MCell3 Quick Tutorial and Reference Guide

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In this document, the main text is in a serif font. Command-line entries, MDL file commands, and code is in a fixed-width font. Values that must be supplied by the user are in an *italicized sans-serif* font.

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1 Running MCell3

MCell3 runs on the command line. The format is

mcell3 options filename

By default, MCell3 sends informational messages, such as simulation progress, to stdout (which will normally appear on the screen); error messages are sent to stderr (which will also normally appear on the screen). Results of simulations are written to files and do not appear as MCell3 is running.

A brief summary of MCell3 optional command-line arguments is given below.

Argument	Explanation
-seed N	Start with random number seed <i>N</i> instead of 1 (the default).
-iterations N	Run the simulation with <i>N</i> timesteps (overrides the value in the mdl file)
-help	Print out a basic help screen.
-logfile <i>filename</i>	Send messages to filename instead of stdout/stderr
-errfile <i>filename</i>	Send error messages to filename instead of stderr
-logfreq N	Print out a message when every <i>N</i> iterations have finished.
-checkpoint_infile filename	Use filename as a checkpoint file for the current simulation (overrides
	any value in the MDL file).

2 Model Description Language overview

MCell3 runs simulations that are specified in *model description language* (MDL) format. These files typically have the extension .mdl, but are not required to. A MDL file is a text file with commands separated by whitespace. The nature and type of whitespace (space, tab, newline) is unimportant to MCell3. You are thus free to use whitespace to clarify the contents of the MDL file.

2.1 The structure of an MDL file

Commands fall into five general groups, which usually should be given in the order presented below. Although this is not always required, there are some commands (e.g. defining a molecule) that must be used before others (e.g.

defining a reaction that uses that molecule). The order below should always be safe:

- 1. Initialization. These commands set global parameters such as the time-step, spatial partitioning, and duration of the simulation.
- 2. Molecule definitions. These commands specify the names and diffusion constants of molecules in the simulation.
- 3. Reaction definitions. These commands specify the reactions that can occur between molecules and the rate at which those reactions occur.
- 4. Geometry specification. These commands describe the membranes and other boundaries within which the simulation occurs, plus where in the world to place molecules initially.
- 5. Output specification. These commands specify what data should be output as the simulation is running; this can include graphical snapshots of the simulation in progress, as well as lists of numbers of molecules or reactions as a function of time.

In addition, there are utility commands-defining variables and including other MDL files-that can appear nearly anywhere.

2.2 How to use this document

This document gives a brief description of every valid MCell3 command. Commands can be specified one after another; it is often convenient to put commands on separate lines but this is not necessary.

Some commands have a scope, delimited by { and } (braces). Within these braces, a different set of commands become available. In this document, each set of commands is given a different title. For example, commands given within a DEFINE_MOLECULE block receive the title Define Molecule Commands.

3 MDL commands

3.1 Initialization commands

The following initialization commands are required in every MDL file.

Command	Explanation
$TIME_STEP = t$	Set the simulation time step to t seconds. 1e-6 is a common value.
	Later commands can change the time steps taken by individual
	molecules, but this time step is still used by all output statements.
ITERATIONS = N	Run the simulation for <i>N</i> iterations.

The following initialization commands are optional.

Command	Explanation
$TIME_STEP_MAX = t$	MCell3 will move longer than the specified simulation time step if it
	seems safe. This command makes sure that the longest possible time
	step is no longer than t seconds, even if MCell3 thinks a longer step
	would be safe. The default is no limit.
SPACE_STEP = N	Have all diffusing molecules take time steps of different duration,
	chosen so that the mean diffusion distance is N microns for each
	molecule. By default, all molecules move the same time step.
CHECKPOINT_INFILE = "filename"	Start the simulation using the conditions specified in the checkpoint file
	<i>filename</i> . This will start at the time that the saved simulation left off,
	and will use molecules stored in the specified file instead of surface
	molecule densities/numbers specified in the MDL file. Release sites can
	add new molecules if the release time is after the time the simulation
	starts.
CHECKPOINT_OUTFILE = "filename"	Save the state of the simulation when CHECKPOINT_ITERATIONS
	(described below) is reached, and stop.
CHECKPOINT_ITERATIONS = N	Used with CHECKPOINT_OUTFILE. This specifies how many iterations to
	run before stopping and writing the checkpoint file. If <i>N</i> is less than
	ITERATIONS, the simulation will terminate normally instead.
SURFACE_GRID_DENSITY = N	Tile all surfaces so that they can hold molecules at N different positions
	per square micron. The default is 10000. For backwards compatability,
	EFFECTOR_GRID_DENSITY works also.
INTERACTION_RADIUS = N	Diffusing molecules will interact with each other when they get within
	<i>N</i> microns of each other. The default is $1/\sqrt{\pi \cdot \sigma_s}$ where σ_s is the
	surface density (default or user-specified).
PARTITION_ $D = [list]$	Subdivide the <i>D</i> 'th axis of space, where <i>D</i> is X, Y, or Z, at the boundaries
	given in <i>list</i> (in microns). In future versions, MCell3 will further
	subdivide space if it is computationally advantageous. By default, each
	axis will be split into between five and fifteen equal partitions. If you do
	not explicitly partition all three axes, MCell3 is likely to ignore your
	request and perform automatic partitioning. The spacing between
	adjacent partitions must be larger than the INTERACTION_RADIUS.
RADIAL_DIRECTIONS = N	Specifies how many different directions to put in the lookup table. The
	default is sensible. Don't use this unless you know what you're doing.
	Instead of a number, you can specify FULLY_RANDOM to generate the
	directions directly from double precision numbers (but this is slower).
RADIAL_SUBDIVISIONS = N	Specifies how many distances to put in the diffusion lookup table.
	Again, the default is sensible. FULLY_RANDOM is not implemented here.

Command	Explanation
ACCURATE_3D_REACTIONS = boolean	Specifies which method to use for computing 3D molecule-molecule interactions. If <i>boolean</i> is TRUE, then molecules will look through partition boundaries for potential interacting partners—this is slower but more accurate. If <i>boolean</i> is FALSE, then molecule interaction disks will be clipped at partition boundaries and probabilities adjusted to get the correct rate—this is faster but can be less accurate. The default is TRUE.
CENTER_MOLECULES_ON_GRID = boolean	If boolean is set to TRUE, then all molecules on a surface will be located exactly at the center of their grid element. If FALSE, the molecules will be randomly located when placed, and reactions will take place at the location of the target (or the site of impact in the case of 3D molecule/surface reactions). The default is FALSE.
VACANCY_SEARCH_DISTANCE = r	Normally, a reaction will not proceed on a surface unless there is room to place all products on the single grid element where the reaction is initiated. By increasing <i>r</i> from its default value of 0, one can specify how far from the reaction's location, in microns, the reaction can place its products. To be useful, <i>r</i> must be larger than the longest axis of the grid element on the triangle in question. The reaction will then proceed if there is room to place its products within a radius <i>r</i> , and will place those products as close as possible to the place where the reaction occurs (deterministically, so small-scale directional bias is possible).
MICROSCOPIC_REVERSIBILITY = value	If <i>value</i> is set to OFF, then binding-unbinding reactions between molecules will be somewhat more efficient but may not be accurate if the probability of binding is high (close to 1). If ON, a more computationally demanding routine will be used to make sure binding-unbinding is more similar in both directions. If <i>value</i> is set to SURFACE_ONLY or VOLUME_ONLY, the more accurate routines will be used only for reactions at surfaces or only for those in the volume. OFF is the default.
NOTIFICATIONS { notification commands }	This block of commands lets you set the informational messages that MCell3 generates. The block can appear multiple times and applies to all MDL below it in the file. It can appear anywhere at the top level (but not inside other blocks).
WARNINGS { warning policy commands }	This block of commands lets you control how MCell3 handles warnings—whether it generates a warning and continues, silently handles the condition, or generates an error and quits. The block can appear multiple times and applies to all MDL below it in the file. It can appear anywhere at the top level (but not inside other blocks).

The following commands can be given in a notifications block; in each case, setting the notification policy to OFF will prevent any informational output regarding that aspect of the simulation. This will not affect warnings.

Notification Command	Explanation
BOX_TRIANGULATION_REPORT = policy	If policy is ON, MCell3 will report how many triangles are generated
	from each box object. Default is OFF.
DIFFUSION_CONSTANT_REPORT = policy	If policy is ON, MCell3 will report four measures of the diffusion
	constant for each molecule. If <i>policy</i> is BRIEF, MCell3 will report just
	one measure (average diffusion distance per step) for each molecule.
	Default is BRIEF.
FILE_OUTPUT_REPORT = policy	If policy is ON, MCell3 will report every time reaction data is written to
	disk. Default is OFF.
FINAL_SUMMARY = policy	If policy is ON, MCell3 will give some information about the CPU time
	used and some of the internal events. Default is ON.

