NAME

AtomTypesFingerprints

SYNOPSIS

use Fingerprints::AtomTypesFingerprints;

use Fingerprints::AtomTypesFingerprints qw(:all);

DESCRIPTION

AtomTypesFingerprints class provides the following methods:

new, GenerateFingerprints, GetDescription, SetAtomIdentifierType, SetAtomTypesSetToUse, SetAtomicInvariantsToUse, SetFunctionalClassesToUse, SetType, StringifyAtomTypesFingerprints

AtomTypesFingerprints is derived from Fingerprints class which in turn is derived from ObjectProperty base class that provides methods not explicitly defined in AtomNeighborhoodsFingerprints, Fingerprints or ObjectProperty classes using Perl's AUTOLOAD functionality. These methods are generated on-the-fly for a specified object property:

```
Set<PropertyName>(<PropertyValue>);
$PropertyValue = Get<PropertyName>();
Delete<PropertyName>();
```

The current release of MayaChemTools supports generation of AtomTypesFingerpritns corresponding to following AtomI dentifierTypes:

```
AtomicInvariantsAtomTypes, DREIDINGAtomTypes, EStateAtomTypes, FunctionalClassAtomTypes, MMFF94AtomTypes, SLogPAtomTypes, SYBYLAtomTypes, TPSAAtomTypes, UFFAtomTypes
```

Based on the values specified for AtomI dentifierType along with other specified parameters such as AtomicInvariantsToUse and FunctionalClassesToUse, initial atom types are assigned to all non-hydrogen atoms or all atoms in a molecule.

Using the assigned atom types and specified Type, one of the following types of fingerprints are generated:

```
AtomTypesCount - A vector containing count of atom types
AtomTypesBits - A bit vector indicating presence/absence of atom types
```

For AtomTypesCount fingerprints, two types of atom types set size is allowed:

```
ArbitrarySize - Corresponds to only atom types detected in molecule FixedSize - Corresponds to fixed number of atom types previously defined
```

For AtomTypesBits fingerprints, only FixedSize atom type set is allowed.

ArbitrarySize corresponds to atom types detected in a molecule where as FixedSize implies a fix number of all possible atom types previously defined for a specific AtomIdentifierType.

Fix number of all possible atom types for supported *AtomIdentifierTypes* in current release of MayaChemTools are:

AtomIdentifier	Total	TotalWithoutHydrogens
DREIDINGAtomTypes	37	34
EStateAtomTypes	109	87
MMFF94AtomTypes	212	171
SLogPAtomTypes	72	67
SYBYLAtomTypes	45	44
TPSAAtomTypes	47	47
UFFAtomTypes	126	124

Combination of Type and AtomTypesSetToUse along with AtomtomI dentifierType allows generation of following different atom types fingerprints:

Type AtomIdentifierType AtomTypesSetToUse

AtomTypesCount	AtomicInvariantsAtomTypes	ArbitrarySize
AtomTypesCount	DREIDINGAtomTypes	ArbitrarySize
AtomTypesCount	DREIDINGAtomTypes	FixedSize
AtomTypesBits	DREIDINGAtomTypes	FixedSize
AtomTypesCount	EStateAtomTypes	ArbitrarySize
AtomTypesCount	EStateAtomTypes	FixedSize
AtomTypesBits	EStateAtomTypes	FixedSize
AtomTypesCount	FunctionalClassAtomTypes	ArbitrarySize
AtomTypesCount	MMFF94AtomTypes	ArbitrarySize
AtomTypesCount	MMFF94AtomTypes	FixedSize
AtomTypesBits	MMFF94AtomTypes	FixedSize
AtomTypesCount	SLogPAtomTypes	ArbitrarySize
AtomTypesCount	SLogPAtomTypes	FixedSize
AtomTypesBits AtomTypesCount	SLogPAtomTypes SYBYLAtomTypes	FixedSize ArbitrarySize
AtomTypesCount	SYBYLAtomTypes	FixedSize
AtomTypesBits	SYBYLAtomTypes	FixedSize
AtomTypesCount	TPSAAtomTypes	FixedSize
AtomTypesBits	TPSAAtomTypes	FixedSize
AtomTypesCount	UFFAtomTypes	ArbitrarySize
AtomTypesCount	UFFAtomTypes	FixedSize
AtomTypesBits	UFFAtomTypes	FixedSize

