

3Worlds user manual

Jacques Gignoux – jacques.gignoux@upmc.fr · Ian D. Davies – ian.davies@anu.edu.au · Shayne R. Flint – shayne.flint@anu.edu.au – Version 1.11, 31 August 2022 |

Table of Contents

1. Purpose and motivation
 - 1.1. Why another simulation environment for ecology?
 - 1.2. Design Concepts
 - 1.2.1. Individual based models
 - 1.2.2. The complex system as a dynamic graph
 - 1.2.3. Categories
 - 1.2.4. Time with simultaneous events
 - 1.2.5. Causality
 - 1.2.6. Space as a mediator for interactions
 - 1.2.7. Modelling made easy
 - 1.2.8. Model comparison: graphs can be compared
 - 1.2.9. Problem upscaling
 - 1.2.10. Portability and performance
2. Getting started - download and installation
 - 2.1. Basics - what you must know before starting
 - 2.2. Prerequisites
 - 2.3. Running ModelMaker
 - 2.4. Setting up a Java development environment for the user code
 - 2.4.1. Setting up the Eclipse Integrated Development Environment (IDE) for 3Worlds
 - 2.4.2. Running ModelMaker from Eclipse
 - 2.4.3. Linking user code with model configuration
 - 2.4.4. Debugging and testing user code
3. Configuration reference: creating and editing a model with ModelMaker
 - 3.1. General concepts: structure of a 3Worlds configuration
 - 3.1.1. A tree structure...
 - 3.1.2. ...with cross-links
 - 3.1.3. What ModelMaker does for you
 - 3.2. Using ModelMaker: software interface and functioning
 - 3.2.1. Main menu
 - 3.2.2. Graph display
 - 3.2.3. Property editors
 - 3.2.4. Message pane
 - 3.3. Configuration options: reference
 - 3.3.1. The *3Worlds* node
 - 3.3.2. The *system* node
 - 3.3.3. The *system/structure* node
 - 3.3.4. The *system/dynamics* node
 - 3.3.5. The *dataDefinition* node
 - 3.3.6. The *experiment* node
 - 3.3.7. The *userInterface* node
 - 3.3.8. The *predefined* sub-tree
 - 3.4. Developing and testing model code
 - 3.4.1. Generated code: the *model main class*

3.4.2. Model main class method arguments

3.4.3. Using space in user code

3.4.4. Accessing complex data structure

3.5. Feeding the model with data

3.5.1. Data uses

3.5.2. Data input methods

3.5.3. Consistency of data files with the model component hierarchy

3.5.4. Data file formats

3.5.5. Import data from external sources

4. Simulation reference: running a simulation experiment with ModelRunner

4.1. General concepts

4.2. Using ModelRunner: software interface and functioning

4.3. Getting output from a simulation experiment

5. Sample models and tutorials

5.1. Tutorial 1: Construct and run a simulation for the first time

5.1.1. Introduction

5.1.2. Installation

5.1.3. Creating a new 3Worlds project

5.1.4. Creating the specifications

5.1.5. Graph layouts

5.1.6. Next

5.2. Tutorial 2: Linking a 3Worlds project to a Integrated Development Environment (IDE)

5.2.1. Introduction

5.2.2. New Java project

5.2.3. Specifications

5.2.4. Writing Java code

5.2.5. Next

5.3. Tutorial 3: Using tables

5.3.1. Introduction

5.3.2. Specifications

5.3.3. Next

5.4. Tutorial 4: Elaborating the model structure: Testing the Intermediate Disturbance Hypothesis

5.4.1. Introduction

5.4.2. Specifications

5.4.3. User interface

5.4.4. Next

5.5. Tutorial 5: Event driven systems - Using the Event Timer

5.5.1. Introduction

5.5.2. Specifications

5.5.3. Next

5.6. Tutorial 6: Event driven systems - Using the Event Timer in a spatial model

5.6.1. Introduction

5.6.2. Specifications

5.6.3. Next

5.7. Tutorial 7: Random number generators

5.7.1. Introduction

5.7.2. Specifications

5.7.3. Simulation

5.7.4. Next

5.8. Tutorial 8: Running an experiment

5.8.1. Introduction
5.8.2. Specifications
5.8.3. Simulation
5.8.4. Results
Bibliography

by **Jacques Gignoux, Ian D. Davies & Shayne R. Flint**

Version: **1.11** (31 August 2022)

1. Purpose and motivation

1.1. Why another simulation environment for ecology?

“Though the organisms may claim our primary interest, when we are trying to think fundamentally we cannot separate them from their special environment, with which they form one physical system” A. G. Tanlsey (1935).

Ecosystems, as first proposed by Arthur Tansley (1935), have both physical and biological aspects. They comprise not only energy and chemical stocks and flows but also living things that are born, reproduce and die, exhibiting particular behaviours over the course of their existence. They are made from and return to the physical world, are part of it and at the same time, constitute a domain where terms such as reproduction, social interaction, communication and life cycle have meaning.

To this day, reconciling these two aspects, where the physical is articulated in SI units and the biological in descriptive terms (e. g. heterotroph, shade-tolerant, obligate seeder), remains a key problem in ecosystem studies: How does an energy flux translate into a number of living individuals or in a higher biodiversity? How might a particular reproduction strategy affect river hydrology and the regional water budget (e. g. the beaver)? How do social interactions affect the frequency of hurricanes (e. g. climate change)? **3Worlds** aims at providing *a tool for building simulations that couple the physical and biological aspects of ecosystems*.

It is our view that existing tools focus predominantly on one or other of these aspects. For example, dynamic models based on differential equations and can only represent reproduction and growth using metaphors for continuous physical quantities (e.g. matrix population models). Multi-agent systems (MAS), on the other hand, model social interactions well but pose difficulties in balancing mass or energy budgets.

In addition to this physical and biological duality, ecosystems are '*multi-scale*' (cf. an extensive discussion on this topic and other questions related to the ecosystem concept in Gignoux et al. 2011). Tansley's ecosystem definition applies equally well to the whole biosphere as to a single bacteria in a drop of water. Currently, there is no software enabling one to consider ecosystems at such different spatial and temporal scales within the same conceptual framework. A second aim of **3Worlds** is to fill this gap. If the ecosystem concept is robust enough, and we believe it is given its success in ecology and its adoption in other scientific fields, then it should be possible to design a conceptually sound and consistent computing framework to deal with a multiplicity of scales.

However, changing the scale of a model is not just a matter of changing spatio-temporal resolution. It also involves the level of detail of the representations used in the model, something which has been formalized in *abstraction theory* (Zucker 2003): the more detailed a model, the less abstract it is. The question arises then as to what level of detail should we model our system to capture its important behaviour? How does changing scale (in this broadest sense) effect model outcomes? There can be no general answer to this question beyond trial and error. This makes the possibility to test and compare different representations of the same system at different scales an essential feature of any software dealing with ecosystems. **3Worlds** provides *great flexibility in ecosystem representation* enabling one to represent *any* ecosystem at *any* spatial and temporal scale with *any* level of detail—all this being determined by the researcher in accordance with the purpose of their study. The researcher has full control on the detailed biological

functions and variables relevant to their study.

How is this flexibility compatible with the strong conceptual backbone? **3Worlds** uses *aspect-oriented thinking (AOT)* (Flint 2006), a method to build complex systems (e.g. software) from independent areas of knowledge. We have designed an *archetype* of what we believe constitutes an ecosystem—a recursive and multi-scale system of interacting entities. We use an implementation of AOT to ensure that any user-defined representation of an ecosystem complies with the archetype. This guarantees that, despite a great freedom left to the modeller, their model will always be compatible with the **3Worlds** software. The great benefit of this is that while we believe it's possible to construct any type of model within this archetype, imposing specification constraints greatly assists in model comparison: why should two models, ostensibly constructed for the same purpose, differ in their outputs? How does a change in temporal or spatial scale affect projections? How does adding or removing sub-systems change model projections? *Models developed in **3Worlds** are always comparable*, unlike many large model intercomparison exercises where the models are independently developed without reference to any common structuring principals.

The ecosystem is a recursive concept (Gignoux et al. 2011): ecosystems can be nested. Parts of an ecosystem can be studied as ecosystems themselves. We use the concept of a hierarchical system, formalized as a graph (Gignoux et al. 2017) to implement this idea. In **3Worlds**, *the modelled ecosystem is a graph* where nodes represent relevant entities: individuals, populations, climate, soil, area, ...; and edges represent relations between these entities. These relations can be of any kind, including a hierarchical relation describing the complex nesting of sub-systems. This provides an elegant solution to the apparent complexity of ecosystems: it allows for various types of *emergence*, enables the comparison of system structures and simulation trajectories, and can represent virtually anything an ecological modeller can conceive of.

3Worlds builds upon more than thirty years of experience of its developers in ecosystem and complex system modelling and simulation. Over this time, we have reached the conclusion that: (1) robust concepts are fundamental for building sound software; (2) some problems, that appear over and over again in the life of a simulation modeller, often have well-established solutions, sometimes in other scientific fields; and (3) complex systems are more easily managed and understood when designed within a sound framework.

Many of the ideas we use are already available, it's just a matter of making them work together, and **3Worlds** is our best attempt to do so!

1.2. Design Concepts

3Worlds is based on a few concepts and techniques that have a very broad application and harnesses them to fit the needs of ecosystem modelling.

1.2.1. Individual based models

Fundamentally, **3Worlds** is designed as a framework for *individual-based models* (IBMs: Grimm & Railsback 2005). IBMs, like multi-agent systems (MASs: Bousquet & Lepage 2004), assume that some important properties of complex systems (like ecosystems and human societies) cannot be understood without representing the detailed behaviour of every individual in the system. In an IBM, a list of 'individuals' (called 'agents' in MASs) is kept in memory, and their interactions through various functions yield the dynamics of the whole system over time. Individuals can come and go, which means that their list is constantly changing in size.

3Worlds simulates a *system* made of interacting *system components*, which are equivalent to the individuals or agents of IBMs and MASs. If required by a simple application, the **3Worlds** *system* can be reduced to zero components, just as in any system dynamics application; but this is not the use **3Worlds** has been designed for, and other more specialised software may be more adapted for this use case.

1.2.2. The complex system as a dynamic graph

Ecosystems are commonly considered as *complex systems*, without much agreement on what this concept means. Complex systems are usually assumed to possess *emergent properties*; again, with little agreement on what this means.

We argued in Gignoux et al. (2017) that the only common feature of all definitions of emergence is that emergence can only appear in systems with a 'macro-state' and 'micro-states'. We called such systems *hierarchical systems* and assumed that they can be ideally represented with a *dynamic graph*.

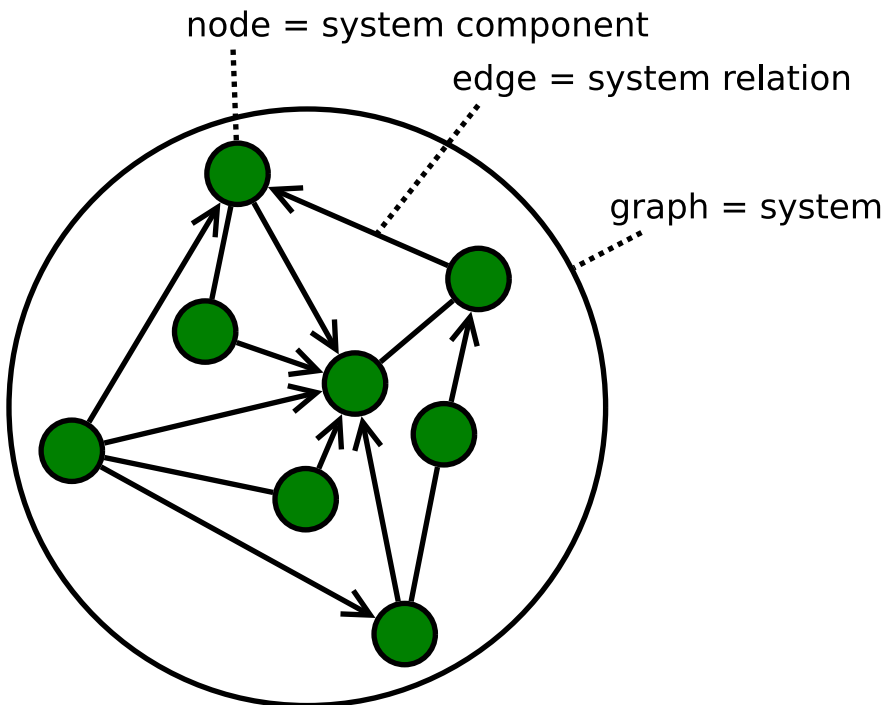


Figure 1. A mathematical graph, showing both the usual mathematical terms and the vocabulary used in 3Worlds. The incidence function defines edges between nodes. Edges can be directed (with arrows) or undirected.

According to graph theory (https://en.wikipedia.org/wiki/Graph_theory), a *graph* is a mathematical construct comprising a set of *nodes*, a set of *edges* linking these nodes defined in an *incidence function* (Figure 1). A *dynamic graph* (Harary & Gupta 1997) is a graph where these three components (the sets of nodes, edges, and the incidence function) can change over time.

With the addition of *descriptors* (e.g. numbers with a specific meaning: Figure 2) to nodes and edges, we can use the graph nodes as the individuals of our IBM, and the graph edges give a concrete existence to the interactions between those nodes. If we define *functions* that can modify nodes and edges and their descriptors, the graph becomes dynamic. A **3Worlds** simulation runs a dynamic graph where nodes, edges, descriptors, functions can be adapted to the user's

particular application. Nodes are called *system components*, edges *system relations*, and descriptors *properties*.

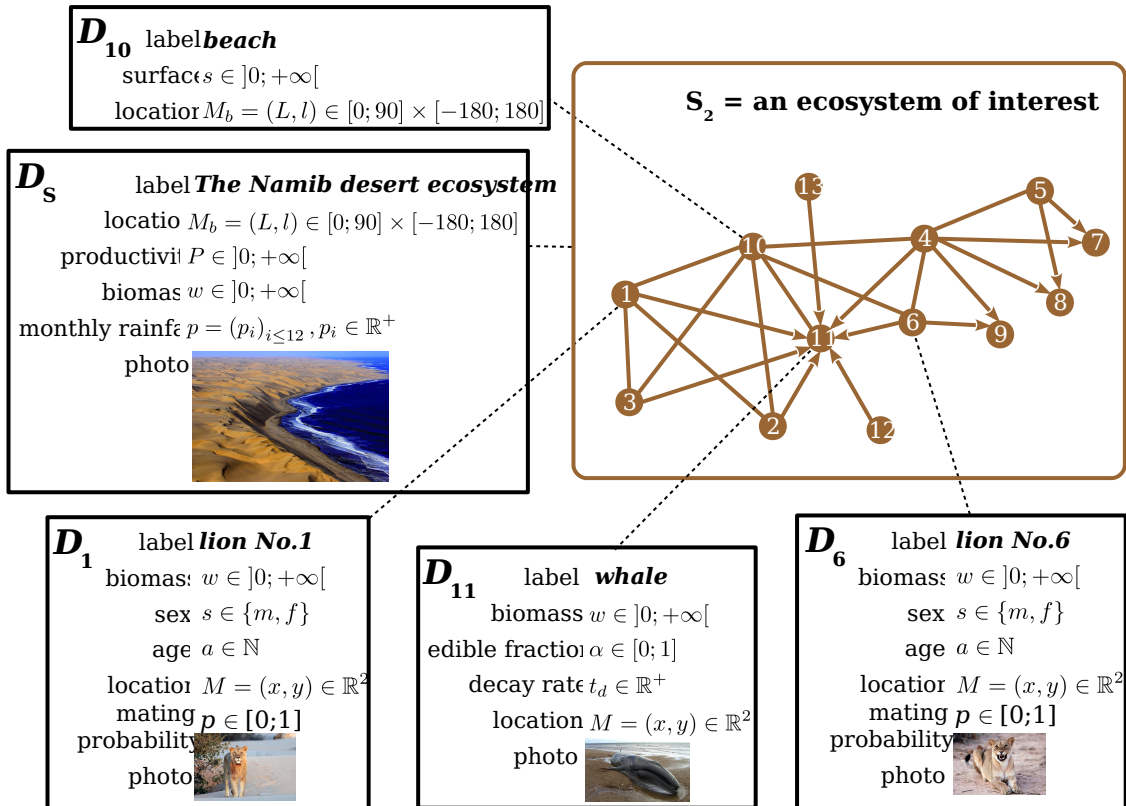


Figure 2. An example of graph descriptors (redrawn from Gignoux et al. 2017).

1.2.3. Categories

In an IBM, every individual differs from all others. But it is common practice to assume that some groups of individuals share some things in common. This is the essence of modelling: finding commonalities within an ocean of particular cases. We use the concept of *category* to group system components that 'look like each other' in some way. Categories are used to express how alike and how different certain groups of components are from each other.

In a dynamic graph, categories are used to specify common descriptors to groups of components (e.g. a plant species average growth rate, an animal cohort survival rate, ...) and functions that operate on a group of components of the same category. They are also used to specify which *type of relation* is possible between components of different categories.

This category concept is similar in many ways to the class concept used in the [UML](http://uml.org/what-is-uml.htm) (<http://uml.org/what-is-uml.htm>) language and in [object oriented programming](https://en.wikipedia.org/wiki/Object-oriented_programming) (https://en.wikipedia.org/wiki/Object-oriented_programming). Categories can be nested, and can be exclusive of each other or not, that is, something can belong to one category of a set of categories but not others within the same set e.g you can be *ephemeral* or *persistant* but not both. They are central to the organisation and execution of a simulation in **3Worlds**.

1.2.4. Time with simultaneous events

Because ecosystems have both biological and physical characteristics, they not only deal with individuals and populations of living organisms, but also with fluxes of matter and energy. To properly compute a mass or energy balance typical of physical questions, we need a time model that insures that all system components are modified synchronously - otherwise, leaks in mass and energy budgets may occur (Figure 3). This is where IBMs differ somewhat from MASs in their most common current implementations: MASs emphasize the *autonomy* of agents by allowing them to modify their state immediately. In other words, MASs assume that no two events occurring in a simulation can be simultaneous, while mass/energy balance requires simultaneity of events. 3Worlds assumes that simultaneous events are the default, but by using particular time models it is possible to relax this constraint.

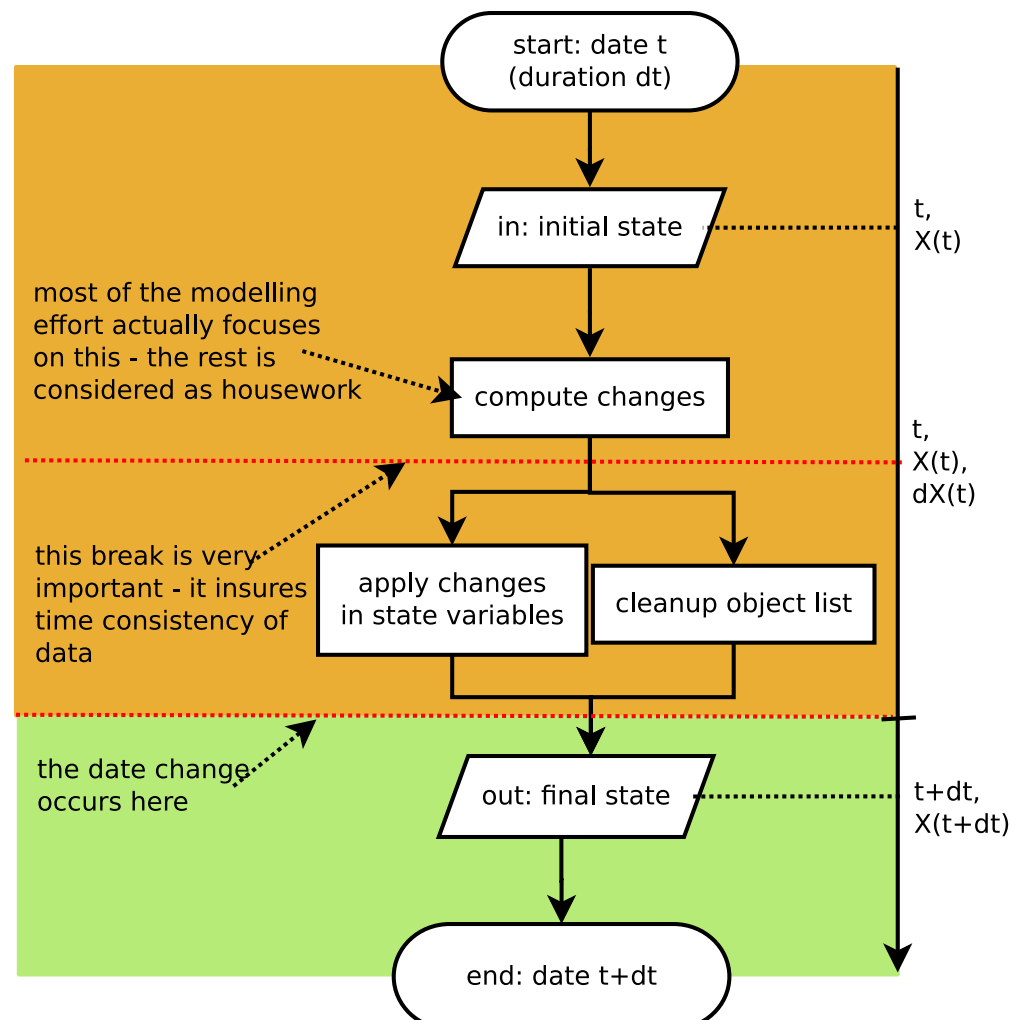


Figure 3. The time model of 3Worlds. The overall dynamics of a system is computed as $X(t+dt) = f(X(t))$ where X represents the state of all system components, t is time and f is the function (or rather, the set of functions) used to compute the change over the time interval dt .

Given the diversity of topics subject to ecological modelling, ecological processes span many orders of magnitude in their rate of action. Demographic models often use a yearly time step, while ecophysiological models may use daily

time steps and physiological models may run with time steps of a second. In **3Worlds**, time steps span the whole scale of time units relevant to ecology, i.e. from milliseconds to millenia. Standard Gregorian calendar time can also be used. **3Worlds** provides three types of *time model* that can be used in interaction: '*clock*' models using a constant time step, *event-driven* time models where time events trigger computations that can generate further events in the future, and *scenarios* where the list of events determines when computations are to be made.

1.2.5. Causality

When designing a model in practice, it appears that some computations can or must be made 'in parallel' while others must be made sequentially. For example:

- before computing how much carbon a plant allocates to its roots or to its stems, one has to know how much carbon was gained from photosynthesis: within a time step, photosynthesis must be computed before allocation.
- before computing how much nitrogen an animal gets from its food according to its digesting system, one has to know how much food was ingested: food intake must be computed before digestion outcome.
- water runoff in a water catchment is triggered by rainfall.
- to compute competition for a shared resource, one has first to compute how much resource is available, how much demand there is for this resource by the organisms having an access to it, and then which share they will get based on some rule. Resource availability must be computed before demand and demand before uptake.

In these examples, 'A must be computed before B' must be understood as 'A is the cause of B'. This 'before' does not refer to time scale: no matter at what scale we measure, think, or compute, a cause must be computed before its effect. It would be nonsense to compute things in another way.

In other words, the order of computations in a simulator does not only depend on time ordering, but also on the logic of causality implied by modelling choices.

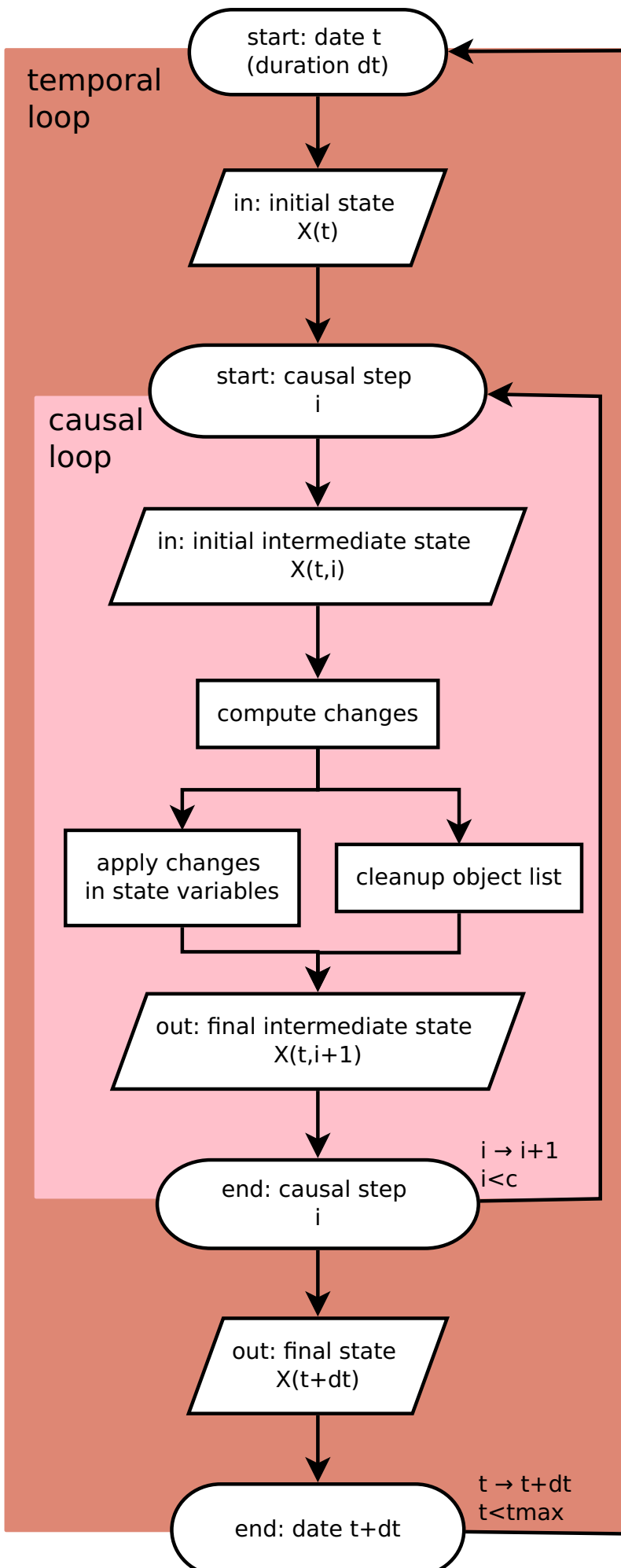
This need for some ordering of computations within a time grain is important and was identified as such in implementations (Quesnel et al. 2009) of versatile simulation timing tools such as **DEVS** (https://fr.wikipedia.org/wiki/Discrete_Event_System_Specification) (Zeigler et al. 2000). In **3Worlds**, the elementary time step of Section 1.2.4 is sub-divided into *causal* steps, enabling to proceed to computations in a meaningful order, and with intermediate update of state and graph structure (Figure 4).

1.2.6. Space as a mediator for interactions

IBMs are often 'spatialized', which means they include some representation of the physical space in which organisms of an ecosystem interact. We argued in Gignoux et al. (2011) that space is not a requirement of the ecosystem definition, but rather an optional feature. Besides the usual 3-dimensional space, we called the place where organisms interact in an ecosystem the *arena*, i.e. the place where things happen and where a public (of ecological modellers) is watching them. This does not mean an Euclidian (or any other kind of) space must be associated with the ecosystem representation. Using a dynamic graph is sufficient.

However, it is often the case that explicitly considering space in a model is necessary for computing ecological interactions. In most ecological process models, there are actually implicit assumptions about space and how it affects organism interactions. For example, seed dispersal in plants is easily computed in a 2-dimensional space where the location of seeds depends on that of parent plants and some simple distance law; water flow in a catchment relies on a 2-dimensional space plus an elevation of ground surface to some x and y resolution; competition between individual trees in a forest assume a vertical distribution of leaves depicted with varying degrees of detail.

Following Gignoux et al. (2011), we optionally provide predefined spatial representations to include in a simulator. Different spaces can be used within a single simulator, depending on the needs of the process computations. They are associated with optimal search algorithms (e.g. Kd-trees) that speed up the search of components with which to form dynamic relations.



1.2.7. Modelling made easy

The community of ecological scientists has been developing an impressively large number of models, yet most of them are poorly designed in terms of programming, as ecologists are not necessarily software engineers. Ecosystem simulators are among the most complex programs (Coquillard & Hill 1997). They require high programming skills and constitute a huge investment in time, which makes their production slow and hazardous. As a result, once built, they tend to be used beyond their initial domain of application (e.g. the overuse and abuse of the CENTURY model: Parton et al. 1988), issues of provenance and repeatability are rarely addressed, shedding some doubt on the discipline as a whole.

With **3Worlds**, we wanted to provide a simulation platform for ecosystem modelling using state-of-the-art concepts and algorithms, and sound programming techniques (e.g. systematic code testing, separated concerns), so that ecological modellers can concentrate on the ecological part of the problem and forget about the computer science part. We used *automatic code generation* to ensure that researchers need only edit one code file to build a simulator for their particular model. We used a *graph editor* to build the configuration and organise the data required for a particular study. In **3Worlds** therefore, an ecosystem model only requires two files: a specification file organised as a graph, and a computer code file where all relevant ecological processes are written. None of this prevents the modeller from using software libraries, either their own or from a third party, to extend their coding capabilities.

When designing a model, it is important to get quick visual feedback system behaviour when one changes equations or their implementation. **3Worlds** comes with a library of user-interface objects (graphs, maps, time series) that can be freely assembled to adapt outputs to the needs of the researcher.

1.2.8. Model comparison: graphs can be compared

Climate change modelling relies on 19 major general circulation models (GCMs) all based on the same equations. When run with identical datasets (initial data plus forcings), they all yield different

Figure 4. The causal loop within the time loop of **3Worlds**. Within a time step, computations are ordered by causal dependency and processed accordingly. Intermediate state

and graph structure changes follow each causal step.

results. This is expected given the size of their code, but what is troubling is that nobody is able to trace within the code where the differences come from (Lim & Roderick 2009). This problem arises again and again in the modelling literature (e.g. Melilo et al. 1995; Roxburgh et al. 2004). The ultimate reason for this impossibility is that all the knowledge invested into these huge models is represented in computer code, which are very difficult to compare for any but the simplest of models.

3Worlds is an attempt to solve this issue *for the future* (there is nothing we can do for past model code). If models are developed within the standard framework of **3Worlds**, the only thing that needs to be compared among models is their specification file (a graph) and their code file - hundreds to thousands of lines, not more. Everything else is equal. In theory this should facilitate model comparison.

1.2.9. Problem upscaling

Developing a simulator is only a small part of the ecological modelling exercise: once the simulator is ready, it is used as a real ecosystem in *simulation experiments*. Designing and running such experiments is a very important part of the job - if not the most important, as it is the one which gives insight and publishable results.

IBMs are often stochastic, as population rates translate into probabilities at the individual level: e.g., the code has to decide which individuals to delete to satisfy a mortality rate of 10%. This is usually based on drawing random numbers. As a result, every simulation is different even when using identical parameters, and an asymptotic behaviour of the system can only be obtained by running multiple simulations. Fortunately, this is easily parallelized with modern computers.

3Worlds is interfaced with [OpenMole](https://openmole.org/) (<https://openmole.org/>) to provide access to big computing power. Through OpenMole, big simulation experiments can be deployed on networks of computers, grids, or supercomputers.

1.2.10. Portability and performance

3Worlds is written in Java to ensure portability between all operating systems. Its code has been carefully optimised, although generality inevitably comes with some performance cost.

2. Getting started - download and installation

2.1. Basics - what you must know before starting

3Worlds is an application designed to develop and launch simulations of ecosystems. It is highly versatile and can simulate any kind of ecosystem using any kind of mathematical logic.

The application and use of 3Worlds to a particular ecosystem for a particular study case is called a *model*—or, more precisely a *simulation* model. The model must first be specified and developed (this involves writing some code in the java programming language) before it can be executed for a particular case study. This execution is called a *simulation experiment*.

3Worlds comprises two main applications:

- `ModelMaker` , to configure a model;
- `ModelRunner` , to run the model.

Creating a model involves creating a configuration with `ModelMaker` and developing some associated Java code to specify details particular to your model. To do this, you must use the [Eclipse](https://www.eclipse.org/downloads/) (<https://www.eclipse.org/downloads/>) programming software (freeware). Later versions of 3Worlds may support other packages, but at the time of writing, 3Worlds will only work with [eclipse](https://www.eclipse.org/downloads/) (<https://www.eclipse.org/downloads/>).

`ModelMaker` will generate Java code for data structures specific to a model (based on the configuration file you have

developed) and *template* java code for each process you have defined. These process templates are where you enter programming code to implement your model. You only need to write code for your processes and for model initialisation. All else is managed by 3Worlds.

3Worlds is written in [java](#) ([https://en.wikipedia.org/wiki/Java_\(programming_language\)](https://en.wikipedia.org/wiki/Java_(programming_language))), which makes it OS-independent. It can be run on MacOS, Linux or Windows computers.

2.2. Prerequisites

You will need the following software installed on your computer before installing 3Worlds:

- Java JDK (Java Development Kit), version 11 or greater ([Oracle](https://www.oracle.com/technetwork/java/javase/downloads/jdk11-downloads-5066655.html) (<https://www.oracle.com/technetwork/java/javase/downloads/jdk11-downloads-5066655.html>) or [open](http://openjdk.java.net/) (<http://openjdk.java.net/>) version)
- Java fx (graphical user interface library for java: [Oracle](https://www.oracle.com/technetwork/java/javase/overview/javafx-overview-2158620.html) (<https://www.oracle.com/technetwork/java/javase/overview/javafx-overview-2158620.html>) or [open](http://openjdk.java.net/projects/openjfx/) (<http://openjdk.java.net/projects/openjfx/>) version)
- an [Eclipse](https://www.eclipse.org/downloads/) (<https://www.eclipse.org/downloads/>) Java development environment and add the [e\(fx\)clipse plugin](https://www.eclipse.org/efxclipse/install.html) (<https://www.eclipse.org/efxclipse/install.html>) required for JavaFX

2.3. Running ModelMaker

This assumes you have downloaded `tw-<os>.jar` (where `<os>` stands for **linux**, **win** or **mac**, according to your operating system) from the last release on github/3W (<https://github.com/3worlds/3w>).

On your computer, in your home directory:

1. Create a new directory and name it `3w`



This is compulsory: 3worlds will **not** run if the directory does not exist, is named otherwise, or is located elsewhere than at the root of the user home directory. To know your user home directory, refer to your [operating system](https://help.dugeo.com/m/faq/l/245652-where-is-my-user-home-directory) (<https://help.dugeo.com/m/faq/l/245652-where-is-my-user-home-directory>).

2. Copy or move `tw-<os>.jar` into `3w`

3. For *linux users*: set the executable file permission of `tw-<os>.jar` to true (`chmod +x tw-<os>.jar` in a terminal, in the directory where the file is) so that it can be executed

4. Double-click on `tw-<os>.jar`. This should launch the ModelMaker application

5. If this doesn't work, open a terminal and type `java -jar tw-<os>.jar`. This should launch the ModelMaker application

It is preferable to run the software from a terminal window. This way, any errors that may arise will be displayed as text in the terminal



to develop your model-specific code, you will need to setup a Java development environment as shown in Section 2.4.

2.4. Setting up a Java development environment for the user code

2.4.1. Setting up the Eclipse Integrated Development Environment (IDE) for 3Worlds

This assumes you have downloaded `tw.jar`.

1. If not yet done, install [Eclipse](https://www.eclipse.org/downloads/) (<https://www.eclipse.org/efxclipse/install.html>)! (don't forget [e\(fx\)clipse](#))
2. Create at *workspace* (a working directory for Eclipse - Eclipse will ask for it when launched). e.g., <*my_workspace*>
3. Within Eclipse, create a *project*:
 - Select menu `File → New → Java project`; this opens a dialog box
 - In the dialog box, type a project name (e.g. <*my_project*>)
 - Click the `Finish` button
4. Import 3Worlds dependencies:
 - Select menu `Project → Properties`; this opens a dialog box
 - In the dialog box, select `Java Build Path`
 - Select the `Libraries` tab
 - Select `ClassPath`
 - Click on the `Add external JARs...` button; this open a file selection dialog box
 - In the file selection dialog box, browse and select `tw-<os>.jar`
 - Click the `Apply and Close` button

2.4.2. Running ModelMaker from Eclipse

ModelMaker can be run from Eclipse or as a standalone application since it is included in the `tw-<os>.jar` library required to develop the user code.

- In the `package explorer` window, expand the `Referenced libraries` entry
- Right-click on the `tw-<os>.jar` entry, select `Run as → Java Application`. This opens a dialog box
- In the dialog box, type *Main*
- In the list of matching items, select `Main - au.edu.anu.twuifx.MMmain` and click `OK`
- If warning errors appear, click `Proceed`. This launches the ModelMaker application

2.4.3. Linking user code with model configuration

This requires the following actions:

1. In ModelMaker,
 - create or open a *3Worlds* project (`Projects` entry of the main menu)
 - select `Preferences → Java Project → Connect...`. This opens a dialog box with a file selector
 - select the root directory of the *Eclipse* project as created above (e.g. <*my_workspace*>/<*my_project*>)

This operation tells `ModelMaker` to generate its code into the user java project. When you want to edit your code in eclipse, you must first **refresh** the eclipse project:

2. In Eclipse,
 - select the project name at the very top of the `package explorer` window
 - right-click on it and select `Refresh`
 - or, alternatively: press the `F5` key



You don't **have** to do this. We provide it as a facility if you want to run ModelMaker from eclipse rather than directly for some reason of your own.

2.4.4. Debugging and testing user code

The user code, first generated by `ModelMaker` and further edited by the user, can be run using `UserCodeRunner.java`. This class can be found in the default `src` directory and was created when linking this project with the 3Worlds project (cf above). It requires three command line arguments (we assume that you know how to setup and run a Run Configuration in Eclipse):

- an instance number (more about this later); leave this at '0' for now.
- the name of the directory of the 3Worlds project as created by `ModelMaker` (e.g. `project_test_model9_2019-09-05-08-50-20-458`). This project directory is located under the `3w` directory automatically created by `ModelMaker` as its working directory
- some optional settings to switch on debugging logs.

With this, the user code should be executed as a test simulation by `UserCodeRunner`.



Further edits and modifications of the configuration can be made in `ModelMaker`, but don't forget to keep the Eclipse project content synchronized with the ModelMaker project by refreshing the Eclipse project as often as necessary.

3. Configuration reference: creating and editing a model with ModelMaker

3.1. General concepts: structure of a 3Worlds configuration

3.1.1. A tree structure...

The configuration of a 3Worlds *simulation experiment* is organised as a tree (Figure 5). Each tree *node* specifies a subset of the parameters of the whole configuration. Each *node* has *child nodes* linked through a *tree edge*, so that large pieces of configuration can be broken down into the relevant details. At each level of this hierarchy, *properties* can be attached to *nodes*.

Nodes have a *label* and a *name* that are displayed in the `ModelMaker` interface as `label:name`:

- The *label* specifies what role this particular *node* plays in the whole configuration. For example, the *node* labelled `experiment` is used to configure a *simulation experiment*.
- The *name* is used to differentiate *nodes*. It must be unique over the whole configuration tree. By default, `ModelMaker` generates unique names (by adding a number to the name if replicates are found).

The *root node* of a configuration is always labelled `3worlds`.

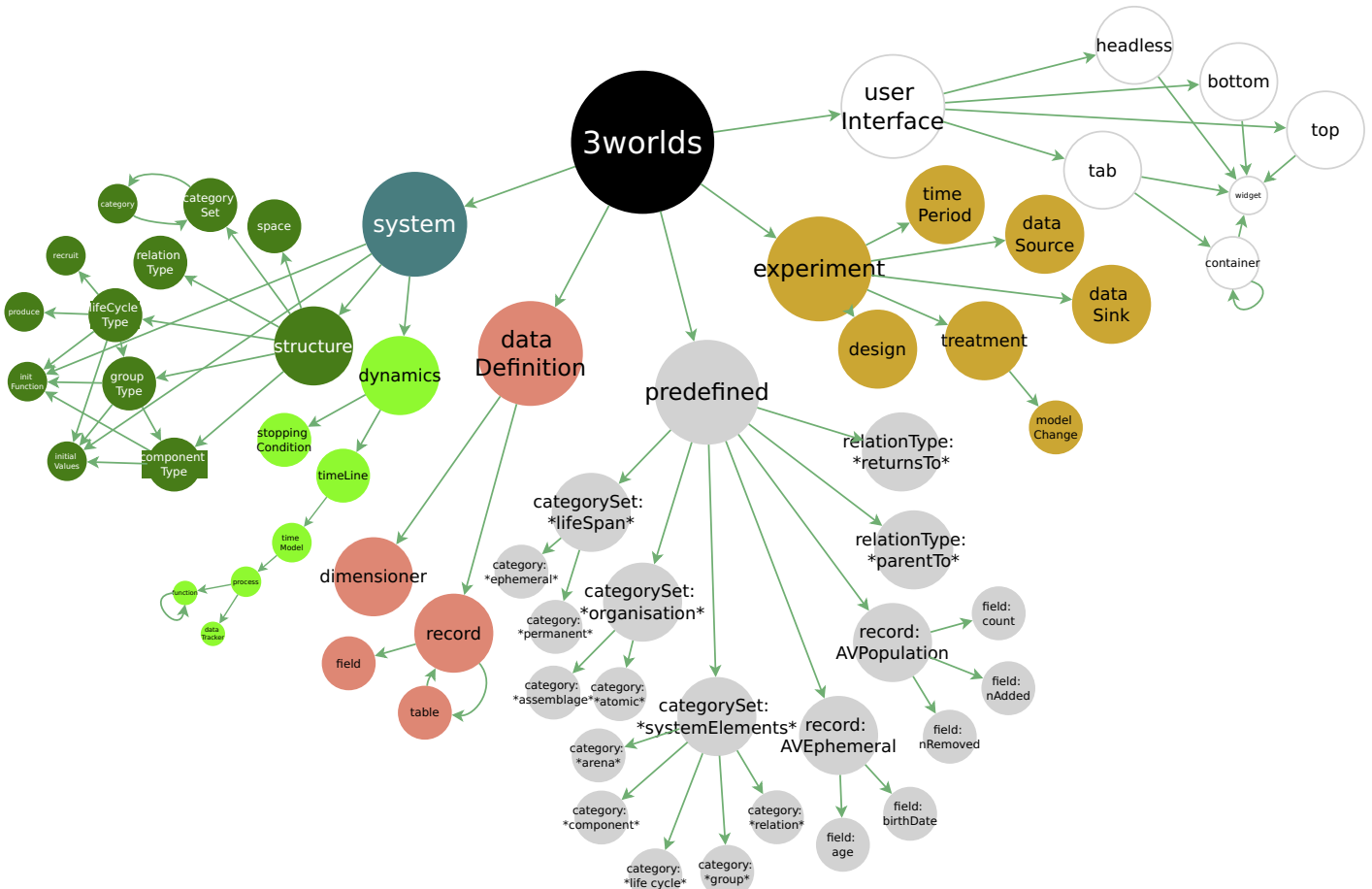


Figure 5. Tree structure of a simulation experiment configuration in 3Worlds

The configuration tree is stored in a file in a specially designed text format, ending with the extension `.ugt` or `.twg`. Such files are produced by `ModelMaker` and can then be exchanged and imported into `ModelMaker` via the `Projects → Import...` menu entry. Their format is human-readable, but they must **never** be edited with another software than `ModelMaker` - **the risk is to corrupt all your configuration** and be unable to run it (or even edit it with `ModelMaker` again).



NEVER edit the 3worlds configuration `.ugt` file 'by hand'! You may make it unreadable to `ModelMaker` and lose all your work.

Each node in the configuration has a particular meaning for `ModelRunner`: the configuration must comply with certain rules and constraints, the first one being the particular set of *nodes* that have been designed and appear on Figure 5. The detailed meaning of all *nodes* and their *properties* is described in Section 3.3.

3.1.2. ...with cross-links

Actually, the 3Worlds configuration is not strictly hierarchical (Figure 6): according to their role in `ModelRunner`, some configuration *nodes* need to gather information from other parts of the configuration tree. This is done by allowing for some cross-reference *edges* to be defined, that overlay with the strict hierarchical structure of the tree. As a result of these cross links, the whole configuration is a *graph* rather than a tree.

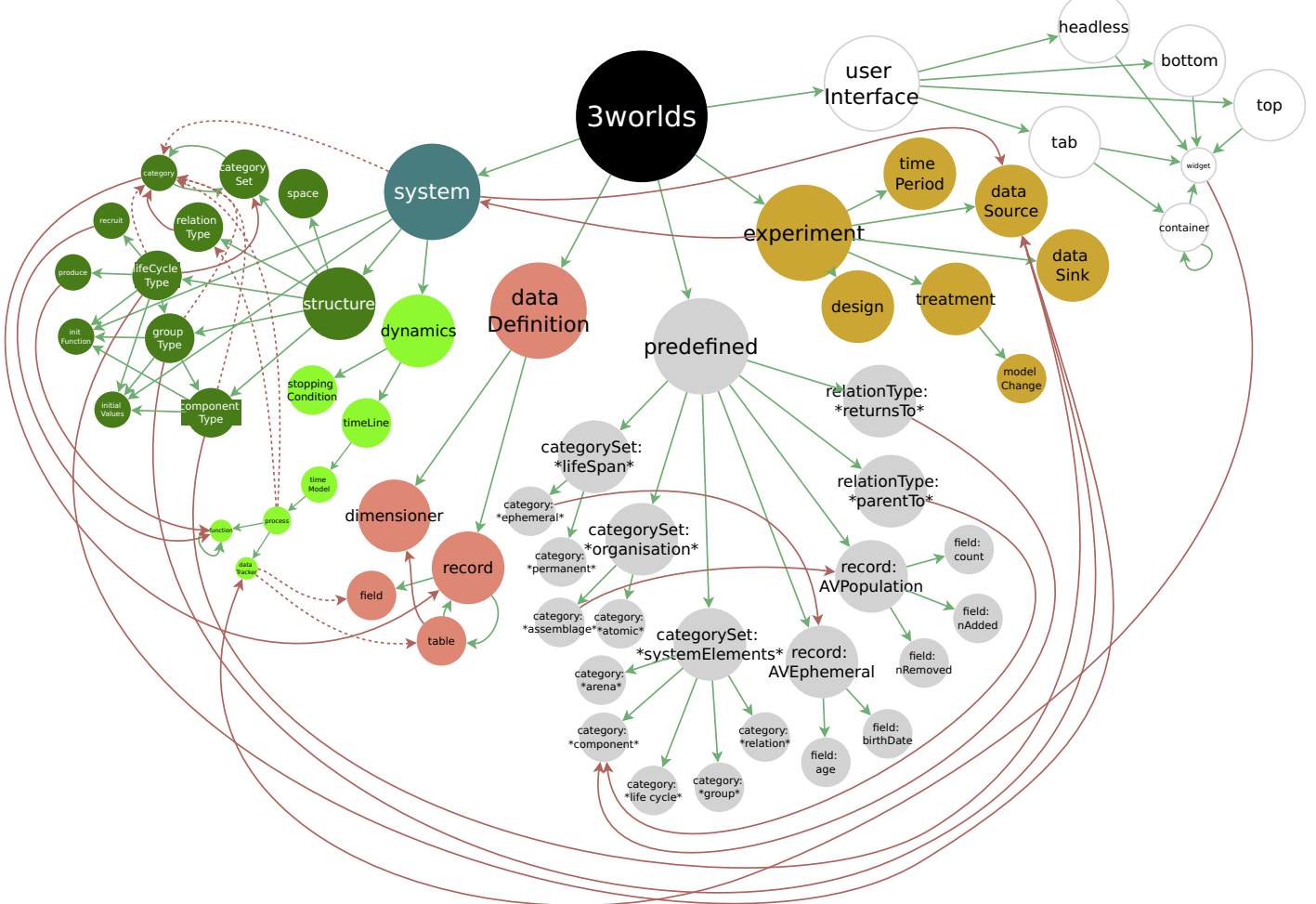


Figure 6. Tree structure of a simulation experiment configuration in 3Worlds, showing the cross-links between tree nodes (NB: not all cross-links are drawn)

Edges representing cross links have a *label*, *name*, and may have *properties* just as *nodes* do. The detailed meaning of all cross-reference edges and their properties is described in Section 3.3.

3.1.3. What ModelMaker does for you

ModelMaker knows the details of the configuration constraints. It facilitates the design of a configuration by only letting you add the *nodes*, *edges* and *properties* that will produce a valid, runnable configuration file. During the configuration building process, it constantly checks the validity of the graph and reports any errors or missing parts in its log panel. ModelMaker is far more than a nice visual editor producing a graph: a configuration graph produced with ModelMaker is **guaranteed** to run with ModelRunner because of all these internal consistency and validity checks.

3.2. Using ModelMaker: software interface and functioning

ModelMaker has the traditional main menu bar, a variety of mouse/keyboard operations and three regions for the construction of the model specifications graph (Figure 7):

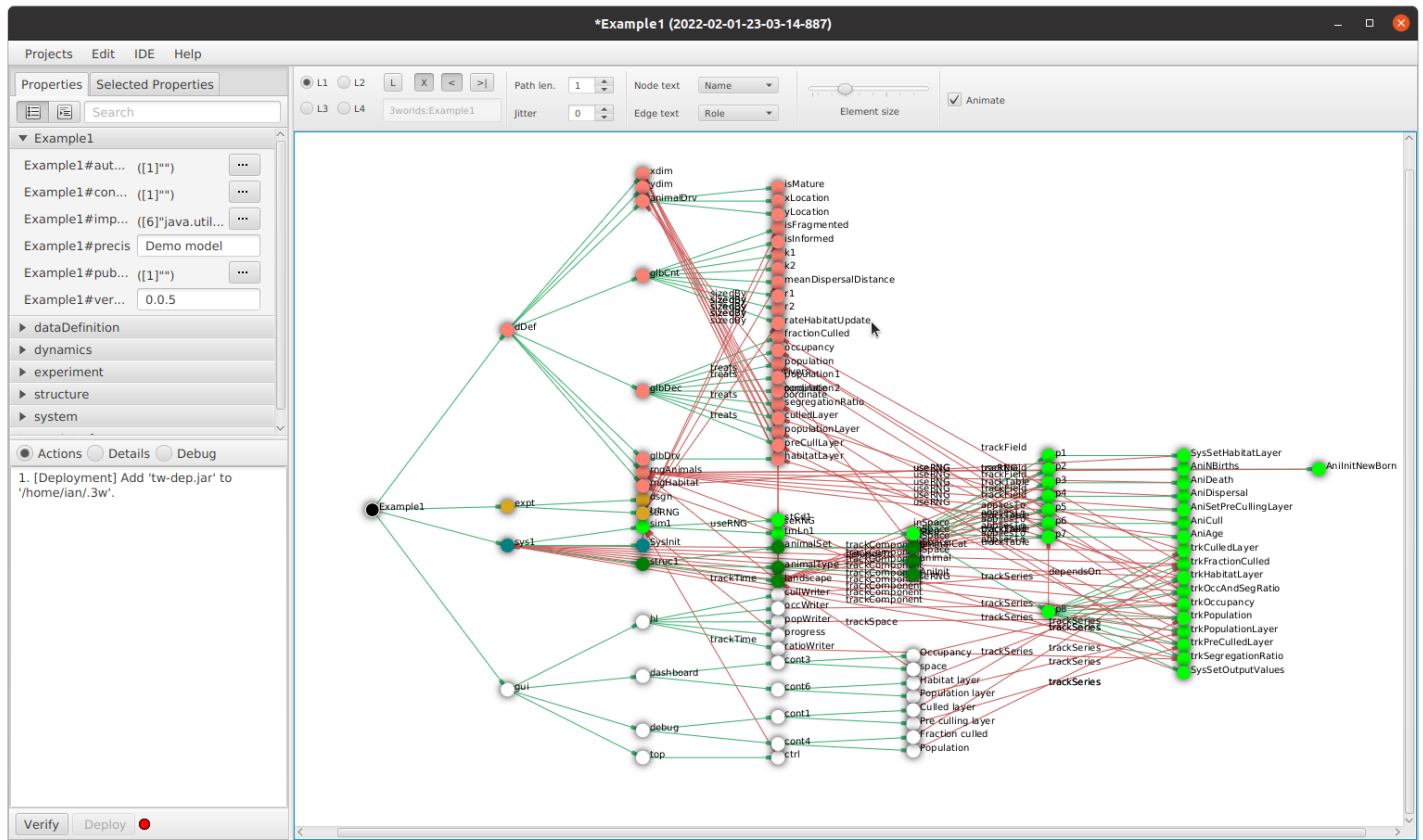


Figure 7. The appearance of ModelMaker displaying an example project.

