Felix2 User Manual

C++ simulation tool for biological neural networks

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Contents

I Getting started 1

1 Introduction 3
    1.1 History of Felix and Felix2 3
    1.2 How to read this book 3

2 Installing Felix2 5

3 Compiling and Running Felix2 simulations 7
    3.1 The structure of a Felix2 simulation program 7
    3.2 Compiling a Felix2 simulation 10
    3.3 Running a Felix2 simulation 10

4 Overview 11
    4.1 Basic architecture of Felix++ 11
        4.1.1 The core modules of Felix++ 12
        4.1.2 Auxiliary modules of Felix++ 13
        4.1.3 Component classes of Felix++ 15
    4.2 Simulation environment and components: Code examples 17
        4.2.1 Simulation environment: Class TSimulationEnvironment 18
        4.2.2 Components: Class TComponent 19
        4.2.3 Class TSSNeuron: a simple spiking neuron model 21
    4.3 Structure of a Felix++ simulation 23
        4.3.1 A skeleton simulation program 23
        4.3.2 The parameter file 25
        4.3.3 Compiling and running simulations 26

II The Graphical User Interface (GUI) of Felix 29

5 The program structure of GUI simulations 31

6 Using the GUI elements of Felix 33

III Fundamentals of Felix2 35

7 Type conventions 37
CONTENTS

8 Layouts and multi-dimensional arrays .................................................. 39
9 Parameters and parsing ......................................................................... 41
10 Parameters and parsing ......................................................................... 43
11 Basic numerics ..................................................................................... 45
   11.1 Time .............................................................................................. 45
   11.2 Random generators ....................................................................... 45
   11.3 Constants, functions, and look-up-tables ....................................... 45
12 Ports ........................................................................................................ 47
13 Kernels ...................................................................................................... 49
14 Simulation environment and components ............................................ 51

IV Modelling the environment: input and output ................................... 53
15 Vectors and patterns ............................................................................. 55
16 Modelling objects and the space around ............................................ 57
17 Recording of simulation data ................................................................. 59

V Further elements of Felix2 ..................................................................... 61
18 Integrators for differential equations ................................................... 63
19 Delays ...................................................................................................... 65
20 Receptors ............................................................................................... 67
21 A library for associative memory .......................................................... 69

VI Components of Felix2 .......................................................................... 71
22 Noise populations .................................................................................. 73
   22.1 TMUniformNoise: the standard noise population ....................... 73
   22.2 TMCorrelatedNoise: noise correlated in space and time ............ 73
23 Neuron populations ................................................................................ 75
   23.1 IFNeuron: a simple integrate-and-fire neuron model .................. 75
   23.2 SGNeuron: ................................................................................... 75
   23.3 SSNeuron: ..................................................................................... 75
   23.4 SSCOscillator: ................................................................................. 75
   23.5 InpNeuron: ..................................................................................... 75
CONTENTS

24 Connections 77
24.1 TopoConnection: ................................. 77
24.2 GaussConnection: ................................. 77
24.3 BlankTopoConnection: ......................... 77
24.4 DemoBlankTopoConnection: .................... 77
24.5 AssoConnection: ................................. 77
24.6 V1Connection: ................................. 77
24.7 RandomConnection: ............................. 77
24.8 DelayKernelConnection: ........................ 77

25 Learner 79
25.1 STDPLearner: ...................................... 79

VII Simulation examples 81
26 Integrating Felix2 and the GUI of Felix1 83
27 A simple network of oscillating neurons 85

VIII Appendices 87
A The GUI reference of Felix 89
B The C++ classes of Felix2 91
C Parameter scopes for Felix2 components 93
  C.1 Noise populations ................................. 93
  C.2 Neuron populations ............................... 93
    C.2.1 Class TIFNeuron .............................. 93
    C.2.2 Class TSSNeuron ............................. 94
  C.3 Connections ........................................ 95
    C.3.1 Template TMGaussConnection .................. 95
    C.3.2 Template TMAssocConnection ................... 97
  C.4 Learner ............................................. 98
Part I

Getting started
Chapter 1

Introduction

1.1 History of Felix and Felix2
1.2 How to read this book
Chapter 2

Installing Felix2

Installing Felix2 on a Linux system is fairly simple:

1. Make sure that you have installed the ‘xview’ package which is contained, for example, in the SuSe Linux distribution.