Notification Command	Explanation
ITERATION_REPORT = policy	If policy is ON, MCell3 will provide a running report of how many
	iterations have completed, chosen based on the total number of
	iterations. If <i>policy</i> is an integer value, MCell3 will report each time that
	number of iterations have elapsed. Default is ON.
PARTITON_LOCATION_REPORT = policy	If <i>policy</i> is ON, MCell3 will print out the locations of the partitions used
	for the simulation. Default is OFF.
PROBABILITY_REPORT = policy	If <i>policy</i> is ON, MCell3 will print out the reaction probabilities for each
	reaction (except special internal surface reactions such as absorptive
	surfaces). Default is ON. This will reset the reporting threshold to a
	probability of zero.
PROBABILITY_REPORT_THRESHOLD = p	MCell3 will print out the probabilities for every reaction with
	probability greater than or equal to <i>p</i> . This will override the policy for
	probability reports.
VARYING_PROBABILITY_REPORT = policy	If <i>policy</i> is ON, MCell3 will print out the reaction probabilities when a
	time-varying reaction updates its reaction rate (regardless of the old or
	new probability). Default is ON.
PROGRESS_REPORT = policy	If policy is ON, MCell3 will print out messages indicating which part of
	the simulation process is underway (initializing, running, etc.). Default
	is ON.
RELEASE_EVENT_REPORT = policy	If policy is ON, MCell3 will print out a message every time molecules
	are released through a release site (indicating how many molecules of
	which type were released and the iteration on which they were
	released). Default is ON.
ALL_NOTIFICATIONS = policy	Set all notification policies to the same value (ON or OFF). This overrides
	the existing probability report threshold, if there is one.

The following commands can be given in a warnings block. Setting the warning policy to IGNORED will prevent any output and the condition will be handled as best it can. WARNING will give a warning message, but the problem will be handled and the simulation will continue. Setting to ERROR will generate an error and the simulation will stop. This will not affect notification policies.

Warning Policy Command	Explanation
DEGENERATE_POLYGONS = policy	Degenerate polygons are polygons with zero area and must be removed
	for the simulation to run. The default policy is WARNING.
HIGH_REACTION_PROBABILITY = policy	Generate warnings or errors if reaction probabilities exceed a certain
	threshold. The default policy is IGNORED. The warnings or errors will be
	generated both at parse time and during runtime if there are time
	varying reaction rates that exceed the threshold.
$HIGH_PROBABILITY_THRESHOLD = p$	If the policy is to generate warnings or errors on high probability
	reactions, have them generated when the probability equals or exceeds
	p. The default value is $1.0 + 10^{-12}$.
LIFETIME_TOO_SHORT = policy	Generate warnings if molecules have short lifetimes (which could affect
	the accuracy of the simulation). This warning occurs after the
	simulation has ended, so ERROR. is not a valid option. The default policy
	is WARNING.
LIFETIME_THRESHOLD = n	If the policy is to generate a warning if molecules have short lifetimes,
	then generate warnings on molecules that have an average lifetime of
	less than <i>n</i> iterations. The default value is 50.
MISSED_REACTIONS = policy	Generate errors or warnings if there are missed reactions (which usually
	is a consequence of an overly high reaction probability). This warning
	occurs after the simulation has ended, so ERROR. is not a valid option.
	The default policy is WARNING.

Warning Policy Command	Explanation
$MISSED_REACTION_THRESHOLD = f$	If the policy is to generate a warning if there are missed reactions, then
	generate a warning for each reaction where a fraction of at least f of
	reactions were missed. The default value is 10^{-3} .
NEGATIVE_DIFFUSION_CONSTANT = policy	Diffusion constants cannot be negative, and will be set to zero if they
	are. The default policy is WARNING.
MISSING_SURFACE_ORIENTATION = policy	Generate errors or warnings if a molecule is placed on a surface or
	reactions occur at a surface without a specified orientation—the code
	will assume you mean that there is no orientation in the warning or
	silent cases. To avoid triggering this condition, if you want to have no
	orientation, you must specify it explicitly with ', or,' or; The
	default policy is ERROR.
NEGATIVE_REACTION_RATE = policy	Reaction rate constants cannot be negative, and will be set to zero if
	they are. The default policy is WARNING.
USELESS_VOLUME_ORIENTATION = policy	Generate errors or warnings if a molecule is placed in a volume or
	reactions occur in free space but an orientation is specified
	anyway—there is no way to impose orientation so the marks will be
	ignored. The default policy is WARNING.
ALL_WARNINGS = policy	Set all warning policies to the same value (IGNORED, WARNING or
	ERROR). If ERROR is not a valid choice, the policy will be set to WARNING
	instead.

3.2 Molecule definition commands

All molecules must be defined by name in a DEFINE_MOLECULES block. For users of MCell 2, note that there is no longer a distinction between a receptor and a ligand. Everything is a molecule, and every different bound state of a receptor must have a unique name (since it must be a unique molecule). The names must be unique in the entire simulation (that is, unique within their own MDL file and any included MDL files that make up the whole simulation).

A define molecule block can be one of the following:

Command	Explanation
DEFINE_MOLECULE name	Define a single molecule called <i>name</i> . The molecule's properties
{	are specified by commands inside braces.
define molecule commands	
}	
DEFINE_MOLECULES	Define a series of molecules by name. Each molecule's properties
{	are specified by commands inside braces.
nameA { define molecule commands }	
nameB { define molecule commands }	
}	

Each molecule must have a diffusion constant set using one of the following commands:

Define Molecule Command	Explanation
DIFFUSION_CONSTANT = D	This molecule diffuses in space with diffusion constant <i>D. D</i> can be
	zero, in which case the molecule doesn't move. Synonyms for this
	command are DIFFUSION_CONSTANT_3D and D_3D. The units of D are cm^2/s .
DIFFUSION_CONSTANT_2D = D	This molecule is constrained to a surface and diffuses with diffusion
	constant D. D_2D is a synonym for this command.

The following optional commands can be applied to each molecule (and must appear in this order, and after the diffusion constant is set):

Define Molecule Command	Explanation
CUSTOM_TIME_STEP = t	This molecule should take timesteps of length <i>t</i> (in seconds). Use either
	this or CUSTOM_SPACE_STEP, not both.
CUSTOM_SPACE_STEP = L	This molecule should take steps of average length L (in microns). If you
	use this directive, do not set CUSTOM_TIME_STEP.
TARGET_ONLY	This molecule will not initiate reactions when it runs into other
	molecules. This setting can speed up simulations when applied to a
	molecule at high concentrations that reacts with a molecule at low
	concentrations (it is more efficient for the low-concentration molecule
	to trigger the reactions). This directive does not affect unimolecular
	reactions.

3.3 Reaction definition commands

All reactions must be defined inside a reaction definition block:

Command	Explanation
DEFINE_REACTIONS	Define a series of reactions inside braces.
{	
reaction commands	
}	

Reactions are specified using arrow notation:

Reaction Command	Explanation
reactants -> products [rate]	Define a reaction that occurs between one or two reactants (names of
	molecules, separated by +) and produces an arbitrary number of <i>prod-</i>
	ucts (also separated by +), with a specified rate. If a molecule is in the
	reactants list and not in the products list, it is destroyed in the reaction.
	The rate can also be a filename, in quotes, that contains two columns:
	the second is the rate, while the first is the time at which that rate should
	start being used. This allows variable reaction rates. If you do not want
	products, use the NULL keyword as a placeholder.
reactants -> products [rate]: name	As above, and call the reaction <i>name</i> so it can be referred to by count
	statements.

This notation is perhaps best explained through examples. In the most basic form, reactants and products are just the names of molecules, separated by +:

Example	Explanation
A -> B [100]	Molecule A changes into molecule B at a rate of 100s^{-1} .
A -> A + B [100]	Molecule A emits molecules of B at a rate of $100 \mathrm{s}^{-1}$.
A -> NULL [100]	Molecule A is destroyed at a rate of $100 \mathrm{s}^{-1}$.
A + B -> A [1e6]	Molecule A destroys molecule B at a rate of $10^6 \text{M}^{-1} \cdot \text{s}^{-1}$.
A + B -> A + C [1e6]	Molecule A catalytically converts B to C at a rate of $10^6 M^{-1} \cdot s^{-1}$.
A+B -> A+B+C [1e6]	Collision of A and B catalytically generates C at a rate of $10^6 M^{-1} \cdot s^{-1}$.

Reactions can take place on surfaces or involve molecules contained therein (surface molecules). Surfaces possess a front and a back side defined by the direction of the surface normal which points from the back toward the front. Surface molecules have an orientation in the form of a top and a bottom domain and are positioned on surfaces with their top domain either on the surfaces' front or back side, or top-front and top-back for short.

Reactions that explicitly involve surfaces are said to occur with an absolute orientation regarding the surface. When reactions involving surface molecules take place in the absence of explicit surfaces they are said to occur without an absolute orientation. Below, we will illustrate both cases.

3.3.1 Reactions without absolute orientation

For reactions without an absolute orientation, the reaction specification lists the required relative orientation of the reactants and products. This allows one to write general reactions that do not depend on the way in which molecules are inserted into surfaces, i.e., either top-front or top-back.

The two possible orientations are specified by ' and , (apostrophe and comma) after the molecule's name. Hence, a surface-bound molecule B can have the orientations B' and B,. The table below provides a few example reactions

Example	Explanation
B' -> B, [10]	Molecule $\[Beta]$ flips (changes its orientation) at a rate of $10\[seps]$ s ⁻¹ .
B' -> B' + A' + C, [10]	Molecule B emits molecules of A on the side it's pointing to and emits C
	on the other side, at a rate of $10s^{-1}$
B, -> B, + A, + C' [10]	This specifies exactly the same reaction as above. B and A end up with
	the same orientation, while $\mathbb C$ has opposite orientation.

The best way to keep the relationships straight is to draw a "before" picture with each reactant facing the direction of the tick mark, and an "after" picture with each product facing in the direction of the tick mark. Clearly, inverting this picture by flipping all tick marks results in the same reaction. One can thus use tick marks that are consistent with ones mental picture.

