# NAME

MACCSKeysFingerprints.pl - Generate MACCS key fingerprints for SD files

### **SYNOPSIS**

MACCSKeysFingerprints.pl SDFile(s)...

MACCSKeysFingerprints.pl [--AromaticityModel AromaticityModelType] [--BitsOrder Ascending | Descending | [--b, --BitStringFormat BinaryString | HexadecimalString] [--CompoundI D DataFieldName or LabelPrefixString] [--CompoundI DLabel text] [--CompoundI DMode DataField | MolName | LabelPrefix | MolNameOrLabelPrefix] [--DataFields "FieldLabel1,FieldLabel2,..."] [-d, --DataFieldsMode All | Common | Specify | CompoundID] [-f, --Filter Yes | No] [--FingerprintsLabel text] [-h, --help] [-k, --KeepLargestComponent Yes | No] [-m, --mode MACCSKeyBits | MACCSKeyCount] [--OutDelim comma | tab | semicolon] [--output SD | FP | text | all] [-o, --overwrite] [-q, --quote Yes | No] [-r, --root RootName] [-s, --size number] [-v, --VectorStringFormat | IDsAndValuesString | IDsAndValuesPairsString | ValuesAndIDsString | ValuesAndIDsPairsString] [-w, --WorkingDir DirName]

## DESCRIPTION

Generate MACCS (Molecular ACCess System) keys fingerprints [ Ref 45-47 ] for *SDFile(s)* and create appropriate SD, FP or CSV/TSV text file(s) containing fingerprints bit-vector or vector strings corresponding to molecular fingerprints.

Multiple SDFile names are separated by spaces. The valid file extensions are .sdf and .sd. All other file names are ignored. All the SD files in a current directory can be specified either by \*.sdf or the current directory name.

For each MACCS 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 -m, --mode 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 | MACCSKeyCount values for -m, --mode option along with two possible 166 | 322 values of -s, --size supports generation of four different types of MACCS keys fingerprint: MACCS166KeyBits, MACCS166KeyCount, MACCS322KeyBits, MACCS322KeyCount.

Example of SD file containing MAACS keys fingerprints string data:

```
. . . . . .
$$$$
. . . . . .
41 44 0 0 0 0 0 0 0 0 0999 V2000
-3.3652 1.4499 0.0000 C 0 0 0 0 0 0 0 0 0 0
2 3 1 0 0 0 0
. . . . . .
M END
> <CmpdID>
Cmpd1
> <MACCSKeysFingerprints>
FingerprintsBitVector; MACCSKeyBits; 166; BinaryString; Ascending; 000000000
11111111110111000
$$$$
. . . . . .
```

Example of *FP* file containing MAACS keys fingerprints string data:

```
#
# Package = MayaChemTools 7.4
# Release Date = Oct 21, 2010
#
```

Example of CSV Text file containing MAACS keys fingerprints string data:

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

FingerprintsBitVector;MACCSKeyBits;166;HexadecimalString;Ascending;000 000000021210210e845f8d8c60b79dffbffffd1

## **OPTIONS**

--AromaticityModel MDLAromaticityModel | TriposAromaticityModel | MMFFAromaticityModel | ChemAxonBasicAromaticityModel | ChemAxonGeneralAromaticityModel | DaylightAromaticityModel | MayaChemToolsAromaticityModel

Specify aromaticity model to use during detection of aromaticity. Possible values in the current release

are: MDLAromaticityModel, TriposAromaticityModel, MMFFAromaticityModel, ChemAxonBasicAromaticityModel, ChemAxonGeneralAromaticityModel, DaylightAromaticityModel or MayaChemToolsAromaticityModel. Default value: MayaChemToolsAromaticityModel.

The supported aromaticity model names along with model specific control parameters are defined in AromaticityModelsData.csv, which is distributed with the current release and is available under lib/data directory. Molecule.pm module retrieves data from this file during class instantiation and makes it available to method DetectAromaticity for detecting aromaticity corresponding to a specific model.