2. Get the Felix2 source code package Felix2.tar.gz, for example, from http://www.informatik.uni-ulm.de/ni/mitarbeiter/ AKnoblauch.html under item “software” and put the file into the directory where you want to install Felix2.

3. In your shell, apply the commands gunzip Felix2.tar.gz and tar xf Felix2.tar which creates a directory “Felix2” containing all files.

4. Add something like

   # Felix2
   setenv FELIX2HOME /private/aknoblaeu/neuro/Felix2
   setenv OPENWINHOME /usr/openwin
   setenv LD_LIBRARY_PATH ${LD_LIBRARY_PATH}:$OPENWINHOME/lib:{$FELIX2HOME}/lib
   alias Felix2 ${FELIX2HOME}/Felix2

   to your .cshrc file in your home directory.

5. Go into directory FELIX2HOME/test and apply at the command-line: Felix2_tdel which compiles a test simulation for Felix1. Start the simulation with ./tdel. If there pop-ups a window with some sliders then all is ok. you can start the simulation by pressing the ‘run’ button.

6. Make a second test for Felix2 by entering directory FELIX2HOME/test and apply at the command-line: Felix2_exampleSS which compiles a test simulation for simple spiking neurons. Start the simulation with ./exampleSS similarly as before.

If problems occur with compiling or linking of the test simulation, you may try to recompile Felix2. For this you have to do the following:

1. enter directory FELIX2HOME/xview, clear all old object files by rm *.o and recompile with make all.
2. enter directory FELIX2HOME/src, clear all old object files by \texttt{rm *.o} and recompile with \texttt{make all}.

3. enter directory FELIX2HOME/src_Felix2, clear all old object files by \texttt{rm *.o} and recompile with \texttt{make all}. This may take some minutes.
Chapter 3

Compiling and Running Felix2 simulations

Every Felix2 simulation is essentially a C++ program which uses the Felix2 library. In so far it is up to the programmer to write a C++ program in an ingenious way and use (or even use not) the Felix2 libraries which deliver just classes for neurons, network connections and other things. However, in this chapter we want to give some guidelines how we do think a typical simulation program should be structured, and moreover, how the whole process of simulating should be structured to make use of Felix2 in an optimal way.

3.1 The structure of a Felix2 simulation program

We recommend that a Felix2 simulation program typically should be structured like this:

```c
0: // Part 1: Felix2 declarations
1: // ------------------------------------
2: #include "F2_simenv.h"
3: ... // further includes
4:
5: #define STANDALONE 1
6: ... // further macro definitions
7:
8: int nArgs;
9: char** args;
10:
11: TSimulationEnvironment* senv;
12: TSSNeuron* popPe;
13: ... // further declarations
14:
15: // Part 2: Felix1 (GUI) declarations
16: // ------------------------------------
17: #if FELIX_GUI==1
18:
19: extern "C" { 
20: #include <stdio.h> // include the Felix headers
21: #include "nn.h"
22: #include "vector.h"
```
CHAPTER 3. COMPILING AND RUNNING FELIX2 SIMULATIONS