1. Right-hand side: **Graph display** window and associated tool bar for graph construction.
2. Top left: **Property editors** for node and edge properties.
3. Bottom left: **Message pane** to display actions that are required to ensure the specifications are valid.

The main window title is the currently open project and any linked Java project associated with it (<3Worlds project name>(<date>) <o-o-o> <Java project name>). Whenever the specifications are edited, an asterisk is prefixed to the window title and cleared upon saving.

3.2.1. Main menu

Project

New

Creates a new project from either a library of **Templates**, **Tutorials**, **Model library** or **Test cases** (Figure 8). You will be prompted for a unique name for the project. The first letter of this name must be upper case as it will eventually be used as the name of a Java class when the project successfully compiles. A sub-directory with the name and creation date will be created under 3w. Once created, the project name and its creation date appear in the window title and the graph root node name is that entered at the prompt. *ModelMaker* will automatically compile the project and list any actions required in the **message pane**.

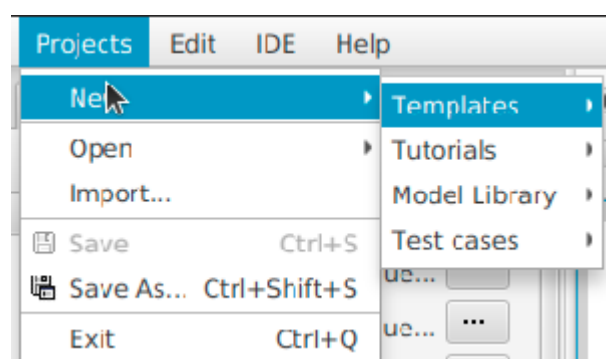


Figure 8. Options for creating a new project.

Open

Opens any project currently in the 3w directory (Figure 9). Projects are listed in alphabetical order.

Import

Imports a tree-graph file from disk (*.utg). Imports are only possible if the tree-graph has a single root. You will be prompted for a unique name for the import.



Tree-graphs with more than one root cannot be imported.

Save

Saves the configuration graph and its associated layout file (`.layout.utg`) to disk.

Save as

Saves the project under a new name in a new directory within `3w`. If the project was linked to a Java project, the new project will maintain this link and generate a new Java file with this name.

Exit

Exits *ModelMaker* after a prompting to save any unsaved work. On exiting, a project specific preferences file is saved automatically (`MM.xml`), recording the current appearance of *ModelMaker*.

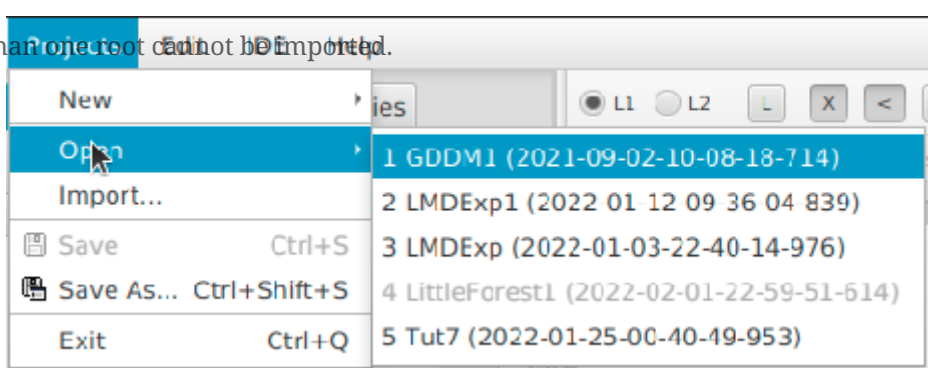


Figure 9. The Open project menu choice.

Edit

The Edit menu provides the Undo/Redo options (Figure 10). All graph editor operations are able to be undo or redone except property edits. The undo/redo system creates temporary history files during a session. These are deleted when the project is closed.

IDE

Import snippets

Enabled when a project is linked to a Java project. This option imports method code from Java into any `snippet` properties in the specifications (`functionSnippet` of the `Function` and `InitFunction` nodes and the `importSnippet` property of the specification root node).



Figure 10. The edit menu choice.



currently, importing code containing double quotes is not supported.

Clear snippets

Clears the Java code in the configuration's `functionSnippet` and `importSnippet` properties. Functions requiring return values will have default return statements inserted.

Link to IDE

Allows connecting to or disconnecting from an external Java project via an Integrated Development Environment. When connecting to a Java project, the java project must have imported `3w/tw<os name>.jar` as an external jar to its class path (Section 2.4.1). Upon first connecting to a Java project, a Java class called `UserCodeRunner.java` is added to the default package. Use this as a main class to debug your model code. If the specifications are valid, additional Java classes are created. One of these has the same name as the `3Worlds` project. Edit this file in an Integrated Development Environment (IDE) to begin developing your model. At the time of writing, only the *Eclipse* IDE is supported for this purpose (Section 2.4).

If the creation times of Java source files and their associated class files become inconsistent, a message to this effect will appear in the **message pane**. In such cases, refresh or clean the Java project.



To compile models, the Java Development Kit (JDK) must be installed on your computer (Section 2.2).



Use 'Refresh' in the IDE to keep *ModelMaker* and the Java project synchronized.

Help

About ModelMaker

A dialog with a succinct description of the purpose of *ModelMaker*. If you're using *ModelMaker* for the first time, it's a good idea to at least read this overview.

3.2.2. Graph display

Construction of the model specification graph takes place in the **Graph display** window by adding or removing nodes and edges (*cross-links*). Above the graph display is a tool bar to enable various display options (Figure 11).

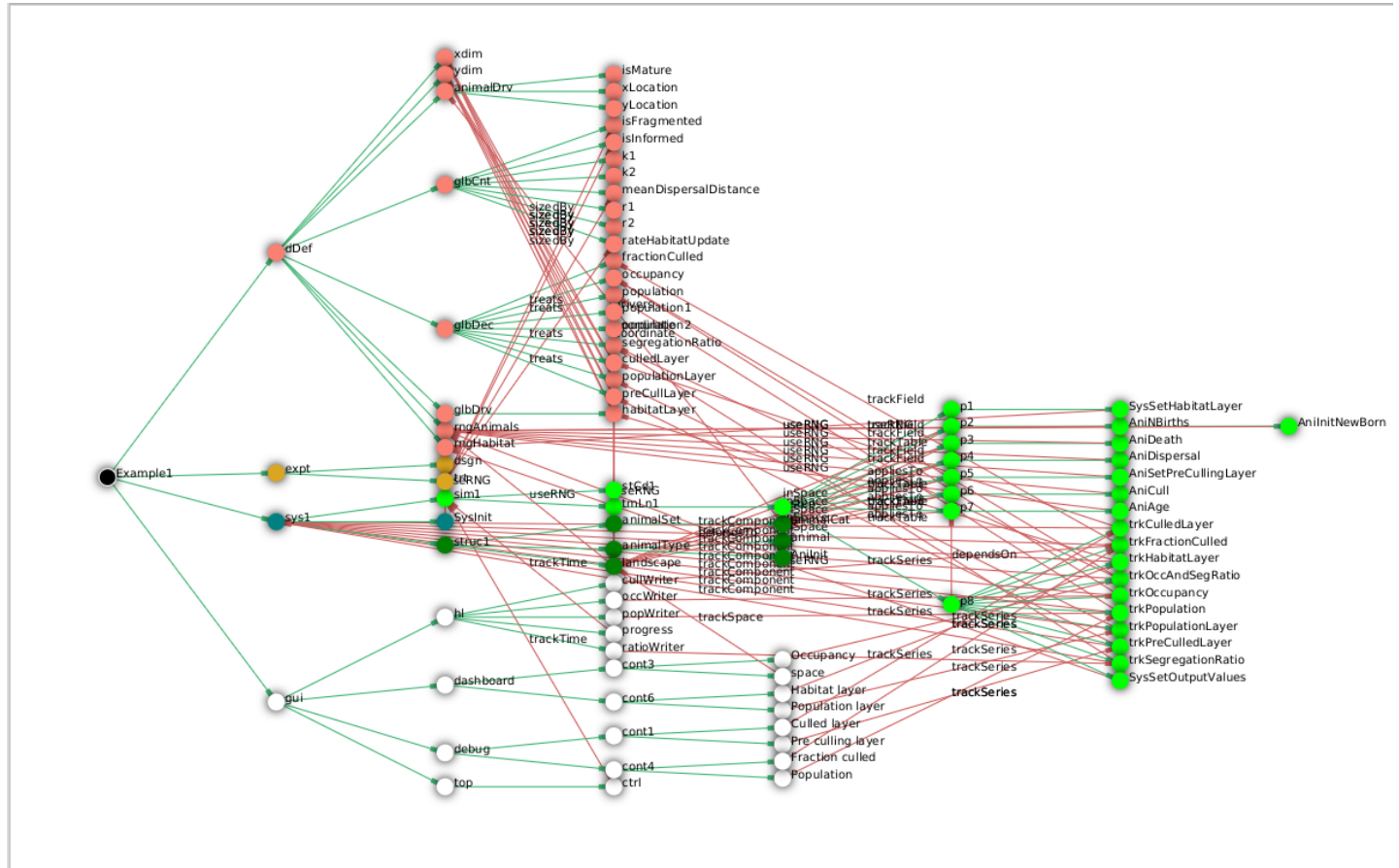


Figure 11. The Graph display window and tool bar. The graph is both a Tree graph and a directed network graph. The tree graph is shown with green parent-child edges. The directed network graph is shown with red, named edges.

The graph is both a tree graph (with a single root) and a directed network graph. The tree structure is shown with green parent-child edges with a thickened end at the child node. The directed network graph is shown with red, named edges with a thickened end at the 'end node'.

Node colours are based on a schema that, for the most part, colours all nodes within a sub-tree the same. The colours are not editable.

Tool bar

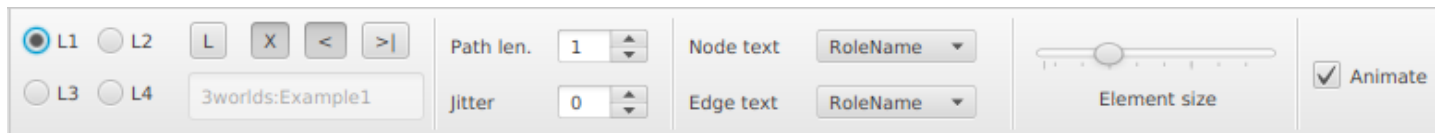


Figure 12. Tool bar options for managing the graph display.

1. Layout selection: The four radio buttons (**L1**, **L2**, **L3**, **L4**) select one of four layout algorithms:

- i. L1 - ordered tree. The nodes are listed vertically in alphabetical order.

- ii. **L2** - radial tree. The radius decreases as the path distance from the root node increases.
- iii. **L3** - radial tree. The radius remains constant.
- iv. **L4** - spring graph. A force directed layout. Parent-child edges (green) and treated in the same way as *cross-links* (red).

2. Layout options

- i. **L**: Applies the current layout options.
- ii. **X**: Toggles the *cross-links*.
- iii. **<**: Toggles the parent-child edges.
- iv. **>|**: Moves any isolated nodes to the right-hand side of the window.
- v. **Current layout root**: The name of the node used for the display root of the graph. The default is the black 3worlds root node. This can be changed by right-clicking on any node while holding down the `Ctrl` key. The display root node is also indicated by a black circle. This option only applies to tree layouts.

3. Miscellaneous

- i. **Path len.**: Sets the path length when the *show local graph* display mode is activated (by moving the mouse over a node while holding down the `Shift` key).
- ii. **Jitter**: Sets the amount of random displacement of a node when applying a layout. This is useful to separate text and/or lines drawn over each other.

4. Text options:

- i. **Node text**: Display the node label and name or any combination of the two including no text.
- ii. **Edge text**: Display the edge label and name or any combination of the two including no text.

5. Element size

Animate: When checked, changes to the shape of the graph by either collapsing/expanding sub-trees or applying a layout function are animated. Switch this off for large graphs - the animation routine can quickly become overloaded and the resulting animation jumpy.

Keyboard and mouse functions

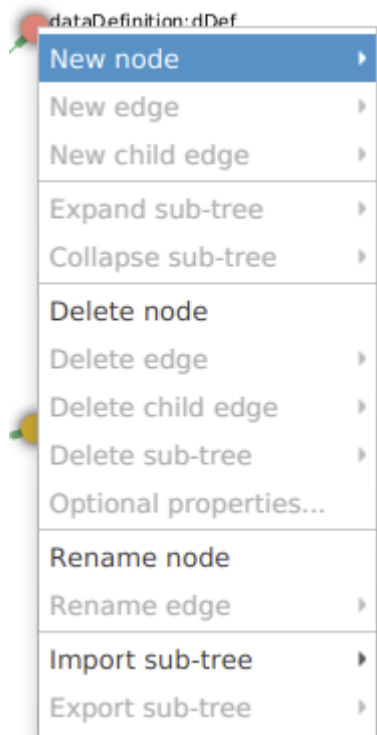
- i. **Pan**: Drag the mouse anywhere other than on a node to pan the graph drawing surface. This assumes the display size is larger than the window.
- ii. **Drag node**: Left-click on any node to change its position. Nodes are shown in red when the mouse passes over them.
- iii. **Show local graph**: Hold down the `Shift` key while moving the mouse over a node. The *path length* for this feature is set in the tool bar.
- iv. **Show node properties**: Right-click on any node to display its properties in the *Selected properties* property editor.
- v. **Zoom**: Hold down the `Ctrl` key while moving the mouse wheel to zoom in and out of the display.
- vi. **Pop-up edit menu**: Right-click on any node to see the edit options available for that node.

Pop-up edit options

This menu is the principle way in which the specification graph is constructed (Figure 13). Note that the `predefined:catoregories*` sub-tree cannot be edited except for any 'in-edges' that may be allowed from outside this sub-tree.

- i. **New node**: Displays a list of valid node labels that can be children of this node.
- ii. **New edge**: Displays a list of valid edges (*cross-links*) from this node to another existing node.

- iii. **New child edge:** Displays a list of (parentless) nodes that can be valid children of this node.
- iv. **Expand sub-tree:** Displays a list of sub-trees that can be expanded from this node. After expanding, any properties of these nodes and edges will appear in the property editor.
- v. **Collapse sub-tree:** Displays a list of sub-trees that can be collapsed from this node. After collapsing, any properties of these nodes and edges will be removed from the property editor.
- vi. **Delete node:** Deletes this node. Note that the `3Worlds` root node cannot be deleted.
- vii. **Delete edge:** Displays a list of 'out-edges' from this node that can be deleted.
- viii. **Delete child edge:** Displays a list of child nodes that can be orphaned. Note that the specifications are not valid until all child nodes have parents.
- ix. **Optional properties:** Displays a dialog with any optional properties that can be added or removed from this node.
- x. **Rename node:** Changes the node name to some other unique name. Note that the name of the root node cannot be changed.
- xi. **Rename edge:** Displays a list of 'out-edges' whose name can be changed to some other unique name.
- xii. **Import sub-tree:** Displays a list of valid child node labels that can form the root of a sub-tree read from file.
- xiii. **Export sub-tree:** Displays a list of the current child nodes of this node that can be exported to file as a sub-tree.



*Figure 13. The pop-up menu.
Available options depend
on the node selected.*

3.2.3. Property editors

Properties		Selected Properties
		Search
Prj1#authors	([1]***)	...
Prj1#contacts	([1]***)	...
Prj1#precis		
Prj1#publication	([1]***)	...
Prj1#version		
ctrl#order	0	
dsgn#type	singleRun	▼
tmLn1#longestTi...	UNSPECIFIED	▼
tmLn1#scale	ARBITRARY	▼
tmLn1#shortestTi...	UNSPECIFIED	▼
tmLn1#timeOrigin	0	...

Figure 14. Property editor showing properties listed by name.

Properties		Selected Properties
		Search
tmLn1#longestTimeUnit	UNSPECIFIED	▼
tmLn1#scale	ARBITRARY	▼
tmLn1#shortestTimeU...	UNSPECIFIED	▼
tmLn1#timeOrigin	0	...

Figure 15. Property editor showing only properties of a selected node (here a TimeLine).

There are two property editors available on separate tabs: one (*Properties*) displays all *editable* properties of the currently displayed nodes and edges (Figure 14); and a second (*Selected properties*) shows *all* properties of a selected node (left-click on any node: Figure 15).

The *Properties* editor has two buttons; one to display properties by category (Figure 16; in this case this means by sub-tree) and the other to display properties by name (Figure 14). There is also a search field that can be used to show only those properties that

match the search text.

Properties are only shown for nodes that are currently visible (expanded) in the graph.

3.2.4. Message pane

The **message pane** displays information about what is required to make the model specifications valid. It grows and shrinks during construction depending on the state of the graph. Information is presented in three levels of detail: action, details and debug (Figure 17). Generally, the 'action' level is all that is required.

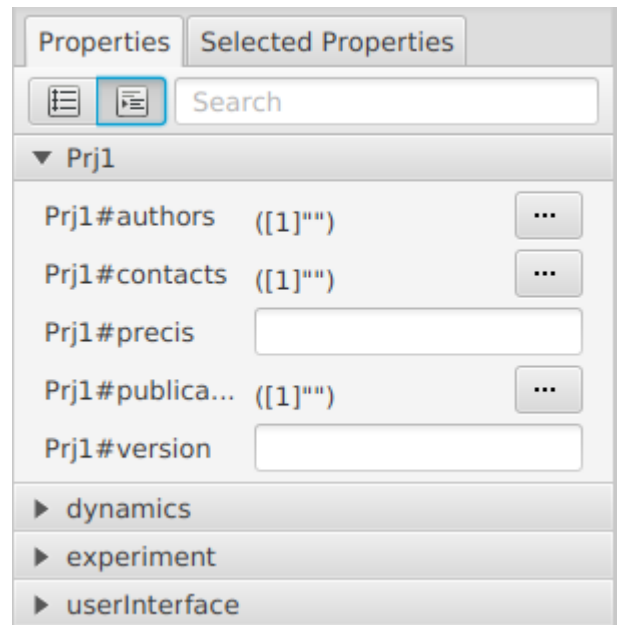


Figure 16. Property editor showing properties listed by category.

The screenshot shows three panels of the Message pane with different display levels:

- Actions:** Shows two numbered messages:
 - [Node] system:sys1: Add edge to one child of 'categorySet:*organisation*'.
 - [Node] timeline:tmLn1: Add node 'timer:' to 'timeline:tmLn1'.
- Details:** Shows detailed logs for each message:
 - Action: Add edge to one child of 'categorySet:*organisation*' Constraint: Expected edge to one child of 'categorySet:*organisation*' but found none. Query class: EdgeToOneChildOfQuery Constraint Specification: mustSatisfyQuery:arenaHasOneCategoryOf organisationSpec Query item: system:sys1
 - Action: Add node 'timer:' to 'timeline:tmLn1'. Constraint: Expected 1..* child nodes with reference 'timer' from parent 'timeline:tmLn1' but found 0. Target: timeline:tmLn1 Constraint Specification: hasNode:timeLineSpec
- Debug:** Shows extremely detailed logs for each message:
 - Action: Add edge to one child of 'categorySet:*organisation*' Constraint: Expected edge to one child of 'categorySet:*organisation*' but found none. Category: [Node] Category class: NODE_QUERY_UNSATISFIED Query class: EdgeToOneChildOfQuery Constraint Specification: mustSatisfyQuery:arenaHasOneCategoryOf organisationSpec Query item: system:sys1=[↑3worlds:Prj1 ↓dynamics:sim1 →category:*permanent* →category:*arena*]
 - Action: Add node 'timer:' to 'timeline:tmLn1'. Constraint: Expected 1..* child nodes with reference 'timer' from parent

Figure 17. Message pane displaying three different message formats.

Messages are displayed in alphabetical order of the Action text no matter what the display level. At the bottom of the **message pane** are two buttons described at the end of this section.

- Action:** This level indicates what action must be taken for the specifications to comply with the 3Worlds archetype. Its format is: a) the message category (usually **[Node]**, **[Edge]** or **[Property]**); b) the target (the label:name of a node or edge or a property name) and the action message (Figure 17).
- Details:** This level shows the action message and the specification constraint that is unsatisfied. The information provided varies with the message but in general includes (Figure 17):
 - Message number. As noted, the message order remains consistent no matter the display option.
 - The action message: usually in the form of "Do this or that".
 - The specification constraint: usually in the form of "Expected this but found that".

D. The Java class of the query issuing the message.

E. The node in the 3Worlds archetype that defines the application of this query class.

F. The target element (node, edge, property or some other item class)

iii. **Debug:** This level shows all information available for this message including a detailed description of the target and archetype elements. It is rarely informative unless you are a contributor to the development of 3Worlds (Figure 17).

Verify and Deploy

i. **Verify:** Verifies that the specification graph conforms with the archetype and compiles the Java code. This process takes place in the background any time the graph is edited. Therefore, there is rarely any need to click the compile button.

ii. **Deploy:** This button is enabled (traffic light green) whenever the **message pane** is clear. When clicked, *ModelRunner* is launched to run the model with the specified experiment design.

3.3. Configuration options: reference

In this section,

- node and edge labels are indicated in **bold**
- text in triangular brackets (`<>`) mean a user-defined value is expected; the text usually specifies what kind of value is expected (e.g. `<name>` for a name, `<int>` for an integer number, etc.). If the text is required, it will be underlined, otherwise it is optional
- a *multiplicity* in curly braces {} tells how many times the item may appear in a configuration:

{1} exactly one item is required

{0..1} the item is optional, i.e. one or zero is required

{1..*} one to many items are required

{0..*} any number of items is possible

- levels in the tree hierarchy are indicated by slashes / .

- a column : separates a node label from its name.

3.3.1. The 3Worlds node

`/3worlds:<name> {1}`

This node is the root of any 3Worlds configuration file (Figure 18). The name will appear in **ModelMaker**'s main window title, in the **project directory name** and in the **configuration graph file**. The name is requested and set when creating a new project (**Projects>New** menu entry in **ModelMaker**).

Properties for 3worlds

authors The list of this project author names.

contacts The list of the author contacts (e.g. e-mail addresses), in the same order as the author names (1-1 match).

precis A short description of the model contained in this project.

publication A list of bibliographic references in relation with the project.

`built-by` The user name under which this project was constructed and saved, together with the date of saving. This property is automatically set by ModelMaker.

`version` A user-defined version identifier for this project.

All these properties appear in the about box of `ModelRunner` and are saved in the ODD file describing the model.

3.3.2. The `system` node

`/3worlds/system:<name> {1..*}`

This node and its sub-tree contains all the ecological concepts used to define a simulation model: what entities are modelled, what biological processes apply to them, at what time step they should run. The name is used to differentiate models as a simulation experiment may involve more than one model.

A 3Worlds ecosystem is constructed as a population of 'biological entities', called *system components*, which establish interactions called *system relations*. Components and relations have *variables* that may change value over time, according to various *ecological processes* attached to components. In other words, an ecosystem is represented as a *dynamic graph* which changes state and structure over time. To reflect this, `system` has two sub-trees called `structure` and `dynamics`.

For very simple models, where the modelled system is not further subdivided into components, the structure sub-tree is not required. In this case, the `system` node will be used as a unique 'component' describing the system (cf tutorial: logistic).

Cross-links for system :

`belongsTo → category:<name> {1..*}`

This link tells to which categories the `system` node belongs. The categories must not belong to the same category set. If there are nested categories, membership is inherited. The categories targets of these links must be specific to the `system`, i.e. no `componentType`, `groupType` or `lifeCycleType` should refer (`belongsTo` cross-link) to any of them.

`belongsTo → category:*arena* {1}`

This link to the predefined category `*arena*` is required in all models (Section 3.3.8). The arena is 'the place where things happen', i.e. it describes the part of the system where components interact. If no components are modelled, then it is the `system`.

`belongsTo → category:*atomic*|*assemblage* {1}`

This link to either the predefined category `*atomic*` or `*assemblage*` is required in all models. Only if the `system` is very simple (no components, no groups, no life cycles) should `*atomic*` be selected. It means the `system` has no further subdivision. In any other case (i.e. as long as there is a `structure` sub-tree), the `system` should be an `*assemblage*`. Assemblages have three automatic variables: the group size (number of components) `count`, the number of new components added during last causal step `nAdded`, and the number of components deleted during the causal step `nRemoved`.

`belongsTo → category:*permanent* {1}`

This link to the predefined category `*permanent*` is required in all models. It specifies that the `system` is going to stay forever (permanent) during a simulation.

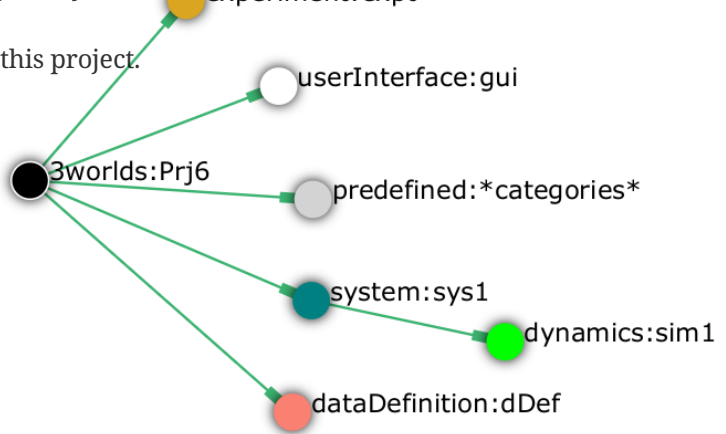


Figure 18. The base tree of any 3Worlds configuration

```
loadFrom → dataSource: {0..*}
```

These optional links refer to data sources from which to read initial values of system-level descriptors (Section 3.5.5).

Optional child for system:

```
/3worlds/system/initFunction:<name> {0..1}
```

This node declares a function of type `SetInitialState` to set or transform the initial values of system-level descriptors (Section 3.5.2).

3.3.3. The *system/structure* node

```
/3worlds/system/structure:<name> {0..1}
```

This node and its sub-tree contains the description of the system component and relation types. It is based on the concept of *category*. A category is a set of components with an identical description, e.g. the same set of state variables or the same growth function.

The *category* and related concepts: specification of groups of entities

3Worlds uses these concepts to specify and generate the ecological entities manipulated during a simulations.

Category

```
.../categorySet/category:<name> {1..*}
```

A *category* is simply a name attached to a set of objects sharing common properties (Section 1.2.3). These common properties are descriptors (Section 1.2.2), and dynamic behaviours (or *processes*). Categories, grouped into `categorySet`s, constitute a user-defined *classification* of *system component types* relevant for a particular model.

Descriptors consist in *drivers*, *decorators*, *constants* and *automatic variables*:

drivers

are *independent* variables (numbers, text, logical values) that characterize the state of a system component at an instant in time (e.g. biomass, age, sex, social status...). They will vary during a simulation. 'Independent' means that these variables cannot be computed from each other within a simulation step, they also depend on their previous *causal* step values. For example, if some category is described by drivers *age* and *biomass*, neither of them can be computed from the other: *age* is computed from its previous value by adding the time step duration; *biomass* is computed in some other way probably involving its previous value. The values of drivers are carried over from a simulation step to the next, i.e. they *drive* the dynamics of the system.

decorators

are *dependent* variables that are computed from drivers and other decorators within a simulation step. For example, *leaf area* could be computed as some constant times *biomass*: its value is completely determined by the value of *biomass*. As such, decorators are not independent from the drivers and are used only for convenience in computations or output display. They are not carried over from a simulation step to the next, they are always automatically reset to zero at the end of each *time* step.

constants

are values (numbers, text, logical values) that do not change during a simulation. For example, *sex* or *number of legs* are not going to change during the lifetime of an animal. They are set at birth and never changed after.

automatic variables

are values internally managed by 3Worlds and available *read-only* for computation. For components representing individuals, these are *age* and *birth date*. For components representing populations, these are *number of individuals*, *number of newly created individuals*, *number of deleted individuals*.

The exact data structures for descriptors are specified under the `dataDefinition` node (Section 3.3.5) and linked to the category through the following:

Cross-links for category :

```
drivers → record:<name> {0..1}  
decorators → record:<name> {0..1}  
constants → record:<name> {0..1}
```

These links tell which data structure in the `dataDefinition` node (Section 3.3.5) is used to store *drivers*, *decorators* and *constants*.

Automatic variables do not need to be specified - because they are automatic.

A category may be defined with no drivers, decorators or constants.

There are *predefined categories* in 3Worlds under the node `/3worlds/predefined` (Section 3.3.8), used to specify particular components of the system. They will be explained in time.

CategorySet

```
/3worlds/system/structure/categorySet:<name> {1..*}
```

Some categories must be exclusive of each other: for example, an ecological entity is either a plant or an animal, but can't be both. For this reason, *exclusive* categories are grouped into `categorySets`. A `categorySet` is a *set of mutually exclusive categories* (mathematically, a *partition* (https://en.wikipedia.org/wiki/Partition_of_a_set)). Categories can be nested, by simply defining a `categorySet` as a child of a `category`:

```
.../category/categorySet:<name> {0..*}
```

RelationType

```
/3worlds/system/structure/relationType:<name> {0..*}
```

A `relationType` is just a name representing a meaningful link between two kinds of categories. It is specified by giving it a name and cross-linking it to the relevant categories with `fromCategory` and `toCategory` cross-links. Note that a `relationType` can link more than one 'from' categories to more than one 'to' categories if required. `RelationType`s are used to implement specific processes acting on ecological entities (for example, a predation process). Notice that relations are *directed* (i.e., they have a *start* and an *end* with potentially different roles).

Cross-links for relationType :

```
fromCategory → category:<name> {1..*}
```

This link tells which categories are at the start of the relation.

```
toCategory → category:<name> {1..*}
```

This link tells which categories are at the end of the relation.

Properties for relationType :

`lifespan` This property specifies if this type of relation will stay attached to its `systemComponents` during all their life, or may get created and deleted during their lifespan.

possible values:

`permanent` relation stays as long as both its ends stay (default value)

`ephemeral` relations are created during a simulation by the means of a `relateTo` function, and kept or deleted by means of a `maintainRelation` decision function. If no `maintainRelation` function is provided, then the relation only lasts for one simulation step.

Example: a category tree

On this diagram (generated with `ModelMaker`), hierarchical links are in green and cross-links are in red.

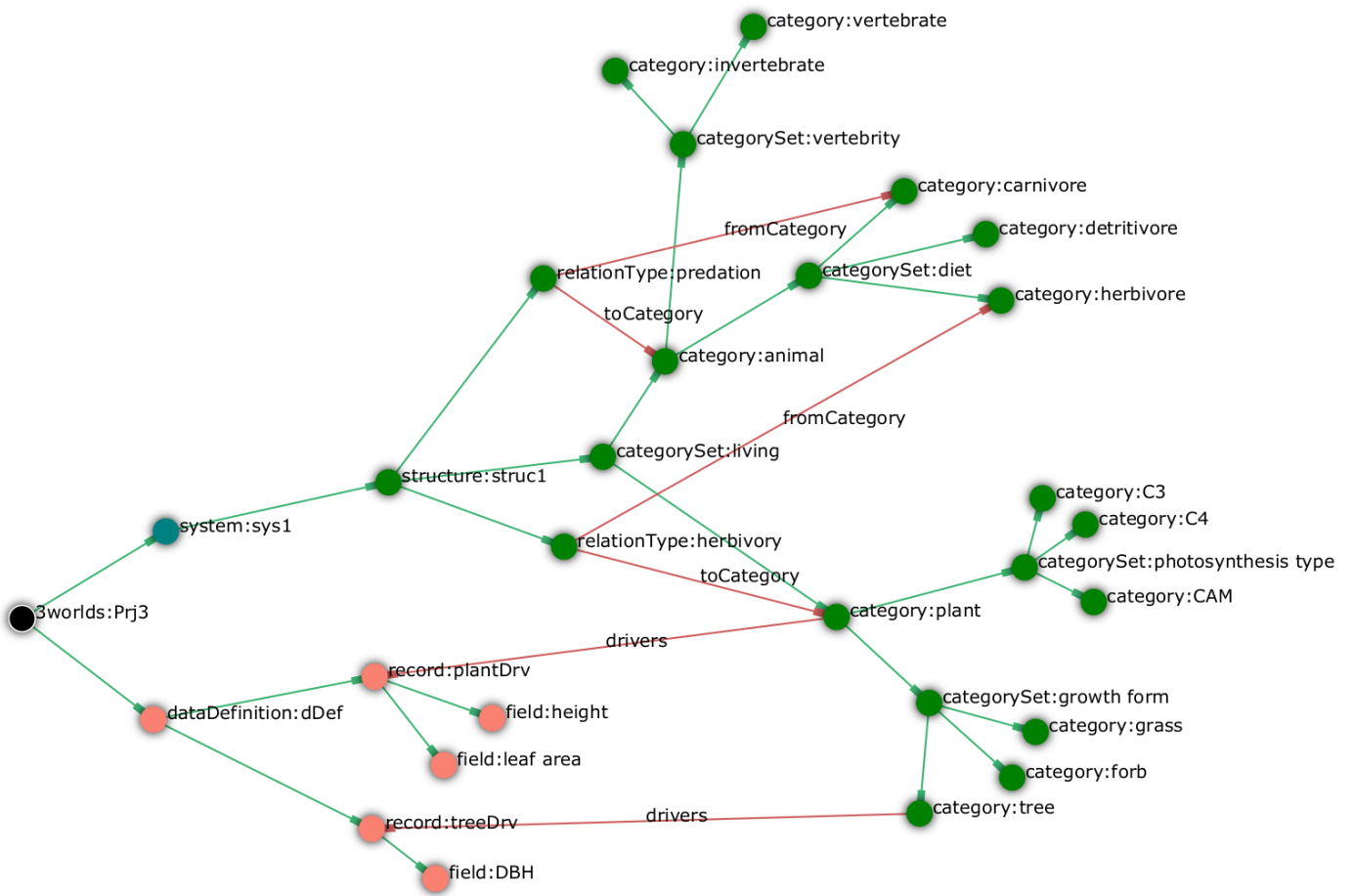


Figure 19. Example of a configuration with category sets, categories and relations

In this example, a *plant* can be a *C3 tree* but cannot be simultaneously a *grass* and a *forb*. Similarly, an *animal* cannot be both *herbivore* and *carnivore*. The *predation* relation links an *animal* of any kind (the prey) to a *carnivore* (its predator). A member of category *plant* will have two driver variables, *height* and *leaf area*. A member of category *tree* will have three driver variables, *height*, *leaf area* and *DBH*, as it inherits all the properties of the *plant* category in which it is nested.

The specification of ecological entities: *system components*

System component

```
/3worlds/system/structure/componentType:<name> {1..*}
```

3Worlds simulates a *system* made of *system components*. These are the things which are instantiated at run time, hold descriptors, and are dynamically changed over the time course of a simulation. When setting up a simulation, one must attach *categories* to *system components*. The rules prevailing to build up category hierarchies mean that a system can belong to a number of non-exclusive categories, as long as the exclusion and nesting rules are respected. For example (Figure 19), we could define a system as belonging to the *plant* and *tree* categories, but not to the *animal* and *tree* categories.

Cross-links for `componentType`:

```
belongsTo → category:<name> {1..*}
```

This link tells to which categories a system component type belongs. The categories must not belong to the same category set. If there are nested categories, membership is inherited (e.g. in the previous example, belonging to the *C3* category automatically implies the system component is also a *plant*). The categories targets of these links must be specific to component types, i.e. no `groupType` or `lifeCycleType` should refer (`belongTo` cross-link) to any of them.

```
belongsTo → category:*component* {1}
```

This link to the predefined category `*component*` is required in all models.

```
belongsTo → category:*atomic* {1}
```

This link to the predefined category `*atomic*` is required in all models. It means that a component has no further subdivisions.

```
belongsTo → category:*ephemeral*|*permanent* {1}
```

This link to either of the predefined category `*ephemeral*` or `*permanent*` is required in all models. It specifies if components of this `componentType` are going to stay forever (permanent) or can be created and deleted (ephemeral) during a simulation. Ephemeral components have two automatic variables, `birthDate` and `age`.

```
loadFrom → dataSource: {0..*}
```

These optional links refer to data sources from which to read initial values of component-level descriptors (Section 3.5.5).

Optional child for `componentType`:

```
/3worlds/.../componentType/initFunction:<name> {0..1}
```

This node declares a function of type `SetInitialState` to set or transform the initial values of component-level descriptors (Section 3.5.2).

Component group

```
/3worlds/system/structure/groupType:<name> {0..*}
```

Sometimes it makes sense to group components of a given `componentType` into *groups*; the best example of this in ecology is the species. Animals of different species may be described by the same descriptors, but nevertheless they will only reproduce within their species. Typically, animals of a species would share some constants, like plumage coloration, average body size, main diet etc. The `groupType` node is meant to fulfill this use case.

To make a `componentType` belong to a `groupType`, you just need to declare it as a child of the `groupType`:

```
.../groupType/componentType:<name> {0..*}
```

Cross-links for `groupType`:

`belongsTo → category:<name> {1..*}`

This link tells to which categories a system component type belongs. The categories must not belong to the same category set. If there are nested categories, membership is inherited. The categories targets of these links must be specific to groups, i.e. no `componentType` or `lifeCycleType` should refer (`belongTo` cross-link) to any of them.

`belongsTo → category:*group* {1}`

This link to the predefined category `*group*` is required in all models.

`belongsTo → category:*assemblage* {1}`

This link to the predefined category `*assemblage*` is required in all models. It means that a group is a population of components. Assemblages have three automatic variables: the group size (number of components) `count`, the number of new components added during last causal step `nAdded`, and the number of components deleted during the causal step `nRemoved`.

`belongsTo → category:*permanent* {1}`

This link to the predefined category `*permanent*` is required in all models. It means that groups will stay forever in the system, even if all components of this groups are gone.

`loadFrom → dataSource: {0..*}`

These optional links refer to data sources from which to read initial values of group-level descriptors (Section 3.5.5).

Optional child for `groupType`:

`/3worlds/.../groupType/initFunction:<name> {0..1}`

This node declares a function of type `SetInitialState` to set or transform the initial values of group-level descriptors (Section 3.5.2).

Life cycle

`/3worlds/system/structure/lifeCycleType:<name> {0..*}`

As *system components* are designed to represent—among other things—individual organisms, they are able to create other system components at runtime, or to transform themselves into a system component of another category assemblage. These abilities are captured in a `lifeCycle`, which describes the possible creations and transitions of system components of a given category set into another.

Since `components` belong to `categories`, different types of system components represented by different state variables, subject to different ecological processes, can coexist in a simulation. It may occur in a particular model that one wishes to represent a transition between, e.g. development stages: think for example of a caterpillar turning into a butterfly. There are chances that you don't want to describe the caterpillar with the same variables and behaviours as the adult butterfly. The operation of transforming a system component from a selection of categories to another is called *recruitment*. Computationally, it means that the simulator must keep track of the system component's identity and age in the first stage and carry these properties on to the new system component of the second stage, and call an appropriate function to transform state variables of the first stage into the new one.

Reproduction is the second process by which system components of a given group of categories may produce other system components belonging to possibly different categories.

A specification of a life cycle requires:

1. a `categorySet` in which the categories represent the various stages of the life cycle;
2. at least one `groupType` definition per `category` of the above `categorySet`;
3. `recruit` and `produce` nodes linking categories within the above `categorySet` to describe possible recruitment

and reproduction transitions;

4. optionnally, `categories` can be linked to a `lifeCycleType` if one wants to attach descriptors to the life cycle.

To define a `groupType` within the context of a `lifeCycleType`, simply make it a child node of the `lifeCycleType`:

```
.../lifeCycleType/groupType:<name> {1..*}
```

Cross-links for lifeCycleType

`appliesTo` → `categorySet:<name> {1}`

This links indicates which categories define the stages of the life cycle. `recruit` or `produce` nodes can only link categories of this set.

`belongsTo` → `category:<name> {0..*}`

These links enable to attach descriptors to a `lifeCycle`, if needed by some ecological processes. The categories targets of these links must be specific to life cycles, i.e. no `componentType` or `groupType` should refer (`belongTo` cross-link) to any of them.

`belongsTo` → `category:*life cycle* {1}`

This link to the predefined category `*life cycle*` is required in all models.

`belongsTo` → `category:*assemblage* {1}`

This link to the predefined category `*assemblage*` is required in all models. It means that a life cycle is a population of components. Assemblages have three automatic variables: the group size (number of components) `count`, the number of new components added during last causal step `nAdded`, and the number of components deleted during the causal step `nRemoved`.

`belongsTo` → `category:*permanent* {1}`

This link to the predefined category `*permanent*` is required in all models. It means that life cycles will stay forever in the system.

`loadFrom` → `dataSource: {0..*}`

These optional links refer to data sources from which to read initial values of life cycle-level descriptors (Section 3.5.5).

Optional child for lifeCycleType :

```
/3worlds/structure/lifeCycleType/initFunction:<name> {0..1}
```

This node declares a function of type `SetInitialState` to set or transform the initial values of life cycle-level descriptors (Section 3.5.2).

Recruitment

```
/3worlds/system/structure/lifeCycle/recruit:<name> {0..*}
```

This node specifies that two categories of the life cycle `categorySet` are linked by a *recruitment* process.

Cross-links for recruit :

`fromCategory` → `category:<name> {1}`

This link tells which system component type is getting changed by the recruitment.

`toCategory` → `category:<name> {1}`

This link tells which system component type is the result of the recruitment.



multiple targets are possible from the same category, i.e. a component of a category may recruit to different categories. E.g., a bee larva can recruit to a worker or a queen.

`effectedBy → process:<name> {1}`

This link tells which ecological process is used to compute the recruitment. This process must implement exactly one `ChangeCategoryDecision` function.

Reproduction

`/3worlds/system/structure/lifeCycle/produce:<name> {0..*}`

This node specifies that two categories of the life cycle `categorySet` are linked by a *reproduction* process.

Cross-links for produce :

`fromCategory → category:<name> {1}`

This link tells which system component type is producing new system components.

`toCategory → category:<name> {1}`

This link tells which system component type is the result of the reproduction.



multiple targets are possible from the same category, i.e. a component of a category may produce components of different categories. E.g., a tree can produce seedlings through sexual reproduction and root suckers through vegetative reproduction.

`effectedBy → process:<name> {1}`

This link tells which ecological process is used to compute the production of new system components. This process must implement exactly one `CreateOtherDecision` function.

Example of a life cycle specification

This life cycle:

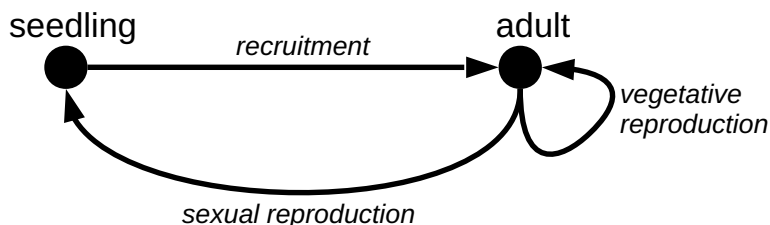


Figure 20. Example of a life cycle

is specified with this graph:

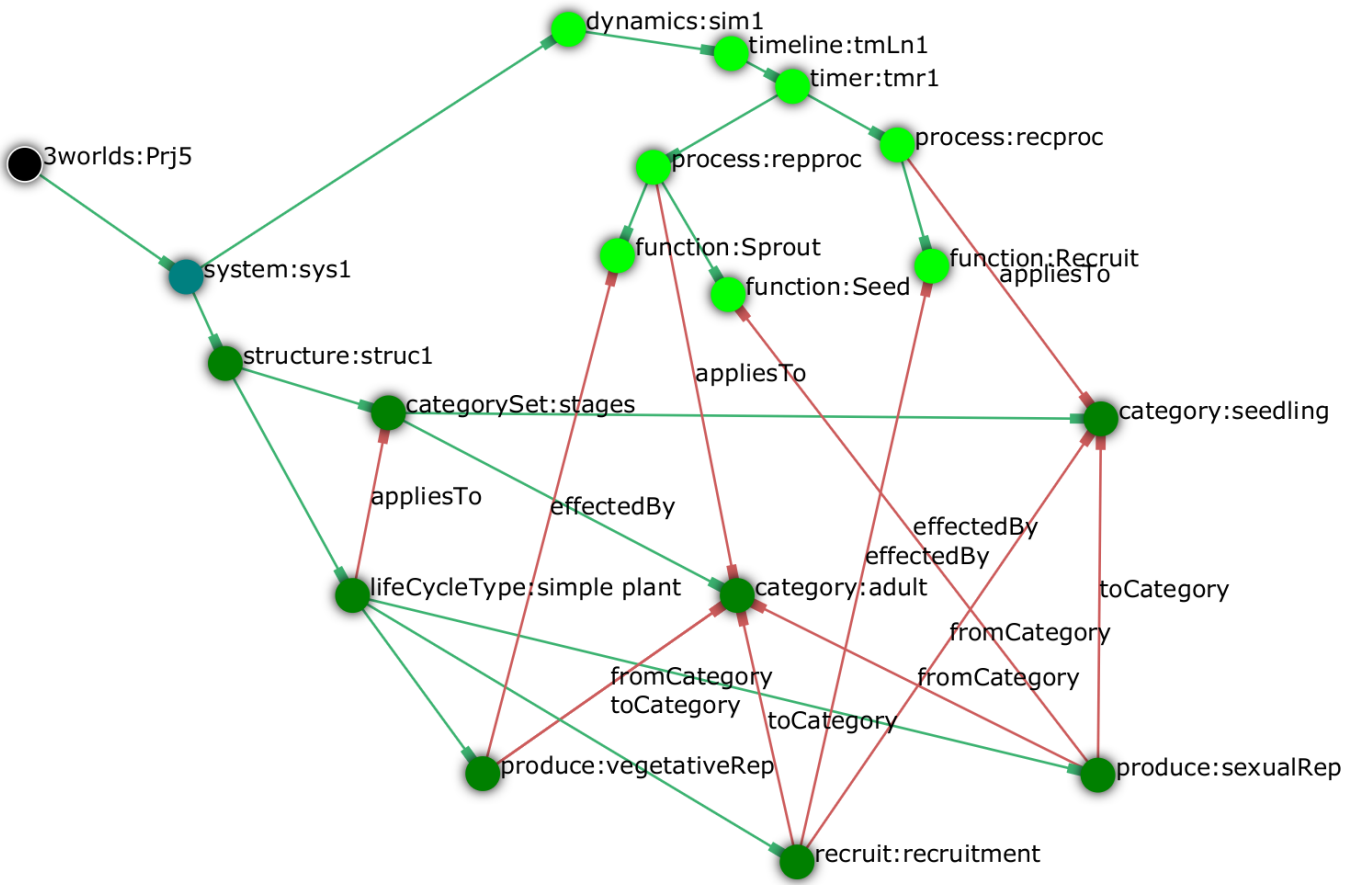


Figure 21. Example of a life cycle configuration

The life cycle has two categories, *seedling* and *adult*, and three transitions: a *recruit* transition called *recruitment*, effected by the function *Recruit*, and two *produce* transitions, effected by the functions *Seed* and *Sprout*.

How is the simulated system implemented?

Internally, system *components* (= ecological entities) are instances of the same class, `SystemComponent`. They are stored in a hierarchy of nested *containers* that represent logical groupings of components based on categories: all components of a container belong to the same categories, in order to facilitate their processing by the user-defined function (remember that a `Process` also applies to some categories).

The nesting of containers is based on the `System > LifeCycle > Group > Component -Type` hierarchy decided by the user. To each container is associated a single particular component instance that represents this container... as one more *component* of the system. It is called the container's *avatar*. Since containers are, by construct, populations of components, they have natural descriptors such as the number of components they contain, and the number of additions and deletions of components during the last simulator iteration. These are stored in the avatar as *automatic variables* (Section 3.3.3.1.1), as specified by the predefined `*assemblage*` category (Section 3.3.8) which must be associated to all containers.

At the top of the container hierarchy is the *arena*, which contains everything else and matches the `System` node. The arena is always present and unique. In a full complexity 3Worlds system, the *arena* container has *life cycle* sub-containers, each of which can have *group* sub-containers which ultimately contain the *components*. Of course this can be simplified: if there are no life cycles or groups defined in a model, these containers will not exist in the simulator and the container hierarchy will be simplified. Possibly to the simplest simulated system in 3Worlds, made of only the arena.

A `LifeCycleType`, `GroupType`, or `ComponentType` does not represent just one life cycle / group / component, but a *class* of those defined by their categories. As all classes, there can be many instances of these at runtime.

When initialising a simulation with data (Section 3.3.3.4), it is possible, and important, to define names that will be used to uniquely identify e.g., various groups as instances of the same `GroupType`.

The representation of space

Sometimes modellers want an explicit representation of space for their model. 3Worlds provides a library of different spaces for this need. Technically, space could just be treated as a component related to other components through particular relations. But since many optimal algorithms have been coded to handle particular aspects of space in relation to other entities, we use space as a kind of container for system components optimized for various operations such as searches for partners for establishing relations (see process).

Space and spatial indexing

```
/3worlds/system/structure/space:<name> {0..*}
```

This defines a space, i.e. a container where system components can be located by their *spatial coordinates* and where distances can be computed. More than one space can coexist in 3Worlds, applying to possibly different types of components and processes. Spaces can be of 1 to 3 dimensions (this is set by choosing a space type).

Properties for space :

type	The type of spatial representation
<p><i>possible values:</i></p>	
continuousFlatSurface	a flat plan with continuous coordinates (default value)
squareGrid	a flat square 2D grid for e.g. cellular automata
topographicSurface	a topographic plan with elevations (not yet implemented)
linearNetwork	a graph of connected segments for e.g., hydrologic networks (not yet implemented)

borderType	The space border behaviour
<p><i>possible values:</i></p>	
wrap	a soft teletransporting border which sends objects crossing it to the other side of the space (default value)
reflection	a hard border on which objects bounce back if they try to escape
sticky	a hard border on which objects stick if they try to escape
oblivion	a soft border into oblivion – objects that cross it disappear into nothingness
infinite	no border – space extends following object movements

edgeEffects

The most common edge effect corrections

possible values:

periodic	wrap-around in all dimensions, i.e. leaving objects enter through the other end (default value)
reflective	all borders are reflective, i.e. objects bounce on borders as on rubber walls
island	a space with oblivious borders in all directions, i.e. leaving objects are lost forever
unbounded	an infinite space adapting to location of items
bounded	a space with sticky borders in all directions, i.e. objects that bump into the border stay there
tubular	wrap around borders in the first dimension, sticky borders in all other dimensions
custom	user-specified border properties – provide a borderType property

precision

Spatial resolution (in space units). Two locations are considered identical if their distance is below this value.

units

The measurement unit for distances and coordinates in this space

guardAreaWidth

The width of a guard area around the space used to define an observation window

observationWindow

A rectangle defining which part of the space is visible

Cross-links for space :

coordinate → space:<name> {1..*}

This link tells which descriptors are used as coordinates to locate system components in this space. This link has a required property rank which tells to which dimension of space a coordinate refers to. For example, you may set to 1 the rank of the x field and to 2 the rank of the y field if you want your 2D space coordinates to be x and y, in this order.

useRNG → rng:<name> {0..1}

Points to the random number generator channel to use with this space (e.g., to generate random locations). If unset, the default random number channel is used.

