Representation of molecular structures

Coutesy of Prof. João Aires-de-Sousa, University of Lisbon, Portugal
### A hierarchy of structure representations

<table>
<thead>
<tr>
<th>Name</th>
<th>(S)-Tryptophan</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>2D Structure</strong></td>
<td><img src="image1" alt="2D Structure" /></td>
</tr>
<tr>
<td><strong>3D Structure</strong></td>
<td><img src="image2" alt="3D Structure" /></td>
</tr>
<tr>
<td><strong>Molecular surface</strong></td>
<td><img src="image3" alt="Molecular surface" /></td>
</tr>
</tbody>
</table>
Storing molecular structures in a computer
Information must be **coded** into interconvertible formats that can be read by software applications.

Applications: visualization, communication, database searching / management, establishment of structure-property relationships, estimation of properties, …
Coding molecular structures

• A non-ambiguous representation identifies a single possible structure, e.g. the name ‘o-xylene’ represents one and only one possible structure.

• A representation is unique if any structure has only one possible representation (some nomenclature isn’t, e.g. ‘1,2-dimethylbenzene’ and ‘o-xylene’ represent the same structure).
IUPAC Nomenclature

- **Advantages:**
  - standardized systematic classification
  - stereochemistry is included
  - widespread
  - unambiguous
  - allows reconstruction from the name

- **Disadvantages:**
  - extensive rules
  - alternative names are allowed (non-unique)
  - long complicated names

IUPAC name: N-[(2R,4R,5S)-5-[[[(2S,4R,5S)-3-acetamido-5-][(2S,4S,5S)-3-acetamido-4,5-dihydroxy-6-(hydroxymethyl)oxan-2-yl]methoxymethyl]-4-hydroxy-6-(hydroxymethyl)oxan-2-yl]methoxymethyl]-2,4-dihydroxy-6-(hydroxymethyl)oxan-3-yl]acetamide
Linear notations

Represent structures by linear sequences of letters and numbers, e.g. IUPAC nomenclature.

Linear notations can be extremely compact, which is an advantage for the storage of structures in a computer (particularly when disk space is limited).

Linear notations allow for an easy transmission of structures, e.g. in a Google-type search, or in an email.
The SMILES notation

1. Atoms are represented by their atomic symbols.
2. Hydrogen atoms are omitted (are implicit).
3. Neighboring atoms are represented next to each other.
4. Double bonds are represented by ‘=’, triple bonds by ‘#’.
5. Branches are represented by parentheses.
6. Rings are represented by allocating digits to the two connecting ring atoms.

Example: 

SMILES representation: CCCO

Example: 

SMILES: CCC(Cl)C=C
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SMILES: C1CCCCC1
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6. Rings are represented by allocating digits to the two connecting ring atoms.
7. Aromatic rings are indicated by lower-case letters.

**SMILES:** \textbf{Nc1ccccc1}
The SMILES notation

• Is unambiguous (a SMILES string unequivocally represents a single structure).

• Is it unique??

Solution: algorithm that guarantees a canonical representation (each structure is always represented by the same SMILES string)

• More at: http://www.daylight.com/dayhtml_tutorials/index.html
SMILES notation in MarvinSketch
SMILES notation in MarvinSketch

The input in MarvinSketch is: C>Cclcc[nH]c1

The SMILES output is: CCclcc[nH]c1
The InChI notation
(IUPAC International Chemical Identifier)

Example:

InChI=1/C6H8O6/c7-1-2(8)5-3(9)4(10)6(11)12-5/h2,5,7-10H,1H2/t2-,5+/m0/s1

L-ascorbic acid

A digital equivalent to the IUPAC name for a compound.

Five layers of information: connectivity, tautomerism, isotopes, stereochemistry, and charge.

An algorithm generates an unambiguous unique notation.

Official web site: http://www.iupac.org/inchi/
The InChI notation
(IUPAC International Chemical Identifier)

Example:

InChI=1/C6H8O6/c7-1-2 (8) 5-3 (9) 4(10) 6(11) 12-5/h2,5,7-10H,1H2/t2-,5+/m0/s1

L-ascorbic acid

Each layer in an InChI string contains a specific class of structural information. This format is designed for compactness, not readability, but can be interpreted manually.

The length of an identifier is roughly proportional to the number of atoms in the substance. Numbers inside a layer usually represent the canonical numbering of the atoms from the first layer (chemical formula) except H.
A molecular structure can be interpreted as a mathematical graph where each atom is a node, and each bond is an edge.

