

SUBSTANCES

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A Chemical Database

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English documentation

The **SUBSTANCES** package allows you to create a database like file that contains data of various chemicals. These data can be retrieved in the document. An index creation of the chemicals used in the document is directly supported.

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1 Licence and Requirements

Permission is granted to copy, distribute and/or modify this software under the terms of the \LaTeX Project Public License, version 1.3 or later (<http://www.latex-project.org/lppl.txt>). The package has the status “maintained.”

SUBSTANCES loads and needs the following packages: `expl3`,¹ `xparse`,² `xtemplate`³ and `l3keys2e`.⁴ It also needs the chemistry package `chemmacros`.⁵

2 About

The **SUBSTANCES** package allows you to create a database like file that contains data of various chemicals. These data can be retrieved in the document. An index creation of the chemicals used in the document is directly supported.

3 Options

SUBSTANCES has only a few options:

- `draft` = `true|false` Default: false
If set to true all warnings will be errors.
- `final` = `true|false` Default: true
The opposite of `draft`.
- `index` = `true|false` Default: false
Add index entries when `\chem` is called, see section 7.
- `style` = `<style>` Default: default
Load specific style, see section 4.3.
- `strict` = `true|false` Default: false
If set to true all warnings will be errors. This option overwrites any `draft` or `final` option that is passed on by the document class.

4 The Database

4.1 Declaring the Chemicals

The data about substances are stored via the command

- `\DeclareSubstance{<id>}{<list of properties>}`

An entry could look like this:

```
1 \DeclareSubstance{NaCl}{  
2   name    = Sodiumchloride ,  
3   sum     = NaCl ,  
4   CAS     = 7647-14-5 ,  
5   mass    = 58.44 ,  
6   mp      = 801 ,  
7   bp      = 1465 ,
```

¹ CTAN: expl3 ² CTAN: xparse ³ CTAN: xtemplate ⁴ CTAN: l3keys2e ⁵ CTAN: chemmacros

```

8   phase    = solid ,
9   density  = 2.17
10  }

```

Such entries can either be declared in the document preamble or probably more useful in a file with the ending `.sub`. Such a file can be input in the document via

- ▶ `\LoadSubstances{<filename>}` → input the file *without* specifying the file ending.

Suppose you have the file `mysubstances.sub` then you input it in the document preamble via `\LoadSubstances{mysubstances}`.

4.2 Available Fields

4.2.1 Always Defined Fields

Below all fields defined by `SUBSTANCES` are listed.⁶

- ▶ **name** required
The IUPAC name of the substance. This is the only field that *has* to be used. The field's input is parsed with chemmacros' command `\iupac`.
- ▶ **sort** optional
If you plan to use the **index** option you should specify this field to get the sorting of the index right. This then creates index entries `\index{<sort field><name field>}`.
- ▶ **alt** optional
An alternative name. The field's input is parsed with chemmacros' command `\iupac`.
- ▶ **altsort** optional
This is the same as the **sort** field but for the alternative name.
- ▶ **CAS** optional
The Chemical Abstract Service (CAS) number. The input needs to be input in the form `<num>-<num>-<num>`.
- ▶ **PubChem** optional
The PubChem number.

The **CAS** field processes the number using the macro `\CAS{<number>}` which is defined like this:

```

1  \def\CAS#1-#2-#3{\iupac{#1\-#2\-#3}}
2  \NewDocumentCommand\CAS{m}{\CAS#1}

```

You're free to redefine it to your needs.

⁶ Look in the file `substances-examples.sub` which is part of this package and should be in the same place as this documentation for example uses.

