MODIAGRAM

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easy creation of molecular orbital diagrams

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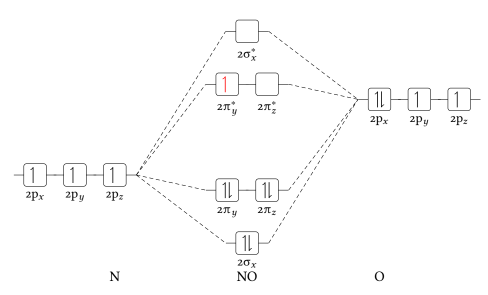


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1 Licence, Requirements

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1 Licence, Requirements

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MODIAGRAM uses l3kernel [L3Pa] and l3packages [L3Pb]. It also uses TikZ [Tan19] and the package chemgreek [Nie16] bundle. Additionally the TikZ libraries calc and arrows are loaded. Knowledge of TikZ is helpful.

2 Motivation

This package has been written as a reaction to a question on http://tex.stackexchange.com/. To be more precise: as a reaction to the question "Molecular orbital diagrams in LaTeX." There it says

I'm wondering if anyone has seen a package for drawing (qualitative) molecular orbital splitting diagrams in LaTeX? Or if there exist any packages that can be easily re-purposed to this task?

Otherwise, I think I'll have a go at it in TikZ.

The problem was solved using TikZ, since no package existed for that purpose. For one thing MODIAGRAM is intended to fill this gap. I also found it very tedious, to make all this copying and pasting when I needed a second, third, ... diagram. MODIAGRAM took care of that.

3 Main Commands

All molecular orbital (MO) diagrams are created using the environment modiagram.

3.1 The \atom Command

```
\atom[\langle name \rangle] \{\langle left \rangle | \langle right \rangle\} \{\langle AO\text{-}spec \rangle\}
```

Place an AO in the diagram. $\langle name \rangle$ is caption of the atom, $\langle left \rangle$ and $\langle right \rangle$ determine the placement in the diagram, $\langle AO\text{-}spec \rangle$ is the specification of the AO.

Let's take a look at an example:

As you can see, the argument $\langle AO\text{-}spec \rangle$ is essential to create the actual orbitals and the electrons within. You can use these key/value pairs to specify what you need:

```
1s = {\langle rel\text{-}energy\rangle; \langle el\text{-}spec\rangle}
Energy level and electron specifications for the 1s orbital.
```

```
2s = {\langle rel\text{-}energy\rangle; \langle el\text{-}spec\rangle}
Energy level and electron specifications for the 2s orbital.
```

```
2p = {\langle rel\text{-}energy \rangle; \langle x el\text{-}spec \rangle, \langle y el\text{-}spec \rangle, \langle z el\text{-}spec \rangle}
Energy level and electron specifications for the 2p orbitals.
```

 $\langle el\text{-}spec \rangle$ can have the values pair, up and down or can be left empty. $\langle rel\text{-}energy \rangle$ actually is the y coordinate and shifts the AO vertically by $\langle rel\text{-}energy \rangle$ cm.

The argument $\langle left \rangle / \langle right \rangle$ is important, when p orbitals are used. For instance compare the following example to the one before:

When both variants are used one can also see, that the right atom is shifted to the right (hence the naming). The right atom is shifted by 4 cm per default and can be adjusted individually, see page 18.

```
1 \begin{modiagram}
  \atom{left}{
     1s = \{ 0; pair \},
     2s = { 1; pair} ,
     2p = \{1.5; up, down \}
   \atom{right}{
     1s = \{ 0; pair \},
     2s = { 1; pair} ,
     2p = \{1.5; up, down \}
12 \end{modiagram}
  + + -
                                 + + -
          #
                                 #
          #
                                 #
```

With the command \molecule (section 3.2) the reason for the shift becomes clear. Any of the arguments for the AO can be left empty or be omitted.

```
Without argument: default height, full:

| Without argument: default height, full:
| :\par
| \begin{modiagram}
| atom{left}{1s, 2s, 2p}
| \end{modiagram}
| \text{#
```

```
using some values:

— + —

using some values:

begin{modiagram}

a \atom{left}{1s, 2s=1, 2p={;,up}}

end{modiagram}

— 

#
```

3.2 The \molecule Command

```
\mbox{molecule}[\langle name \rangle] \{\langle MO\text{-}spec \rangle\}
```

Place a MO in the diagram. $\langle name \rangle$ is caption of the molecule, $\langle MO\text{-}spec \rangle$ is the specification of the MO.