The current release of MayaChemTools generates the following types of atom types fingerprints bit-vector and vector strings:

FingerprintsVector;AtomTypesCount:AtomicInvariantsAtomTypes:ArbitrarySize;10;NumericalValues;IDsAndValuesString;C.X1.B01.H3 C.X2.B02.H2 C.X2.B03.H1 C.X3.B03.H1 C.X3.B04 F.X1.B01 N.X2.B02.H1 N.X3.B03 O.X1.B01.H1 O.X1.B02;2 4 14 3 10 1 1 1 3 2

FingerprintsVector;AtomTypesCount:DREIDINGAtomTypes:ArbitrarySize;8;Nu mericalValues;IDsAndValuesString;C_2 C_3 C_R F_ N_3 N_R O_2 O_3;2 9 22 1 1 1 2 3

FingerprintsVector;AtomTypesCount:DREIDINGAtomTypes:FixedSize;34;Order edNumericalValues;IDsAndValuesString;B_3 B_2 C_3 C_R C_2 C_1 N_3 N_R N _2 N_1 O_3 O_R O_2 O_1 F_ Al3 Si3 P_3 S_3 Cl Ga3 Ge3 As3 Se3 Br In3 Sn 3 Sb3 Te3 I_ Na Ca Fe Zn;0 0 9 22 2 0 1 1 0 0 3 0 2 0 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0

FingerprintsVector;AtomTypesCount:EStateAtomTypes:ArbitrarySize;11;Num ericalValues;IDsAndValuesString;aaCH aasC aasN dO dssC sCH3 sF sOH ssC H2 ssNH sssCH;14 8 1 2 2 2 1 3 4 1 3

FingerprintsVector;AtomTypesCount:FunctionalClassAtomTypes:ArbitrarySize;8;NumericalValues;IDsAndValuesString;Ar Ar.HBA HBA HBA.HBD HBD Hal NI None;22 1 2 3 1 1 1 10

FingerprintsVector;AtomTypesCount:MMFF94AtomTypes:ArbitrarySize;13;Num ericalValues;IDsAndValuesString;C5A C5B C=ON CB COO CR F N5 NC=O O=CN O=CO OC=O OR;2 2 1 18 1 9 1 1 1 1 1 1 2

FingerprintsVector;AtomTypesCount:SLogPAtomTypes:ArbitrarySize;16;Nume ricalValues;IDsAndValuesString;C1 C10 C11 C14 C18 C20 C21 C22 C5 CS F N11 N4 O10 O2 O9;5 1 1 1 14 4 2 1 2 2 1 1 1 1 3 1

FingerprintsVector;AtomTypesCount:SYBYLAtomTypes:ArbitrarySize;9;Numer
icalValues;IDsAndValuesString;C.2 C.3 C.ar F N.am N.ar 0.2 0.3 0.co2;2
9 22 1 1 1 1 2 2

```
FingerprintsVector;AtomTypesCount:UFFAtomTypes:ArbitrarySize;8;Numeric
alValues;IDsAndValuesString;C_2 C_3 C_R F_ N_3 N_R O_2 O_3;2 9 22 1 1
1 2 3
```

METHODS

new

```
$NewAtomTypesFingerprints = new AtomTypesFingerprints(%NamesAndValues);
```