4.2.2 Style-dependent Fields

SUBSTANCES defines the style ‘default’ which is loaded if no other style has been specified. It defines the following additional fields and loads the packages chemfig⁷ and siunitx.⁸

- ▶ **formula** optional
The molecular formula of the substance. The field’s input is parsed with chemmacros’ command `\ch`.
- ▶ **structure** optional
The structural formula of the substance. The field’s input is parsed with chemfig’s command `\chemfig`.
- ▶ **mp** optional
The boiling point. The field’s entry is input into the siunitx command `\SI` in the following way: `\SI{<field>}{\celsius}`.
- ▶ **bp** optional
The melting point. The field’s entry is input into the siunitx command `\SI` in the following way: `\SI{<field>}{\celsius}`.
- ▶ **density** optional
The density. The field’s entry is input into the siunitx command `\SI` in the following way: `\SI{<field>}{\gram\per\cmc}`.
- ▶ **phase** optional
The state of aggregation.
- ▶ **pKa** optional
The pK_A value. The field’s entry is input into the siunitx command `\num`.
- ▶ **pKa1** optional
The first of several pK_A values. The field’s entry is input into the siunitx command `\num`.
- ▶ **pKa2** optional
The second of several pK_A values. The field’s entry is input into the siunitx command `\num`.
- ▶ **pKb** optional
The pK_B values. The field’s entry is input into the siunitx command `\num`.
- ▶ **pKb1** optional
The first of several pK_B values. The field’s entry is input into the siunitx command `\num`.
- ▶ **pKb2** optional
The second of several pK_B values. The field’s entry is input into the siunitx command `\num`.
- ▶ **pictograms** optional
The GHS pictograms. This field takes a list of pictogram names as they’re input into chemmacros’ command `\ghspic`.

⁷ CTAN: chemfig ⁸ CTAN: siunitx

- **H** optional
The H statements. This field takes a list of numbers as they're input into chemmacros' command `\ghs{h}{<number>}`.
- **P** optional
The P statements. This field takes a list of pictogram names as they're input into chemmacros' command `\ghs{p}{<number>}`.
- **EUH** optional
The EUH statements. This field takes a list of pictogram names as they're input into chemmacros' command `\ghs{h}{<number>}`.
- **LD50** optional
The Median Lethal Dose (LD50) in mg kg⁻¹. The field's entry is input into the siunitx command `\SI` in the following way: `\SI{<field>}{\milli\gram\per\kilo\gram}`.

4.3 Define Custom Styles

4.3.1 Background

You might have other needs for fields than the ones defined by **SUBSTANCES** and the 'default' style. You can easily define your own style which means that you save a file with the name `substances-<style>.def`. In it you save the commands you need and use the command `\DeclareSubstanceProperty`, which is explained in the next section, to declare your own fields.

Please beware that this style file is loaded by the package when the `expl3` namespace is active. This means that all spaces are ignored and you need to use `~` if you need a space. It also means that `_` and `:` are letters (as is `@`). Should you ever decide to call `\ExplSyntaxOff` to turn off the `expl3` namespace *don't* forget to use `\ExplSyntaxOn` at the end of the file!

4.3.2 Declare New Fields or Change Existing Fields

You might want other fields or change the definition of the predefined ones. For this there's

- `\DeclareSubstanceProperty*{<field name>}[<pre code>][<post code>]`

This command declares a new property field for a substance. The star makes the property a required one which means an error will be issued if a substance is declared without it. The optional arguments `<pre code>` and `<post code>` specify any code that should be input directly before or after the field entry, respectively. The `<pre code>` may end with a command that takes one mandatory argument. In this case the field entry will be its argument.

The following example would define a field **EC** which uses a custom command to parse the field entry. The European Commission Number (EC) is assigned to chemical substances for regulatory purposes within the European Union by the regulatory authorities.

```

1 \makeatletter
2 \def\@EC#1-#2-#3{#1-#2-#3}
3 \newcommand*\EC[1]{\@EC#1}
4 \makeatother
5 \DeclareSubstanceProperty{EC}[\EC]

```

For further examples of the usage of pre and post code look at the definition of the `name` and the `mp` field:

```

1 \DeclareSubstanceProperty*{name}[\iupac]
2 \DeclareSubstanceProperty{mp}[\SI][\celsius]

```

5 Retrieving the Data

There are two commands defined by `SUBSTANCES` that allow the retrieving of the data. The command `\chem` is intended as user command, the command `\GetSubstanceProperty` can be used to define your own user command (perhaps in your own style file, see section 4.3).