### --BitsOrder Ascending | Descending

Bits order to use during generation of fingerprints bit-vector string for *MACCSKeyBits* value of -m, --mode option. Possible values: *Ascending, Descending.* Default: *Ascending.* 

Ascending bit order which corresponds to first bit in each byte as the lowest bit as opposed to the highest bit.

Internally, bits are stored in *Ascending* order using Perl vec function. Regardless of machine order, big-endian or little-endian, vec function always considers first string byte as the lowest byte and first bit within each byte as the lowest bit.

# -b, --BitStringFormat BinaryString | HexadecimalString

Format of fingerprints bit-vector string data in output SD, FP or CSV/TSV text file(s) specified by --output used during *MACCSKeyBits* value of -m, --mode option. Possible values: *BinaryString*, *HexadecimalString*. Default value: *BinaryString*.

BinaryString corresponds to an ASCII string containing 1s and 0s. HexadecimalString contains bit values in ASCII hexadecimal format.

### Examples:

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

# --CompoundI D DataFieldName or LabelPrefixString

This value is --CompoundIDMode specific and indicates how compound ID is generated.

For *DataField* value of --CompoundIDMode option, it corresponds to datafield label name whose value is used as compound ID; otherwise, it's a prefix string used for generating compound IDs like LabelPrefixString<Number>. Default value, *Cmpd*, generates compound IDs which look like Cmpd<Number>.

Examples for DataField value of --CompoundI DMode:

MolID ExtReg

Examples for LabelPrefix or MolNameOrLabelPrefix value of --CompoundI DMode:

Compound

The value specified above generates compound IDs which correspond to Compound<Number> instead of default value of Cmpd<Number>.

# --CompoundI DLabel text

Specify compound ID column label for FP or CSV/TSV text file(s) used during *CompoundID* value of --DataFieldsMode option. Default: *CompoundID*.

#### --CompoundI DMode DataField | MolName | LabelPrefix | MolNameOrLabelPrefix

Specify how to generate compound IDs and write to FP or CSV/TSV text file(s) along with generated fingerprints for FP | text | all values of --output option: use a SDFile(s) datafield value; use molname line from SDFile(s); generate a sequential ID with specific prefix; use combination of both MolName and LabelPrefix with usage of LabelPrefix values for empty molname lines.

Possible values: DataField | MolName | LabelPrefix | MolNameOrLabelPrefix. Default: LabelPrefix.

For *MolNameAndLabelPrefix* value of --CompoundIDMode, molname line in *SDFile(s)* takes precedence over sequential compound IDs generated using *LabelPrefix* and only empty molname values are replaced with sequential compound IDs.

This is only used for CompoundID value of -- DataFieldsMode option.

### -- DataFields "FieldLabel1, FieldLabel2,..."

Comma delimited list of *SDFiles(s)* data fields to extract and write to CSV/TSV text file(s) along with generated fingerprints for *text | all* values of --output option.

This is only used for *Specify* value of --DataFieldsMode option.

Examples:

Extreg

MolID, CompoundName

# -d, --DataFieldsMode All | Common | Specify | CompoundID

Specify how data fields in *SDFile(s)* are transferred to output CSV/TSV text file(s) along with generated fingerprints for *text | all* values of --output option: transfer all SD data field; transfer SD data files common to all compounds; extract specified data fields; generate a compound ID using molname line, a compound prefix, or a combination of both. Possible values: *All | Common | specify | CompoundID*. Default value: *CompoundID*.

# -f, --Filter Yes | No

Specify whether to check and filter compound data in SDFile(s). Possible values: Yes or No. Default value: Yes.

By default, compound data is checked before calculating fingerprints and compounds containing atom data corresponding to non-element symbols or no atom data are ignored.

# --FingerprintsLabel text

SD data label or text file column label to use for fingerprints string in output SD or CSV/TSV text file(s) specified by --output. Default value: MACCSKeyFingerprints.