[3Worlds library of spaces](#)

1. **FlatSurface** : This is a 2 dimensional, continuous space with rectangular limits. Internally, it is coupled to a Quadtree (ref) to optimise the search of nearby points when establishing relations (`relateToDecision` function).

Additional properties for space type FlatSurface :

x-limits An Interval representing the minimum and maximum values of the x (first) dimension.

`y-limits` An `Interval` representing the minimum and maximum values of the y (second) dimension.

2. `SquareGrid`: This is a 2 dimensional, discrete space with rectangular limits and square cells. The locations are only known at the cell level, which helps optimize searches for nearby locations.

Additional properties for space type `SquareGrid`:

`cellSize` The length of the side of the (square) cells in space units

`x-nCells` The number of cells in the x (first) dimension.

`y-nCells` The number of cells in the y (second) dimension.

Shapes and topology

This part is still under construction.

The setup of an initial state for a simulation

To run a simulation, an initial population of system component, group, life cycle, and system *instances* must be constructed from their respective *types* (e.g. `groupType`, `componentType`, etc.). They constitute the *initial state* of a simulation. The initial state is kept in memory by the simulator, and is re-loaded at every simulator reset prior to a new simulation run. To set descriptor values in initial instances, two methods are possible: through the use of a specific *function* (`setInitialState`), or through external data files listed as *data sources*.

Component

`/3worlds/.../componentType/component:<name> {0..*}`

Use this node to instantiate components of a given `componentType`.

Properties for component:

`nInstances` Number of instances to create. If not set, the default behaviour is to create only one instance of this component. Otherwise, `nInstances` components are created, all with identical descriptors - unless a `setInitialState` function is used that can use random numbers to differentiate instances.

Cross-links for component:

`instanceOf → group:<name> {0..1}`

This link tells to which `group` these component instances belong. It is required if `groupTypes` have been defined in the model, optional otherwise. This property is ignored if a `loadFrom` link (see below) is present.

Group

`/3worlds/.../groupType/group:<name> {0..*}`

Use this node to instantiate a group of components. Use groups to set properties that are shared by a set of component instances, e.g. a species name or group-level constants.

Cross-links for group:

`groupOf → componentType:<name> {0..1}`

If no `component` node is defined for this group, this link is required to specify which `componentType` should be used for instantiation of components of this group. In any other case it is not required. You can view this as a replacement for the `instanceOf` link above: either must be present for a group, but not both.

`cycle → lifeCycle:<name> {0..1}`

This link tells in which `lifeCycle` instance this group is involved. The `groupType` of this `group` must be compatible with the `lifeCycleType` category requirements.

Life cycle

`/3worlds/.../lifeCycleType/lifeCycle:<name>`

Use this node to instantiate a life cycle applying to a set of groups.

3.3.4. The `system/dynamics` node

`/3worlds/system/dynamics:<name> {1}`

This node and its sub-tree contains the description of the processes that will change the state of the system and create its dynamics. It is based on the concepts of *time model*, *ecological process*, and *life cycle*. Internally, the `dynamics` node is the *simulator*, i.e. the object which, when kicked to do so, will make all the computations necessary to run a simulation.

The representation of time

Simulation is about mimicking the dynamics of a real system. Here, dynamics is specified by attaching particular behaviours (called processes) to either categories or relation types. Processes may act at a different rhythm or rate in nature, so we need to have a great flexibility in the way time is represented.

Time line

`/3worlds/system/dynamics/timeLine:<name> {1}`

Every simulator has a reference *time line*. Since different ecological processes may run according to different time models, they must refer to a common time frame for interaction to be possible among them. A `timeLine` defines what kind of time scale and time units can be used in a simulation. In 3Worlds, time is always discrete in the end, so that the selected values of time scale and time unit define the time *grain* of the simulation, i.e. the duration below which events are considered simultaneous. Internally, the `ModelRunner` uses integers to represent time, with 1 = one time grain.

Properties for timeLine :

scale

This property specifies the type of time scale to use. The usual time units pose many problems, because years, months, weeks and days are not integer multiples of each other. The option is either to use a real calendar time scale – but this is not needed in most simulation studies – or to use approximations which enable year, months, weeks and days to be integer multiples of each other (e.g. an easy approximation is to assume 30-day months, but this means years must be only 360-day long). This property proposes a set of such simplified, compatible sets of units, denoted as time scales.

possible values:

ARBITRARY	arbitrary time units with no predefined name (default value)
GREGORIAN	real calendar time
YEAR_365D	365-days years, no weeks, no months
YEAR_13M	28-days months, 13-months/52-weeks years
WMY	28-days months, 12-months/48-weeks years
MONTH_30D	30-days months, weeks replaced by 15-days fortnights
YEAR_366D	366-days year, months replaced by 61-days bi-months
LONG_TIMES	long time units only (month or longer), calendar-compatible
SHORT_TIMES	short time units only (week or shorter), calendar-compatible
MONO_UNIT	single time unit, calendar-compatible

`shortestTimeUnit` The shortest time unit used in this model. Note that the time scale constraints the time units compatible with each other for this property.

possible values:

<code>UNSPECIFIED</code>	an arbitrary time unit (default value)
<code>MICROSECOND</code>	microsecond
<code>MILLISECOND</code>	millisecond = 1000 microseconds
<code>SECOND</code>	second = 1000 milliseconds
<code>MINUTE</code>	minute = 60 seconds
<code>HOUR</code>	hour = 60 minutes
<code>DAY</code>	day = 24 hours
<code>WEEK</code>	week = 7 days
<code>FORTNIGHT_15</code>	French-style fortnight = 15 days
<code>MONTH_28</code>	month = 4 weeks of 7 days
<code>MONTH_30</code>	month = 30 days
<code>MONTH</code>	calendar month (= 1/12 of a calendar year), <i>i.e.</i> approx. 30,44 days, but with irregular durations (28,29, 30 or 31 days)
<code>BIMONTH_61</code>	2 months = 61 days
<code>YEAR_336</code>	year = 12 months of 4 weeks of 7 days
<code>YEAR_360</code>	year = 12 months of 30 days
<code>YEAR_364</code>	year = 52 weeks of 7 days = 13 months of 28 days
<code>YEAR_365</code>	year = 365 days
<code>YEAR</code>	calendar year, <i>i.e.</i> approx. 365.25 days, but with irregular durations (365 or 366 days)
<code>YEAR_366</code>	year = 6 bimonths of 61 days
<code>DECADE</code>	decade = 10 years
<code>CENTURY</code>	century = 10 decades
<code>MILLENNIUM</code>	millennium = 10 centuries

`longestTimeUnit` The longest time unit used in this model. *cf.* `shortestTimeUnit` for valid values

`timeOrigin` The value of time at simulation start, either as a long (number of `shortestTimeUnit`) or as a date; 0 by default.

Timers

```
/3worlds/system/dynamics/timeLine/timer:<name> {1..*}
```

Ecological processes may be run following different times. A timer is a particular way of representing time in the simulator. Timers may differ in parameters, like e.g. two timers using different time steps; but they can also be radically different in their logic: e.g. clock-like ticking vs. event-driven simulation.

How do timers work?

Internally, each timer is an instance of the class `Timer`, which is able to return the next time where computation must be made according to its own logic. Every iteration of 3Worlds starts with the simulator asking all timers for their next time, then keeping all of those with the next time closest to last time, and executing them *in unpredictable order*. 3Worlds uses long integers to measure time internally, which means that simultaneous events are possible (times may be strictly equal). The finest time interval is given by `TimeLine.shortestTimeUnit`.

Properties for timer :

`subclass` the type of `timer` to use.

possible values:

`ClockTimer` Time is incremented by a constant amount dt . This is commonly used to simulate regular processes like growth.

`EventTimer` Model dynamics generates *events* and computes the date in the future at which they are going to occur. This is commonly used to generate irregular processes like fire occurrence.

`ScenarioTimer` **Not yet implemented.**

Additional properties when `class = ClockTimer`

`timeUnit` the base time unit used by this timer. cf. `timeLine.shortestTimeUnit` for the valid values of this property

`nTimeUnits` the number of base time units in the time unit of this model (e.g., a model may have a 2 year time unit)

dt

The constant time increment used in this `ClockTimer`, expressed as an integer number of `timer` base unit (= `timer.nTimeUnits × timer.timeUnit`). For example, if the `timer` has `timeUnit = DAY` and `nTimeUnits = 3`, `dt` is expressed in units of 3 days (e.g. `dt = 2` means the time increment is 6 days).



if calendar time is used (`timeLine.scale = GREGORIAN`), then `dt` will sometimes not be constant (e.g. if `dt = 2 MONTH`, `dt` will vary in duration between 59 and 62 days according to the exact date).

offset

In some cases it is desirable to offset a `ClockTimer` relative to another one running at the same time step, for example to be certain of their order of execution or to simulate cyclic phenomena (e.g. seasons). The timer will start after `offset × timeUnit` units, where `offset` is a fraction between 0.0 and 1.0.

Additional cross-links when `class = EventTimer`

An `EventTimer` maintains a queue of time events that gets populated by user-defined ecological functions. This way, future events depend on the dynamics of the system. Functions that can populate an event queue will have an `EventQueue` argument in their heading, usable to create future time events. Events must set a time in the future, i.e. their time must be larger than current time. Event times are always expressed in `Timeline.shortestTimeUnits` in an `EventTimer`.

`fedBy → initFunction:<name> {1}`

This link is used to set the first event of an `EventTimer` at simulation start, hence from an `initFunction`. Without this initial event, the timer will never start. More than one event can be created here.

`fedBy → function:<name> {0..*}`

These links indicate which functions can populate the event queue with time events. Multiple events can be created here.

Simulation stopping condition

`/3worlds/system/dynamics/stoppingCondition:<name> {0..*}`

A simulation may be run indefinitely (interactively), but in big simulation experiment it is useful to automatically stop the simulations according to some criterion. Besides the simplest stopping condition, reaching a maximal time value, 3Worlds provides many other possibilities to stop a simulation (e.g. based on a population size, on a variable passing a threshold value, etc.).

When no stopping condition is defined, the simulation will run indefinitely.

Properties for `stoppingCondition`

`subclass` The type of stopping condition to use

possible values:

`SimpleStoppingCondition`

Simulation stops when a maximal time value is reached.

`ValueStoppingCondition`

Simulation stops when a variable in a reference system component reaches a given value.

`InRangeStoppingCondition`

Simulation stops when a variable in a reference system gets within the given range.

`OutRangeStoppingCondition`

Simulation stops when a variable in a reference system gets out of the given range.

`MultipleOrStoppingCondition`

Compound stopping condition: simulation stops when *any* of the elementary stopping conditions within this multiple condition's list is true.

`MultipleAndStoppingCondition`

Compound stopping condition: simulation stops when *all* of the elementary stopping conditions within this multiple condition's list are true.

Additional properties when `class = SimpleStoppingCondition`

`duration` The duration of the simulation beyond time line `timeOrigin`, in time line `shortestTimeUnits`.

Additional cross-links when `class = ValueStoppingCondition, InRangeStoppingCondition, OutRangeStoppingCondition`

`stopSystem → component:<name> {1}`

The system component in which some variable will be checked to stop the simulation.

Additional properties when `class = ValueStoppingCondition, InRangeStoppingCondition, OutRangeStoppingCondition`

`stopVariable` The name of the variable in `stopSystem` which values are used to decide to stop the simulation.

Additional properties when `class = ValueStoppingCondition`

`stopValue` The value of `stopVariable` at which to stop the simulation.

Additional properties when `class = InRangeStoppingCondition, OutRangeStoppingCondition`

`upper` The upper value of the `stopVariable` range. Only `double` values are accepted.

`lower` The lower value of the `stopVariable` range. Only `double` values are accepted.

Additional cross-links when class = MultipleOrStoppingCondition, MultipleAndStoppingCondition

condition → stoppingCondition:<name> {1}

These links point to the stopping conditions that will be used as elementary stopping conditions by the multiple and/or stopping condition. Use these links to construct complex stopping conditions.

The transformations of a system component

Changes in a *system component* through time may be of different kinds: changes in *state*, i.e. in its driver and decorator variables; or more radical changes where the component actually changes *category*, so becomes represented by a different set of variables. Plus, a component may have an ephemeral life (*lifespan* property), which means component objects are dynamically created or deleted during a simulation.

Process

/3worlds/system/dynamics/timeLine/timer/process:<name> {1..*}

Processes are used in 3Worlds to compute change in *system components*. Each process acts on system components of a particular group of categories and is scheduled by a particular timer. Processes contain user-defined code that represents ecological processes. This gives 3Worlds its versatility: one can mix in a single model completely different ecological entities (system components of different categories), implement any ecological process depending on user needs, and put them to work on different time scales (times). A `process` is run according to its parent `timer`.

Cross-links for process

A process can act on a single system component at a time (called the *focal* system component), or on a pair of components linked by a relation (called the *focal* and the *other* system components). This is specified using the `appliesTo` cross-link (one at least must be present):

appliesTo → category:<name> {0..*}

These links indicate the categories of system components that will be acted on by the process.

appliesTo → relation:<name> {0..1}

This link indicates to which relation type between system component the process applies.



A process **must** apply either to categories (*category process*) or to a single relation type (*relation process*), but cannot apply to both.

dependsOn → process:<name> {0..*}

This link tells that the process must be activated *after* the processes targeted by the links. Use this link to organize computations when there are dependencies between them.

inSpace → space:<name> {0..1}

This link indicates that this process will make use of this space for its optimisation of neighbour searches. Use this link when you want fast search for candidates to establish a relation, i.e. in association with the `relateToDecision` function. The required property `searchRadius` in the `inSpace` link tells to which *maximal* distance candidates for establishing a relation must be searched. 3Worlds will only present components at a shorter distance than `searchRadius` (possibly none) to the `relateToDecision` function. A value of zero means the algorithm searches for the closest neighbour only (possibly more than one if all exactly at the same distance), whatever its distance to the focal component.



Forgetting to set the value of `searchRadius` is a common source of unexpected behaviour in spatial models.

How do processes work?

Internally, an instance of the `Process` class implements a '*for each*' loop on system components during an iteration step, as scheduled by its `Timer`. The order in which the components are processed in a loop is unpredictable (not fixed, nor random); the only safe assumption is to consider that they are all processed simultaneously.

If the process applies to a list of categories, it is a single loop on all components belonging to these categories. This is the `ComponentProcess` sub-class.

If it applies to a relation type, it is either (1) a double loop, possibly optimized by using a space, on components belonging to the 'from' list of categories of the relation type, and components belonging to the 'to' list of categories of the relation type (`SearchProcess` sub-class); or (2) a single loop on relations instances of its relation type (`RelationProcess` sub-class).

Processes of a same timer are run in any (= unpredictable) order unless explicitly ordered by the use of the `dependsOn` cross-link. Ordering processes in such ways defines *causal steps* (Section 1.2.5) within the time steps.

In short: process = (parallel) loop on components (within one causal step) within one time step.

Function

```
/3worlds/system/dynamics/timeLine/timer/function:<name> {1..*}
```

This node is used to specify the details of the computations made in a `process`. The `process` defines which system components are going to be activated and at what time in the simulation course; the `function` defines which computations, in detail, will be applied to the system components of that process. This enables to build complex computations applying to one component (a series of `functions` within a `process`) in the context of a particular subset of components (`process`).

There are different types of functions, differing by the way they affect system components and relations. The selection of a function type will trigger the generation in the model java source file of a method having the name specified in the `function` node name. This method is expected to be edited by the modeller in order to implement her/his favourite version of the ecological process modelled by the function.



The name of a function must be a valid java class name, starting with an uppercase letter.

Properties for function :

type This property specifies which kind of biological function will be implemented within the linked `Process` object.

possible values:

<code>ChangeState</code>	change the state, i.e. the values of the descriptors of a system component (default value)
<code>ChangeCategoryDecision</code>	change category of a system component according to life cycle (has no effect if no life cycle is specified)
<code>CreateOtherDecision</code>	create another system component, of the same categories if no life cycle is present, otherwise as specified by the life cycle
<code>DeleteDecision</code>	delete self
<code>ChangeOtherState</code>	<i>focal</i> changes the state of <i>other</i>
<code>RelateToDecision</code>	<i>focal</i> establishes a new relation to <i>other</i>
<code>MaintainRelationDecision</code>	decision to maintain or remove an existing relation
<code>ChangeRelationState</code>	change the state of a relation, i.e. possibly both the state of <i>focal</i> and <i>other</i> at the same time
<code>SetInitialState</code>	sets the initial state of a newly created <code>SystemComponent</code>
<code>SetOtherInitialState</code>	sets the initial state of a newly created <code>SystemComponent</code> given a parent component

! The `relateToDecision` function has a special status among relation processes as it is used to establish a relation, whereas all other functions for relation processes use an already established relation. As a result, a process parent to a `relateToDecision` function cannot be parent to any other type of function. For computation efficiency, it is recommended to associate a *space* to the process parent to a `relateToDecision` function.

functionSnippet This optional property is used in 3Worlds tutorial and test models to store the java code of user-defined function. This code is inserted in the body of the generated user-code methods.



Use snippets only for simple models as there no Java compiler checking your code for errors.

Cross-links for `function`:

`useRNG → rng:<name> {0..1}`

Points to the random number generator channel to use in this function. If unset, the default random number channel is used.

Additional properties when `function = createOtherDecision`:

`relateToProduct` A logical value (`false` by default) which tells to establish a permanent `parentTo` relation (defined in the `predefined` sub-tree: Section 3.3.8) between a parent component and its offspring created by calling `createOtherDecision`.

How do functions work?

Functions are called within the loop on components of their parent process. The processing order of functions within their process is constant:

- In a `ComponentProcess` :
 - execute all `changeState` functions
 - then, all `deleteDecision` functions,
 - then, all `createOtherDecision` functions,
 - then, all `changeCategoryDecision` functions,
 - then send data to all data trackers
- In a `SearchProcess` :
 - execute all `relateToDecision` functions
- In a `RelationProcess`
 - execute all `changeOtherState` functions
 - then, all `maintainRelationDecision` functions,
 - then, all `changeRelationState` functions

The order of these computations does not matter for *driver* descriptors, nor for component creation and deletion, because all the changes are postponed until after the process loop is over. For example, a component may in the same time step reproduce an die. The order may only matter for *decorators*, which are provided here as a convenience for complex computations.

A way to make your model code efficient is to group functions run with the same timer in a single process, so that they are called in a single pass on components instead of many in case you use different processes. This assumes that your functions can be processed in any order without consequences on the outcome - which is not always true. If you need a precise ordering of function calls because, e.g. one of them requires that a decorator variable has been set by another, then you must place your functions in different processes linked by a `dependsOn` cross-link to tell which is computed first.

At the end of a process loop (causal step, Section 1.2.5):

- the values of all *drivers* are updated (i.e. values computed by functions are stored in a *next* driver data structure, which now replaces the *current* driver structure);
- *components* to be deleted and to be created as decided by `-Decision` functions, and *ephemeral* relations to create/delete, are now inserted in the current list of components;
- *spaces* are updated according to changes in the component community;
- *permanent* relations between components are updated.

At the end of all process loops (time step, Section 1.2.5):

- all *decorators* are set to zero;

- *automatic* descriptors (e.g. component age, population numbers) are updated;

Function consequences

/3worlds/.../process/function/consequence:<name> {0..*}

Some functions may imply consequences: for example, a decision to delete another system component may be followed by a change in state based on the deleting component's state at the time it is deleted. Such functions that are only activated when certain events take place are called *consequences* and may be specified by a child node to a function. Here also, rules apply:

function	consequence	use to
changeState		
changeRelationState		
changeOtherState		
deleteDecision	changeOtherState	carry over values to another component linked by a returnsTo relation
createOtherDecision	setOtherInitialState	set the initial state of the new component
relateToDecision		
changeCategoryDecision	setOtherInitialState	carry over and compute values from the former component (<i>focal</i>) to the new recruit
maintainRelationDecision		
relateToDecision		

Consequence functions have the same properties and cross-links as functions (cf. above).

How do consequence functions work?

Internally, consequence functions are just functions which are called *immediately after* their parent function. E.g., in a `ComponentProcess` loop on `deleteDecision` functions, each `deleteDecision` function is immediately followed by a call to its consequence `changeOtherState` function, if any.

Data tracking

/3worlds/.../process/dataTracker:<name> {0..*}

Data trackers are used to get output from a simulation. This output can then be redirected either to graphic windows of `ModelRunner` or to a data file for later processing with other software. Using a data tracker is similar to what happens in the real world: you set a sensor into the system you want to monitor and wire it to a data logger which translates the signal into human-readable data. There are different kinds of data trackers according to the output format and the sampling method you want to use; and different ways of wiring these trackers to the simulated system and its components.

A data tracker will record values of descriptors (except constants) of a sample of system components or groups, or of the system itself and send them to any listener (usually a widget) linked to it. The frequency of data sending is determined by the *timer* of the parent *process* of the data tracker.



A `dataTracker` node can only be specified in a *category process*, not in a *relation process*.

Properties for `dataTracker` :

<code>samplingMode</code>	This property specifies how to pick system components for data tracking <i>within a group</i> . The data will be either (1) selected for one particular system component within each group, or (2) taken from that group's own data, if any (e.g. species population size), or (3) aggregated using some statistical method ; all this, depending on the cross-links of the <code>dataTracker</code> . For (1), the default behaviour is that once a system component is selected, it will be tracked until its deletion by the simulator. In all cases, remember that the maximal number of data tracking channels is set by the <code>sampleSize</code> property; this property only tells the software how to fit the data coming from possibly many system components into the requested number of output channels.
---------------------------	---

possible values:

- `RANDOM` selects system components randomly to constitute the sample (default value)
- `FIRST` selects the first system components of their group (as internally stored) to constitute the sample. Notice that components may come in any order and that this order may change unpredictably during a simulation.
- `LAST` selects the last system components of their group (as internally stored) to constitute the sample. Notice that components may come in any order and that this order may change unpredictably during a simulation.

<code>sampleSize</code>	How many system components at most will be tracked. Valid values are any positive integer or the keyword <code>ALL</code> to sample all the components of a given group.
-------------------------	--

statistics

This property lists transformations of the raw data to compute when a group contains more than one system component.

possible values:

`mean` mean (default value)

`var` variance

`se` standard error

`cv` coefficient of variation (%)

`sum` sum

`N` count

`min` minimum

`max` maximum

tableStatistics

This property lists transformations of the raw data to compute in case of a table variable. The grouping is determined by the index specification in the track variable list. cf. property `statistics` for valid values.

subclass

The type of data tracker to use

possible values:

`DataTracker0D`

time series tracker; returns (t, x_1, \dots, x_n) values (where t is time and x_i the descriptors of interest).

`DataTrackerXY`

raw tracker; returns (x,y) values at each time step.

`DataTracker2D`

unimplemented yet

Cross-links for `dataTracker`

`sampleComponent` → `componentType:<name>` {0..1}

The `dataTracker` will track components of the `componentType` pointed by this link, following the sampling strategy specified by properties `samplingMode`, `sampleSize`, `statistics` and `tableStatistics`. *Permanent* components will be tracked until the end of the simulation. *Ephemeral* components are tracked all their life long: a new sampling is done every time a formerly tracked component is deleted. Sampling strategy can further be refined using two optional properties (see below).



The `componentType` categories must match those of the `process` parent of the `dataTracker`.

Optional properties for sampleComponent cross-links:

`idLifecycle` The identifier of the *life cycle* which components are to be sampled from, as a `String`; only required if a life cycle has been specified.

`idGroup` The identifier of the *group* which components are to be sampled from, as a `String`; only required if a group has been specified

sampleGroup → groupType:<name> {0..1}

The `dataTracker` will track groups of the `groupType` pointed by this link.



The `groupType` categories must match those of the `process` parent of the `dataTracker`.

Optional property for sampleGroup cross-links:

`idGroup` The list of identifiers of the groups to track, as a `StringTable`. If not set, all groups of the `groupType` are tracked.

sampleLifeCycle → lifeCycleType:<name> {0..1}

The `dataTracker` will track life cycles of the `lifeCycleType` pointed by this link.



The `lifeCycleType` categories must match those of the `process` parent of the `dataTracker`.

Optional property for sampleLifeCycle cross-links:

`idLifecycle` The list of identifiers of the life cycles to track, as a `StringTable`. If not set, all life cycles of the `lifeCycleType` are tracked.

sampleArena → system:<name> {0..1}

The `dataTracker` will only track the *system* data.

trackField → field:<name> {0..*}

The `dataTracker` will track all fields pointed by these cross-links in all objects (*components, groups, life cycles or system*) it is sampling.

trackTable → table:<name> {0..*}

The `dataTracker` will track all tables pointed by these cross-links in all objects (*components, groups, life cycles or system*) it is sampling.



the `trackField` or `trackTable` cross-links must refer to fields or tables present in the categories of the parent `process` of the `dataTracker` (`process appliesTo` cross-links).

Properties for trackField and trackTable cross-links:

`index` A list of `String`s giving the range of table indices to track. The following rules are used to specify which cells of the table should be picked:

1. an index specification for a table is enclosed in square brackets: `[]`;
2. vertical bars separate dimensions: e.g. `[||]` refers to a table with three dimensions;
3. a single number is used to specify a single cell in one dimension: e.g. `[1||]` refers to the cell 1 in dimension 1 (Note: this is the *second* cell in this dimension as all dimension indices start at 0 = first cell);
4. a list of cells in one dimension is specified using commas: e.g. `[1|0,2,7|]` refers to the cells 0, 2 and 7 in the second dimension;
5. a range of cells in one dimension is specified using a column: e.g. `[1|0,2,7|2:4]` refers to the cells 2,3 and 4 in the third dimension;
6. a negative number means that all cells in a dimension must be tracked *except* this number: e.g. `[-1|0,2,7|2:4]` indicates that all cells in the first dimension except cell 1 must be tracked.
7. a minus sign in front of a range means that all cells in a dimension must be tracked *except* this whole range: e.g. `[1|0,2,7|-2:4]` indicates that all cells in the third dimension except cells 2, 3 and 4 must be tracked.
8. an empty dimension means all cells in this dimension are to be tracked: e.g. `[1||2:4]` indicates that all cells in the second dimension will be tracked;
9. an empty index string `[]`, or no index string, means that the whole table (all cells) is to be tracked.

The descriptors may be elaborate hierarchical data structure which include a table at some level. For each table in this hierarchy, an index `String` as described above must be given in the `index` property, in a comma-separated list, e.g.: `[0:2|0],[4,6]` means that index `[0:2|0]` should be used for the top-level table of the descriptor, and index `[4,6]` for the bottom-level table.

[TO DO: a few examples]

3.3.5. The `dataDefinition` node

```
/3worlds/dataDefinition {1}
```

This node and its sub-tree contains the detailed specifications of descriptors used in a model. This information is used by 3Worlds to generate java code for the data structures that will be used in simulations of this particular model. These data structures are then made available to descendants of 3Worlds functions that users will edit and modify to describe their particular model.

The descriptor tree

Every *system component* of 3Worlds potentially holds four kinds of *descriptors*. Each of these (*drivers*, *decorators*, *constants* and *automatic variables*) is a *tree* of two kinds of data structures: *records* and *tables*. Records contain *fields*, i.e. atomic (i.e. not breakable into smaller parts) pieces of data of different types (numbers, character strings, logical values), each field having a name to access its value (i.e. field = (name,value) pair). Tables are multi-dimensional arrays of values of the same atomic type.

The data tree is constructed by allowing records to contain tables as fields (but not records), and by allowing tables to contain records all made of the same fields (but not tables) instead of atomic types. This gives end-user modellers great flexibility to organise their data into elaborate structures.

Record

```
/3worlds/dataDefinition/record:<name> {0..*}
```

```
/3worlds/.../table/record:<name> {0..*}
```

A *record* is a data structure made of *fields* (=name,value) pairs or *tables* that represent the *descriptors* of a *category* of *components*. All descriptor trees must start with a record as their root. The record name will be turned into a java class name by the 3Worlds code generator for use in end-user code. A record may be nested in a *table*.



A record cannot be empty, i.e. it **must** have at least one child node (field or table), see below.

Field

```
/3worlds/.../record/field:<name> {0..*}
```

A *field* is an atomic piece of data accessed by its name within a record.

Properties for field :

type The data type of this field.

possible values:

Double a double precision floating point number (4.9406564584124654 10^-324 to 1.7976931348623157 10^308, symmetric for negative numbers) with 16 significant digits (default value)

Integer an integer [-2147483648; 2147483647]

Long a long integer [-9223372036854775808; 9223372036854775807]

Float a single precision floating point number (1.40239846 10^-45 to 3.40282347 10^38, symmetric for negative numbers) with 8 significant digits

Boolean a logical value {true, false}

String a text string

Short a short integer [-32768; 32767]

Char a character value (16-bit Unicode = UTF16, i.e. 65535 different values)

Byte a very, very short integer [-128 ; 127]

Object anything else **TODO: should we keep this? it's probably useless**

description A long description of the field

units Measurement units of field values

precision Precision used for display of field values

interval Range of possible values for this field (real number types)

range Range of possible values for this field (integer number types)

Dimensioner

```
/3worlds/dataDefinition/dimensioner:<name> {0..*}
```

A *dimensioner* is used to define the number of entries in a table.

Properties for dimensioner :

size The number of table cells in this dimension.

Table

```
/3worlds/.../record/table:<name> {0..*}
```

A table is a multi-dimensional data structure containing either atomic data types or *records*, all of the same type.

Properties for table :

dataElementType The data type of the elements of the table. cf property `field.type` for valid values.

description A long description of the table

units Measurement units of table values

precision Precision used for display of table values

interval Range of possible values for this table (real number types)

range Range of possible values for this table (integer number types)



These properties are not needed when the table is a table of records (cf. above).

Cross links for table :

```
sizedBy → dimensioner:<name> {1..*}
```

This link sets one dimension of a table to be of the length found in the dimensioner `size` property. The required property `rank` in the `sizedBy` edge tells the rank of the dimension. All `sizedBy` links of a table should have a different rank, in increasing order starting from 0 to the number of dimensions -1.

Random number channels

```
/3worlds/dataDefinition/rng:<name> {0..*}
```

Simulation models make extensive use of random numbers to make decisions on components (e.g. decision to delete a component or to create new ones). 3Worlds defines random number channels, that can be attached to specific functions. These channels can be held constant by using the same seed in a series of simulations, thus enabling to conduct factorial experiments controlling for a source of random numbers or another.

Properties for rng :

`algorithm` Algorithm used to produce random numbers

possible values:

- `PCG32` Implementation of a permuted congruential generator with 32-bit output ([PGC32](https://en.wikipedia.org/wiki/Permuted_congruential_generator) (https://en.wikipedia.org/wiki/Permuted_congruential_generator)); fast (56% faster than `JAVA`) and good quality (default value)
- `JAVA` The default Java random number generator of `java.util.Random`; medium speed, poor quality
- `XSRANDOM` Implementation of a Xorshift random number generator as found [here](http://demesos.blogspot.com/2011/09/replacing-java-random-generator.html) (<http://demesos.blogspot.com/2011/09/replacing-java-random-generator.html>); very fast (76% faster than `JAVA`), medium quality

`seedSource` How to create the seed of the random number generator

possible values:

- `PSEUDO` The random number seed is produced from a call to a unique instance of `java.util.Random.Random()`. It uses time to the nanosecond to produce a 'very likely to be distinct' seed (default value)
- `NATURAL` The random number seed is taken as an element in a table of 1000 natural random numbers that have been obtained from atmospheric noise. Use the property `tableIndex` to specify which item in this table should be taken for the seed
- `CONSTANT` The random number seed is a constant (either 0 or 1, depending on the algorithm)

`resetTime` When to reset the random number channel seed

possible values:

- `NEVER` The random number seed is never reset after initialisation of the random number channel (default value)
- `ONRUNSTART` The random number seed is reset to its former value for every simulation run (producing the same series of random numbers for every simulation)

`tableIndex` Index into an `array[0..999]` of naturally generated random numbers to act as seeds for resetting



If no random number channel is specified, 3Worlds uses a single default channel with algorithm `XSRANDOM`, `seedSource PSEUDO` and `resetTime NEVER`.

3.3.6. The experiment node

```
/3worlds/experiment:<name> {1}
```

This node and its sub-tree describe the experimental design to run using a given *model* and external data sets. Typically, it will tell `ModelRunner` how many simulations should be run, possibly varying some parameters of the model according to some plan, where data should be read and saved. The name is used to differentiate simulation experiments in a meaningful way.

The default, simplest, simulation experiment is just to run a single simulation of the *baseline* model.

Properties for experiment :

`nReplicates` The number of times all elementary treatments must be replicated. Use this only if there is an internal source of variation in simulations, like random numbers.

Cross-links for experiment :

```
baseLine → system:<name> {1}
```

This link points to a model setup (Section 3.3.2) that will be used as a *base line* simulation. A base line simulation is the equivalent to a control in a real experiment, i.e. a reference case that serves as a basis to which other treatments are compared. Often, the base line is the setup for which data is available to compare simulation outputs to.

Simulation duration

```
/3worlds/experiment/timePeriod {0..*}
```

The duration of a particular simulation is specified using a `timePeriod` node.

[TO DO: sort out the conflict between this and the stopping conditions]

Properties for timePeriod

`start` The starting time of a simulation in `timeLine` shortest time units.

`end` The ending time of a simulation in `timeLine` shortest time units.



Both properties are optional. If none is set, the simulation will start at time 0 and run indefinitely.

Cross-links for timePeriod :

```
stopOn → stoppingCondition:<name> {0..1}
```

This link tells how to stop the simulation in case no `end` property is given.



The stopping condition has the priority over the `end` property. **TO DO: check this**

Experimental design

```
/3worlds/experiment/design:<name> {1}
```

An experimental design specifies the method used to perform the simulations, e.g. number of replicate simulations, treatments as changes in parameter values or initial states, etc. An experimental design can be specified by using standard designs, or by passing a design description file.

For more information on experimental designs for simulation experiments, we recommend reading the documentation of the R software *planor* and *mtk* packages (e.g. these packages could be used to generate design files for use in 3Worlds).

[TO DO: OpenMole integration]

Properties for design :

type This property specifies an experimental design for the simulation experiment. It only provides basic, standard experimental designs. For more elaborate or specialized designs, use an ad-hoc file description of the design (`file` property, cf. below)

possible values:

`singleRun` an experiment consisting of a single simulation run (default value)

`crossFactorial` a cross-factorial experiment based on a limited set of parameter values (factors)

file This property gives the name of an experimental design file **TODO : expand on this description**

Experimental treatments

Treatments

`/3worlds/experiment/treatment:<name> {0..*}`

An experimental treatment records a particular set of parameter values and initial state to run a simulation or a series of replicated simulations. It is the basic block of the experiment, just as in real-world experimentation.

Treatments may be specified

- in full detail: this is done by specifying more than one `system` node, each `system` being used for a different treatment;
- as (minor) changes relative to the experiment `baseLine`.

NB: not fully implemented yet

Treatments as changes relative to `baseLine`

`/3worlds/experiment/treatment/modelChange {0..*}`

NB: not fully implemented yet

Managing experiment data

Inputs: `dataSource`

`/3worlds/experiment(dataSource:<name> {0..*})`

This node describes a source of data to use to instantiate a model: name of the data source, access method, etc.

Properties for dataSource

file A valid data file name

`subclass`

A data format to use to read that file

possible values:

	<code>CsvFileLoader</code>	<code>comma</code>	<code>separated</code>	<code>value</code>
		(https://en.wikipedia.org/wiki/Comma-separated_values)	.csv	text file
	<code>OdfFileLoader</code>	OpenDocument (https://en.wikipedia.org/wiki/OpenDocument)		.ods

`read`

(optional) A list of the column headers to read. Each column name must match a declared `field` (Section 3.3.5.2) or `table` (Section 3.3.5.2.2). If this property is absent or not set, all columns will be read.

`idLifeCycle`

(optional) Header of the column containing the *life cycle* names (instances of `lifeCycleType`). If only this property is set, each row of the file/spreadsheet will be used to generate a different *life cycle* instance (with associated component container, constants, drivers and decorators). If it is used in conjunction with `idGroup` and/or `idComponent` (see below), it will be used to search for or create the proper *life cycle* container in which to place the requested items (*groups* or *components*). Values of this property will become the life cycle instance unique identifiers. They must be consistently used across `dataSource` / `initialValues` nodes when initial items are constructed from more than one source.

`idGroup`

(optional) Header of the column containing the *group* names (instances of `groupType`). If only this property is set, each row of the file/spreadsheet will be used to generate a different *group* instance (with associated component container, constants, drivers and decorators). If it used in conjunction with `idComponent` (see below), it will be used to search for or create the proper *group* container in which to place the requested *components*. Values of this property will become the group instance unique identifiers. They must be consistently used across `dataSource` / `initialValues` nodes when initial items are constructed from more than one source. If the `groupType` is declared as the child of a `lifeCycleType`, then the `idLifeCycle` property (cf. above) is also required, either in `dataSource` or `initialValues` nodes.

`idComponent`

(optional) Header of the column containing system *component* unique identifiers (instances of `componentType`). Every different value will generate a different *component*. Contrary to the two previous properties, the value is *not* used to generate the component instance identifier, which is specifically generated by 3Worlds. Nevertheless, they must be used consistently across `dataSource` / `initialValues` nodes when initial items are constructed from more than one source. If the `componentType` is declared as the child of a `groupType`, then the `idGroup` property (cf. above) is also required, either in `dataSource` or `initialValues` nodes.

`dim`

(optional) A list of column headers containing dimension indices. This must only be present when other column headers refer to tables. When this property is present, every column name must match a declared `table` `table` with as many dimensions as listed here, and compatible maximal index values.

Additional properties when `subclass = CsvFileLoader`

`separator` the field separator used for this `.csv` file (default: tabulation "\t")

Additional properties when subclass = OdfFileLoader

sheet the name of the spreadsheet to load from this `.ods` file (different spreadsheets in the same `.ods` file must be specified as different `dataSource` nodes). If this property is absent or not set, the first spreadsheet will be loaded.

`.csv` and `.ods` file formats both assume the data come in 2 dimensional tables with cross-references between the tables. Table columns must match parameter and driver field or table names. Table rows must match species, stage or system component instances.

The following rules must be respected when preparing the data files:

- The data must not contain any missing value or structural empty cells.
- Empty lines are permitted (they are skipped).
- Text data must not be quoted.
- The first data line of any file or spreadsheet must contain column headers. They must match field names as defined under the `dataDefinition` node (Section 3.3.5).

Outputs: `dataSink`

`/3worlds/experiment/dataSink:<name> {0..*}`

This node describes a 'sink' where data resulting from simulation output is to be stored for later processing by other software.

[Not yet implemented]

Properties for dataSink

file A valid data file name

Cross-links for dataSink :

`source → dataTracker:<name> {1}`

This link the origin of the data to save into the sink. Usually, a data tracker

3.3.7. The *userInterface* node

`/3worlds/userInterface:<name> {1}`

This node and its sub-tree specifies the look of the `ModelRunner` user interface. `ModelRunner` is highly configurable and can show many graphs during a simulation run, as help when debugging a new model; or only show a progress bar to improve computing performance when running a big simulation experiment.

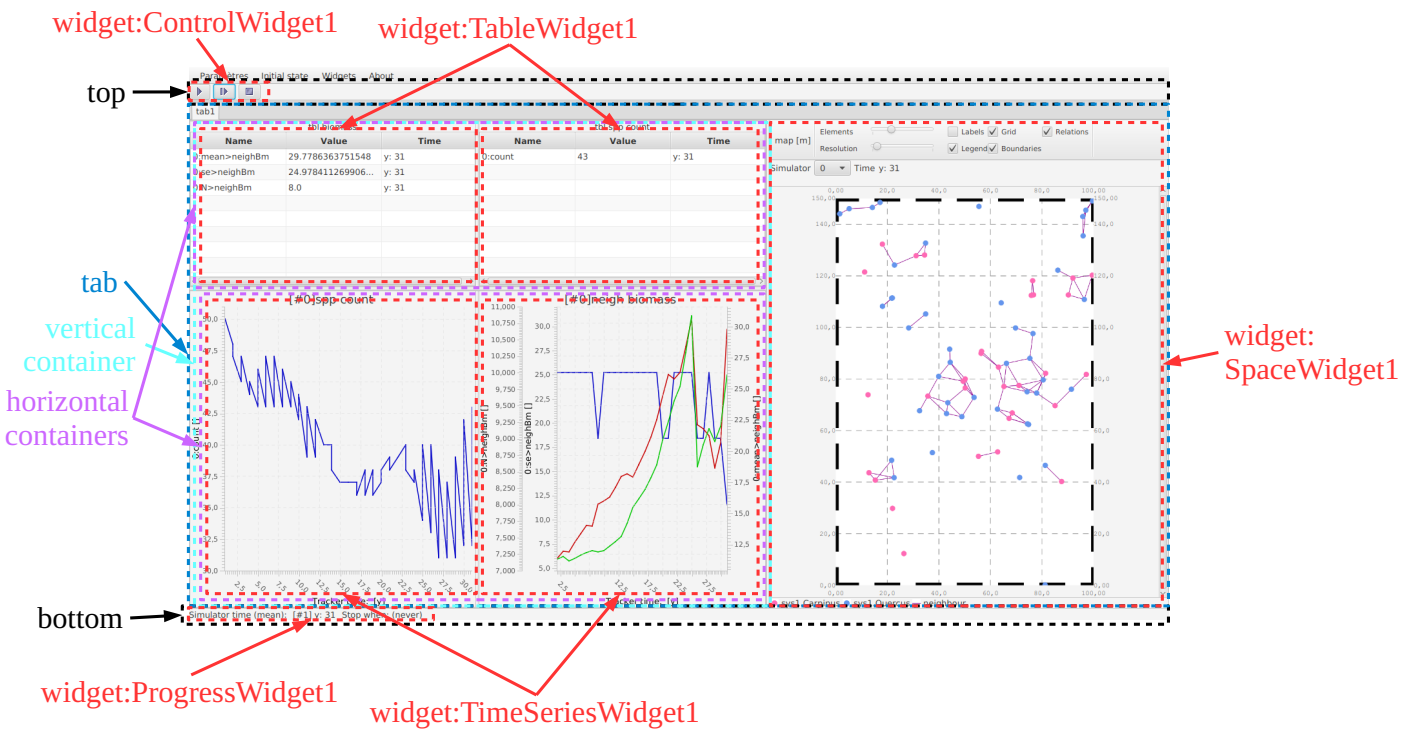


Figure 22. The ModelRunner graphical user interface.

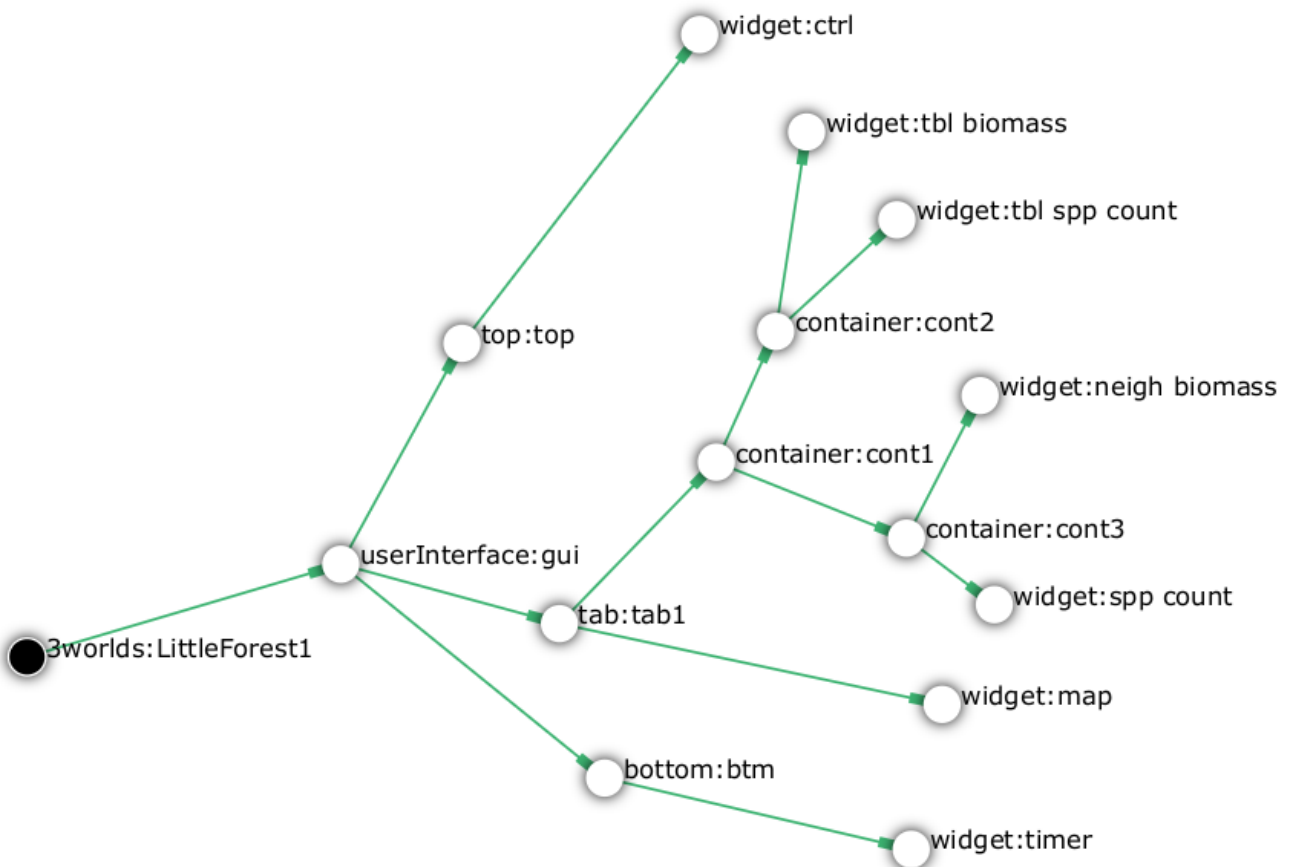


Figure 23. The configuration sub-tree specifying the GUI of Figure 22.

3Worlds provides a series of *widgets*, i.e. classes that can manage output from the simulator and either show it graphically or perform some other non-graphic task (e.g. save data to disk). Examples of graphic widgets are: time series or scatter plots, simulation control buttons, progress bars, maps etc... The basic ModelRunner GUI just provides places to place widgets: a *top*, a *bottom* and any number of *tabs*. Any number of widgets can be placed within these (Figure 22, Figure 23).

Top

```
/3worlds/userInterface/top:<name> {1}
```

This is a tool bar located at the top just below the menu bar. Widgets placed here will appear beside each other in an order specified in `ModelMaker` properties (*Order*). The top is typically a place where small widgets are added that remain accessible no matter where other widgets are placed. Typically a controller or progress widgets are placed here. Checks are made by `ModelMaker` to ensure only small widgets can be placed here (Figure 22).

Bottom

```
/3worlds/userInterface/bottom:<name> {0..1}
```

The *bottom* is a status bar located at the bottom of the main window. It has the same constraints as the *top*. Widgets such as a progress bar can be placed here if desired (Figure 22).

Tabs and containers

```
/3worlds/userInterface/tab:<name> {0..*}
```

```
/3worlds/userInterface/tab/container:<name> {0..*}
```

```
/3worlds/userInterface/.../container/container:<name> {0..*}
```

There can be any number of *Tabs* and each *Tab* can contain any number of widgets through the use of *containers*. *Tabs* are intended for large widgets such as charts and maps. Since only one tab is visible at a time when the model is run, large GUIs can be assembled without the limits imposed by window size (Figure 22). *Tab* contents are structured as a binary tree made of a combination of widgets or containers of widgets. A *Tab* can have:

- One or two widgets;
- One widget and one container; or,
- Two containers.

Containers have the same constraints as tabs - i.e. each container can have:

- One or two widgets;
- One widget and one container; or,
- Two containers.

Only widgets can be leaf nodes in this binary tree i.e you can't have a container or tab that contains nothing.

The widget/container pairs in this binary tree can be arranged vertically or horizontally. This allows all possible arrangements of widgets in the GUI.

Properties for tab :

orientation This property specifies the way widgets and containers are placed: side by side or on top of each other

possible values:

horizontal Display panel is split by a vertical splitter into two panels, first on left and second on right (default value)

vertical Display panel is split by a horizontal splitter into two panels, first above second

Properties for container :

orientation This property specifies the way widgets and containers are placed: side by side or on top of each other

order Rank of placement in its container (smaller value: to the left/top; larger: to the right/bottom)

Headless UI

/3worlds/userInterface/headless {0..*}

If you want to run a simulation unattended, you may not need a GUI at all. Instead, you may want the simulation to begin immediately and use widgets to write data to disk. Such widgets are call `headless`. All headless widgets are children of the `gui:headless:` node.

Widgets

/3worlds/userInterface/top/widget:<name> {0..*}

/3worlds/userInterface/bottom/widget:<name> {0..*}

/3worlds/userInterface/tab/widget:<name> {0..*}

/3worlds/userInterface/.../container/widget:<name> {0..*}

/3worlds/userInterface/headless/widget:<name> {0..*}

Widgets are the interesting part of the GUI configuration as they provide feedback and control of a simulation. A widget may be added as a child of `top`, `bottom`, `tab`, `container` or `headless` nodes.

Properties for widget :

order Rank of placement in its container (smaller value: to the left/top; larger: to the right/bottom)

subclass A widget class;

possible values:

ControlWidget1

A simple simulator controller with run/pause, step and reset buttons. Add this as a child of a `top` or `bottom` node. A GUI definition can only have one controller.

ControlWidget2

As for `ControlWidget1`, but also displays the time taken to complete each simulation step and the total simulation time. These timings are only accurate when deploying a single simulation. They are not accurate when deploying multiple simulations. Therefore, this widget is best used when developing a model to check its performance. Use `ControlWidget1` otherwise.

ControlWidget3

As for `ControlWidget2`, but displays a time series plot of simulation execution time. Add this widget as a child of a `tab` or `container` node.

ProgressWidget1

A simple display of simulation time (average time if deploying multiple simulations) and stopping condition. Add this widget as a child of a `top` or `bottom` node.

ProgressWidget2

A bar graph of simulation progress intended for use only when deploying multiple simulations. Add this widget as a child of a `tab` or `container` node.

ProgressWidget3

As for `ProgressWidget2` but using a time series chart instead of a bar chart.

TimeseriesWidget1

A graph of $x_i = f(t)$, where x_i are component descriptors and t is time. Add as a child of a `tab` or `container` node. This widget uses the fast charting library from [Chartfx](#) (<https://github.com/GSI-CS-CO/chart-fx>). The widget has a number of additional facilities accessed by moving the mouse to the top of the chart display. Here, data can be viewed as a table and exported to file. At the time of writing we have not found this feature very reliable (chartfx version 11.2.6). A better option to save time series data, is to use `HLEperimentWidget`.

ScatterplotWidget1

A graph of $y = f(x)$ where x and y are two descriptors of a component. Add as a child of a `tab` or `container` node. This widget also uses the fast charting library from [Chartfx](#) (<https://github.com/GSI-CS-CO/chart-fx>).

SpaceWidget1

A map of a space. Add as a child of a `tab` or `container` node.

TableWidget1

A table of number values $x_i(t)$. Add as a child of a `tab` or `container` node. Add as a child of a `tab` or `container` node.