► `\chem*{<pre>}[<post>]{<id>}[<property>]`

► `\GetSubstanceProperty{<id>}{<property>}`

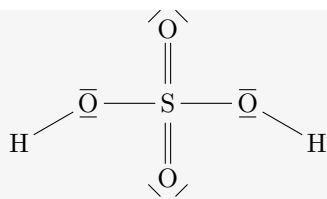
If the command `\chem` is called without the `[<property>]` argument the name entry will be called. The starred version calls the alt entry if it is defined and the name entry otherwise. The arguments `[<pre>]` and `[<post>]` add arbitrary input before or after the output, respectively.

All of the next examples use the data defined in the file `substances-examples.sub` that is part of this package, see section 8.

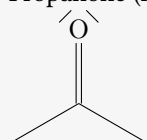
```

1 \chem{H2SO4}[structure] \newline
2 \chem{H2SO4} has the boiling point \(\chem[T_b=]{H2SO4}[bp]\) and a
3 density of \(\chem[\rho=]{H2SO4}[density]\).
4
5 Compare the melting points of methane and ethane,
6 \(\chem[T_m=]{methane}[mp]\) and \(\chem[T_m=]{ethane}[mp]\),
7 with the boiling points \(\chem[T_b=]{methane}[bp]\) and
8 \(\chem[T_b=]{ethane}[bp]\).
9
10 \chem{NaCl} has the \ac{CAS} number \chem{NaCl}[CAS].
11
12 \chem{acetone} (\chem*{acetone}) is the most simple ketone:
13
14 \chem{acetone}[structure]

```



Sulfuric Acid has the boiling point $T_b = 279.6\text{ }^{\circ}\text{C}$ and a density of $\rho = 1.8356\text{ g cm}^{-3}$.
 Compare the melting points of methane and ethane, $T_m = -182\text{ }^{\circ}\text{C}$ and $T_m = -183\text{ }^{\circ}\text{C}$, with
 the boiling points $T_b = -162\text{ }^{\circ}\text{C}$ and $T_b = -89\text{ }^{\circ}\text{C}$.
 Sodiumchloride has the CAS number 7647-14-5.
 Propanone (Acetone) is the most simple ketone:



The following code creates table 1.

```

1 \begin{table}[htp]
2 \centering\chemsetup[ghsystem]{hide}
3 \sisetup{scientific-notation=fixed,fixed-exponent=0,per-mode=
  symbol}
4 \begin{tabular}{l>\raggedright\arraybackslashp{.6\linewidth}}
5 \toprule
6 name & & \chem{methane} \\
7 formula & & \chem{methane}[formula] \\
8 & & \chem{methane}[structure] \\
9 \midrule
10 CAS & & \chem{methane}[CAS] \\
11 PubChem & & \chem{methane}[PubChem] \\
12 \midrule
13 boiling point & & \chem{methane}[bp] \\
14 melting point & & \chem{methane}[mp] \\
15 density & & \chem{methane}[density] \\
16 molar mass & & \chem{methane}[mass] \\
17 \midrule
18 & & \chem{methane}[pictograms] \\
19 H statements & & \chem{methane}[H] \\
20 P statements & & \chem{methane}[P] \\
21 \bottomrule
22 \end{tabular}
23 \caption{\label{tab:methane}All properties of \chem{methane} that
  have been
24 saved in the example database.}
25 \end{table}

```


name	Methane
formula	CH ₄ $ \begin{array}{c} \text{H} \\ \\ \text{H} - \text{C} - \text{H} \\ \\ \text{H} \end{array} $
CAS	74-82-8
PubChem	297
boiling point	−162 °C
melting point	−182 °C
density	0.000 72 g/cm ³
molar mass	16.04 g/mol
	
H statements	H220
P statements	P210, P377, P381, P410 + P403

Table 1: All properties of Methane that have been saved in the example database.