An example first:

```
begin{modiagram}
    \atom{left} { 1s = { 0; up } }
    \atom{right}{ 1s = { 0; up } }
    \molecule { 1sM0 = {.75; pair } }
    \end{modiagram}
```

The command \molecule connects the AOs with the bonding and anti-bondung MOs. \molecule can only be used *after* one has set *both* atoms since the orbitals that are to be connected must be known.

The argument $\langle MO\text{-}spec \rangle$ accepts a comma separated list of key/value pairs:

```
1sM0 = {\langle energy gain \rangle / \langle energy loss \rangle; \langle s el-spec \rangle, \langle s^* el-spec \rangle} connects the AOs specified by 1s.

2sM0 = {\langle energy gain \rangle / \langle energy loss \rangle; \langle s el-spec \rangle, \langle s^* el-spec \rangle} connects the AOs specified by 2s

2pM0 = {\langle s energy gain \rangle / \langle s energy loss \rangle, \langle p energy gain \rangle / \langle p energy loss \rangle; \langle s el-spec \rangle, \langle p el-spec \rangle, \langle pz^* el-spec \rangle, \langle s^* el-spec \rangle \rangle s^* el-spec \rangle \rangle s el-spec \rangle \rangle \rangle s el-spec \rangle \rangle \rangle s el-spec \rangle \rangle \rangle s el-spec \rangl
```

Obviously the regarding AOs must have been set in order to connect them. This for example won't work:

```
1 \begin{modiagram}
2 \atom{left} { 1s = 0 }
3 \atom{right}{ 1s = 0 }
4 \molecule { 2sM0 = .75 }
5 \end{modiagram}
```

The value used in $\langle energy \ gain \rangle$ determines how many cm the bonding MO lies below the lower AO or how many cm the anti-bondung MO lies above the higher AO.

If you specify *(energy loss)* you can create non-symmetrical splittings. Then, the first value

 $(\langle energy \ gain \rangle)$ is used for the bonding MO and the second value $(\langle energy \ loss \rangle)$ is used for the anti-bonding MO.

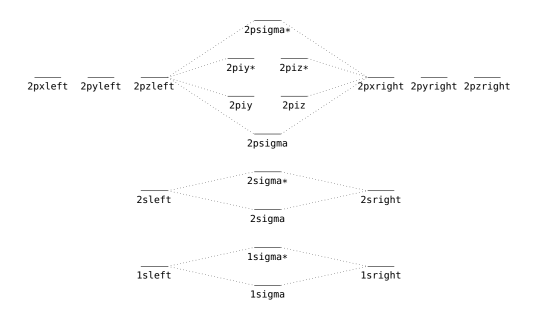
Please be aware, that you have to specify *two* such values or pairs with 2pM0: the splitting of the σ orbitals and the splitting of the π orbitals.

```
1 \begin{modiagram}
2 \atom{left} { 2p = { 0; up, up } }
3 \atom{right}{ 2p = { 1; up, up } }
4 \molecule { 2pM0 = { 1.5, .75; pair, up, up } }
5 \end{modiagram}
+ + -
```

The complete MO diagram for triplett dioxygen now could look something like that:

3.3 The Naming Scheme

Since one wants to be able to put labels to the orbitals and since they are nodes in a tikzpicture, the internal naming scheme is important. It closely follows the function:



With these names it is possible to reference the orbitals with the known TikZcommands:

```
1 \begin{modiagram}
2 \atom{left} { 1s = 0 }
3 \atom{right}{ 1s = 0 }
4 \molecule { 1sM0 = .75 }
5 \draw[<-->, red, semithick]
6 (1sigma.center) -- (1sigma*.center
        );
7 \draw[red]
8 (1sigma*) ++ (2cm, .5cm) node {
        splitting};
9 \end{modiagram}
```

```
begin{modiagram}

atom{left} { 1s = 0 }

atom{right}{ 1s = 0 }

molecule { 1sM0 = .75 }

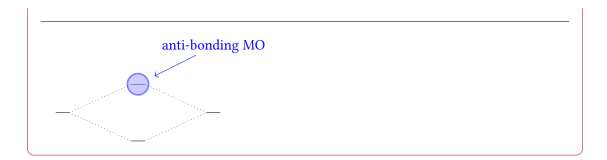
draw[draw=blue,very thick,fill=blue!40,opacity=.5]

(1sigma*) circle (8pt);

draw[<-,shorten <=8pt,shorten >=15pt,blue]

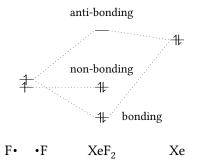
(1sigma*) --++(2,1) node {anti-bonding M0};

end{modiagram}
```