Below are additional reaction examples involving a molecule A diffusing in 3D and surface molecules B and C:

Example	Explanation
A' + B' -> C' [1e5]	Molecule A binds to B if it is on the side that B is pointing to, producing
	a C facing the same way as B, at a rate of $10^5 \mathrm{M}^{-1}\cdot\mathrm{s}^{-1}$.
A, + B, -> C, [1e5]	The same reaction again—everything occurs on the same side, but we
	wrote it on the bottom this time.
A' + B, -> C' [1e5]	Molecule A binds when it hits the opposite side of B, producing a C facing
	the opposite way as B (i.e. towards the side A came from), at a rate of
	$10^5 \mathrm{M}^{-1}\cdot\mathrm{s}^{-1}$.
A, + B' -> C, [1e5]	Same as above.

So far, all examples have used the first orientation class, specified with ' and , . The second orientation class is specified by '' and , , . The third is ''' and , , , and so on. Molecules in different orientation classes do not pay attention to each other's orientation. In a reaction with orientation, every molecule must be explicitly given an orientation class otherwise an error is generated. This behavior can be adjusted to generate warnings or no messages instead; in this case, molecules without an orientation class act without regard to orientation. Orientation classes are a fundamentally new concept introduced in MCell3. They replace the MCell2 idea of POSITIVE_POLE, NEGATIVE_POLE, and BOTH_POLES specifications for receptors. Several examples follow:

Example	Explanation
A'' + B, -> C' [1e5]	Molecule A binds to either side of B (since they are in different orientation
	classes); this produces a C facing the opposite way as B, at a rate of
	$10^5 \mathrm{M}^{-1}\cdot\mathrm{s}^{-1}$.
A,, + B, -> C' [1e5]	This is the same reaction—since A is the only molecule in the second
	orientation class, it doesn't matter which way we specify things.
A,, + B' -> C, [1e5]	Same again—B and C still have opposite orientations.
A, + B' -> C,, [1e5]	Molecule A hits the opposite side of B and produces C that is equally
	likely to point either way, at a rate of $10^5 \mathrm{M}^{-1}\cdot\mathrm{s}^{-1}$.
A, + B' -> C'' [1e5]	Same as above, since C is still not in the same orientation class as the
	others.
A'+B'' -> A,+B''' [1e5]	Molecule A hits molecule B on either side; A keeps traveling (goes to the
	other side) and B tumbles to a random orientation, at a rate of $10^5\mathrm{M}^{-1}$ ·
	s^{-1} .
A'+B''-> C'''+D'''' [1e5]	A and B react in any orientation and produce C and D in random orien-
	tations. All orientation classes are different, so there are no geometrical
	constraints here.

There are more examples of how one would use this syntax to model well-known biological reactions at the end of this document in section 5.

3.3.2 Reactions with absolute orientation

Reactions can specify an absolute orientation with respect to the surface on which they take place via including a surface class specification in the reaction definition. The general form for defining reactions with absolute orientations is accomplished via the "@" character as shown below

Reaction Command	Explanation
reactants @ surf_class_name -> products [rate]	Define a reaction that occurs between one or two oriented <i>reac-</i>
	tants (names of molecules, separated by +) on a set of surface
	regions identified by <i>surf_class_name</i> . The reaction produces
	an arbitrary number of oriented <i>products</i> (also separated by +),
	with a specified <i>rate</i> . If a molecule is in the <i>reactants</i> list and
	not in the <i>products</i> list, it is destroyed in the reaction. The rate
	can also be a filename, in quotes, that contains two columns:
	the second is the rate, while the first is the time at which that
	rate should start being used. This allows variable reaction rates.
	If you do not want products, use the NULL keyword as a place-
	holder.
reactants @ surf_class_name -> products [rate]: name	As above, and call the reaction <i>name</i> so it can be referred to by
	count statements.

A reaction defined in this way takes place on all surface regions which specify SURFACE_CLASS = surf_class_name. The relative orientation of reactants and products is specified as explained in 3.3.1 but now the reaction takes place with respect to the orientation given for surf_class_name indicating the front or back of the selected surface regions. Please note that all reactants have to be listed to the left of surf_class_name and no surface class specifications can occur on the product side of the reaction definition. Furthermore, for bi-molecular reactions at least one of the two reactants has to be a surface molecule.

The table below lists several examples of oriented reactions involving a surface class SURF, a 3D molecule A, and surface molecules B and C.

Example	Explanation
A' + B' @ SURF' -> C, [1e5]	The reaction affects surface molecules B located on surface regions iden-
	tified by surface class SURF which have their top domain at the front
	of the surface. B reacts with A approaching from the front at a rate of
	$10^5 \mathrm{M}^{-1}\cdot\mathrm{s}^{-1}$ to yield surface molecule C whose orientation is flipped
	with respect to B, i.e., C has its top domain aligned to the back of the
	surface regions.
A' + B, @ SURF' -> C, [1e5]	Same as above, but B now has its top domain at the back of the surface
	and reaction product C assumes the same orientation.
A,, + B, @ SURF' -> C' [1e5]	Since A is in an orientation class different from both B and SURF, A can
	react from both sides. B has its top domain at the back of the surface and
	the reaction product C has its orientation flipped, i.e., its top domain is at
	the front of the surface.
A' + B' @ SURF' -> C,, [1e5]	Same as the first reaction, but since product C is in a orientation class
	different from either A, B, and SURF, its orientation is random with respect
	to the surface regions, i.e., its top domain can be either on the front or
	back.

Tick marks add, so that ', and ,' mean no orientation. Reactions will occur from either orientation when given reactants with no orientation, and products will orient randomly. A semicolon, ;, can be used instead of two opposite tick marks. Orientations can also be specified numerically inside $\{\}$ after the molecule name. For example, $A\{1\}$ and $A\{-1\}$ are synonyms for A' and A, and $A\{0\}$ is a synonym for A;

There are several variants of the normal reaction arrow ->. One can use an arbitrary number of dashes in the arrow, i.e., ->, -->, and -----> all mean the same thing. In addition, the following arrows have different meanings:

Reaction Arrow	Explanation
->	A unidirectional reaction going from reactants (on the left) to products
	(on the right).
<->	A bidirectional reaction going in either direction; at most two molecule
	names can appear on each side. A rate must be given for each direction
	using the notation $[>k_+, < k]$, where k_+ is the forward rate constant
	and k_{-} is the backward rate constant.
reactant catalyst -> products	This specifies a catalytic reaction where <i>reactant</i> is converted to <i>products</i>
	in the presence of <i>catalyst</i> . This is the same as the reaction <i>catalyst</i> +
	reactant -> catalyst + products. Presently, there can only be one reactant.
reactant <- catalyst -> product	A bidirectional catalytic reaction. There can only be one reactant and
	one product.

Finally, a few special cases deserve particular mention

• For catalytic reactions, if a catalyst is a surface class, the latter is not copied to the list of products, i.e.:

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A' -- SURF' -> C, [rate] is equivalent to A' @ SURF' -> C, [rate]
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- Reversible reactions of the form A' @ SURF' <--> C, [>rate1, <rate2] or
 - A' <-- SURF'--> C, [>rate1, <rate2] are equivalent to the following two reactions:
 - (i) A' @ SURF' \rightarrow C, [rate1]
 - (ii) C, @ SURF' -> A' [rate2]

3.4 Geometry definition commands

3.4.1 Surface properties

MCell3 allows the user to specify properties of the surfaces of objects. For example, one may wish to specify that a surface does not block the diffusion of molecules. Each type of surface is defined by name, and each surface name must be unique in the simulation and should not match any molecule names. Surface properties are specified inside a surface definition block:

Command	Explanation
DEFINE_SURFACE_CLASS name	Define a single surface type called <i>name</i> . The properties are specified by
{	zero or more commands inside braces.
surface property commands	
}	
DEFINE_SURFACE_CLASSES	Define a series of surface types by name.
{	
nameA { surface property commands }	
nameB { surface property commands }	
}	

To define surface properties, use the following commands:

Surface Property Command	Explanation
REFLECTIVE = name	The molecule called <i>name</i> is reflected by this surface. This is the de-
	fault behavior for volume molecules; it prevents surface molecules from
	crossing triangle boundaries. Tick marks on the name allow selective
	passage of molecules in one orientation relative to the surface.
TRANSPARENT = name	The molecule called <i>name</i> passes through this surface. This is only
	meaningful for volume molecules; surface molecules assume that their
	surface can be traveled in unless it is labeled REFLECTIVE. Tick marks
	allow the creation of one-way transparent surfaces.
ABSORPTIVE = name	The molecule called <i>name</i> is destroyed if it touches this surface. Tick
	marks allow destruction from only one side for volume molecules, or
	destruction of only one orientation of surface molecules.
CLAMP_CONCENTRATION name = value	The molecule called <i>name</i> is destroyed if it touches the surface (as if it
	had passed through), and new molecules are created at the surface, as
	if molecules had passed through from the other side at a concentration
	<i>value</i> (units = M). Orientation marks may be used; in this case, the other
	side of the surface is reflective. Note that this command is only used to
	set the effective concentration of a volume molecule at a surface; it is not
	valid to specify a surface molecule. This command can be abbreviated
	as CLAMP_CONC.
MOLECULE_DENSITY	Add the named molecules at the specified densities D1, D2,, (units =
{	μ m ⁻²) to every surface with this surface class. Use orientation marks
name1 = D1	after the name to specify the direction relative to the surface normal. For
name2 = D2	example, A' specifies a molecule in the same orientation as the surface,
	while A, specifies the opposite orientation. Using both marks indicates
}	that the molecule should be assigned an orientation randomly.