6 Additional Commands

SUBSTANCES provides a few commands that maybe are useful in building custom macros for styles. A field exists if it has been defined with `\DeclareSubstanceProperty` regardless if it has been used or not. A substance exists if it has been defined with `\DeclareSubstance`.

- ▶ `\GetSubstanceProperty{<id>}{<field>}` → You know that already: retrieve the property specified in <field> for a given substance. This command is *not* expandable.
- ▶ `\RetrieveSubstanceProperty{<id>}{<field>}` → Same as `\GetSubstanceProperty` but expandable.
- ▶ `\ForAllSubstancesDo{<code>}` → Loops through all existing substances. Inside <code> #1 may be used to refer to the <id> of the current substance. This command is expandable.
- ▶ `\AllSubstancesSequence` → A sequence of all substances. This is a sequence of balanced groups each containing the <id> of a substance. This command is expandable.
- ▶ `\AllSubstancesClist` → A comma separated list of all substances. Every <id> is separated from the next with a comma. This command is expandable.
- ▶ `\IfSubstancePropertyTF{<id>}{<field>}{<true code>}{<false code>}` → Tests if the property <field> is defined for the substance <id> and returns either <true code> or <false code>.

code>. This command is expandable.

- ▶ `\IfSubstancePropertyT{<id>}{<field>}{<true code>}` → Tests if the property <field> is defined for the substance <id> and returns <true code> if it is. This command is expandable.
- ▶ `\IfSubstancePropertyF{<id>}{<field>}{<false code>}` → Tests if the property <field> is defined for the substance <id> and returns <false code> if it isn't. This command is expandable.
- ▶ `\IfSubstanceFieldTF{<field>}{<true code>}{<false code>}` → Tests if the property <field> exists and returns either <true code> or <false code>. This command is expandable.
- ▶ `\IfSubstanceFieldT{<field>}{<true code>}` → Tests if the property <field> exists and returns <true code> if it does. This command is expandable.
- ▶ `\IfSubstanceFieldF{<field>}{<false code>}` → Tests if the property <field> exists and returns <false code> if it doesn't. This command is expandable.
- ▶ `\IfSubstanceExistTF{<id>}{<true code>}{<false code>}` → Tests if the property <field> exists and returns either <true code> or <false code>. This command is expandable.
- ▶ `\IfSubstanceExistT{<id>}{<true code>}` → Tests if the substance <id> exists and returns <true code> if it does. This command is expandable.
- ▶ `\IfSubstanceExistF{<id>}{<false code>}` → Tests if the substance <id> exists and returns <false code> if it doesn't. This command is expandable.

```
1 Just to demonstrate how these commands can be used:\par
2 \newcounter{substances}
3 \ForAllSubstancesDo{%
4   \ifnum0=\value{substances}\relax
5   \else, \fi
6   \stepcounter{substances}%
7   \chem{#1}%
8   \IfSubstancePropertyT{#1}{alt}{
9     (\chem*{#1})}}
```

Just to demonstrate how these commands can be used:

Sodiumchloride, Hydrochloric Acid, Nitric Acid, Sulfuric Acid, Methane, Ethane, Propane, Butane (*n*-Butane), Pentane (*n*-Pentane), Hexane (*n*-Hexane), Heptane (*n*-Heptane), Octane (*n*-Octane), Nonane (*n*-Nonane), Decane (*n*-Decane), Propanone (Acetone)

7 Create an Index

When `SUBSTANCES` is called with `index = true` the command `\chem` will add index entries each time it is used. In this case the entries of the fields `name`, `sort`, `alt` and `altsort` will be

expanded during the process. You should keep that in mind if some error arises. It might be due to a `\textbf` or similar in your database. In this case you either need to replace it with some robust command or put a `\noexpand` in front of it.

