

Package: rdkitpyr (via r-universe)

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Type Package

Title Task-Oriented Cheminformatics in R Using 'RDKit' via 'Python'

Version 0.2.1

Description A task-oriented R interface to the 'RDKit' <<https://www.rdkit.org>> library through its 'Python' API via 'reticulate'. The package offers high-level cheminformatics functionality, including molecule parsing, descriptor calculation, and fingerprint generation without replicating the native structure of 'RDKit'.

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URL <https://mass-spec.ru/projects/cheminformatics/rdkitpyr/eng/>

BugReports <https://github.com/AndreySamokhin/rdkitpyr/issues>

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<code>.IsRdkitAvailable</code>	<i>Check whether RDKit is available</i>
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Description

Used in examples and tests to determine whether the Python module `rdkit` is available via **reticulate**.

This function is exported to support examples and tests. It is not part of the stable user-facing API and may change without notice.

The result is cached for the duration of the R session to avoid repeated calls to `reticulate::py_module_available()`.

Usage

```
.IsRdkitAvailable(initialize = TRUE)
```

Arguments

<code>initialize</code>	A logical value. If <code>FALSE</code> and Python is not initialized, RDKit is assumed to be unavailable and the corresponding tests and examples are skipped.
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Value

A logical value indicating whether the `rdkit` Python module is available.

`CalculateAllDescriptors`*Calculate all RDKit molecular descriptors*

Description

Calculate all molecular descriptors available in RDKit for a set of molecules.

The descriptors are calculated using the `CalcMolDescriptors()` function from the `rdkit.Chem.Descriptors` module in RDKit.

The set of returned descriptors may depend on the installed RDKit version.

Each molecule is represented by a full set of descriptor values returned as a data frame. Invalid molecules are represented by rows containing NA values. Row order is preserved so that the output aligns with the input.

Usage

```
CalculateAllDescriptors(mols, verbose = FALSE)
```

Arguments

<code>mols</code>	A character vector of SMILES or InChI strings, or a list of RDKit Mol objects.
<code>verbose</code>	A logical value. If TRUE, display warning and error messages emitted by the RDKit backend. Default is FALSE.

Value

A data frame with one row per molecule and one column per descriptor. Elements corresponding to invalid molecules are returned as NA. Original RDKit descriptor names are used.

Additionally, the "valid" attribute is attached to indicate which molecules were successfully processed.

Examples

```
# Calculate all RDKit descriptors
smiles <- c("CCO", "c1ccccc1", "invalid_molecule")
desc <- CalculateAllDescriptors(smiles)

# Inspect names of first three descriptors
names(desc)[1:3]
#> "MaxAbsEStateIndex" "MaxEStateIndex" "MinAbsEStateIndex"

# Display Molecular weight, LogP, TPSA, and molar refractivity
desc[c("MolWt", "MolLogP", "TPSA", "MolMR")]
#>   MolWt MolLogP  TPSA  MolMR
#> 1 46.069 -0.0014 20.23 12.7598
#> 2 78.114  1.6866  0.00 26.4420
#> 3    NA      NA    NA     NA
```

```
# Check which molecules were successfully processed
attr(desc, "valid")
#> TRUE TRUE FALSE
```

CalculateExactMass *Calculate the exact mass of molecules*

Description

Calculate the exact mass (monoisotopic mass) for a set of molecules.

The calculation is performed using the `ExactMolWt()` function from the `rdkit.Chem.Descriptors` module in RDKit.

Usage

```
CalculateExactMass(mols, verbose = FALSE)
```

Arguments

<code>mols</code>	A character vector of SMILES or InChI strings, or a list of RDKit Mol objects.
<code>verbose</code>	A logical value. If TRUE, display warning and error messages emitted by the RDKit backend. Default is FALSE.

Value

A numeric vector containing the exact mass for each molecule. Elements corresponding to invalid molecules are returned as NA.

Examples

```
# Calculate exact mass for a set of molecules
smiles <- c("CCO", "c1ccccc1", "invalid_molecule")
CalculateExactMass(smiles)
#> 46.04186 78.04695 NA
```

`CalculateMaccsFingerprints`*Calculate MACCS fingerprints for a set of molecules*

Description

Calculate MACCS (Molecular ACCess System) fingerprints for a set of molecules. Each fingerprint is a fixed-length binary vector representing the presence or absence of predefined structural features.

The fingerprints are calculated using the `GetMACCSKeysFingerprint()` function from the `rdkit.Chem.rdMolDescriptors` module in RDKit.