Surface Property Command	Explanation
MOLECULE_NUMBER	Add the exact numbers N1, N2,, of molecules onto any region that is
{	made out of this surface class. Note: this usage is not recommended; it
name1 = N1	is better to add exact numbers of molecules to the region. Orientation
name2 = N2	marks after the name must be used to specify the direction the molecules
	are facing.
}	

Note that surface normals are defined by the right-hand rule applied to the vertices in order as listed (see section 3.4.2). Box objects are converted internally into triangles and the surface normals point outwards.

3.4.2 Geometrical objects

Two types of geometrical objects are supported in MCell3. Objects should not have coincident surfaces unless neither surface contains a molecule that any moving molecule can react with. Also, all coincident surfaces should agree on whether they are transparent, reflective, or absorptive to each molecule that might strike them. Geometrical objects can be defined using:

Command	Explanation
name BOX { box commands region commands transformation commands }	This defines a box object called <i>name</i> . The shape and position of the box is defined by . Optionally, additional commands can create regions and perform geometrical transformations on the box. Internally, a box is represented as a set of triangles.
name POLYGON_LIST { polygon commands region commands transformation commands }	This defines a polygon list object called <i>name</i> . Polygon list objects explicitly give their triangular surface elements.

A variety of optional commands can be used inside a geometrical object definition block, after corners or vertex list / element connections are specified, to modify the basic composition of the object and its surface properties. These are described below. Geometrical transformations are described later, in section 3.4.5.

Box Command	Explanation
CORNERS = $[x1, y1, z1], [x2, y2, z2]$	The box object has corners as specified. The first coordinates should be
	less than the second set of coordinates, although MCell3 may fix it if you
	do it incorrectly.
ASPECT_RATIO = a	Make sure that the ratio of the long to short side of each triangle making
	up the box is no more than a. The smallest allowed value is 2. The
	default is to not care about triangle shape.

Polygon Command	Explanation
VERTEX_LIST	Specify the vertices of the triangles inside a polygon list object inside
{	braces. Each vertex is given by its triple $[x, y, z]$. This command must
[x0, y0, z0]	be given before the ELEMENT_CONNECTIONS command.
[x1,y1,z1]	
}	

Polygon Command	Explanation
ELEMENT_CONNECTIONS	Specify the triangles by vertex indices. The vertices are numbered from
{	0 upwards in the order they were given in the vertex list. The direction of
[a0,b0,c0]	the surface normal is determined by the right-hand rule while following
[a1, b1, c1]	the vertices. Each triangle is given by a triple [a, b, c] of vertex num-
	bers. This command must be given after the VERTEX_LIST command.
}	

Region Command	Explanation
DEFINE_SURFACE_REGIONS	Define regions on the object. The extent of a region is given by the el-
{	ement specifier commands (at least one is required). Molecules can be
nameA {	added and surface properties can be set with the optional regional surface
element specifier commands	commands. You can have an arbitrary number of regions on an object,
regional surface commands	and they may overlap if you wish. Molecules added to overlapping re-
}	gions accumulate; surface properties are those of the last region applied.
name2 { }	Every BOX and POLYGON_LIST object has a pre-defined ALL region which
	consists of the entire object and has no special properties.
}	
REMOVE_ELEMENTS	Remove the portion of the object specified by the element specifiers. You
{	can think of this as a special type of region that defines the removed
element specifier commands	portions of the object. No real region exists on any part of the object that
}	has been removed. You can use a list of element numbers/names instead
	of element specifiers if you wish, but you cannot mix a list of element
	numbers/names with the element specifier syntax. It is an error to remove
	all elements in an object or region.

Element Specifier Command	Explanation
INCLUDE_ELEMENTS = [list]	Include the elements specified by number or name. For polygon objects,
	these refer to the triangles defined by the element connections, count-
	ing from zero upwards in the order given. For boxes, the side names
	LEFT, RIGHT, FRONT, BACK, BOTTOM, and TOP can be used to refer to the
	sides, where left/right corresponds to the x axis (left is lower x values),
	front/back to y, and bottom/top to z. ALL_ELEMENTS refers to the entire
	object. Numbers can be specified individually (separated by commas) or
	in ranges with the format N TO M. The two styles can be mixed (separated
	by commas).
EXCLUDE_ELEMENTS = [list]	Exclude the elements listed. If this is the first element specifier, assume
	that all elements not listed are included. If not, subtract from the existing
	list.
INCLUDE_REGION = name	Include the existing region on this object called <i>name</i> into this region,
	too.
EXCLUDE_REGION = name	Exclude the existing region on this object called <i>name</i> from this new
	region.
INCLUDE_PATCH=[$x1, y1, z1$], [$x2, y2, z2$]	This specifier is only valid on box objects, and the corners must define a
	rectangular patch that is on exactly one side of the box. The box will be
	divided into triangles in such a way that this patch consists of separate
	triangles and will form a region.
EXCLUDE_PATCH= $[x1, y1, z1]$, $[x2, y2, z2]$	Exclude the patch from this region.

After element specifiers, regions can specify a surface type and add extra molecules using:

Regional Surface Command	Explanation
SURFACE_CLASS = name	Set the surface type of this region to the previously defined surface class
	called <i>name</i> .
MOLECULE_DENSITY {}	This is the same as the Surface Property Command of the same name.
MOLECULE_NUMBER {}	This is the same as the Surface Property Command of the same name.
	Its usage is recommended here, as a regional surface command, rather
	than as a surface property command, so that the number of molecules
	is specified in the same place as the geometry, thus making the density
	easier to figure out.

3.4.3 Release objects

Release objects place molecules into the world. Release objects provide the only means of placing molecules in a three dimensional space, but some release shapes can place molecules on surfaces as well. Release objects are defined using the following commands:

Command	Explanation
name release_site	Create a release site called <i>name</i> . The shape and method of release is
{	specified by the release site commands. Optionally, geometrical trans-
release site commands	formations can be applied also.
transformation commands	
}	
<pre>name CUBIC_RELEASE_SITE {}</pre>	Create a cubic release site called <i>name</i> . Molecules are released in a box
	as specified by the radius. (This is the same as using the SHAPE=CUBIC
	command inside RELEASE_SITE.)
<pre>name SPHERICAL_RELEASE_SITE {}</pre>	Create a spherical release site called <i>name</i> . Molecules are released uni-
	formly within the sphere depending on the defined radius of the ob-
	ject. (This is the same as using the SHAPE=SPHERICAL command inside
	RELEASE_SITE.)
<pre>name SPHERICAL_SHELL_SITE {}</pre>	Create a spherical shell release site called <i>name</i> . Molecules are dis-
	tributed on a spherical shell at the defined radius of the object. For
	now, you must specify the number to distribute, not a concentration.
	(This is the same as using the SHAPE=SPHERICAL_SHELL command in-
	side RELEASE_SITE.)
DEFINE_RELEASE_PATTERN name	Define a new release pattern according to the commands given. A release
{	pattern must be defined for anything other than release at the beginning
release pattern commands	of the simulation. Release patterns must be defined before they are used.
}	Multiple release sites can use the same pattern.

The following commands define where, what, and when a release object releases molecules:

Release Site Command	Explanation
SHAPE = geometry	Release molecules in the specified shape. Valid shapes are CUBIC, SPHERICAL, SPHERICAL_SHELL, and LIST; or the name of region(s) on
	which to release. Each region must already be instantiated or be inside
	the same OBJECT as the release site (see OBJECT command). Region
	names can be combined with + to indicate release on both regions, -
	to indicate the release occurs on the first and not the second, and * to
	indicate the release occurs only where the two regions overlap. Paren-
	theses may be used for grouping. Volume molecules will be released in
	the volume bounded by the regions (each region must be closed); sur-
	face molecules will be released on the surface (and regions need not be
	closed). If the region name is omitted and only the name of a BOX or
TOGATION []	POLYGON_LIST object is specified, the object's ALL region will be used.
LOCATION = $[x, y, z]$	The release occurs centered at this location. Only used for geometrical shapes.
MOLECULE = name	The named molecule is the one that will be released. Not used for the
MODECOLE - Manie	LIST shape. You must specify an orientation if the molecule is a surface
	molecule.
MOLECULE_POSITIONS	The named molecules are added in the locations given. The molecule
{	names must be followed by orientation marks if they have a 2D diffusion
name1 [x1, y1, z1]	constant. If a molecule has a 2D diffusion constant, it will be placed on
name2 [x2, y2, z2]	the surface closest to the coordinate given. This command is used for the
	LIST shape only.
} SITE_DIAMETER = d	For a geometrical release site, this releases molecules uniformly within a
$SITE_RADIUS = r$	diameter d or a radius r . Not used for releases on regions. With the LIST
	shape, this is the distance that surface molecules search for a surface
	before giving up; free molecules pay no attention to this value for the
	LIST shape.
$SITE_DIAMETER = [x, y, z]$	Release is asymmetric with a different diameters in different directions,
$SITE_RADIUS = [x, y, z]$	as indicated by the vector. Not used for releases on regions or with the LIST shape.
RELEASE_PROBABILITY = p	This release does not occur every time, but rather with probability p . (If
	omitted, the default is to release without fail.) Either the whole release
	occurs or none of it does; the probability does not apply molecule-by-
	molecule. p must be in the interval [0,1].
NUMBER_TO_RELEASE = n	Release n molecules. For releases on regions, n can be negative, and
	the release will then remove molecules of that type from the region. To remove all molecules of a type, just make n large and negative. It is
	unwise to both add and remove molecules on the same timestep—the
	order of addition and removal is not defined in that case. This directive
	is not used for the LIST shape, as every molecule is specified.
CONCENTRATION = c	Release molecules at concentration c molar for volumes and d molecules
DENSITY = d	per square micron for surfaces. Neither can be used for the LIST shape;
	DENSITY is only valid for regions.
GAUSSIAN_RELEASE_NUMBER	Release molecules according to a Gaussian distribution with the specified
{	mean and standard deviation.
MEAN_NUMBER = n	
STANDARD_DEVIATION = s	
	I