3.4 Placing AOs and MOs Arbitrarily

The standard orbitals are not always sufficient in order to draw a correct MO diagram. For example in the MO diagram of XeF_2 one would need the part that illustrates the interaction between the bonding and anti-bonding combination of two p orbitals of Flourine with one p orbital of Xenon:



To create diagrams like this there is the following command, which draws a single AO:

```
AO[\langle name \rangle](\langle xshift \rangle) \{\langle type \rangle\} [\langle options \rangle] \{\langle energy \rangle; \langle el-spec \rangle\}
```

Place an AO in the diagram. $\langle name \rangle$ (optional) is the name of the node; if not specified, AO# is used where # is a consecutive number. $\langle xshift \rangle$ is the vertical position of the orbital(s), a TeX dimension. $\langle type \rangle$ can be s or p. $\langle options \rangle$ is a list of key/value pairs with which the AO can be customized, see section 4.3. $\langle AO\text{-}spec \rangle$ is the specification of the AO.

Depending on the $\langle type \rangle$ one s or three p orbitals are drawn.

```
1 \begin{modiagram}
2 \A0{s}{0;}
3 \A0(-20pt){p}{1;pair,up,down}
4 \end{modiagram}
```

If one wants to place such an AO at the position of an atom, one has to know their $\langle xshift \rangle$. They have predefined values (also see section 3.5):

atom left: 1 cmmolecule: 3 cmatom right: 5 cm

Within the p orbitals there is an additional shift by 20 pt per orbital. This is equivalent to a double shift by the length A0-width (see section 4.1.3):

```
1 \begin{modiagram}
    \atom{left} {2p=0}
    \arrowvert a tom{right}{2p=0}
    % above the left atom:
    \A0(1cm)
                {s}{ .5}
   A0(1cm-20pt){s}{1;up}
    A0(1cm-40pt){s}{1,5;down}
   % above the right atom:
    \A0(1cm)
                 {s}{ .5}
    A0(5cm+20pt){s}{1;up}
    A0(5cm+40pt){s}{1.5;down}
12 \end{modiagram}
  +
           #
```

The AOs created with \land also can be connected. For this you can use the TikZ command \land draw, of course. You can use the predefined node names...

```
1 \begin{modiagram}
2 \A0{s}{0} \A0(2cm){s}{1}
3 \A0{s}{2} \A0(2cm){s}{1.5}
4 \draw[red] (A01.0) -- (A02.180) (A03.0) -- (A04.180);
5 \end{modiagram}
```

... or use own node names

```
1 \begin{modiagram}
2 \A0[a]{s}{0} \A0[b](2cm){s}{1}
3 \A0[c]{s}{2} \A0[d](2cm){s}{1.5}
4 \draw[red] (a.0) -- (b.180) (c.0) -- (d.180);
5 \end{modiagram}
```

The predefined names are A01, A02 etc.for the type s and A01x, A01y, A01z, A02x etc. for the type p. Nodes of the type p get an x, y or z if you specify your own name, too.

```
1 \begin{modiagram}
2 \A0{p}{0}
3 \draw[<-,shorten >=5pt] (A01y.-90) -- ++ (.5,-1) node {y};
4 \end{modiagram}
5 and
6 \begin{modiagram}
7 \A0[A]{p}{0}
8 \draw[<-,shorten >=5pt] (Ay.-90) -- ++ (.5,-1) node {y};
9 \end{modiagram}
```



However, if you want the lines to be drawn in the same style as the ones created by \molecule, you should use the command \connect.

```
\connect{\langle AO\text{-}connect \rangle}
```

Connects the specified AOs. $\langle AO\text{-}connect \rangle$ is comma separated list of node name pairs connected with &.

