NAME

MACCSKeys

SYNOPSIS

```
use Fingerprints::MACCSKeys;
use Fingerprints::MACCSKeys qw(:all);
```

DESCRIPTION

MACCSKeys [Ref 45-47] class provides the following methods:

new, GenerateFingerprints, GenerateMACCSKeys, GetDescription, SetSize, SetType, StringifyMACCSKeys

MACCSKeys is derived from Fingerprints class which in turn is derived from ObjectProperty base class that provides methods not explicitly defined in MACCSKeys, 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>();
```

For each MACCS (Molecular ACCess System) keys definition, atoms are processed to determine their membership to the key and the appropriate molecular fingerprints strings are generated. An atom can belong to multiple MACCS keys.

For *MACCSKeyBits* value of Type option, a fingerprint bit-vector string containing zeros and ones is generated and for *MACCSKeyCount* value, a fingerprint vector string corresponding to number of MACCS keys [Ref 45-47] is generated.

MACCSKeyBits or MACCSKeyCount values for Type along with two possible 166 | 322 values of Size supports generation of four different types of MACCS keys fingerprint: MACCS166KeyBits, MACCS166KeyCount, MACCS322KeyBits, MACCS322KeyCount.

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

FingerprintsBitVector;MACCSKeyBits;166;HexadecimalString;Ascending;000 000000021210210e845f8d8c60b79dffbfffd1

METHODS

new

```
$NewMACCSKeys = new MACCSKeys(%NamesAndValues);
```

Using specified *MACCSKeys* 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 = ''
Size = ''
```

Examples:

GetDescription

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

Returns a string containing description of MACCS keys fingerprints.

GenerateMACCSKeys or GenerateFingerprints

```
$MACCSKeys = $MACCSKeys->GenerateMACCSKeys();
```

Generates MACCS keys fingerprints and returns MACCSKeys.

For *MACCSKeyBits* value of Type, a fingerprint bit-vector string containing zeros and ones is generated and for *MACCSKeyCount* value, a fingerprint vector string corresponding to number of MACCS keys is generated.

MACCSKeyBits or MACCSKeyCount values for Type option along with two possible 166 | 322 values of Size supports generation of four different types of MACCS keys fingerprint: MACCS166KeyBits, MACCS166KeyCount, MACCS322KeyBits, MACCS322KeyCount.

Definition of MACCS keys uses the following atom and bond symbols to define atom and bond environments:

```
Atom symbols for 166 keys [ Ref 47 ]:
A : Any valid periodic table element symbol
Q : Hetro atoms; any non-C or non-H atom
X : Halogens; F, Cl, Br, I
  : Others; other than H, C, N, O, Si, P, S, F, Cl, Br, I
Atom symbols for 322 keys [ Ref 46 ]:
A : Any valid periodic table element symbol
Q : Hetro atoms; any non-C or non-H atom
X : Others; other than H, C, N, O, Si, P, S, F, Cl, Br, I
Z is neither defined nor used
Bond types:
- : Single
  : Double
  : Triple
  : Triple
  : Single or double guery bond
% : An aromatic query bond
None : Any bond type; no explicit bond specified
$ : Ring bond; $ before a bond type specifies ring bond
! : Chain or non-ring bond; ! before a bond type specifies chain bond
  : A ring linkage and the number following it specifies the
     atoms position in the line, thus @1 means linked back to the first
     atom in the list.
Aromatic: Kekule or Arom5
Kekule: Bonds in 6-membered rings with alternate single/double bonds
       or perimeter bonds
Arom5: Bonds in 5-membered rings with two double bonds and a hetro
        atom at the apex of the ring.
```