Invalid molecules are represented by NA vectors. Row order is preserved so that the output aligns with the input.

Usage

```
CalculateMaccsFingerprints(mols, verbose = FALSE)
```

Arguments

<code>mols</code>	A character vector of SMILES or InChI strings, or a list of RDKit Mol objects.
<code>verbose</code>	A logical value. If TRUE, display warning and error messages emitted by the RDKit backend. Default is FALSE.

Value

A matrix of integers (0 or 1) with one row per molecule and 167 columns (MACCS keys). Rows corresponding to invalid molecules contain NA.

Additionally, the "valid" attribute is attached to indicate which molecules were successfully processed.

Examples

```
# Calculate MACCS fingerprints
smiles <- c("CCO", "c1ccccc1", "invalid_molecule")
fps <- CalculateMaccsFingerprints(smiles)

# Get the number of fingerprints (columns)
ncol(fps)
#> 167

# Check which molecules were successfully processed
attr(fps, "valid")
#> TRUE TRUE FALSE
```

`CalculateMolecularWeight`*Calculate the average molecular weight of molecules*

Description

Calculate the molecular weight (average mass) for a set of molecules.

The calculation is performed using the `MolWt()` function from the `rdkit.Chem.Descriptors` module in RDKit.

Usage

```
CalculateMolecularWeight(mols, verbose = FALSE)
```

Arguments

<code>mols</code>	A character vector of SMILES or InChI strings, or a list of RDKit Mol objects.
<code>verbose</code>	A logical value. If TRUE, display warning and error messages emitted by the RDKit backend. Default is FALSE.

Value

A numeric vector containing the molecular weight for each molecule. Elements corresponding to invalid molecules are returned as NA.

Examples

```
# Calculate average molecular weight for a set of molecules
smiles <- c("CCO", "c1ccccc1", "invalid_molecule")
CalculateMolecularWeight(smiles)
#> 46.069 78.114 NA
```

`CalculateMorganFingerprints`*Calculate Morgan fingerprints for a set of molecules*

Description

Calculate Morgan (circular) fingerprints for a set of molecules.

The fingerprints are calculated using the `GetMorganGenerator()` function from the `rdkit.Chem.rdFingerprintGenerator` module in RDKit.

Invalid molecules are represented by NA vectors. Row order is preserved so that the output aligns with the input.

Usage

```
CalculateMorganFingerprints(  
  mols,  
  radius = 3L,  
  fp_size = 2048L,  
  count_simulation = FALSE,  
  include_chirality = FALSE,  
  use_bond_types = TRUE,  
  include_ring_membership = TRUE,  
  verbose = FALSE  
)
```

Arguments

<code>mols</code>	A character vector of SMILES or InChI strings, or a list of RDKit Mol objects.
<code>radius</code>	An integer value. Bond radius defining the size of circular substructures.
<code>fp_size</code>	An integer value. Number of bits in the fingerprint.
<code>count_simulation</code>	A logical value. If set, use count simulation while generating the fingerprint.
<code>include_chirality</code>	A logical value. If set, chirality information will be added to the generated fingerprint.
<code>use_bond_types</code>	A logical value. If set, bond types will be included as a part of the default bond invariants.
<code>include_ring_membership</code>	A logical value. If set, whether or not the atom is in a ring will be used in the invariant list.
<code>verbose</code>	A logical value. If TRUE, display warning and error messages emitted by the RDKit backend. Default is FALSE.

Value

A matrix of integers (0 or 1) with one row per molecule and `fp_size` columns. Rows corresponding to invalid molecules contain NA.

Additionally, the "valid" attribute is attached to indicate which molecules were successfully processed.

Examples

```
# Calculate Morgan fingerprints  
smiles <- c("CCO", "c1ccccc1", "invalid_molecule")  
fps <- CalculateMorganFingerprints(smiles)  
  
# Get the number of fingerprints (columns)  
ncol(fps)  
#> 2048
```

```
# Check which molecules were successfully processed
attr(fps, "valid")
#> TRUE TRUE FALSE
```

CalculateRdkitFingerprints

Calculate RDKit topological fingerprints for a set of molecules

Description

Calculate RDKit topological (path-based) fingerprints for a set of molecules.

The fingerprints are calculated using the `RDKFingerprint()` function from the `rdkit.Chem.rdmlops` module in RDKit.

Invalid molecules are represented by NA vectors. Row order is preserved so that the output aligns with the input.