Alternative names as specified in the `alt` also get an index entry with a reference to the one of the corresponding `name` field. The entry of the `name` field in this case gets the `alt` name appended in braces.

This behaviour is not customizable for the time being. It is planned for future versions of this package, though.

As a demonstration an index for all chemicals used in this documentation is created with the help of the package `imakeidx`.⁹

7.1 Formatting Commands

The index entries are formatted with the following commands. You can redefine them to your needs. If you do make sure they have the same number of required arguments and are expandable!

- ▶ `\SubstanceIndexNameEntry#1#2{#1@#2}` → Formats the name if no `alt` field is given. #1 refers to the `sort` field entry and #2 refers to the `name` field entry.
- ▶ `\SubstanceIndexNameAltEntry#1#2#3{#1@#2 (#3)}` → Formats the name if also the `alt` field is given. #1 refers to the `sort` field entry, #2 refers to the `name` field entry, and #3 to the `alt` field entry.
- ▶ `\SubstanceIndexAltEntry#1#2#3{#1@#3|see#2}` → Formats the entry for the `alt` field. #1 refers to the `altsort` field entry, #2 refers to the `name` field entry, and #3 to the `alt` field entry.

7.2 Using `makeidx`

Using the option `index = true` with the standard way to create an index will add the entries `\index{<name>}` to the index. This means you would mix them with other entries if you have any. Below a sample document is shown that needs to be compiled with `pdflatex`, `makeindex` and again with `pdflatex`.

```
1 \documentclass{article}
2 \usepackage[T1]{fontenc}
3 \usepackage[index]{substances}
4 \LoadSubstances{substances-examples}
5
6 \usepackage{makeidx}
7 \makeindex
8 \begin{document}
9
10 \newcounter{substances}
11 \ForAllSubstancesDo{%
12   \ifnum0=\value{substances}\relax
13   \else, \fi
```

⁹ CTAN: `imakeidx`

```

14 \stepcounter{substances}\chem{#1}}
15
16 \printindex
17 \end{document}

```

7.3 Using splitidx

Maybe a separate index for the chemicals will make more sense. In this case you could use the package `splitidx`.¹⁰ `SUBSTANCES` will recognize this and create `\sindex[\jobname-chem]{<name>}` entries each time `\chem` is used. The sample document below needs to be compiled with `pdflatex`, `splitindex` and again with `pdflatex`.

```

1 \documentclass{article}
2 \usepackage[T1]{fontenc}
3 \usepackage[index]{substances}
4 \LoadSubstances{substances-examples}
5
6 \usepackage{splitidx}
7 \makeindex
8 \newindex[Chemicals]{\jobname-chem}
9 \begin{document}
10
11 \newcounter{substances}
12 \ForAllSubstancesDo{%
13   \ifnum0=\value{substances}\relax
14   \else, \fi
15   \stepcounter{substances}\chem{#1}}
16
17 \printindex[\jobname-chem]
18 \end{document}

```

7.4 Using imakeidx

Another way to create multiple indexes is the package `imakeidx`. `SUBSTANCES` recognizes its usage and creates index entries `\index[\jobname-chem]{<name>}`. The sample document below needs to be compiled with `pdflatex -shell-escape`.

```

1 \documentclass{article}
2 \usepackage[T1]{fontenc}
3 \usepackage[index]{substances}
4 \LoadSubstances{substances-examples}
5
6 \usepackage{imakeidx}
7 \makeindex[name=\jobname-chem,title=Chemicals]
8 \begin{document}

```

¹⁰ CTAN: `splitidx`

```

9
10 \newcounter{substances}
11 \ForAllSubstancesDo{%
12   \ifnum0=\value{substances}\relax
13   \else, \fi
14   \stepcounter{substances}\chem{#1}}
15
16 \printindex[\jobname-chem]
17 \end{document}