## -h, --help

Print this help message.

# -k, --KeepLargestComponent Yes | No

Generate fingerprints for only the largest component in molecule. Possible values: Yes or No. Default value: Yes

For molecules containing multiple connected components, fingerprints can be generated in two different ways: use all connected components or just the largest connected component. By default, all atoms except for the largest connected component are deleted before generation of fingerprints.

# -m, --mode MACCSKeyBits | MACCSKeyCount

Specify type of MACCS keys [ Ref 45-47 ] fingerprints to generate for molecules in *SDFile(s)*. Possible values: *MACCSKeyBits*, *MACCSKeyCount*. Default value: *MACCSKeyBits*.

For *MACCSKeyBits* value of -m, --mode 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 is generated.

MACCSKeyBits | MACCSKeyCount values for -m, --mode option along with two possible 166 | 322 values of -s, --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
   Z : 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
   T : Triple
   # : Triple
~ : Single or double query 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 QAA@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 OQ(O)O
- 49 CHARGE
- 50 C=C(C)C 51 CSO
- 52 NN
- 53 QНАААОН
- 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 QN
- 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 0 > 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:

- . 26 atom properties of type P, as listed in Table 1
- . 32 one-atom environments, as listed in Table 3
- . 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
1
2
      Q - heteroatom
      Anot%not-A - atom involved in one or more multiple bonds, not aromatic
3
      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
6
      A(QQQ) or QA(Q)Q - atom with at least three heteroatom neighbors
      QH - heteroatom with at least one hydrogen attached
8
      CH2(AA) or ACH2A - carbon with at least two single bonds and at least
      two hydrogens attached
9
      CH3(A) or ACH3 - carbon with at least one single bond and at least three
      hydrogens attached
10
      Halogen
      A(-A-A-A) or A-A(-A)-A - atom has at least three single bonds
11
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
      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
```

Atomic environments properties-based keys (32):

more neighbors (including the central atom).

CHACH2 - atom has two hydrocarbon (CH2) neighbors

```
Key
      Description
27
      C(CC)
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)
```

26

