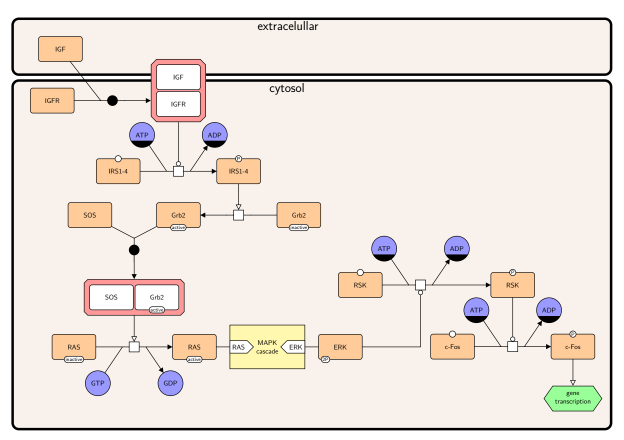
sbgntikz

manual for version 0.1

Adrien Rougny

May 9, 2018



 ${\it Map adapted from SBGN PD language L1V1.3, Journal of Integrative Bioinformatics, 2015 Sep~4;} 12(2):263$

Contents

1	Introduction			
	1.1	About	2	
	1.2	Installation and usage	2	
	1.3	A first map	3	
2	Dra	wing glyphs	5	
	2.1	Nodes	5	
	2.2	Arcs	9	
	2.3	Nodes' and arcs' attributes	11	
3	Customizing and drawing maps effectively			
	3.1	Useful options for nodes and arcs	14	
	3.2	Positioning of nodes, arcs, and their attributes	15	
	3.3	Bended arcs, multi-part arcs	16	
		Nodes along paths for ER maps		

1 Introduction

1.1 About

sbgntikz is a TikZ [1] library to draw SBGN PD, AF or ER maps [2] directly into LATEX documents. It basically encodes SBGN glyphs into TikZ shapes and arrowheads named by keywords, making them straightforwardly drawable within a TikZ picture. Drawing a specific glyph is then as simple as specifying its corresponding keyword in some TikZ command.

The present manual is intended for an audience that knows SBGN but not particularly TikZ. The rest of the present section is dedicated to the first steps in using sbgntikz: installing the library and drawing a first map (while introducing some basic TikZ syntax). Section 2 references all glyphs and their associated keywords, whereas section 3 gives some TikZ options and syntaxes that I find most useful to draw SBGN maps. I believe users already familiar with TikZ will mostly be interested in reading section 2, and might have different (and maybe better) solutions to the issues presented in section 3.

1.2 Installation and usage

The directory tikz-sbgn/ should be copied to a directory where it can be found by the TFX engine:

- in the directory of your LATEX source file
- in your local texmf directory (/home/<user>/texmf/ under Linux, /Users/Library/texmf/ under MacOS).

Usually, TikZ is installed within your TEX distribution, so TikZ and sbgntikz can be imported directly into your IATEX source file with no further installation adding the following two commands to your preamble:

```
\usepackage{tikz}
\usetikzlibrary{sbgn}
```

An SBGN map can then be drawn within a TikZ picture using the sbgn key:

```
\begin{tikzpicture}[sbgn]
% tikz code to draw an SBGN map
\end{tikzpicture}
```

1.3 A first map

SBGN is all about drawing nodes with specific shapes and arcs with specific arrow heads. Fortunately, drawing TikZ pictures is not different, making it pretty straightforward to draw SBGN maps using sbgntikz: the \node command is used to draw nodes, while the \draw command is used to draw arcs. The code to draw an SBGN node (or an attribute) will usually look like the following:

```
\node[<sbgn node>, ...] (name) at (point) {LABEL};
```

where

- <sbgn node> is a keyword corresponding to the type of node to be drawn (e.g. simple chemical for a simple chemical);
- ... is a list of other options for the node (e.g. its relative positioning towards another node, color, line width ...);
- (name) specifies the name of the node (optional);
- at (point) specifies the point on the canvas where to draw the node (optional, by default (0,0) if no relative positioning is specified in the nodes' options);
- {LABEL} specifies the label of the node that will be displayed (mandatory but can be empty).