23: # include "random.h"
24: # include "delay.h"
25: }
26: #endif
27:
28: char *outPe;  // field to be displayed by GUI
29: ...            // further declarations
30:
31: #if FELIX_GUI==1
32: #if STANDALONE
33: NO_DISPLAY
34: #else
35: BEGIN_DISPLAY
36: reparseSWITCH((*senv));    // declaration of reparse-switch
37: ...                      // further switches and sliders
38:
39: WINDOW("out")     // declare image in window
40:  IMAGE("outPe",NR,AC,&outPe,POINTER TO bMATRIX,PeX,PeY,-0.1f,1.2f,1);
41: ...                  // further GUI declarations
42: END_DISPLAY
43: #endif
44:
45: NO_OUTPUT            // no Felix1 output
46: #endif
47:
48: // Part 3: main_init()
49: // ------------------------------------
50: int main_init() {
51:  senv = new TSimulationEnvironment(parameterFile,nArgs,args);
52:  ...                        // further declarations/creations
53:  popPe = new TSSNeuron(*senv,"popPe",popLT_Pe,0,integrator1,1);
54:  ...                      // further creation of objects
55:  senv->allocate();
56:  outPe  = (char*)popPe->y;
57:  ...                    // further assigning GUI variables
58:  cout << "maininit completed...\n";
59:  return 0;
60: }
61:
62: // Part 4: init()
63: // ------------------------------------
64: int init() {
65:  senv->init();
66:  ...                         // further initialization
67:  cout << "init completed...\n";
68:  return 0;
69: }
70:
71: // Part 5: step()
72: // ------------------------------------
73: int step() {
74:  senv->step();
75:  ...                        // further step()-stuff
3.1. THE STRUCTURE OF A FELIX2 SIMULATION PROGRAM

As illustrated the typical Felix2 simulation consists of 6 parts:

- **Part 1** (lines 0-14) contains all the declarations for the simulation except the declarations for the graphical user interface (GUI). It should include at least the F2_simenv.h but also the headers of other Felix2 components that are used in the program. The macro definition STANDALONE serves to switch between two simulation modes: For STANDALONE=0 the GUI is used and simulation variables can be displayed online and simulation parameters can be modified online e.g. via the switches, sliders, or most easily using the reparse-switch and the parameter file (see also section ?? on page ??). The variables nArgs and args are used for the transfer of parameters from the call of the C++ program to the simulation environment in standalone simulations (STANDALONE=1, cf. Part 6). Although it is possible to declare the Felix2 components as the simulation environment or neurons statically as global variables, it is strongly recommended that the Felix2 components are declared dynamically as pointers in Part 1, while the creation of the actual objects is placed in main_init() (Part 3). One reason why to do so is the handling of the parameter transfer via nArgs and args. If the Felix2 components are declared as static global objects then they are already created and parsed(!) before the call to main(nArgs, args) is performed. Consequently it is not possible to adjust constants in the parameter file via a the nArgs_/args_parameters of main(), see also section ?? on page ???. Another reason why to use pointer declarations in part 1 is that certain free parameters can only be adequately reparsed if they were declared as pointers (see section ?? on page ??).

- **Part 2** (lines 15-47) contains the GUI declarations of Felix1. First the Felix1 headers must be included as extern C. Then there should be declared variables which are used to display simulation data online. Finally Part 2 contains the declaration of the GUI components of Felix1, such as the reparse-switch, sliders, images, graphs, functions, etc. See section ?? on page ?? for more details.
• **Part 3** (lines 48-61) is essentially the `main_init()` procedure as required by Felix1. First the simulation environment should be created which initiates the parsing of the remaining objects using correctly possibly modified constant of the parameter file. The remaining objects are created and parsed from the parameter file as soon as they allocated. After creating the Felix2 objects, a call to `senv->allocate()` may be necessary to allocate certain integrator objects (see section ?? on page ??). Finally the display pointer variables (like `outPe` in line 56) must connected with the correct component fields.

• **Part 4** (lines 62-70) is the `init()` procedure as required by Felix1. In the simplest case it may only contain the call to `senv->init();` which will call the `init()` methods of all the previously created Felix2 objects.

• **Part 5** (lines 71-79) is the `step()` procedure as required by Felix1. In the simplest case it may only contain the call to `senv->step();` which will call the `step()` methods of all the previously created Felix2 objects. In addition `senv->step();` will check sliders and the reparse-switch (line 36), and react to possible events as parameter updates.

• Finally **Part 6** (lines 80-96) contains the `main(nArgs, args)` procedure which is only used for standalone simulations with `STANDALONE=1`.

### 3.2 Compiling a Felix2 simulation

### 3.3 Running a Felix2 simulation
Chapter 4
Overview

The following is unedited text from my dissertation "Synchronization and pattern separation in spiking associative memories and visual cortical areas."

