights were obtained from

Development of Improved Chemical Speciation Database for Processing Emissions of Volatile Organic Compounds for Air Quality Models <https://intra.engr.ucr.edu/~carter/emitdb/>

Some mappings were obtained from:

Carter, W. P. (2015). Development of a database for chemical mechanism assignments for volatile organic emissions. *Journal of the Air & Waste Management Association*, 65(10), 1171-1184.

Lopez-Norena, Ana and Fernandez, Rafael & Puliafito, SALVADOR. (2019). ESPECIACION DE INVENTARIOS DE EMISIONES DE AEROSOL Y COMPUESTOS ORGANICOS VOLATILES PARA EL MODELO WRF-CHEM, APLICADO A LOS ESQUEMAS RADM-2, CBM-Z Y MOZART-4.

Examples

```
# Downloading EDGAR data ####
p <- tempdir()
tot <- paste0(p, "/TOT")
dir.create(tot)

get_edgar(
  dataset = "v432_VOC_spec",
  destpath = tot,
  sector = c("TOTALS"),
  type = "nc",
  year = 2012,
```

```

    ask = FALSE
  )

  get_edgar(
    dataset = "v50_AP",
    destpath = tot,
    sector = c("TOTALS"),
    type = "nc",
    year = 2014,
    ask = FALSE
  )

  nc <- paste0(p, "/nc")

  totals <- list.files(
    path = tot,
    full.names = TRUE,
    pattern = ".zip"
  )

  lapply(totals, unzip, exdir = nc)

  r <- chem_edgar(path = nc,
                 chem = "radm")
  library(raster)
  plot(r$E_CO)
  r$E_CO[] <- ifelse(r$E_CO[] <= 0, NA, r$E_CO[])
  br <- c(1.133594e-10, 0.001105391, 0.002727838, 0.007006367, 0.0146665,
         0.03192566, 0.08026344, 0.5739333, 4.110863, 16.47688, 6.641064e+05)
  plot(r$E_CO, col = cptcity::cpt(rev = TRUE), breaks = br)

```

 edgar

Emissions EDGAR

Description

Several datasets

Usage

```
data(edgar)
```

Format

A data.frame with links to download EDGAR data

data datasets

pol pollutants

sector Sector

links description
url URL
year Year
type nc, txt or NA
note Notes data(edgar)

Source

<https://edgar.jrc.ec.europa.eu/>

emisco	<i>Emissions from VEIN examples</i>
--------	-------------------------------------

Description

Emissions for street models such as munich. They need to be splitted using st_explode

Usage

data(emisco)

Format

A sf object of type LINESTRING with 1505 rows and 24 variables:

V8 Emissions for 08:00-09:00 in Easth Sao Paulo, Brazil (g/h)

geometry Geometry class sfc_LINESTRING sfc data(emisco)

Source

<https://github.com/atmoschem/vein>

emis_opt	<i>List of WRF emission species</i>
----------	-------------------------------------

Description

Emission package definitions from WRF 4.0.1, for use in wrf_create function.

Usage

data(emis_opt)

Format

A list of emission variables names, same number as emis_opt in namelist.

Note

look to the number of aerosol of the emis_opt in WRF domumentation / code.

Author(s)

Daniel Schuch

Source

<https://github.com/wrf-model/WRF>

See Also

[wrf_create](#)

Examples

```
data(emis_opt)
names(emis_opt)
emis_opt[["eradm"]]
```

gCO

Gridded emissions from VEIN demo

Description

Emissions in g/h for morning rush hour.

Usage

```
data(gCO)
```

Format

A sf object of POLYGON with 437 rows and 2 variables:

V9 Emissions of CO (g/h) for 08:00-09:00

geometry geometry data(gCO)

Source

<https://github.com/atmoschem/vein>

get_edgar

Download datasets of EDGAR emissions

Description

The Emissions Database for Global Atmospheric Research (EDGAR) is a project from the Joint Research Centre. They provide provides global past and present day anthropogenic emissions of greenhouse gases and air pollutants by country and on spatial grid. [get_edgar](#) provide functions to download any of the EDGAR datasets.

Usage

```
get_edgar(
  dataset = "v50_AP",
  pol,
  sector,
  year,
  n = parallel::detectCores()/2,
  destpath = tempdir(),
  type = "nc",
  ask = TRUE,
  copyright = TRUE,
  verbose = TRUE
)
```

Arguments

dataset	Character; name of the datasets: "v50_AP", "v432_AP", "v432_VOC_spec", "htap_v2_2", "v50_GHG"
pol	Character; one of the pollutants shown on note.
sector	Character; one of the sectors shown on note.
year	Integer; years.
n	Integer; Number of cores in the machine, by default the half.
destpath	Character: Path to create the directory for downloads datasets
type	Extension, character/ Indicates if the file should be "txt", "nc" or "other". txt with untis t/year and nc with units are ug/m2/s
ask	Logical; Are these URL ok?
copyright	Logical; to show copyright information.
verbose	Logical, to print more information.