As for arcs, they can be drawn using the following piece of code:

```
\draw[<sbgn arc>, ...] (a) -- (b);
```

where

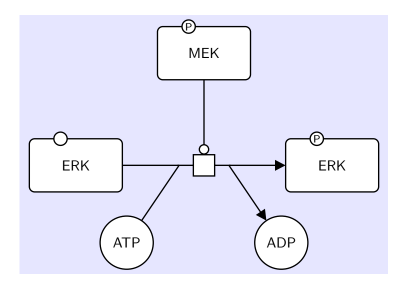
• <sbgn arc> is a keyword corresponding to the type of arc to be drawn (e.g. necessary stimulation for a necessary stimulation);

- ... is a list of other options for the arc (e.g. its color, line width ...);
- (a) is a point on the canvas or the name of a node from which the arc will depart;
- (b) is a point on the canvas or the name of a node on which the arc will arrive.

Knowing those two basic syntaxes, one can draw pretty much any desired SBGN map. Following is an example of code to draw a simple PD map. It relies on relative positioning provided by TikZ's positioning library, as positioning all nodes with absolute coordinates would be too cumbersome (see section 3 for few more details, or the PGF/TikZ manual for lot more details).

```
\documentclass{standalone}
\usepackage{tikz}
\usetikzlibrary{positioning, sbgn}
\begin{document}
\begin{tikzpicture}[sbgn]
\node[macromolecule] (erk) {ERK};
                                    % this node has no absolute nor relative positioning, so it
\hookrightarrow is placed at (0,0) by default
\node[sv] at (erk.120) {}; % the state variable is placed on the border of the node, at an angle
\hookrightarrow of 120 deg
\node[generic process, connectors = horizontal, right = of erk] (p) {}; % we add connectors, and
\hookrightarrow use relative positioning
\node[macromolecule, right = of p] (perk) {ERK};
\node[sv] at (perk.120) {P};
\node[simple chemical, below left = of p] (atp) {ATP};
\node[simple chemical, below right = of p] (adp) {ADP};
\node[macromolecule, above = 2cm of p] (pmek) {MEK};
\node[sv] at (pmek.120) {P};
\draw[consumption] (erk) -- (p.west); % p being the name of the process node, p.west is the tip
\hookrightarrow of its left connector
\draw[consumption] (atp) -- (p.west);
\draw[production] (p.east) -- (perk);
\draw[production] (p.east) -- (adp);
\draw[catalysis] (pmek) -- (p);
\end{tikzpicture}
\end{document}
```

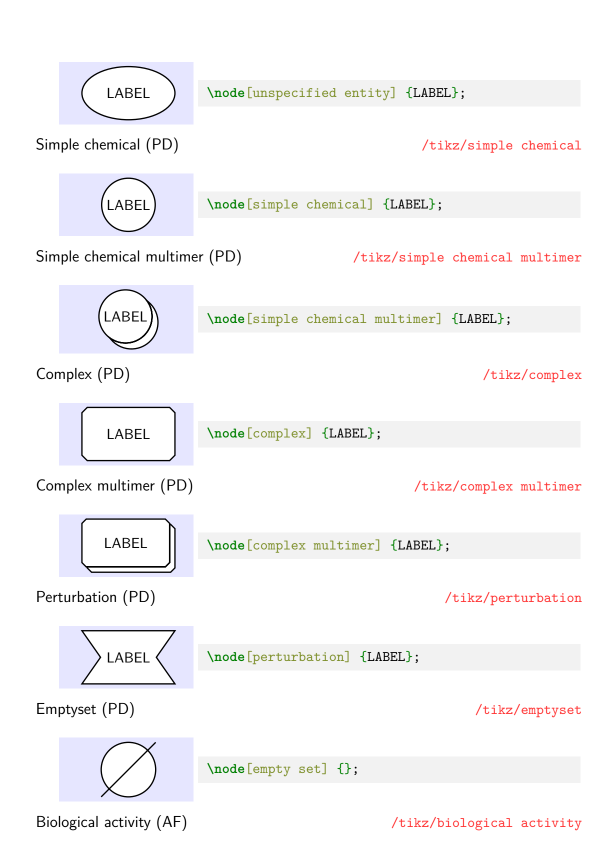
Compiling the above code would produce the following figure:

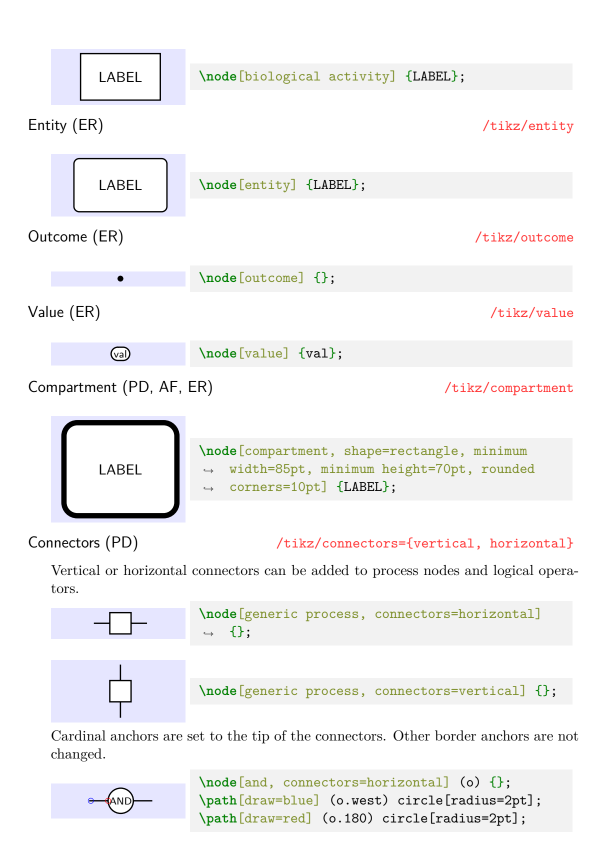


2 Drawing glyphs

2.1 Nodes



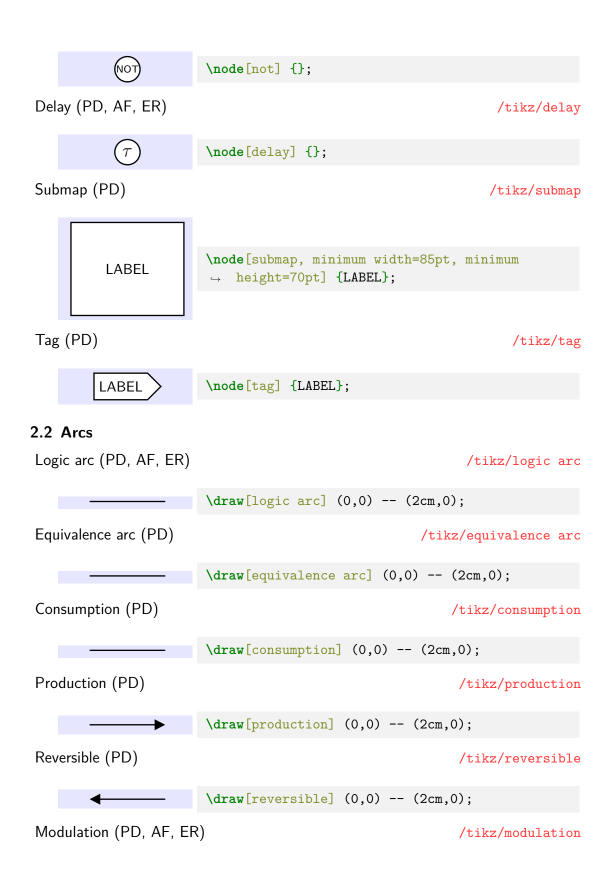




Clone (PD) /tikz/clone **LABEL** \node[unspecified entity, clone] {LABEL}; Generic process (PD) /tikz/generic process \node[generic process] {}; Omitted process (PD) /tikz/omitted process $\overline{\mathbb{M}}$ \node[omitted process] {}; Unknown process (PD) /tikz/unknown process ? \node[unknown process] {}; Association (PD) /tikz/association \node[association] {}; Dissociation (PD) /tikz/dissociation \bigcirc \node[dissociation] {}; Phenotype (PD, AF, ER) /tikz/phenotype \node[phenotype] {LABEL}; LABEL And (PD, AF, ER) /tikz/and (AND) \node[and] {}; Or (PD, AF, ER) /tikz/or (OR) \node[or] {};

/tikz/not

Not (PD, AF, ER)



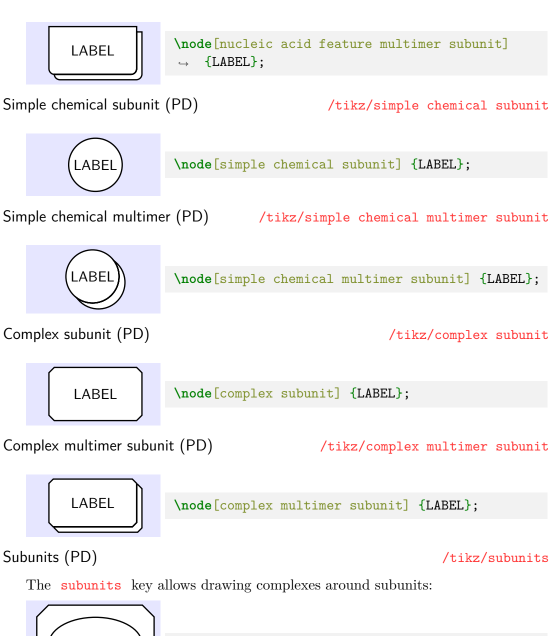
```
\draw[modulation] (0,0) -- (2cm,0);
Stimulation (PD, AF, ER)
                                                          /tikz/stimulation
                       \draw[stimulation] (0,0) -- (2cm,0);
Necessary stimulation (PD, AF, ER)
                                               /tikz/necessary stimulation
                       \draw[necessary stimulation] (0,0) -- (2cm,0);
Absolute stimulation (ER)
                                                /tikz/absolute stimulation
               \draw[absolute stimulation] (0,0) -- (2cm,0);
Inhibition (PD, AF, ER)
                                                           /tikz/inhibition
                        \draw[inhibition] (0,0) -- (2cm,0);
Absolute inhibition (ER)
                                                 /tikz/absolute inhibition
                        \draw[absolute inhibition] (0,0) -- (2cm,0);
Assignment (ER)
                                                           /tikz/assignment
                        \draw[assignment] (0,0) -- (2cm,0);
Interaction (ER)
                                                          /tikz/interaction
                       \draw[interaction] (0,0) -- (2cm,0);
   N-ary interactions can be drawn using the nary node:
                       \draw[interaction] (0,0) -- node[nary, pos=0.5]
                        \rightarrow (a) {} (2cm,0);
                        \node[outcome] at (a.100) {};
Anchor point (ER)
                                                         /tikz/anchor point
                       \draw[interaction] (0,0) -- coordinate[anchor
                        \rightarrow point, pos = 0.5] (a) (2,0);
                        \draw[stimulation] (1,1) -- (a);
```