MatrixWidget1

A colour display of a 2-dimensional table. Add as a child of a `tab` or `container` node.

GraphWidget1	A display of the system graph. Add as a child of a <code>tab</code> or <code>container</code> node. not yet implemented
HLEExperimentWidget1	Saves data to file. Add as a child of a <code>headless</code> node. The data written depends on the experiment design. A <code>SingleRun</code> produces a time series of each simulator and an average of replicates. <code>CrossFactorial</code> experiments produce the results of an ANOVA and <code>svg</code> (TODO) files of each factor's sensitivity. All statistical analysis uses <code>R</code> .
HLSimpleControlWidget	As <code>ControlWidget1</code> , but for unattended simulations. Add as a child of a <code>headless</code> node.
HLProgressWidget1	Writes simulator progress to the console.

Many widgets have settings which can be adjusted from the *Widget* menu of *ModelRunner*. These settings are automatically saved to a preferences file when *ModelRunner* is closed thus preserving the look and feel of the interface between uses.

Additional properties for widget sub-classes

for `ProgressWidget1`:

`refreshRate` (optional) The rate at which this widget updates its display (ms) (*default* 250 ms). This property is for the purpose of efficiency. Because this widget can display the average progress of a very large number of simulations (e.g 1 million parallel simulations), it would be very inefficient, not to mention uninformative, if the display was updated every time any one of the simulators sent a message.

for `TimeSeriesWidget1` and `TableWidget1`:

`nSenders` (optional) The number of parallel simulators to display (*default* 1). As noted, there can be far too many simulators to reasonably display their output. In such cases, this property limits that number to whatever you think you can handle.

`firstSender` (optional) The lowest sender (simulator id) to display (*default* 0). Output from simulators is shown for contiguously numbered simulators. This property sets the lowest number from which to display *n* simulator outputs.

for `TimeSeriesWidget1`:

`maxAxes` Sets the maximum number of y axes to display on the chart. As this widget can show time series for any number of variables, it is important to assign them to their own axis. However, if you have so too many outputs the chart will be all axes and no data. There, the number of axes can be constrained with this property.

`bufferSize` Number of time intervals to plot on the time axis, e.g. a value of 100 will display times 900-1000 when reaching time 1000. That is, what you see is a rolling buffer of data.

for `SpaceWidget1`:

`nViews` This widget records data produced from any number of simulators. However, it's often unreasonable to view every one of what may be a very large number. However, being able to visually compare independent parallel simulations is useful. This setting allows more than one (at the moment this is unlimited) view. A drop-down list for each view allows you to select which data set to display.

Additional cross-links for widget sub-classes

`for ControlWidget2, ControlWidget3, ProgressWidget1 and ProgressWidget2:`

`trackTime → dynamics:<name> {1}`

This link identifies the `dynamics` system the widget is listening to.

`for TimeSeriesWidget1, TableWidget1 and HLSimpleTimeSeriesWidget:`

`trackSeries → dataTracker:<name> {1}`

This link identifies the `dataTracker` the widget is listening to.

`for SpaceWidget1:`

`trackSpace → space:<name> {1}`

This link identifies the `space` the widget is listening to.

`for GraphWidget1:`

`trackSystem → system:<name> {1}`

This link identifies the `system` the widget is listening to

3.3.8. The *predefined* sub-tree

`/3worlds/predefined:*categories* {1}`

This sub-tree is not editable. It specifies predefined categories, relation types and automatic variables that are used in any 3Worlds model in association with particular objects in the simulator. During the setup of a configuration, ModelMaker will sometimes request you to link some of your nodes to items found in this hierarchy.

Predefined categories are grouped in 3 `categorySet`s:

- `*systemElements*` represent hierarchical levels in the system:
 - `*arena*` is a category attached to the whole system
 - `*component*` is a category attached to any system component
 - `*group*` is a category attached to any group of system components
 - `*life cycle*` is a category attached to any life cycle
 - `*relation*` is currently not in use
- `*organisation*` represent levels of organisation in the system:
 - `*atomic*`, is a category attached to single, indivisible (hence *atomic*), components
 - `*assemblage*` is a category attached to populations of components. Members of this category have the following automatic variables:
 - `count`, the population size
 - `nAdded`, the number of components added to the population during the last simulation step
 - `nRemoved`, the number of components removed from the population during the last simulation step

- ***atomic*** is a category attached to elementary (atomic) components that cannot be further divided
- ***lifespan*** represent the lifespan of a component in the system:
 - ***permanent*** , is a category for components that stay forever
 - ***ephemeral*** , is a category for components that can be created or deleted during a simulation. Members of this category have the following automatic variables:
 - `age` , the age of the component
 - `birthDate` , the simulation time when this component was created.

There are two predefined relation types:

- ***parentTo***: `*component* → *component*` may be used to relate a component to its offspring, i.e. it enables to track genealogic links between components (see the `CreateOtherDecision` function).
- ***returnsTo***: `*component* → *component*` is not yet in use.

3.4. Developing and testing model code

A model of an ecological system in 3Worlds consists in a *configuration file* constructed with the `ModelMaker` application (Section 3) and *java code files* that must be edited by the end-user, here a modeller. Some basic knowledge of `java` (https://en.wikiversity.org/wiki/Java_Programming/Introduction) is required before going on here.

3.4.1. Generated code: the *model main class*

During the specification of a model, ModelMaker generates java classes meant to be further edited in order to implement the specific behaviours imagined by the modeller. Among these classes, one is meant to be edited; the others are interface code, that you can see as 'glueing' code between the 3Worlds main code and the user-defined one. The file to be edited is named after the 3Worlds root node of the specification file (Section 3.3.1): e.g., if your 3Worlds node has the name `myModel` , the java file to edit will be called `MyModel.java` . In what follows, we call it the *model main class*.

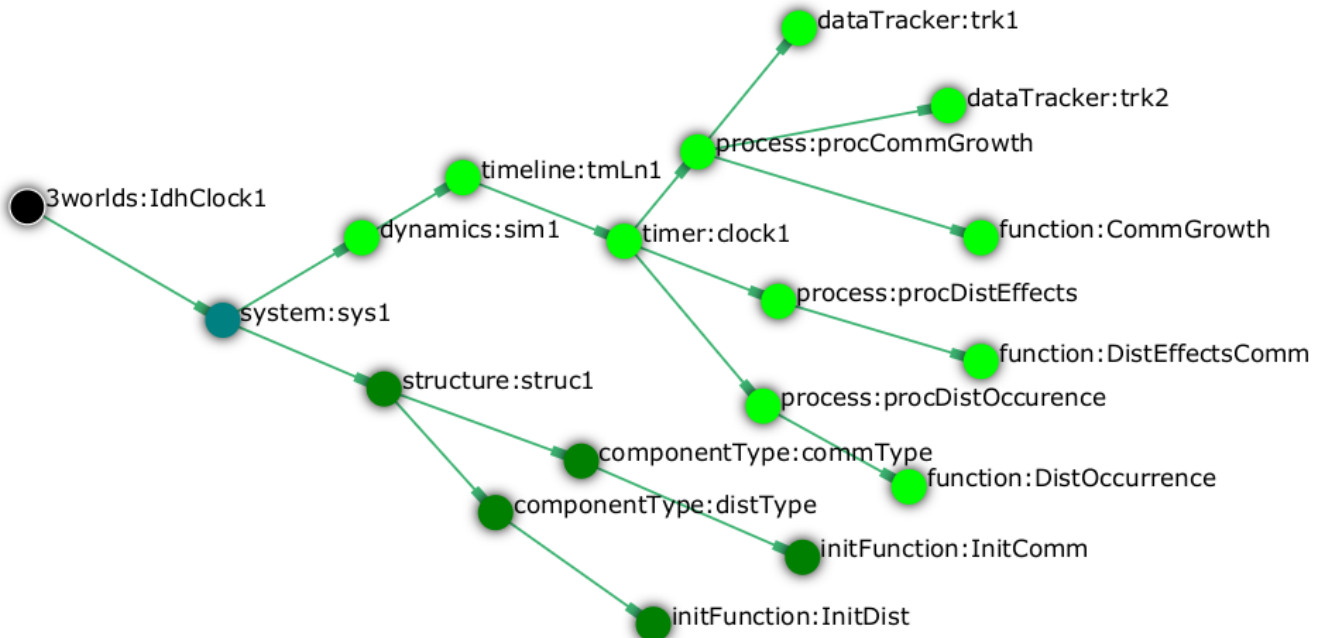


Figure 24. The functions defined in the model of the IDHClock tutorial.

This section gives general rules to follow to successfully edit the generated *model main class* in order for your model to behave as you wish. Section 2.4 describes how to link your `ModelMaker` session with an `eclipse` development environment where you can edit the generated *model main class*. It is important to read about functions before starting

writing your code, in order to master all the options you can setup in `ModelMaker` to specify your function.

For each `function` node of the specification, you will find one static method with the same name in the *model main class*. For example, the **IDHClock** tutorial (Section 5.4) specifies 3 functions, 2 `initFunctions` and 2 data trackers (Figure 24).

This generates a *model main class* with corresponding methods (code stripped of its javadoc comments):

JAVA

```

package code.sys1;

import static java.lang.Math.*;
import au.edu.anu.rscs.aot.collections.tables.DoubleTable;
import au.edu.anu.twcore.ecosystem.runtime.biology.DecisionFunction;
import java.util.Random;
import code.sys1.generated.*;
// Hey, model developer! You may add your own imports here as needed

public interface IdhClock1 {

    public static void distEffectsComm(
        double t,                                     // current time
        double dt,                                    // current time step
        int count,                                   // whole system autoVar count (#) ε[0..*]
        int nAdded,                                  // whole system autoVar nAdded (#) ε[0..*]
        int nRemoved,                                // whole system autoVar nRemoved (#) ε[0..*]
        double freq,                                 // focal component constants ± 0.0 ε]-∞,+∞[
        double inten,                                // focal component constants ± 0.0 ε]-∞,+∞[
        DoubleTable other_x,                         // other component drivers population size dim = [40] ± 0.0 ε]-∞,+∞[
        DistEffectsComm.OtherDrv otherDrv,          // next drivers for other component
        double other_div,                           // other component decorators ± 0.0 ε]-∞,+∞[
        DistEffectsComm.OtherDec otherDec,           // new decorators for other component
        DoubleTable other_K,                         // other component constants carrying capacity dim = [40] ± 0.0 ε]-∞,+∞[
        DoubleTable other_alpha,                     // other component constants interspecific competition coefficient
        DoubleTable other_r,                         // other component constants growth rate dim = [40] ± 0.0 ε]-∞,+∞[
        Random random) {                            // random number generator
        // distEffectsComm ---- Code insert Begin-->
        // distEffectsComm ---- Code insert End----<
    }

    public static boolean distOccurrence(
        double t,                                     // current time
        double dt,                                    // current time step
        int count,                                   // whole system autoVar count (#) ε[0..*]
        int nAdded,                                  // whole system autoVar nAdded (#) ε[0..*]
        int nRemoved,                                // whole system autoVar nRemoved (#) ε[0..*]
        double freq,                                 // focal component constants ± 0.0 ε]-∞,+∞[
        double inten,                                // focal component constants ± 0.0 ε]-∞,+∞[
        DoubleTable other_x,                         // other component drivers population size dim = [40] ± 0.0 ε]-∞,+∞[
        double other_div,                           // other component decorators ± 0.0 ε]-∞,+∞[
        DoubleTable other_K,                         // other component constants carrying capacity dim = [40] ± 0.0 ε]-∞,+∞[
        DoubleTable other_alpha,                     // other component constants interspecific competition coefficient
        DoubleTable other_r,                         // other component constants growth rate dim = [40] ± 0.0 ε]-∞,+∞[
        Random random,                               // random number generator
        DecisionFunction decider) {                  // decision function
        // distOccurrence ---- Code insert Begin-->
        // distOccurrence ---- Code insert End----<
    }

    public static void initDist(
        double freq,                                // focal component constants ± 0.0 ε]-∞,+∞[
        double inten,                                // focal component constants ± 0.0 ε]-∞,+∞[
        InitDist.FocalCnt focalCnt,                 // new constants for focal component
        Random random) {                            // random number generator
        // initDist ---- Code insert Begin-->
        // initDist ---- Code insert End----<
    }

    public static void commGrowth(
        double t,                                     // current time
        double dt,                                    // current time step
        int count,                                   // whole system autoVar count (#) ε[0..*]
        int nAdded,                                  // whole system autoVar nAdded (#) ε[0..*]
        int nRemoved,                                // whole system autoVar nRemoved (#) ε[0..*]
        DoubleTable x,                               // focal component drivers population size dim = [40] ± 0.0 ε]-∞,+∞[
        CommGrowth.FocalDrv focalDrv,              // next drivers for focal component
        double div,                                 // focal component decorators ± 0.0 ε]-∞,+∞[
        CommGrowth.FocalDec focalDec,              // new decorators for focal component

```

```

        DoubleTable K,
        DoubleTable alpha,
        DoubleTable r,
        Random random) {
    // focal component constants carrying capacity dim = [40] ± 0.0 ε]-∞,+∞[
    // focal component constants interspecific competition coefficient
    // focal component constants growth rate dim = [40] ± 0.0 ε]-∞,+∞[
    // random number generator

    // commGrowth ---- Code insert Begin-->
    // commGrowth ---- Code insert End----<
}

public static void initComm(
    DoubleTable x,
    InitComm.FocalDrv focalDrv,
    double div,
    DoubleTable K,
    DoubleTable alpha,
    DoubleTable r,
    InitComm.FocalCnt focalCnt,
    Random random) {
    // focal component drivers population size dim = [40] ± 0.0 ε]-∞,+∞[
    // next drivers for focal component
    // focal component decorators ± 0.0 ε]-∞,+∞[
    // focal component constants carrying capacity dim = [40] ± 0.0 ε]-∞,+∞[
    // focal component constants interspecific competition coefficient
    // focal component constants growth rate dim = [40] ± 0.0 ε]-∞,+∞[
    // new constants for focal component
    // random number generator

    // initComm ---- Code insert Begin-->
    // initComm ---- Code insert End----<
}

}

```

In this example, you can see that:

- the package name `code.sys1` is constructed from the name of the `system` node (Figure 24);
- the interface name `IDhClock1` is constructed from the name of the `3Worlds` node (Figure 24);
- each method name is constructed from a matching `function` or `initFunction` node (Figure 24).
- the argument lists are partly constructed from the `categories` the process declaring the `function` applies to;
- the comments documenting the method arguments are constructed from the `field` or `table` node properties (`description`, `precision`, `interval`, `units`, etc.).



Do not neglect these metadata: coding errors due to discrepancies in measurement units between equations are frequent and yield wrong computation results that are difficult to trace. Most publicly available model code does not document the units, and hence is not easily re-usable.

When ModelRunner is launched on the IDHClock tutorial model, it will include the generated `IDhClock1` class and call each of its methods for all system components they are dealing with as specified in the model configuration file.

As you can see in this example, the body of each method is empty, only containing two comments:

```
// initComm ---- Code insert Begin-->
// initComm ---- Code insert End----<
```

JAVA

These are the *code insertion markers*. The user-defined code must be inserted between these two lines.



Never remove the *code insertion markers* as they are used by 3Worlds when using code snippets (as in all tutorial and test models).

As *model main class* is a java **interface**, all data is passed as arguments to its static methods. As you can see in the example above, there may be many arguments. If you look closely, you will see that these arguments match the *descriptors* that were attached to the *categories* to which the *processes* apply. All this information is provided in the *model main class* as javadoc comments. For example, the javadoc comment of the `commGrowth` method above produces this output:

commGrowth

```
static void commGrowth(double t, double dt, int count, int nAdded, int nRemoved,
au.edu.anu.rscs.aot.collections.tables.DoubleTable x,
code.sys1.generated.CommGrowth.FocalDrv focalDrv, double div,
code.sys1.generated.CommGrowth.FocalDec focalDec,
au.edu.anu.rscs.aot.collections.tables.DoubleTable K,
au.edu.anu.rscs.aot.collections.tables.DoubleTable alpha,
au.edu.anu.rscs.aot.collections.tables.DoubleTable r, java.util.Random random)
```

CommGrowth method of type *ChangeState*: change the state, i.e. the values of the descriptors of a system component

- applies to categories { *commCat* }
- follows timer *clock1* of type *ClockTimer*, with time unit = 1 t.u
- called before function *distEffectsComm*(...).

Parameters:

t - current time

dt - current time step

count - whole system autoVar count (#) ε[0..*]

nAdded - whole system autoVar nAdded (#) ε[0..*]

nRemoved - whole system autoVar nRemoved (#) ε[0..*]

x - focal component drivers population size dim = [40] ± 0.0 ε]-∞,+∞[

focalDrv - next drivers for focal component

div - focal component decorators ± 0.0 ε]-∞,+∞[

focalDec - new decorators for focal component

K - focal component constants carrying capacity dim = [40] ± 0.0 ε]-∞,+∞[

alpha - focal component constants interspecific competition coefficient dim = [40,40] ± 0.0 ε]-∞,+∞[

r - focal component constants growth rate dim = [40] ± 0.0 ε]-∞,+∞[

random - random number generator

This comment recalls the categories to which the `commGrowth` method applies, which timer it follows and which time units it uses, and any other useful information like precedence between methods as specified by `dependsOn` cross-links between processes.

Finally, the *model main class* itself has a general javadoc description that gives some information about how to insert useful code into its methods:

Package code.sys1

Interface **IdhClock1**

public interface **IdhClock1**

Model-specific code for model **IdhClock1**

version demo - Tue Apr 27 15:07:13 CEST 2021

Authors:

Jacques Gignoux

Ian D. Davies

Contacts:

jacques.gignoux@upmc.fr

Ian.Davies@anu.edu.au

Reference publication:

https://en.wikipedia.org/wiki/Competitive_Lotka%20Volterra_equations

Instructions to model developers:

1. Non 3worlds-generated extra methods should be placed in other files linked to the present file through imports.
2. **Do not** alter the code insertion markers. They are used to avoid losing your code when managing this file.
3. For convenience, all the static methods of the **Math** and **Distance** classes are directly accessible here
4. The particular random number stream attached to each **TwFunction** is passed as the *random* argument.
5. For all *Decision*- functions, a *decider* argument is provided to help make decisions out of probabilities.
decider.decide(double) returns true with probability equal to the argument.
6. For *ChangeCategoryDecision* functions, a *selector* argument is provided to select among different possible outcomes.
selector.select(double...) returns an integer between 0 and *n* (the number of arguments) using the arguments as weights for probabilities (ie the argument do not need to sum to 1).
7. For *ChangeCategoryDecision* functions, a *recruit* argument is provided that must be used to return the proper category name as a String. **recruit.transition(boolean)** will return the category to recruit to if the argument is true, triggering the change in category of the focal SystemComponent. **recruit.transition(int)** will return the category name matching the index using alphabetical order, 0 index meaning no change in category. For example, if the decision may result in category "young" or "juvenile", 0 will map to no change, 1 to change to juvenile and 2 to change to young.

3.4.2. Model main class method arguments

The list of arguments of each method is defined by its function type, the organisation level to which it applies (system, life cycle, group or component), the categories or relation types it applies to, and the user-defined data structures attached to these. Some arguments are read-only, others are writeable so that computation output can be passed back to the 3Worlds main code.

Read-only arguments

Arguments present for all *functions* and *initFunctions*

random The random number generator channel associated to this function. For details of how to use an instance of the class `java.util.Random`, see the [javadoc](https://docs.oracle.com/en/java/javase/11/docs/api/java.base/java/util/Random.html) (<https://docs.oracle.com/en/java/javase/11/docs/api/java.base/java/util/Random.html>) for this class. Most of the time, you will be calling `random.nextDouble()` which returns a random double value between 0.0 and 1.0.

Arguments present for all *functions* but not for *initFunctions*

t the *current time* passed by the simulator as a double value in units of the `timer` of the parent `process` of the function.

`dt` the *current time step*, passed by the simulator as a double value in units of the `timer` of the parent process of the `function`.



In the case of multiple timers, for `ClockTimers`, the current time step may be different from the timer's `dt` property because it is the time since last simulator iteration, which may have been triggered by a different timer.

Arguments present for all '*decision*' functions

'Decision' functions are: `ChangeCategoryDecision`, `DeleteDecision`, `CreateOtherDecision`, `RelateToDecision` and `MaintainRelationDecision`. They all return a result that is a decision: a number of components to create (`CreateOtherDecision`), the name of a category (`ChangeCategoryDecision`), or a boolean (all others).

`decider` This argument of class `DecisionFunction` **[TODO: ref to javadoc]** is provided as a helper for transforming probabilities into decisions. This class comprises only one method `decide(...)` which given a probability, returns a `boolean`. More precisely: it returns `true` with the probability passed as argument, ie `decider.decide(0.7)` will return `true` in 7 calls out of 10. It uses the function random number generator (the `random` argument) to make the decision. Technically, this is the realisation of a Bernoulli distribution (https://en.wikipedia.org/wiki/Bernoulli_distribution).

`selector` This argument is only present for the `ChangeCategoryDecision` function type. The `SelectionFunction` class **[TODO: ref to javadoc]**, of which it is an instance, only has one method `select(...)` which, given a list of weights w , returns an integer i with probability $w[i]/\sum_j w[j]$, i.e. a realisation of a single trial of a multinomial distribution (https://en.wikipedia.org/wiki/Multinomial_distribution).

`recruit` This argument is only present for the `ChangeCategoryDecision` function type. The `RecruitFunction` class **[TODO: ref to javadoc]**, of which it is an instance, has one method `transition(...)` which returns a category name (`String`), or `null` if the component does not change category. Its argument is either a `boolean` or an `int`, typically the result of a call to `selector.select(...)` or `decider.decide(...)`. Example of use:

```
public static String recruitSeedling(
    double group_recruitRate,
    ...
    Random random,                                // random number generator
    DecisionFunction decider,                      // decision function
    SelectionFunction selector,                    // selection function
    RecruitFunction recruit) {                     // recruitment function

    return recruit.transition(decider.decide(group_recruitRate));
}
```

JAVA

Arguments that represent a component in function types which process applies to categories

These function types are `ChangeCategoryDecision`, `ChangeState`, `DeleteDecision`, `CreateOtherDecision`, and `SetInitialState`.

Internally, the system component which is the target of such functions is called *focal*.

The argument list will contain all the fields and tables declared in the root record of the descriptors (drivers, automatic variables, constants and decorators) of the *focal* component. The argument comments will indicate that these arguments are descriptors of the *focal* component, as in this example from the **IDHClock** tutorial for a function of type `SetInitialState`:

```

public static void initComm(
    DoubleTable x,                                     // focal component drivers population size dim = [40] ± 0.0 €]-∞,+∞[
    double div,                                       // focal component decorators ± 0.0 €]-∞,+∞[
    DoubleTable K,                                     // focal component constants carrying capacity dim = [40] ± 0.0 €]-∞,+∞[
    DoubleTable alpha,                                // focal component constants interspecific competition coefficient dim
    DoubleTable r,                                     // focal component constants growth rate dim = [40] ± 0.0 €]-∞,+∞[
    Random random) {                                 // random number generator
    ...
}

```

Arguments that represent the two components of a relation in function types which process applies to relation types

These function types are `ChangeOtherState`, `ChangeRelationState`, `MaintainRelationDecision`, `RelateToDecision`, and `SetOtherInitialState`.

These functions apply to a pair of components linked by a relation. The first of these components (the one at the 'from' end of the relation) is called *focal*, as before. The second one (the one at the 'to' end of the relation) is called *other*.

To distinguish the descriptors of *other* from those of *focal* (since these might belong to the same categories and hence have the same descriptors), all the descriptors of *other* are prefixed with 'other_-, as in this example from the **IDHClock** tutorial for a function of `ChangeOtherState` type:

```

public static void distEffectsComm(                                     JAVA
    double t,                                         // current time
    double dt,                                        // current time step
    double freq,                                       // focal component constants ± 0.0 €]-∞,+∞[
    double inten,                                      // focal component constants ± 0.0 €]-∞,+∞[
    DoubleTable other_x,                                // other component drivers population size dim = [40] ± 0.0 €]-∞,+∞[
    double other_div,                                  // other component decorators ± 0.0 €]-∞,+∞[
    DoubleTable other_K,                                // other component constants carrying capacity dim = [40] ± 0.0 €]-∞,+∞[
    DoubleTable other_alpha,                             // other component constants interspecific competition coefficient dim
    DoubleTable other_r,                                // other component constants growth rate dim = [40] ± 0.0 €]-∞,+∞[
    Random random) {                                 // random number generator ...
}

```

Arguments that represent the local context of a component

The local context of a component is the part of the system it always sees. The dynamic graph of a simulated system in 3Worlds **always** has:

- an object representing the whole system, called the *arena* ('the place where things happen': [Gignoux et al. 2011](https://doi.org/10.1007%2Fs10021-011-9466-2) (<https://doi.org/10.1007%2Fs10021-011-9466-2>)). As 3Worlds uses a dynamic graph to represent the whole system (Section 1.2.2), this object actually represents the whole graph. It matches the *system* node in the configuration graph of the `.ugt` file.

And it **may** (0..* multiplicity) have the following other kinds of nodes:

- objects representing the individual entities of the system, which are nodes in the dynamic graph and are just called *components*.
- objects that represent the common properties of a group of components, which are called *groups*.
- objects that represent the transitions that can occur during the life of a component, and are therefore called *life cycles*.

These four types of objects may belong to *categories* and have *descriptors*, and as a consequence can be passed to *functions* of a *process* referencing their categories.

With regard to processes and functions, these objects are treated like components, except they cannot establish *relations*; only true components can. This limits the set of functions compatible with them: groups, life cycles and the

arena can only be affected by `ChangeState`, `SetInitialState` and `CreateOtherDecision` (not life cycles) function types. When these functions apply to the categories of a `group`, `lifeCycle` or `Arena` object, they are treated as components above, i.e. they become the *focal* 'component' of the user-defined methods.

The arena, groups and life cycles play a particular role in the dynamic graph, as indicated by their names. They also have implicit, 'ontological' relations with system components: a component always knows about its group, life cycle, and arena because they describe part of its own behaviour. As such, they are always accessible as arguments in the function calls of any component.

Just as for the *other* component above, the descriptors of arena, life cycles and groups are prefixed when they appear in a method argument list, with a comment giving more information on the argument, as in this example from the **Palms** show-case model for a function of `CreateOtherDecision` type:

```
public static double reproduction()
    double t,                                     // current time
    double dt,                                    // current time step
    int count,                                   // whole system autoVar count (#) ∈[0..*]
    int nAdded,                                  // whole system autoVar nAdded (#) ∈[0..*]
    int nRemoved,                                // whole system autoVar nRemoved (#) ∈[0..*]
    int lifeCycle_count,                         // focal life cycle autoVar count (#) ∈[0..*]
    int lifeCycle_nAdded,                        // focal life cycle autoVar nAdded (#) ∈[0..*]
    int lifeCycle_nRemoved,                      // focal life cycle autoVar nRemoved (#) ∈[0..*]
    int group_count,                             // focal group autoVar count (#) ∈[0..*]
    int group_nAdded,                            // focal group autoVar nAdded (#) ∈[0..*]
    int group_nRemoved,                          // focal group autoVar nRemoved (#) ∈[0..*]
    double group_aGinc,                           // focal group constants adult BudHeight growth coefficient (m yr-1) ±
    double group_aPdead,                          // focal group constants Mortality : int ± 0.01 ∈[0.0,1.0]
    double group_deadNbLeaves,                   // focal group constants Mortality : nbf slope ± 0.01 ∈[0.0,1.0]
    double group_decay,                           // focal group constants decay ± 0.0 ∈]-∞,+∞[
    double group_dis,                            // focal group constants dispersal parameter ± 0.001 ∈[0.0,1.0]
    double group_fec,                            // focal group constants fecundity * # leaves ± 0.1 ∈[0.0,+∞[
    double group_jPdNNeg,                         // focal group constants Adult P(dN=-1) ± 0.01 ∈[0.0,1.0]
    double group_remanence,                      // focal group constants seedling remanenc (y) ± 0.0 ∈]-∞,+∞[
    double group_sirec0,                          // focal group constants rect els to sls alive ± 0.01 ∈[0.0,1.0]
    double group_sirec1,                          // focal group constants rect els to sls dead ± 0.01 ∈[0.0,1.0]
    double budHt,                               // focal component currentState ± 0.0 ∈]-∞,+∞[
    int dead,                                   // focal component currentState dead ∈[MIN_INTEGER..*]
    double nELSeedlings,                         // focal component currentState Nb. EL seedlings ± 0.0 ∈[0.0,+∞[
    int nleaves,                                // focal component currentState nleaves ∈[0..*]
    double neighbourhoodIndexAdults,             // focal component decorators neighbourhoodIndexAdults ± 0.01 ∈]-∞,+∞[
    double neighbourhoodIndexJuveniles,           // focal component decorators neighbourhoodIndexJuveniles ± 0.0 ∈]-∞,+∞[
    double neighbourhoodIndexMounds,              // focal component decorators neighbourhoodIndexMounds ± 0.0 ∈]-∞,+∞[
    double neighbourhoodIndexTrees,               // focal component decorators neighbourhoodIndexTrees ± 0.0 ∈]-∞,+∞[
    boolean sex,                                 // focal component constants female?
    double x,                                    // focal component constants x spatial coordinate (m) ± 0.1 ∈[0.0,300.
    double y,                                    // focal component constants y spatial coordinate (m) ± 0.1 ∈[0.0,300.
    Random random,                             // random number generator
    DecisionFunction decider) {                  // decision function
    ...
}
```

The code generator decides which arguments must be made available to a method based on this hierarchy: **arena > life cycle > group > component**. When the method applies to a component, then all four hierarchical levels are accessible; when it applies to a group, only arena and life cycle descriptors are available, and the group arguments are treated as the *focal* 'component'.

Space data

When a space is attached to a process through a `inSpace` cross-link, the following argument is added to the method list:

limits The limits of the space used with this process. This is an immutable object of class `Box` [TO DO: link to javadoc]. It returns the coordinates of the lower and upper ends of the space in all its dimensions through the `limits.lowerBound(int i)` and `limits.upperBounds(int i)` methods, where `i` is the dimension index.

Writable arguments and method return values

User-defined functions are meant to modify the state of the graph, which technically means induce changes in descriptor values and creation/deletion of graph elements, namely components and relations. This cannot be done with read-only arguments.

Method return values

Decision functions all have return values that are interpreted as follows:

function type	return type	return value
<code>ChangeCategoryDecision</code>	<code>String</code>	<code>null</code> or a new Category
<code>DeleteDecision</code>	<code>boolean</code>	<code>true</code> if <code>focal</code> is to be deleted
<code>CreateOtherDecision</code>	<code>double</code>	the number of new components to create; the decimal part is interpreted as a probability to create an extra component.
<code>RelateToDecision</code>	<code>boolean</code>	<code>true</code> if <code>other</code> is to be related to <code>focal</code>
<code>MaintainRelationDecision</code>	<code>boolean</code>	<code>true</code> if the existing relation between <code>focal</code> and <code>other</code> is maintained

They all have read-only helper arguments. Example from the **IDHClock** tutorial for a `RelateToDecision` function:

```
public static boolean distOccurrence(
    double t,                                     // current time
    double dt,                                     // current time step
    ...
    double freq,                                    // focal component constants ± 0.0 ∈]-∞,+∞[
    ...
    Random random,                                // random number generator
    DecisionFunction decider) {                   // decision function
    // distOccurrence ---- Code insert Begin-->
    return decider.decide(1.0/freq);
    // distOccurrence ---- Code insert End---<
}
```

JAVA

Method writeable arguments

Change/Set-State functions have the following writeable arguments as output:

function type	writeable arguments
<code>setInitialState</code>	<code>focal</code> constants & drivers
<code>changeState</code>	<code>focal</code> drivers & decorators
	decorators of <code>arena</code> , <code>life cycle</code> & <code>group</code>
<code>setOtherInitialState</code>	<code>other</code> drivers & constants

function type	writeable arguments
ChangeOtherState	<i>other</i> drivers & decorators
	decorators of <i>arena</i> , <i>life cycle</i> , <i>group</i> , <i>other life cycle</i> & <i>other group</i>
ChangeRelationState	<i>focal</i> drivers & decorators
	<i>other</i> drivers & decorators
	decorators of <i>arena</i> , <i>life cycle</i> , <i>group</i> , <i>other life cycle</i> & <i>other group</i>

These arguments appear in the argument list as specific inner classes instances with self-explained names: `focalDrv`, `focalCnt`, `focalDec`, `groupDec`, `groupCnt`, `arenaDrv`, `otherGroupDrv`, etc... Each of these arguments will contain the same fields as the original data structure it comes from. Any value set in these arguments will be carried back to the component when the method returns. For example, this is the generated code from the **IDHClock** tutorial for a function of type `ChangeOtherState`:

```
public static void distEffectsComm(
    double t,                                     // current time
    double dt,                                    // current time step
    int count,                                   // whole system autoVar count (#) ∈[0..*]
    int nAdded,                                  // whole system autoVar nAdded (#) ∈[0..*]
    int nRemoved,                                // whole system autoVar nRemoved (#) ∈[0..*]
    double freq,                                 // focal component constants ± 0.0 ∈]-∞,+∞[
    double inten,                                // focal component constants ± 0.0 ∈]-∞,+∞[
    DoubleTable other_x,                         // other component drivers population size dim = [40] ± 0.0 ∈]-∞,+∞[
    DistEffectsComm.OtherDrv otherDrv,           // next drivers for other component
    double other_div,                            // other component decorators ± 0.0 ∈]-∞,+∞[
    DistEffectsComm.OtherDec otherDec,           // new decorators for other component
    DoubleTable other_K,                         // other component constants carrying capacity dim = [40] ± 0.0 ∈]-∞,+∞[
    DoubleTable other_alpha,                     // other component constants interspecific competition coefficient dim
    DoubleTable other_r,                         // other component constants growth rate dim = [40] ± 0.0 ∈]-∞,+∞[
    Random random) {                            // random number generator
    ...
}
```

JAVA

In this example, the writeable arguments are `otherDrv` and `otherDec`. Both of them are of ad-hoc inner classes defined in the generated sub-directory of the eclipse project for **IDHClock** (file `DistEffectsComm.java`):

```
public class OtherDec {
    public double div;
}

public class OtherDrv {
    public DoubleTable x;
}
```

JAVA

Here, to set the `div` field of the *other* decorators to 3.2, simply write `otherDec.div=3.2;` in your method code.



As you can see above, the writeable arguments are also present as read-only arguments.

You may also have noticed in the example above that the comment besides `otherDec` is `// new decorators for other component`, while that besides `otherDrv` is `// next drivers for other component`. What does this mean? Well, this is just a reminder that decorators and drivers are not handled in the same way by 3Worlds (cf. categories):

- **Decorator values** are only valid within a time step and are immediately changed, which means that any method

writing a decorator value will change it for all other methods called during the same time step. Hence the 'new' adjective.

- **Driver values** are carried over to the next time step, and thus are subject to a synchronized modification. To keep all component states consistent, the read-only argument for drivers store the values that were set at the previous time step, i.e. they represent the *current* state of all components. When a modification of their value is computed, it is stored in the proper writeable argument that will be copied into a separate driver state representing the *next* state of the component. *This makes sure that all components within a time step are viewed by each other in the same, consistent state.* It is only after all methods on all components of the current time step have been called that 3Worlds will replace the *current* driver state with the *next* driver state. Hence the 'next' adjective.



Forgetting about the difference of treatment between *decorators* and *drivers* can be the source of **major** but difficult to detect computation errors. Always know what you do!

3.4.3. Using space in user code

3Worlds assumes that an ecosystem representation can require a detailed description of zero, one or *many* spaces ([Gignoux et al. 2011](https://doi.org/10.1007%2Fs10021-011-9466-2) (<https://doi.org/10.1007%2Fs10021-011-9466-2>)) - depending on the question addressed. It further assumes that a given space is best designed to suit the needs of one particular process, in other words, there is a `1..*` relation between a space and various processes (a space can be used by many processes, but a process only uses one space). The relation between a process and a space is specified with the `inSpace` cross-link (see processes).

Spaces are used:

1. to *locate* system components and help *compute geometric quantities* within their space. This is done by passing spatial data as arguments to all functions of a process having an `inSpace` link to a space (ie the limits argument). A component location in its space is given by the value of specific driver or constant descriptors (see cross-links for space). Any change in these coordinates will result in a movement of the component into this space.



When component coordinates get out of the space limits, the selected *edge effect correction* rules apply and will either result in the return of the component within space limits or its exclusion from the model. This is done internally by 3Worlds just after the user method call, so that you don't need to bother if your computed coordinates get out of range.

2. to optimise the search of candidate components to establish a relation with a `RelateToDecision` function. This is an old computing problem: how to find closest neighbours in a geometric space *efficiently*? It has been solved long ago by using [Kd-trees](https://en.wikipedia.org/wiki/K-d_tree) (https://en.wikipedia.org/wiki/K-d_tree). Kd-trees are the most efficient algorithms to find nearest neighbours ($O(\log(n))$ time). In continuous spaces (cf. space library), 3Worlds uses Kd-trees to search for closest neighbour first to establish relations. Depending on the property defined in the `inSpace` cross-link (`searchRadius` or `searchNeighbours`) 3Worlds will either present:

- the list of all components found within `searchRadius` space units of the *focal* component; or
- the list of the `searchNeighbours` first closest components found around the *focal* component;

...to the `RelateToDecisionFunction`. In other words, the use of the space will restrict the number of candidates for establishing a relation by looping only on those *others* which are close enough to *focal*, using two different search methods.



Whenever you have to search for items to construct relations, associate a space to your process, it will make computations far more efficient.

3.4.4. Accessing complex data structure

The code generated by `ModelMaker` also comprise specific data structure classes for component, group, life cycle and arena descriptors. These classes are put in the `generated` sub-directory of the project `code/<system-name>/`

directory in the user-defined java project. Have a look at these files (but do not edit them!) to understand how to access data in complex hierarchical structures.

In all cases, those data structures have getters and setters for fields and tables that use the name of the field as method name. For example, if `x` is a field in record `B`, it will be accessed as `B.x()` and set with `B.x(value)`.

3.5. Feeding the model with data

Simulators usually rely on possibly large data sets to initialise and run a simulation.

3.5.1. Data uses

According to their use during a simulation, we can classify data into:

Parameters	Values usually visible to a large number of system components that remain constant over a simulation or over the life time of an ephemeral component. These are stored in the constants of components, groups, life cycles and the arena.
-------------------	---

Examples: the *average solar radiation at the top of the canopy* for the arena component; the *spatial coordinates* of a plant; the *species name* for a group of components.

Initial community	In most cases a simulation must start with a non-empty initial system. This implies that components be instantiated with specific initial values for their drivers and proper connections between them.
--------------------------	---

Examples: an initial forest reconstructed from LIDAR data; a monitored population of marked animals within an area, where each animal has a measured age.

Forcing data	Sometimes a simulation requires that data be fed in during its time course, at some time step, replacing values of a driver that cannot be computed by the model.
---------------------	---

Examples: a series of meteorological data used to force a fire spread or a vegetation growth model.

Data for *parameters* and *initial community* are directly used to instantiate components and fill their descriptors with values. *Forcing data* must be loaded into some initial data structure (probably most often a `DoubleTable`, cf. Section 3.3.5.2.2, as a constant in the `system` instance), made accessible to some user-defined function that will read its content as the simulation proceeds.

3.5.2. Data input methods

There are three different ways of feeding `ModelRunner` with data to start a simulation experiment (cf. Section 3.3.3.4 for the specification of these):

1. Entering parameter and initial driver values in the model configuration tree. This is useful only for small quantities of data, (i) with models requiring only a few parameters, or (ii) during the initial building/testing file of a model; typically when one does not want to bother with formatting an input file with only two parameter values. This is specified using `initialValues` nodes as children of `system`, `componentType`, `groupType` or `lifeCycleType`, depending on where the data should go (Section 3.3.2 and Section 3.3.3).
2. Loading data from an external source, typically, one or more files. This is recommended whenever a large number of initial values and items must be used. This is specified using `loadFrom` cross-links to `dataSource` nodes (Section 3.3.6.4.1).
3. Using a user-defined initialiser function to set or modify initial values. This complements both points above and enables to compute transformations of initial drivers or constants prior to run the simulation. This is specified using `initFunction` nodes of the `SetInitialState` type (Section 3.3.4.2.2) as children of `system`,

`componentType`, `groupType` or `lifeCycleType`, depending on where the data should go (Section 3.3.2 and Section 3.3.3).

These three methods can be used together within the same model. However, since an initialiser function must be run on an existing instance of a component, it can only be used (i) on the `system` node, which is always instantiated and unique, or (ii) as a complement to the two other methods which can specify the component instances to create.

The `loadFrom` cross-link

`loadFrom` → `dataSource:<name>` {0..*}

A `dataSource` node specifies the information required to access some data source: for example, a file name, a file format, and all other requested information that the java code just needs to read the file.

A `loadFrom` cross-link links the `system`, a `componentType`, `groupType` or `lifeCycleType` a to a data source.

The `initialValues` node

/3worlds/.../componentType/initialValues:<name>

/3worlds/.../groupType/initialValues:<name>

/3worlds/.../lifeCycleType/initialValues:<name>

/3worlds/system/initialValues:<name>

An `initialValues` node enables one to directly specify the number, driver and constant values, belonging to groups and life cycle instances, of initial instances of components, groups, life cycle, and the arena.

Properties for initialValues :

`nInstances` The number of instances of `componentType`, `groupType` or `lifeCycleType` to create at the start of the simulation. For groups and life cycle, the instance name will be the name of the `initialValues` node, plus a number if more than one instance is required. For components, the unique identifier is automatically generated. Note that a value of zero will not create any instance. Ignored for the arena (always present and unique).

`idGroup` (only applies to `.../groupType/componentType/initialValues` nodes): The `groupType` instance name of these `ComponentType` instances

`idLifeCycle` (only applies to `.../lifeCycleType/groupType/componentType/initialValues` and `.../lifeCycleType/groupType/initialValues` nodes): The `lifeCycleType` instance name of these `ComponentType` or `groupType` instances

`<name>` driver and constant (name,value) pairs to populate the instances. The property name must match a declared field. Tables are not supported.



The latter property is only partly supported and at the time of writing, `ModelMaker` does not enable one to write such properties. One has to edit the configuration file by hand. Only do that if you are an expert.

The `initFunction` node

/3worlds/.../componentType/initFunction:<name>

/3worlds/.../groupType/initFunction:<name>

/3worlds/.../lifeCycleType/initFunction:<name>

/3worlds/system/initFunction:<name>

Properties for initFunction :

type The type of function used: only `setInitialState` is possible.

functionSnippet The java code to be inserted into the body of the `initFunction` method in the generated code.

Cross-links for initFunction :

useRNG → `rng:<name> {0..1}`

Points to the random number generator channel to use in this function. If unset, the default random number channel is used.

3.5.3. Consistency of data files with the model component hierarchy

The simulated system is a hierarchy with up to 4 levels: arena > life cycle > group > component (cf. Box How is the simulated system implemented?). As a consequence, input data files must reflect this hierarchy.

3.5.4. Data file formats

The data file formats currently supported as input to 3Worlds are listed in Section 3.3.6.4.1. At the moment, `.csv` text files and `.ods` spreadsheet files are supported. Their content must comply with constraints listed in Section 3.3.6.4.1.

For a practical understanding of how this works, we recommend to run the `TestDataLoading` model (accessible in **ModelMaker** through the `Projects > New > Test cases` menu).

Here are examples extracted from this test model:

Example 1: using a *dataSource* only

This `.ods` spreadsheet:

Screenshot of a LibreOffice Calc spreadsheet application showing a table of climate data.

The menu bar includes: Fichier, Édition, Affichage, Insertion, Format, Styles, Feuille, Données, Outils.

The toolbar includes: File, Print, Find, Cut, Copy, Paste, Undo, Redo, Font, Alignment, Number, Cell Style, and others.

The formula bar shows cell C12 selected with value 17.

The table has columns A through F:

	A	B	C	D	E	F
1	lat	long	temperature	otherVar:windspeed	otherVar:windDir	otherVar:rainfall
2	0	0	21.8	3.1	C	8
3	0	1	21.4	2.9	C	0
4	0	2	23	5.3	C	2
5	0	3	24	3.1	C	0
6	0	4	19.4	3.9	C	16
7	0	5	20.5	3.9	C	17
8	0	6	22	2	C	5
9	0	7	18	2.6	C	17
10	1	0	21.3	1.2	C	0
11	1	1	22.5	0.4	C	0
12	1	2	17	1.6	C	4
13	1	3	23.4	3.6	C	0
14	1	4	23.5	6.6	C	13
15	1	5	19	8.4	C	0
16	1	6	24	0.3	NW	11
17	1	7	23.6	0.6	C	0
18	2	0	19	0.9	C	0
19	2	1	25.2	1.3	N	23
20	2	2	24.5	0	C	0
21	2	3	19.9	0.1	C	35
22	2	4	25.4	1.9	C	0
23	2	5	25	4	C	0
24	2	6	21.3	3.1	C	2
25	2	7	25.3	21	C	0
26	2	8	25.3	0.3	C	2

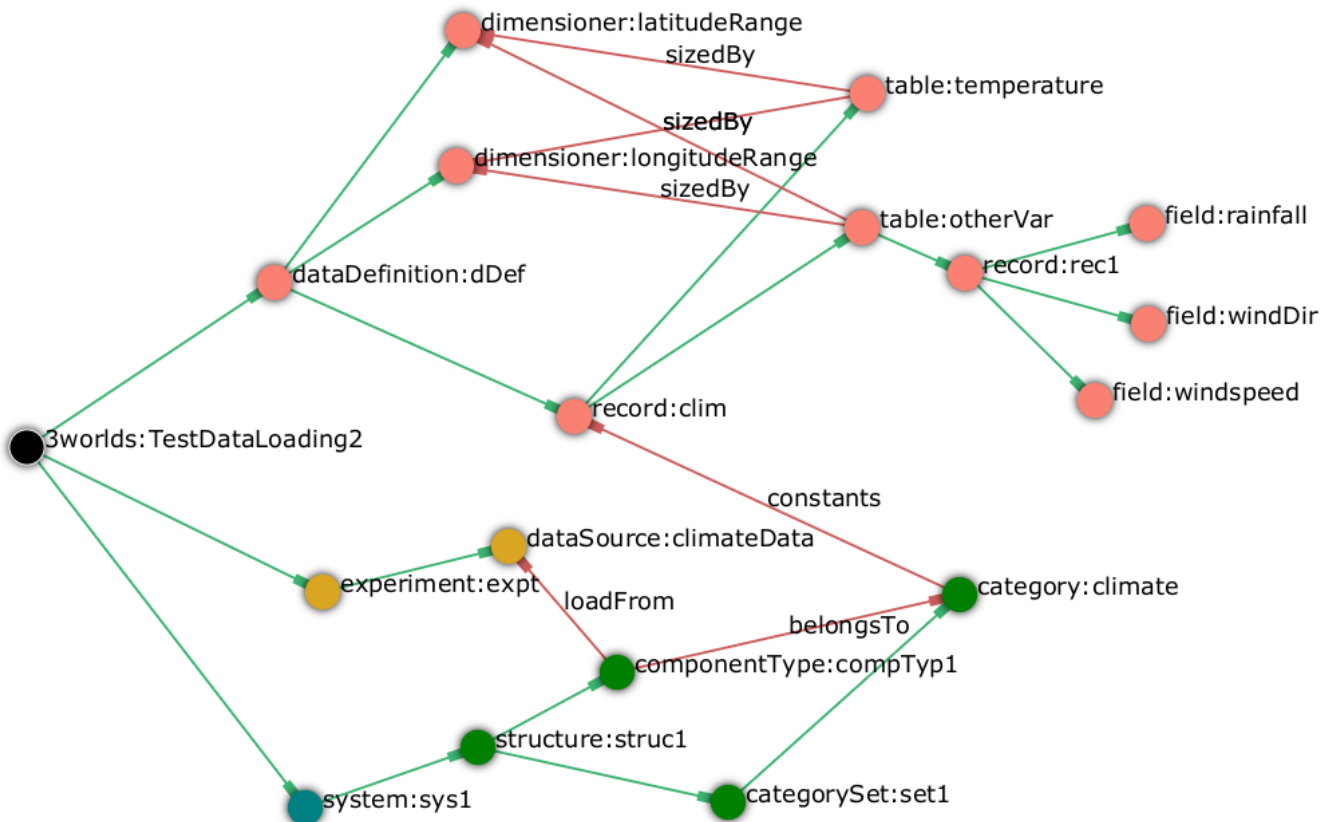
Bottom navigation bar: Back, Forward, New, Feuille1, Feuille 1 sur 1, Par défaut, Français (France), Zoom, 120%.

with the following specifications:

```

climateData#dim      ([2]"lat","long") ...
climateData#file      /climat.ods ...
climateData#subclass  au.edu.anu.twcore.experiment.runtime.io.OdfFileLoader

```



will result in the instantiation of a single system component with id **49**, belonging to category **climate**, with an elaborate data structure as shown on the graph, containing a table **otherVar** of dimensions **[12,8]** with values read from the **climat.ods** file (from debugger):

```
container:sys1[  
    categories:[category:*arena*], ...]  
    local_items:{  
        49=[SystemComponent:49  
            otherVar={[12,8],(8 3.1 C),(0 2.9 C),(2 5.3 C),(0 3.1 C),  
                      (16 3.9 C),(17 3.9 C),(5 2.0 C),(17 2.6 C),  
                      (0 1.2 C),(0 0.4 C)...}  
            temperature={[12,8],21.8,21.4,23.0,24.0,19.4,20.5,22.0,  
                        18.0,21.3,22.5...}  
        ]  
    }  
    ...  
]
```

How?

- property `subClass` in `dataSource:climateData` tells **ModelMaker** that the file to be read is in `.odf` format. Since there is no `sheet` property, only the first (actually, only) sheet is read;
 - property `file` gives the name of the file to read, here `climat.ods`.
 - since `componentType:compTyp1` is child of the `structure` node and has no `initialValues` child node, the number of components to instantiate must come from the data source;
 - since the `dataSource` node does not have a `componentId` property, only one component can be instantiated (there is no way to uniquely identify it in the spreadsheet). As a result, all data in `climat.ods` will be used to instantiate a single instance of `componentType:compTyp1`;
 - property `dim` tells ModelMaker that indices for the first table dimensions are to be read in column *lat*, and for the second dimension in column *long*;

- column headings prefixed with "otherVar:" indicate that these columns are sub-fields of the **otherVar** table (cf. graph);
- finally, the 96 (=12×8) different lines with different indices (CAUTION: the indices start at 0 for 3Worlds table data structures) will all go into the same constants table as specified by the **climate** category of the **componentType:compTyp1**, of which only one instance automatically identified as #49 by **ModelRunner** has been created.