Value

Downloads data

Note

I will create a package to download all the emissions files, dedicated to that:

I recommend 2 ways:

1. include 'sector' and do not include 'pol', which download all pollutants as default

```
get_edgar(dataset = "v432_AP", destpath = tempdir(), sector = c("TRO", "TOTALS"), year = 2012)
```

2. include 'pol' and do not include 'sector', which download all sectors as default

```
get_edgar(dataset = "v432_AP", destpath = tempdir(), pol = c("CO", "NOx"), year = 2012)
```

dataset	pollutant
v50_AP	BC, CO, NH3, NMVOC, NOx, OC, PM10, PM2.5, SO2
v50_GHG	CH4, CO2_excl_short-cycle_org_C, CO2_org_short-cycle_C, N2O
v432	CH4, CO2_excl_short-cycle_org_C, CO2_org_short-cycle_C, N2O
v432_AP	BC, CO, NH3, NMVOC, NOx, OC, PM10, PM2.5_bio, PM2.5_fossil, SO2
v432_VOC_spec	voc1, voc2, voc3, voc3, voc5, voc6, voc7, voc8, bvoc9, voc10, voc11, voc12, voc13, voc14, voc15, voc16,
htap_v2_2	BC, CO, NH3, NMVOC, NOx, OCm, PM10, PM2.5, SO2

voc11 only 2008

References

- **v50_AP** Crippa, M., Solazzo, E., Huang, G., Guizzardi, D., Koffi, E., Muntean, M., Schieberle, C., Friedrich, R., Janssens-Maenhout, G.: High resolution temporal profiles in the Emissions Database for Global Atmospheric Research, Nature Scientific Data, 2020, in press.
- **v432:** Muntean, M., Guizzardi, D., Schaaf, E., Crippa, M., Solazzo, E., Olivier, J.G.J., Vignati, E. Fossil CO2 emissions of all world countries - 2018 Report, EUR 29433 EN, Publications Office of the European Union, Luxembourg, 2018, ISBN 978-92-79-97240-9, doi:10.2760/30158, JRC113738.
- **v432_AP:** Crippa, M., Guizzardi, D., Muntean, M., Schaaf, E., Dentener, F., van Aardenne, J. A., Monni, S., Doering, U., Olivier, J. G. J., Pagliari, V., and Janssens-Maenhout, G.: Gridded emissions of air pollutants for the period 1970–2012 within EDGAR v4.3.2, Earth Syst. Sci. Data, 10, 1987–2013, <https://doi.org/10.5194/essd-10-1987-2018>, 2018.
- **v432_VOC_spec:** Huang, G., Brook, R., Crippa, M., Janssens-Maenhout, G., Schieberle, C., Dore, C., Guizzardi, D., Muntean, M., Schaaf, E., and Friedrich, R.: Speciation of anthropogenic emissions of non-methane volatile organic compounds: a global gridded data set for 1970–2012, Atmos. Chem. Phys., 17, 7683-7701, <https://doi.org/10.5194/acp-17-7683-2017>, 2017.
- **htap_v2_2:** Janssens-Maenhout, G., Crippa, M., Guizzardi, D., Dentener, F., Muntean, M., Pouliot, G., Keating, T., Zhang, Q., Kurokawa, J., Wankmüller, R., Denier van der Gon, H., Kuenen, J. J. P., Klimont, Z., Frost, G., Darras, S., Koffi, B., and Li, M.: HTAP_v2.2: a mosaic of regional and global emission grid maps for 2008 and 2010 to study hemispheric transport of air pollution, Atmos. Chem. Phys., 15, 11411–11432, <https://doi.org/10.5194/acp-15-11411-2015>, 2015.

MNM is MNN for NOx v432_AP

Examples

```
# see all the links:
data(edgar)
head(edgar)
# Download all pollutants for sector
get_edgar(dataset = "v50_AP",
          pol = "CO",
          sector = "TOTALS",
          year = 2014,
          destpath = tempdir(),
          type = "nc",
          ask = FALSE)
```

Lights

Spatial distribution example

Description

Spatial distribution for vehicular emissions based on an image of persistent lights of the Defense Meteorological Satellite Program (DMSP) for 5 Brazilian states (Sao Paulo, Rio de Janeiro, Mato Grosso, and Santa Catarina e Parana).

Usage

```
data(Lights)
```

Format

A matrix of spatial distribution

Details

https://en.wikipedia.org/wiki/Defense_Meteorological_Satellite_Program

Author(s)

Daniel Schuch

Source

<https://ngdc.noaa.gov/eog/dmsp/downloadV4composites.html>

See Also

[to_wrf](#)

Examples

```

dir.create(file.path(tempdir(), "EMISS"))
wrf_create(wrfinput_dir = system.file("extdata", package = "eixport"),
           wrfchemi_dir = file.path(tempdir(), "EMISS"),
           frames_per_auxinput5 = 24)

# get the name of created file
files <- list.files(path = file.path(tempdir(), "EMISS"),
                   pattern = "wrfchemi",
                   full.names = TRUE)

data(Lights)

perfil <- c(0.010760058, 0.005280596, 0.002883553, 0.002666932,
           0.005781312, 0.018412838, 0.051900411, 0.077834636,
           0.067919758, 0.060831614, 0.055852868, 0.052468599,
           0.050938043, 0.051921718, 0.052756244, 0.052820165,
           0.058388406, 0.072855890, 0.075267137, 0.063246412,
           0.042713523, 0.029108975, 0.022091855, 0.015298458)

plot(perfil,
     ty = "l",
     col= "purple",
     xlab = "Hour",
     main = "Time profile",
     ylab = "Weight",
     axes = FALSE,
     xlim = c(0, 24))
axis(2)
axis(1,
     at = c(0, 6, 12, 18, 24),
     labels = c("00:00", "06:00", "12:00", "18:00", "00:00"))

to_wrf(Lights,
       files[1],
       total = 1521983,
       profile = perfil,
       name = "E_CO")