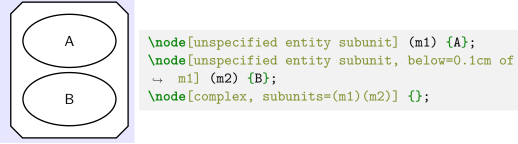
2.3 Nodes' and arcs' attributes

State variable (PD, ER) /tikz/sv (val@var) \node[entity, draw=gray!60] (m) {}; \node[sv] at (m.north) {val@var}; Existence state variable (ER) /tikz/sv existence lacksquare\node[entity, draw=gray!60] (m) {}; \node[sv existence] at (m.north) {}; Location state variable (ER) /tikz/sv location 0 \node[entity, draw=gray!60] (m) {}; \node[sv location] at (m.north) {}; Unit of information (PD, ER) /tikz/ui pre:label \node[entity, draw=gray!60] (m) {}; \node[ui] at (m.north) {pre:label}; In PD, stoichiometry can be drawn using a unit of information along an arc: \draw[production] (0,0) -- node[ui, above, N:5 \rightarrow pos=0.5] {N:5} (2cm,0); Unit of information simple chemical (AF) /tikz/ui simple chemical LABEL \node[biological activity, draw=gray!60] (m) {}; \node[ui simple chemical] at (m.north) {LABEL}; Unit of information nucleic acid feature (AF) /tikz/ui nucleic acid feature LABEL \node[biological activity, draw=gray!60] (m) {}; \node[ui nucleic acid feature] at (m.north)

Unit of information macromolecule (AF) /tikz/ui macromolecule LABEL \node[biological activity, draw=gray!60] (m) {}; \node[ui macromolecule] at (m.north) {LABEL}; Unit of information perturbation (AF) /tikz/ui perturbation LABEL (\node[biological activity, draw=gray!60] (m) {}; \node[ui perturbation] at (m.north) {LABEL}; Unit of information complex (AF) /tikz/ui complex LABEL \node[biological activity, draw=gray!60] (m) {}; \node[ui complex] at (m.north) {LABEL}; Unspecidied entity subunit (PD) /tikz/unspecified entity subunit \node[unspecified entity subunit] {LABEL}; LABEL Macromolecule subunit (PD) /tikz/macromolecule subunit **LABEL** \node[macromolecule subunit] {LABEL}; Macromolecule multimer subunit (PD) /tikz/macromolecule multimer subunit LABEL \node[macromolecule multimer subunit] {LABEL}; Nucleic acid feature subunit (PD) /tikz/nucleic acid feature subunit \node[nucleic acid feature subunit] {LABEL}; LABEL Nucleic acid feature multimer subunit (PD) /tikz/nucleic acid feature

multimer subunit





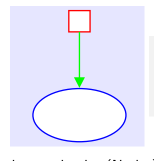
3 Customizing and drawing maps effectively

3.1 Useful options for nodes and arcs

The style of nodes and arcs (and their attributes) can be customized at will using the numerous options offered by TikZ. Following are a few options that might be useful for customizing SBGN maps.

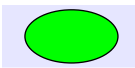
Foreground color (Nodes, Arcs)

/tikz/draw



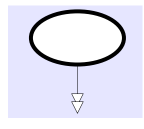
Background color (Nodes)

/tikz/fill



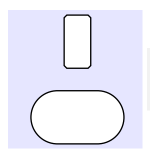
Line width (Nodes, Arcs)

/tikz/line width



Minimum width (Nodes)

/tikz/minimum width



Minimum height (Nodes)

/tikz/minimum height

```
\node[complex, minimum height = 20pt] (m) {};
\node[simple chemical, minimum height = 70pt] at

(0cm,-2cm) (m) {};
```

3.2 Positioning of nodes, arcs, and their attributes

TikZ offers two ways of positioning nodes: with absolute coordinates and relatively to other nodes. We discourage using absolute coordinates as those cannot be changed easily through the drawing process. Drawing maps is far easier using relative positioning. For example, state variables can be placed using the border anchor of the entity pool node they decorate:

```
\node[macromolecule] at (0,0) (m) {}; \node[sv] at (m.140) {val@var};
```

Here, we decided to place the state variable at an angle of 140° on the border of the entity pool node. Now, if we want to change the position of the entity pool node, we don't have to change the position of the state variable:

```
\node[macromolecule] at (0,2) (m) {}; \node[sv] at (m.140) {val@var};
```

Nodes can also be easily be placed relatively to other nodes using the *positioning* TikZ library, with the following syntax:

```
\node[<sbgn_node>, <direction> = <distance> of <node_name>] ...;
```