All simulations described in this work have been implemented using the Felix or Felix++ simulation tools. Originally the C based simulation tool Felix has been developed by Thomas Wennekers at the University of Ulm [?] as a universal simulation environment for physical and, in particular, neural systems. The development of Felix was motivated by the need for a fast implementation of multi-layer one- or two-dimensional neural structures such as neuron populations. For this purpose, Felix provides elementary algorithms for single-cell dynamics, inter-layer connections, and learning. Additionally, there exist also libraries for non-neural applications, e.g., for general dynamical systems and elementary image processing.

Simulations can be observed and influenced online via the X11/XView-based graphical user interface (GUI) of Felix. The Felix GUI provides elements such as switches for conditional execution of code fragments, sliders for online-manipulation of simulation parameters (like connection strengths, time constants, etc.), and graphs for the online observation of the states of a simulated system in xy-plots or gray-scale images (see [?, ?] for more details).

During this work the simulation tool Felix++ has been developed as a C++ based object-oriented extension of Felix. Felix++ provides additionally classes for neuron models, n-dimensional connections, pattern generation, and data recording. Current installations of Felix++ are running on PC/Linux as well as on 64bit-SunFire/Solaris9 systems. In the following the architecture of Felix++ is briefly sketched (for more details see [?]).

4.1 Basic architecture of Felix++

Essentially Felix++ is a collection of C++ libraries supporting fast development of neural networks in C++ [?, ?]. Thus Felix++ comprises a number of modules each consisting of a header (with the suffix " .h") and a corpus (with the suffix " .cpp" for Felix++/C++ or " .c" for Felix/C). The header files contain declarations of classes, types, and algorithms, whereas in the corpus files the declarations are implemented. Figure 4.1 illustrates the architecture of Felix++ by classifying all the modules of Felix++ and Felix in a hierarchy.
Figure 4.1: Architecture of Felix++: A simulation is a C++ program that includes headers of Felix and Felix++. Felix++ contains core modules (e.g., F2_simenv.h/cpp; cf. Fig. 4.2), auxiliary modules, and modules for simulation components such as neuron populations and connections between neuron populations (cf. Fig. 4.3). The Felix modules implement a graphical user interface and elementary algorithms (see [?] for more details).

4.1.1 The core modules of Felix++

The core of Felix++ contains the most important modules required by all other Felix++ modules.

- **F2.types.h/cpp** declares some elementary type conventions and some global objects.
- **F2.time.h/cpp** declares classes for time, for example to evaluate the time necessary for computing a simulation.
- **F2.random.h/cpp** provides several different random number generators (see [?]).
- **F2.layout.h/cpp** declares so-called layouts. A layout can be used to define the topology of a vector (or in terms of C++, an array). For example a population of 1000 neurons can be arranged as a $10 \times 10 \times 10$ cuboid. Apart from cuboid layouts also ellipsoid layouts are defined which are useful in particular for saving memory when modeling isotropic local connectivity (in three dimensions, for example, an ellipsoidal kernel saves almost 50 percent of the memory required by a cuboid kernel).
- **F2.parameter.h/cpp** declares classes for simulation parameters. For example, the membrane time constant field of a neuron class is usually declared as such a parameter. This allows conveniently parsing and online reparsing of the parameters from a parameter
file as well as online manipulation via sliders of the Felix GUI (including updating of other dependent parameters). The parameters declared in this module are essentially arrays equipped with a cuboid layout (see above). Therefore they can be used not only for single parameter values, but as well for multi-dimensional parameter collections such as parameter vectors or matrices.

- **F2.parser.h/cpp** provides the classes for parsing parameters from a parameter file. Usually, in Felix++ a component class (e.g., a neuron class) is designed in such a way that a construction of an object is paralleled by parsing the corresponding parameters from a file. Furthermore, during the simulation the parameter file can be modified and reparsed by pressing the reparse-button.