```

rawprofile

Raw profile

Description

Raw profile

Usage

```
data(rawprofile)
```

Format

A matrix with 1 column and 168 rows

```
data(rawprofile)
```

sfx_explode	<i>splits line by vertex</i>
-------------	------------------------------

Description

`sfx_explode` splits line by vertex

Usage

```
sfx_explode(x)
```

Arguments

x sf LINESTRING.

Value

spatial lines

Examples

```
{
  data(emisco)
  dim(emisco)
  dfco <- sfx_explode(emisco)
  dim(dfco)
}
```

st_explode	<i>Split line by vertex (experimental)</i>
------------	--

Description

`st_explode` split a lines data.frame into each vertex. It to mimic the function explode from qgis, that the reason for the name `st_explode`

Usage

```
st_explode(net)
```

Arguments

net	A spatial dataframe of class "sp" or "sf". When class is "sp" it is transformed to "sf".
-----	--

Value

a spatial data.frame

Note

All variables are transformed into numeric.

Examples

```
# do not run
require(vein)
data(net)
net2 <- st_explode(net)
dim(net)
dim(net2)
```

to_as4wrf	<i>Generates emissions dataframe to generate WRF-Chem inputs</i>
-----------	--

Description

`to_as4wrf` returns a dataframes with columns lat, long, id, pollutants, local time and GMT time. This dataframe has the proper format to be used with WRF assimilation system: "Another Assimilation System 4 WRF (AAS4WRF)" as published by Vera-Vala et al (2016)

Usage

```
to_as4wrf(sdf, nr = 1, dmyhm, tz, crs = 4326, islist)
```

Arguments

sdf	Gridded emissions, which can be a SpatialPolygonsDataFrame, or a list of SpatialPolygonsDataFrame, or a sf object of "POLYGON". The user must enter a list with 36 SpatialPolygonsDataFrame with emissions for the mechanism CBMZ. When there are no emissions available, the SpatialPolygonsDataFrame must contain 0.
nr	Number of repetitions of the emissions period
dmyhm	String indicating Day Month Year Hour and Minute in the format "d-m-Y H:M" e.g.: "01-05-2014 00:00" It represents the time of the first hour of emissions in Local Time
tz	Time zone as required in for function as.POSIXct
crs	Coordinate reference system, e.g: "+init=crs:4326". Used to transform the coordinates of the output
islist	logical value to indicate if sdf is a list or not

Value

data-frame of gridded emissions g/h

Note

The user must produce a text file with the data-frame resulting of this function. Then, use this file with the NCL script AAS4WRF.ncl

The reference of the emissions assimilation system is Vara-Vela, A., Andrade, M. F., Kumar, P., Ynoue, R. Y., and Munoz, A. G.: Impact of vehicular emissions on the formation of fine particles in the Sao Paulo Metropolitan Area: a numerical study with the WRF-Chem model, Atmos. Chem. Phys., 16, 777-797, doi:10.5194/acp-16-777-2016, 2016. A good website with timezones is <http://www.timezoneconverter.com/cgi-bin/tzc> The crs is the same as used by [sp](#) package It returns a dataframe with id,, long, lat, pollutants, time_lt, time_utc and day-UTC-hour (dutch) The pollutants for the CBMZ are: e_so2, e_no, e_ald, e_hcho, e_ora2, e_nh3 e_hc3, e_hc5, e_hc8, e_eth, e_co, e_ol2, e_olt, e_oli, e_tol, e_xyl, e_ket e_csl, e_iso, e_no2, e_ch3oh, e_c2h5oh, e_pm25i, e_pm25j, e_so4i, e_so4j e_no3i, e_no3j, e_orgi, e_orgj, e_eci, e_ecj, e_so4c, e_no3c, e_orgc, e_ecc

See Also

[wrf_create to_wrf](#)

Examples

```
{
data(gCO)
df <- to_as4wrf(sdf = gCO,
                dmyhm = "29-04-2018 00:00",
                tz = "America/Sao_Paulo")
```

```
head(df)
df2 <- to_as4wrf(sdf = list(co = gCO, pm = gCO),
                dmyhm = "29-04-2018 00:00",
                tz = "America/Sao_Paulo")
head(df2)
}
```

to_brams_spm

Inputs for BRAMS-SPM

Description

Create inputs for BRAMS-SPM. The inputs consist of a data-frame or a list of data-frames with daily emissions (mol/day), lat, long. Also, including a functions describing the hourly profile.

Usage

```
to_brams_spm(sdf, epsg = 4326)
```

Arguments

sdf	Grid emissions, which can be a SpatialPolygonsDataFrame or polygon grid class 'sf' including the hourly emissions in mol/h for 24 hours. The object can also be a list of objects SpatialPolygonsDataFrame or Spatial Features polygon grid class 'sf'.
epsg	Coordinate reference system, e.g: "4326". Used to transform the coordinates of the output.