Such a representation allows for the mathematical processing of molecular structures using the graph theory.
Topological Graph Theory

branch of mathematics particularly useful in chemical informatics and in computer science generally

study of “graphs” which consist of

a set of “nodes”
a set of “edges” joining pairs of nodes
Properties of graphs

graphs are only about connectivity
spatial position of nodes is irrelevant
length of edges are irrelevant
crossing edges are irrelevant
Properties of Graphs

nodes and edges can be “coloured” to distinguish them
Structure Diagrams as Graphs

2D structure diagrams very like topological graphs

- atoms ↔ nodes
- bonds ↔ edges

Terminal hydrogen atoms are not normally shown as separate nodes ("implicit" hydrogens)
- reduces number of nodes by ~50%
- "hydrogen count" information used to colour neighbouring "heavy atom" atom

Separate nodes sometimes used for "special" hydrogens

- deuterium, tritium
- hydrogen bonded to more than one other atom
- hydrogens attached to stereocentres
Advantages of using graphs

- mathematical theory is well understood
- graphs can be easily represented in computers
  - many useful algorithms are known
- identical graphs ⇔ identical molecules
- different graphs ⇔ different molecules
Matrix representations

A molecular structure with \( n \) atoms may be represented by an \( n \times n \) matrix (H-atoms are often omitted).

**Adjacency matrix** : indicates which atoms are bonded.

```
1 0 1 0 0 0 0
2 1 0 1 0 0 0
3 0 1 0 1 1 0
4 0 0 1 0 0 0
5 0 0 1 0 0 1
6 0 0 0 0 1 0
```
Matrix representations

A molecular structure with $n$ atoms may be represented by an $n \times n$ matrix (H-atoms are often omitted).