Here, <direction> defines in what direction the node should be place relatively to a previously defined node named <node_name>. This direction can be unitary (left, right, above, below) or composed of two values (vertical direction first, e.g. above left). The node will be placed in that direction at the distance <distance> of the border of node <node_name>. In case of a composed direction, one can define a distance for each sub-direction: <distance1> and <distance2>. If no distance is provided, the node will be placed at a distance of 1.5cm (default in sbgntikz). Following are some usage examples:

```
\node[unspecified entity] (m) {};
\node[generic process, below = of m] {};

\node[unspecified entity] (m) {};
\node[generic process, above left = 0.5cm of m]

→ {};

\node[unspecified entity] (m) {};
\node[generic process, below left = 2cm and 0.5cm

→ of m] {};
```

To make the node closer to the other one, a negative distance can be defined:

3.3 Bended arcs, multi-part arcs

TikZ offers a simple way to bend arcs with the following syntax:

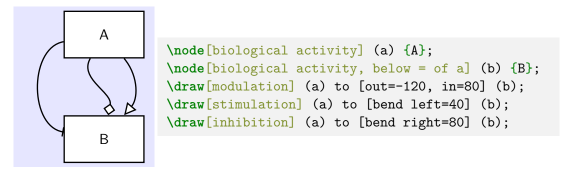
```
\draw (a) to [in=<in_angle>, out=<out_angle>] (b);
```

where <in_angle> specifies the angle at which the arc leaves the source point or node and <out_angle> the angle at which the arc arrives on the target point or node. Both angles are defined relatively to the picture's coordinate.

One can also use the following shortcut:

```
\draw (a) to [bend <direction>=<angle>] (b);
```

where <direction>={left, right} specifies the direction where to bend the arc and <angle> the angle at which the arc leaves the source point or node. The angle is this time defined relatively to the line passing through both points/nodes.



It is often necessary to break arcs into muliple parts for improved readability. TikZ offers very simple operations to break arcs into horizontal and vertical sub-parts, that replace the default -- operation. The |- operation will produce an horizontal sub-part followed by a vertical one, and the -| a vertical sub-part followed by a horizontal one. It can also be convenient to use the --+ and --++ to draw arcs with more than two sub-parts.

```
A \ \node[biological activity] (a) {A}; \ \node[biological activity, below right = 1.5cm and \ \to 0.3 cm of a.center] (b) {B}; \ \draw[stimulation] (a.240) |- (b); \ \draw[inhibition] (a) -| (b); \ \draw[modulation] (b.240) --++ (0,-1) --++ (1,0) \ \to \ --++ (0,1);
```

3.4 Nodes along paths for ER maps

In ER maps, outcomes should be drawn along interaction or assignment arcs. TikZ provides a handy syntax to do so:

Here, the outcome is added along the path using the node command (with no \), as if it was a normal node. The pos option allows defining the distance of the node from the source of the arc relatively to the length of the arc. Here, we placed the outcome at a distance of 70%. More than one outcome can be placed along an arc, by repeating the node syntax:

```
\draw[interaction] (0,0) -- (2,0)

→ node[outcome, pos = 0.3] (o1) {}

→ node[outcome, pos = 0.5] (o2) {}

→ node[outcome, pos = 0.7] (o3) {};
```

One other particularity of drawing ER maps is that one might have to draw arcs targeting other arcs. This is not straightforwardly possible in TikZ, as arcs must target points or nodes. One solution provided by sbgntikz is to draw anchor point coordinates along arcs, as in the following example:

```
\draw[interaction] (0,0) -- (2,0)

\to coordinate[anchor point, pos = 0.5] (a);
\draw[stimulation] (1,1) -- (a);
```

Here, the anchor point coordinate (which is a circle with no drawing, no label and a radius of 2pt) is placed halfway along the interaction arc.