Value

data-frame of daily gridded emissions, lat, long and a message with function.

Note

When the input is class 'Spatial', they are converted to 'sf'. If the input is a data-frame, the output is a data-frame. If the input is a list, the output is a list.

Author(s)

Sergio Ibarra and Edmilson Freitas

References

SPM BRAMS: FREITAS, E. MARTINS, L., SILVA, P. and ANDRADE, M. A simple photochemical module implemented in rams for tropospheric ozone concentration forecast in the metropolitan area of são paulo, brazil: Coupling and validation. Atmospheric Environment, Elsevier, n. 39, p. 6352–6361, 2005.

Examples

```

data(gCO)
df1 <- to_brams_spm(sdf = gCO,
                    epsg = 4326)

head(df1)
df2 <- to_brams_spm(sdf = list(co = gCO, pm = gCO),
                    epsg = 4326)

lapply(df2, head)

```

to_munich	<i>Export emissions to Model of Urban Network of Intersecting Canyons and Highways (MUNICH)</i>
-----------	---

Description

`to_munich` Export spatial emissions objects according the format required by MUNICH. This function was designed to read street emissions from VEIN by it can be used to read any other.

Usage

```
to_munich(sdf, idbrin, typo, width, height, crs = 4326)
```

Arguments

sdf	Street Emissions object class 'sf' LINESTRING or "SpatialLinesdataFrame". The columns are the emissions.
idbrin	Integer; id.
typo	Integer; id2.
width	Integer; width.
height	Integer; heigth.
crs	Numeric; Coordenade Reference System to project data or not.

Value

A list with a data frame with columns "i", "idbrin", "typo", "xa", "ya", "xb", "yb" and the pollutants; and another data.frame with "i", "length" (m), "width" (with value 0) and "height" (with value 0). Width and height must be obtained by the user.

Note

The user must ensure that the spatial object has one line feature per vertex and lines with more than one vertex must be previously splitted. the resulting units must be **ug/km/h**

References

Kim, Y., Wu, Y., Seigneur, C., and Roustan, Y.: Multi-scale modeling of urban air pollution: development and application of a Street-in-Grid model (v1.0) by coupling MUNICH (v1.0) and Polair3D (v1.8.1), *Geosci. Model Dev.*, 11, 611-629, <https://doi.org/10.5194/gmd-11-611-2018>, 2018.

Examples

```
# Not run
library(vein)
library(units)
library(sf)
data(net)
data(pc_profile)
data(profiles)
data(fkm)
PC_G <- c(33491,22340,24818,31808,46458,28574,24856,28972,37818,49050,87923,
         133833,138441,142682,171029,151048,115228,98664,126444,101027,
         84771,55864,36306,21079,20138,17439, 7854,2215,656,1262,476,512,
         1181, 4991, 3711, 5653, 7039, 5839, 4257,3824, 3068)
pc1 <- my_age(x = net$lvd,
             y = PC_G,
             name = "PC")

# Estimation for morning rush hour and local emission factors and speed
speed <- data.frame(S8 = net$ps)
lef <- EmissionFactorsList(ef_cetesb("CO",
                                   "PC_G",
                                   agemax = ncol(pc1)))

E_CO <- emis(veh = pc1,
            lkm = net$lkm,
            ef = lef,
            speed = speed)

# rowSums drop units
net$CO <- set_units(rowSums(E_CO), g/h)

# selecting only CO and exploding lines and updating emissions
df <- st_explode(net["CO"])

# st_explode should not drop units, must fix
df$CO <- set_units(df$CO, g/h)

# now we have split line in vertex
# selecting 1000 links
dfco <- df[1:1000,"CO"]

#####
#MUNICH relies in a python script that reads emissions with units ug/km/h
# Therefore
dfco$CO <- set_units(dfco$CO, ug/h)
dfco$CO <- dfco$CO/set_units(st_length(dfco), km)
```



```
etm <- to_munich(sdf = dfco)

names(etm)
class(etm)
head(etm$Emissions)
head(etm$Street)

write.table(x = etm$Emissions,
           file = paste0(tempfile(), "_Emissions.txt"),
           row.names = FALSE,
           sep = " ",
           quote = FALSE)

write.table(x = etm$Street,
           file = paste0(tempfile(), "_Street.txt"),
           row.names = FALSE,
           sep = " ",
           quote = FALSE)
```

to_rline

Export emissions to other formats

Description

Export emissions object according to format of file 'Sources.txt' of the model R-LINE

Usage

```
to_rline(
  Emis,
  Z_b,
  Z_e,
  dCL,
  sigmaz0,
  lanes,
  Hw1,
  dw1,
  Hw2,
  dw2,
  Depth,
  Wtop,
  Wbottom,
  experimental = FALSE,
  crs
)
```

Arguments

Emis	Column with the emissions whose unit must be g/ms.
Z_b	initial meters above sea level (m).
Z_e	final meters above sea level (m).
dCL	offset distance for each source relative to the centerline.
sigmaz0	vertical dispersion (m).
lanes	number of lanes at each street.
Hw1	Height of the barrier 1 (m).
dw1	Distance to barrier 1 (m).
Hw2	height of the barrier 2 (m).
dw2	Distance to barrier 2 (m).
Depth	Depth of the depression. Used for depressed roadway (m).
Wtop	width of the opening at the top of the depression (m).
Wbottom	width of the roadway at the bottom of the depression (m).
experimental	Boolean argument to denote the use of the experimental features (TRUE) or not (FALSE).
crs	Numeric; Coordenade Reference System to project data or not.