Release Site Command	Explanation
RELEASE_PATTERN = name	Use the named release pattern instead of the default. The default is to
	release the specified number of molecules at the beginning of the sim-
	ulation. If <i>name</i> is the name of a reaction pathway, the release event
	will happen every time that reaction happens. The location will then be
	relative to the site of the reaction, and the z-axis will be rotated to align
	with the surface normal if the reaction was at a surface. This is much
	slower than creating products within a reaction, so only use it for special
	cases (e.g. synaptic vesicle release with a random or very large number
	of neurotransmitter molecules).

Release patterns are defined as follows.

Release Pattern Command	Explanation
DELAY = t	The release pattern will start at time <i>t</i> . (Default is to start at time zero.)
RELEASE_INTERVAL = t	During a train of releases, release molecules after every t seconds. De-
	fault is to release only once $(t = \infty)$.
TRAIN_DURATION = t	The train of releases lasts for t seconds before turning off. Default is to
	never turn off $(t = \infty)$.
TRAIN_INTERVAL = t	A new train of releases happens every <i>t</i> seconds. Default is to never have
	a new train $(t = \infty)$. The train interval must not be shorter than the train
	duration.
NUMBER_OF_TRAINS = n	Repeat the release process for <i>n</i> trains of releases. Default is one train.
NUMBER_OF_TRAINS = UNLIMITED	Repeat trains forever.

3.4.4 Instantiation, grouping, and modification of objects

An object is a box, polygon, release site, or a metaobject which contains other objects. Metaobjects are defined and modified using

Command	Explanation
name OBJECT	Define a new object called <i>name</i> . Inside the braces, list other objects one
{	at a time to be added (see below).
object specifier commands	
transformation commands	
}	
INSTANTIATE name OBJECT { }	Same as above, except we also insert the object into the world. A simu-
	lation must have at least one INSTANTIATEd object.
MODIFY_SURFACE_REGIONS	This modifies surface regions on existing objects via their name and re-
{	gion name. Element lists may not be changed, but otherwise all regional
nameA[regA1] {	surface commands are available. The full name must be given in the
regional surface commands	case of separate objects (using name1.name2 to refer to objects inside
}	metaobjects). If an object is included in a metaobject, then has a sur-
nameB[regB1] { }	face region modified, and is included in another metaobject, the surface
	regions will differ in those the two metaobjects.
}	

You can define release sites, boxes, and polygon objects inside another object, as well as placing previously defined objects into existing ones:

Object Specifer Command	Explanation
newname OBJECT oldname	Add the existing object called <i>oldname</i> into the existing object and label
{	it <i>newname</i> . You can add extra commands (e.g. transformation) inside
transformation commands	the braces. The old and new names can be the same thing. Thereafter,
}	this object can be referred to in the world as <i>name</i> . <i>newname</i> .
name BOX {}	Create a box inside the existing object (using the same syntax as previ-
	ously defined).
name POLYGON_LIST {}	Create a polygon list object inside the existing object (using the same
	syntax as previously defined).
name RELEASE_SITE {}	Create a release site inside the existing object.
newname OBJECT {}	Create an object inside the existing object.

3.4.5 Geometrical transformations

At the end of the definition of a release object or geometrical object, or in the block where an object is instantiated, it can be moved using the following transformation commands (placed at the end of the block before the closing brace).

Transformation Command	Explanation
TRANSLATE = $[x, y, z]$	Move the object by the specified vector.
SCALE = [x, y, z]	Scale the object by multiplying each coordinate by the corresponding value in the vector.
DOMARIN [
ROTATE = $[x, y, z]$, A	Rotate A degrees about the axis defined by the supplied vector.

3.5 Output specification commands

There are two forms of output in MCell3, visualization output and count output. Visualization output typically contains the molecules and/or geometry of the model in a form suitable for visualization or analysis that requires knowledge of the precise location of particles. Count output reports running totals of summary statistics such as the total number of molecules of a certain type in the world, the number of times a reaction has occurred inside some object in the world, and so on. Count output can also be written when triggered by a specific event such as a reaction taking place.

3.5.1 Visualization Output

Command	Explanation
VIZ_OUTPUT	Define a new visualization output block. MDL files can have multiple
{	VIZ_OUTPUT blocks.
viz output commands	
}	

Each viz output block consists of the following commands:

Viz Output Command	Explanation
MODE = viz_mode	Specifies the mode of the visualization output. The mode defines the di-
	rectory structure and number of files comprising the visualization output.
	The valid values for are DREAMM_V3, DREAMM_V3_GROUPED, and DX. The
	DX mode requests the old MCell2 style of output format for compatibility
	purposes. The default is DREAMM_V3.
FILENAME = "filename_specifier"	Name of the master header file containing all information for DREAMM
	and references to the multiple binary data files.
MESHES	Defines meshes visualization data output block.
\ {	
data output block	
}	
MOLECULES	Defines molecules visualization data output block.
{	
data output block	
}	

Each data output block consists of the following commands:

Data Output Block Commands	Explantation
NAME_LIST	Defines a valid name list. The valid values are either names separated
{	by any type of whitespace, strings with wildcards (in quotes) that match
name list commands	names, or keywords defined below. All children of the named objects
}	are included by default. If this statement occurs in a MESHES block, the
	names should be names of objects; in a MOLECULES block they should be
	names of molecules.
TIME_POINTS	Defines what data should be output at what times. The data types
{	are given below and valid notations for time_points_list are [time1],
data type @ time_points_list	or [time1, time2,, time_end], or [time1, time2, [time3 TO
}	time_end STEP delta_time]], or ALL_TIMES. Mutually exclusive with
	ITERATION_NUMBERS.

Data Output Block Commands	Explantation
ITERATION_NUMBERS	Defines what data should be output at what iterations. The data
{	types are given below and valid notations for iteration_numbers_list
data type @ iterations_numbers_list	are [iteration1], or [iteration1, iteration2,, iteration_end],
}	or [iteration1, iteration2, [iteration3 TO iteration_end STEP
	delta_iteration]], or ALL_ITERATIONS. Mutually exclusive with
	TIME_POINTS.

The following name list commands for MESHES and MOLECULES are available:

Name list Commands (MESHES)	Explanation
ALL_MESHES	All mesh object names should be included in the NAME_LIST sub-block
	inside a MESHES block. ALL_MESHES is equivalent to naming the top-level
	mesh object (assuming that only a single INSTANTIATE block is present).

Name list Commands (MOLECULES)	Explanation
ALL_MOLECULES	All molecule names should be included in the NAME_LIST sub-block in-
	side MOLECULES block.

The following data type commands for MESHES and MOLECULES are available:

Data types (MESHES)	Explanation
GEOMETRY	Mesh vertex and connectivity information should be written at the spec-
	ified time/iteration.
REGION_DATA	Mesh region information should be written at the specified time/iteration.
ALL_DATA	Equivalent to using both GEOMETRY and REGION_DATA.

Data types (MOLECULES)	Explanation
POSITIONS	Molecule position information should be written at the specified
	time/iteration.
ORIENTATIONS	Molecule orientation information should be written at the specified
	time/iteration.
ALL_DATA	Equivalent to using both POSITIONS and ORIENTATION.

There are two possible visualization output file formats. DREAMM_V3 mode is the default, and creates files in native DX format. This mode is optimized for speed of visualization, but creates many individual files. It has a directory structure with the top-level directory given by adding _viz_data to the filename, i.e. filename_viz_data. For example if FILENAME = "./viz_data/diffusion_box" then the directory diffusion_box_viz_data will be created inside the ./viz_data directory. Inside the filename_viz_data directory there is the data directory called frame_data and three files:

• filename.dx (the header file)

• filename.iteration_numbers.bin

• filename.time_values.bin

The directory frame_data contains a number of sub-directories named by combining the word iteration_ with the iteration number of the simulation, such as iteration_0, iteration_20, etc. Each of these iteration sub-directories by itself contains up to nine files:

- meshes.dx (header file for meshes)
- mesh_positions.bin
- mesh_states.bin (optional)
- region_indices.bin
- surface_molecules.dx (header file for surface molecules)
- surface_molecules_orientations.bin
- surface_molecules_positions.bin
- surface molecules states.bin (optional)
- volume_molecules.dx (header file for volume molecules)
- volume_molecules_orientations.bin
- volume_molecules_positions.bin
- volume_molecules_states.bin (optional)

Visualization data output for the DREAMM_V3_GROUPED mode is in native DX format and includes one master header file and seven binary data files, plus up to two optional data files if state values are specified in the NAME_LIST blocks:

- *filename*.dx (the master header file)
- filename.mesh_positions.bin
- filename.mesh_states.bin (optional)
- *filename*.region_indices.bin
- filename.molecule_positions.bin
- filename.molecule_orientations.bin
- filename.molecule_states.bin (optional)
- filename.iteration_numbers.bin
- filename.time_values.bin

Because the DREAMM_V3_GROUPED mode produces a small number of files, they each may become very large. Hence, reading the files may be slow, but this mode may be best for use on production (supercomputer) machines to avoid transferring large number of files.