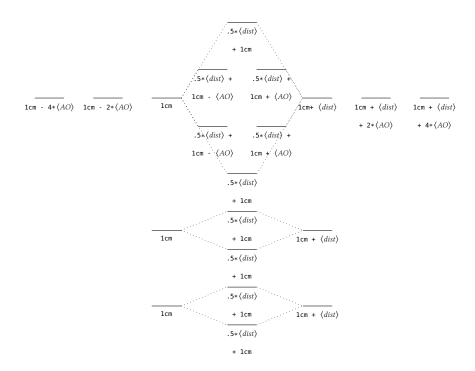
This command expects a comma separated list of node name pairs that are to be connected. The names have to be connected with a &:

Some things still need to be said: \connect adds the anchor east to the first name and the anchor west to the second one. This means a connection only makes sense from the left to the right. However, you can add own anchors using the usual TikZ way:

^{1.} which can be customized, see page 21

3.5 The Positioning Scheme

The figure below shows the values of the x coordinates of the orbitals depending on the values of $\langle distance \rangle$ ($\langle dist \rangle$) and $\langle AO-width \rangle$ ($\langle AO \rangle$). In sections 4.1.2 and 4.1.3 these lengths and how they can be changed are discussed.



3.6 Default Values

If you leave the arguments (or better: values) for the specification of the AO or MO empty or omit them, default values are used. The table below shows you, which ones.

	AO/MO	omitted	empty
syntax:		1s	1s=
	1s	{0;pair}	{0;}
	2s	{2;pair}	{2;}
	2p	{5;pair,pair,pair}	{5;,,}
	1sMO	{.5;pair,pair}	{.5;,}
	2sM0	<pre>{.5;pair,pair}</pre>	{.5;,}
	2pM0	{1.5,.5;pair,pair,pair,pair,pair,pair}	{1.5,.5;,,,,,}

This is similar for the A0 command (page 10); it needs a value for $\langle energy \rangle$, though.

(type)	⟨el-spec⟩
S	pair
p	pair,pair,pair

Compare these examples:

4 Customization

The options of the section 4.1 can be set global as package option, *i. e.*, with $\usepackage[\langle options \rangle] \{ modiagram \}$, or via the setup command $\setmodiagram \{ \langle options \rangle \}$.

4.1 Environment Options

There are some options with which the layout of the MO diagrams can be changed:

```
style = \{\langle type \rangle\}
   change the style of the orbitals and the connecting lines, section 4.1.1.
distance = \{\langle dim \rangle\}
   distance betwen left and right atom, section 4.1.2.
A0-width = \{\langle dim \rangle\}
   change the width of orbitals, section 4.1.3.
el-sep = \{\langle num \rangle\}
   distance between the electron pair arrows, section 4.1.4.
up-el-pos = \{\langle num \rangle\}
   position of the spin-up arrow, section 4.1.4.
down-el-pos = \{\langle num \rangle\}
   position of the spin-down arrow, section 4.1.4.
lines = \{\langle tikz \rangle\}
   change the TikZ style of the connecting lines, section 4.1.5.
names = true|false
   add captions to the atoms and the molecule, section 4.1.6.
```

```
names-style = \{\langle tikz \rangle\} change the TikZ style of the captions, section 4.1.7.

names-style-add = \{\langle tikz \rangle\} change the TikZ style of the captions, section 4.1.7.

labels = \underline{\text{true}}|\text{false} add default labels to the orbitals, section 4.1.8.

labels-fs = \{\langle cs \rangle\} change the font size of the labels, section 4.1.9.

labels-style = \{\langle tikz \rangle\} change the TikZ style of the labels, section 4.1.10.
```

They all are discussed in the following sections. If they're used as options for the environment, they're set locally and only change that environment.

```
1 \begin{modiagram}[options]
2 ...
3 \end{modiagram}
```

4.1.1 Option style

There are five different styles which can be chosen.