Often in ER maps, one wants to draw horizontal or vertical modulation arcs departing from outcomes. These particular cases, on the other hand, can be easily achieved using TikZ's |- syntax:

```
\draw[interaction] (0,0) -- (2,0);
\draw[stimulation] (1,1) -- (1,1 |- 2,0);
```

Additional space between the tip of the modulation arc and the target arc can be added (here 2pt, that is the anchor point's default radius):

```
\draw[interaction] (0,0) -- (2,0);
\draw[inhibition] (1,1) -- ($(1,1 |-

\to 2,0)+(0,2pt)$);
```

Drawing vertical or horizontal modulation arcs can also be achieved using anchor points and TikZ's let syntax, which allows to place the anchor point at the X-coordinate (for a vertical arc) or Y-coordinate (for an horizontal arc) of the outcome. Following is an example:

```
\draw[interaction] (0,0) -- (2,0) node[outcome,

pos = 0.7] (o1) {};
\draw[interaction] let \p1=(o1) in (0,-2) --

(2,-2) coordinate[anchor point] (a) at

(\x1,-2) node[outcome, pos = 0.3] (o2) {};
\draw[stimulation] (o1) -- (a);
```

In the second interaction, we define the point p1 as being the outcome of the second interaction, and we use its X-coordinate defined as x1 to place the anchor point. More coordinates can be defined and accessed using the let command, always using the syntax pi and xi,

```
\draw[interaction] (0,0) -- (2,0) node[outcome,

pos = 0.7] (o1) {};

\draw[interaction] (-0.7,-0.5) -- (1.3,-0.5)

node[outcome, pos = 0.7] (o2) {};

\draw[interaction] let \p1=(o1), \p2=(o2) in

(0,-2) -- (2,-2) coordinate[anchor point] (a1)

at (\x1,-2) coordinate[anchor point] (a2) at

(\x2,-2);

\draw[stimulation] (o1) -- (a1);

\draw[absolute inhibition] (o2) -- (a2);
```

References

- [1] Till Tantau. The TikZ and PGF Packages.
- [2] Nicolas Le Novere, Michael Hucka, Huaiyu Mi, Stuart Moodie, Falk Schreiber, Anatoly Sorokin, Emek Demir, Katja Wegner, Mirit I Aladjem, Sarala M Wimalaratne, et al. The systems biology graphical notation. *Nature biotechnology*, 27(8):735, 2009.

Index

absolute inhibition, 10	nucleic acid feature subunit, 12
absolute stimulation, 10	omitted process, 8
anchor point, 10	or, 8
and, 8	outcome, 7
assignment, 10	perturbation, 6
association, 8	phenotype, 8
background color, 14	production, 9
biological activity, 6	reversible, 9
clone, 8	simple chemical, 6
compartment, 7	simple chemical multimer, 6, 13
complex, 6	simple chemical subunit, 13
complex multimer, 6	state variable, 11
complex multimer subunit, 13	stimulation, 10
complex subunit, 13	submap, 9
connectors, 7	subunits, 13
consumption, 9	tag, 9
delay, 9	unit of information, 11
dissociation, 8	unit of information complex, 12
emptyset, 6	unit of information macromolecule, 12
entity, 7	unit of information nucleic acid feature,
equivalence arc, 9	11
existence state variable, 11	unit of information perturbation, 12
foreground color, 14	unit of information simple chemical, 11
generic process, 8	unknown process, 8
inhibition, 10	unspecidied entity subunit, 12
interaction, 10	unspecified entity, 5
line width, 14	value, 7
location state variable, 11	
logic arc, 9	bended arc, 16
macromolecule, 5	1.1
macromolecule multimer, 5	multi-part arc, 17
macromolecule multimer subunit, 12	
macromolecule subunit, 12	
minimum height, 14	
minimum width, 14	
modulation, 9	
necessary stimulation, 10	
not, 8	
nucleic acid feature, 5	
nucleic acid feature, 5 nucleic acid feature multimer, 5	
nucleic acid feature multimer, 3 nucleic acid feature multimer subunit, 12	
nucleic acid feature multimer subumit, 12	