Using specified *AtomTypesFingerprints* property names and values hash, new method creates a new object and returns a reference to newly created PathLengthFingerprints object. By default, the following properties are initialized:

```
Molecule = '';
   Type = ''
   AtomIdentifierType = ''
   AtomTypesSetToUse = ''
    IgnoreHydrogens = 1
    AtomicInvariantsToUse = ['AS', 'X', 'BO', 'H', 'FC', 'MN']
    FunctionalClassesToUse = ['HBD', 'HBA', 'PI', 'NI', 'Ar', 'Hal']
Examples:
    $AtomTypesFingerprints = new AtomTypesFingerprints(
                               'Molecule' => $Molecule,
                               'Type' => 'AtomTypesCount',
                               'AtomIdentifierType' =>
                                               'AtomicInvariantsAtomTypes');
    $AtomTypesFingerprints = new AtomTypesFingerprints(
                               'Molecule' => $Molecule,
                               'Type' => 'AtomTypesCount',
                               'AtomIdentifierType' =>
                                               'AtomicInvariantsAtomTypes',
                               'AtomicInvariantsToUse' =>
                                               ['AS', 'X', 'BO', 'H', 'FC'] );
    $AtomTypesFingerprints = new AtomTypesFingerprints(
                               'Molecule' => $Molecule,
                               'Type' => 'AtomTypesCount',
                               'AtomIdentifierType' =>
                                               'DREIDINGAtomTypes');
    $AtomTypesFingerprints = new AtomTypesFingerprints(
                               'Molecule' => $Molecule,
                               'Type' => 'AtomTypesCount',
                               'AtomIdentifierType' =>
```

'AtomTypesSetToUse' =>

'EStateAtomTypes',

```
'ArbitrarySize');
```

```
$AtomTypesFingerprints = new AtomTypesFingerprints(
                           'Molecule' => $Molecule,
                           'Type' => 'AtomTypesCount'
                           'AtomIdentifierType' =>
                                           'SLogPAtomTypes',
                           'AtomTypesSetToUse' =>
                                           'FixedSize');
$AtomTypesFingerprints = new AtomTypesFingerprints(
                           'Molecule' => $Molecule,
                           'Type' => 'AtomTypesBits',
                           'AtomIdentifierType' =>
                                           'MMFF94AtomTypes',
                           'AtomTypesSetToUse' =>
                                           'FixedSize');
$AtomTypesFingerprints->GenerateFingerprints();
print "$AtomTypesFingerprints\n";
```

GenerateFingerprints

```
$AtomTypesFingerprints->GenerateFingerprints();
```

Generates atom types fingerprints and returns AtomTypesFingerprints.

GetDescription

```
$Description = $AtomTypesFingerprints->GetDescription();
```

Returns a string containing description of atom types fingerprints.

SetAtomI dentifierType

```
$AtomTypesFingerprints->SetAtomIdentifierType($IdentifierType);
```

Sets atom *IdentifierType* to use during atom types fingerprints generation and returns *AtomTypesFingerprints*.

Possible values: AtomicInvariantsAtomTypes, DREIDINGAtomTypes, EStateAtomTypes, FunctionalClassAtomTypes, MMFF94AtomTypes, SLogPAtomTypes, SYBYLAtomTypes, UFFAtomTypes.

SetAtomTypesSetToUse

```
$AtomTypesFingerprints->SetAtomTypesSetToUse($Value);
```

Sets Value of AtomTypesSetToUse and returns AtomTypesFingerprints. Possible values: ArbitrarySize or FixedSize. Default for AtomTypesCount value of AtomTypesSetToUse: ArbitrarySize.

SetAtomicInvariantsToUse

```
$AtomTypesFingerprints->SetAtomicInvariantsToUse($ValuesRef);
$AtomTypesFingerprints->SetAtomicInvariantsToUse(@Values);
```

Sets atomic invariants to use during *AtomicInvariantsAtomTypes* value of *AtomIdentifierType* for atom neighborhood fingerprints generation and returns *AtomTypesFingerprints*.

Possible values for atomic invariants are: AS, X, BO, LBO, SB, DB, TB, H, Ar, RA, FC, MN, SM. Default value: AS, X, BO, H, FC.

The atomic invariants abbreviations correspond to:

```
AS = Atom symbol corresponding to element symbol
```

```
X<n> = Number of non-hydrogen atom neighbors or heavy atoms  
BO<n> = Sum of bond orders to non-hydrogen atom neighbors or heavy atoms  
LBO<n> = Largest bond order of non-hydrogen atom neighbors or heavy atoms  
SB<n> = Number of single bonds to non-hydrogen atom neighbors or heavy atoms  
DB<n> = Number of double bonds to non-hydrogen atom neighbors or heavy atoms  
TB<n> = Number of triple bonds to non-hydrogen atom neighbors or heavy atoms  
H<n> = Number of implicit and explicit hydrogens for atom  
Ar = Aromatic annotation indicating whether atom is aromatic
```

```
RA = Ring atom annotation indicating whether atom is a ring
FC<+n/-n> = Formal charge assigned to atom
MN<n> = Mass number indicating isotope other than most abundant isotope
SM<n> = Spin multiplicity of atom. Possible values: 1 (singlet), 2 (doublet) or
3 (triplet)
```