- style = {plain} # (default)
- style = {square} 1
- style = {circle}
- style = {round}
- style = {fancy} -11-

Let's take the MO diagram of H₂ to illustrate the different styles:

```
1 % use package `chemmacros'
2 \begin{modiagram}[style=plain]%
    default
3   \atom[H]{left} { 1s = {;up} }
4   \atom[H]{right}{ 1s = {;up} }
5   \molecule[\ch{H2}]{ 1sM0 = {.75;pair}
} }
6 \end{modiagram}
```

```
1 % use package `chemmacros'
2 \begin{modiagram}[style=square]
3 \atom[H]{left} { 1s = {;up} }
4 \atom[H]{right}{ 1s = {;up} }
5 \molecule[\ch{H2}]{ 1sM0 = {.75;pair } }
6 \end{modiagram}
```

```
1 % use package `chemmacros'
2 \begin{modiagram}[style=circle]
3  \atom[H]{left} { 1s = {;up} }
4  \atom[H]{right}{ 1s = {;up} }
5  \molecule[\ch{H2}]{ 1sM0 = {.75;pair } }
6 \end{modiagram}
```

```
1 % use package `chemmacros'
2 \begin{modiagram}[style=round]
3  \atom[H]{left} { 1s = {;up} }
4  \atom[H]{right}{ 1s = {;up} }
5  \molecule[\ch{H2}]{ 1sM0 = {.75;pair } }
6 \end{modiagram}
```

```
1 % use package `chemmacros'
2 \begin{modiagram}[style=fancy]
3 \atom[H]{left} { 1s = {;up} }
4 \atom[H]{right}{ 1s = {;up} }
5 \molecule[\ch{H2}]{ 1sM0 = {.75;pair }
} }
6 \end{modiagram}
```

4.1.2 Option distance

Depending on labels and captions the 4 cm by which the right and left atom are separated can be too small. With distance = $\{\langle dim \rangle\}$ the length can be adjusted. This will change the position

of the right atom to 1cm + $\langle dim \rangle$ and the position of the molecule is changed to 0.5*(1cm + $\langle dim \rangle$), also see page 10 and section 3.5.

```
1 % use package `chemmacros'
2 \begin{modiagram}[distance=6cm]
3  \atom[H]{left} { 1s = {;up} }
4  \atom[H]{right}{ 1s = {;up} }
5  \molecule[\ch{H2}]{ 1sM0 = {.75;pair } }
6 \end{modiagram}
```

4.1.3 Option A0-width

The length A0-width sets the length of the horizontal line in a orbital displayed with the plain style. It's default value is 10 pt.

```
1 % use package `chemmacros'
2 \begin{modiagram}[A0-width=15pt]
3   \atom[H]{left} { 1s = {;up} }
4   \atom[H]{right}{ 1s = {;up} }
5   \molecule[\ch{H2}]{ 1sM0 = {.75;pair } }
6 \end{modiagram}
```

By changing the value of A0-width the positions of the p and the π orbitals also change, see section 3.5.

4.1.4 Optionen el-sep, up-el-pos und down-el-pos

These three options change the horizontal positions of the arrows representing the electrons in an AO/MO. The option el-sep = $\{\langle num \rangle\}$ needs a value between 0 and 1. 0 means *no* distance

between the arrows and 1 *full* distance (with respect to the length AO-width, see section 4.1.3).

```
1 % use package `chemmacros'
2 \begin{modiagram}[el-sep=.2]% default
3  \atom[H]{left} { 1s = {;up} }
4  \atom[H]{right}{ 1s = {;up} }
5  \molecule[\ch{H2}]{ 1sM0 = {.75;pair } }
6 \end{modiagram}
```

```
1 % use package `chemmacros'
2 \begin{modiagram}[el-sep=0]
3  \atom[H]{left} { 1s = {;up} }
4  \atom[H]{right}{ 1s = {;up} }
5  \molecule[\ch{H2}]{ 1sM0 = {.75;pair } }
6 \end{modiagram}
```

```
1 % use package `chemmacros'
2 \begin{modiagram}[el-sep=1]
3 \atom[H]{left} { 1s = {;up} }
4 \atom[H]{right}{ 1s = {;up} }
5 \molecule[\ch{H2}]{ 1sM0 = {.75;pair }
} }
6 \end{modiagram}
```