Usage

```
CalculateRdkitFingerprints(
  mols,
  min_path = 1L,
  max_path = 7L,
  fp_size = 2048L,
  n_bits_per_hash = 2L,
  use_hydrogens = TRUE,
  target_density = 0,
  min_size = 128L,
  branched_paths = TRUE,
  use_bond_order = TRUE,
  verbose = FALSE
)
```

Arguments

<code>mols</code>	A character vector of SMILES or InChI strings, or a list of RDKit Mol objects.
<code>min_path</code>	An integer value. Minimum number of bonds to include in the subgraphs.
<code>max_path</code>	An integer value. Maximum number of bonds to include in the subgraphs.
<code>fp_size</code>	An integer value. Number of bits in the fingerprint.
<code>n_bits_per_hash</code>	An integer value. Number of bits to set per path.
<code>use_hydrogens</code>	A logical value. Include paths involving hydrogens in the fingerprint if the molecule has explicit hydrogens.
<code>target_density</code>	A numeric value. Fold the fingerprint until this minimum density has been reached.

min_size	An integer value. The minimum size the fingerprint will be folded to when trying to reach target_density.
branched_paths	A logical value. If set, both branched and unbranched paths will be used in the fingerprint.
use_bond_order	A logical value. If set, both bond orders will be used in the path hashes.
verbose	A logical value. If TRUE, display warning and error messages emitted by the RDKit backend. Default is FALSE.

Value

A matrix of integers (0 or 1) with one row per molecule and fp_size columns. Rows corresponding to invalid molecules contain NA.

Additionally, the "valid" attribute is attached to indicate which molecules were successfully processed.

Examples

```
# Calculate RDKit fingerprints
smiles <- c("CCO", "c1ccccc1", "invalid_molecule")
fps <- CalculateRdkitFingerprints(smiles)

# Get the number of fingerprints (columns)
ncol(fps)
#> 2048

# Check which molecules were successfully processed
attr(fps, "valid")
#> TRUE TRUE FALSE
```

ConvertToInchi

Convert molecules to InChI strings

Description

Convert molecules to InChI strings.

InChI identifiers can technically be provided as input. In this case, the output is expected to be identical to the input. This can be useful only to test RDKit's consistency with challenging molecules.

Usage

```
ConvertToInchi(mols, verbose = FALSE)
```

Arguments

mols	A character vector of SMILES or InChI strings, or a list of RDKit Mol objects.
verbose	A logical value. If TRUE, display warning and error messages emitted by the RDKit backend. Default is FALSE.

Value

A character vector. InChI strings. Elements that cannot be converted are returned as NA.

Examples

```
# Convert a vector of SMILES to InChI identifiers
smiles <- c("CC", "CCC")
ConvertToInchi(smiles)
#> "InChI=1S/C2H6/c1-2/h1-2H3"
#> "InChI=1S/C3H8/c1-3-2/h3H2,1-2H3"

# Providing InChI as input returns identical output
ConvertToInchi("InChI=1S/CH4/h1H4")
#> "InChI=1S/CH4/h1H4"
```

ConvertToInchikey *Convert molecules to InChIKey strings*

Description

Convert molecules to InChIKey strings.

Conversion of an InChI string to an InChIKey relies on the IUPAC library, allowing conversion without creating intermediate RDKit Mol objects.

Usage

```
ConvertToInchikey(mols, verbose = FALSE)
```

Arguments

mols A character vector of SMILES or InChI strings, or a list of RDKit Mol objects.

verbose A logical value. If TRUE, display warning and error messages emitted by the RDKit backend. Default is FALSE.

Value

A character vector. InChIKey strings. Elements that cannot be converted are returned as NA.

Examples

```
# Convert a vector of InChI to InChIKey identifiers
inchi <- c("InChI=1S/C2H6/c1-2/h1-2H3",
           "InChI=1S/C6H6/c1-2-4-6-5-3-1/h1-6H")
ConvertToInchikey(inchi)
#> "OTMSDBZUPAUEDD-UHFFFAOYSA-N"
#> "UHOVQNZJYSORNB-UHFFFAOYSA-N"
```

```
# Convert a vector of SMILES to InChIKey identifiers
smiles <- c("CC", "c1ccccc1")
ConvertToInchikey(smiles)
#> "OTMSDBZUPAUEDD-UHFFFAOYSA-N"
#> "UHOVQNZJYSORNB-UHFFFAOYSA-N"
```

ConvertToSmiles

Convert molecules to SMILES strings

Description

Convert molecules to SMILES strings.

SMILES strings can be provided as input to obtain their canonical form or to remove stereochemistry.

