BOHR

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simple atom representation according to the Bohr model

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This package provides means for the creation of simple Bohr models of atoms up to the atomic number 112. Additionally commands are provided to convert atomic numbers to element symbols or element names and vice versa.

The package is inspired by a question on http://tex.stackexchange.com/: Draw Bohr atomic model with electron shells in TEX?

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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 (LPPL), version 1.3 or later (http://www.latex-project.org/lppl.txt). The software has the status "maintained."

The BOHR package loads and needs the packages pgf¹ [Tan10] and cnltx-base² [Nie13].

2 Options

Every option described in the manual can also be used as package option. Options are indicated as option and are all key/value like options. Some options can be set without value, too. Then

^{1.} on CTAN as pgf: http://mirrors.ctan.org/graphics/pgf/

^{2.} on CTAN as cnltx: http://mirrors.ctan.org/macros/latex/contrib/cnltx/

the underlined value is used.

3 Usage

BOHR is used like any other LATEX 2ε package:

```
ı \usepackage{bohr}
```

The main command, \bohr, creates the models:

```
\box{bohr}[\langle num\ of\ shells \rangle] \{\langle num\ of\ electrons \rangle\} \{\langle atom\ name \rangle\}
```

The main command. The mandatory arguments take the number of electrons to be printed and the atom symbol that is printed in the center.

This is described best by an example:

```
ı \bohr{3}{Li}
```

There is not much more to it. Another example using the optional argument:

```
1 \bohr[2]{2}{$\mathrm{Li^+}$}
```

4 Customization

BOHR provides a handful of options to customize the appearance:

```
\strut \strut
```

Options are set in a key/value syntax using this command.

```
insert-symbol = true|false
```

Default: false

If set to true **BOHR** will insert the atomic symbol suiting to the given electron number if *no* third argument is given.

```
insert-number = true|false
```

Default: false

If set to true **BOHR** will use the appropriate number of electrons for the given element symbol in the third argument if *no* second argument is given. This of course only works if the third argument is one of the 112 element symbols.

4 Customization

Default: false insert-missing = true|false Sets both insert-symbol and insert-number. atom-style = $\{\langle code \rangle\}$ (initially empty) This code will be placed immediatly before the third argument of \bohr. The last macro in it may need one argument. $name-options-set = \{\langle tikz \ option \rangle\}$ (initially empty) This value is passed to the options of the \node the third argument of \bohr is placed in. $name-options-add = \{\langle tikz \ options \rangle\}$ (initially empty) This value will be added to options set with name-options-set. $nucleus-options-set = \{\langle tikz \ options \rangle\}$ Default: draw=black!80, fill=black!10, opacity=.25 This value is passed to the options of the \draw command that draws the circle around the name-node. $nucleus-options-add = \{\langle tikz \ options \rangle\}$ (initially empty) This value will be added to options set with nucleus-options-set. $nucleus-radius = \{\langle dimension \rangle\}$ Default: 1em The radius of the circle around the name-node. electron-options-set = $\{\langle tikz \ options \rangle\}$ Default: blue!50!black!50 This value is passed to the options of the \fill command that draws the electrons. electron-options-add = $\{\langle tikz \ options \rangle\}$ (initially empty) This value will be added to options set with electron-options-set. electron-radius = $\{\langle dimension \rangle\}$ Default: 1.5pt The radius of the circles that represent the electrons. $shell-options-set = \{\langle tikz \ options \rangle\}$ Default: draw=blue!75,thin This value is passed to the options of the \draw command that draws the circles that represent the shells. $shell-options-add = \{\langle tikz \ options \rangle\}$ (initially empty) This value will be added to options set with shell-options-set. Default: 1em $shell-dist = \{\langle dimension \rangle\}$ The distance between the nucleus and the first shell and between subsequent shells. $language = \{\langle language \rangle\}$ (initially empty)

Introduced in version 0.2d

Select the language used for the element names manually rather than letting BOHR select it automatically. This option must be used as package option or in the preamble in order to have an effect. Currently provided languages are English, French and German. If this option is not used the document language at the end of the preamble (as set by babel or polyglossia) is used.

distribution-method = periodic|quantum

Default: quantum

Introduced in version 0.3

Determines how the electrons are distributed on the shells. periodic distributes the electrons 2-8-8-18-18-32-32, *i. e.*, according to the place of the corresponding atom in the periodic table of elements. quantum distributes the electrons according the the electron configuration of the corresponding atom where each shell represents the main quantum number. Pd for example has the configuration $1s^22s^22p^63s^23p^63d^{10}4s^24p^64d^{10}$ and would get two electrons on the first shell, 8 electrons on the second, and 18 electrons each on the forth and fifth.

```
1 \setbohr{name-options-set={font=\footnotesize\sffamily}}
2 \bohr{2}{He} \bohr{7}{N}
```

```
1 % uses package 'chemmacros'
2 \setbohr{atom-style={\footnotesize\sffamily\ch}}
3 \bohr{0}{H+} \bohr{10}{F-}
```

```
1 \setbohr{
2   shell-options-add = dashed,
3   shell-dist = .5em,
4   insert-missing
5 }
6 \bohr{6}{} \bohr{}{K}
```

5 Additional Commands

BOHR provides some additional commands that return the element symbol or the element name to a given atomic number and vice versa.

Returns the element symbol for a given atomic number or element name. \elementsymbol{80}: Hg; \elementsymbol{rhenium}: Rh.

```
\ensuremath{\mbox{\mbox{element name}}} \arrowvert \a
```

Returns the element name for a given atomic number or element symbol. \elementname{80}: Mercury; \elementname{Rh}: Rhenium.

```
\arrowvert a tomic number { \langle element name \rangle | \langle element symbol \rangle }
```

Returns the atomic number for a given element name or element symbol. \atomicnumber{Hg}: 80; \atomicnumber{rhenium} 75.

\Z

If this command isn't defined by some other package it is available as an alias of \atomicnumber.

```
1 The elements \elementname{F}, \elementname{Cl}, \elementname{Br},
2 \elementname{I} and \elementname{At} are called \emph{halogens}.
```

The elements Fluorine, Chlorine, Bromine, Iodine and Astatine are called *halogens*.

6 Internal Commands

The element properties used by **BOHR** are defined through the following commands:

```
\DeclareAtomSymbol{\langle atomic\ number\rangle}{\langle atom\ symbol\rangle}
```

The package file contains 112 lines like the following: \DeclareAtomSymbol{29}{Cu}. They associate atomic number with the corresponding atom symbol. This command could be used to change an element's symbol.

```
\DeclareAtomName{\langle atomic\ number \rangle}{\langle element\ name \rangle}
```

BOHR comes with a few language files named bohr_elements_ $\langle language \rangle$. def. They contain of 112 entries like \DeclareAtomName{29}{Copper} which associate atomic number and element name. This command could be used to change an element's name.

```
\DeclareElectronDistribution{\langle atomic number \rangle}{\langle electron distribution \rangle}
```

The ⟨electron distribution⟩ is a comma separated list of integers which determines how the electrons are distributed on the shells when distribution-method = quantum. The package file contains 112 entries like \DeclareElectronDistribution{29}{2,8,18,1}. This command could be used to change an element's electron configuration.

References

```
[Nie13] Clemens Niederberger. cnltx. version 0.7a, Oct. 4, 2013.

URL: http://mirror.ctan.org/macros/latex/contrib/cnltx/.
```

[Tan1o] Till Tantau. TikZ/pgf. version 2.10, Oct. 25, 2010. URL: http://mirror.ctan.org/graphics/pgf/.

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