```

8 The Example Database

The following code shows the example database `substances-examples.sub` that is part of this package.

```

1  % -----
2  % the SUBSTANCES package
3  %
4  %   A Chemical Database
5  %
6  % -----
7  % Clemens Niederberger
8  % Web:   https://bitbucket.org/cgnieder/substances/
9  % E-Mail: contact@mychemistry.eu
10 % -----
11 % Copyright 2012 Clemens Niederberger
12 %
13 % This work may be distributed and/or modified under the
14 % conditions of the LaTeX Project Public License, either version 1.3
15 % of this license or (at your option) any later version.
16 % The latest version of this license is in
17 %   http://www.latex-project.org/lppl.txt
18 % and version 1.3 or later is part of all distributions of LaTeX
19 % version 2005/12/01 or later.
20 %
21 % This work has the LPPL maintenance status 'maintained'.
22 %
23 % The Current Maintainer of this work is Clemens Niederberger.
24 % -----
25 % The substances package consists of the files
26 % - substances.sty, substances-default.def, substances-examples.sub,
27 %   substances_en.tex, substances_en.pdf, README
28 % -----
29 % If you have any ideas, questions, suggestions or bugs to report, please
30 % feel free to contact me.
31 % -----
32 %
33 % example database to the package 'substances'
34 %
35 \DeclareChemIUPAC\normal{\textit{n}}{
36 \DeclareSubstance{NaCl}{
37   name      = Sodium\|chloride ,

```

```

38     sort      = Sodiumchloride ,
39     formula   = NaCl ,
40     CAS       = 7647-14-5,
41     mass      = 58.44 ,
42     mp        = 801 ,
43     bp        = 1465 ,
44     phase     = solid ,
45     density   = 2.17
46 }
47
48 \DeclareSubstance{HCl}{
49     name      = Hydro\|chloric Acid ,
50     formula   = HCl ,
51     CAS       = 7647-01-0 ,
52     pictograms = {acid,exclam} ,
53     H         = {314,335} ,
54     P         = {260,301+330+331,303+361+353,305+351+338,405,501} ,
55     mass      = 36.46 ,
56     density   = 1.19 ,
57     mp        = -30
58 }
59
60 \DeclareSubstance{HNO3}{
61     name      = Nitric Acid ,
62     sort      = Nitric Acid ,
63     formula   = HNO3 ,
64     CAS       = 7697-37-2 ,
65     PubChem   = 944 ,
66     mass      = 63.01 ,
67     density   = 1.51 ,
68     mp        = -42 ,
69     bp        = 86 ,
70     pKa       = -1.37 ,
71     pictograms = {flame-0,acid} ,
72     H         = {272,314} ,
73     P         = {220,280,305+351+338,310}
74 }
75
76 \DeclareSubstance{H2SO4}{
77     name      = Sulfuric Acid ,
78     sort      = Sulfuric Acid ,
79     formula   = H2SO4 ,
80     structure = {H-[:30]\Lewis{26,0}-S(=[2]\Lewis{13,0})(=[6]\Lewis{57,0})-\Lewis{26,
81     0}-[:30]H} ,
82     CAS       = 7664-93-9 ,
83     PubChem   = 1118 ,
84     mass      = 98.08 ,
85     density   = 1.8356 ,
86     mp        = 10.38 ,
87     bp        = 279.6 ,
88     phase     = liquid ,
89     pKa       = -3.0 ,
90     pKa1      = -3.0 ,
91     pKa2      = 1.9 ,