The options $up-el-pos = \{\langle < num > \rangle\}$ and $down-el-pos = \{\langle < num > \rangle\}$ can be used alternatively to place the spin-up and spin-down electron, respectively. Again they need values between 0 and 1. This time 0 means on the left and 1 means on the right.

```
1 % use package `chemmacros'
2 \begin{modiagram}[up-el-pos=.4,down-el-pos=.6]% default
3 \atom[H]{left} { 1s = {;up} }
4 \atom[H]{right}{ 1s = {;up} }
5 \molecule[\ch{H2}]{ 1sM0 = {.75;pair} }
6 \end{modiagram}
```



```
1 % use package `chemmacros'
2 \begin{modiagram}[up-el-pos=.333,down-el-pos=.667]
3 \atom[H]{left} { 1s = {;up} }
4 \atom[H]{right}{ 1s = {;up} }
5 \molecule[\ch{H2}]{ 1sM0 = {.75;pair} }
6 \end{modiagram}
```

```
1 % use package `chemmacros'
2 \begin{modiagram}[up-el-pos=.7,down-el-pos=.3]
3  \atom[H]{left} { 1s = {;up} }
4  \atom[H]{right}{ 1s = {;up} }
5  \molecule[\ch{H2}]{ 1sM0 = {.75;pair} }
6 \end{modiagram}
```

4.1.5 Option lines

The option lines can be used to modify the TikZ style of the connecting lines:

```
1 % use package `chemmacros'
2 \begin{modiagram}[lines={gray,thin}]
3  \atom[H]{left} { 1s = {;up} }
4  \atom[H]{right}{ 1s = {;up} }
5  \molecule[\ch{H2}]{ 1sM0 = {.75;pair } }
6 \end{modiagram}
```

4.1.6 Option names

If you use the option names the atoms and the molecule get captions provided you have used the optional $\langle name \rangle$ argument of \atom and/or \molecule.

4.1.7 Options names-style and names-style-add

These options enable to customize the style of the captions of the atoms and of the molecule. By default this setting is used: names-style = $\{\langle anchor = base \rangle\}$.²

```
1 % use package `chemmacros'
2 \begin{modiagram}[names,names-style={
    draw=blue}]
3    \atom[p]{left} { 1s = {;up} }
4    \atom[b]{right}{ 1s = {;up} }
5    \molecule[\ch{X2}]{ 1sM0 = {.75;pair } }
6    \end{modiagram}
p
X2
b
```

With this the default setting is overwritten. As you can see it destroys the vertical alignment of the nodes. In order to avoid that you can for example specify text height and text depth yourself ...

^{2.} Please see "TikZ and PGF - Manual for Version 2.10" p. 183 section 16.4.4 (pgfmanual.pdf) for the meaning

..., add the anchor again ...

```
1 % use package `chemmacros'
2 \begin{modiagram}[names,names-style={anchor=base, draw=blue}]
3 \atom[p]{left} { 1s = {;up} }
4 \atom[b]{right}{ 1s = {;up} }
5 \molecule[\ch{X2}]{ 1sM0 = {.75;pair} }
6 \end{modiagram}
```

... or use the option names-style-add = $\{\langle . \rangle\}$ It doesn't overwrite the current setting but appends the new declaration:

```
1 % use package `chemmacros'
2 \begin{modiagram}[names,names-style-add={draw=blue}]
3 \atom[p]{left} { 1s = {;up} }
```

```
4 \atom[b]{right}{ 1s = {;up} }
5 \molecule[\ch{X2}]{ 1sM0 = {.75;pair} }
6 \end{modiagram}

The state of th
```

```
1 % use package `chemmacros'
2 \setmodiagram{
   names,
    names-style = {
     text height = 2.5ex,
     text depth = .5ex,
     draw = blue!80,
      rounded corners
9
10 }
\begin{modiagram}
    \\atom[p]{left} { 1s = {;up} }
    \\atom[b]{right}{ 1s = {;up} }
    \mbox{molecule}[\ch{X2}]{ 1sM0 = {.75;pair} }
_{15} \setminus end\{modiagram\}
\atom[p]{left} { 1s = {;up} }
    \atom[b]{right}{ 1s = {;up} }
18
    \mbox{molecule}[\ch{X2}]{ 1sM0 = {.75;pair} }
20 \end{modiagram}
                          b
                                                   b
              X_2
                                       X_2
   p
                             p
```

4.1.8 Option labels

If you use the option labels predefined labels are written below the orbitals. These labels can be changed, see section 4.2.1.

```
1 % use package `chemmacros'
2 \begin{modiagram}[labels]
3 \atom[H]{left} { 1s = {;up} }
4 \atom[H]{right}{ 1s = {;up} }
5 \molecule[\ch{H2}]{ 1sM0 = {.75;pair }
} }
6 \end{modiagram}
```