Example 2: using multiple *dataSource* nodes

These .ods spreadsheets:

termites.ods - LibreO... (Sheet 1)

	A	B	C	D	E
1	xx	yy	type	depth	id
2	12	27	pond	-3.5	P1
3	24	87.2	pond	-1.02	P2
4	36	53	pond	-0.578	P4
5					
6					
7					
8					
9					

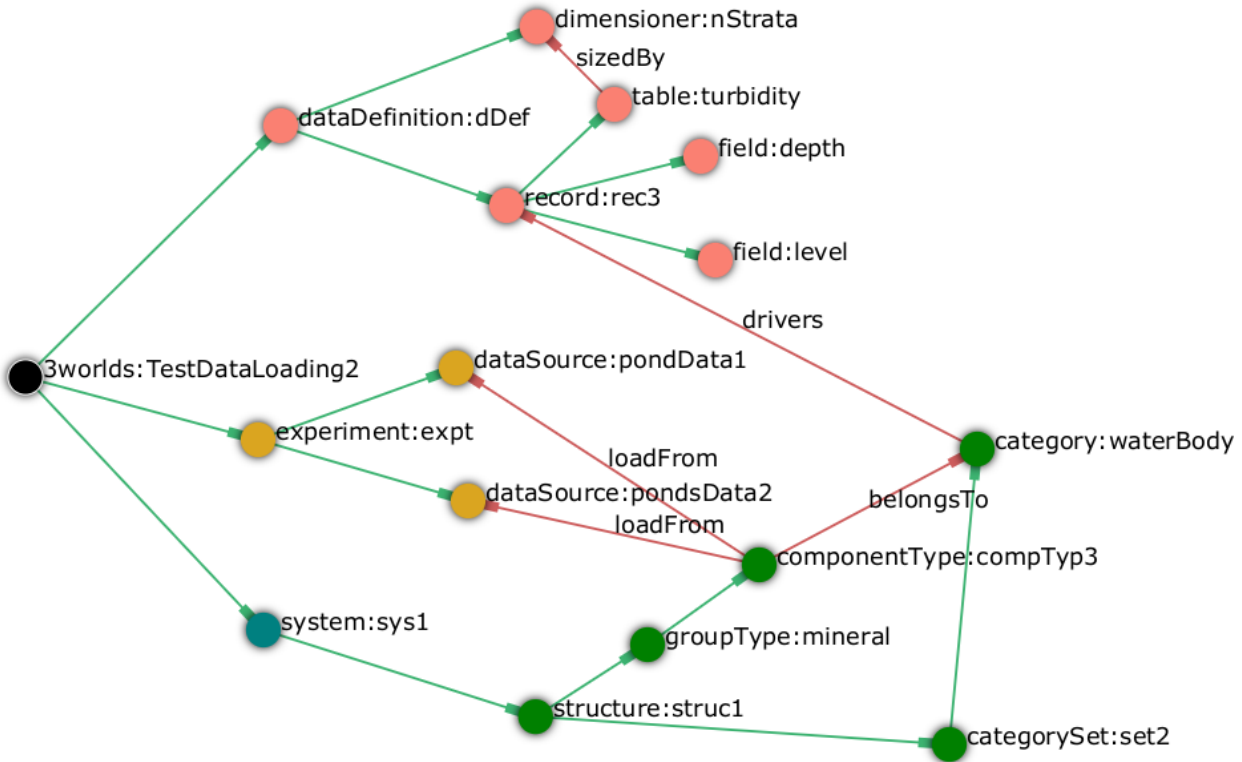
termites.ods : 2 - LibreO... (Sheet 2)

	A	B	C	D	E
1	type	pid	d	turbidity	
2	pond	P1	0	0.21	
3	pond	P1	1	0.74	
4	pond	P4	0	0.91	
5	pond	P4	1	0.61	
6	pond	P4	2	0.18	
7	pond	P3	1	0.37	
8	pond	P3	2	0.14	
9					

with the following specifications:

experiment

pondData1#file	/termites.ods	
pondData1#idComponent	id	
pondData1#idGroup	type	
pondData1#sheet	ponds1	
pondsData2#dim	([1]"d")	
pondsData2#file	/termites.ods	
pondsData2#idComponent	pid	
pondsData2#idGroup	type	
pondsData2#sheet	ponds2	



will result in the instantiation of *four* system components with ids 77 to **80**, belonging to category **waterBody**, with an elaborate data structure as shown on the graph, containing a table **turbidity** of dimensions **3** with values read from the **termites.ods** file, sheets **ponds1** and **ponds2** (from debugger):

```

container:sys1[
  categories:[category:*arena*, ...]
  local_items:{...}
  sub_containers:[
    container:pond[
      local_items:{
        77=[SystemComponent:77 (xx=36.0 yy=53.0 turbidity={[3],0.91,0.61,0.18} depth=-0.578 level=0.0)],
        78=[SystemComponent:78 (xx=12.0 yy=27.0 turbidity={[3],0.21,0.74,0.0} depth=-3.5 level=0.0)],
        79=[SystemComponent:79 (xx=0.0 yy=0.0 turbidity={[3],0.0,0.37,0.14} depth=0.0 level=0.0)],
        80=[SystemComponent:80 (xx=24.0 yy=87.2 turbidity={[3],0.0,0.0,0.0} depth=-1.02 level=0.0)]
      }
    ]
    ...
  ]
]

```

How?

- property `subClass` in `dataSource:pondData1` and `dataSource:pondData2` (not shown) tell **ModelMaker** that the file to be read is in `.odf` format.
- property `file` gives the name of the file to read, here **termites.ods**.
- property `sheet` in `dataSource:pondData1` and `dataSource:pondData2` tell **ModelMaker** that the data are contained in, respectively, sheets **ponds1** and **ponds2** of this file.
- since `componentType:compTyp3` is child of a `groupType`, an `idGroup` property must be provided in the `dataSource` nodes to identify into which group container components instances will be stored; and an `idComponent` property to identify how many components must be instantiated;
- four different components identifiers **P1**, **P2**, **P3**, **P4** (columns `id` and `pid` according to `idComponent` properties of the two `dataSource` nodes) and only one group identifier **pond** (column `type` according to `idGroup` properties of the two `dataSource` nodes) are found in the data sources. They trigger the instantiation of one group container

with four components;

- as `dataSource:pondData2` contains tabular data, a `dim` property tells which column contains the table indices (**d**);
- as `dataSource:pondData1` contains no data for component **P3** (instantiated with identifier **79**), its `xx`, `yy` and `depth` fields are empty (= 0.0);
- as `dataSource:pondData2` contains no data for component **P2** (instantiated with identifier **80**), for `turbidity[2]` in **P1** (instantiated with identifier **78**), and for `turbidity[0]` in **P3** (instantiated with identifier **79**), all these table celles are empty (= 0.0).



If two different data sources contain data for the same field / table cell in the same component, only the last value read will be kept. Since the order of `dataSource` nodes processing is unpredictable, this should be avoided. Take care!

Example 3: using a mixture of `dataSource` and `initialValues` nodes

This .csv text file:

species.csv	
~/3w/project_TestDataLoading2_2022-06-20-1...	
1	species;stage;id;age
2	elephant;e_adult;robert;28
3	elephant;e_adult;jasmine;65
4	elephant;e_adult;pepe;12
5	elephant;e_adult;groumpf;8
6	rat;r_adult;squick;1
7	rat;r_adult;squack;1

with the following specifications:

animalData#file /species.csv

animalData#idComponent id

animalData#idGroup stage

animalData#idLifeCycle species

animalData#separator ;

animalData#subklass au.edu.anu.twcore.experiment.runtime.io.CsvFileLoader

▼ structure

eoffspring#idLifeCycle elephant

eoffspring#nInstances 1

eyoung#idLifeCycle elephant

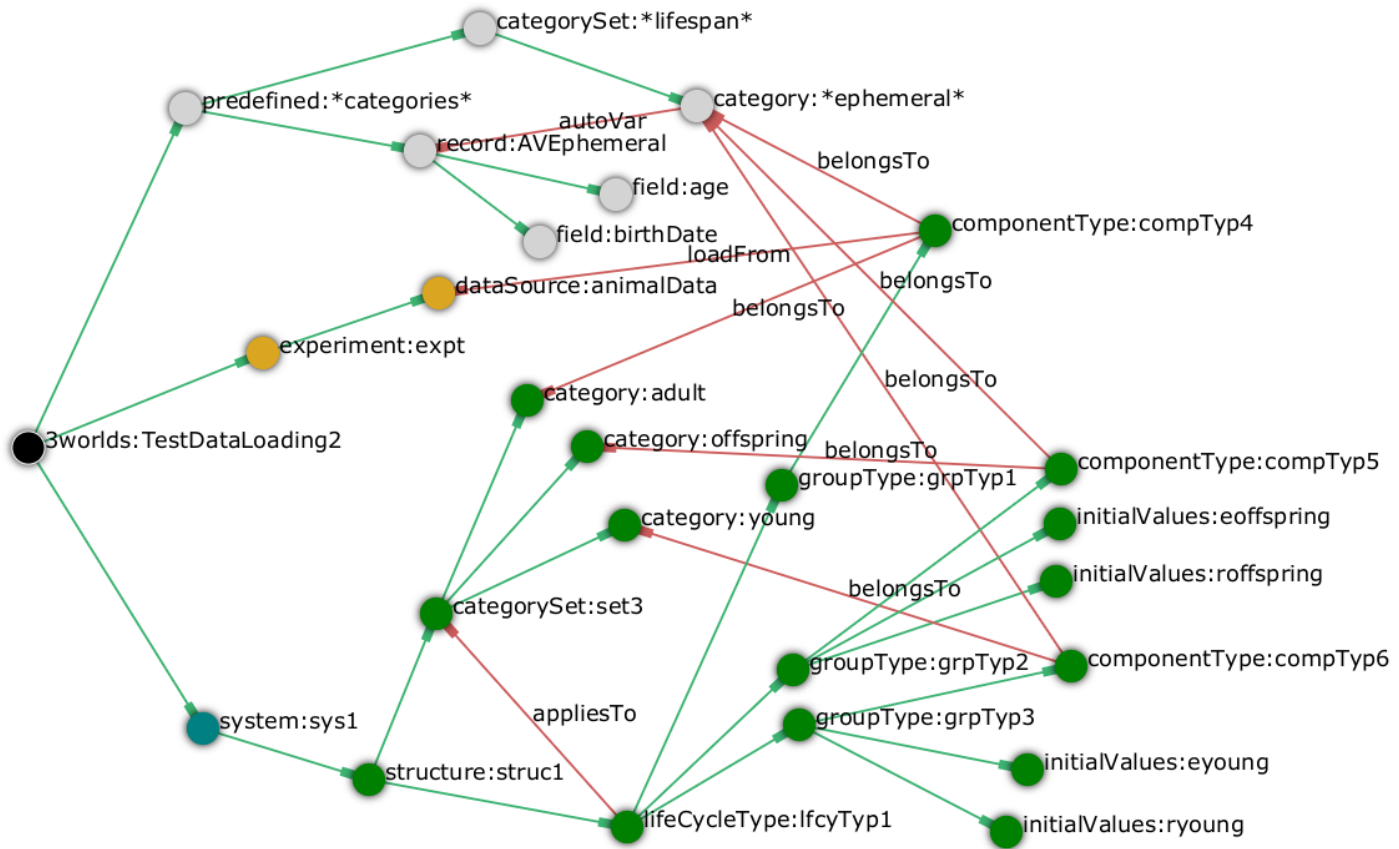
eyoung#nInstances 1

roffspring#idLifeCycle rat

roffspring#nInstances 1

ryoung#idLifeCycle rat

ryoung#nInstances 1



will result in the instantiation of:

- six system components with ids **50** to **55**, belonging to category **adult**, with their `age` automatic variable read from the `dataSource:animalData` node;
- six group containers with ids **e_adult**, **e_offspring**, **e_young**, **r_adult**, **r_offspring** and **r_young**. Components **50** to **53** belong to group **e_adult** while components **54** to **55** belong to group **r_adult** of life cycle **rat**.
- two life cycles with ids **elephant** (containing the three former groups) and **rat** (containing the three latter groups). with the hierarchy of containers is described by all the `initialValues` nodes visible on the graph (from debugger):

```

container:sys1[
  categories:[[category:*arena*], ...]
  local_items:{...}
  sub_containers:[
    elephant=container:elephant[
      categories:[[category:*life cycle*], ...]
      sub_containers:[
        eoffspring=container:eoffspring[
          categories:[[category:*group*], ...]
          local_items:{}
        ],
        eyoung=container:eyoung[
          categories:[[[category:*group*], ...]
          local_items:{}
        ],
        e_adult=container:e_adult[
          categories:[[[category:*group*], ...]
          local_items:{
            50=[SystemComponent:50 (birthDate=0 age=28)],
            51=[SystemComponent:51 (birthDate=0 age=65)],
            52=[SystemComponent:52 (birthDate=0 age=12)],
            53=[SystemComponent:53 (birthDate=0 age=8)]
          }
        ],
        ...
      ]
    ]
  ]
]

rat=container:rat[
  categories:[[[category:*life cycle*], ...]
  sub_containers:[
    roffspring=container:roffspring[
      categories:[[[category:*group*], ...]
      local_items:{}
    ],
    ryoung=container:ryoung[
      categories:[[[category:*group*], ...]
      local_items:{}
    ],
    r_adult=container:r_adult[
      categories:[[[category:*group*], ...]
      local_items:{
        54=[SystemComponent:54 (birthDate=0 age=1)],
        55=[SystemComponent:55 (birthDate=0 age=1)]
      }
    ],
    ...
  ]
]
]

...
]

```

How?

- property `subclass` in `dataSource:animalData` tells **ModelMaker** that the file to be read is in `.csv` format.
- property `file` gives the name of the file to read, here **species.csv**.
- property `separator` in `dataSource:animalData` tells **ModelMaker** that the fields are separated with ";".
- since `componentType` nodes in this graph are all children of a `groupType` node, themselves children of a `lifeCycleType` node, properties `idLifeCycle`, `idGroup` and `idComponent` are required in data files to properly identify the components to instantiate;
- six different components identifiers **robert**, **jasmine**, **pepe**, **groumpf**, **squick** and **squack** (column `id` according to the `idComponent` property of `dataSource:animalData`); two group identifiers **e_adult** and **r_adult** (column `stage`

according to the `idGroup` property of `dataSource:animalData`); and two life cycle identifiers **elephant** and **rat** (column **species** according to the `idLifeCycle` property of `dataSource:animalData`); are found in the data source. They trigger the instantiation of two life cycle containers, each containing a single group container (from the data structure in the file), the former containing the first 4 components and the latter the last 2 components;

- four `initialValues` nodes are also defined in this graph. They are all children of `groupType` nodes, meaning that they apply to groups and not to components. Since the groups in this model do not declare any data, the `initialValues` nodes only tell **ModelRunner** to instantiate one container for each of them (`nInstances` property set to 1). Without these declarations, **ModelRunner** would not instantiate thee containers and would not know where to store instances of components of categories `young` and `offspring` (as no instance of these is required in the data source);
- as `dataSource:animalData` only contains data for the `age` automatic variable, all the instances of adult rats and elephants have their `birthDate` value set to 0; you can also verify that elephants have a much longer life span than rats.



if `dataSource` and `initialValues` nodes are used to instantiate the same components, groups, life cycles, or the arena, all `dataSource` nodes are processed first (in unpredictable order), followed by all `initialValues` nodes (in unpredictable order too).

3.5.5. Import data from external sources

Not yet implemented.

4. Simulation reference: running a simulation experiment with ModelRunner

4.1. General concepts

ModelRunner can be used with or without a GUI. If using without a GUI, all `widget` nodes must be specified as children of a `headless` node. Alternatively, *ModelRunner* will have a GUI if any `widget` nodes are children of `top`, `bottom'`, `'tab` or `container` nodes. You can freely mix these two types of widgets.

At the time of writing, two major aspects of *ModelRunner*'s functionality have not been developed: the ability to change constants during a simulation (i.e to play around with a model while it's running) and to save and restart simulations from particular states (e.g create a save a spin-up state from which to start an experiment).

4.2. Using ModelRunner: software interface and functioning

The layout of widgets in *ModelRunner* is determined by the structure and properties of the `userInterface` sub-tree constructed with *ModelMaker* (ref to UI sub-tree).

The two functioning menu items are **widgets** and **about**.

- **widgets**:: Edit properties of widgets (ref to UI sub-tree). These settings are saved automatically when *ModelRunner* quits.
- **about**:: Information about the experiment design, *ModelRunner* and provision to create model documentation (ODD).

The following widgets have run-time properties, saved automatically by *ModelRunner* so that its appearance remains as selected between each deployment of the model:

- **TimeseriesWidget** :
 - Legend visible: Toggles the legend display on or off.
 - Legend position: The placement of the legend relative to the chart.

- Max legend items: The maximum number of items in the legend. This limits the number of items to display when running experiments with a large number of simulators running in parallel.

4.3. Getting output from a simulation experiment

Outputs from a simulation are detailed in the section on widget properties (ref to UI sub-tree).

5. Sample models and tutorials

5.1. Tutorial 1: Construct and run a simulation for the first time

5.1.1. Introduction

This tutorial explains the procedure for developing and running a simulation with *ModelMaker*. We start by using a simple [logistic growth model](https://en.wikipedia.org/wiki/Logistic_map) (https://en.wikipedia.org/wiki/Logistic_map) as a test case and gradually introduce more capabilities in the tutorials that follow.

In *3Worlds*, the specification of a simulator is developed by constructing a graph of nodes and edges and setting their respective properties. The nodes represent elements of interest in the model while the edges show the relationships between them: the entire graph then constituting the simulator specification. The simulation can be run as soon as the specification is complete and Java code has been written to define the behaviour of the functions you have specified.

As you construct the graph, *ModelMaker* is constantly checking and providing feedback on how the specification complies with the *specification archetype*. *ModelMaker* guides you through this process by managing the available options at each stage in the graph construction and by providing an updated list of tasks requiring attention.

While this process greatly accelerates model development, a more important aim is to provide a firm basis for model comparison. As all models developed with *ModelMaker* must comply with the *specification archetype*, structural differences reduce to a comparison of graphs. That is, while you can construct any model, the design, being constrained by the *specification archetype*, will not be arbitrary. Of course, this is not the only difference between models. The code for each function defined in the specifications will be particular but at least these differences are encapsulated at specific places in the model implementation.

The specification graph is both a tree and a directed graph (Section 3.1). Aside from the root, each node has a parent and may have additional edges between nodes to define particular relationships. We refer to these edges as *cross-links* to distinguish them from edges that represent *parent-child* relationships.

5.1.2. Installation

Installing *ModelMaker* is very simple: create a directory called `3w` in your home directory and copy `tw<os name>.jar` to it where `os name` is one of `'linux', 'mac' or 'win'`.

To start, open a terminal window, navigate to `3w` and enter:

```
java -jar tw<os name>.jar .
```



To compile your own models, the Java Development Kit (JDK) must be installed on your computer (Section 2.2).

5.1.3. Creating a new 3Worlds project

1. Start *ModelMaker* as above. If you haven't read any other *3Worlds* documentation, reading "About ModelMaker" from the main menu (*Help* → *About*) will provide some preliminary context.
2. Create a new project (Main menu: *Projects* → *New* → *Templates* → *Blank*)
3. When prompted enter 'Tut1' (first letter upper case). A small graph of ten nodes appears in the graph window.

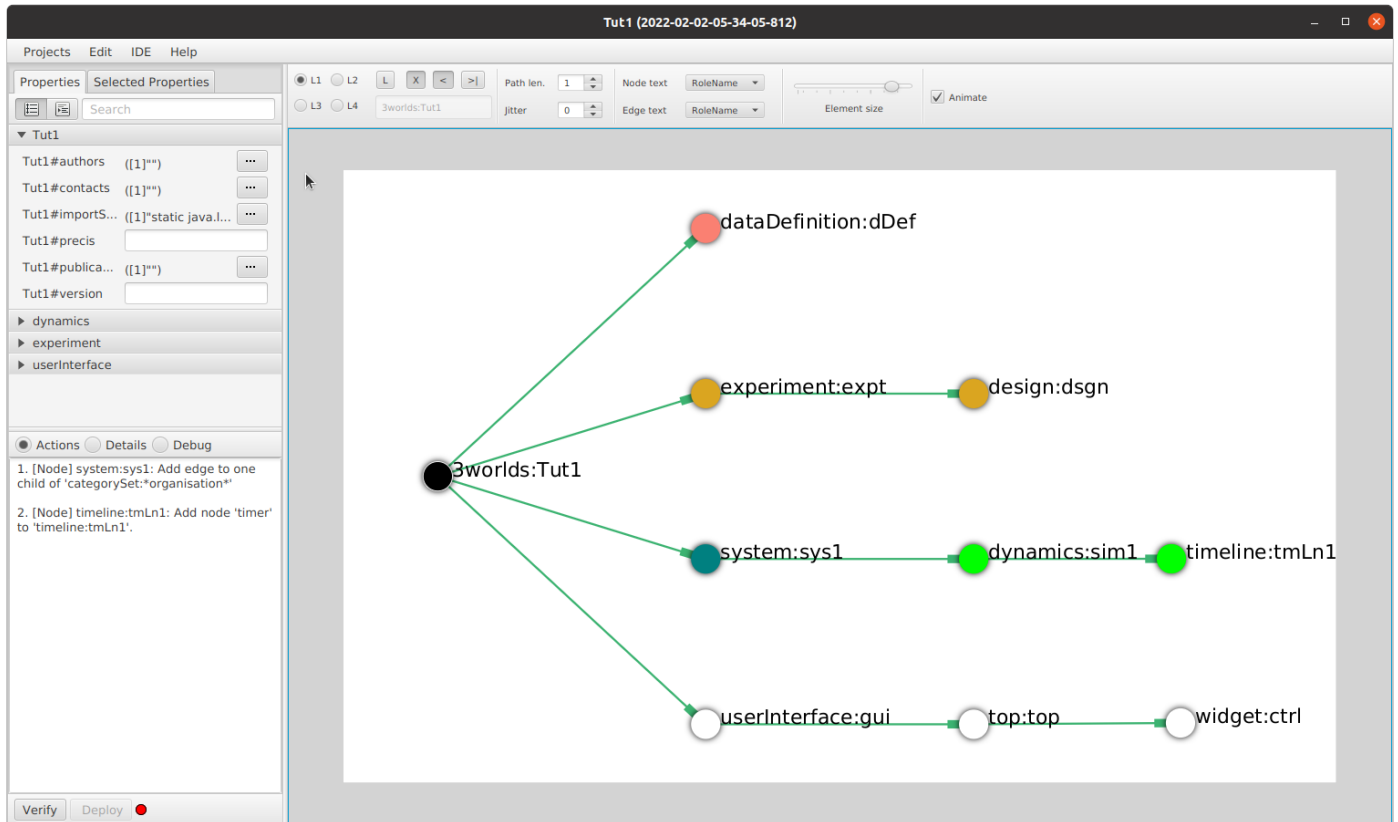


Figure 25. The appearance of *ModelMaker* after creating a new project from the 'Blank' template.

The black node (`3worlds:Tut1`) is the *root* of a graph that forms the simulator's specification. The other nodes are, for the most part, colour-coded roots of sub-trees of the graph wherein various aspects of the specification are defined (Figure 25, Section 3.3).

All nodes are identified by a *label:name* pair. The *label* is the type of node, representing its role, while the *name* is a unique identifier. A few node types, of which the root node is one, must have names beginning with an upper case letter. These nodes are used to generate Java classes when the simulation compiles.

At this stage you can try out a few basic edit operations:

- Move a node around the graph display by dragging it with the left mouse button;
- Zoom in and out in the graph display using the mouse wheel while holding down the `ctrl` key (or equivalent on your operating system). Zooming is centered on the mouse position;
- Pan the drawing surface by dragging with the left-mouse button anywhere outside a node (assuming the drawing area is larger than the display);
- Change the node and text size with the *element size* slider in the tool bar;
- Right-click on a node to display a pop-up menu of edit options for that particular node. It is via these pop-up menus that the graph is constructed; and,
- While holding down the `shift` key, left-click on a node to see just this node and its nearest neighbours. What constitutes 'nearest' is set by the 'path len.' control in the tool bar.

A list of tasks remaining to be completed appears at the bottom-left of the main window (Figure 25). This list grows and shrinks as the developing graph is checked against the specification archetype. A check takes place in the background every time the graph structure or its properties change.

At the top-left of the main window are the two property editors - *Properties* and *Selected Properties* (Figure 25). These show properties for the entire graph or for a single selected node (selected by clicking on the node with the left button) respectively. It's here that values are entered for node and edge properties created in the following steps.

At this stage, there are five properties under the section `Tut1`. These are properties of the root node (`3worlds:Tut1`) intended to provide a limited amount of documentation for this project. If you click on the root node and then select the *Selected Properties* editor tab, an additional non-editable property is shown called 'built-by'. This property is filled out automatically with your computer login name and the creation date of the project. Viewing non-editable properties can be useful at times. These are always shown in the *Selected Properties* editor but for brevity, not in the *Properties* editor. Two icons at the top left of the *Properties* editor allow you to display properties by category (by sub-tree) or by name. When this last option is selected, all editable properties are displayed in one list.

Directory structure

All projects are located within the `3Worlds` root directory called `3w`.

(Figure 26).

The `3w` directory contains a Jar file called `tw<os name>.jar` (Figure 26). This file contains all dependent libraries for `3Worlds`. It's a good idea to keep a backup of this file.

Within `3w` there is now a sub-directory called something like `project_Tut1_2020-06-07-01-47-44-708`. Every project directory begins with the key word `project`, followed by the name of the project you entered (`Tut1` in this case) and the creation date and time. The purpose of naming project directories in this way is to make it almost impossible to inadvertently overwrite a project.

Inside this directory are a number of text files:

- i. `Tut1.utg` is the model configuration graph we're constructing;
- ii. `.layout.utg` contains information used to display the graph;
- iii. `MM.xml` contains the session's preference settings.
- iv. `_StateA1` and sequences of similarly named files are temporary files to support the program's undo/redo of editing operations. These files are created during a session and deleted when the session ends.

All these files are text files and you can open and inspect them with a simple text editor. However, as they are written by *ModelMaker* you should avoid editing them directly as this will likely cause problems for your project.

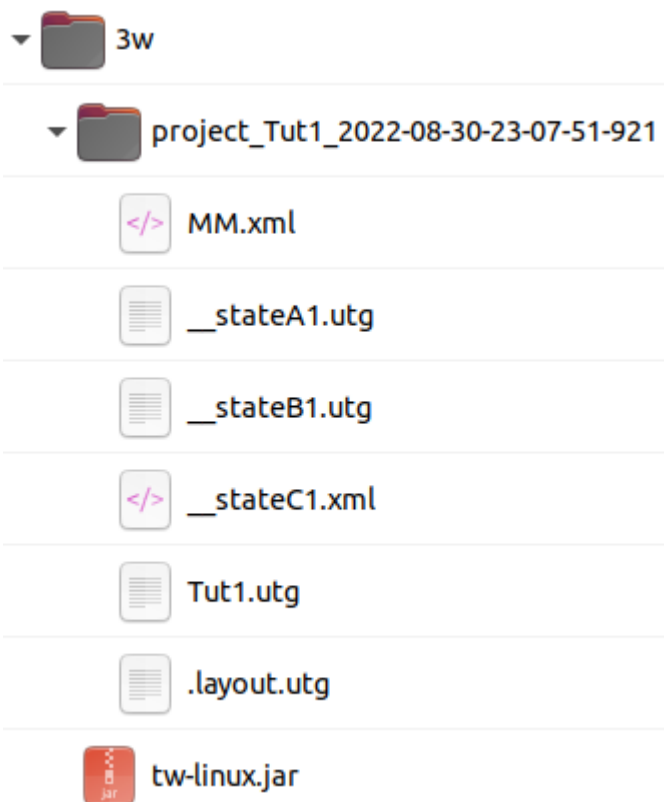


Figure 26. The initial directory structure after creating a new project from the 'Blank' template. Later, after all tasks have been completed and the model is deployed, another sub-directory is added for run-time files.



Editing any project graph files (*.utg) with a text editor is likely to corrupt the project.

On the other hand, you can delete project directories at any time you wish. If you accidentally delete the project of a currently open session, it will be recreated automatically by *ModelMaker* (apart from the *undo-redo* history). However, if you delete the entire `3w` directory you will have to restore `tw<os name>.jar` from backup.

5.1.4. Creating the specifications

Having created `Tut1`, the task list shows two actions are required (Figure 25). These tasks can be dealt with in any order but in general, a logical approach is to build the specification as follows:

- i. **data definition**: the data structures required (cf. Section 3.3.5 for full reference);
- ii. **structure**: define the organisation of components - their roles and relationships (cf. Section 3.3.3 for full reference).
- iii. **dynamics**: define how the modelled system evolves over time and the parts of the **structure** that dynamic processes apply to (cf. Section 3.3.4 for full reference).

For the most part, these tutorials will proceed in this order and leave defining the simulation's **user interface** (Section 3.3.7) and **experiment** design (Section 3.3.6) until last. In fact, the Blank template has provided a minimum specification for the **experiment** sub-tree (a single run of the model) and the **user interface** (a simulation controller).

Data definition

The logistic equation we will implement is: $x(t+1) = rx(t)(1-x(t))$. Though simple, it has interesting chaotic behaviour for values of r between 3.7 and 4.0. All we need do for the data definition section is to define the parameter r and the state variable x .

From here on and throughout these tutorials, parameters are called *constants* (data that does not change over the course of a simulation) and state variables are called *drivers* (data that drives the simulation from one time to the next).

Note that at this stage, the task list says nothing about adding anything to the **data definition** sub-tree. This is because it is possible to define a model without data. Nevertheless, once data is defined, various actions will be required and displayed in the task list to correctly define its use.

1. Right-click on the data definition node (`dataDefinition:dDef`) (pale red) and select `New node → record` from the popup menu. You're then prompted for a name. The default name is `rec1`. Change this to '`cnsts`' (constants) and click `ok`. The mouse pointer immediately becomes a cross-hair: *ModelMaker* is asking where to place this node. Move to some place within the graph display and left-click the mouse.

You can name nodes and edges anything you like but accepting the recommended names and edges will make these tutorials easier to follow. Note that *ModelMaker* will prevent naming nodes or edges with duplicate names.

All nodes in the configuration graph are children of some parent (apart from the root node). You can only create nodes by right-clicking on a parent and selecting a child to create from the available options provided by the pop-up menu. The items in this menu vary according to the possibilities allowed by the *specification archetype*. This is one way *ModelMaker* ensures the developing configuration conforms with the *archetype*, greatly simplifying an otherwise complex workflow.

2. Create a `field` node as child of `record:cnsts`, name it '`r`' and when prompted, set its type as `Double`.

All `fields` (and later `tables`) must be children of some `record`.

3. Create another `record` as child of `dataDefinition:dDef` and name it '`drv`s' (drivers).
4. Create a `field` node as child of `record:drv`s, name it '`x`' and again set the type to `Double` (Figure 27).

Note that the names '`drv`s' and '`cnsts`' don't imply any meaning to the simulation specification - they're just names. Their *roles* as drivers and constants will be defined later.

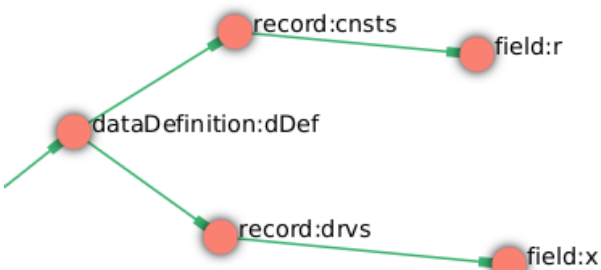


Figure 27. Defining constant r and driver x within the data definition sub-tree.

This is all the data required for this tutorial. The task list has now grown to four because the roles of this data remains undefined.

You can tidy up the graph display by clicking the L button (re-apply layout) in the tool bar.

Structure

At this time, the task list displays four actions. Three of these actions relate to the structure of the model. The `structure` sub-tree describes how the modelled system is organised into separate components playing particular roles. In an elaborate model, there can be many components but in the present case,

we need only one, and for convenience, the `system` node can act as this single component without the necessity of actually creating a `component` within a `structure` sub-tree.

Here, the component's *role* will be defined as:

- lifetime: *permanent*;
- organisation: *atomic*;
- systemElements: *arena*;
- Using *r* as a constant and *x* as a driver.

The component has a *permanent* lifespan because it persists throughout the simulation; it's *atomic* simply because it is a single indivisible component and not an assemblage of sub-components; and, it belongs to something called the *arena*. The *arena* is a unique top level component - it's more or less a global component accessible to all other components. No matter how many components a model has, exactly one of them must belong to the *arena* category,

While this is complicated for such a simple model, later tutorials will show how this arrangement can be a powerful approach to structuring any complex hierarchical dynamic system composed of interacting physical and biological components.

To create this role, we use nodes of the type `categorySet` and `category`. A `categorySet` is a set of mutually exclusive categories. By that we mean a `component` can only be associated with one category of a given `categorySet`. So for example *permanent* and *ephemeral* are two categories within a set called *lifespan* and clearly, a component can only be one or the other. Categories and CategorySets are recursive: a CategorySet contains Categories and Categories can contain CategorySets without limit.

Apart from the `system` node doubling as a `component`, an additional convenience is provided: a sub-tree of predefined category sets and categories. We use these nodes to define the role described above. To see this sub-tree:

1. Right-click on the root node and select `Collapse → All`.
2. Right-click again on the root node and select `Expand → predefined:*categories*`.
3. Re-apply the layout ('L' [Alt+L])

The `predefined:*categories*` sub-tree is created with every new project (collapsed by default) and is *immutable* apart from allowing edges to be added between it and other sub-tree nodes.

There are two `record` nodes within this sub-tree for default handling of average population and ephemeral data. Since the single component used here will be neither of these we can ignore this section:

4. Right-click on `predefined:*categories*` and collapse both the `AVPopulation` and `AVEphemeral` sub-trees.
5. Right-click on the root node, expand the `system` and re-apply the layout.

We are now in a position to define the *role* of the system node (a.k.a. component in this case). The system node always belongs to the arena and permanent categories. This has already been done by the 'Blank' template. It just remains to make the system atomic.

6. Right-click on `system:sys1` and select New edge → `belongsTo` → `category:*atomic*`.

The system's role of belonging to the *atomic*, *permanent* and *arena* categories is indicated by three cross-link edges (Figure 28).

All cross-links are red - thin at the *start node* and thick at the *end node*. Unlike parent-child links, they have names. Generally, the names of cross links are not much use. They can be hidden by selecting the drop-down list E text in the tool bar, and selecting Role. The relationship can be read as, for example: `system:sys1` `belongsTo` `category:*arena*`.

We have yet to relate `system:sys1` to the driver 'x' and the constant 'r'. These are global data and therefore are defined as part of the arena.

7. Right-click on `category:*arena*` and select New edge → `constants` → `record:csts`.

8. Right-click again on `category:*arena*` and select New edge → `drivers` → `record:drvs` (Figure 29).

There is now one message remaining which refers to the dynamics of the model.

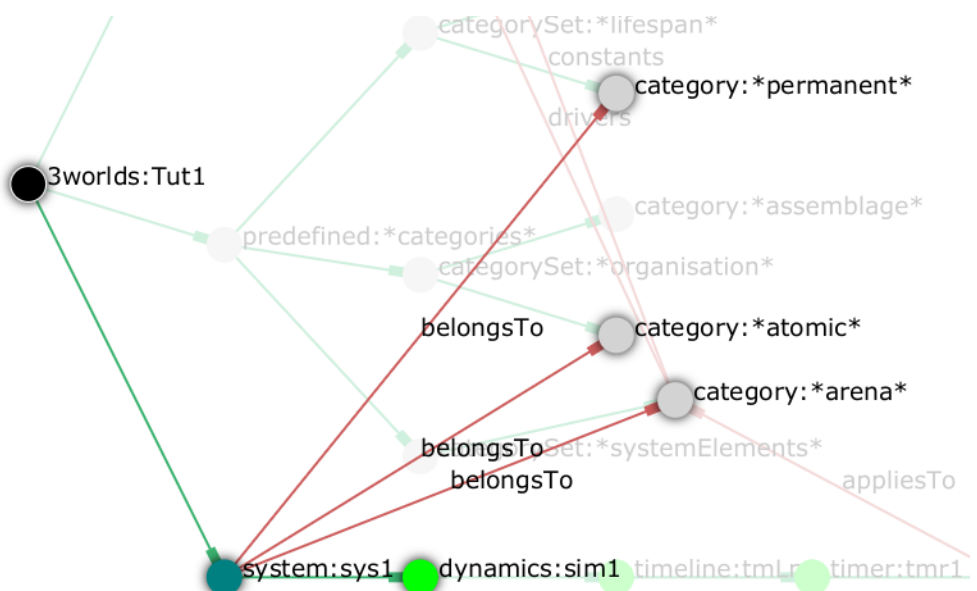


Figure 28. The specification of the system component in belonging to the three categories; permanent, atomic and arena.

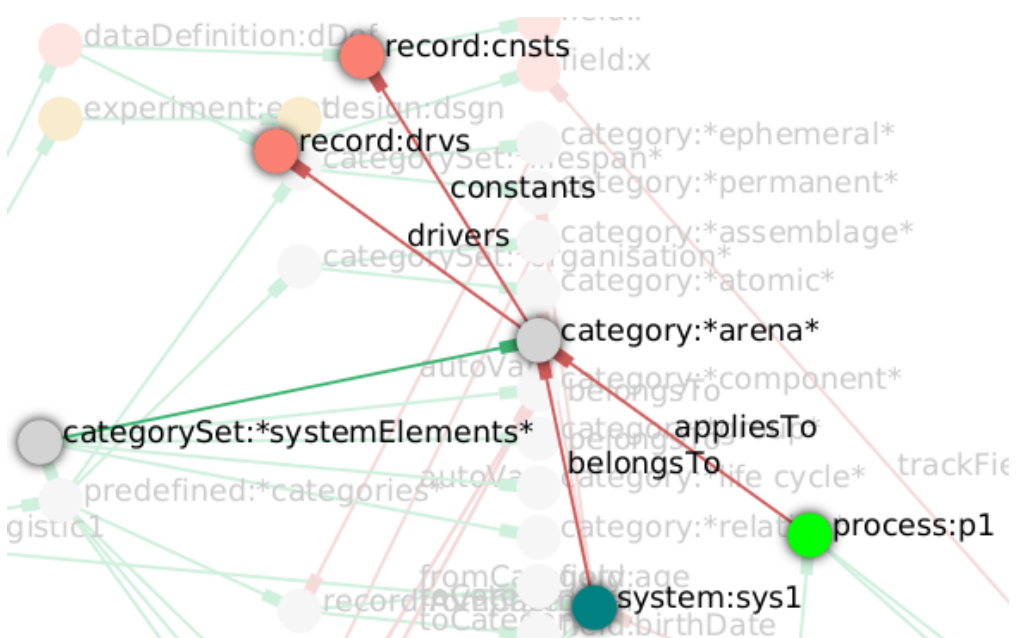


Figure 29. The specification of the constants and drivers for the arena, a category to which the only component (aka `system:sys1`) in this tutorial belongs.

Dynamics

The `dynamics` sub-tree specifies how the modelled system will evolve over time. It determines the temporal order of function calls, their type, the categories they apply to (cf above), the conditions under which the simulation will stop (if any) and what and when data will be tracked for output.

In the present case, the main task is to call the logistic equation a set number of times and present the result from each iteration to the equation at the next time step.

The `dynamics` sub-tree (lime green) is a child of the `system` node - the root of the modelled system that defines both its dynamics and structure. These nodes are already present in the "Blank" template we started with. The `dynamics` node is the specification of a type of simulator. There can be many simulators (instances) of this specification running in parallel depending on the experimental design.

A dynamic system must have a single definition of the time scale. There are ten different types of time scale available: all of them define exact sub-divisions of time except for the `GREGORIAN` scale type which implements the standard Gregorian calendar. The 'Blank' template has already defined a `timeline` with an `ARBITRARY` time scale which is sufficient for this tutorial.

In the task list is an action asking that a `timer` be added to `timeline:tmLn1`.

1. Create a `timer` as a child of `timeline`. Here an extra prompt appears asking for the class of the timer: `{ClockTimer, EventTimer, ScenarioTimer}`. Select `ClockTimer`. This timer class increments time by a constant step during simulation, unless the timeline uses a Gregorian scale in which case irregularities such as leap years are managed.

Two new tasks have been added to the task list indicating that `dt` and `nTimeunits` must be greater than 1.0.

2. In the property editor, locate these two properties (properties of `timer:tmr1`) and set them to 1.0.

Clock timers use their own time units to count time. The time unit is specified by:

- the `timeUnit` property, which defines in which measurement unit time is to be expressed. There are 22 time unit types available ranging from microseconds to millennia, constrained by the choice of the time scale. The current default value of `UNSPECIFIED` is fine for this tutorial, as here, time is just a sequence of steps;
- `nTimeUnits`, i.e. the number of measurement units per timer unit - for example, a timer could use '2 days' as its basic time unit. It means that an elementary time tick in this timer represents 2 days, so (time = 11) means 22 days, etc. It is the finest *grain* at which time can be measured with this timer. In this tutorial, 1 is fine as we simply use a sequence of steps.

Finally, `dt` is the number of time *grains* per time step. Here, 1 will do. This tutorial runs with time steps of 1 unspecified unit, i.e just simple ticks.

Note that a model can have any number of `timers` using any of the available time steps and time units as long as the time units selected are compatible with the parent `timeline`. The task messages will indicate if this is not the case. Because the specification archetype allows for more than one system, it follows there can be many dynamics sub-trees, each with their own time system.

There is currently a task asking that a `process` be added to `timer:tmr1`.

3. Create a `process` node as child of `timer:tmr1`.

This process will operate on `r` and `x` that we defined in the Data definition section. These data are associated with the `arena` as constants (`r`) and drivers (`x`). Therefore `process:p1` will applyTo the arena.

4. Right-click on `process:p1` and select 'New edge > appliesTo → category:`*arena*`'.

A `process` defines a set of computations acting on model components driven at the rate of the parent `timer`. A component is some unit of simulation. It can be any physical or biological entity represented in the model that has dynamic behaviour (plants, animals, nutrient pools, lakes, the atmosphere or the rhizosphere etc).

Processes can be composed of any number of functions of various types (much more on this later). We need just one function to implement the logistic equation - a `ChangeState` function that takes the current state of a component and

calculates the next state.

5. Create a `function` as a child of `process:p1`, name it `Chaos` and select `ChangeState` as its type.

The function type can't be changed after creating node, so if you make a mistake, delete and recreate it (`Delete` from the pop-up menu or `Undo` from the main menu).

ModelMaker can link to an Integrated Development Environment (IDE) such as *Eclipse*, to write code for these functions. However, in this tutorial the function is only one line of code and we can just associate a code snippet with the function without the need to link to an IDE. The snippet will be inserted in the function when the simulation is compiled.

6. Edit the `functionSnippet` property of `function:Chaos`, and enter the following text: `focalDrv.x = r*x*(1-x);`

This line sets the next value of `x` (`focalDrv.x`) in terms of the current value of `x`.

Save your work (`Ctrl+s`) and the task list will be empty indicating the simulation is ready to deploy.

Deployment: launching *ModelRunner*

Before deploying the simulation, it's useful to display the overall specifications by hiding the predefined sub-tree and showing the experiment and user interface sub-trees.

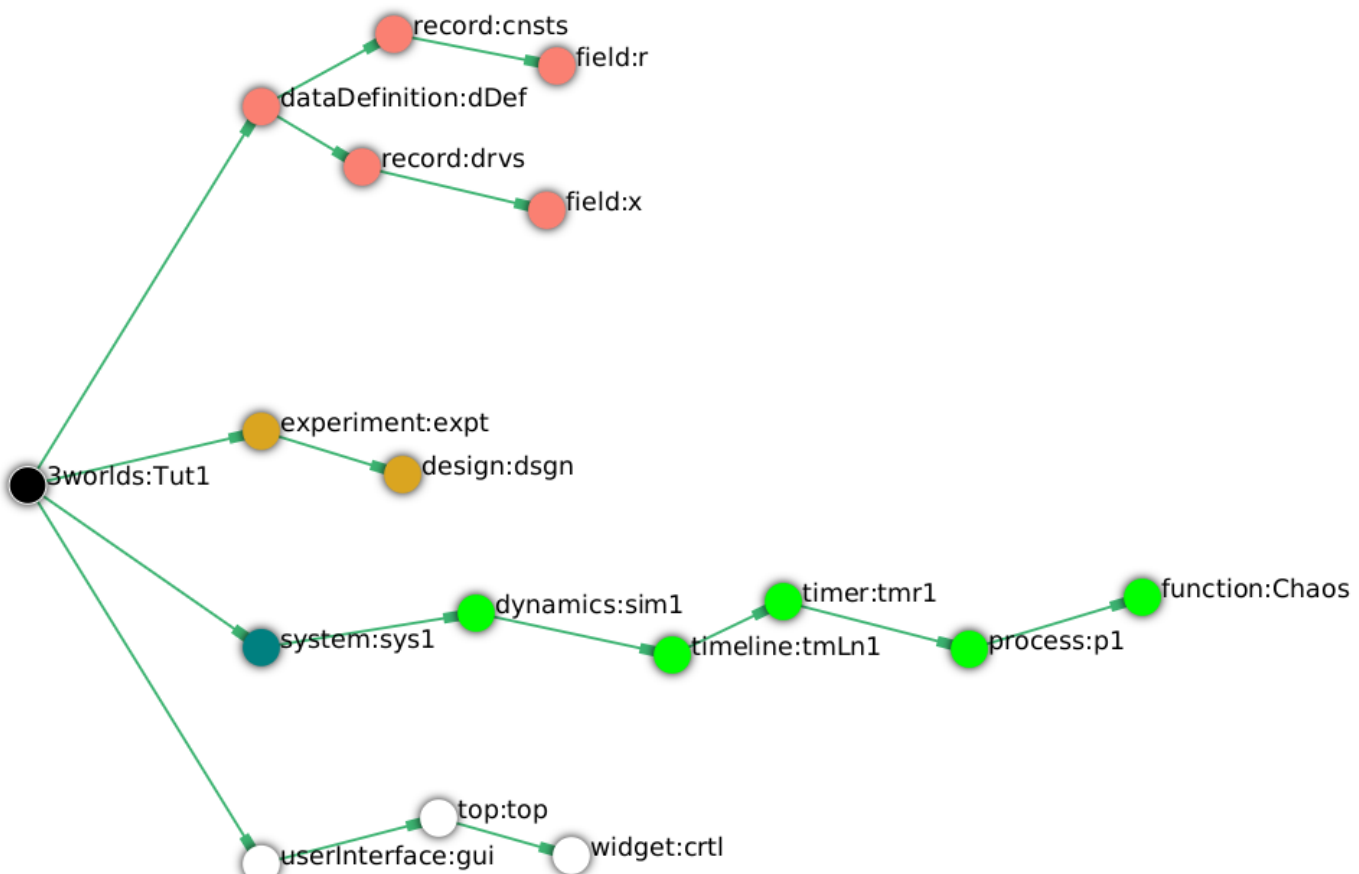


Figure 30. A view of the specification graph showing principal nodes of interest.

1. Right-click the root node and select 'Collapse → predefined:*categories*'.
 2. Again, right-click the root node and select 'Expand → experiment:expt' and 'UserInterface:gui'.
 3. Re-apply the layout (L) (Figure 30).
 4. Click the `Deploy` button. *ModelMaker* now launches *ModelRunner* to start the run-time application: *ModelRunner* (cf. Section 4 for a full reference).

At the top of *ModelRunner* are some control buttons to start, step and stop the simulation. This is the `ControlWidget1` shown in the figure above that was included by default in the 'Blank' template. The `run` button becomes a `pause` button while running and the `stop` button resets the simulator to its starting state.

However, as expected, there's nothing to see so the next step is to add a time series widget. This is an optional requirement so the task list didn't complain about this.

You can move easily between design and execution of the specifications simply by deploying *ModelRunner*, checking the simulation and quitting to return to *ModelMaker*.

To add a time series for `x`:

5. Quit *ModelRunner* and return to *ModelMaker*.
6. Create a `tab` node from the `userInterface:gui` node.

A `tab` is a container that can contain either widgets or other containers. The task list indicates this now asking that one of these be added.

7. Create a `widget` node from `tab:tab1`, name it 'srsx' and select `TimeSeriesWidget1` from the drop-down list.

A new task has been added to the list requiring an edge from this widget to a `dataTracker`.

For this widget to receive values of `x`, something must post values of `x` to the widget at the same rate as the `Chaos` function is executed. This is the job of a `dataTracker` and it properly belongs in the `dynamics` sub-tree.

8. Create a `dataTracker` as a child of `process:p1` and choose `DataTracker0D` as its class. This class of data tracker is suitable for scalar data such as `x`.
9. Create an edge from the `dataTracker` to `x` by selecting `New edge → trackField -> field:x`.
10. Create an edge from `widget:srsx` to the new data tracker.
11. Create an edge from `dataTracker:trk1` to a `component` i.e. in this case `system:sys1`. A data tracker must not only track some data but also the `component` that uses this data.
12. Save the graph (`Ctrl+s`) and click the 'Deploy' button again.

The display is still uninteresting because we haven't set an initial value for `x` or parameterised `r`. This can be done in a number of ways but for this tutorial we will add an initialisation function and a code snippet.

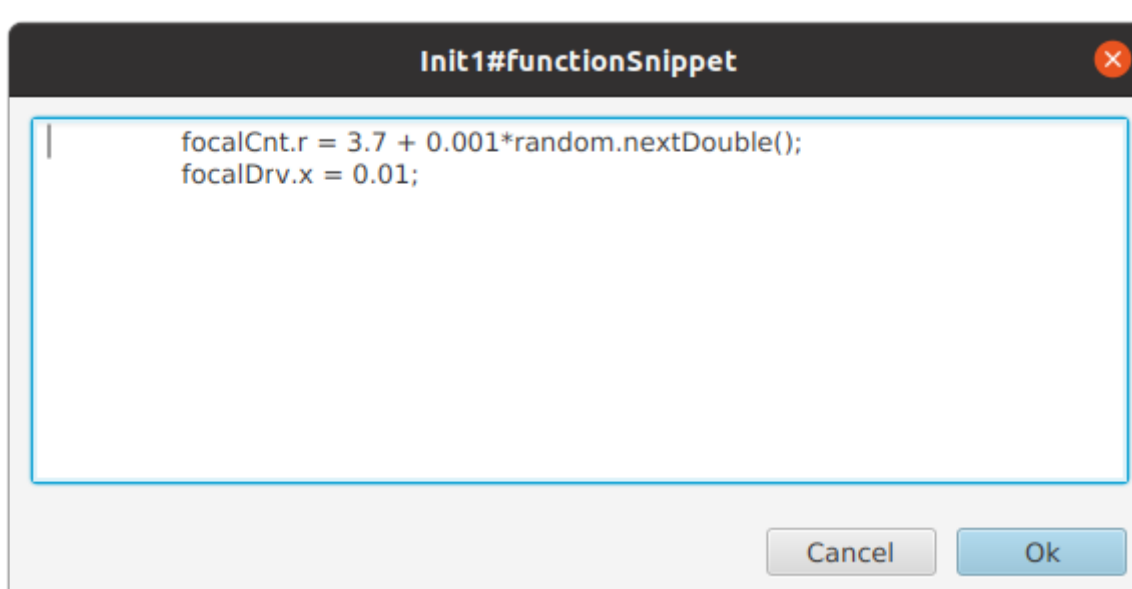


Figure 31. Entering Java code snippet for the `initFunction` (for the

system:sys1 *component*) in the property editor.

i. focalDrv.x =
0.001;

ii. focalCnt.r = 3.7;

If you make a typo, the task list will show the details of the compile error.

Stopping conditions

If you examine the graph and all its properties, you may notice that there is no indication as to how long the simulation should run. This means that when we run it we should expect it to continue indefinitely. You may or may not want this. If your model contains an unconstrained exponential function, it may eventually crash unless your code takes measures to handle this. You can add a variety of simple or complicated stopping conditions to the `dynamics` node. These will be discussed in later tutorials.

When we first ran this model it had no output. Now that we have a time series chart, displaying data of unlimited length will make the *ModelRunner* fairly unresponsive because the simulator has little to do but the UI must update the graph continuously. Note: the `TimeSeriesWidget1` displays data in a rolling buffer. The default buffer size is 1,000 data points.

If you press the run button and then the stop or pause button of the controller, it may take a while for the simulator to respond. So for now, it's best to test the simulation with the `Step` button.