```
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
      O(CO)
51
      Q(CCO)
52
      Q(NO)
53
      Q(CNO)
54
      Q(NNO)
55
      Q(00)
56
      Q(C00)
      Q(NOO)
57
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
82
      O-Cl
83
      O-P
84
      O-F
85
      O-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
      S-I
96
      S-X
97
      Cl-Cl
98
      Cl-P
99
      Cl-F
100
      Cl-Br
101
      Cl-Si
102
      Cl-I
103
      Cl-X
```

104 P-P 105 P-F 106 P-Br P-Si 107 108 P-I 109 P-X 110 F-F111 F-Br 112 F-Si 113 F-I114 F-X115 Br-Br Br-Si 116 117 Br-I 118 Br-X 119 Si-Si 120 Si-I 121 Si-X 122 I-I 123 I-X124 X-X125 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=N137 N=O 138 N=S 139 N=Cl 140 N=P 141 N=F142 N=Br 143 N=Si 144 N=I145 N=X 146 0=0 147 0=S 148 O=Cl 149 O=P 150 O=F 151 O=Br 152 O=Si 153 O=I 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=I169 Cl=X170 P=P P=F

171 172

P=Br

173 P=Si 174 P=I 175 P=X 176 F=F 177 F=Br 178 F=Si 179 F=I180 F=X 181 Br=Br 182 Br=Si 183 Br=I 184 Br=X 185 Si=Si 186 Si=I 187 Si=X 188 I = I189 I=X190 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 0#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#X257 C\$C 258 C\$N 259 C\$0 260 C\$S 261 C\$Cl 262 C\$P 263 C\$F 264 C\$Br 265 C\$Si 266 C\$I 267 C\$X 268 N\$N 269 N\$O 270 N\$S 271 N\$Cl 272 N\$P 273 N\$F 274 N\$Br 275 N\$Si 276 N\$I 277 N\$X 278 0\$0 279 0\$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 S\$I 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 P\$X 308 F\$F 309 F\$Br

310

F\$Si

311 FŠI 312 F\$X 313 Br\$Br 314 Br\$Si 315 Br\$I 316 Br\$X 317 Si\$Si 318 SiŠI 319 Si\$X 320 I\$I 321 Ι\$X 322 X\$X

### --OutDelim comma | tab | semicolon

Delimiter for output CSV/TSV text file(s). Possible values: comma, tab, or semicolon Default value: comma.

# --output SD | FP | text | all

Type of output files to generate. Possible values: SD, FP, text, or all. Default value: text.

### -o, --overwrite

Overwrite existing files.

# -q, --quote Yes | No

Put quote around column values in output CSV/TSV text file(s). Possible values: Yes or No. Default value: Yes.

#### -r, --root RootName

New file name is generated using the root: <Root>.<Ext>. Default for new file names: <SDFileName><MACCSKeysFP>.<Ext>. The file type determines <Ext> value. The sdf, fpf, csv, and tsv <Ext> values are used for SD, FP, comma/semicolon, and tab delimited text files, respectively. This option is ignored for multiple input files.

## -s, --size number

Size of MACCS keys [ Ref 45-47 ] set to use during fingerprints generation. Possible values: *166 or 322*. Default value: *166*.

# -v, --VectorStringFormat ValuesString | IDsAndValuesString | IDsAndValuesPairsString | ValuesAndIDsString | ValuesAndIDsPairsString

Format of fingerprints vector string data in output SD, FP or CSV/TSV text file(s) specified by --output used during MACCSKeyCount value of -m, --mode option. Possible values: ValuesString, IDsAndValuesString | IDsAndValuesPairsString | ValuesAndIDsString | ValuesAndIDsPairsString. Defaultvalue: ValuesString.

# Examples:

# -w, --WorkingDir DirName

Location of working directory. Default: current directory.

# **EXAMPLES**

To generate MACCS keys fingerprints of size 166 in binary bit-vector string format and create a SampleMACCS166FPBin.csv file containing sequential compound IDs along with fingerprints bit-vector strings data, type:

% MACCSKeysFingerprints.pl -r SampleMACCS166FPBin -o Sample.sdf

To generate MACCS keys fingerprints of size 166 in binary bit-vector string format and create SampleMACCS166FPBin.csv and SampleMACCS166FPBin.csv files containing sequential compound IDs in CSV file along with fingerprints bit-vector strings data, type:

```
% MACCSKeysFingerprints.pl --output all -r SampleMACCS166FPBin
-o Sample.sdf
```

To generate MACCS keys fingerprints of size 322 in binary bit-vector string format and create a SampleMACCS322FPBin.csv file containing sequential compound IDs along with fingerprints bit-vector strings data, type:

```
% MACCSKeysFingerprints.pl -size 322 -r SampleMACCS322FPBin -o Sample.sdf
```

To generate MACCS keys fingerprints of size 166 corresponding to count of keys in ValuesString format and create a SampleMACCS166FPCount.csv file containing sequential compound IDs along with fingerprints vector strings data, type:

```
% MACCSKeysFingerprints.pl -m MACCSKeyCount -r SampleMACCS166FPCount
-o Sample.sdf
```

To generate MACCS keys fingerprints of size 322 corresponding to count of keys in ValuesString format and create a SampleMACCS322FPCount.csv file containing sequential compound IDs along with fingerprints vector strings data, type:

```
% MACCSKeysFingerprints.pl -m MACCSKeyCount -size 322
-r SampleMACCS322FPCount -o Sample.sdf
```

To generate MACCS keys fingerprints of size 166 in hexadecimal bit-vector string format with ascending bits order and create a SampleMACCS166FPHex.csv file containing compound IDs from MolName along with fingerprints bit-vector strings data, type:

```
% MACCSKeysFingerprints.pl -m MACCSKeyBits --size 166 --BitStringFormat
HexadecimalString --BitsOrder Ascending --DataFieldsMode CompoundID
--CompoundIDMode MolName -r SampleMACCS166FPBin -o Sample.sdf
```

To generate MACCS keys fingerprints of size 166 corresponding to count of keys in IDsAndValuesString format and create a SampleMACCS166FPCount.csv file containing compound IDs from MolName line along with fingerprints vector strings data, type:

```
% MACCSKeysFingerprints.pl -m MACCSKeyCount --size 166
--VectorStringFormat IDsAndValuesString --DataFieldsMode CompoundID
--CompoundIDMode MolName -r SampleMACCS166FPCount -o Sample.sdf
```

To generate MACCS keys fingerprints of size 166 corresponding to count of keys in IDsAndValuesString format and create a SampleMACCS166FPCount.csv file containing compound IDs using specified data field along with fingerprints vector strings data, type:

```
% MACCSKeysFingerprints.pl -m MACCSKeyCount --size 166
  --VectorStringFormat IDsAndValuesString --DataFieldsMode CompoundID
  --CompoundIDMode DataField --CompoundID Mol_ID -r
  SampleMACCS166FPCount -o Sample.sdf
```

To generate MACCS keys fingerprints of size 322 corresponding to count of keys in ValuesString format and create a SampleMACCS322FPCount.tsv file containing compound IDs derived from combination of molecule name line and an explicit compound prefix along with fingerprints vector strings data in a column labels MACCSKeyCountFP, type:

```
% MACCSKeysFingerprints.pl -m MACCSKeyCount -size 322 --DataFieldsMode
CompoundID --CompoundIDMode MolnameOrLabelPrefix --CompoundID Cmpd
--CompoundIDLabel MolID --FingerprintsLabel MACCSKeyCountFP --OutDelim
Tab -r SampleMACCS322FPCount -o Sample.sdf
```

To generate MACCS keys fingerprints of size 166 corresponding to count of keys in ValuesString format and create a SampleMACCS166FPCount.csv file containing specific data fields columns along with fingerprints vector strings data, type:

```
% MACCSKeysFingerprints.pl -m MACCSKeyCount --size 166
--VectorStringFormat ValuesString --DataFieldsMode Specify --DataFields
Mol_ID -r SampleMACCS166FPCount -o Sample.sdf
```

To generate MACCS keys fingerprints of size 322 corresponding to count of keys in ValuesString format and create a SampleMACCS322FPCount.csv file containing common data fields columns along with fingerprints vector strings data, type:

```
% MACCSKeysFingerprints.pl -m MACCSKeyCount --size 322
  --VectorStringFormat ValuesString --DataFieldsMode Common -r
  SampleMACCS322FPCount -o Sample.sdf
```

To generate MACCS keys fingerprints of size 166 corresponding to count of keys in ValuesString format and create SampleMACCS166FPCount.sdf, SampleMACCS166FPCount.fpf and SampleMACCS166FPCount.csv files containing all data fields columns in CSV file along with fingerprints vector strings data, type:

```
% MACCSKeysFingerprints.pl -m MACCSKeyCount --size 166 --output all
  --VectorStringFormat ValuesString --DataFieldsMode All -r
  SampleMACCS166FPCount -o Sample.sdf
```

### **AUTHOR**

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# SEE ALSO

InfoFingerprintsFiles.pl, SimilarityMatricesFingerprints.pl, AtomNeighborhoodsFingerprints.pl, ExtendedConnectivityFingerprints.pl, PathLengthFingerprints.pl, TopologicalAtomPairsFingerprints.pl, TopologicalAtomTorsionsFingerprints.pl, TopologicalPharmacophoreAtomPairsFingerprints.pl, TopologicalPharmacophoreAtomTripletsFingerprints.pl

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