4.1.9 Option labels-fs

Labels are set with the font size \small. If you want to change that you can use the option labels-fs.

```
1 % use package `chemmacros'
2 \begin{modiagram}[labels,labels-fs=\footnotesize]
3 \atom[H]{left} { 1s = {;up} }
4 \atom[H]{right}{ 1s = {;up} }
5 \molecule[\ch{H2}]{ 1sM0 = {.75;pair} }
6 \end{modiagram}
```

This also allows you to change the font style or font shape of the labels.

```
1 % use package `chemmacros'
2 \begin{modiagram}[labels,labels-fs=\sffamily\footnotesize]
3 \atom[H]{left} { 1s = {;up} }
4 \atom[H]{right}{ 1s = {;up} }
5 \molecule[\ch{H2}]{ 1sM0 = {.75;pair} }
6 \end{modiagram}
```



4.1.10 Option labels-style

The option labels-style changes the TikZ style of the nodes within which the labels are written.

```
1 % use package `chemmacros'
2 \begin{modiagram}[labels,labels-style={blue,yshift=4pt}]
3 \atom[H]{left} { 1s = {;up} }
4 \atom[H]{right}{ 1s = {;up} }
5 \molecule[\ch{H2}]{ 1sM0 = {.75;pair} }
6 \end{modiagram}
```

4.2 \atom and \molecule Specific Customizations

4.2.1 The label Key

If you don't want to use the predefined labels, change single labels or use only one or two labels, you can use the key label. This option is used in the \atom and \molecule commands in the $\langle AO\text{-}spec \rangle$ or $\langle MO\text{-}spec \rangle$ argument, respectively. The key awaits a comma separated key/value list. The names mentioned in section 3.3 are used as keys to specify the AO that you want to label.

```
1 \begin{modiagram}[style=square,
   distance=6cm]
   \atom{left} { 1s = {;up} }
   \displaystyle \operatorname{tom}{right}{1s = {;up}}
   \molecule{
                                                                      anti-bonding
      1sM0 = \{.75; pair\},
     label = {
        1sigma = \chemsigma,
        1sigma* = \chemsigma$^*$
                                                                   bonding
     }
   }
   \node[right] at (1sigma.-45) {
   bonding);
   \node[right] at (1sigma*.45) {anti-
   bonding;
\end{modiagram}
```

If the option is used together with the labels option (page 24) single labels are overwritten:

```
1 % use package `chemmacros'
2 \begin{modiagram}[labels]
3 \atom[H]{left} { 1s = {;up} }
4 \atom[H]{right}{ 1s = {;up} }
5 \molecule[\ch{H2}]{
6    1sM0 = {.75;pair},
7    label = { 1sigma = \textcolor{red} }{???} }
8  }
9 \end{modiagram}
```

4.2.2 The color Key

Analogous to the label key the color key can be used to display coloured electrons:

4.2.3 The up-el-pos and down-el-pos keys

The options up-el-pos and down-el-pos allow it to shift the arrows representing the electrons in a single AO or MO individually. You need to use values between 0 and 1, also see section 4.1.4.

4.3 \A0 Specific Customizations

These keys enable to customize orbitals created with \AO .

4.3.1 The label Key

The key label [$\langle x \rangle / \langle y \rangle / \langle z \rangle$] allows you to put a label to the AO/MO. If you use the type p you can specify the orbital you want to label in square brackets:

```
| The property of the control of the
```

4.3.2 The color Key

Analogous to the label key there is the key color $[\langle x \rangle / \langle y \rangle / \langle z \rangle]$ which enables you to choose a color for the electrons. If you use the type p you can specify the orbital in square brackets:

4.3.3 The up-el-pos and down-el-pos Keys

Then there are the keys up-el-pos[$\langle x \rangle / \langle y \rangle / \langle z \rangle$] and down-el-pos[$\langle x \rangle / \langle y \rangle / \langle z \rangle$] with which the electrons can be shifted horizontally. You can use values between 0 and 1, also see section 4.1.4. If you use the type p you can specify the orbital in square brackets:

4.4 Energy Axis

Last but not least one might want to add an energy axis to the diagram. For this there is the command \EnergyAxis.

\EnergyAxis[\langle option \rangle]

Adds an energy axis to the diagram. *(options)* are key/value pairs to modify the axis.

```
1 \begin{modiagram}
2 \atom{left} { 1s = {;up} }
3 \atom{right}{ 1s = {;up} }
4 \molecule{ 1sM0 = {.75;pair} }
5 \EnergyAxis
6 \end{modiagram}
```

For the time being there are two options to modify the axis.