**Adjacency matrix**: indicates which atoms are bonded.

```
1 1
2 1 1
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6
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```
Matrix representations

A molecular structure with \( n \) atoms may be represented by an \( n \times n \) matrix (H-atoms are often omitted).

**Adjacency matrix** : indicates which atoms are bonded.
Matrix representations

Distance matrix: encodes the distances between atoms.

The distance is defined as the number of bonds between atoms on the shortest possible path.

Distance may also be defined as the 3D distance between atoms.
Matrix representations

**Bond matrix**: indicates which atoms are bonded, and the corresponding bond orders.
A disadvantage of matrix representations is that the matrix size increases with the square of the number of atoms.

A **connection table** lists the atoms of a molecule, and the bonds between them (may include or not H-atoms).
The MDL Molfile format


Nr of atoms

Nr of bonds

Description of an atom

Description of a bond

H₃C

Cl

CH₂
### The atom block

**L-Alanine**

<table>
<thead>
<tr>
<th>Field</th>
<th>Meaning</th>
<th>Values</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>x y z</td>
<td>atom coordinates</td>
<td></td>
<td>(Generic)</td>
</tr>
<tr>
<td>aaa</td>
<td>atom symbol</td>
<td></td>
<td>(Generic, Query, 3D, Rgroup)</td>
</tr>
<tr>
<td>dd</td>
<td>mass difference</td>
<td>-3, -2, -1, 0, 1, 2, 3, 4 (0 if value beyond these limits)</td>
<td>(Generic) Difference from mass in periodic table. Wider range of values allowed by M ISO line, below. Retained for compatibility with older Ctabs. M ISO takes precedence.</td>
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<tr>
<td>ccc</td>
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![Diagram of L-Alanine]
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<tr>
<td>aaa</td>
<td>atom symbol</td>
<td>entry in periodic table or L</td>
<td>(Generic, Query, 3D, Rgroup)</td>
</tr>
<tr>
<td></td>
<td>for atom list, A, Q, * for unspecified atom, and LP for one pair, or R# for Rgroup label</td>
<td></td>
<td></td>
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## The atom block

- L-Alanine

### Table: Atom Block

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<td>x y z</td>
<td>atom coordinates</td>
<td>entry in periodic table or L for atom list, A, Q, * for unspecified atom, and LP for lone pair, or R# for Rgroup label</td>
<td>(Generic)</td>
</tr>
<tr>
<td>aaa</td>
<td>atom symbol</td>
<td></td>
<td>[Generic, Query 3D, Rgroup]</td>
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### Diagram: L-Alanine Structure

- **Chiral Center**: A chiral center is indicated by a flexible line with two lines, each of which is numbered with a circle. The flexibility indicates that the molecule can rotate about the bond, giving rise to different stereochemistries.

- **Bonding**: The bonds are represented by straight lines connecting the atoms, with arrows indicating the direction of the bond. The atoms are labeled with their respective symbols: 
  - C (Carbon) 
  - N (Nitrogen) 
  - O (Oxygen)

- **Coordination**: The structure shows the coordination around the chiral center, indicating the orientation of the bonds relative to each other.

The structure is a representation of L-Alanine, highlighting the chiral center and the bond orientation.
# The atom block

L-Alanine

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<tr>
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<td>may be any atom symbol</td>
<td><strong>[Generic, Query, 3D, Rgroup]</strong></td>
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## The atom block

-0.6622  0.5342  0.0000  C  0  0  2  0  0  0  L-Alanine
0.6622  -0.3000  0.0000  C  0  0  0  0  0  0
-1.8622  -0.3695  0.0000  C  0  3  0  0  0  0
1.9464  0.4244  0.0000  O  0  0  0  0  0  0

### Field Meaning Values Notes

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### Diagram

![3D Structure of L-Alanine](image)
The bond block

L-Alanine

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<tr>
<td>111</td>
<td>first atom number</td>
<td>1 - number of atoms</td>
<td>[Generic]</td>
</tr>
<tr>
<td>222</td>
<td>second atom number</td>
<td>1 - number of atoms</td>
<td>[Generic]</td>
</tr>
<tr>
<td>tt</td>
<td>bond type</td>
<td>1 = Single, 2 = Double, 3 = Triple, 4 = Aromatic, 5 = Single or Double, 6 = Single or Aromatic, 7 = Double or Aromatic, 8 = Any</td>
<td>[Query] Values 4 through 8 are for SSS queries only.</td>
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<td>[Generic] The wedge (pointed end of the stereo bond is at the first atom (Field 111 above)</td>
</tr>
<tr>
<td>xxx</td>
<td>not used</td>
<td></td>
<td></td>
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<tr>
<td>rrr</td>
<td>bond topology</td>
<td>0 = Either, 1 = Ring, 2 = Chain</td>
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<td>ccc</td>
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### The bond block

#### Diagram of L-Alanine

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![L-Alanine structure](image)
### The bond block

![Chemical structure of L-Alanine](image)

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The MDL Molfile format

L-Alanine (13C)
GSMACCS-II10169115362D 1 0.00366 0.00000 0

6 5 0 0 1 0 3 V2000
-0.6622 0.5342 0.0000 C 0 0 2 0 0 0
0.6622 -0.3000 0.0000 C 0 0 0 0 0 0
-0.7207 2.0817 0.0000 C 1 0 0 0 0 0
-1.8622 -0.3695 0.0000 N 0 3 0 0 0 0
0.6220 -1.8037 0.0000 O 0 0 0 0 0 0
1.9464 0.4244 0.0000 O 0 5 0 0 0 0

1 2 1 0 0 0
1 3 1 1 0 0
1 4 1 0 0 0
2 5 2 0 0 0
2 6 1 0 0 0

M CHG 2 4 1 6 -1
M ISO 1 3 13
M END
The properties block

2 charged atoms
The properties block

2 charged atoms

atom 4: charge +1
atom 6: charge -1
The properties block

M  CHG  2  4  1  6  -1
M  ISO  1  3  13
M  END

1 entry for an isotope
The properties block

1 entry for an isotope

atom 3: mass=13
The SDFile (.SDF) format

Includes structural information in the Molfile format and associated data items for one or more compounds.

Molfile1
Associated data
$$$$
Molfile2
Associated data
$$$$
...

The SDF File (.SDF) format

Example

Molfile1
Associated data

Molfile2
Associated data

Example

Associated data (molecular)
### The SDFFile (.SDF) format

**Example**