All of the keywords in the VIZ_OUTPUT block are optional except FILENAME. If the user does not specify the FILENAME keyword an error message is printed and the simulation aborted. Some of the binary files for both formats may be empty. For example, if no regions are defined the file region_indices.bin will be empty. Similarly, if no meshes or molecules are defined the corresponding mesh_positions.bin or all molecules related binary files will be empty. This avoids unintentional mixing of pre-existing and new files that could result during several runs if incomplete file sets were to be generated with the same names. In DReAMM, the user will only need to point to the *filename*.dx file, and the data from the binary files will be imported as needed for different frames. While using checkpointing in case of the DREAMM_V3_GROUPED format the resulting visualization output files add the checkpoint sequence number to their names, like *filename*.mesh_positions.1.bin. In the case of the DREAMM_V3 format only three files - *filename*.dx, *filename*.iteration_numbers.bin and *filename*.time_values.bin will add the checkpoint sequence number to their names since all the data for each iteration is stored separately in the corresponding iteration # subdirectory.

Examples of VIZ_OUTPUT statements are given below.

Short-hand #1 (time style):

```
VIZ_OUTPUT {
  FILENAME = "viz_data/output_example"
  MESHES {
    NAME_LIST { ALL_MESHES /* or list of object names */ }
    TIME_POINTS { ALL_DATA @ [0] }
}
MOLECULES {
    NAME_LIST { ALL_MOLECULES /* or list of molecule names */ }
    TIME_POINTS { ALL_DATA @ ALL_TIMES }
}
```

Short-hand #2 (iterations style):

```
VIZ_OUTPUT {
  FILENAME = "viz_data/output_example"
  MESHES {
    NAME_LIST { ALL_MESHES /* or list of object names */ }
    ITERATION_NUMBERS { ALL_DATA @ [0] }
}
MOLECULES {
    NAME_LIST { ALL_MOLECULES /* or list of molecule names */ }
    ITERATION_NUMBERS { ALL_DATA @ ALL_ITERATIONS }
}
```

Expanded case:

```
VIZ_OUTPUT {
  FILENAME = "viz_data/output_example"
  MESHES {
    NAME_LIST { ALL_MESHES /* or list of object names */ }
    TIME_POINTS {
        GEOMETRY @ [0]
        REGION_DATA @ [0]
    }
}

MOLECULES {
    NAME_LIST { ALL_MOLECULES /* or list of molecule names */ }
    TIME_POINTS {
        POSITIONS @ ALL_TIMES
        ORIENTATIONS @ ALL_TIMES
    }
}
```

Usual UNIX-style wildcards like "*" and "?" are allowed in the *name_list* but must be enclosed in quotes. For example in the case of MOLECULES the following NAME_LIST statements are all valid:

```
NAME_LIST{A B C1 C2 C3}
NAME_LIST{A B "C*"}
NAME_LIST{A B "C?"}
```

Each MESHES / NAME_LIST statement may contain a single mesh object name or multiple mesh object names with optional state values. It is left to the user to avoid possible confusion arising from overlapping object trees within a single master header file and its associated data files.

3.5.2 Reaction Data Output

Command	Explanation
REACTION_DATA_OUTPUT	Define a new count data output block which contains the commands be-
{	low. Each MDL file can have multiple reaction data output blocks.
reaction output commands	
}	

Each reaction data output block consists of the following commands:

Reaction Output Command	Explanation
OUTPUT_BUFFER_SIZE = N	Write output to disk after every N lines. The default is N =10000. This command is optional, but must be first if it is used. The output will also always be written when the simulation terminates, regardless of N .
STEP = t	Output this block every <i>t</i> seconds. Exactly one of STEP or the follow-
	ing two commands should be used. Triggered output ignores the values
	specified, but some value must still be given.
TIME_LIST = [list]	Output this block at the times specified in the list.
ITERATION_LIST = [list]	Output this block at the iteration numbers specified in the list (i.e. after that number of timesteps).
HEADER = setting	Output blocks by default have no header but can optionally have a header line that states the output (name of molecule, reaction, etc.) in each column. This command can set the behavior of that header line; it applies to all output files until the next HEADER line. A <i>setting</i> of ON turns on the header line; OFF prevents any header. A string, in quotes, will turn the header on and prepend the string to the line; this is useful to add comment character(s). For example, "//" would add a C++-style comment prefix to the line. For TRIGGER statements (see below), the column label (plus comment character if specified) is appended to each line of output when headers are on.
SHOW_EXACT_TIME = setting	TRIGGER statements (see below) can report timing information more precisely than by iteration. However, if only iteration timing is of interest, this can be set OFF. The default is ON. It applies to all output files until the next SHOW_EXACT_TIME line.
{ value } => "file"	Output the value in braces to the filename in quotes. The first column will be the time (in seconds) of the iteration unless the ITERATION_LIST specifier is used, in which case the first column will be the iteration number. For COUNT values, the second column will be the value of the count; other possibilities appear later in this document. This command, and the variants listed below, can be repeated to send different output to many files. The output symbol => has several variants which are described below.
{ value : "name" } => "file"	Output the value in braces with the column header string <i>name</i> to the filename <i>file</i> . Not valid if <i>value</i> is found using wildcards. Trigger outputs put this header in the rightmost column on each line; count outputs put the name at the top of the appropriate column.
{ value , value , } => "file"	For counts, output the list of values in braces, one to a column, in the order listed. The first column will be the time/iteration number; successive columns will be the values in the order listed. If headers are on, each column header can be customized by specifying: "name" after the value. For triggers, all the specified events will be combined into one file.

The values specified in braces are either TRIGGER statements, COUNT statements, or mathematical operations applied to COUNT statements (e.g. you can add, subtract, etc. COUNT statements to each other and to constants and so on). Wildcards can be used to select multiple molecules or reactions by name, but in this case mathematical operations cannot be used. The wildcards? and * can be used to match any single character and any sequence of characters, respectively; internally, this will generate one count/trigger statement per matching name. Having headers on is convenient in this case, so one can tell which column (for COUNT statements) or row (for TRIGGER statements) corresponds to which name.

If a simulation starts from a checkpoint file, it will add to any existing output files. Otherwise, the output files will be overwritten if they already exist.

COUNT statements are either *instantaneous*, and give information about the state of the model at the instant the count is output—the number of molecules in a region, for example—or are *cumulative*, and count the number of events that have occurred since the beginning of the simulation. Alternatively, they can output the time and location of each reaction or molecular collision of the type specified. In all cases, if a region or object is referred to, it should be the fully qualified name starting with the name of the instantiated object.

The COUNT statements themselves have the following syntax:

Count Statement	Explanation
COUNT[name, WORLD]	Count molecules or reactions in the world. If <i>name</i> refers to a molecule, this is an instantaneous count of the number of copies that molecule in the world. If <i>name</i> refers to a reaction, count how many times that reaction has occurred since the beginning of the simulation. If " <i>name</i> " is in quotes, in this command or any of the following commands, the string in quotes can contain wildcards which will be matched to molecule and reaction names and will be listed in alphabetical order. It is usually a good idea when using wildcards to turn on headers so one can see which column is which.
COUNT[name, object]	Count molecules or reactions inside the object called <i>object</i> . This must be an instantiated object. For example, if you have instantiated an object called my_world with a box called my_box inside it, <i>object</i> would be my_world.my_box. If you are counting surface molecules or reactions at a surface, only the ones that actually occur on <i>object</i> will be counted (not those inside which are on a different object). Molecules with a 3D diffusion constant will be counted inside the object, but the object must be closed. All counts are instantaneous.
COUNT[name, region]	Count molecules or reactions inside the named region. For a grid molecule, <i>name</i> can also specify its surface orientation and in such a case has to be enclosed in quotes, e.g., "A,". The surface orientation may be given by an arbitrary number of either ', , or ;. Mixing is not possible. Equivalently, the numerical orientation specifiers {-1}, {0}, or{1} can be used. Clearly, the specification A; or A{0} is equivalent to A since; and {0} both specify no orientation. The named region must be referenced fully. E.g. if my_box (from above) has a region called my_region, the name would be my_world.my_box[my_region]. The count is instantaneous. As with the object syntax, molecules and reactions on surfaces must be on the named region, while volume molecules and reactions must be inside.