Value

Data frame with format for R-LINE model.

Note

Michelle G. Snyder, Akula Venkatram, David K. Heist, Steven G. Perry, William B. Petersen, Vlad Isakov, RLINE: A line source dispersion model for near-surface releases, In Atmospheric Environment, Volume 77, 2013, Pages 748-756, ISSN 1352-2310, <https://doi.org/10.1016/j.atmosenv.2013.05.074>.

Examples

```
{
data(emisco)
emisco <- st_explode(emisco)
emisco$V8 <- units::set_units(emisco$V8, "g/ms")
Source <- to_rline(Emis = emisco["V8"],
                  Z_b = 0,
                  Z_e = 0,
                  dCL = 0,
                  sigmaz0 = 2,
                  lanes = 1)
head(Source)
write.table(x = Source,
           file = paste0(tempdir(), "/Sources.txt"),
           row.names = FALSE,
           sep = " ",
           quote = FALSE)
}
```

to_wrf

Combine total/spatial/temporal/split and write emission to file

Description

Function to expand, split and write emissions. The input is expanded into time by profile and split between variables with different weights.

Usage

```
to_wrf(
  POL,
  file = file.choose(),
  name = NA,
  total = NA,
  norm = FALSE,
  profile = 1,
  weights = 1,
  verbose = TRUE
)
```

Arguments

POL	matrix or array of emissions of spatial weights
file	emission file name
name	species to be write
total	total of emitted species (modifier)
norm	if the spatial weights need to be normalized (modifier)
profile	temporal profile to expand the emissions (modifier)
weights	weight of each species (modifier)
verbose	display additional information

Value

Write files to a wrf netcdf file

Note

length(profile) must be the number of times in the emission file (value of frames_per_auxinput5 if wrf_create() was used to create this file).

total is an additional way to calculate or correct the total emissions

sum(profile) = 1 and sum(weights) = 1 to conserve mass

names and weights must have the same length

Author(s)

Daniel Schuch

See Also[wrf_create](#), [wrf_get](#), [wrf_profile](#) and [wrf_plot](#)**Examples**

```
dir.create(file.path(tempdir(), "EMISS"))
wrf_create(wrfinput_dir = system.file("extdata", package = "elexport"),
          wrfchemi_dir = file.path(tempdir(), "EMISS"),
          frames_per_auxinput5 = 24)

# get the name of created file
files <- list.files(path = file.path(tempdir(), "EMISS"),
                  pattern = "wrfchemi",
                  full.names = TRUE)

data(Lights)

perfil <- c(0.010760058, 0.005280596, 0.002883553, 0.002666932,
           0.005781312, 0.018412838, 0.051900411, 0.077834636,
           0.067919758, 0.060831614, 0.055852868, 0.052468599,
           0.050938043, 0.051921718, 0.052756244, 0.052820165,
           0.058388406, 0.072855890, 0.075267137, 0.063246412,
           0.042713523, 0.029108975, 0.022091855, 0.015298458)

plot(perfil,
     ty = "l",
     col= "purple",
     xlab = "Hour",
     main = "Time profile",
     ylab = "Weight",
     axes = FALSE,
     xlim = c(0, 24))
axis(2)
axis(1,
     at = c(0, 6, 12, 18, 24),
     labels = c("00:00", "06:00", "12:00", "18:00", "00:00"))

to_wrf(Lights,
       files[1],
       total = 1521983,
       profile = perfil,
       name = "E_CO")
```

wrf_add	<i>Function to add values for variables on emission files</i>
---------	---

Description

Add values to a variable in a netCDF file, the main use is to combine different emissions like top-down emission (EmissV emissions) and inventory emission (such as EDGAR, GAINS, RETRO, etc).

Usage

```
wrf_add(file = file.choose(), name = NA, POL)
```

Arguments

file	name of file interactively (default) or specified
name	name of the variable (any variable)
POL	variable to be written

Value

Adds vars too a NETCDF WRF file

Note

this function might be deprecated in future

Author(s)

Daniel Schuch

Examples

```
{
# create the folder and emission file
dir.create(file.path(tempdir(), "EMISS"))
wrf_create(wrfinput_dir = system.file("extdata", package = "elexport"),
           wrfchemi_dir = file.path(tempdir(), "EMISS"))

# get the name of created file
files <- list.files(path = file.path(tempdir(), "EMISS"),
                   pattern = "wrfchemi",
                   full.names = TRUE)

# open, put some numbers and write
CO <- wrf_get(file = files[1], name = "E_CO")
CO[] = rnorm(length(CO), mean = 5, sd = 1)
wrf_put(file = files[1], name = "E_CO", POL = CO)
```

```
# open, put some different numbers and write
CO[] = rnorm(length(CO),mean = 10,sd = 1)
wrf_add(file = files[1], name = "E_CO", POL = CO)
}
```

wrf_create

Create emission files for the WRF-Chem model

Description

Create WRF-chem emission files using information from the WRF initial conditions (wrfinput) file(s). The wrfinput file of the corresponding domain is read from the current folder or from the wrfinput_dir.