```
61203-01-8
Marvin 02130718303D

11 10 0 0 0 0 0 0 0 0 999 02800
 1.6947 -0.2675 -0.0016 C 0 0 0 0 0 0 0 0 0 0 0 0
 0.5843  0.6242  0.0001 C 0 0 0 0 0 0 0 0 0 0 0 0
 3.0673  0.3883 -0.0014 C 0 0 0 0 0 0 0 0 0 0 0 0
 1.5555 -1.4702  0.0019 O 0 0 0 0 0 0 0 0 0 0 0 0
 0.7695  1.9478  0.0018 C 0 0 0 0 0 0 0 0 0 0 0 0
-1.2132 -0.0983 -0.0002 Br 0 0 0 0 0 0 0 0 0 0 0 0
 3.5067 -0.7060  0.0000 H 0 0 0 0 0 0 0 0 0 0 0 0
 4.1016  0.7277  0.0000 H 0 0 0 0 0 0 0 0 0 0 0 0
 2.6679  1.3226  0.0000 H 0 0 0 0 0 0 0 0 0 0 0 0
 1.7238  2.3671  0.0000 H 0 0 0 0 0 0 0 0 0 0 0 0
-0.1088  2.6166  0.0000 H 0 0 0 0 0 0 0 0 0 0 0 0

1 2 1 0 0 0 0
1 3 1 0 0 0 0
1 4 2 0 0 0 0
2 5 2 0 0 0 0
2 6 1 0 0 0 0
3 7 1 0 0 0 0
3 8 1 0 0 0 0
3 9 1 0 0 0 0
5 10 1 0 0 0 0
5 11 1 0 0 0 0
```

Associated data (atomic)

```
> (AMES test categorisation)
mutagen

> (CHARGE)
0.11; 0.05; -0.03; -0.29; -0.02; -0.05; 0.63; 0.03; 0.03; 0.06; 0.06

> (EXACTMASS)
147.952377

$$$
598-55-6
Marvin 02130718303D

18 9 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 999 02800
```

Molfile1
Associated data $$$
Molfile2
Associated data $$$
...
```
The SDFFile (.SDF) format

Example

Molfile1
Associated data

Molfile2
Associated data

Associated data (molecular)
The SDFFile (.SDF) format

Molfile1
Associated data
$$$$
Molfile2
Associated data
$$$$
...

Example

Beginning of Molfile2

Delimiter

Delimiter
Beginning of Molfile2
The SDFFile (.SDF) format

Example

Molfile1
Associated data

Molfile2
Associated data

...
The ChemAxon Standardize program

- Conversion of file formats
- Generation of unique SMILES strings
- Standardization of structures
- Addition of H-atoms, removal of H-atoms, assignment of aromatic systems, cleaning of stereochemistry, …
The ChemAxon Standardize program

Aromatize

Replaces the alternating single/double bonds of aromatic rings with aromatic bond types. The aromatic rings of the current compound are determined by the Hückel's rule. Two methods are provided, the GENERAL approach operates on the SSSR (smallest set of smallest rings) and considers some mesomers as well (for example, pyridone is aromatic). The BASIC method detects all aromatic rings (not just the SSSR) and operates on the current structure (pyridone is non aromatic).
A Markush structures diagram is a type of representation specific for a SERIES of chemical compounds.

The diagram can describe not only a specific molecule, but several families of compounds.

It includes a core and substituents, which are listed as text separately from the diagram.

These are mostly used in databases of patents.
Representation of molecular fragments

Just like a text document may be indexed on the basis of specified keywords, a chemical structure may be indexed on the basis of specific chemical characteristics, usually fragments.

Fragments may be, e.g., small groups of atoms, functional groups, rings. These are defined beforehand.

It is an ambiguous representation: different structures may have common fragments.

Fragments:

- -OH
- -COOH
- >C=O
- -NH2
- -3-indole
Fingerprints

Fingerprints encode the presence or absence of certain features in a compound, e.g., fragments.

If 20 fragments are defined, the fingerprint has a length of 20.
It is an ambiguous representation.
Allows for similarity searches.
‘Hashed Fingerprints’

Encode the presence of sub-structures. **These are not previously defined.**

All patterns are listed consisting of
- 1 atom
- 2 bonded atoms and their bond
- Sequences of 3 atoms and their bonds
- Sequences of 4 atoms and their bonds
- …

Patterns up to 3 atoms
- C, N, O
- C-C, C-N, C=O, C-O
- C-C-C, C-C-N, C-C=O, C-C-O, O=C-O
‘Hashed Fingerprints’

Each pattern activates a certain number of positions (bits) in the fingerprint, in the following example two bits / pattern:

An algorithm determines which bits are activated by a pattern. The same pattern always activates the same bits. The algorithm is designed in such a way that it is always possible to assign bits to a pattern.