Count Statement	Explanation
COUNT [name, region, ALL_ENCLOSED]	Count all molecules or reactions that occur in the area enclosed by re-
	gion (not counting those that occur on the surface of the region). For a
	grid molecule, name can also specify its surface orientation and in such
	a case has to be enclosed in quotes, e.g., "A,". The surface orientation
	may be given by an arbitrary number of either ', , or ;. Mixing is not
	possible. Equivalently, the numerical orientation specifiers $\{-1\}$, $\{0\}$,
	or $\{1\}$ can be used. Clearly, the specification A ; or $A\{0\}$ is equivalent
	to A since; and $\{0\}$ both specify no orientation. This COUNT statement
	lets you count surface molecules contained on surfaces that lie within a
	box, for example. This will work with object names as well as region
	names, but the object or region must be closed. It is only useful for sur-
	face molecules and reactions at surfaces; adding ALL_ENCLOSED is valid
	for volume molecules and reactions, but ALL_ENCLOSED is the default
	behavior. The count is instantaneous.
COUNT [molecule, region, ESTIMATE_CONC]	Estimate the concentration of the molecule at that region, averaged since
	the beginning of the simulation (output has units of μ M). A single ob-
	ject can be used instead of a region. The region/object does not need
	to be closed. To find the average concentration during one count in-
	terval, let t_i be the time of the <i>i</i> th output, let t_j be some earlier output,
	and let $\bar{c}(t)$ be the concentration averaged up to time t . Then the aver-
	age concentration between times t_j and t_i is $\bar{c}(t_j \rightarrow t_i) = \frac{t_i \bar{c}(t_i) - t_j \bar{c}(t_j)}{t_i - t_j}$.
	Note that this is the concentration all around the surface, so if the
	molecule can only reach one side, the concentration on that side will
	be twice what is reported here. The command can be given verbosely
	as ESTIMATE_CONCENTRATION. The estimate is based on a cumulative
	count.
COUNT [molecule, region, hits]	Output the number of times the named molecule has hit the named region
	(or object). The <i>hits</i> specifier should be one of FRONT_HITS, BACK_HITS,
	ALL_HITS, FRONT_CROSSINGS, BACK_CROSSINGS, and ALL_CROSSINGS.
	The count is cumulative.
EXPRESSION[expression]	Evaluate and output a mathematical expression. This can be mixed with
	COUNT statements but not with TRIGGER statements.

Cumulative counts are reset when a simulation is started from a checkpoint. This breaks ESTIMATE_CONC, but the other cumulative counts can be recovered by adding the last report before the checkpoint to the first one after the checkpoint.

TRIGGER statements output the time and location each time the number of molecules changes or a reaction happens. Most COUNT statements have a corresponding TRIGGER, but TRIGGER statements are not compatible with the WORLD or the ESTIMATE_CONC directives. Within output statements pointing to the same output file, there can only be TRIGGER commands, i.e., they cannot be mixed with COUNT or EXPRESSION statements.

 ${\tt TRIGGER}\ statements\ obey\ the\ following\ syntax:$

Trigger Statement	Explanation
TRIGGER[molecule, region]	Generates output each time the number of molecules inside the specified
	region changes. The output has the format <i>iteration_time exact_time X Y</i>
	Z orientation number [name] as described below. The sixth column, ori-
	entation, gives the molecule orientation, i.e., it is 0 for volume molecules
	and +/-1 for surface molecules according to their orientation with respect
	to the surface containing them. The seventh column, <i>number</i> , can take
	on values of +/-1 depending on if the molecule was added or removed,
	respectively, from the region. If HEADER is on, the eighth column lists the
	molecule name.

Trigger Statement	Explanation
TRIGGER[reaction, region]	Generates output each time the named reaction takes place inside the
	specified closed region. The output has the format iteration_time ex-
	act_time X Y Z [name]. The fields are described below. Note that since
	reactions do not have an orientation and always occur one at a time the
	orientation and number fields are omitted. If HEADER is on, the sixth
	column lists the reaction name.
TRIGGER[name, object]	This is equivalent to specifying a list of TRIGGER statements which con-
	sist of all regions in that object.
TRIGGER[name, region, ALL_ENCLOSED]	Generates output each time the named reaction takes place, or number
	of named molecules changes, inside the specified closed region. As with
	COUNT statements, this is only useful for surface reactions and molecules,
	and does not include the surface of the named region, only events wholly
	inside it. The output has the format appropriate for molecules or reac-
	tions.
TRIGGER[molecule, region, hits]	Generates output each time the molecule hits or crosses the named re-
	gion. The hits specifier should be one of FRONT_HITS, BACK_HITS,
	ALL_HITS, FRONT_CROSSINGS, BACK_CROSSINGS, and ALL_CROSSINGS.
	The output has the format iteration_time exact_time X Y Z orientation
	[name]. The orientation column can take on values of +/-1 depending
	on if the region was hit or crossed from the front or the back, respec-
	tively; other columns are described below. Note that the <i>number</i> column
	is omitted. If HEADER is on, the seventh column lists the molecule name.

The output contains one row of data for each even that happened. The format of the columns is:

iteration_time exac	ct_time X	Y	Z	[orientation]	[number]	[name]
---------------------	-----------	---	---	---------------	----------	--------

Iteration_time is the time of the iteration during which the event happened, or the iteration number if ITERATION_LIST was specified for the block.

Exact_time is the time at which the event was scheduled, between iteration_time and the time of the next iteration. Since events within one iteration are not ordered precisely, exact_time values will not always increase. This column can be turned off by using the SHOW_EXACT_TIME=OFF directive inside the REACTION_DATA_OUTPUT block. These values are always times, even if ITERATION LIST is specified for the block.

X, Y, and Z are the coordinates at which the event took place. Reactions and hits always report their coordinates precisely. Volume molecules that disappear at a surface will report their final position as slightly inside the surface along their last trajectory (so that it is possible to tell which side of the surface they were on); if they react with another volume molecule they will report the position they reached when their interaction disk intersected the target molecule, not the position of the target. Surface molecules diffuse by hopping rather than raytracing, so when a surface molecule leaves a region of interest, the position reported is the last position where the molecule was located inside the region, not the boundary of the region where it crossed out (and conversely, when entering, it's the first position where the molecule stopped at the end of its time-step).

Orientation and number are only provided for certain types of triggers and are described above.

Name is the name of the molecule or a user-defined string, and present in the last column (6, 7, or 8 depending on which type of trigger is used) if headers are on.

The following output symbols can be used in place of => and give the behaviors described below. All output symbols will create files if none exist. No output symbols will create directories—if the files that are referred to cannot be created as specified, MCell3 will quit with an error message. Output may create empty files if the simulation ends without producing output (either because of an error condition or because the simulation did not run long enough to reach the time/iteration of any reaction data output).

Output Symbol	Explanation
=>	-
=>	If a checkpoint file is not used, overwrite the existing file (with headers if
	requested). If a checkpoint file is used, discard any of the output file that
	appears to be a later time than the start of the current run, and append
	to the file from that point. Headers are not written unless the file has
	to be created or is empty to begin with. This command generally does
	"what you expect"—after the simulation has run, it will contain data from
	earlier in the simulation that the current run, plus the data created in the
	current simulation. If you switch between ITERATION_LIST and other
	output time specifiers, this command won't know whether output is by
	time or by iteration number, so don't use this command if you switch
	from one to the other after checkpointing.
>	Always overwrite the file, whether or not a checkpoint is used. If headers
	are requested, they will appear at the beginning of the file.
+>	Always create a new file, whether or not a checkpoint is used. If a file
	of the given name already exists and is not empty, MCell3 will print an
	error message and exit. If headers are requested, they will appear at the
	beginning of the file.
>>	Always append to an existing file without removing any previous data.
	Headers are only written if the file starts out empty or has to be created.
>>>	Always append to an existing file without removing any previous data
	and if headers are requested, write them even into the middle of the file.

3.5.3 Other Output Commands

The MCell 2 style $\tt VIZ_DATA_OUTPUT$ block is also supported (a maximum of one per MDL file) for backwards compatibility. It is no longer explicitly supported, however, so the format is not described here.

3.6 Utility commands

MCell3 understands the standard numeric operations +	_ *	/ as well as the following standard numerical functions:

Numerical Command	Explanation	
SQRT (X)	Return the square root of <i>x</i>	
EXP(x)	Return the value of e raised to the x^{th} power	
LOG(X)	Return the natural logarithm of x	
LOG10(x)	Return the base 10 logarithm of x	
SIN(x)	Return the sine of <i>x</i>	
COS(X)	Return the cosine of <i>x</i>	
TAN(X)	Return the tangent of <i>x</i>	
ASIN(X)	Return the inverse sine of <i>x</i>	
ACOS (X)	Return the inverse cosine of <i>x</i>	
ATAN (X)	Return the inverse tangent of <i>x</i>	
ABS(x)	Return the absolute value of <i>x</i>	
CEIL(X)	Return the smallest integer at least as big as x	
FLOOR(x)	Return the largest integer at no bigger than x	
MAX(x,y)	Return the larger of x and y	
MIN(x, y)	Return the smaller of <i>x</i> and <i>y</i>	
RAND_UNIFORM	Return a random number uniformly distributed between 0 and 1	
RAND_GAUSSIAN	Return a random number from a Gaussian distribution with mean 0 and	
	standard deviation 1.	
PI	The numeric value $\pi = 3.14159265358979323846$	
SEED	The value of the random number generator seed	

At any outer block in MCell3, one can define variables simply by assigning a value to the name of the variable. E.g. my_lucky_number=13 would be a valid (if unusual) way to define a variable. Variables can take numeric, array, or string values. String values consist of text between double quotes. Strings can be combined with the & operator. Array values are lists of numbers inside brackets separated by commas, or starting and ending values plus a step size, as exemplified below (note the double brackets):

```
my_lucky_number = 13
my_favorite_array = [1,3,5,7,11,17]
my_second_favorite_array = [[1.3 TO 2.75 STEP 0.331]]
my_boring_string = "la la la, la la la" & ", la la"
```

The C-style printf and sprintf commands work too, pretty much the way you'd expect them to. Note that MCell3 variables are doubles, so the integer formats may not give the expected results. To turn the random number generator seed into a string that you can use as part of a filename, use sprintf (my_string_name, "%g", SEED). If you want it to be a fixed width, e.g. 3 characters padded with zeros, use the appropriate format string, e.g. "%03g".