MACCS 166 keys [Ref 45-47] are defined as follows:

```
Key Description
1 ISOTOPE
2 103 < ATOMIC NO. < 256
3 GROUP IVA, VA, VIA PERIODS 4-6 (Ge...)
4 ACTINIDE
5 GROUP IIIB, IVB (Sc...)
6 LANTHANIDE
7 GROUP VB, VIB, VIIB (V...)
8 QAAA@1
9 GROUP VIII (Fe...)
10 GROUP IIA (ALKALINE EARTH)
11 4M RING
12 GROUP IB, IIB (Cu...)
13 ON(C)C
14 S-S
15 OC(0)0
16 OAA@1
17 CTC
18 GROUP IIIA (B...)
19 7M RING
20 SI
21 C=C(Q)Q
22 3M RING
23 NC(0)0
24 N-O
25 NC(N)N
26 C$=C($A)$A
27 I
28 QCH2Q
29 P
30 CQ(C)(C)A
31 QX
32 CSN
33 NS
34 CH2=A
35 GROUP IA (ALKALI METAL)
36 S HETEROCYCLE
37 NC(O)N
38 NC(C)N
39 OS(O)O
40 S-O
41 CTN
42 F
43 QHAQH
44 OTHER
45 C=CN
46 BR
47 SAN
48 00(0)0
49 CHARGE
50 C=C(C)C
51 CSO
52 NN
53 ОНАААОН
54 QHAAQH
55 OSO
56 ON(O)C
57 O HETEROCYCLE
58 QSQ
59 Snot%A%A
60 S=0
61 AS(A)A
62 A$A!A$A
63 N=O
64 A$A!S
65 C%N
66 CC(C)(C)A
67 QS
68 QHQH (&...)
69 QQH
70 QNQ
71 NO
72 OAAO
73 S=A
74 CH3ACH3
```

75 A!N\$A 76 C=C(A)A

```
77 NAN
78 C=N
```

79 NAAN

- 80 NAAAN
- 81 SA(A)A
- 82 ACH2QH
- 83 QAAAA@1
- 84 NH2
- 85 CN(C)C
- 86 CH2QCH2
- 87 X!A\$A
- 88 S
- 89 OAAAO
- 90 QHAACH2A
- 91 QHAAACH2A
- 92 OC(N)C
- 93 QCH3
- 94 ON
- 95 NAAO
- 96 5M RING
- 97 NAAAO
- 98 QAAAAA@1
- 99 C=C
- 100 ACH2N
- 101 8M RING
- 102 QO
- 103 CL
- 104 QHACH2A
- 105 A\$A(\$A)\$A
- 106 QA(Q)Q
- 107 XA(A)A
- 108 CH3AAACH2A
- 109 ACH20
- 110 NCO
- 111 NACH2A
- 112 AA(A)(A)A
- 113 Onot%A%A
- 114 CH3CH2A
- 115 CH3ACH2A
- 116 CH3AACH2A
- 117 NAO
- 118 ACH2CH2A > 1
- 119 N=A
- 120 HETEROCYCLIC ATOM > 1 (&...)
- 121 N HETEROCYCLE
- 122 AN(A)A
- 123 OCO
- 124 QQ
- 125 AROMATIC RING > 1
- 126 A!O!A
- 127 A\$A!O > 1 (&...)
- 128 ACH2AAACH2A
- 129 ACH2AACH2A
- 130 QQ > 1 (&...)
- 131 QH > 1
- 132 OACH2A
- 133 A\$A!N
- 134 X (HALOGEN)
- 135 Nnot%A%A
- 136 O=A > 1
- 137 HETEROCYCLE 138 QCH2A > 1 (&...)
- 139 OH
- 140 0 > 3 (&...)
- 141 CH3 > 2 (&...)
- 142 N > 1
- 143 A\$A!O
- 144 Anot%A%Anot%A
- 145 6M RING > 1
- 146 0 > 2
- 147 ACH2CH2A
- 148 AQ(A)A
- 149 CH3 > 1 150 A!A\$A!A
- 151 NH
- 152 OC(C)C
- 153 OCH2A
- 154 C=O

```
155 A!CH2!A

156 NA(A)A

157 C-O

158 C-N

159 O > 1

160 CH3

161 N

162 AROMATIC

163 6M RING

164 O

165 RING

166 FRAGMENTS
```

MACCS 322 keys set as defined in tables 1, 2 and 3 [Ref 46] include:

- o 26 atom properties of type P, as listed in Table 1
- o 32 one-atom environments, as listed in Table 3
- o 264 atom-bond-atom combinations listed in Table 4

Total number of keys in three tables is: 322

Atom symbol, X, used for 322 keys [Ref 46] doesn't refer to Halogens as it does for 166 keys. In order to keep the definition of 322 keys consistent with the published definitions, the symbol X is used to imply "others" atoms, but it's internally mapped to symbol X as defined for 166 keys during the generation of key values.