 $title = \{\langle title \rangle\}$ Default: energy

the axis label. If used without value the default is used.

 $head = \{\langle tikz \ arrow \ head \rangle\}$ Default: >

the arrow head; you can use the arrow heads specified in the TikZ library arrows (pgfmanual v2.10 pages 256ff.)

```
energy

1 \begin{modiagram}
2 \atom{left} { 1s = {;up} }
3 \atom{right}{ 1s = {;up} }
4 \molecule{ 1sM0 = {.75;pair} }
5 \EnergyAxis[title]
6 \end{modiagram}
```

```
begin{modiagram}
    \begin{modiagram}
    \atom{left} { 1s = {;up} }
    \atom{right}{ 1s = {;up} }
    \molecule{ 1sM0 = {.75;pair} }
    \EnergyAxis[title=E,head=stealth]
    \end{modiagram}
```

5 Examples

The example from the beginning of section 3.4.

```
1 % use package `chemmacros'
2 \begin{modiagram}[names]
    \atom[\chlewis{0.}{F}\hspace*{5mm}\chlewis{180.}{F}]{left}{
      1s=.2;up,up-el-pos={1sleft=.5}
5
   \atom[Xe]{right}{1s=1.25;pair}
   \molecule[\ch{XeF2}]{1sM0={1/.25;pair}}
   A0(1cm){s}{0;up}
   A0(3cm){s}{0;pair}
   \connect{ A01 & A02 }
    \node[right,xshift=4mm] at (1sigma) {\footnotesize bonding};
    \node[above] at (A02.90) {\footnotesize non-bonding};
    \node[above] at (1sigma*.90) {\footnotesize anti-bonding};
14 \end{modiagram}
              anti-bonding
             non-bonding
              -----1
                     bonding
      •F
                             Xe
   F•
                XeF<sub>2</sub>
```

```
1 % use package `chemmacros'
2 \begin{figure}[p]
3  \centering
4  \begin{modiagram}[style=square,labels,names,A0-width=8pt,labels-fs=\footnotesize]
5  \atom[\ch{0_a}]{left}{
6    ls, 2s, 2p = {;pair,up,up}
7  }
8  \atom[\ch{0_b}]{right}{
9    ls, 2s, 2p = {;pair,up,up}
10  }
11  \molecule[\ch{02}]{
```

5 Examples

```
_{\scriptscriptstyle 1} % use package `chemmacros'
2 \begin{figure}[p]
    \centering
    \setmodiagram{style = fancy, distance = 7cm, AO-width = 15pt, labels}
    \begin{modiagram}
      \atom[N]{left}{
        2p = \{0; up, up, up\}
      \atom[0]{right}{
        2p = \{2; pair, up, up\}
10
11
      \molecule[NO]{
12
        2pM0 = {1.8,.4;pair,pair,pair,up},
13
         color = { 2piy*=red }
14
      }
15
      \EnergyAxis
   \end{modiagram}
    \caption{Part of the MO diagram of \chlewis{180.}{NO}.}
19 \end{figure}
```

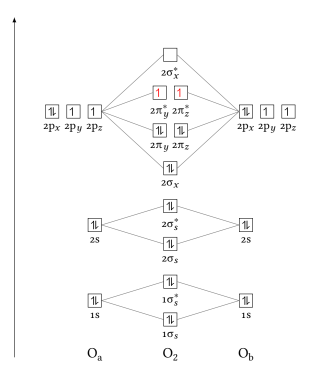


FIGURE 1: MO diagram of $^3\Sigma$ -O₂.

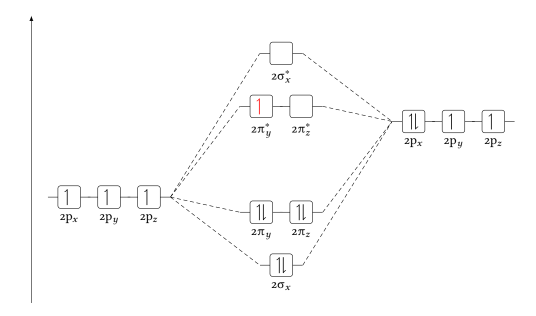


FIGURE 2: Part of the MO diagram of •NO.

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