There may be collisions. Pre-definition of fragments is not required. But it is not possible to interpret fingerprints.
‘Hashed Fingerprints’

Parameters to define: fingerprint length, size of patterns, and number of bits activated by each pattern.

Main application: similarity search in large databases.

H-atoms are omitted. Stereochemistry is not considered.
‘Hashed Fingerprints’
Influence of parameters

Length of fingerprint:
• too short ⇒ almost all bits=1, poor discrimination of molecules.
• too large ⇒ too many bits=0, too much disk space required.

Maximum size of patterns:
• too short ⇒ poor discrimination of molecules.
• too large ⇒ ability to discriminate molecules, but many bits=1.

Nr of bits a pattern activates:
• too few ⇒ poor ability to discriminate between patterns.
• too many ⇒ ability to discriminate between patterns, but many bits=1.

‘Hashed Fingerprints’

or Daylight fingerprints

Can be calculated with several software packages, e.g. the `generfp` command of the program JCHEM (Chemaxon).

Length (in bytes)

Maximum size of patterns

Nr of bits activated by a pattern

Input file

Output file

`C:\Documents and Settings>cd ...

C:\>generfp -fl 16 -pl 5 -hc 5 <abcd.smi> abcd.fp`
‘Hashed Fingerprints’
or Daylight fingerprints

Can be calculated with the `generfp` command of the program JCHEM (Chemaxon).

![Command Prompt showing `generfp` command output]

![Programmer's File Editor showing chemical structures and fingerprint output]
**Similarity measures based on fingerprints**

Similarity between compounds $X$ and $Y$ can be calculated from the similarity between their fingerprints.

- $a =$ nr of bits ‘on’ in $X$ but not in $Y$.
- $b =$ nr of bits ‘on’ in $Y$ but not in $X$.
- $c =$ nr of bits ‘on’ both in $X$ and in $Y$.
- $d =$ nr of bits ‘off’ both in $X$ and in $Y$.

$n = (a + b + c + d)$ is the total number of bits

**Euclidean coefficient :**

\[
\frac{c + d}{n} \quad \text{(common bits in $X$ and $Y$)}
\]

**Tanimoto coefficient :**

\[
c / (a + b + c)
\]
‘Hash codes’

Hash codes result from an algorithm that transforms a molecular structure into a sequence of characters or numbers encoding the presence of fragments in the molecule.

They have a fixed length.

Hash codes are not interpretable. They’re used as unique identifiers of structures, e.g. in large databases of compounds hash codes allow for the fast perception of an exact match between two molecules.

Hash codes can also be defined for atoms, or bonds.
**Representation of stereochemistry**

The Cahn-Ingold-Prelog (CIP) rules

Useful for nomenclature but difficult to implement: assignment of priorities.

But in a Molfile…
Atoms are ranked. Priorities can easily be assigned corresponding to the atoms’ ranks in the Molfile.

(R) - lactic acid

(S) - lactic acid

CIP priorities: OH > CO$_2$H > CH$_3$ > H
1. Number the atoms surrounding the stereo center with 1, 2, 3, and 4 in order of increasing atom number (position in the atom block) (a hydrogen atom should be considered atom 4).

2. View the center from a position such that the bond connecting the highest-numbered atom (4) projects behind the plane formed by atoms 1, 2, and 3.

3. Parity ‘1’ if atoms 1-3 are arranged in clockwise direction in ascending numerical order, or parity ‘2’ if counterclockwise.
**Representation of stereochemistry**

**Molfile**

```
xxxx.xxxxxx.yyyyyy.zzzz aaaaaaasssshhbhhbvvvHHrrriiiimumnnnnnee

-0.6622  0.5342  0.0000  C  0  0  2  0  0  0  0
0.6622  -0.3000  0.0000  C  0  0  0  0  0  0  0
-0.7207  2.0817  0.0000  C  1  0  0  0  0  0  0
-1.8622  -0.3695  0.0000  N  0  3  0  0  0  0  0
0.6220  -1.8037  0.0000  O  0  0  0  0  0  0  0
1.9464  0.4244  0.0000  O  0  5  0  0  0  0  0
```

<table>
<thead>
<tr>
<th>Field</th>
<th>Meaning</th>
<th>Values</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>sss</td>
<td>atom stereo parity</td>
<td>0 = not stereo, 1 = odd, 2 = even, 3 = either or unmarked stereo center</td>
<td>[Generic] Ignored when read.</td>
</tr>
</tbody>
</table>

**Chiral center: atom 1.** Ligands: atoms 2, 3, 4 and H. H is the last. Looking at the chiral center with the H-atom pointing away (as in the figure) atoms 2, 3, and 4 are arranged counterclockwise. Therefore parity = 2.
Representation of stereochemistry

Molfile

1. Number the atoms surrounding the stereo center with 1, 2, 3, and 4 in order of increasing atom number (position in the atom block) (a hydrogen atom should be considered atom 4).