Atom properties-based keys (26):

```
Key
      Description
      A(AAA) or AA(A)A - atom with at least three neighbors
      O - heteroatom
2
      Anot*not-A - atom involved in one or more multiple bonds, not aromatic
4
      A(AAAA) or AA(A)(A)A - atom with at least four neighbors
5
      A(QQ) or QA(Q) - atom with at least two heteroatom neighbors
      A(QQQ) or QA(Q)Q - atom with at least three heteroatom neighbors
6
      QH - heteroatom with at least one hydrogen attached
      \ensuremath{\mathtt{CH2}}\xspace(\mathtt{AA}) or \mathtt{ACH2A} - carbon with at least two single bonds and at least
8
      two hydrogens attached
9
      CH3(A) or ACH3 - carbon with at least one single bond and at least three
      hydrogens attached
10
      Halogen
11
      A(-A-A-A) or A-A(-A)-A - atom has at least three single bonds
12
      AAAAAA@1 > 2 - atom is in at least two different six-membered rings
13
      A(\$A\$A\$A) or A\$A(\$A)\$A - atom has more than two ring bonds
14
      A$A!A$A - atom is at a ring/chain boundary. When a comparison is done
      with another atom the path passes through the chain bond.
15
      Anot%A%Anot%A - atom is at an aromatic/nonaromatic boundary. When a
      comparison is done with another atom the path
      passes through the aromatic bond.
16
      A!A!A - atom with more than one chain bond
17
      A!A$A!A - atom is at a ring/chain boundary. When a comparison is done
      with another atom the path passes through the ring bond.
      A%Anot%A%A - atom is at an aromatic/nonaromatic boundary. When a
18
      comparison is done with another atom the
      path passes through the nonaromatic bond.
19
      HETEROCYCLE - atom is a heteroatom in a ring.
20
      rare properties: atom with five or more neighbors, atom in
      four or more rings, or atom types other than
      H, C, N, O, S, F, Cl, Br, or I
21
      rare properties: atom has a charge, is an isotope, has two or
      more multiple bonds, or has a triple bond.
22
      N - nitrogen
      S - sulfur
23
      0 - oxygen
24
25
      A(AA)A(A)A(AA) - atom has two neighbors, each with three or
      more neighbors (including the central atom).
26
      CHACH2 - atom has two hydrocarbon (CH2) neighbors
```

Atomic environments properties-based keys (32):

```
Description
Key
      C(CC)
27
28
      C(CCC)
29
      C(CN)
30
      C(CCN)
31
      C(NN)
32
      C(NNC)
33
      C(NNN)
34
      C(CO)
35
      C(CCO)
36
      C(NO)
37
      C(NCO)
```

```
38
      C(NNO)
39
      C(00)
40
      C(C00)
41
      C(NOO)
42
      C(000)
43
      Q(CC)
44
      Q(CCC)
45
      Q(CN)
46
      Q(CCN)
47
      Q(NN)
48
      Q(CNN)
49
      Q(NNN)
50
      Q(CO)
51
      Q(CCO)
52
      Q(NO)
53
      Q(CNO)
54
      Q(NNO)
55
      0(00)
56
      Q(C00)
57
      Q(NOO)
58
      Q(000)
```

Note: The first symbol is the central atom, with atoms bonded to the central atom listed in parentheses. Q is any non-C, non-H atom. If only two atoms are in parentheses, there is no implication concerning the other atoms bonded to the central atom.

Atom-Bond-Atom properties-based keys: (264)

```
Key
      Description
59
      C-C
60
      C-N
61
      C-0
62
      C-S
63
      C-Cl
64
      C-P
65
      C-F
66
      C-Br
67
      C-Si
68
      C-I
69
      C-X
70
      N-N
71
      N-O
72
      N-S
73
      N-Cl
74
      N-P
75
      N-F
76
      N-Br
77
      N-Si
78
      N-I
79
      N-X
80
      0-0
81
      0-S
      0-C1
82
83
      O-P
84
      O-F
85
      0-Br
86
      O-Si
87
      0-I
88
      O-X
89
      S-S
90
      S-Cl
91
      S-P
92
      S-F
93
      S-Br
94
      S-Si
95
96
      S-X
97
      Cl-Cl
98
      Cl-P
99
      Cl-F
100
      Cl-Br
      Cl-Si
101
102
      Cl-I
      Cl-X
103
104
      P-P
105
      P-F
106
      P-Br
107
      P-Si
```