Usage

```
ConvertToSmiles(  
  mols,  
  isomeric = TRUE,  
  kekulize = FALSE,  
  canonical = TRUE,  
  explicit_bonds = FALSE,  
  explicit_hydrogens = FALSE,  
  verbose = FALSE  
)
```

Arguments

mols	A character vector of SMILES or InChI strings, or a list of RDKit Mol objects.
isomeric	A logical value. If TRUE, include stereochemistry in the output.
kekule	A logical value. If TRUE, kekulize aromatic bonds.
canonical	A logical value. If TRUE, generate canonical SMILES.
explicit_bonds	A logical value. If TRUE, make all bonds explicit.
explicit_hydrogens	A logical value. If TRUE, make all hydrogens explicit.
verbose	A logical value. If TRUE, display warning and error messages emitted by the RDKit backend. Default is FALSE.

Value

A character vector. SMILES strings. Elements that cannot be converted are returned as NA.

Examples

```
# Convert a vector of InChI identifiers to canonical SMILES
inchi <- c("InChI=1S/C2H6/c1-2/h1-2H3",
          "InChI=1S/C6H6/c1-2-4-6-5-3-1/h1-6H")
ConvertToSmiles(inchi)
#> "CC"
#> "c1ccccc1"

# Convert a vector of SMILES to SMILES with Kekulized aromatic bonds
smiles <- c("c1ccccc1", "c1ccc2ccccc2c1")
ConvertToSmiles(smiles, kekule = TRUE)
#> "C1=CC=CC=C1"
#> "C1=CC=C2C=CC=CC2=C1"
```

GetPythonInfo

Get information about the currently used Python environment

Description

Get details about the Python interpreter, numpy and rdkit packages.

Usage

```
GetPythonInfo(verbose = TRUE)
```

Arguments

verbose A logical value. If TRUE (default), function prints information about currently used Python environment.

Value

Invisibly return a named list with the following components:

python_path Full path to the Python executable.

python_version Version of the Python interpreter.

numpy_version Installed version of the numpy package.

rdkit_version Installed version of the rdkit package.

forced_by If applicable, indicates whether the Python interpreter was forced via RETICULATE_PYTHON, use_*(), or py_require().

Examples

```
# Print information about the Python environment
GetPythonInfo()

# Access programmatically
py_env <- GetPythonInfo(verbose = FALSE)
py_env$python_version
py_env$rdkit_version
```

ParseMolecules

Parse SMILES and InChI strings into RDKit Mol objects

Description

Parse SMILES and InChI strings into RDKit Mol objects

This function converts a character vector of molecular representations (SMILES or InChI) into a list of RDKit Mol objects (Python-backed pointers via *reticulate*). The resulting objects can be reused in subsequent operations without repeated conversion from SMILES or InChI. This is particularly useful when multiple cheminformatics tasks are performed on the same set of molecules, improving efficiency by avoiding repeated parsing steps.

Usage

```
ParseMolecules(mols, verbose = FALSE)
```

Arguments

mols	A character vector of SMILES or InChI strings.
verbose	A logical value. If TRUE, display warning and error messages emitted by the RDKit backend. Default is FALSE.

Value

A list of RDKit Mol objects.

Examples

```
# Convert a vector of SMILES to RDKit Mol objects
mols <- ParseMolecules(c("CC", "CCC"))
print(mols[[1L]])
#> <rdkit.Chem.rdchem.Mol object at 0x000001CC4D60F4C0>

# Convert a list of RDKit Mol objects to InChI identifiers
ConvertToInchi(mols)
#> "InChI=1S/C2H6/c1-2/h1-2H3"
#> "InChI=1S/C3H8/c1-3-2/h3H2,1-2H3"
```

test_compounds	<i>Test compounds</i>
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Description

A small reference set of chemical compounds extracted from the PubChem database. Despite its limited size, the dataset covers a broad range of chemical properties, including:

- neutral, charged, and radical species;
- aromatic and aliphatic compounds;
- molecules containing heteroatoms;
- isotopically labeled compounds;
- stereochemistry;
- species with disconnected fragments.

Usage

test_compounds

Format

A data frame.

pubchem_cid PubChem compound identifier (CID).

name Compound name (as reported in the PubChem database).

smiles SMILES identifier.

inchi InChI identifier.

inchikey InChIKey identifier.

n_labeled_atoms Number of isotopically labeled atoms.

charge The total charge of a molecule.

exact_mass Exact monoisotopic mass.

molecular_weight Average molecular weight.

formula Molecular formula.

formula_isotopes Molecular formula with explicit isotopes.

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