MCell3 comments are delimited by /* and */ and can be nested.

MDL files can include other MDL files using the following syntax:

Command	Explanation
INCLUDE_FILE = "filename"	Parse the text in <i>filename</i> as if it were inserted into this MDL file at this
	point.

Paths are relative to the location that MCell was run from, not relative to the MDL file being parsed.

4 Technical details affecting simulation speed and accuracy

4.1 Partitioning

In future releases, MCell3 will automatically partition space to improve execution speed. Currently, however, this must be performed manually. In general, partitions should be chosen to avoid having too many surfaces and molecules in one subvolume defined by the partitions. Molecules that are specified as TARGET_ONLY or which do not interact with other molecules diffusing in 3D need only have relatively few surfaces in one subvolume.

If there are few surfaces and/or molecules in a subvolume, it is advantageous to have the subvolume as large as possible. Crossing partition boundaries takes a small amount of time, so it is rarely useful to have partitions more finely spaced than the average diffusion distance of the faster-moving molecules in the simulation.

In cases where the diffusing molecules do not interact with each other, they can safely take extended time-steps by measuring how far they are from things they could interact with. In this case, the partitions with no surfaces should be as large as possible. For example, a box works well with partitions just inside its outer walls.

Finally, note that partition placement is not exact. The model is divided into 16384 possible partition boundaries, so partitions may shift by up to about one part in twenty thousand of the size of the model. For instance, if the model has a structure that is 6μ m long, partitions may vary by about 0.0003μ m. Thus, do not place partitions too close to objects in your model or they may not appear on the side you expect them to appear.

4.2 Mean diffusion distance

Diffusion in MCell3 (and in earlier versions of MCell) is modeled as a series of motions in a straight line. This is a good approximation around geometry that is of a larger scale than the mean diffusion length for the time-step of the molecule in question. For accurate results around intricate geometry, it may be necessary to reduce the time step (or space step).

4.3 Reaction probabilities

MCell3 assigns a probability of reaction for each collision. These probabilities are chosen to match the bulk reaction rate specified in the MDL file. The match will not occur, however, if the probability goes above 1.0. Internal correction factors can also raise the actual probability above the typical probability specified at the beginning. Therefore, MCell3 will output a warning if the reaction probability goes above 0.8 for reactions where a volume molecule hits a surface, or if the probability goes above 0.3 for a collision between pairs of volume molecules. !! Is this even true? !!

If warnings are given (and possibly even if they are not), one should reduce the time step to lower the probabilities and see if the same results are generated. If not, simulations should be run with shorter time steps in order to avoid overly high probabilities.

Unimolecular reactions with half-lives of less than one time step are also not perfectly accurate. Although unimolecular transitions will always occur at the right rate, other molecules may not experience the right effective concentration of each state, since a short-lifetime species may not be converted to another species until the end of the time step after which many other molecules may have had a chance to interact with it. Thus, the shortest-lifetime species in a series of unimolecular transitions should not have a half-life of less than approximately one time step if other molecules can interact with that state.

4.4 Interaction radii

Bimolecular reactions occur within a distance specified by the INTERACTION_RADIUS command. In many cases, one may want to increase or decrease this value. In particular, in order to get the right probability of reaction, MCell3 increases the probability of reaction when near surfaces.

If ACCURATE_3D_REACTIONS is set to FALSE, MCell3 also treats partition boundaries as opaque and increases the probability of reaction rather than looking for molecules on the other side of the partition. This speeds execution time but can lead to error, the reaction rate has approximately 1-2% error if the average spacing between surfaces is at least 10 times the interaction radius, and the reaction probabilities are 0.3 or less. For example, if one has partitions spaced $0.02\mu m$ apart, simulation accuracy will be poor with the default interaction radius of $0.01\mu m$. Thus, one might wish to specify INTERACTION RADIUS=0.001.

4.5 Placing molecules in the world

There are two ways to place molecules on surfaces: with a release site on a region, and as part of the property of a surface or region. Release sites are more flexible but slower; if you do not need the flexibility of release site notation, you're better off defining a region and using the MOLECULE_DENSITY or MOLECULE_NUMBER commands to add molecules at initialization.

All placement of molecules in volumes is done with release sites. However, the geometrical release sites (CUBIC and SPHERICAL) require less computation to place each molecule. Thus, these should be used preferentially for simple geometry. To release many particles at a one point, use a cubic release site and set the diameter to 0. To release many particles at different points, use the LIST release type.

5 Example models

5.1 Ligand-gated ion channel

Below are a set of molecule definitions and reactions that specify an ion channel that is gated by the binding of a single ligand.

```
DEFINE_MOLECULES {
  channel_unbound { D_2D=0 }
  channel_bound { D_2D=0 }
 channel_open { D_2D=0 }
 ligand
                \{ D 3D=2e-8 \}
 ion
                \{ D_3D=3e-8 \}
DEFINE_REACTIONS {
 channel_unbound' + ligand' -> channel_bound'
                                                         [1e7]
 channel_bound' -> channel_unbound' + ligand' [2e2]
 channel bound'
                           -> channel open'
                                                         [5e2]
                           -> channel open'
 channel open'
                                               + ion,
                                                         [8e4]
```

We have defined a reaction where a ligand binds to one end of a channel (presumably the extracellular face), which causes the channel to be in its bound state. In that state it can either release the ligand or become open. While open, it will emit ions on the other end (presumably the intracellular face). This would be suitable if the ion concentration is much higher outside than inside, or the membrane potential makes it highly favorable for the ion to move inside, so that we don't have to worry about the reverse reaction. If there is no electrical driving force, we might have to model ions both inside and outside:

```
DEFINE_REACTIONS {
  channel_unbound' + ligand' -> channel_bound' [1e7]
  channel_bound' -> channel_unbound' + ligand' [2e2]
  channel_bound' -> channel_open' [5e2]
  channel_open'' + ion' -> channel_open'' + ion, [4e7]
}
```

Here, the ion travels in either direction just as easily since it pays no attention to the orientation of the channel. However, if there was a modest driving force, traveling in might be easier than traveling out, which would be reflected in the rates.

```
DEFINE REACTIONS {
 channel unbound' + ligand' -> channel bound'
                                                           [1e7]
 channel bound'
                            -> channel unbound' + ligand' [2e2]
                            -> channel_open'
 channel bound'
                                                           [5e2]
                         -> channel_open'
 channel_open' + ion'
                                                 + ion,
                                                           [4e8]
 channel_open' + ion,
                           -> channel_open'
                                                 + ion'
                                                           [1e8]
```

In this case, the ion is four times as likely to travel from outside to inside as inside to outside.

5.2 Example bimolecular reaction

Here's a complete MDL file that implements a simple bimolecular reaction that should achieve equilibrium at 482 molecules of each species.

```
time\_step = 1.0e-6
TIME STEP = time step
TIME_STEP_MAX = time_step
ITERATIONS = 1e-2/time_step
EFFECTOR_GRID_DENSITY = 10000
INTERACTION_RADIUS = 0.001
PARTITION_X = [ [-0.1 \text{ TO } 0.1 \text{ STEP } 0.01] ]
PARTITION_Y = [-0.1 \text{ TO } 0.1 \text{ STEP } 0.01]
PARTITION_Z = [ [-0.1 TO 0.1 STEP 0.01] ]
DEFINE_MOLECULES
  A \{ D_3D = 100e-8 \}
  B \{ D_3D = 100e-8 \}
 C \{ D 3D = 100e-8 \}
/* Your basic reversible binding reaction */
DEFINE_REACTIONS
 A + B -> C [1e7]
 C -> A + B [1e3]
small_box BOX
  CORNERS = [-0.1, -0.1, -0.1] , [0.1, 0.1, 0.1]
 /* REMOVE_ELEMENTS { TOP, LEFT } */ /* Could remove sides ... */
  /* REMOVE_ELEMENTS { INCLUDE_PATCH = [0.1,0,0] , [0.1,0.05,0.05] } /*... or patch*/
INSTANTIATE my_world OBJECT
  A_release CUBIC_RELEASE_SITE {
    LOCATION=[0,0,0]
   MOLECULE=A
    NUMBER TO RELEASE=482
    SITE_DIAMETER=0.196
  B_release CUBIC_RELEASE_SITE {
    LOCATION=[0,0,0]
    MOLECULE=B
   NUMBER_TO_RELEASE=482
    SITE_DIAMETER=0.196
  C_release CUBIC_RELEASE_SITE {
    LOCATION=[0,0,0]
    MOLECULE=C
    NUMBER_TO_RELEASE=482
    SITE DIAMETER=0.196
my_box OBJECT small_box {}
```

```
REACTION_DATA_OUTPUT
{
   STEP = 1e-5
   { COUNT [A,WORLD] } => "eq_A.dat"
   { COUNT [B,WORLD] } => "eq_B.dat"
   { COUNT [C,WORLD] } => "eq_C.dat"
}
```

6 Authors

The following authors have contributed to this document, as described.

- Rex Kerr
- Markus Dittrich