```

```

91 pictograms = acid ,
92 H          = 314 ,
93 P          = {280,301+330+331,309,310,305+351+338} ,
94 LD50       = 510
95 }
96
97 \DeclareSubstance{methane}{
98 name       = Methane ,
99 sort       = Methane ,
100 formula    = CH4 ,
101 structure   = H-C(-[2]H)(-[6]H)-H ,
102 CAS        = 74-82-8 ,
103 PubChem    = 297 ,
104 pictograms = {flame,bottle} ,
105 H          = 220 ,
106 P          = {210,377,381,410+403} ,
107 mass       = 16.04 ,
108 density    = 0.72e-3 ,
109 mp         = -182 ,
110 bp         = -162 ,
111 phase      = gaseous
112 }
113
114 \DeclareSubstance{ethane}{
115 name       = Ethane ,
116 sort       = Ethane ,
117 formula    = C2H6 ,
118 structure   = H-C(-[2]H)(-[6]H)-C(-[2]H)(-[6]H)-H ,
119 CAS        = 74-84-0 ,
120 PubChem    = 6324 ,
121 pictograms = {flame,bottle} ,
122 H          = 220 ,
123 P          = {210,377,381,403} ,
124 mass       = 30.07 ,
125 density    = 0.72e-3 ,
126 mp         = -183 ,
127 bp         = -89 ,
128 phase      = gaseous
129 }
130
131 \DeclareSubstance{propane}{
132 name       = Propane ,
133 sort       = Propane ,
134 formula    = C3H8 ,
135 structure   = H-C(-[2]H)(-[6]H)-C(-[2]H)(-[6]H)-C(-[2]H)(-[6]H)-H ,
136 CAS        = 74-98-6 ,
137 pictograms = {flame,bottle} ,
138 H          = 220 ,
139 P          = {201,377,381,403} ,
140 mass       = 44.10 ,
141 density    = 2.01e-3 ,
142 mp         = -188 ,
143 bp         = -42 ,
144 phase      = gaseous