There are two emission styles available: the 12 hour pair of emissions (that will be recycled by the model) using io_style_emissions = 1 and the date_hour format using io_style_emissions = 2 (default), see notes for more detail.

The initial time is the original (wrfinput file) adjusted by the day_offset argument, this argument can be useful for split the emissions into several files or for a restarted simulation. The emissions are recorded at the interval of 60 minutes (or the auxinput5_interval_m argument) for 1 time (or frames_per_auxinput5 argument times).

The variables created on output file is based on emis_opt data or a character vector contains the species, any change in variables need to be followed by a change in the n_aero for the correspondent number of aerosol species in the emission file (the n_aero last variables).

Title argument will be written on global attribute TITLE, from the version 4.0 the model checks if the TITLE version contains "V4.", this can be disabled setting 'force_use_old_data = .true.' on WRF namelist.input.

Usage

```
wrf_create(
  wrfinput_dir = getwd(),
  wrfchemi_dir = wrfinput_dir,
  domains = 1,
  frames_per_auxinput5 = 1,
  auxinput5_interval_m = 60,
  day_offset = 0,
  io_style_emissions = 2,
  kemit = 1,
  variables = "ecb05_opt2",
  n_aero = 15,
  COMPRESS = NA,
  force_ncdf4 = FALSE,
  title = "Anthropogenic emissions for WRF V4.0",
  separator = "default",
  prefix = "wrfchemi",
  overwrite = TRUE,
```

```

    return_fn = FALSE,
    verbose = FALSE
)

```

Arguments

wrfinput_dir	input folder with the wrfinput file(s)
wrfchemi_dir	output folder
domains	domain / domains to be process
frames_per_auxinput5	value from wrf &time_control namelist.input, number of times (frames) in a single emission file
auxinput5_interval_m	value from wrf &time_control namelist.input, interval in minutes between different times (frames) see Details
day_offset	number of days (can be a fraction) see Details
io_style_emissions	from wrf &chem namelist.input, 1 for 12z/00z style and 2 to date_hour style, see Details
kemit	from wrf &chem namelist.input number of vertical levels of the emission file
variables	emission species, can be used emis_opt
n_aero	number of aerosol species
COMPRESS	integer between 1 (least comp.) and 9 (most comp.) or NA for no compression
force_ncdf4	force NetCDF4 format
title	TITLE attribute for the NetCDF
separator	filename alternative separator for hour:minutes:seconds with io_style_emission=2
prefix	file name prefix, default is wrfchemi (wrf default)
overwrite	logical, default is true, if FALSE check if the file exist
return_fn	logical, return the name of the last file created
verbose	print file info

Value

Creates NetCDF file

Note

Using io_style_emissions = 1, the wrfchemi_00z will be generated with day_offset = 0 and wrfchemi_12z with day_offset = 0.5 (frames_per_auxinput5 and auxinput5_interval_m will have no effect).

Windows users may need to rename the emission files or change in namelist the default filename before run wrf.exe with these emission files.

The separator argument can be useful for write in NTSF format discs on linux systems, for 'default' the separator is ':' for linux-like systems and '%3A' for windows.

Author(s)

Daniel Schuch

See Also[to_wrf](#) and [emis_opt](#)**Examples**

```
# Do not run

# emissions for a 1 day forecast for domains 1 and 2

dir.create(file.path(tempdir(), "EMISS"))

# emissions on date_hour style
wrf_create(wrfinput_dir = system.file("extdata", package = "eixport"),
          wrfchemi_dir  = file.path(tempdir(), "EMISS"),
          domains       = 1:2,
          frames_per_auxinput5 = 25,
          auxinput5_interval_m = 60,
          verbose       = TRUE)

# emissions on 00z / 12z style, create the 00z
wrf_create(wrfinput_dir = system.file("extdata", package = "eixport"),
          wrfchemi_dir  = file.path(tempdir(), "EMISS"),
          domains       = 1:2,
          io_style_emissions = 1,
          day_offset    = 0,
          verbose       = TRUE,
          )

# emissions on 00z / 12z style, create the 12z
wrf_create(wrfinput_dir = system.file("extdata", package = "eixport"),
          wrfchemi_dir  = file.path(tempdir(), "EMISS"),
          domains       = 1:2,
          io_style_emissions = 1,
          day_offset    = 0.5,
          verbose       = TRUE)
```

wrf_get*Function to read variables of emission files*

Description

Read a variable

Usage

```
wrf_get(  
  file = file.choose(),  
  name = NA,  
  as_raster = FALSE,  
  raster_crs = "WRF",  
  raster_lev = 1,  
  k = NA,  
  verbose = FALSE,  
  ...  
)
```

Arguments

file	name of file interactively (default) or specified
name	name of the variable (any variable) or time to return a POSIXlt object from model
as_raster	return a raster instead of an array
raster_crs	crs of output if as_raster is TRUE, see details
raster_lev	level for rasters from a 4D variable
k	multiplier
verbose	display additional information
...	additional parameters passed to wrf_raster

Format

array or raster object

Details

wrf_get can return a raster object with the option as_raster = TRUE, raster_crs can be used to specify the output crs of the raster object, raster_crs = 'latlon' can be specified to use latlon option in wrf_raster. If raster_crs is 'WRF' (default), the output projection is equivalent to the WRF grid.