16. Deploy *ModelRunner* (saving first if prompted)

17. Click the `step` button a few times. A time series of zeros is shown.

18. Click the `run` button twice in rapid succession. The time (x axis) now reads approximately 30,000 or so depending on the speed of your computer.

To complete this tutorial, add a simple stopping condition:

19. Close *ModelRunner* and return to *ModelMaker*.

20. Create a `stoppingCondition` as a child of `dynamics:sim1`. When prompted, select `SimpleStoppingCondition` from the drop-down list.

21. Select this new node and in the properties editor, set the value of `stCd1#endTime` to 100 (Figure 32).

22. Save, re-deploy and run the simulation. You'll now see a time series of the chaos function of 100 time steps (Figure 33).

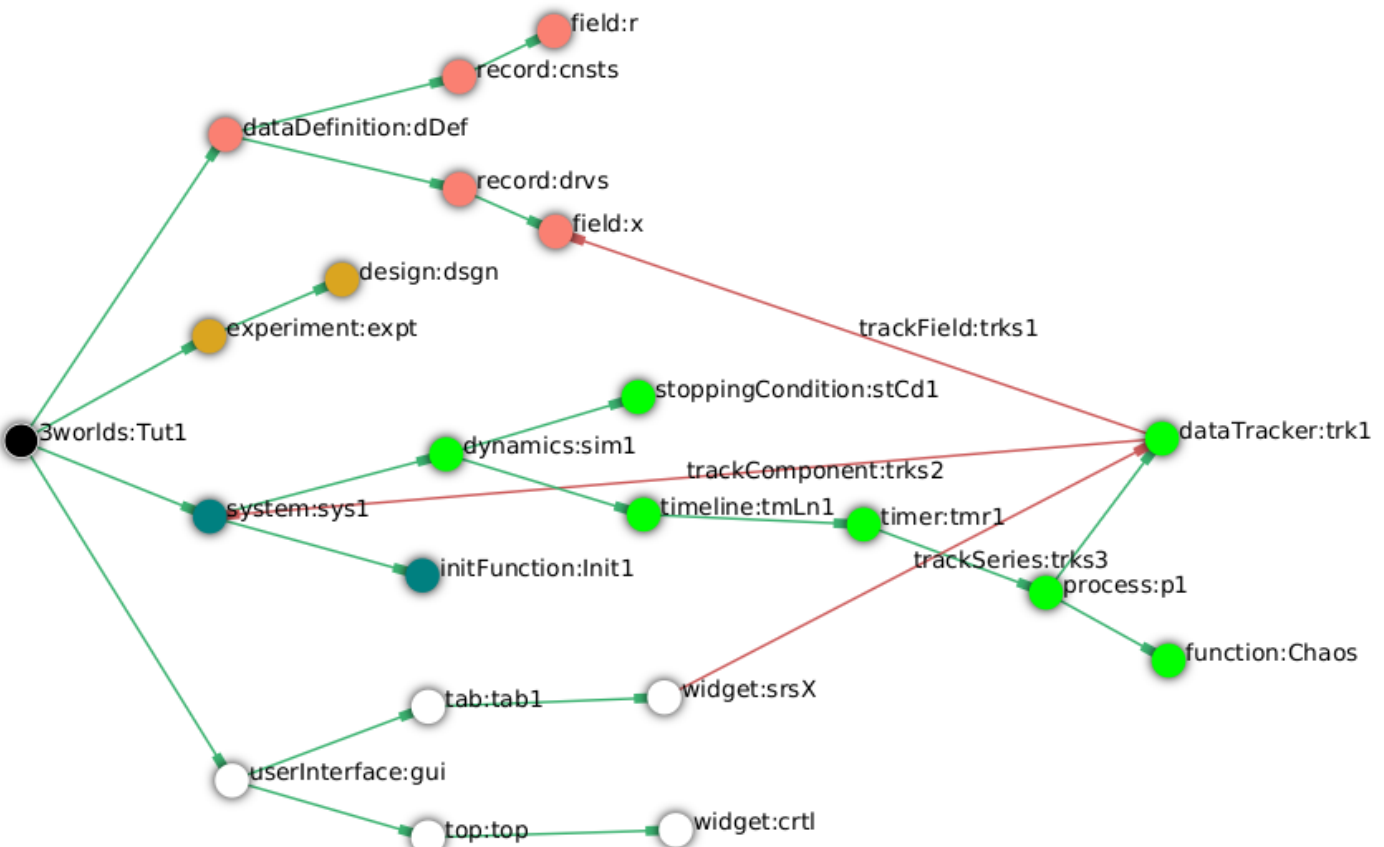


Figure 32. The specification graph for Tutorial 1.

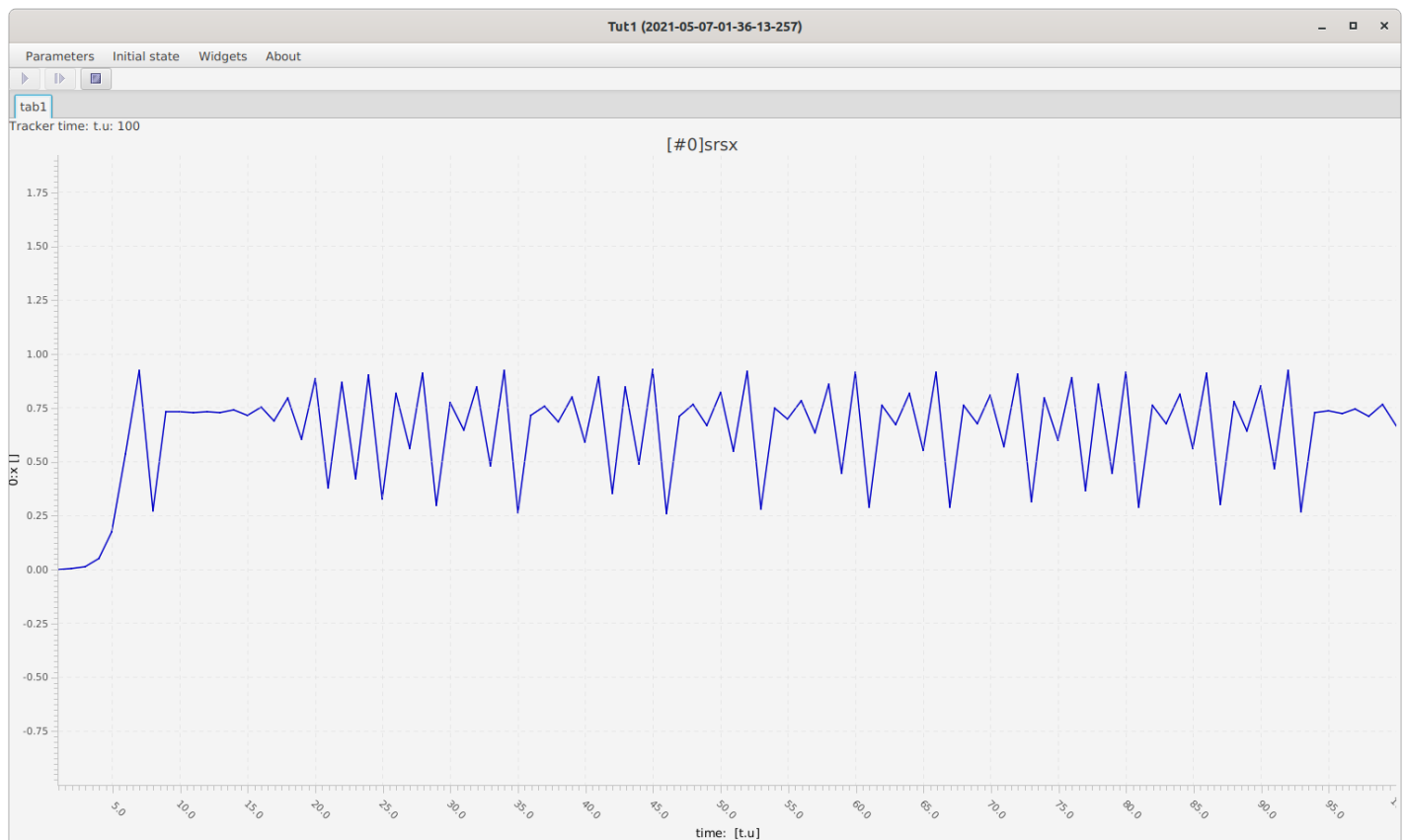


Figure 33. The appearance of ModelRunner while running Tutorial 1.

You can now generate documentation for this model from *ModelRunner*.

23. From *ModelRunner* select 'About → Create documentation' from the main menu.

This creates an ODD template (Overview, Design concepts and Details) (*Tut1.odt* , Figure 34), an established standard for documenting simulation models. This file can be generated any time from *ModelRunner*. When you're satisfied with

the specification, edit this file as a basis for the complete documenting of the model. The file can be found in the Tut1 project directory Figure 35:

~/3w/project_Tut1_2021-05-07-01-36-13-257/local/runTime/documentation1/Tut1.odt .

Most word processors can read this standard format, native to LibreOffice, including Microsoft Word.



Every time documentation is created, it is placed in a incrementally numbered directory to prevent overwrites.

Overview, Design concepts and Details

Tut1 (2021-05-22-21-03-29-813) (Version: demo)

Jacques Gignoux

Ian D. Davies

Date: 22-May-2021

Purpose

[Explanation: Every model has to start from a clear question, problem, or hypothesis. Therefore, ODD starts with a concise summary of the overall objective(s) for which the model was developed. Do not describe anything about how the model works here, only what it is to be used for. We encourage authors to use this paragraph independently of any presentation of the purpose in the introduction of their article, since the ODD protocol should be complete and understandable by itself and not only in connection with the whole publication (as it is also the case for figures, tables and their legends). If one of the purposes of a model is to expand from basic principles to richer representation of real-world scenarios, this should be stated explicitly.]

Entities, state variables, and scales

Table 1. Component (Entity) description

Component type	Categories	Component	Drivers	Time	Space
sys1	permanent atomic arena	none	drv1	tmr1 [ClockTimer]	non-spatial

Table 2. Driver (state variable) details

Driver	name	Description	Dimensions	Type	Units	Range
drv1	x		scalar	Double] -∞, +∞ [

Figure 34. A section of the automatically generated documentation for Tutorial 1. The text in italics is a quote from Grimm et al. (2010) supplementary material ODD template file with suggestions as to how to write this section of the documentation. This quote is replaced by the `precis` property of the root node for convenience.

Once the specification has been compiled without errors and deployed and the documentation generated, a number of files will have been created Figure 35:

- i. `local/java/code/` : This directory contains the generated Java code. The entire model is contained within the `Tut1.java` class.
- ii. `Tut1.jar` : A stand-alone jar of the model that can be run independently of *ModelMaker* (Section 4).
- iii. `runTime/MR.xml` : A preferences file for *ModelRunner*. These settings are saved whenever *ModelRunner* quits so it

will have the same appearance between successive uses.

- iv. runTime/documentation1/Tut1.odt : This is the documentation file generated in the previous steps.

5.1.5. Graph layouts

An aspect of *ModelMaker* we have only touched on so far, is the graph layout system.

While using a graph to construct model specifications has many advantages, you can quickly become lost in a confusion of nodes and edges. The advantage in presenting the specification as a graph is that the huge number of options possible can be constrained by context. For example, to have a dynamic process, it makes sense that it's associated with a particular timer, that other processes working at the same rate are associated with the same timer and that all timers are coordinated by the one timeline. The user interface for problems such as this would be very error-prone if presented say, as a series of dialog boxes.

ModelMaker has a number of features to help arrange the graph display. Please refer to Section 3.2.2.

As an exercise if you wish, try displaying just the nodes with *cross-links* using the SpringGraph layout (L4) (Figure 36). This is a common way to look at just the *cross-link* relationships between nodes. Generally, adding a screen capture of this and a second screen capture of just the relevant *parent-child* relationships make useful additions to the ODD appendix.

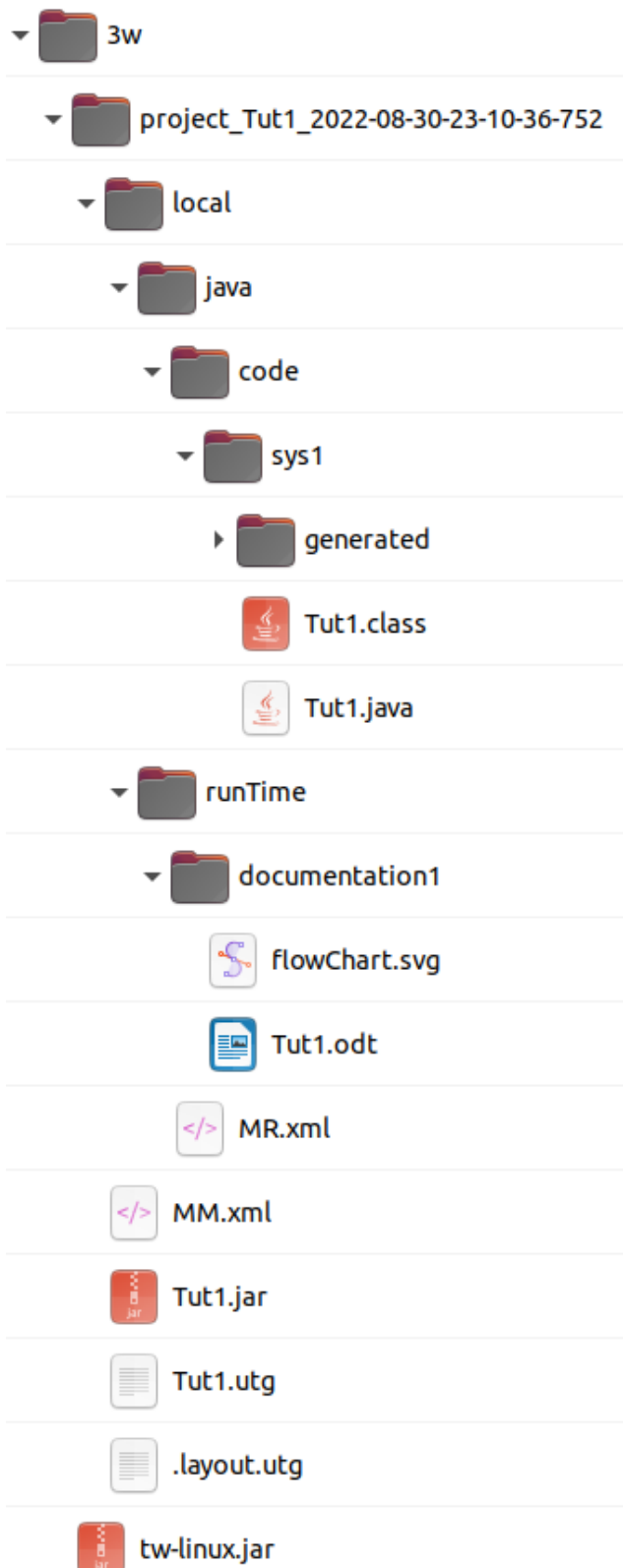


Figure 35. The directory structure after deploying ModelRunner and creating the ODD documentation.

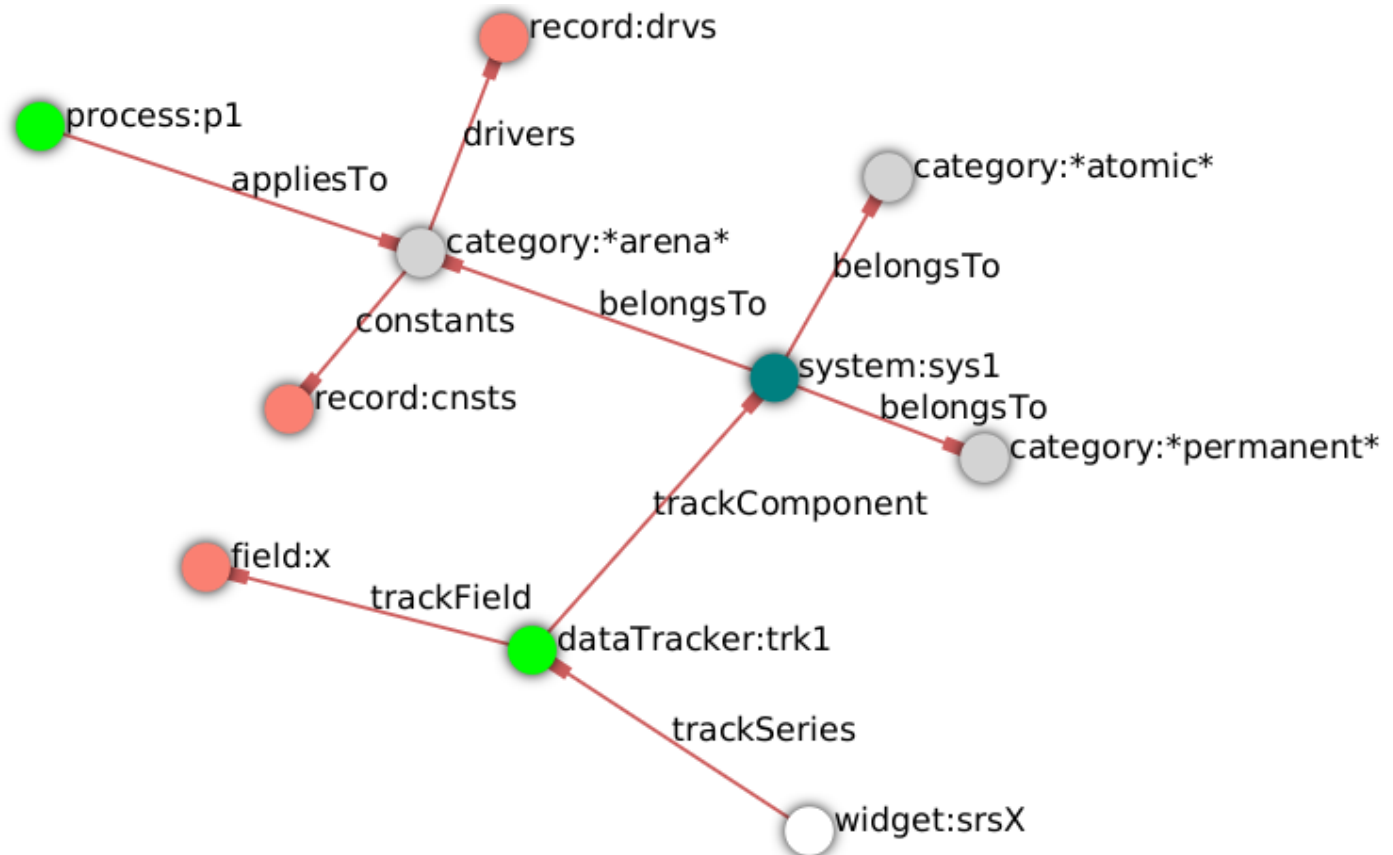


Figure 36. A view of the specification graph for Tutorial 1 showing only relevant cross-links using the Spring layout (L4).

That's the end of this tutorial. Recreate this project at any time from the main menu (Project → New → Tutorials → 1 Logistic).

5.1.6. Next

The next tutorial (Tutorial 2) will demonstrate linking this project to an IDE and adding some Java program code.

5.2. Tutorial 2: Linking a 3Worlds project to a Integrated Development Environment (IDE)

5.2.1. Introduction

The previous tutorial used code snippets to insert Java code into the model specifications. In this tutorial we show how to use an Integrated Development Environment (IDE) to achieve the same end. Generally, the only way to develop any but the simplest of models is to code in a professional IDE such as *Eclipse* (<https://www.eclipse.org/downloads/>), *IntelliJ* (<https://www.jetbrains.com/idea/>) or *NetBeans* (<https://netbeans.apache.org/>). At the time of writing, linking specifications to a Java project has only been tested with the *Eclipse* IDE.

5.2.2. New Java project

First create a Java project in *Eclipse*.

1. In *Eclipse*, create a new Java project and name it `tut2` (first letter lower case).
2. Right-click on the new project and select `properties` → `Java Build Path`.
3. Under the `Libraries` tab, highlight `Classpath` and click the `Add External JARs...` button to the right.
4. Navigate to the `3w` directory and select `tw<os name>.jar`.
5. Click `Apply` and `close`.

This installs the `3Worlds` dependencies for this java project.

5.2.3. Specifications

Now create a `3Worlds` project using tutorial 1 as a starting point.

1. From *ModelMaker* main menu select `New → Tutorials → 1 Logistic` name it `Tut2`

Next link this project to the Java project created above.

2. From the *ModelMaker* main menu, select `Edit → Java project → Connect...`.
3. Navigate to your *Eclipse* workspace and select `tut2`.
4. In the *Eclipse Project Explorer* view, right-click on `tut2` and select `Refresh`.
5. In the *Eclipse* main menu, select `Project -> clean`, flag `tut2` and click `OK`.

While editing in *Eclipse*, a task message may appear in *ModelMaker* from time to time complaining about file dates. If so, make sure your Java files are saved, refreshed and the project clean. The message will disappear when the project is re-compiled in *ModelMaker* using the *Verify* button in the bottom left-hand corner.

5.2.4. Writing Java code

In the `src` folder of `tut2`, you'll now find a new package called `code.sys1` and `code.sys1.generated` (Figure 37). These have been generated by *ModelMaker* upon linking. In the `code.sys1` package is a Java file with the same name as the `3Worlds` project: `Tut2.java`. This single class is where you add all your code. The other classes in the `generated` package are not intended to be edited. If you do edit these, your edits will be overwritten whenever the specifications are edited in *ModelMaker*.

1. Open `Tut2.java` in the *Eclipse* editor.

Near the bottom of the file are two methods (Figure 38): `chaos` and `init1`, mapped from the names of the function and initFunction nodes we created in `Tut2` with *ModelMaker*. Within the body of these methods are a pair of insertion markers:

- Code insert `Begin-- >`; and
- Code insert `End---- <`.

The code snippet property that was present in the original `Logistic` model this tutorial derives from has already been added for both the `chaos` and `init` methods.

`chaos :`

```
focalDrv.x = r*x*(1-x);
```

JAVA

`init1 :`

```
focalDrv.x = 0.001;
focalCnt.r = 3.7;
```

JAVA

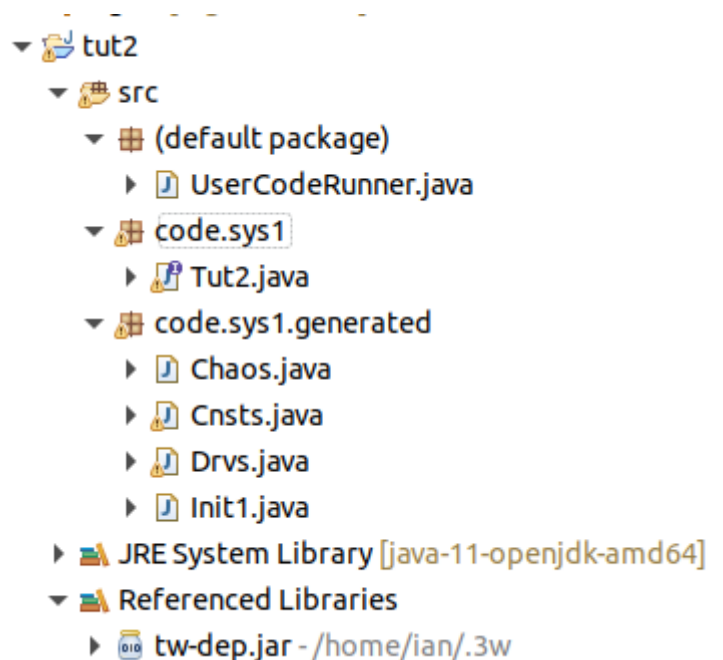


Figure 37. The package structure in *Eclipse* after linking Tutorial 2 to a Java project.

```

package code.sys1;

* import static java.lang.Math.*;□
// Hey, model developer! You may add your own imports here as needed

* * <h2>Model-specific code for model <em>Tut2</em></h2>□
public interface Tut2 {

*   * <p><strong>Chaos</strong> method of type <em>ChangeState</em>: change the state, i.e.
*   public static void chaos(
    double t,                                     // current time
    double dt,                                    // current time step
    double x,                                     // focal component drivers ± 0.0 ∈]-∞,+∞[
    Chaos.FocalDrv focalDrv,                     // next drivers for focal component
    double r,                                     // focal component constants ± 0.0 ∈]-∞,+∞[
    Random random) {                             // random number generator
    // chaos ---- Code insert Begin-->
    focalDrv.x = r * x * (1 - x);
    // chaos ---- Code insert End----<
}

*   * <p><strong>Init1</strong> method of type <em>SetInitialState</em>: sets the initial :
*   public static void init1(
    double x,                                     // focal component drivers ± 0.0 ∈]-∞,+∞[
    Init1.FocalDrv focalDrv,                     // next drivers for focal component
    double r,                                     // focal component constants ± 0.0 ∈]-∞,+∞[
    Init1.FocalCnt focalCnt,                     // new constants for focal component
    Random random) {                            // random number generator
    // init1 ---- Code insert Begin-->
    focalCnt.r = 3.7 + 0.001*random.nextDouble();
    focalDrv.x = 0.01;
    // init1 ---- Code insert End----<
}

}

```

Figure 38. The main source code file ('Tut2.java').

You can also create other Java classes or import third-party libraries to extend you model code. This will be discussed in later tutorials. There are two "import" markers at the top of the main source code file. Add any dependencies to third-party or other java classes you use.

! When editing the main java source code file, any imports your IDE might insert automatically will not necessarily appear between the import markers. Take care to edit these statements so they do appear between the import statements. These imports are in fact a code snippet property found in the specification root node.

Because these two methods are mapped to this generated file from the specifications created in *ModelMaker*, it follows that changes to the specifications may result in changes to these methods. So the question arises, how does *ModelMaker* manage this without losing the Java code you write?

If there have been changes to the number, name or type of `function` used in the specification, changes to anything in the `dataDefinition` sub-tree, or changes to the `roles` of `components`, *ModelMaker* will back up your old file as a text file with a unique name (e.g. `_Tut2_1.txt`, `_Tut2_2.txt` etc...) within the current java package. This way you can copy and paste you code back to the proper place. Otherwise *ModelMaker* will never overwrite this file.

! When working simultaneously with an IDE and *ModelMaker*, it is useful to keep the code snippets in *ModelMaker* up-to-date by re-importing them from time to time [ctrl+I]. If changes made in the specifications by *ModelMaker* result in a new file being created, code snippets will be carried over to the new file.

When *ModelMaker* searches for differences between previous and current versions of this Java file, it will ignore any

comments and the code you have added between the insertion markers. Therefore, although you can do as you please with the generated comments, you should take care not to alter the text of the markers themselves.

If the task list is empty, you can now execute the specification by deploying them from *ModelMaker*.

Running the simulation directly from *ModelMaker* is a convenient way to speed up the turn-around time in developing and testing specifications. Once the specifications are stable, you can run your model from a jar: the jar file that is created in your project directory in this case `Tut2.jar`. To do this:

2. Open a terminal, change to your project directory and enter the following text: `java -jar Tut2.jar`

Finally we discuss how to debug your equations by running your model from the IDE.

Debugging your code

Once the simulator specification is valid (i.e no tasks in the task list) you can quit *ModelMaker* and debug your code from the IDE.

In the default package of the linked java project (`tut2`) is a java file called `UserCodeRunner.java`. This was created the first time you linked the project made with *ModelMaker* to the Java project. By placing breakpoints in the methods you added to `Tut2.java` above, you can debug by running from this `main` class. The correct command-line arguments have already been provided.

5.2.5. Next

The next tutorial will introduce the use of tables and some additional output `widgets` to add to the simulator's user interface.

5.3. Tutorial 3: Using tables

5.3.1. Introduction

This tutorial introduces the use of tables in *ModelMaker* by implementing a multi-species competitive Lotka-Volterra equation (https://en.wikipedia.org/wiki/Competitive_Lotka%20Volterra_equations) (CLV).

The equation has three *constants* and one *driver*; all of which are tables:

- constants:
 - K : carrying capacity;
 - r : growth rate;
 - α : inter-specific competition coefficient;
- driver:
 - x : population size.

5.3.2. Specifications

Apart from the use of tables and a more elaborate model equation, these specifications are very similar to those of Tutorial 1. *ModelMaker* provides a library of model templates for common model patterns such this. One of these, the `SimpleClock` template, has already defined a basic structure, dynamics and a minimal user interface ([fig-simple-clock-template]).

1. Start *ModelMaker* and create a new: `Projects → New → Templates → 2 SimpleClock` and name the project 'Tut3'.

Data definition

Data is defined through the recursive use of records and tables: records can contain tables and tables can contain records without limit. Records can also contain fields of any primitive Java type. Tables can also contain Java primitive

types as well as user-defined records (Figure 39).

Tables can have any number of dimensions.

There is usually more than one way of defining the data for any model. For the CLV, we will use a 2-dimensional table for *alpha*, a 1-dimensional table for *r* and *K* and another 1-dimensional table for the population *x* (Figure 40).

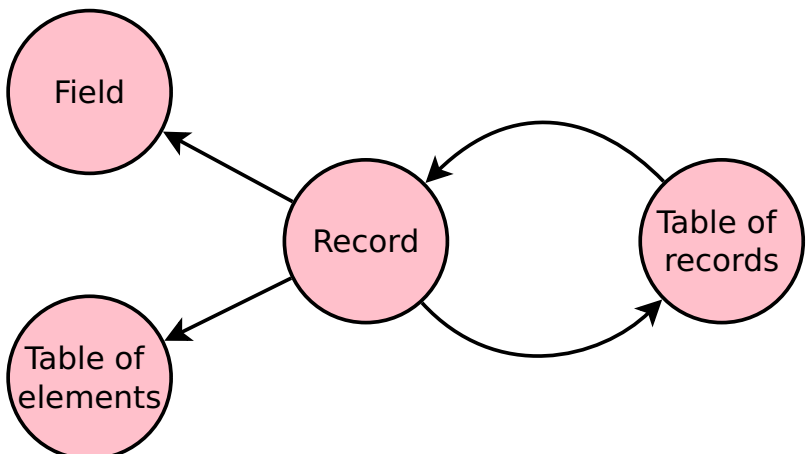


Figure 39. Data is defined through the recursive use of records and tables. A definition begins with a record. Leaf nodes of the resulting graph must be either Fields or Tables that define a standard Java type (element) such as Double, Integer etc.

2. Collapse all sub-trees from the root node and then expand `dataDefinition:dDef`.
3. Create a `table` as a child of `record:csts`, name it 'alpha', choose `dataElementType` and then set its type to `Double` (the default).
4. Create a second `table` as child of `record:csts`, name it 'r' and choose `dataElementType` and then set its type to `Double`.
5. Create a third `table` as child of `record:csts`, name it 'K' and choose `dataElementType` and then set its type to `Double`.
6. Create a `table` as child of `record:drvs`, name it 'x', choose `dataElementType` and set its type to `Double`.

To complete the definition, we must set the table dimensions and indexing order for the 2-dim table 'alpha'.

Only one dimensioner is needed: the number of species. This is done by creating a single `dimensioner` node to be associated with all these tables.

7. Create a `dimensioner` as a child of `dataDefinition:dtDef` and name it 'nspp' (number of species).

A new task appears in the task list saying the 'size' property must be greater than zero.

8. Click on `dimensioner:nspp` and go to the `Selected properties` tab. Set a value of `npp#size` to 4 (four species).
9. Right-click on each table node in turn and select `New edge → dimensioner:nspp`. Repeat this for `table:alpha` to get the second dimension.

Now set the rank order for the indexing of `table:alpha`.

10. Click on `table:alpha` and examine its properties in the `Selected properties` editor.
11. Set the value of one of its rank properties to 1 - it doesn't matter which.

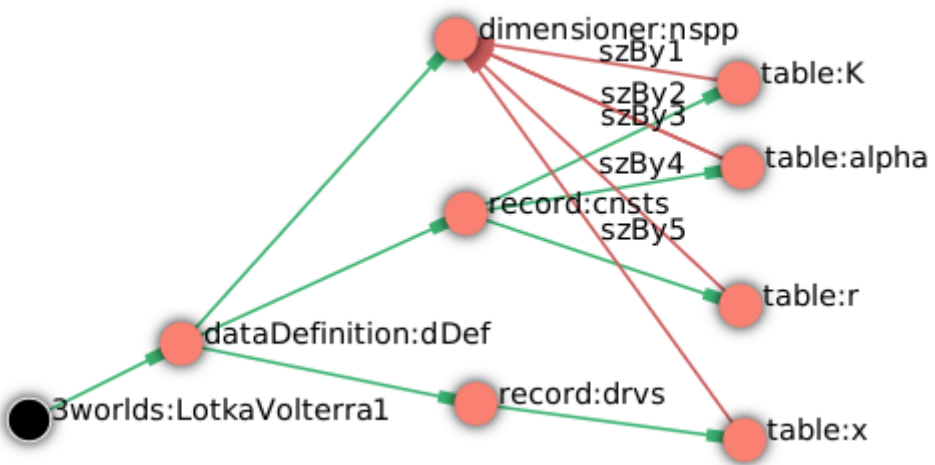


Figure 40. The `dataDefinition` specifications for Tutorial 3: three tables of one dimension and one of two dimensions ('alpha'). This size of each table is the same so only one dimensioner is required.

Finally, add some meta-data to each table by adding suitable text to the `description` property of each table or field.

Structure

The `SimpleClock1` template has already defined the role of `system.sys` as *permanent*, *atomic* and belonging to the *arena* - the definition of a non-hierarchical model. There is nothing additional needed for this section.

Dynamics

Although the task list is now empty (i.e. the specifications are now deployable), we have yet to write the model equation and provide some output. We can now rename the function provided by the template to something meaningful in the present context and add a data tracker to follow *x*.

1. Expand the `system.sys1` sub-tree from the root node and re-apply the layout [Alt+L].
2. Right-click on `Function:F1`, select `Rename node` and name it 'Growth'. If the prompt is unresponsive, remember to begin the name with an upper case letter.
3. Create a `dataTracker` as a child of `process:p1` and select `DataTrackerD0` as the type (the default).
4. Create an edge from the data tracker to *x*: `trackTable → table:x`.

There are now two tasks in the list, one about 'indexing' and the other asking which 'component' to track. As with the previous tutorials, the component to track is the `system.sys1` node.

5. Create the edge `trackComponent → system:sys1` from the data tracker node.

As we are using a scalar data tracker (`DataTrackerD0`) to follow a table, we can specify which elements of the table to track. If we want to track all elements of the table, we need do nothing as a blank entry for the `trks#index` property assumes this. Indexing is a property of the edge between data tracker and the table. Edge properties appear in the property list of nodes that are at the start of the edge (thin line) - in this case `dataTracker:trk1`.

This indexing will provide four data outputs. Indexing statements can select any number of contiguous or discontiguous table elements. The syntax is similar to that found in the **R statistical software** (<https://www.r-project.org/>).

User interface

We can now add some additional widgets to the user interface as the template provided only a controller.

1. Hide all nodes and expand the `userInterface/gui` node.
2. Add a `tab` as a child of `userInterface/gui`.

3. Add a `widget` as a child of `tab:tab1`, name it 'srsx' and select `TimeseriesWidget1` as the widget class.

4. Add a second `widget` as a child of `tab:tab1`, name it 'tblx' and select `TableWidget1`.

Both these widgets are compatible with this data tracker class: the `TimeseriesWidget1` produces a chart while the `TableWidget1` displays the data as a continuously updated table. The task list requires these widgets to be connected to a data tracker.

5. Right-click on each of these widgets in turn and select `New edge → trackSeries → dataTracker:trks`.

6. Set the property `tblx#order` to '1' to arrange their positions in `ModelRunner` as `srsx` on the left (`order = 0`) and `tblx` to the right (`order = 1`).

Save the specifications (`Ctrl+s`) and they're now ready to run (Figure 41, Figure 42). The next step is to create a Java project to write the 'Growth' and 'Init1' functions, the latter having been provided by the `SimpleClock` template we started this tutorial with.

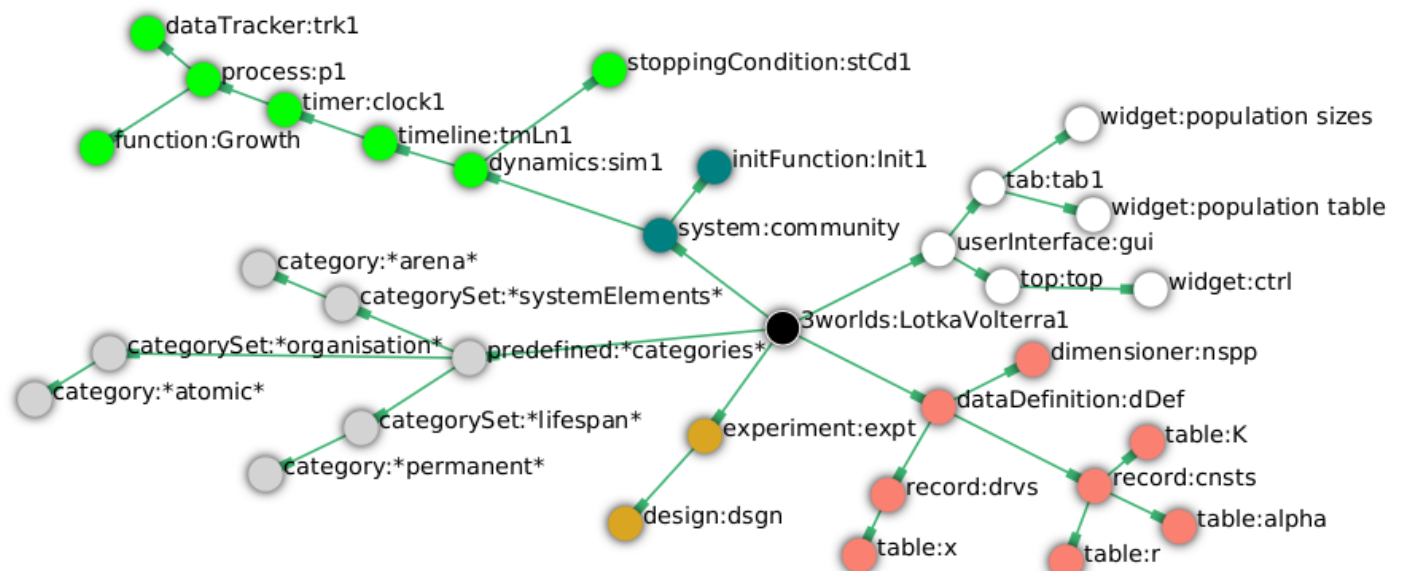


Figure 41. Tree graph of the final specification graph for Tutorial 3.

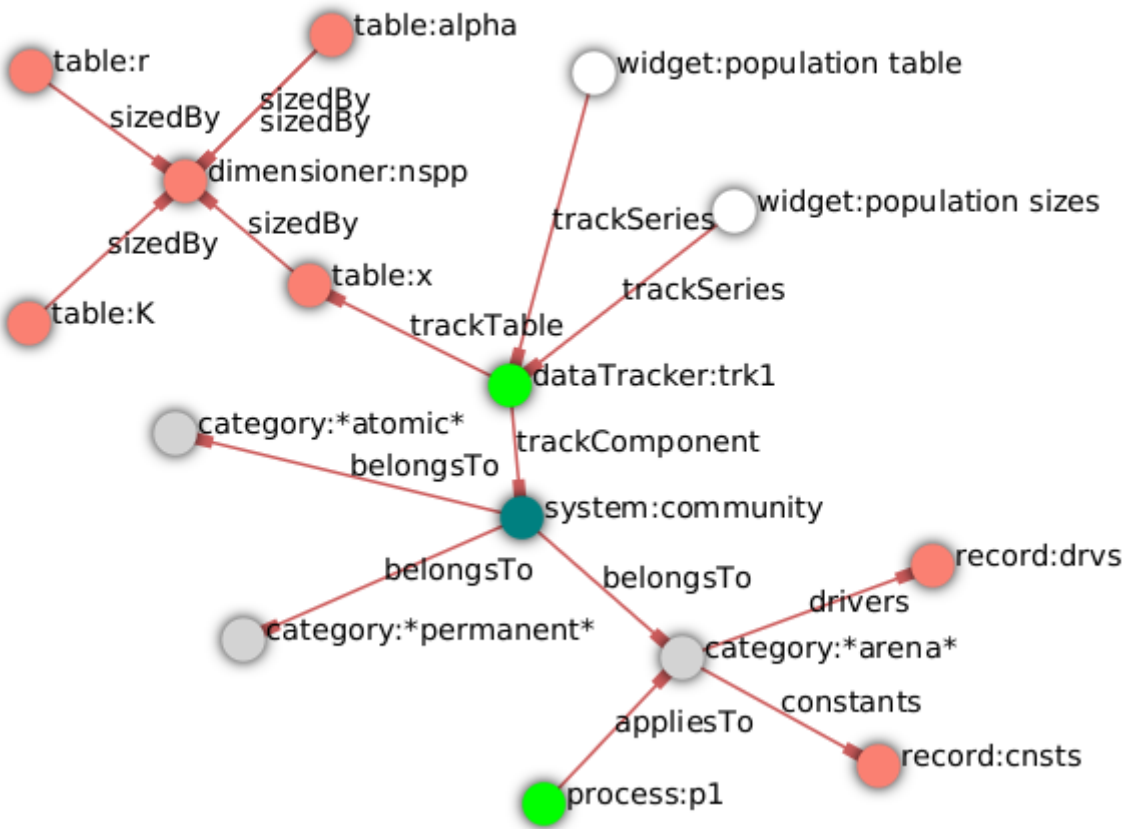


Figure 42. Cross-links of the final specification graph for Tutorial 3.

Link to a Java project

1. Follow the steps in Tutorial 2 to create a Java project with *Eclipse* and name it 'tut3'.
2. Link it to this *ModelMaker* project.
3. Open *Tut3.java* in the *Eclipse* editor and enter the following source code between the relevant insertion markers:

`init:`

```

for (int i = 0; i < r.size(0); i++) {
    focalCnt.r.setByInt(random.nextDouble() * 2.0, i);
    focalCnt.K.setByInt(0.2 + random.nextDouble(), i);
    for (int j = 0; j < alpha.size(1); j++) {
        if (i == j)
            focalCnt.alpha.setByInt(1.0, i, j);
        else
            focalCnt.alpha.setByInt(random.nextDouble(), i, j);
    }
}
for (int i = 0; i < x.size(0); i++)
    focalDrv.x.setByInt(0.2, i);

```

JAVA

The above method simply initialises the equation constants: growth rate (*r*), carrying capacity (*K*) and the interspecific competition coefficient (*alpha*) to random values and the population size (*x*) to 0.2. Notice that the generated code includes a detailed javadoc that reminds the forgetful user of all the table dimensions, driver and constant units, etc. (Figure 43).

There is a default random number generator (RNG) available to all functions. In later tutorials we will show how the specifications can factor any number of RNGs into groups. For example, one RNG can be assigned to functions of a particular type such as those effecting reproduction or mortality.

ModelMaker has two types of RNG classes in addition to the standard Java RNG. These two are faster and produce streams of higher quality than the standard Java RNG. There are also various ways of seeding RNGs to ensure their uniqueness and to help with debugging.

growth :

```
double integrationStep = 0.01;
double[] dxdt = new double[x.size(0)];
for (int i = 0; i < x.size(0); i++) {
    double sum = 0;
    for (int j = 0; j < alpha.size(1); j++)
        sum += alpha.getByInt(i, j) * x.getByInt(j);
    dxdt[i] = r.getByInt(i) * x.getByInt(i) * (1 - sum / K.getByInt(i));
}
for (int i = 0; i < dxdt.length; i++)
    focalDrv.x.setByInt(x.getByInt(i) + dxdt[i] * dt * integrationStep, i);
```

JAVA

void code.sys1.Tut3.growth(double t, double dt, DoubleTable x, FocalDrv focalDrv, DoubleTable K, DoubleTable alpha, DoubleTable r, Random random)

Growth method of type *ChangeState*: change the state, i.e. the values of the descriptors of a system component

- applies to categories { *arena* }

- follows timer *clock1* of type [ClockTimer](#), with time unit = 1 t.u

Parameters:

t current time

dt current time step

x focal component drivers population size dim = [4] ± 0.0 €]0.0,1.0[

focalDrv next drivers for focal component

K focal component constants carrying capacity dim = [4] ± 0.0 €]0.2,1.0[

alpha focal component constants inter-specific competition coefficient dim = [4,4] ± 0.0 €]0.0,1.0[

r focal component constants growth rate dim = [4] ± 0.0 €]1.0,2.0[

random random number generator

Figure 43. Example of the generated javaDoc for the `growth` function of Tutorial 3. The figure assumes correct meta-data has been added to the definitions of `K`, `r`, `alpha` and `x`.

Following a note from 'Tutorial 1', it is a good idea to keep the code snippets in *ModelMaker* up-to-date with changes in the source code.

4. In *ModelMaker* select `Edit → Import snippets from IDE` and save (`Ctrl-s`).

The model is now ready to run (Figure 44). However, you may want to change the time duration of the simulation from the template default of 100 to 1,000 steps.

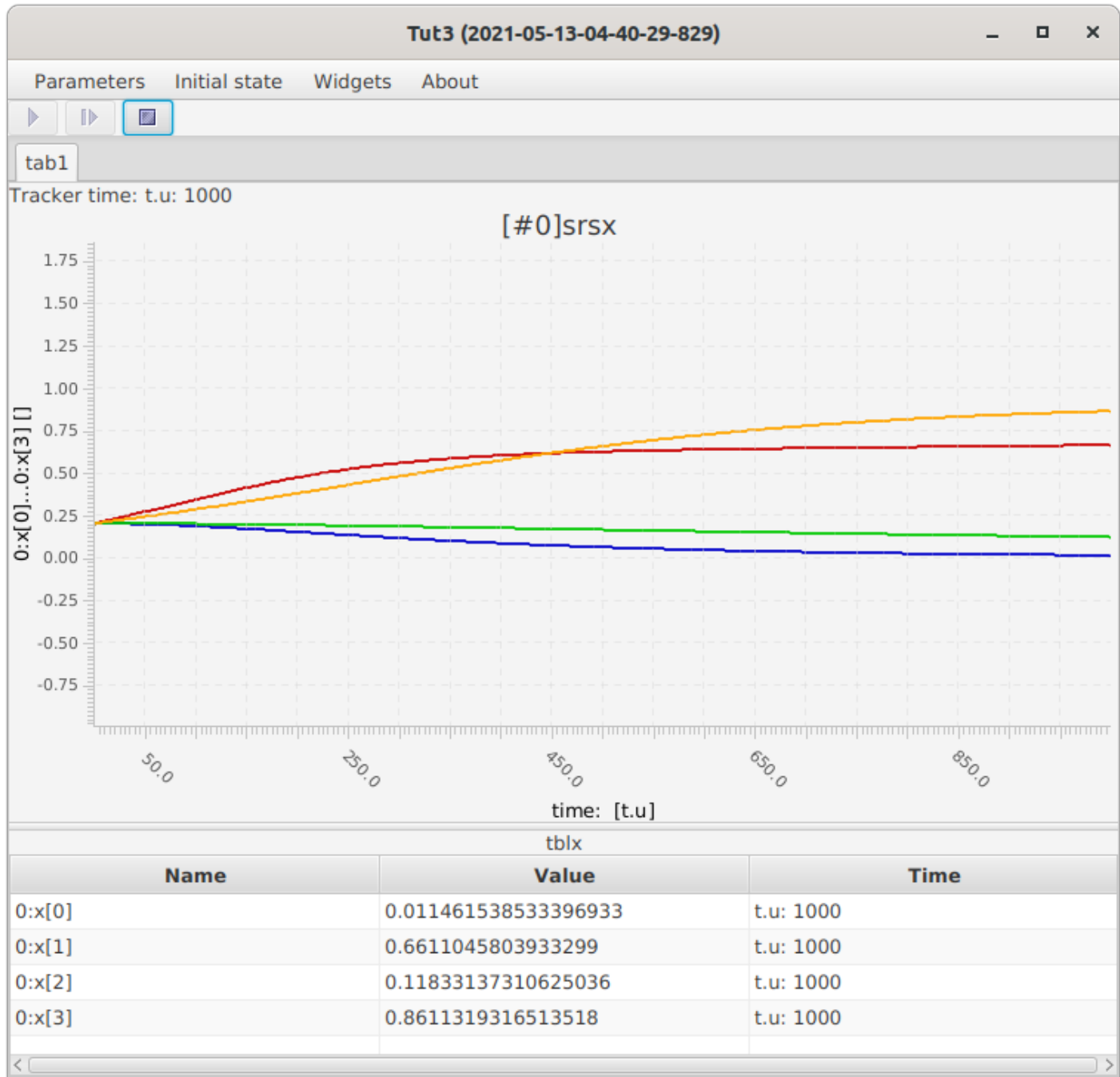


Figure 44. ModelRunner executing Tutorial 3. Note that the property `tab1#orientation` has been set to `vertical` in this figure. This displays the time series graph above the table.

5.3.3. Next

The next tutorial elaborates the Lotka-Volterra model by developing the `structure` sub-tree through the addition of a disturbance component.

5.4. Tutorial 4: Elaborating the model structure: Testing the Intermediate Disturbance Hypothesis

5.4.1. Introduction

In this tutorial we add a disturbance to effect a community of species modelled with the competitive Lotka-Volterra equation (CLV) from Tutorial 3. This model could then be used to examine the Intermediate Disturbance Hypothesis (IDH: Connell, 1978).

5.4.2. Specifications

Begin this tutorial using the Lotka-Volterra specifications from the `Tutorials` menu as a starting point.

1. Create a new project from `Tutorials → 2 LotkaVolterra` and name it 'Tut4'.

As the tutorial title suggests, we will be *elaborating* the model structure. The Lotka-Volterra model has the simplest of structures: so simple that its `System` node can be named `community`. In the model we are developing here, this name would be misleading since we will have not only a 'community' but also a 'disturbance' (i.e. the system node will no longer be 'atomic' but an 'assemblage'). Therefore it will be clearer if we change the name of the system node back to its nondescript default name of `sys1`.

2. Right-click on `system:community` and rename 'community' to the default name 'sys1'.

Data definition

Data can play one of three *roles* in the *3Worlds archetype*; as:

- constants*,
- drivers*, or
- decorators*.

Thus far we have only made use of data as *drivers* and *constants*. The third role, *decorators*, are values derived in some way from the other two. Importantly, they are re-initialised at the beginning of each time step. We need a species diversity index and this is a good example of a *decorator* being derived from relative abundance of each species (`'x[n]'`).



Decorators are re-initialised (zero, false, null etc.) at the start of each time step.

1. Collapse all nodes except for the `dataDefinition` sub-tree, hide the *cross-links* ('X'), and re-apply the layout ('L').
2. Add a new `record` as a child of `dataDefinition:dDef` and name it 'decs'.
3. Add a `field` as a child of `record:decs`, name it 'di' (diversity index) of type `Double`.

As noted previously, these names are just for clarity: they don't impose roles as *drivers*, *constants* or *decorators*. Here, the records are named to indicate their intended roles just for clarity. Their role will later be formally defined by a relationship with a `category` which we create when building the `structure` specifications.

While building the data definition (Figure 45), expect task messages to accumulate until these roles have been defined.

We now define some data for the simple disturbance model: frequency and intensity.

4. Add a new `record` to `dataDefinition:dDef` and name it 'distCnsts' (disturbance constants).
5. Add two fields to this record, both of type `Double` called 'freq' and 'inten' respectively.

The records carried over from the previous tutorial should now be given better names. A trick with the Tree layout functions is that they order children alphabetically. To arrange nodes so that related nodes of a particular role are close together, prefix the name with the same characters - in this case 'dist'. We should also prefix all the data associated with the CLV with 'comm' (community).