2. View the center from a position such that the bond connecting the highest-numbered atom (4) projects behind the plane formed by atoms 1, 2, and 3.

3. Parity ‘1’ if atoms 1-3 are arranged in clockwise direction in ascending numerical order, or parity ‘2’ if counterclockwise.

Chiral center: atom 4. Ligands: atoms 1, 3, 5, and H. H is the last. Looking at the chiral center with the H-atom pointing away (as in the figure) atoms 1, 3, and 5 are arranged clockwise. Therefore parity = 1.
### Representation of stereochemistry

**Molfile - bond block**

111222ttttssssxxxxrrrrccc

L-Alanine

![L-Alanine Structure](image)

<table>
<thead>
<tr>
<th>Field</th>
<th>Meaning</th>
<th>Values</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>111</td>
<td>first atom number</td>
<td>1 = number of atoms</td>
<td>(Generic)</td>
</tr>
<tr>
<td>222</td>
<td>second atom number</td>
<td>1 = number of atoms</td>
<td>(Generic)</td>
</tr>
<tr>
<td>ttt</td>
<td>bond type</td>
<td>1 = Single, 2 = Double, 3 = Triple, 4 = Aromatic, 5 = Single or Double,</td>
<td>(Query) Values 4 through 8 are for SSS</td>
</tr>
<tr>
<td></td>
<td></td>
<td>6 = Single or Aromatic, 7 = Double or Aromatic, 8 = Any</td>
<td>queries only.</td>
</tr>
<tr>
<td>sss</td>
<td>bond stereo</td>
<td>Single bonds: 0 = not stereo, 1 = Up, 4 = Either, 6 = Down, Double</td>
<td>(Generic) The wedge (pointed) end of the</td>
</tr>
<tr>
<td></td>
<td></td>
<td>bonds: 0 = Use x-, y-, z-coords from atom block to determine cis or</td>
<td>stereo bond is at the first atom</td>
</tr>
<tr>
<td></td>
<td></td>
<td>trans, 3 = Cis or trans (either) double bond</td>
<td>(Field 111 above)</td>
</tr>
<tr>
<td>xxx</td>
<td>not used</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ttt</td>
<td>bond topology</td>
<td>0 = Either, 1 = Ring, 2 = Chain</td>
<td>(Query) SSS queries only.</td>
</tr>
<tr>
<td>ccc</td>
<td>reacting center status</td>
<td>0 = unmarked, 1 = a center, -1 = not a center, Additional: 2 = no</td>
<td>(Reaction, Query)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>change, 4 = bond made/broken, 8 = bond order changes, 12 = 4+8</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>(both made/broken and changes); 5 = (4 + 1), 9 = (8 + 1), and 13 =</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>(12 + 1) are also possible.</td>
<td></td>
</tr>
</tbody>
</table>
**Representation of stereochemistry in SMILES notation**

Chirality in a tetrahedral center is specified by ‘@’ (clockwise direction) or ‘@@’ (counterclockwise direction). Looking to the chiral center from the ligand appearing first in the SMILES string, the other three ligands are arranged clockwise or counterclockwise in the order of appearance in the SMILES string.
**Representation of cis-trans stereochemistry in double bonds**

**Stereochemistry around a double bond** (cis/trans) is specified with characters ‘\’ and ‘/’.

Example: *trans*-1,2-dichloroethene - **Cl/C=C/Cl** (starting at the 1st Cl, a bond goes up (/) to C=C, and from here goes up (/) to the 2nd Cl).

*cis*-1,2-dichloroethene - **Cl/C=C\Cl** (starting at the 1st Cl, a bond goes up (/) to C=C, and from here goes down (\) to the 2nd Cl).
**Representation of cis-trans stereochemistry in double bonds**

Stereochemistry around a double bond (cis/trans) is specified with characters ‘\’ and ‘/’.

\[ \text{Bond goes down} \downarrow \text{Bond goes up} \]

\[ \text{Two cis substituents} \]

\[ C\backslash C(F)=C(/C)Cl \]
**Representation of the 3D structure**

The most obvious (and common) representation consists of a Cartesian system, i.e. the x, y, and z coordinates of each atom.