108

P-I

- 109 P-X
- 110 F-F
- 111 F-Br
- 112 F-Si
- 113 F-I
- 114 F-X
- 115 Br-Br
- 116 Br-Si
- 117 Br-I
- 118 Br-X
- 119 Si-Si
- 120 Si-I
- Si-X 121
- 122 I-I
- 123 I-X
- 124 X-X
- 125 C=C
- 126 C=N
- 127 C=O
- 128 C=S
- 129 C=Cl
- 130
- C=P
- 131 C=F
- 132 C=Br
- 133 C=Si
- 134 C=I
- 135 C=X
- 136 N=N
- 137 N=O
- 138 N=S
- 139 N=Cl
- 140 N=P
- 141 N=F
- 142 N=Br
- 143 N=Si
- 144 N=I
- 145 N=X 146 0=0
- 147 0=S
- 148 O=Cl
- 149 O=P
- 150 O=F
- 151 O=Br
- 152 O=Si
- O=I 153
- 154 O=X 155 S=S
- 156 S=Cl
- 157 S=P
- 158 S=F
- 159 S=Br
- 160 S=Si
- 161 S=I
- 162 S=X
- 163 Cl=Cl 164 Cl=P
- 165 Cl=F
- 166 Cl=Br
- 167 Cl=Si
- 168 Cl=I
- 169 Cl=X
- 170 P=P
- 171 P=F
- 172 P=Br
- 173 P=Si
- 174 P=I
- 175 P=X 176 F=F
- 177 F=Br
- 178 F=Si
- 179 F=I
- 180 F=X181 Br=Br
- 182 Br=Si
- 183 Br=I
- 184 Br=X 185 Si=Si
- 186 Si=I

```
187
      Si=X
188
      I = I
189
      I=X
190
      X=X
191
      C#C
192
      C#N
193
      C#0
194
      C#S
195
      C#Cl
196
      C#P
197
      C#F
198
      C#Br
199
      C#Si
200
      C#I
201
      C#X
202
      N#N
203
      N#O
204
      N#S
205
      N#Cl
206
      N#P
207
      N#F
208
      N#Br
209
      N#Si
210
      N#I
211
      N#X
212
      0#0
213
      O#S
214
      O#Cl
215
      O#P
216
      O#F
217
      O#Br
218
      O#Si
219
      O#I
220
      О#Х
221
      S#S
222
      S#Cl
223
      S#P
224
      S#F
225
      S#Br
226
      S#Si
227
      S#I
228
      S#X
229
      Cl#Cl
230
      Cl#P
231
      Cl#F
232
      Cl#Br
233
      Cl#Si
234
      Cl#I
235
      Cl#X
236
      P#P
237
      P#F
238
      P#Br
239
      P#Si
240
      P#I
241
      P#X
242
      F#F
243
      F#Br
244
      F#Si
245
      F#I
246
      F#X
247
      Br#Br
248
      Br#Si
249
      Br#I
250
      Br#X
251
      Si#Si
252
      Si#I
253
      Si#X
254
      I#I
255
      I#X
256
      X#X
257
      C$C
258
      C$N
259
      C$0
260
      C$S
261
      C$C1
262
      C$P
263
      C$F
```

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C\$Br

```
265
                 C$Si
           266
                 CŚI
           267
                 C$X
           268
                 NSN
           269
                 N$O
           270
                 NSS
           271
                 N$Cl
           272
                 N$P
           273
                 N$F
           274
                 N$Br
           275
                 NSSi
           276
                 N$I
           277
                 N$X
           278
                 0$0
           279
                 O$S
           280
                 O$Cl
           281
                 O$P
           282
                 O$F
           283
                 O$Br
           284
                 O$Si
           285
                 O$I
           286
                 O$X
           287
                 S$S
           288
                 S$Cl
           289
                 S$P
           290
                 S$F
           291
                 S$Br
           292
                 S$Si
           293
                 SŠI
           294
                 S$X
           295
                 Cl$Cl
           296
                 Cl$P
           297
                 Cl$F
           298
                 Cl$Br
           299
                 Cl$Si
           300
                 Cl$I
           301
                 Cl$X
           302
                 P$P
           303
                 P$F
           304
                 P$Br
           305
                 P$Si
           306
                 P$I
           307
                 Р$X
           308
                 F$F
           309
                 F$Br
           310
                 F$Si
           311
                 F$I
           312
                 F$X
           313
                 Br$Br
           314
                 Br$Si
           315
                 Br$I
           316
                 Br$X
           317
                 Si$Si
           318
                 Si$I
           319
                 Si$X
           320
                 I$I
           321
                 I$X
           322
                 X$X
           $MACCSKeys->SetSize($Size);
      Sets size of MACCS keys and returns MACCSKeys. Possible values: 166 or 322.
           $MACCSKeys->SetType($Type);
      Sets type of MACCS keys and returns MACCSKeys. Possible values: MACCSKeysBits or MACCSKeysCount.
StringifyMACCSKeys
           $String = $MACCSKeys->StringifyMACCSKeys();
```

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Returns a string containing information about MACCSKeys object.

SetSize

SetType

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SEE ALSO

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

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