Atom type generated by AtomTypes::AtomicInvariantsAtomTypes class corresponds to:

```
AS.X<n>.BO<n>.LBO<n>.<SB><n>.<DB><n>..TB><n>.H<n>.Ar.RA.FC<+n/-n>.MN<n>.SM<n>
```

Except for AS which is a required atomic invariant in atom types, all other atomic invariants are optional. Atom type specification doesn't include atomic invariants with zero or undefined values.

In addition to usage of abbreviations for specifying atomic invariants, the following descriptive words are also allowed:

```
X : NumOfNonHydrogenAtomNeighbors or NumOfHeavyAtomNeighbors
BO : SumOfBondOrdersToNonHydrogenAtoms or SumOfBondOrdersToHeavyAtoms
LBO : LargestBondOrderToNonHydrogenAtoms or LargestBondOrderToHeavyAtoms
SB : NumOfSingleBondsToNonHydrogenAtoms or NumOfSingleBondsToHeavyAtoms
DB : NumOfDoubleBondsToNonHydrogenAtoms or NumOfDoubleBondsToHeavyAtoms
TB : NumOfTripleBondsToNonHydrogenAtoms or NumOfTripleBondsToHeavyAtoms
H : NumOfImplicitAndExplicitHydrogens
Ar : Aromatic
RA : RingAtom
FC : FormalCharge
MN : MassNumber
SM : SpinMultiplicity
```

AtomTypes::AtomicInvariantsAtomTypes module is used to assign atomic invariant atom types.

SetFunctionalClassesToUse

```
$AtomTypesFingerprints->SetFunctionalClassesToUse($ValuesRef);
$AtomTypesFingerprints->SetFunctionalClassesToUse(@Values);
```

Sets functional classes invariants to use during FunctionalClassAtomTypes value of AtomIdentifierType for atom types fingerprints generation and returns AtomTypesFingerprints.

Possible values for atom functional classes are: Ar, CA, H, HBA, HBD, Hal, NI, PI, RA. Default value [Ref 24]: HBD, HBA, PI, NI, Ar, Hal.

The functional class abbreviations correspond to:

```
HBD: HydrogenBondDonor
HBA: HydrogenBondAcceptor
PI : PositivelyIonizable
NI : NegativelyIonizable
Ar : Aromatic
Hal : Halogen
H : Hydrophobic
RA : RingAtom
CA : ChainAtom
```

Functional class atom type specification for an atom corresponds to:

```
Ar.CA.H.HBA.HBD.Hal.NI.PI.RA or None
```

AtomTypes::FunctionalClassAtomTypes module is used to assign functional class atom types. It uses following definitions [Ref 60-61, Ref 65-66]:

```
HydrogenBondDonor: NH, NH2, OH
HydrogenBondAcceptor: N[!H], O
PositivelyIonizable: +, NH2
NegativelyIonizable: -, C(=0)OH, S(=0)OH, P(=0)OH
```

SetType

```
$AtomTypesFingerprints->SetType($Type);
```

Sets type of AtomTypes fingerpritns and returns *AtomTypesFingerprints*. Possible values: *AtomTypesFingerprintsBits or AtomTypesFingerprintsCount*.

StringifyAtomTypesFingerprints

\$String = \$AtomTypesFingerprints->StringifyAtomTypesFingerprints();

Returns a string containing information about AtomTypesFingerprints object.

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SEE ALSO

Fingerprints.pm, FingerprintsStringUtil.pm, AtomNeighborhoodsFingerprints.pm, EStateIndiciesFingerprints.pm, ExtendedConnectivityFingerprints.pm, MACCSKeys.pm, PathLengthFingerprints.pm, TopologicalAtomPairsFingerprints.pm, TopologicalAtomTripletsFingerprints.pm, TopologicalAtomTorsionsFingerprints.pm, TopologicalPharmacophoreAtomPairsFingerprints.pm,

Topological Pharmacophore Atom Triplets Finger prints.pm

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