Tree layout algorithms sort child nodes in alphabetical order. Consider this behaviour when choosing node names.

6. Rename `record:csts` to `record:commCnsts`, `record:drv`s to `record:commDrv`s and `record:decs` to `record:commDecls`.
7. Re-apply the layout and the nodes should now be arranged in a more sensible order.

To display an xy plot of diversity by time since disturbance, we can add a driver to the `commDrv`s record to record this.

8. Create a `field` of type `Double` as a child of `record:commDrv`s and name it 'tsd'.

Fill out the meta-data properties with at least the description, range and units if relevant. This information will be incorporated in the comments of the generated Java code.

9. Add the following descriptions to the new data definitions properties:

- `freq#description` : 'average return time'
- `inten#description` : '% population decrease'
- `di#description` : 'Shannon's diversity index'
- `tsd#description` : 'Time since disturbance'

Finally, change the number of species in the model to a larger value.

10. Set the value of `nspp#size` to 40.

For simplicity, we have defined `inten` as a constant rather than a driver.

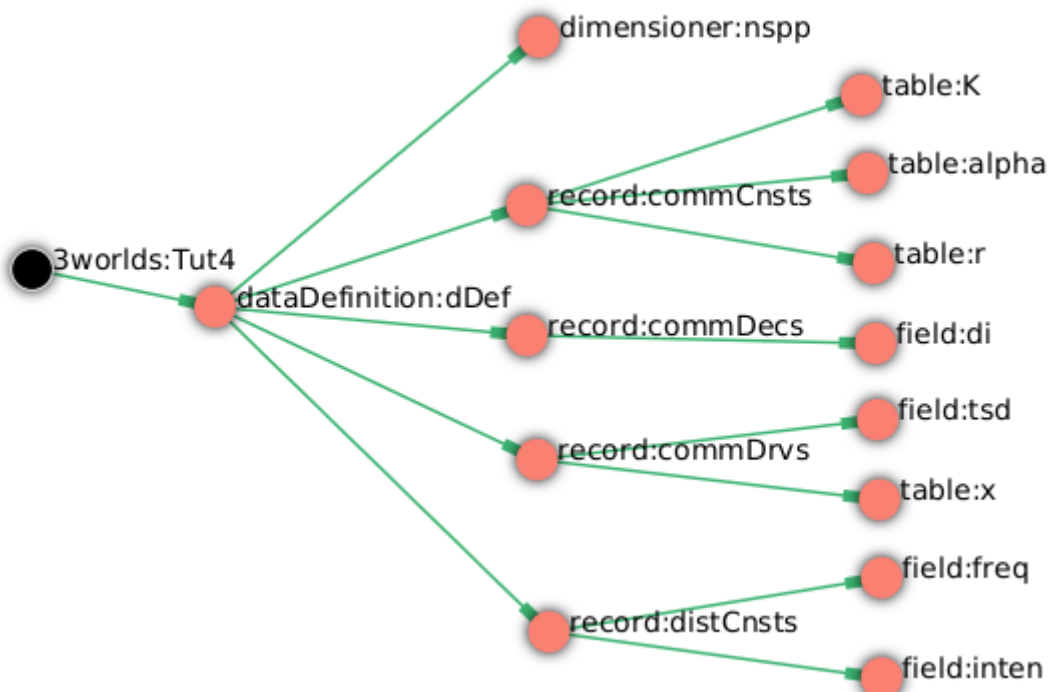


Figure 45. The data definition sub-tree for Tutorial 4. Tree layout algorithms sort nodes in alphabetical order. Keeping this in mind when naming nodes can help keep your specifications in useful order.

That's all the data needed for this model. We can now specify the system structure.

Structure

Recall from the previous tutorial, that `system:sys1` is acting as a single component. Its `role` was defined as a permanent, indivisible global component:

- i. `permanent`;
- ii. `atomic`;
- iii. belonging to the `arena`;

The `arena` is a special category. One and only one component must belong to this category. Any data associated with it (`constants`, `drivers` and `decorators`) is available to all `functions`: the data is global. The system (`system:sys1`) was also being tracked by the data tracker. This relationship was removed above because `x`, the population being tracked,

will now be part of a category related to the community component of the model i.e 'x' will no longer be global data.

The *role* of a `component` is defined by its `componentType`. We'll need two `ComponentTypes`; one for the community (the CLV equation) and one for the disturbance. We also need a `relationType` to define the relationship between the disturbance and the effected community.

1. Right-click on `system.sys1` and collapse the `dynamics:sim` sub-tree.
2. Create a `structure` as child of `system:sys1`.
3. Create a `componentType` as a child of `structure:struc1` and name it 'distType'.
4. Create a `component` as child of `componentType:distType` and name it 'dist'.
5. Create a second `componentType` as child of `structure:struc1` and name it 'commType'.
6. Create a `component` as child of `componentType:commType` and name it 'comm'.
7. Create a `relationType` as child of `structure:struc1` and name it 'distEffectComm'.

Before proceeding to the definition of the *roles* of these component types, we should redefine the *role* of `system:sys1` that we inherited from Tutorial 3 when starting this project.

The node `system:sys1`, in aliasing as a `componentType`, must now be redefined as an `assemblage` with no functions or data associated with it - that is, it is no longer `atomic`.

8. Right-click on `system:sys1` and select `Delete edge` → `belongsTo` → `category:*atomic*`.
9. Right-click again on `system:sys1` and select `New edge` → `belongsTo` → `category:*assemblage*`.

Remove the data associated with the *arena*.

10. Right-click on the root node (`3worlds:Tut4`), expand the `predefined:*categories*` and re-apply the layout.
11. Right-click on `category:*arena*` and select `Delete edge` → `drivers` → `record:commDrvs`.
12. Right-click again on `category:*arena*` and select `Delete edge` → `constants` → `record:commCnsts`.
13. Right-click on the root node and collapse `predefined:*categories*` and re-apply the layout [Alt+L].

The node `system:sys` is now defined as simply a *permanent assemblage* belonging to the *arena*.

Now define a `categorySet` to partition data between the disturbance and community. Category Sets contain mutually exclusive categories: something can belong to one or the other but not both. Since disturbance and community is all there is in this model, this is the 'world' of the model so this seems a reasonable name for this set of categories i.e the 'world' comprises a community and a disturbance category: something can belong to one or other of these but not both.

14. Create a `categorySet` as child of `structure:struc1` and name it 'world'.
15. Create a `category` as child of `categorySet:world` and name it 'distCat'.
16. Create another `category` as child of `categorySet:world` and name it 'commCat'.

Define the data for these new categories.

16. Right-click on `category:commCat` and create the following edges:
 - i. `drivers` → `record:commDrvs`.
 - ii. `constants` → `record:commCnsts`.
 - iii. `decorators` → `record:commDecs`.
17. Right-click on `category:commDist` and select `New edge` → `constants` → `record:distCnsts`.

18. Expand the `dataDefinition:dDef` sub-tree, toggle the *cross-links* to confirm you have done the links as above.

Now define the *roles* of the new component types for the community and disturbance. Both belong to the *permanent*, *atomic*, and *component* categories.

18. Right-click on `componentType:commType` and create `belongsTo` edges to:

- i. `category:*permanent*`,
- ii. `category:*atomic*`,
- iii. `category:*component*` and
- iv. `category:commCat`.

19. Right-click on `componentType:distType` and create `belongsTo` edges to:

- i. `category:*permanent*`,
- ii. `category:*atomic*`,
- iii. `category:*component*` and
- iv. `category:distCat`.

Now define the relation between disturbance and the community.

20. Right-click on `relationType:distEffectComm` and create edges:

- i. `fromCategory` → `distCat`.
- ii. `toCategory` → `commCat`.

Finally, add initialisation functions for the disturbance and community component types:

22. Create an `initFunction` as child of `componentType:commType` and name it 'InitComm'.

23. Create an `initFunction` as child of `componentType:distType` and name it 'InitDist'.

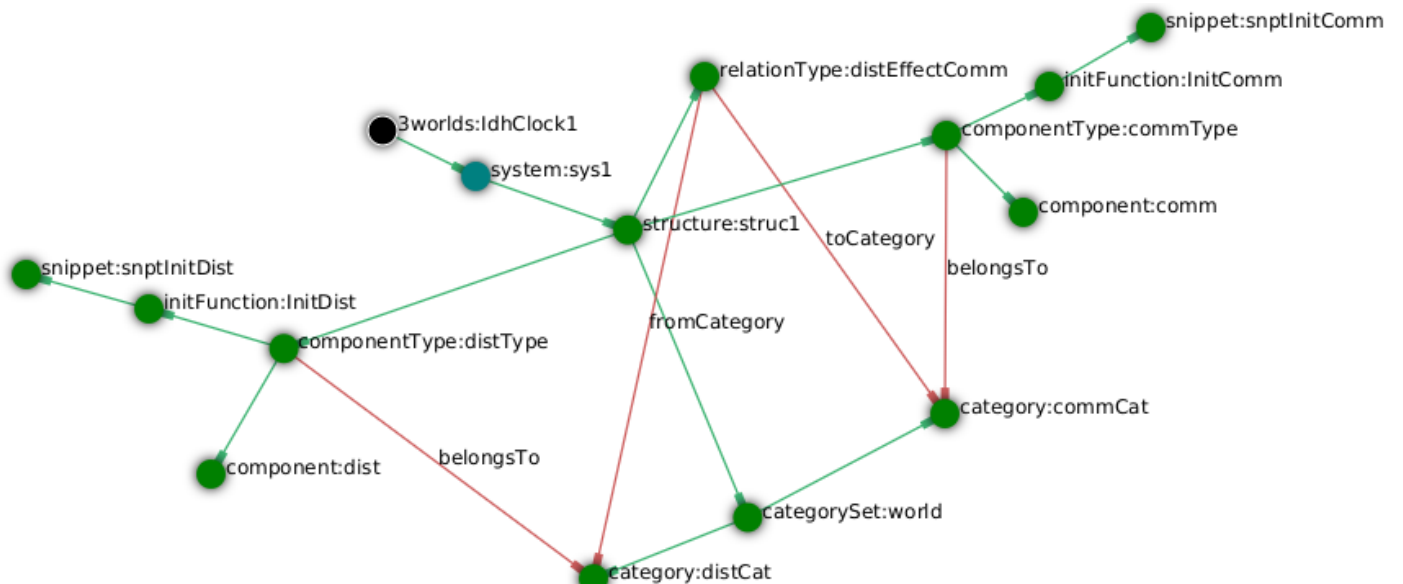


Figure 46. The structure sub-tree for Tutorial 4. .

That's all that is required in this section. To finish up, we now need to connect various sub-trees of the graph to each other. Foremost among these is to associate processes with the new structure (Figure 46).

In this section we add two additional functions: one to initiate a disturbance and a second to apply the disturbance to the community.

Before proceeding, some explanations are required to make sense of the following steps. Ignoring `dataTrackers` for the moment, a `process` can have one or more `functions`. However, the relationships (`cross_links`) between the model's dynamics, data and structure are defined at the level of `processes` not `functions`. If we don't care what order the functions are called in, and they all apply to the same data and structural elements, then we need only one process to contain all the functions. However, this is not the case here. The three functions: community growth, disturbance occurrence, and disturbance effects, must have access to different data, components and component relationships. Specifically, whether or not a disturbance is to occur depends only on disturbance data (in this model). Also order is important: it must come before disturbance effects are applied to the community. The disturbance effects applies to a relationship: that of the disturbance and the community. The growth of the community only depends on community data (the Lotka-Volterra equation). In addition, we make the decision (in this model) that the disturbance can only occur after the growth of the community, mimicking a late growing season disturbance.

For clarity, we will prefix the names of `processes` in such a way that the layout algorithm will place them in execution order.

1. Collapse the `dataDefinition` sub-tree and expand `system:sys1`.
2. Create a `process` as child of `timer:clock1` and name it 'p1DistOcc'.
3. Create a `function` as child of `process:p1DistOcc` and name it 'DistOcc' and set its type as `RelateToDecision`.
4. Create a second `process` as child of `timer:clock1` and name it 'p3DistEffectComm'.
5. Create a `function` as child of `process:p3DistEffectComm`, name it 'DistEffectComm' and set its type to `ChangeOtherState`.

The name the growth process and function should now be changed to something more appropriate:

6. Rename `process:p1` to `process:p2CommGrowth`.
7. Rename `function:Growth` to `function:CommGrowth`.

The function `function:CommGrowth` will perform the same task as it did in the previous tutorial: calculate the next state of x from its current state. At the same time we can calculate the diversity index and increment a counter to record the time since the last disturbance.

The function `function:DistOcc`, a `RelateToDecision` function, will decide if a disturbance is to occur. If it returns `true`, a relation will be formed between the, yet to be specified, disturbance and community categories.

If a disturbance occurs, `function:DistEffectComm` will implement the consequences of the disturbance on the community **but at the next time step**.



All methods read data at the current time step and write data of the next time step. That is, always be mindful that $x[t+1] \leftarrow f(x[t])$. In code, $x[t+1]$ is `focalDrv.x` and $x[t]$ is just `x`.

As a reminder of this ordering, we have prefixed the process that creates the temporary (ephemeral) relation between disturbance and the community with 'p1' so that the layout (if using a *Tree* layout) will display processes in a sensible order.

First, show the *cross-links* and set `process:p2DistEffectComm` to depend on `process:p1DistOcc`.

8. Show the *cross-links* (X).
9. Right-click on `process:p2DistEffectComm` and select `New edge` → `dependsOn` → `process:p1DistOcc`.

To force the disturbance effects to take place *after* the community growth (i.e. a late growing season disturbance), make `process:p2DistEffectComm` depend on `p2CommGrowth`.

10. Right-click on `process:p3DistEffectComm` and select New edge → dependsOn → `process:p2CommGrowth`.

After re-applying the layout (assuming a Tree layout is in use), it's clear why these process names were prefixed by p1, p2 and p3.

At this time, `dataTracker:trk1` is set to track the component `system.sys1`. There will be no global data to track in this model as there was in the previous Lotki-Volterra so we should remove this edge.

11. Right-click on `dataTracker:trk` and select Delete edge → `trackComponent → system.sys1`.

Finally, because there will be no global data in this model, `system:sys` will not need an initialisation function. Therefore we can delete this node.

12. Right-click on `initFunction:Init1` and select 'Delete node'.

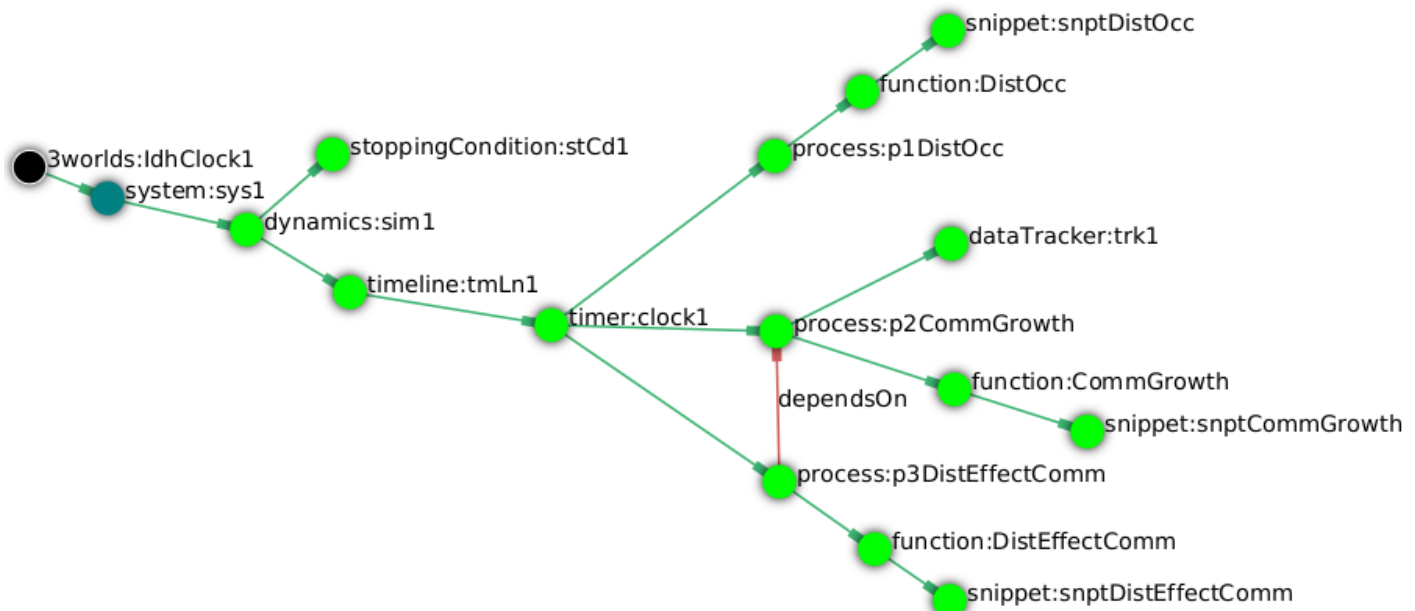


Figure 47. The system dynamics sub-tree for Tutorial 4. .

That's all that's required in this section (Figure 47). The task list indicates that the new processes must belong to some categories. To do that we need to move on to the structure of the specifications.

Relations between sub-trees

Currently, `process:p3CommGrowth` is applied to `category:*arena*`. We want re-apply this process to `category:commCat`.

1. Expand `dynamics:sim1` from `system:sys1`.
2. Expand `predefined:*categories*` from the root node and re-apply the layout.
3. Show *cross-links (X)*.
4. Delete the `appliesTo` edge between `process:p3CommGrowth` and `category:*arena*` and re-apply it by creating an `appliesTo` edge to `category:commCat`.

The task list now has two tasks asking to connect both disturbance processes to either a `category` or a `relationType`.

5. Create `appliesTo` edges from both `process:p2DistEffects` and `process:p1DistOccurrence` to

```
relationType:distEffectComm.
```

Note here that we expect this relation between the disturbance and the community to be reset every time step depending on whether or not `DistOccur` is true. That is, the relationship is ephemeral. This is achieved by setting the `lifespan` property of the `relationType:distEffectComm` to `ephemeral` (the default is `permanent`).

7. Change the property `distEffectComm#lifespan` to `ephemeral`.

The simulation can now be run but, of course, we have yet to add code to the various functions. Here, we'll just add code snippets but if you prefer, you can create a java project and add the code there instead (Section 5.2).

Java code

1. Copy and paste the code below to all the relevant `functionSnippet` properties. A convenient way to do this is to enter "func" into the search field in the *All properties* property editor and edit each in turn. Note: click the 'By Name' button at the top-left of the editor, otherwise the search function will only show properties containing 'func' within a property category. If you received compile error messages, check that you have pasted into the property of the correct function:

```
function:InitComm:
```

```
double initFreq = 1.0 / x.size();
focalDrv.x.fillWith(initFreq);
for (int i = 0; i < r.size(0); i++) {
    focalCnt.r.setByInt(random.nextDouble(), i);
    focalCnt.K.setByInt(5.0 + initFreq + random.nextDouble(), i);
    for (int j = 0; j < alpha.size(1); j++) {
        if (i == j)
            focalCnt.alpha.setInt(1.0, i, j);
        else
            focalCnt.alpha.setByInt(max(0.0001, random.nextDouble()), i, j);
    }
}
```

JAVA

```
function:InitDist:
```

```
focalCnt.freq = 5 + random.nextInt(50);
focalCnt.inten = random.nextDouble()*100;
```

JAVA

```
function:CommGrowth:
```

JAVA

```

// growth
double[] dxdt = new double[x.size(0)];
for (int i = 0; i < x.size(0); i++) {
    double sum = 0;
    for (int j = 0; j < alpha.size(1); j++)
        sum += alpha.getByInt(i, j) * x.getInt(j);
    dxdt[i] = r.getInt(i) * x.getInt(i) * (1 - sum / K.getInt(i));
}
for (int i = 0; i < dxdt.length; i++)
    focalDrv.x.setByInt(Math.max(x.getInt(i) + dxdt[i] * dt, 0.0), i);

// compute diversity
double xtot = 0.0;
for (int i = 0; i < focalDrv.x.size(0); i++)
    xtot += focalDrv.x.getInt(i);

focalDec.di=0.0;
for (int i = 0; i < focalDrv.x.size(0); i++)
    if (focalDrv.x.getInt(i) > 0.0)
        focalDec.di -= (focalDrv.x.getInt(i) / xtot) * log(focalDrv.x.getInt(i) / xtot);

// increment 'time since disturbance'
// use focalDrv.ts not tsd as the order of calling of this method and DistEffectsComm could be altered
focalDrv.ts = focalDrv.ts +1;

```

function:DistOcc :

```

if (random.nextDouble() < 1.0 / freq)
    return true;
else
    return false;

```

JAVA

function:DistEffectsComm :

```

for (int i = 0; i < other_x.size(); i++)
    if (other_x.getInt(i) > other_K.getInt(i) * inten / 100000.0) {
        otherDrv.x.setByInt(otherDrv.x.getInt(i) * other_K.getInt(i) * inten / 100000.0, i);
        // reset time since disturbance
        otherDrv.ts = 0;
    }
    // compute diversity
    double xtot = 0.0;
    for (int i = 0; i < other_x.size(0); i++)
        xtot += other_x.getInt(i);
    otherDec.di = 0.0;
    for (int i = 0; i < other_x.size(0); i++)
        if (other_x.getInt(i) > 0.0)
            otherDec.di -= (otherDrv.x.getInt(i) / xtot) * log(otherDrv.x.getInt(i) / xtot);

```

JAVA

5.4.3. User interface

We now need to modify the user interface, provided courtesy of the previous model (Section 5.3), to display `di` and `tsd`. We are simulating 40 species so their abundance (`x[n]`) is best placed in a separate time series chart and `di` in another. In addition, we need an xy plot of `di` by `tsd`. For this we need two additional `dataTrackers`.

1. For clarity, rename `dataTracker:trk1` to `dataTracker:trkx`.
2. Create another `dataTracker`, as a child of `process:p2CommGrowth`, name it 'trkdi' and select `DataTrackerD0` to follow this scalar value.
3. Right-click on `dataTracker:trkdi` and select `Add edge → trackField → field:di`.
4. Right-click again on `dataTracker:trkdi` and select `Add edge → trackComponent → component:comm`.

Now create a data tracker for `di` and `tsd` as an xy pair.

5. Create another `dataTracker`, as a child of `process:p2CommGrowth`, name it 'trkditsd' and select `DataTrackerXY`.
6. Right-click on `dataTracker:trkditsd` and select `Add edge → trackField → field:di`.
7. Right-click on `dataTracker:trkditsd` and select `Add edge → trackField → field:tsd`.
8. Right-click again on `dataTracker:trkditsd` and select `Add edge → trackComponent → component:comm`.

Now create the widgets to listen to these data trackers. First we delete the table widget inherited from the previous tutorial and then add time series and scatter plot widgets.

9. Collapse all sub-trees and expand the `userInterface` sub-tree.
10. Right-click on `widget:population table` and select `Delete node`.
11. Create a new `container` as a child of `tab:tab1`.

Containers form a binary tree of `widgets`. Each `container` can contain one or two `widgets` or one or two `containers` or a combination of both with a limit of two. These binary trees must end in a `widget` as a leaf node (i.e. you can't have empty `containers`). The `tab` node also behaves as a `container` and the same rules apply. However, you can have as many `tabs` but of course only one tab is visible at a time in the *ModelRunner* interface.

12. Add a `widget` as a child of `container:cont1`, name it 'srdsdi' and select `TimeseriesWidget1` as its class.
13. Add another `widget` as a child of `container:cont1`, name it 'diversity time since dist' or something similar, and select `ScatterPlotWidget1` as its class.

Set these `widgets` to track the appropriate `dataTrackers` (Figure 48).

14. From `widget:srdsdi`, add a `trackSeries` edge to `dataTracker:trkdi`.
15. From `widget:diversity time since dist`, add a `trackSeries` edge to `dataTracker:trktsddi`.

All that remains now is to arrange the widgets in the *ModelRunner* interface. Between each pair of widgets/containers is a divider which can be moved as required (Figure 49). You can specify the pairs to be divided horizontally or vertically and the order of the pairs (left/right or top/bottom).

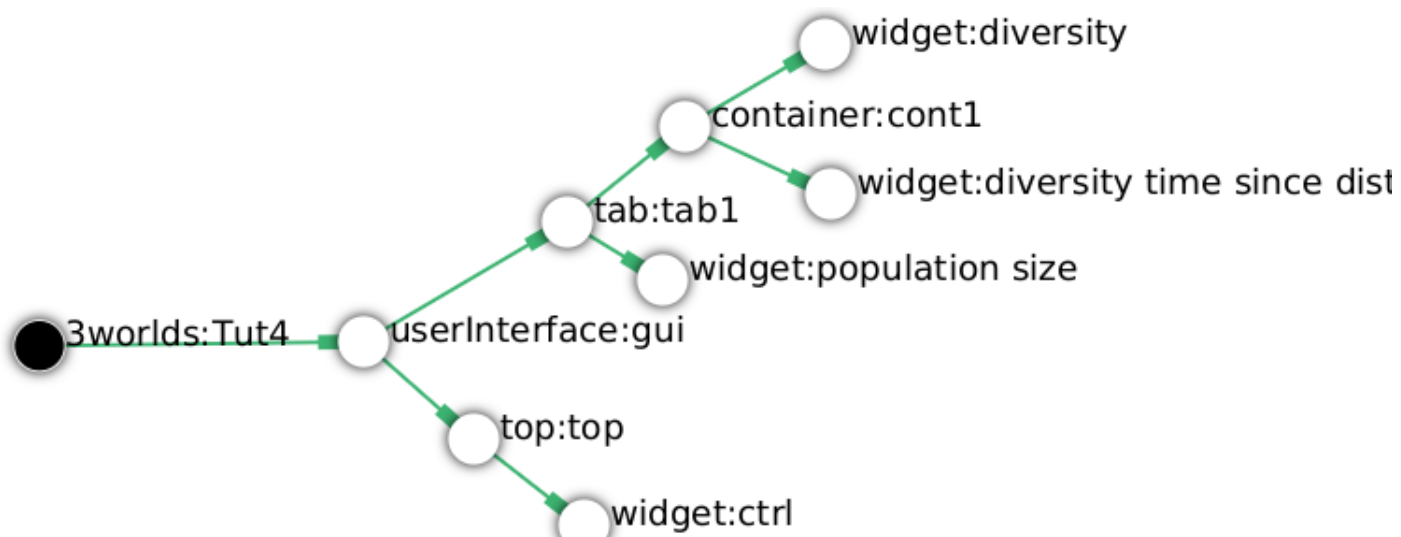


Figure 48. The user-interface sub-tree for Tutorial 4. .

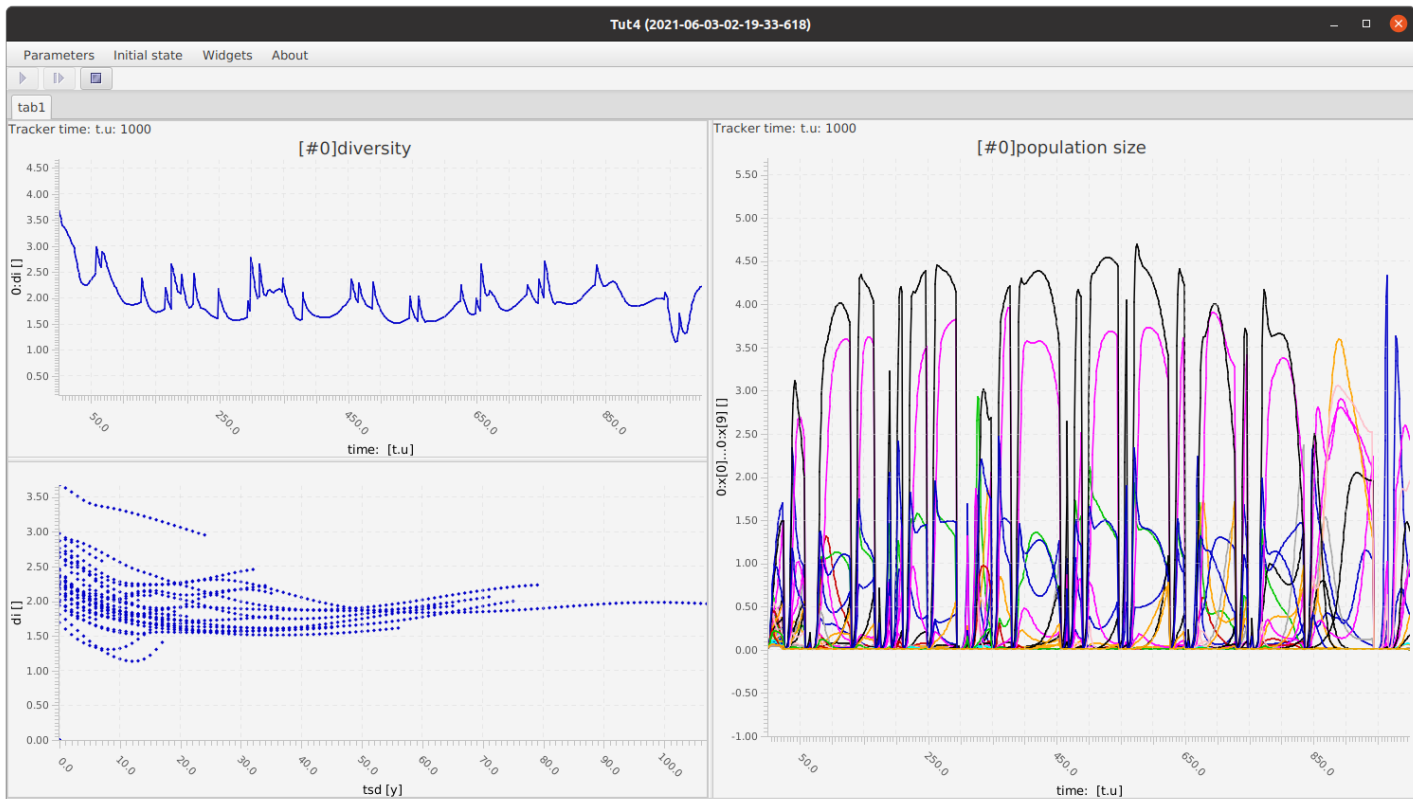


Figure 49. Running Tutorial 4.

The axis order of the xy plot 'diversity time since dist' can be set using the appropriate dialog from the widget menu in *ModelRunner* (Figure 51).

The arrangement of widgets in the tab binary sub-tree is controlled by the `order` and `orientation` properties (Figure 50).

5.4.4. Next

The next tutorial introduces the event timer to simulate episodic systems such as pulse germination or irregular disturbance regimes.

5.5. Tutorial 5: Event driven systems - Using the Event Timer

5.5.1. Introduction

Many processes in nature are episodic and can be triggered when some other process reaches a particular state. For example, in arid environments, organisms may reproduce en masse when a rain event occurs; an epidemic may break out when a population reaches a certain density or; smouldering fire may spread if weather conditions cross some threshold.

The previous tutorials have used a clock timer to drive the simulation dynamics. In essence, a clock timer simply issues regular time events autonomously, at intervals determined by its properties – its time step. An event timer, on the other hand, is not autonomous. It drives its child processes through events posted to it by other processes.

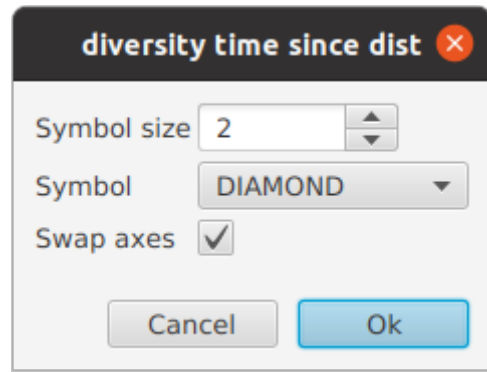


Figure 51. Scatter plot properties editor in ModelRunner.

▼ userInterface	
cont1#order	0
cont1#orientation	vertical
diversity time since dist#order	1
diversity#order	0
population size#order	1
tab1#orientation	horizontal

Figure 50. User-interface properties for Tutorial 4.

In this tutorial we mimic the dynamics of an episodic system with an [Individual-based Model](https://en.wikipedia.org/wiki/Agent-based_model#In_biology) (https://en.wikipedia.org/wiki/Agent-based_model#In_biology) (IBM). We use an event timer to drive pulse creation of entities and a clock timer for their deletion. Here, we refer to the 'entities' as 'items' so as not to be distracted by ecological details.

5.5.2. Specifications

1. Start *ModelMaker* and create a new project from the `SimpleClock1` template and name the project 'Tut5'.

Data definition

Three fields are required: a deletion rate, pulse rate and a ceiling to provide a limit to item creation. These will be global constants (Figure 52).

2. Collapse all sub-trees from the root node and then expand `dataDefinition:dDef`.
3. Delete `record:drv`. There are no drivers needed.
4. Rename `record:csts` to '`glbCnsts`'.
5. Create three `field` nodes as children of `record:glbCnsts`, name them 'ceiling', 'deleteRate' and 'pulseRate' - all of type `Double`.

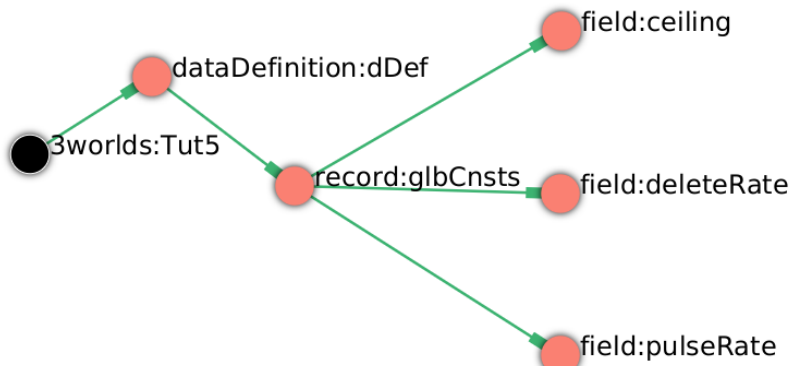


Figure 52. Data definition specifications.

Structure

We need 2 component types for this model: the `system` (which already exists i.e. `system:sys`) and the 'item' component type which we've yet to define (Figure 53). These have a hierarchical relationship: the `system` is an assemblage of 'items'. However, `system:sys1` currently belongs to the *atomic* category. This must be changed to the *assemblage* category to signify this hierarchical structure.

1. Collapse the `dataDefinition` sub-tree and expand the `system` sub-tree.
2. From `system:sys1`, collapse the `dynamics` sub-tree.
3. Select the `system.sys1` node and delete the edge to `category:*atomic*`.
4. From the same node, create a `belongsTo` edge to `category:*assemblage*`.

An 'itemType' will be a component type with a role defined as belonging to the *ephemeral*, *atomic*, *component* and *item* categories. It's *ephemeral* because it can be created and destroyed; it's *atomic* because it has no sub-components; it's a *component* because it is a sub-component of a larger system and it's a *item* because processes that apply to it must reference a category unique to it. All these categories already exist in the `predefined:*categories*` sub-tree except for the 'item' category which we must create.

5. Create a `structure` node as a child of `system.sys1` to begin building the model structure sub-tree.
6. Create a `categorySet` node as a child of `structure:struc1` and name it 'itemSet'.
7. Create a `category` node as a child of `categorySet:itemSet` and name it 'itemCat'.

We now build the item component.

8. Create a `componentType` as a child of `structure:struc1` and name it 'itemType'.
9. Create a `component` as a child of `componentType:itemType` and name it 'item'.

We now define the *role* of `itemType` by specifying the categories it belongs to.

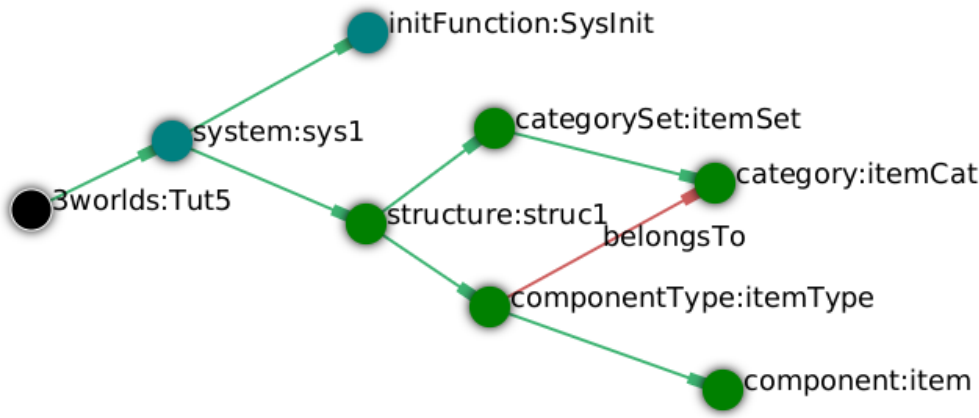


Figure 53. Model structure specifications.

just add the optional property.

10. Create 4 `belongsTo` edges from `componentType:itemType` to `category:*ephemeral*`, `category:*atomic*`, `category:*component*` and `category:itemCat`.

A message now appears suggesting two ways we can create an initial number of items: by the addition of an optional `nInstances` property or loading from a file. For this tutorial we will

11. Right-click on `component:item` and select *optional properties* from the popup menu. Check the `nInstances` box.
12. In the properties editor, set the value of `nInstances` to 100.
13. Before ending this section, rename `initFunction:init1` to 'SysInit'.

Dynamics

The `SimpleClock1` template we started with contains a clock timer (`timer:clock1`) which we can use to call the delete function to remove items at a regular rate. Later we add `eventTimers` to implement the episodic dynamics. It will make the following steps clearer if we rename two nodes first (Figure 54).

1. Collapse all sub-trees from the root node and then expand `system:sys1`.
2. From `system:sys1`, collapse all sub-trees and then expand `dynamics:sim` and re-apply the layout [Alt+L].
3. Rename `timer:clock1` to `timer:t3Clock`.
4. Rename `process:p1` to `process:p3`.

We need a `deleteDecision` function associated with this process rather than the current `ChangeState` function.

5. Delete node `function:F1`.
6. Create a `function` node as a child of `process:p3`, name it 'DeleteItem' and select `deleteDecision` as its type.

`process:p3` is currently applied to the `arena` category. This edge was present from the template we started with. We must reassign this to apply `p3` to `category:itemCat`.

7. Select `process:p3` and delete the edge to the `arena` category and create an `appliesTo` edge to `category:itemCat`.

We now need an event timer to drive 'item' creation events.

8. Create a `timer` node as a child of `timeline:tmLn`, name it 't2Event' and select `EventTimer` as its type.
9. Create a `process` node as child of `timer:t2Event` and name it 'p2'.
10. Add an `appliesTo` edge from `process:p2` to `category:itemCat`.
11. Create a `function` node as child of `process:p2`, name it 'Nitems' and select `CreateOtherDecision` as its type.

This function returns a floating point number that will be the number of items to create for each existing item. The fractional part is rounded probabilistically to give a discrete value.

A new message has now appeared asking that `timer:t2Event` have an edge to a `function` or `initFunction`. Every event timer must have at least one function that feeds events to it (specified by a `fedBy` edge). Therefore, we need a third function to generate these events.

12. Create a `timer` node as a child of `timeline:tmLn`, name it 't1Event' and select `EventTimer` as its type.
13. Create a `process` node as child of `timer:t1Event` and name it 'p1'.
14. Create a `function` node as child of `process:p1`, name it 'Pulse' and select `ChangeState` as its type.
15. Add an `appliesTo` edge from `process:p1` to `category:*arena*`.

This function operates at a global level (the arena) not at the level of individual items as does `function:Nitems`. Its job is to send an event to `timer:t2Event` that will cause `function:Nitems` to be called in a loop on all existing items. The function `function:Pulse` will also post a future event to call itself so the system will run indefinitely. However, since `function:Pulse` is also driven by an `eventTimer` we will get into an infinite regress unless we somehow feed at least one event to `timer:t1Event`. To do this we just add a `fedBy` edge to an `initFunction`.

To do all this, we must add 3 `fedBy` edges: one from `function:Pulse` to drive item creation; one from `function:Pulse` to `timer:t1Event` to drive pulse creation; and a third from `initFunction:SysInit` to start the chain of events.

16. Add an `fedBy` edge from `timer:t1Event` to `initFunction:SysInit`. `SysInit` will post a first event to begin the process.
17. Add another `fedBy` edge from `timer:t1Event` to `function:Pulse`. `Pulse` will continue calls to itself at some random time.
18. Add a third `fedBy` edge this time from `timer:t2Event` to `function:Pulse`. `Pulse` will also generate the item creation events.

There can be any number of event timers in a specification and each timer can be driven by any number of functions posting events at any time in 'present' or future. This design allows specifying very complex causal chains.

The template we began with includes a `stoppingCondition`. This is not needed as we can allow the simulation continue indefinitely for now.

19. Select `stoppingCondition:stCd1` (a child of `dynamics:sim1`) and delete it.

The next step is to record the number of 'items' at the end of the time step for display. A state variable already exists for this purpose in the `predefined:*categories*` sub-tree called `count`. To make use of this we must add another `process` with a `dataTracker` child.

20. Create a `process` as a child of `timer:t3clock` and name it 'p4'.
21. Add an `appliesTo` edge from `process:p4` to `category:*assemblage*`. This category is associated with `record:AVPopualation` that contains the `count` field.
22. Create a `dataTracker` as a child of `process:p4`.
23. From the `dataTracker`, add two edges: one a `trackField` edge to `field:count` and the second a `trackComponent` edge to `system:sys`.

Now add `dependsOn` edges between some of the processes to set their order of execution.

24. Right-click on `process:p4` and create a `dependsOn` edge to `process:p3`. This ensures `count` is recorded as the last step.
25. Right-click on `process:p3` and create a `dependsOn` edge to `process:p2`. This ensures `function:Delete` always

occurs after `function:Nitems` should they occur simultaneously.

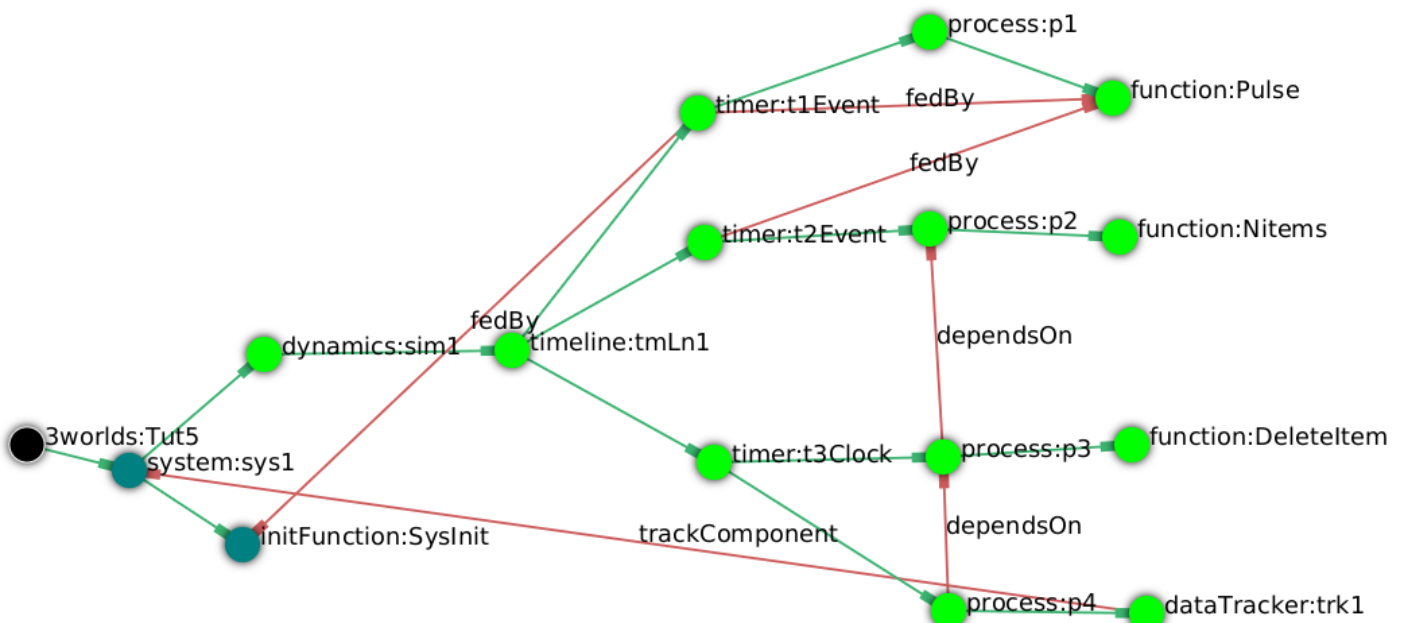


Figure 54. Model dynamics specification.

Coding

We now add the program code to all `functionSnippet` properties for all the functions we have created.

1. Ensure that the `system` sub-tree is fully expanded. To do this *Collapse sub-tree > All* from `system:sys` node and then select *Expand sub-tree > All* from the same node.
2. In the `Properties` editor, select the 'By name' button in the top-left corner and type 'func' in the search bar. All 4 `functionSnippet` properties should now be listed.
3. Edit `SysInit#functionSnippet`, and enter the following text:

```
// Set values of global constants
focalCnt.deleteRate = 0.01;
focalCnt.pulseRate = 0.002;
focalCnt.ceiling = 5000.0;
// post event to the pulse time to kick start it. After this it posts events to itself.
t1Event.postTimeEvent(1);
```

4. Edit `Pulse#functionSnippet` and enter:

```
// post a 'item' creation event for this time
t2Event.postTimeEvent(t);
// schedule a call to this method at some future time
t1Event.postTimeEvent(t + random.nextInt((int)(1.0/pulseRate)) + 1);
```

5. Edit `Nitems#functionSnippet` and enter:

```
// limit the recruitment rate of items to the ceiling
double rate = (ceiling-count)/(double)count;
return rate;
```

6. Edit `Delete#functionSnippet` and enter:

```
// return the deleteRate probabilistically converted to true or false
return decider.decide(deleteRate);
```

User Interface

The template we are using has only provided a control widget (`widget:ctrl`) to run the model. To see some output we will add a widget to display a time series of `count` (Figure 55).

1. Create a `tab` as a child of `userInterface/gui`.
2. Create a `widget` as a child of `tab:tab1` and name it `Number of items` and select `TimeSeriesWidget1` as its type.
3. Add a `trackSeries` edge from this widget to `dataTracker:trk1`.
4. Run the simulator (Deploy [Alt+D]).

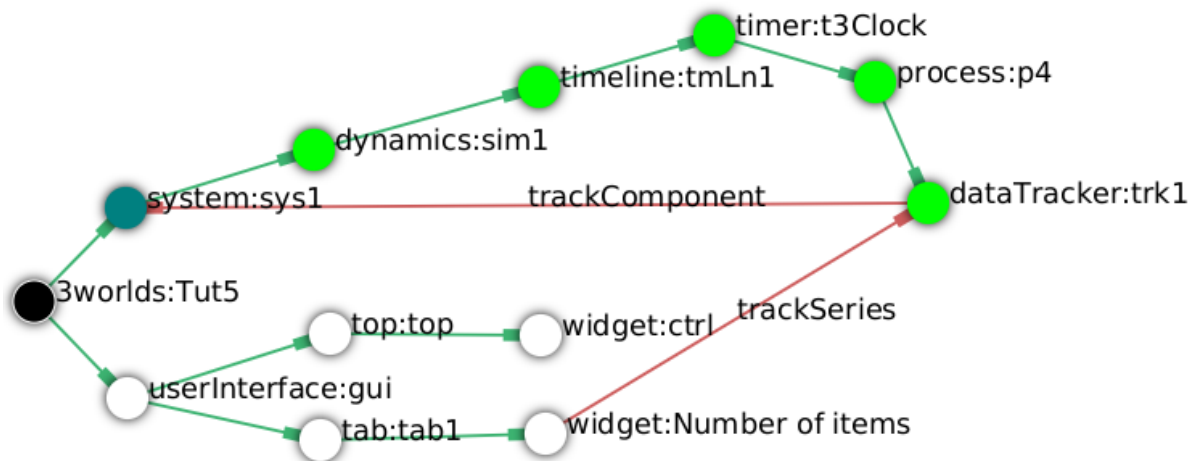


Figure 55. User interface specifications.

The simulator runs indefinitely (assuming you have deleted the `stoppingCondition` node) with exponentially declining numbers and a creation pulses up to 500 time steps apart (Figure 56).

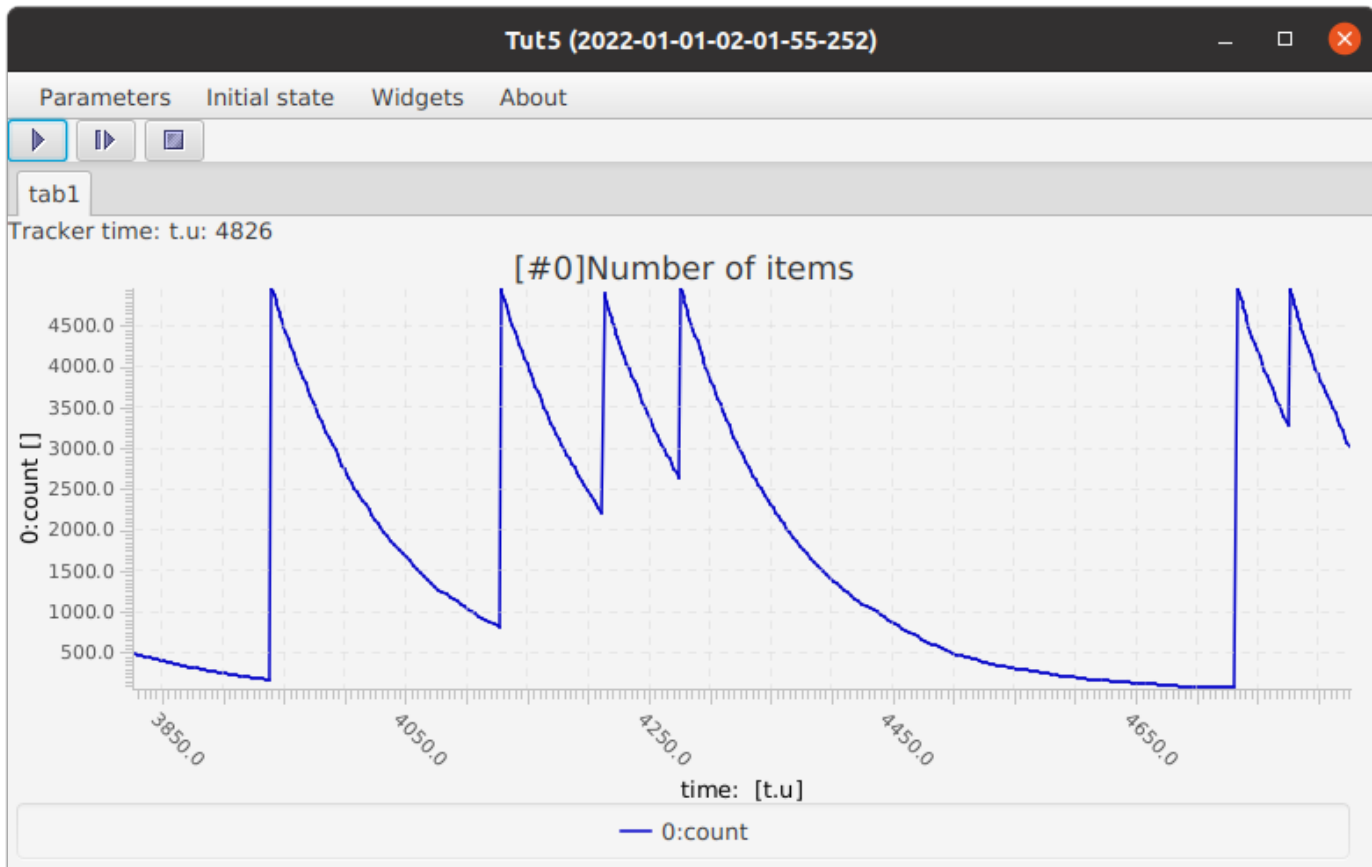


Figure 56. Running Tutorial 5.