Value

Read vars from NetCDF WRF file.

Author(s)

Daniel Schuch

See Also

[wrf_plot](#) and [wrf_put](#)

Examples

```

{

# create the folder and emission file
dir.create(file.path(tempdir(), "EMISS"))
wrf_create(wrfinput_dir = system.file("extdata", package = "elexport"),
           wrfchemi_dir = file.path(tempdir(), "EMISS"))

# get the name of created file
files <- list.files(path = file.path(tempdir(), "EMISS"),
                   pattern = "wrfchemi",
                   full.names = TRUE)

# open, put some numbers and write
CO <- wrf_get(file = files[1],
              name = "E_CO")

CO[] = rnorm(length(CO))

wrf_put(file = files[1],
        name = "E_CO",
        POL = CO)

COr <- wrf_get(file = files[1],
               name = "E_CO",
               as_raster = TRUE)

}

```

wrf_grid

Creates grid from wrf file

Description

Return a Spatial Feature multipolygon or matrix

Usage

```
wrf_grid(filewrf, type = "wrfinput", matrix = FALSE, as_raster = FALSE)
```

Arguments

filewrf	wrf file
type	Type of wrf file: "wrfinput" or "geo". When type is "geo", lat long comes from mass grid, XLONG_M and XLAT_M
matrix	if the output is matrix or polygon (sf)
as_raster	logical, to return a raster

Value

Returns a sf polygon grid from a NetCDF wrf file

Note

The default crs is 4326 (see <http://spatialreference.org/ref/epsg/>)

Examples

```
{
# Do not run
wrf <- paste(system.file("extdata", package = "elexport"),
             "/wrfinput_d02", sep="")
gwrf <- wrf_grid(wrf)
plot(gwrf, axes = TRUE)
}
```

wrf_meta

Returns metadata (attributes) of wrf file in a data.frame

Description

`wrf_meta` returns the attributes of a wrf NetCDF file in a data.frame. Therefore, there is no need to use `ncdump -h "wrf_file"`

Usage

```
wrf_meta(file = file.choose())
```

Arguments

`file` Character; name of file interactively (default) or specified

Value

a list of data.frames describing a NetCDF WRF file.

Examples

```
{
file = paste0(system.file("extdata", package = "elexport"),"/wrfinput_d02")
wrf_meta(file)
}
```

`wrf_plot`*Simple plot from wrf emission file*

Description

Create a quick plot from wrf emission file

Usage

```
wrf_plot(  
  file = file.choose(),  
  name = NA,  
  time = 1,  
  nivel = 1,  
  barra = TRUE,  
  lbarra = 0.2,  
  col = cptcity::cpt(n = 20, rev = TRUE),  
  map = NULL,  
  skip = FALSE,  
  no_title = FALSE,  
  verbose = TRUE,  
  ...  
)
```

Arguments

<code>file</code>	emission file name
<code>name</code>	pollutant name
<code>time</code>	time from emission file
<code>nivel</code>	level from the emission file
<code>barra</code>	barplot if TRUE
<code>lbarra</code>	length of barplot
<code>col</code>	color vector
<code>map</code>	function call to plot map lines, points and annotation (experimental)
<code>skip</code>	logical, skip plot of constant values
<code>no_title</code>	no title plot
<code>verbose</code>	if TRUE print some information
<code>...</code>	Arguments to be passed to plot methods

Value

a plot of a WRF var.

Note

If the file contains levels (kemit>1), and one frame (auxinput5_interval_m = 1) time with control the level which will be plotted

In case of an error related to plot.new() margins lbarra must be adjusted

Author(s)

Daniel Schuch

See Also

[Lights](#), [to_wrf](#) and [wrf_create](#)

Examples

```
dir.create(file.path(tempdir(), "EMISS2"))
wrf_create(wrfinput_dir = system.file("extdata",
                                     package = "eixport"),
          wrfchemi_dir = file.path(tempdir(), "EMISS2"))

# get the name of created file
files <- list.files(path = file.path(tempdir(), "EMISS2"),
                  pattern = "wrfchemi",
                  full.names = TRUE)

# load end write some data in this emission file
data(Lights)
to_wrf(Lights, files[1], total = 1521983, name = "E_CO")

wrf_plot(files[1], "E_CO", col = cptcity::cpt(n = 14))
```

wrf_profile

Create a spatial profile from a wrf emission file and a data frame with

Description

returns a traffic intensity profile (based on wrf file Times) and a traffic intensity data frame

Usage

```
wrf_profile(x, file, adjust = 0, verbose = TRUE)
```

Arguments

x	data.frame of intensity of traffic by hours (rows) and weekdays (columns)
file	emission file name
adjust	number of hours to advance (positive value) or delay (negative value)
verbose	display additional information

Format

a numeric vector

Value

a vector with a profile

Note

It might be deprecated in future release

Author(s)

Daniel Schuch

See Also

[wrf_create](#) and [to_wrf](#)