```

```

145 }
146
147 \DeclareSubstance{butane}{
148   name      = Butane ,
149   sort      = Butane ,
150   alt       = \normal\ -Butane ,
151   altsort   = n-Butane ,
152   formula   = C4H10 ,
153   structure = H-C(-[2]H)(-[6]H)-C(-[2]H)(-[6]H)-C(-[2]H)(-[6]H)-C(-[2]H)(-[6]H)-H ,
154   CAS       = 106-97-8 ,
155   PubChem   = 7843 ,
156   pictograms = {flame,bottle} ,
157   H         = {220,280} ,
158   P         = {201,377,381,403} ,
159   mass      = 58.12 ,
160   density   = 2.71e-3 ,
161   mp        = -138.3 ,
162   bp        = -0.5 ,
163   phase     = gaseous
164 }
165
166 \DeclareSubstance{pentane}{
167   name      = Pentane ,
168   sort      = Pentane ,
169   alt       = \normal\ -Pentane ,
170   altsort   = n-Pentane ,
171   formula   = C5H12 ,
172   structure = H-C(-[2]H)(-[6]H)-C(-[2]H)(-[6]H)-C(-[2]H)(-[6]H)-C(-[2]H)(-[6]H)-C
173   (-[2]H)(-[6]H)-H ,
174   CAS       = 109-66-0 ,
175   PubChem   = 8003 ,
176   pictograms = {flame,health,exclam,aqpol} ,
177   H         = {225,304,336,411} ,
178   EUH       = 066 ,
179   P         = {273,301+310,331,403+235} ,
180   mass      = 72.15 ,
181   density   = 0.63 ,
182   mp        = -130 ,
183   bp        = 36 ,
184   phase     = liquid
185 }
186
187 \DeclareSubstance{hexane}{
188   name      = Hexane ,
189   sort      = Hexane ,
190   alt       = \normal\ -Hexane ,
191   altsort   = n-Hexane ,
192   formula   = C6H14 ,
193   structure = -[:30]-[:30]-[:30]-[:30]-[:30] ,
194   CAS       = 110-54-3 ,
195   PubChem   = 8058 ,
196   pictograms = {flame,health,exclam,aqpol} ,
197   H         = {225,361f,304,373,315,336,411} ,
198   P         = {210,240,273,301+310,331,302+352,403+235} ,

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198     mass      = 86.18 ,
199     density   = 0.66 ,
200     mp        = -95 ,
201     bp        = 69 ,
202     phase     = liquid
203 }
204
205 \DeclareSubstance{heptane}{
206     name      = Heptane ,
207     sort      = Heptane ,
208     alt       = \normal\ -Heptane ,
209     altsort   = n-Heptane ,
210     formula   = C7H16 ,
211     structure = -[:30]-[: -30]-[:30]-[: -30]-[:30]-[: -30] ,
212     CAS       = 142-82-5 ,
213     PubChem   = 8900 ,
214     pictograms = {flame,health,exclam,aqpol} ,
215     H         = {225,304,315,336,410} ,
216     P         = {210,273,301+310,331,302+352,403+235} ,
217     mass      = 100.21 ,
218     density   = 0.68 ,
219     mp        = -91 ,
220     bp        = 98 ,
221     phase     = liquid
222 }
223
224 \DeclareSubstance{octane}{
225     name      = Octane ,
226     sort      = Octane ,
227     alt       = \normal\ -Octane ,
228     altsort   = n-Octane ,
229     formula   = C8H18 ,
230     structure = -[:30]-[: -30]-[:30]-[: -30]-[:30]-[: -30]-[:30] ,
231     CAS       = 111-65-9 ,
232     PubChem   = 356 ,
233     pictograms = {flame,health,exclam,aqpol} ,
234     H         = {225,304,315,336,410} ,
235     P         = {210,273,301+330+331,302+352} ,
236     mass      = 114.23 ,
237     density   = 0.70 ,
238     mp        = -56.8 ,
239     bp        = 126 ,
240     phase     = liquid
241 }
242
243 \DeclareSubstance{nonane}{
244     name      = Nonane ,
245     sort      = Nonane ,
246     alt       = \normal\ -Nonane ,
247     altsort   = n-Nonane ,
248     formula   = C9H20 ,
249     structure = -[:30]-[: -30]-[:30]-[: -30]-[:30]-[: -30]-[:30]-[: -30] ,
250     CAS       = 111-84-2 ,
251     PubChem   = 8141 ,

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252 pictograms = {flame,exclam,health} ,
253 H           = {226,304,315,319,332,336,413} ,
254 P           = {261,301+310,305+351+338,331} ,
255 mass        = 128.26 ,
256 density     = 0.72 ,
257 mp          = -54 ,
258 bp          = 151 ,
259 phase       = liquid
260 }
261
262 \DeclareSubstance{decane}{
263   name        = Decane ,
264   sort        = Decane ,
265   alt         = \normal-Decane ,
266   altsort     = n-Decane ,
267   formula     = C10H22 ,
268   structure   = -[:30]-[:30]-[:30]-[:30]-[:30]-[:30]-[:30]-[:30]-[:30] ,
269   CAS         = 124-18-5 ,
270   PubChem     = 15600 ,
271   pictograms  = {flame,health} ,
272   H           = {226,304} ,
273   P           = {210,260,262,301+310,331} ,
274   mass        = 142.29 ,
275   density     = 0.73 ,
276   mp          = -29.7 ,
277   bp          = 174 ,
278   phase       = liquid
279 }
280
281 \DeclareSubstance{acetone}{
282   name        = Propanone ,
283   sort        = Propanone ,
284   alt         = Acetone ,
285   altsort     = Acetone ,
286   formula     = C3H6O ,
287   structure   = {-[:30](=[2]\Lewis{13,0})-[:30]} ,
288   CAS         = 67-64-1 ,
289   PubChem     = 180 ,
290   mass        = 58.08 ,
291   density     = 0.79 ,
292   mp          = -95 ,
293   bp          = 56 ,
294   pictograms  = {flame,exclam} ,
295   H           = {225,319,336} ,
296   EUH         = {066} ,
297   P           = {210,233,305+351+338} ,
298   LD50        = 5800
299 }
300
301 \endinput

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