5.5.3. Next

The next tutorial introduces spatial models by modifying this model to display 'items' in a 2-dimensional continuous surface.

5.6. Tutorial 6: Event driven systems - Using the Event Timer in a spatial model

5.6.1. Introduction

This tutorial introduces 2-dimensional space by converting Tutorial 5 to a spatial model.

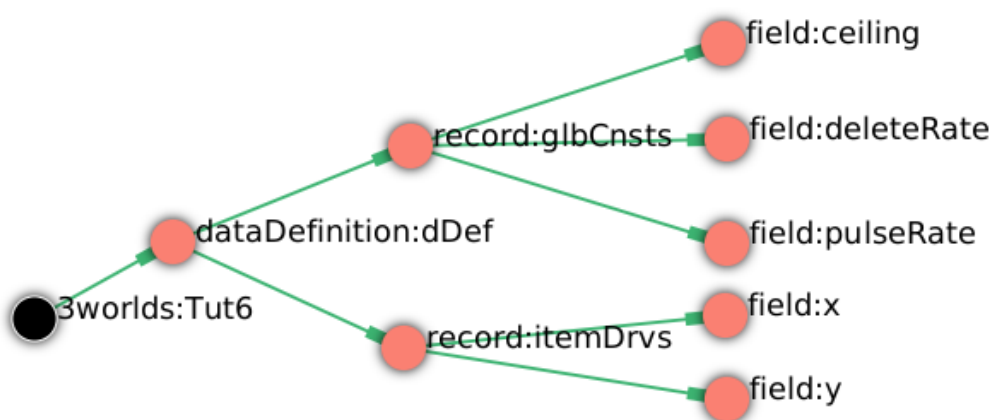
5.6.2. Specifications

1. Start *ModelMaker* and create a new project from *Tutorials* → *Event timer 1* and name it 'Tut6'.

Data definition

We must first specify x,y coordinates for 'items' - the modelled entity (Figure 57).

1. Create a record as a child of *dataDefinition:dDef* and name it 'itemDrvs'.
2. Add 2 fields to this record named 'x' and 'y' of type Double.



A message appears indicating this new record must be associated with a category. This is done in the next section.

Figure 57. Data definition specifications.

Structure

In the **structure** sub-tree we specify the type of space to use and associate it, and the `category:itemCat`, with the `xy` coordinates specified above (Figure 58). We can also set the initial values of the coordinates at start up time.

Associate the item drivers with the item category.

1. Right-click on

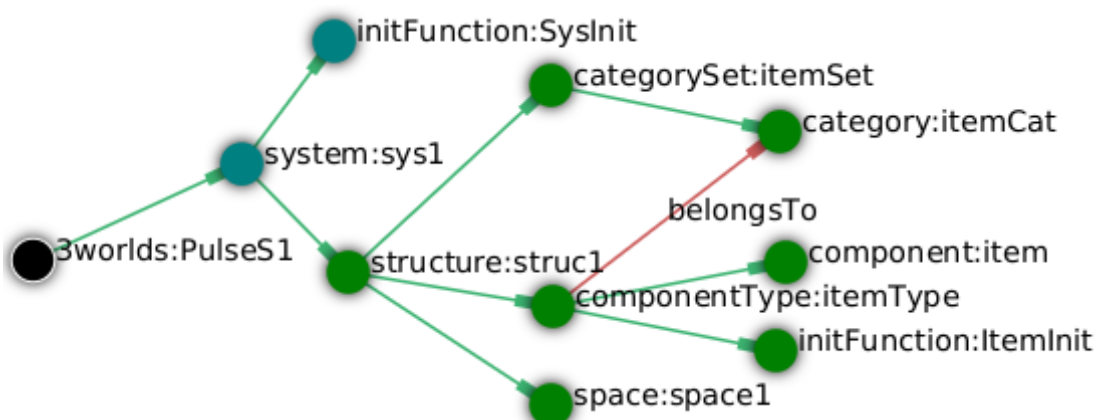


Figure 58. Model structure specifications.

`category:itemCat` and create an edge to the `xy` coordinates by selecting `New edge → drivers → record:itemDrvs`.

Create the space and set its properties.

2. Create a `space` node as a child of `structure:struc1` and select `ContinuousFlatSurface` as its type.
3. Right-click on `space:space` and create a new edge to `coordinate → field:x` and name it 'x1'.
4. Repeat the above but select `coordinate → field:y` and name it 'y1'.
5. In the properties editor for `space:space1`, set `y1#rank` to 1 and `space1#precision` to 0.01.

Set the dimensions of the space to 1,000 by 1,000 meters.

6. Set the properties of `space1#x-limits` and `space1#y-limits` to 0 to 1000 inclusive.
7. Add the optional property `units` to `space:space1` and set its value to 'm' (meters). This is just for documentation.

Create a function to set initial values (the `xy` coordinates) for the 100 items (a property of `component:item`) at start up.

8. Create an `initFunction` as a child of `componentType:itemType` and name it 'ItemInit'.

Dynamics

When new items are created by `process:Nitems` we need a function to set `xy` coordinates of the newly created 'items'. This is a 'consequence' function. Consequence functions are children of other functions rather than a process (Figure 59).

1. Create a `function` as a child of `function:Nitems`, name it 'MakeItem' and select `setOtherInitialState` as its type.

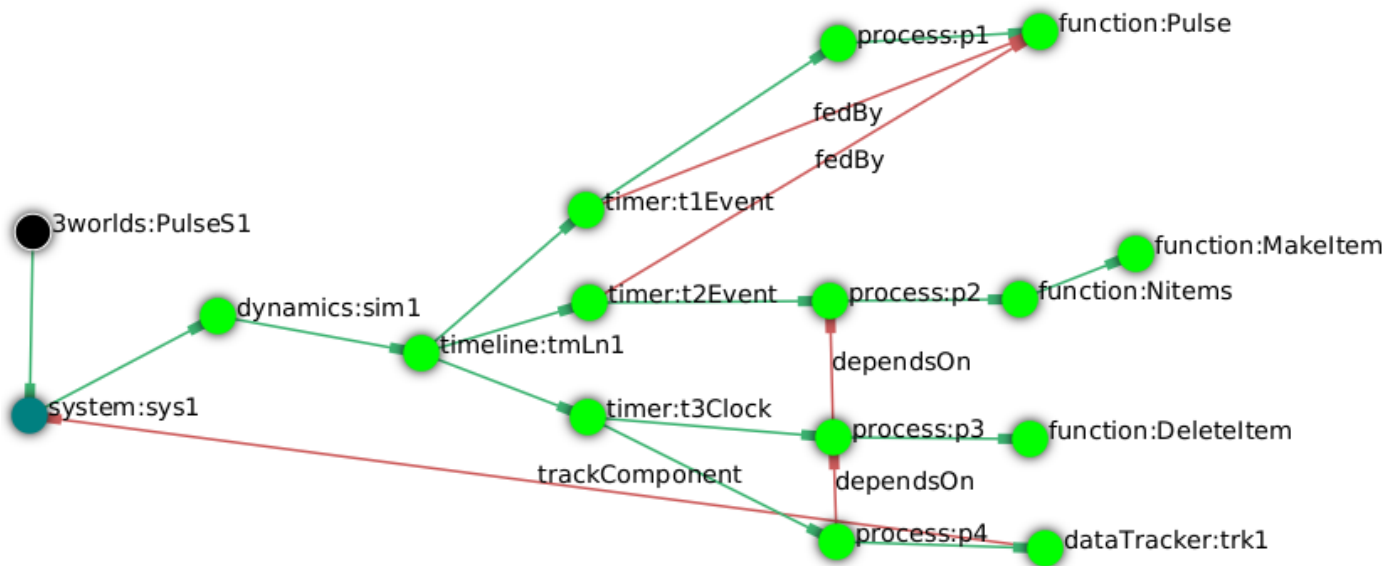


Figure 59. Model dynamics specifications.

Coding

We now add some Java code to set values for the *xy* coordinates. We will just set random values for the state up items and set values in newly created items to be clumped near their parent item.

1. Add the following code to the `ItemInit#functionSnippet` property:

```
// place dots in random locations (assuming these are the dimensions of the map)
focalDrv.x = random.nextDouble() * 1000;
focalDrv.y = random.nextDouble() * 1000;
```

2. Add the following code to the `MakeItem#functionSnippet` property:

```
// place dot randomly near 'parent' dot.
double distance = -50.0 * Math.log(1 - random.nextDouble());
double direction = random.nextDouble() * (2 * Math.PI);
otherDrv.x = x + distance * Math.cos(direction);
otherDrv.y = y + distance * Math.sin(direction);
```

User Interface

Add a widget to view the items in space (Figure 60).

1. Create a `widget` as a child to `tab:tab`, name it 'Item landscape' and select `SpaceWidget1` as its type.
2. Add a `trackSpace` edge from `widget:Item landscape` to `space:space` (`trackSpace → space:space1`).

Arrange the two widget horizontally (the default) - time series first.

3. Click on `widget:Item landscape` and in the *Selected properties* editor, set `Item landscape#order` to 1.

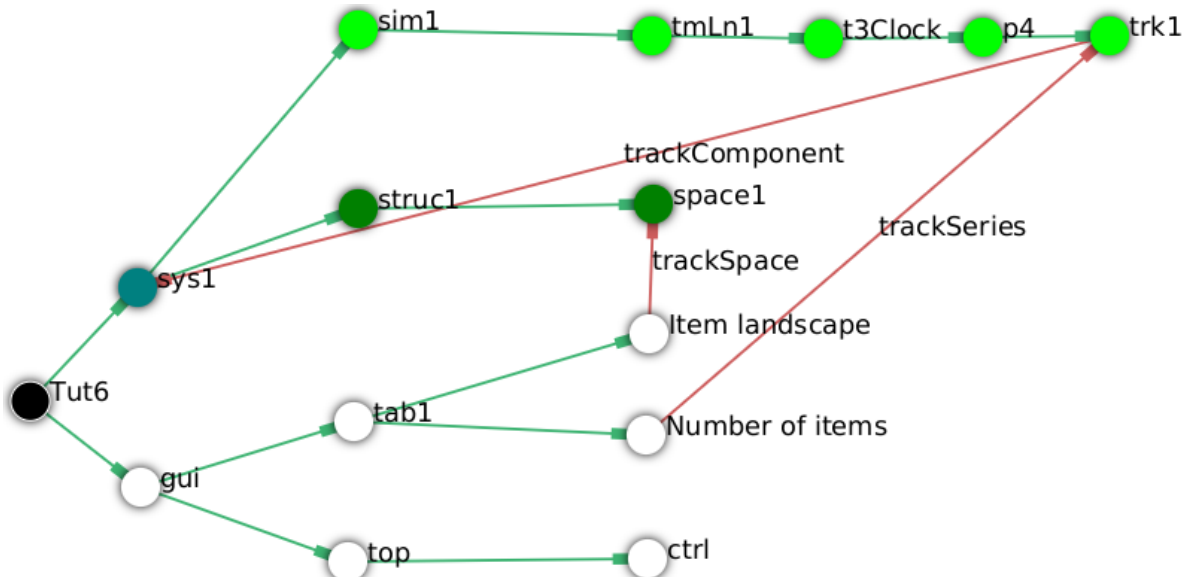


Figure 60. Simulator's user-interface specifications showing cross-links to nodes in the structure and dynamics sub-trees.

4. Deploy the simulation [Alt+D] (Figure 61).

Information about `SpaceWidget1` can be found at Section 3.3.7.5.

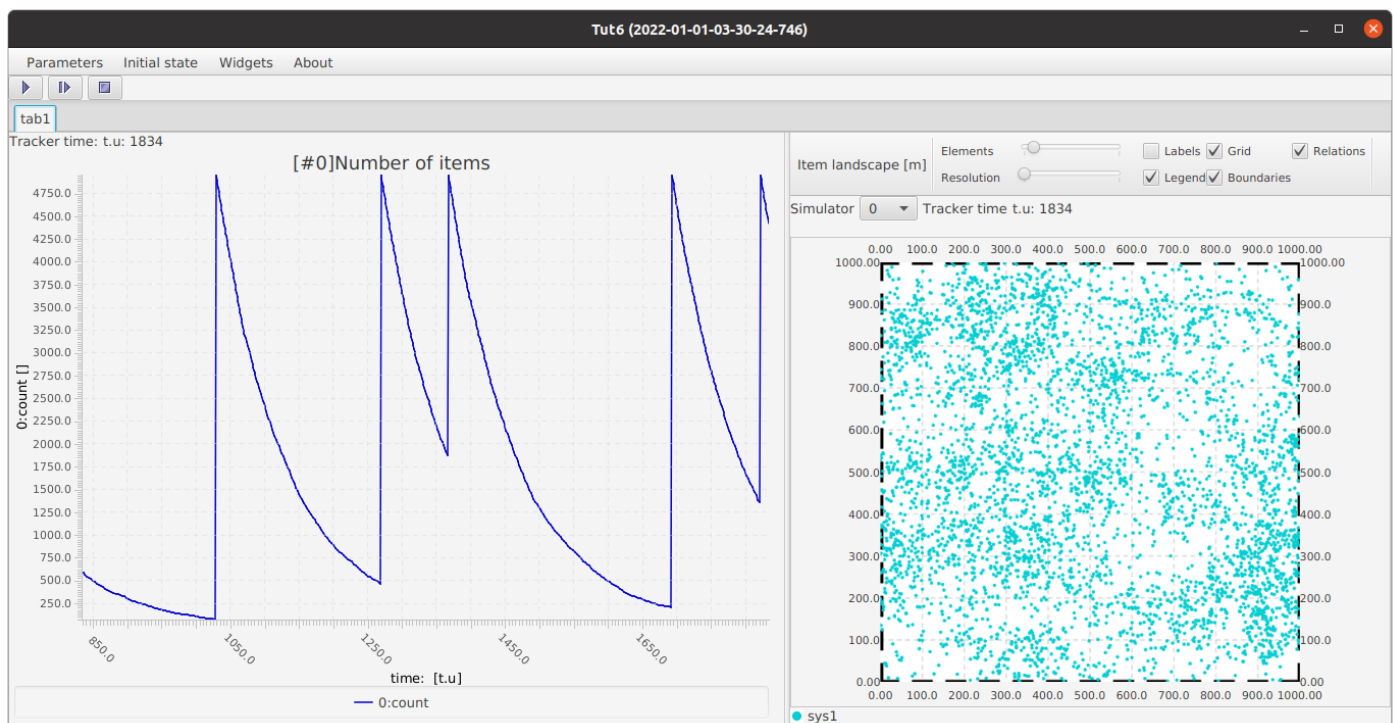


Figure 61. Running Tutorial 6.

5.6.3. Next

The next tutorial will explain how to implement and use random numbers in models that require stochasticity.

5.7. Tutorial 7: Random number generators

5.7.1. Introduction

This tutorial introduces the use of [random number generators](https://en.wikipedia.org/wiki/Random_number_generation) (https://en.wikipedia.org/wiki/Random_number_generation) (RNG).

3Worlds provides for:

- A single default RNG
- Any number of independent RNGs;
- Three different RNG algorithms;
- Management for seeding the RNGs; and,
- Assignment of RNGs to particular `Functions`.

A single default RNG

In many cases, management of random number generation is not required. For this reason, a single default RNG (PCG32) is available to all functions and can be used by the modeller in their code. Which RNG is available and how it is managed is defined in the model's specifications using *ModelMaker*. These choices are transparent to the model so no coding effort is required by the modeller to utilise the various features.

Any number of independent RNGs

You can create any number of independent RNGs and apply them individually or to groups of `Functions`. For example, you could assign an RNG to all functions that apply to reproduction and another to all functions that concern landscape dynamics.

Three different RNG algorithms

Three RNG algorithms are available: the standard Java RNG, Pcg32 ([Permuted Congruential Generator](https://www.pcg-random.org/pdf/hmc-cs-2014-0905.pdf) (<https://www.pcg-random.org/pdf/hmc-cs-2014-0905.pdf>)) and XSRandom ([XorShift](http://demesos.blogspot.com/2011/09/replacing-java-random-generator.html) (<http://demesos.blogspot.com/2011/09/replacing-java-random-generator.html>)). Two important properties of RNGs are their quality (the degree to which they cannot be predicted better than by a random chance) and the computer resources required to produce them (memory and speed). Both Pcg32 and XSRandom are much faster (50% and 75% respectively) and of higher quality than the native Java RNG. Pcg32 is of higher quality than XSRandom (the default).

Management for seeding the RNGs

For the purpose of debugging and experimental design it is often necessary to manage the RNG seeds.

There are two ways to set the source of the RNG seed:

- **TABLE:** From [a table of 1,000 independently generated random number seeds](https://www.random.org/) (<https://www.random.org/>); and,
- **RANDOM:** From a randomly generated seed using current time to the nano second.

In addition, the seed can be set at the time of model instantiation (**NEVER**) or upon every reset of the simulation (**ONMODELSTART**). Using the **RANDOM** option means there is a new seed created at every instantiation. This option is here just so you don't need to delete all the random number generators from the specifications - it effectively switches off reproducible RNG streams.

In this tutorial, we build a model with two functions to generate random numbers from two independent streams.

5.7.2. Specifications

1. Create a new project starting from the `SimpleClock` template.

Data definition

1. Delete the node `record:cnsts`
2. Create two `field` nodes as children of `record:drv`s accept their default names ('fld1', 'fld2') and use the default type of `Double`.
3. Create two `rng` nodes as children of `dataDefinition:dDef` and accept their default names ('gen1', 'gen2').

Structure

1. Delete node `initFunction:init1`: it's no needed for this tutorial.

Dynamics

1. Create a `Function` as child of `process:p1` accept the default name ('F2") and select `ChangeState` as the type.
2. Add the following code to `F1#functionSnippet` property: `'focalDrv.fld1 = random.nextDouble();'`.
3. Add the following code to `F2#functionSnippet` property: `'focalDrv.fld2 = random.nextDouble();'`.
4. Create a `useRNG` edge from `function:F1` to `rng:gen1`.
5. Create a `useRNG` edge from `function:F2` to `rng:gen2`.

Now the value of 'fld1' is set by the random number generator 'gen1' and the value of 'fld2' by 'gen2'.

6. Create a `dataTracker` as a child of `process:p1` and select `DataTrackerD0` as its type.
7. Create two `trackField` edges from `dataTracker:trk1` to `field:fld1` and `field:fld2` respectively.
8. Create a `trackComponent` edge from `dataTracker:trk1` to `system.sys1`.

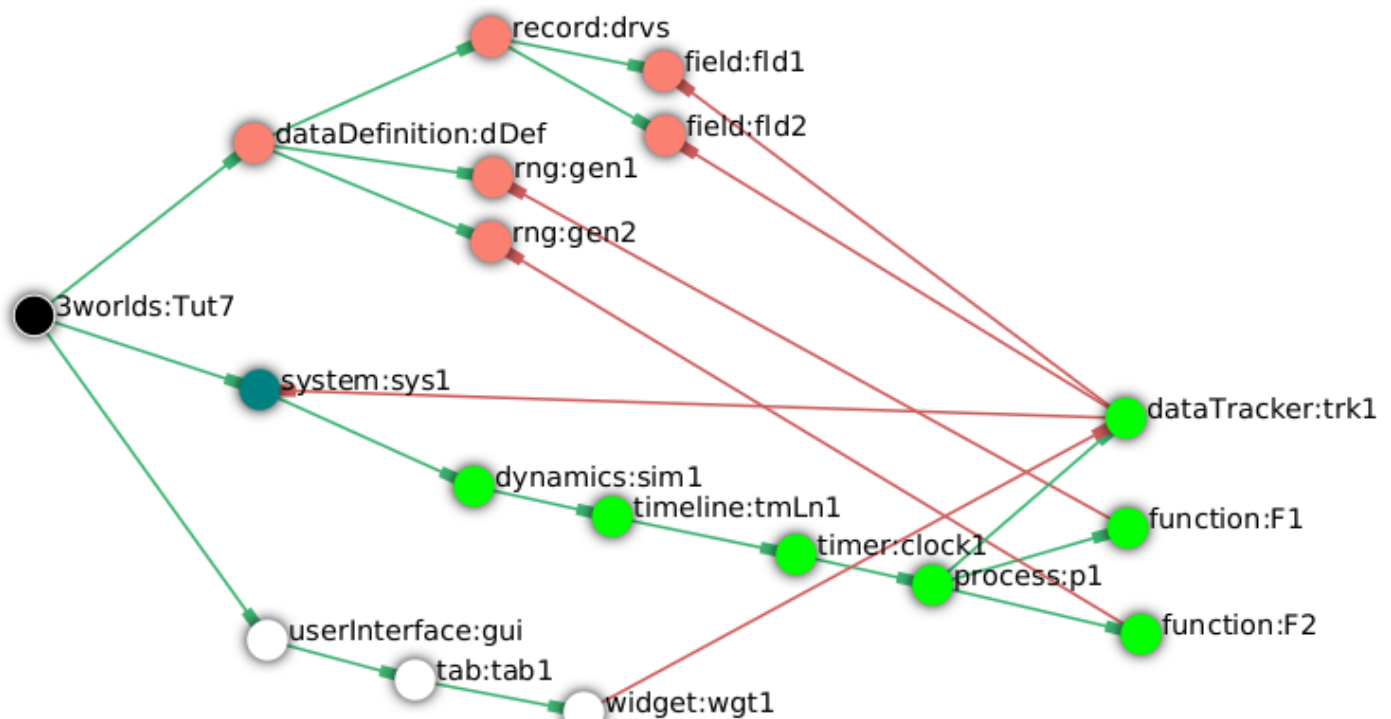


Figure 62. Essential elements of the configuration for tutorial 7.

User interface

1. Create a `tab` node as a child of `userInterface:gui`.
2. Create a `widget` node as a child of `tab:tab1` and select `TimeseriesWidget1` as its type.
3. Create a `trackSeries` edge from `widget:wgt1` to `dataTracker:trk1`.

Once the configuration is finished (Figure 62), deploy the simulator.

5.7.3. Simulation

When the simulator is run, two independent random number series (Figure 64) are produced. If this is not the case, check that the code snippets (steps 2 & 3 in *Dynamics* above) are correct and that `function:F1` sets the value of `fld1` and `function:F2` sets the value of `fld2`.

1. Re-run the simulator a few times to ensure a different series is produced each time

We will now hold the random number series constant. To see all the relevant RNG properties:

2. Select the *Properties* tab in *ModelMaker* and click the *by name* button (top left).
3. Enter 'gen' in the search field. Now only properties of nodes `gen1` and `gen2` are displayed (Figure 63).
4. Set properties `gen1#seedSource` and `gen2#seedSource` to **TABLE** and re-run the simulator.

Both series are now identical as they are set by the same seed (index 0 of the table of seeds). Of course if you change the RNG algorithm to one of the other algorithms, different streams will result.

Properties		Selected Properties
<input type="button"/> <input type="button"/>		x
gen		
gen1#algorithm	PCG32	▼
gen1#resetTime	NEVER	▼
gen1#seedSource	RANDOM	▼
gen1#tableIndex	0	
gen2#algorithm	PCG32	▼
gen2#resetTime	NEVER	▼
gen2#seedSource	RANDOM	▼
gen2#tableIndex	0	

Figure 63. Displaying just the properties of the random number generators.

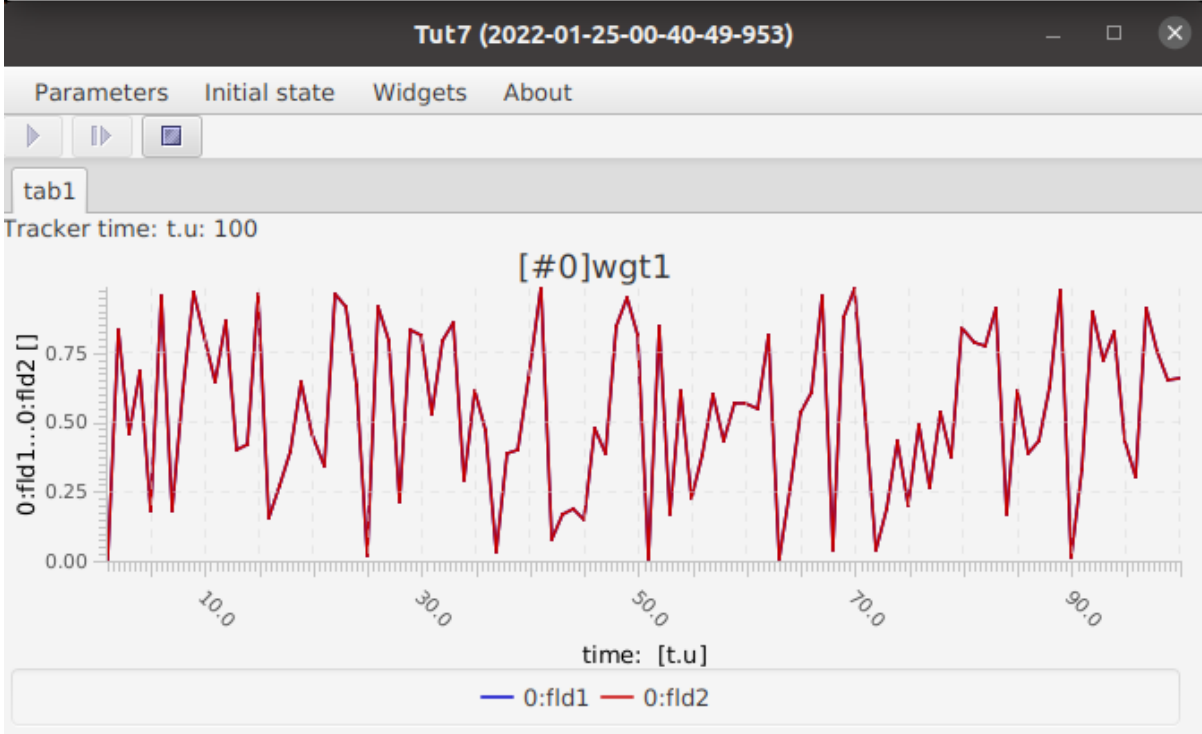
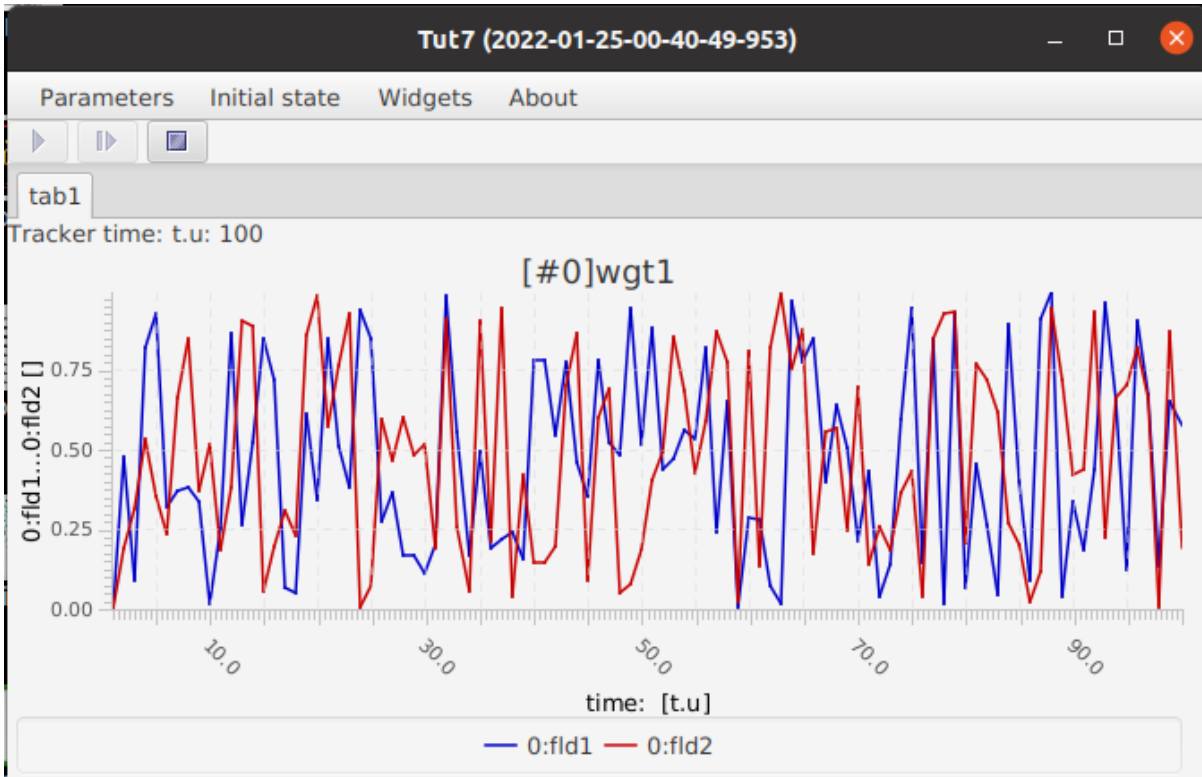


Figure 64. Independent and dependent random number streams.

5.7.4. Next

This next and last tutorial covers running simulation experiments.

5.8. Tutorial 8: Running an experiment

5.8.1. Introduction

This tutorial introduces the current experiment design system (22/06/2022). This system has a major limitation in that it can only work with arena (i.e. global) constants. We will update this tutorial when OpenMOLE is integrated with 3Worlds, something we are currently working on.

NB: R must be installed to run this tutorial.

We demonstrate the current system using GDDM (Genetics, Demography and Disturbance Model). GDDM is an individual-based spatial model of a population of animals in an environment experiencing recurrent disturbance. The disturbance return interval is short (about every two years) because the study is intended for the North Australian savannah. Animals mate, give birth, disperse and die on an annual time step. Their longevity is about 12 years.

One of the aims of the study is to examine the population and genetic consequences of informed dispersal. In the present case, informed dispersal means tending to select habitat with the maximum space considering the carrying capacity and current population at this location.

We perform a simple ANOVA experiment to examine the variance explained in Fst and total population by just two parameters (constants): dispersal distance (long/short) and dispersal method (informed/uninformed). Most of the processes in this model depend on random number generation. Therefore, we will run five replicates.

5.8.2. Specifications

1. Create a new project by selecting `new → model library → GDDM` and name it 'Tut8'.
2. Run (Deploy) the model and adjust the widget settings (widget menu) to suit to get a feel for what's going on.

The model runs for 100 years and displays many variables such as Fst and population size. The habitat pattern changes every year with approximately half being 'good' habitat and half 'bad'. 'Good' habitat has a carrying capacity of 3 animals per ha. and bad 1.5.

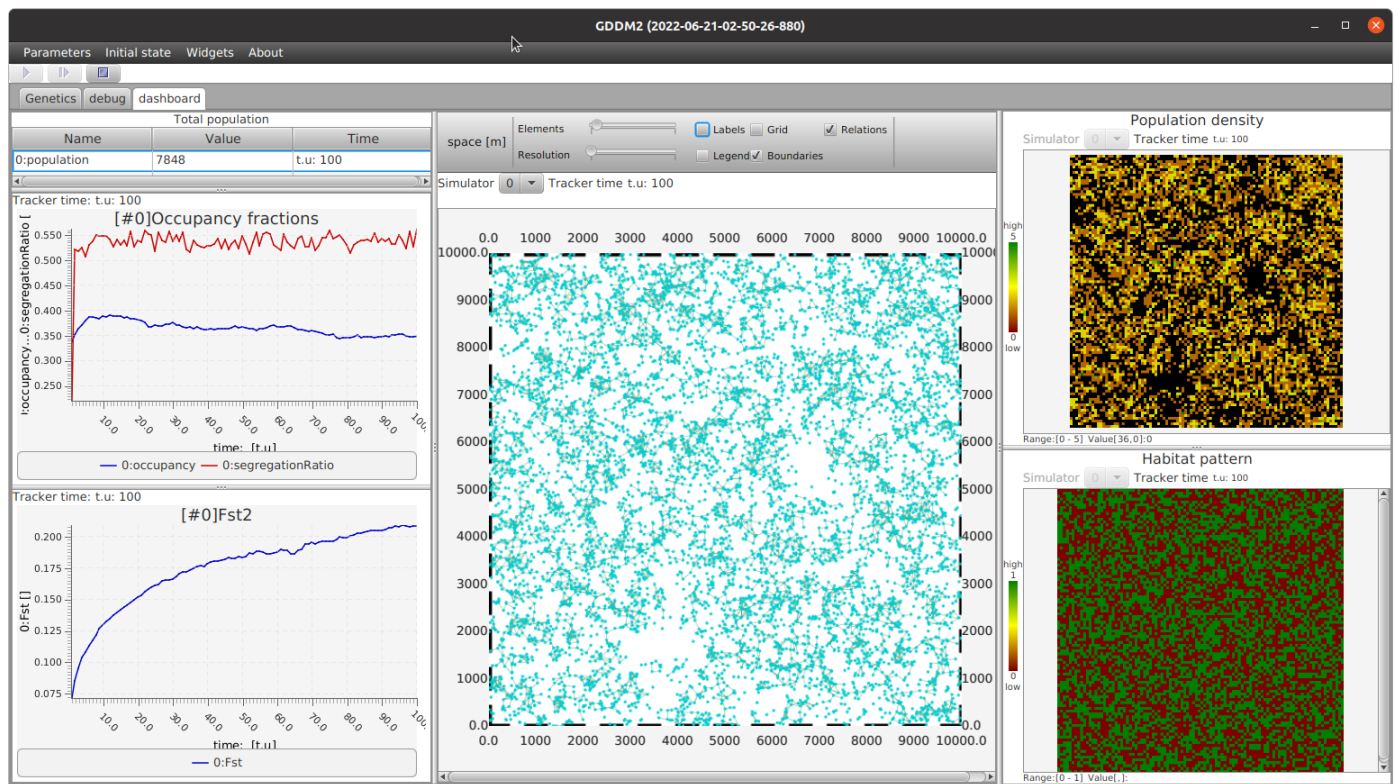


Figure 65. A view of GDDM running in ModelRunner.

Return to *ModelMaker* and arrange the graph as follows:

3. Collapse `system.sys1`.
4. Collapse all sub-trees from `DataDefintion:dDef` and then expand `record:glbCnt`.

User interface

To reduce the running time of the experiment, we can replace the graphic displays and just have text output to file.

5. Select the root node (`3worlds:Tut8`) and delete the entire `UserInterface/gui` sub-tree.

6. Add a `Headless` node as a child of `UserInterface:gui`.

Widgets that are children of `Headless` have no visible presence except for console output. Note: you won't see this output unless you are running `ModelMaker` from a terminal window.

7. Add a `widget` as a child of `headless:h1`, name it 'ctrl' and select `HLControlWidget1` as its sub-class.

This control widget can be used even if you have visible widgets. It begins the simulation without user-intervention.

8. Add another `widget` as a child of `headless:h1` name it 'progress' and select `HLProgressWidget1` as its sub-class.

9. Add a `trackTime` edge from `widget:progress` to `dynamics:sim1`.

This progress widget will produce progress information to the console as the experiment proceeds.

9. Add two more widgets as a children of `headless:h1`, name them 'fstWriter' and 'popWriter' respectively and select `HLTimeSeriesAnalysisWidget1` as their type.

10. Add a `trackSeries` edge from `widget:fstWriter` to `dataTracker:trkFst`.

11. Add another `trackSeries` edge, this time from `widget:popWriter` to `dataTracker:trkMetaPop`.

We can also add headless widgets to save spatial images of various matrices to give some visual feedback of these data for each of the factor combinations (i.e. dispersal distance and dispersal method). To do this:

12. Add two more widgets as a children of `headless:h1`, name them 'HsMapWriter' and 'popMapWriter' respectively and select `HLMATRIXWidget1` as their type.

13. Add a `trackSeries` edge from `widget:HsMapWriter` to `dataTracker:trkHeterozygosity`.

14. Add another `trackSeries` edge, this time from `widget:popMapWriter` to `dataTracker:trkPopulationLayer`.

The newly added `HLMATRIXWidget1` widgets have some properties that should be set.

For the `HsMapWriter` set:

15. `imageMagnify` to 6.

16. `palette` to 'OrangeMauveBlue'.

17. `sampleTimes` to 100 to write an image at the last (100th) time step.

18. `zRange` to ']0.4,0.89[

For the `popMapWriter` set:

19. `imageMagnify` to 2.

20. `palette` to 'BrownYellowGreen'.

21. `sampleTimes` to 100.

22. `zRange` to ']1.0,4.0[

23. Now add the optional properties to these two widgets (`missingValueColour` and `missingValueMethod`).

24. For both widgets set `missingValueColour` to 'BLACK' and `missingValueMethod` to 'LT_MIN'. This means all values in the matrix less than the minimum range will be drawn as black.

Experiment

We now define some treatments of `fields` that are children of `record:glbCnt`.

1. Add a `treatment` node as a child of `experiment:exp`.

2. Add a `treats` edge from `treatment:trt` to `field:isDispersalInformed` and name it 'dm' (dispersal method).

This field (`isDispersalInformed`) is a boolean constant so the only treatments are 'true' and 'false'.

3. Click on the `treatment:trt` node and, in the selected properties editor, edit the `dm#values` property by adding 'true' and 'false' on two separate lines.

4. Edit the `dm#valueName` property and enter 'inf' and 'uninf', again on two separate line. These are the names by which the treatment is described in outputs.

5. Add another `treats` edge from `treatment:trt` to `field:isDispersalShort` and name it 'dd' (dispersal distance).

This field (`isDispersalShort`) is also a boolean constant.

6. Click on the `treatment:trt` node and, in the selected properties editor, edit the `dd#values` property by adding 'true' and 'false' on two separate lines.

7. Edit the `dd#valueName` property and enter 'short' and 'long', again, on two separate line.

To run an ANOVA:

8. Select the `design:dsgn` node, and in the selected properties editor set `design:dsgn` to `crossFactorial`.

A message now appears indicating that treatments must have a designated rank order for this experiment design.

9. Select the `treatment:trt` node again and in the selected properties editor set `dd#rank` to 1 and `dm#rank` to 2.

This model employs random number generation in many of its processes so replicates must be used.

10. Right-click on the `experiment:exp` node, select `Optional properties...`, check `experiment:exp#nReplicates` and click 'ok'.

11. Select `experiment:exp` and in the selected properties editor set `exp#nReplicates` to 5.

This experiment will create 5 x 2 x 2 simulators (20) running in parallel.

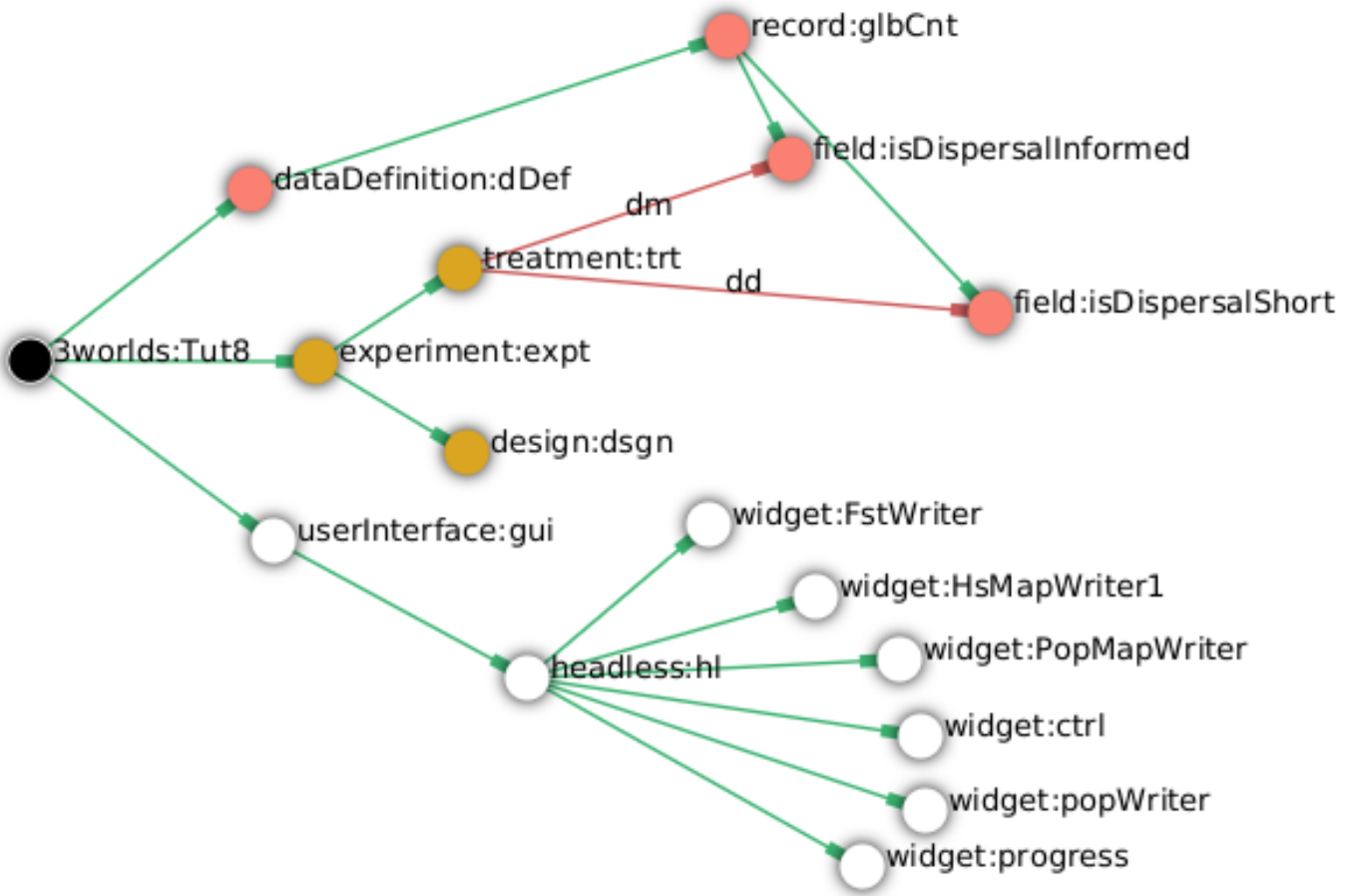


Figure 66. Relevant parts of the configuration graph ready for running.

5.8.3. Simulation

1. Save changes (Ctrl-S) and click the deploy button (Alt+D).

The date and experiment design type are now written to the console followed by a message from each simulator as it is initialised. The headless controller then starts the simulators and a running message is produced from each. The number of simulators running in parallel depends on the number of cores on your machine. The experiment takes about 5 minutes on a 16 core machine. The ODD documentation is generated automatically at the end of the experiment.

```

Running... [Project: Tut8 (2022-06-22-01-50-25-964);
Date: 2022/06/22 01:57:08:903]
Initialising... [Simulators: 20]
Initialising [done]
Starting... [experiment:expt(crossFactorial)]
[1/20] ready...
[2/20] ready...
[3/20] ready...
[4/20] ready...
[5/20] ready...
[6/20] ready...
[7/20] ready...
[8/20] ready...
[9/20] ready...
[10/20] ready...
[11/20] ready...
[12/20] ready...
[13/20] ready...
[14/20] ready...
[15/20] ready...
[16/20] ready...
[17/20] ready...
[18/20] ready...
[19/20] ready...
[20/20] ready...
waiting          01:57:14 22/06/2022
[6]   running...    01:57:14 22/06/2022
[7]   running...    01:57:14 22/06/2022
[4]   running...    01:57:14 22/06/2022
[8]   running...    01:57:14 22/06/2022
[5]   running...    01:57:14 22/06/2022
[2]   running...    01:57:14 22/06/2022
[9]   running...    01:57:14 22/06/2022
[1]   running...    01:57:14 22/06/2022
[12]  running...    01:57:14 22/06/2022
[11]  running...    01:57:14 22/06/2022
[10]  running...    01:57:14 22/06/2022
[3]   running...    01:57:14 22/06/2022
[13]  running...    01:58:39 22/06/2022
[14]  running...    01:58:43 22/06/2022
[15]  running...    01:58:47 22/06/2022
[16]  running...    01:59:37 22/06/2022
[17]  running...    01:59:39 22/06/2022
[18]  running...    01:59:44 22/06/2022
[19]  running...    02:00:05 22/06/2022
[20]  running...    02:00:44 22/06/2022
finished        02:02:56 22/06/2022
Experiment [done; Instance: 0; Duration: 342513 ms]
Running [done]
Writing documentation...
Writing [done]
Finished

```

Figure 67. Console output produced when running Tutorial 8 experiment.

5.8.4. Results

The results of the experiment can be found in 4 directories with the names of the experiment widgets `fstWriter`, `popWriter`, `HsMapWriter` and `popMapWriter`.

The widgets `fstWriter` and `popWriter` have produced the following files:

- **Design.csv**: Experiment design details.
- <Field name>.csv: time series of the data from each simulator
- <Field name>_avg.csv: time series of the data averaged over simulators
- <Field name>_anova.R: R script for the anova computation.
- <Field name>_boxplots.R: R script for creating box plots of the output.
- <Field name>_barplots.R: R script for creating a bar plot of the output.
- <Field name>_RVEplot.svg: A plot, created by R, of the relative variance explained by each of the terms of the ANOVA.
- <Field name>_boxplots.svg: A box plot, created by R, response variable to each of the treatments.
- <Field name>_barplots.svg: A bar chart, created by R, of the response variable to each of the factor combinations (i.e. each simulation averaged over replicates).
- <Field name>_AnovaInput.csv: table of data by treatment value name presented to R.
- <Field name>_anovaResults.csv: raw results produced by R.
- <Field name>_RelSumSq.csv: variance explained relative to the total explained.
- <Field name>_var.csv: variance in time series over replicates.

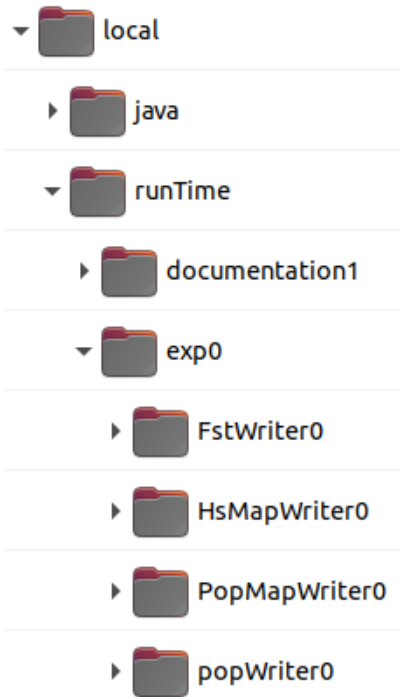


Figure 68. Directory structure created by the `FstWriter` and `popWriter` widgets.

The widgets `HsMapWriter` and `popMapWriter` will also have produced (a duplicate) of the 'Design.csv' file and one *.svg image for each of the simulations. These files will have the following naming convention:

`r<replicate number>_s<simulator number>\t<time step>\<factor level>`

The factors will be in the order set by the `rank` property (step 9 in the Experiment section above).

Examining `Fst_RVEplot.svg` and `population_RVEplot.svg` we can see that most of the variation in `Fst` is explained by dispersal distance (`dd`) (63%) and Dispersal method (`dm`) is the most important factor in explaining variation in population size (67%).

`Fst` was increased by short distance dispersal and to a lesser extent by uninformed dispersal (`Fst_boxplots.svg` and `Fst_barplot.svg`).

Population was increased by informed dispersal and to a lesser extent by long distance dispersal (`population_boxplots.svg` and `population_barplot.svg`)

You can recreate this tutorial by selecting new → tutorials → 12 Running experiments .

Bibliography

Bousquet, F., & Le Page, C. (2004). Multi-agent simulations and ecosystem management: a review. *Ecological Modelling*, 176:313–332. <https://doi.org/10.1016/j.ecolmodel.2004.01.011>

Connell, J.H. (1978). Diversity in tropical rain forests and coral reefs - high diversity of trees and corals is maintained only in a nonequilibrium state. *Science* 199:1302–1310

Coquillard, P., & Hill, D. (1997). *Modélisation et simulation d'écosystèmes. Des modèles déterministes aux simulations à événements discrets*. Masson, Paris.

Flint, S. R. (2006). *Aspect-Oriented Thinking - An approach to bridging the disciplinary divides*. PhD, Australian National University. <https://openresearch-repository.anu.edu.au/handle/1885/49328>

Gignoux, J., I.D. Davies, S.R. Flint, & J.D. Zucker (2011). The Ecosystem in Practice: Interest and Problems of an Old Definition for Constructing Ecological Models. *Ecosystems* 14: 1039-54. <https://doi.org/10.1007/s10021-011-9466-2>.

Gignoux, J., G. Chérel, I.D. Davies, S.R. Flint, & E. Lateltin (2017). Emergence and Complex Systems: The Contribution of Dynamic Graph Theory. *Ecological Complexity* 31: 34-49. <https://doi.org/10.1016/j.ecocom.2017.02.006>.

Grimm, V., & Railsback, S. (2005). *Individual-based modelling and ecology*. Princeton University Press.

Grimm V, Berger U, DeAngelis DL, Polhill JG, Giske J, Railsback SF (2010) The ODD protocol: A review and first update. *Ecological Modelling* 221:2760–2768.

Harary, F., & Gupta, G. (1997). Dynamic graph models. *Mathematical and Computer Modelling*, 25(7), 79–87. [https://doi.org/10.1016/S0895-7177\(97\)00050-2](https://doi.org/10.1016/S0895-7177(97)00050-2)

Lim, W. H., & Roderick, M. L. (2009). *An atlas of the global water cycle based on the IPCC AR4 climate models*. ANU E Press.

Melilo, J. M., Borchers, J., Chaney, J., Fisher, H., Fox, S., Haxeltine, A., Janetsos, A., Kicklighter, D. C., Kittel, T. G. F., McGuire, A. D., McKeown, R., Neilson, R., Nemani, R., Ojima, D. S., Painter, T., Pan, Y., Parton, W. J., Pierce, L., Pitelka, L., ... Woodward, F. I. (1995). Vegetation/ecosystem modeling and analysis project: comparing biogeography and biogeochemistry models in a continental-scale study of terrestrial ecosystem responses to climate change and CO₂ doubling. *Global Biogeochemical Cycles*, 9(4), 407–437.

Parton, W., Stewart, J., & Cole, C. (1988). Dynamics of C,N, P and S in grassland soils: a model. *Biogeochemistry*, 5, 109–131.

Quesnel G, Duboz R, Ramat E (2009). The Virtual Laboratory Environment - An operational framework for multi-modelling, simulation and analysis of complex dynamical systems. *Simulation Modelling Practice and Theory* 17:641–653.

Roxburgh, S. H., Barrett, D. J., Berry, S. L., Carter, J. O., Davies, I. D., Gifford, R. M., Kirschbaum, M. U. E., McBeth, B. P., Noble, I. R., Parton, W. G., Raupach, M. R., & Roderick, M. L. (2004). A critical overview of model estimates of net primary productivity for the Australian continent. *Functional Plant Biology*, 31(11), 1043–1059.

Tansley, A G. (1935). The use and abuse of vegetational concepts and terms. *Ecology* 16: 284-307.

Zeigler B, Kim TG, Praehofer H (2000) *Theory of Modeling and Simulation: integrating discrete event and continuous complex dynamic systems*. Academic Press

Zucker, J.D. (2003). A Grounded Theory of Abstraction in Artificial Intelligence. *Philosophical Transactions of the Royal Society B: Biological Sciences* 358: 1293-1309. <https://doi.org/10.1098/rstb.2003.1308>.