Examples

```
# Profile based on Sao Paulo tunnel experiments
data(rawprofile)
rawprofile <- matrix(rawprofile, nrow = 24, byrow = TRUE)
rawprofile <- as.data.frame(rawprofile)
names(rawprofile) <- c("Sunday", "Monday", "Tuesday", "Wednesday", "Thursday",
  "Friday", "Saturday")
row.names(rawprofile) <- c("00:00", "01:00", "02:00", "03:00", "04:00", "05:00",
  "06:00", "07:00", "08:00", "09:00", "10:00", "11:00",
  "12:00", "13:00", "14:00", "15:00", "16:00", "17:00",
  "18:00", "19:00", "20:00", "21:00", "22:00", "23:00")

print(rawprofile)

# create the folder and emission file
dir.create(file.path(tempdir(), "EMISS"))
wrf_create(wrfinput_dir = system.file("extdata", package = "eixport"),
  wrfchemi_dir = file.path(tempdir(), "EMISS"),
  frames_per_auxinput5 = 24)

files <- list.files(path = file.path(tempdir(), "EMISS"),
```

```

        pattern = "wrfchemi",
        full.names = TRUE)

profile <- wrf_profile(rawprofile,files[1])

plot(profile,
      ty="l",
      lty = 2,
      axe = FALSE,
      main = "Traffic Intensity for Sao Paulo", xlab = "hour")
axis(2)
axis(1,
      at = 0.5 + c(0, 6, 12, 18, 24),
      labels = c("00:00","06:00","12:00","18:00", "00:00"))

```

wrf_put

Function to write variables in emission files

Description

Extract variable

Usage

```

wrf_put(
  file = file.choose(),
  name = NA,
  POL,
  k,
  check = FALSE,
  verbose = FALSE
)

```

Arguments

file	Character; name of file interactively (default) or specified
name	Character; name of the variable (any variable)
POL	Numeric; emissions input or string/POSIXlt time
k	Numeric; multiplier. If the length is more than 1, it multiplies POL for each value of k. It can be used if you want to add an hourly profile to your emissions.
check	logic (default is FALSE), TRUE to check for NA and negative values and replace with zeros
verbose	display additional information

Value

Add vars to a NetCDF WRF file.

Author(s)

Daniel Schuch and Sergio Ibarra-Espinosa

See Also

[wrf_plot](#) and [wrf_get](#)

Examples

```
{
# create the folder and emission file
dir.create(file.path(tempdir(), "EMISS"))
wrf_create(wrfinput_dir = system.file("extdata", package = "elexport"),
           wrfchemi_dir = file.path(tempdir(), "EMISS"))

# get the name of created file
files <- list.files(path = file.path(tempdir(), "EMISS"),
                   pattern = "wrfchemi",
                   full.names = TRUE)

# open, put some numbers and write
CO <- wrf_get(file = files[1],
              name = "E_CO")

CO[] = rnorm(length(CO))

wrf_put(file = files[1],
        name = "E_CO",
        POL = CO)
}
```

wrf_raster

Creates raster from a variable from a wrf file

Description

Return a Raster

Usage

```
wrf_raster(
  file = file.choose(),
  name = NA,
  latlon = FALSE,
  level = 1,
  use_sf = FALSE,
  as_polygons = FALSE,
  verbose = FALSE
)
```


Arguments

file	wrf file
name	variable name
latlon	project the output in "+proj=longlat +datum=WGS84 +no_defs"
level	only for 4d data, default is 1 (surface)
use_sf	set TRUE to use sf package instead of rgdal to project XLAT/XLONG
as_polygons	logical, true to return a poligon instead of a raster
verbose	display additional information

Value

return a raster from a NetCDF WRF file

Note

newer versions of gdal/rgdal can present a issue related to lat/lon set use_sf to TRUE use functions of sf instead of rgdal

Examples

```
{
wrf <- paste(system.file("extdata", package = "elexport"),
             "/wrfinput_d02", sep="")

r <- wrf_raster(file=wrf, name='XLAT')

library(raster)
plot(r, axes = TRUE)
}
```

wrf_summary

Summary of variables inside WRF files

Description

This return returns a summary for each variable.

Usage

```
wrf_summary(file, vars, clean = FALSE)
```

Arguments

file	String path to the wrf.
vars	String of WRF variables. If missing, all variables.
clean	logical, default is FALSE, TRUE for remove Times, XLAT and XLONG

Value

data.frame

Examples

```
# do not run
file = paste0(system.file("extdata", package = "elexport"), "/wrfinput_d02")
wrf_summary(file = file)
```

Index

* datasets

- edgar, 4
- emis_opt, 5
- emisco, 5
- gCO, 6
- Lights, 9
- rawprofile, 10

as.POSIXct, 13

chem_edgar, 2

edgar, 4

emis_opt, 5, 24

emisco, 5

gCO, 6

get_edgar, 7, 7

Lights, 9, 29

rawprofile, 10

sfx_explode, 11, 11

sp, 13

st_explode, 12, 12

to_as4wrf, 12

to_brams_spm, 14

to_munich, 15, 15

to_rline, 17

to_wrf, 9, 13, 19, 24, 29, 30

wrf_add, 21

wrf_create, 6, 13, 20, 22, 29, 30

wrf_get, 20, 24, 32

wrf_grid, 26

wrf_meta, 27, 27

wrf_plot, 20, 25, 28, 32

wrf_profile, 20, 29

wrf_put, 25, 31

wrf_raster, 32

wrf_summary, 33