For a given conformation the coordinates depend on the orientation of the structure relative to the reference axes.

In a Molfile, 3D coordinates can be listed.
Representation of the 3D structure in a Molfile
Representation of the 3D structure

It is also possible to represent only coordinates, with no specification of bonds. Bonds may be inferred with reasonable confidence from the 3D interatomic distances. But demands some kind of computer processing.

```
  The Source - XYZ

File     Edit     Format

12

C    -0.85493   -1.08954   -0.31429
C    0.04142    -0.28622    0.63989
C    1.48250    -0.07374    0.08904
Cl   2.47193    0.83884     1.22712
Cl   1.52514    0.79699    -1.44386
O    -2.15604   -1.23574    0.23262
H    -0.45118   -2.10113    -0.35634
H    -0.99610   -0.52044    -1.23316
H    0.13479    -0.63867    1.54749
H    -0.39662    0.74055    0.66996
H    1.74581   -1.12601   -0.00780
H    -2.10655   -1.22985    1.19117
```
Another representation of the 3D structure is the Z matrix, in which internal coordinates are specified (bond lengths, bond angles and dihedral angles). It is mostly used for the input to quantum chemistry software. Example for cyclopropane:

```
C   0.00    0.00      0.00       0  0  0
C   1.35    0.00      0.00       1  0  0
C   1.35    60.00    0.00       2  1  0
H   1.10    110.00  120.00   3  2  1
H   1.10    110.00  240.00   3  2  1
H   1.10    110.00  120.00   2  1  3
H   1.10    110.00  240.00   2  1  3
H   1.10    110.00  120.00   1  2  3
H   1.10    110.00  240.00   1  2  3
```
**Generation of a 3D structure**

Theoretical methods:

*ab initio* (e.g. Gaussian)

semi-empirical (e.g. Mopac)

molecular mechanics (e.g. Mopac, Chem3D)

Empirical methods (e.g. CONCORD, CORINA):

use fragments with predefined geometries

use rules

use databases of geometries

use simple optimizations
Generation of the 3D structure

Chemaxon’s Marvin
Generation of the 3D structure - CORINA

http://www.mol-net.com/online_demos/corina_demo.html
Representation of molecular surfaces

The 3D structure presented up to here is just the skeleton of the molecule, but a molecule also has a ‘skin’… the molecular surface.

The molecular surface divides the 3D space in an internal volume and an external volume. This is just an analogy with macroscopic objects, since molecules cannot rigorously be approached with classical mechanics. The electronic density is continuous, and there are probabilities of finding electrons at certain locations (it tends to zero at infinite distance from nuclei).

The electronic distribution “at the surface” determines the interactions a molecule can establish with others (e.g. docking to a protein).
Representation of molecular surfaces

A molecular surface can express different properties, such as charge, electrostatic potential, or hydrophobicity, by means of colors.

Such properties may be experimentally determined (2D NMR, x-ray crystallography and electronic cryomicroscopy give indications about 3D molecular properties), or theoretically calculated.

There are several ways of defining a surface. The most used are: van der Waals surface, surface accessible to a solvent, and Connolly surface.
van der Waals surface

It is the simplest surface. It can be determined from the van der Waals radius of all atoms. Each atom is represented by a sphere. The spheres of all atoms are fused – the total volume is the van der Waals volume, and the envelop defines the van der Waals surface. It is fast to be calculated.
Connolly surface

It is generated by simulating a sphere rolling over the van der Waals surface. The sphere represents the solvent. The radius of the sphere may be chosen (typically it is set at 1.4 Å, the effective radius of water). The Connolly surface has two regions: the convex contact surface (it is a segment of the van der Waals surface) and the concave surface (where the sphere touches two or more atoms).
Surface accessible to the solvent

The path of the center of the sphere that generates the Connolly surface defines the surface accessible to the solvent.
Molecular surfaces with ChemAxon MarvinSpace
Molecular surfaces with ChemAxon MarvinSpace
Molecular surfaces with ChemAxon MarvinSpace
Molecular surfaces with ChemAxon MarvinSpace