- **F2.port.h/cpp** declares interfaces for the communication between different simulation components, so-called ports. For example, a neuron class may contain an output port representing the spikes of the neuron, and an input port representing synaptic input to the neuron. Correspondingly, the constructor of a connection component class requires as parameters the output port of a neuron population and the input port of another neuron population such that the spikes from the first population can be propagated to the dendrites of the second population.

- **F2.simenv.h/cpp** declares the simulation environment class TSimulationEnvironment and the component base class TComponent as well as some base classes for special components such as neurons (TNeuron) and connections (TConnection). This module should be included by any simulation program using Felix++. The simulation environment is essentially a container for the simulation components (see below for more details; cf. Fig. 4.2), but provides also additional infrastructure such as look-up tables (for example for Gaussians), random number generators, and much more. Usually, the construction of a simulation component requires a TSimulationEnvironment as an argument, such that the component is automatically inserted. After construction of all the components, calls to methods allocate() will allocate memory shared by multiple components (for example when integrating differential equations via TIntegrator objects; see below). Before starting the simulation all the components can be initialized by calling the init() method of the simulation environment. Similarly, during the simulation a call to the step() method will compute one simulation step.

### 4.1.2 Auxiliary modules of Felix++

Besides the core modules there are a number of auxiliary modules that provide additional functionality required by only some of the Felix++ component modules, and perhaps also by the programmer developing a simulation.

- **F2.numerics.h/cpp** provides a number of useful constants (e.g., \( \pi \), \( e \), and \( \ln 2 \)) and functions (e.g., density function of Binomials or Gaussians, information and transformation functions for binary random variables, etc.). Further declarations provide classes for look-up-tables and interpolation.

- **F2.kernel.h/cpp** declares classes for kernels that can be used, for example, for implementing synaptic connections. Kernels are essentially arrays (e.g., of synaptic weights
CHAPTER 4. OVERVIEW

Simulation Environment

<table>
<thead>
<tr>
<th>Components:</th>
<th>Methods:</th>
</tr>
</thead>
<tbody>
<tr>
<td>neuron populations</td>
<td>allocate()</td>
</tr>
<tr>
<td>connections</td>
<td>init()</td>
</tr>
<tr>
<td>etc.</td>
<td>step()</td>
</tr>
</tbody>
</table>

Figure 4.2: The simulation environment object (of class TSimulationEnvironment) is essentially a container object containing all the simulation components such as neuron populations or connections. The components are inserted during construction. Before starting a simulation a call to method allocate() is necessary to allocate memory. A call to init() initializes the components, and each call to step() results in the computation of one simulation step.

or delays) that have been assigned a topology via layouts (see F2_layout.h/cpp). The classes defined in this module enable, for example, the coordination of a neuron population (layout) to a set of kernels. This happens in a rather flexible manner such that each neuron can be assigned individually a kernel index, where also certain regularities of kernel arrangements can be exploited (such as the regularities occurring for the orientation modules in our visual model; cf. Fig. ??a).

- **F2_vector.h/cpp** provides basic vector functionality. This module also declares classes for numerical vector parameters (cf. F2_parameter.h/cpp).

- **F2_pattern.h/cpp** implements classes for various types of patterns. From the pattern base type (TMPattern) which corresponds simply to a multi-dimensional array there are derived specialized pattern types such as binary patterns (TMbPattern), sparse binary patterns (TMsbPattern), sparse patterns (TMsPattern), or sparse binary spatio-temporal patterns (TMsbSTPattern). Additionally, further auxiliary classes have been implemented in order to facilitate the use of patterns. For example, pattern container classes are declared (TMPatternStock and derivatives of TMPatternGroup) for convenient construction and parsing of pattern groups from parameter files. The TMPatternRanking class can be used for analyzing neural activity with respect to a set of patterns (i.e., to determine the pattern in the set that is most similar to the neural activity pattern). Similarly, the TMPatternHistogram class can be used to create pattern-specific histograms of state variables (as used, for example, for the threshold distance histograms in Fig. ??d-h).

- **F2_delay.h/cpp** provides classes based on the definitions in F2_kernel.h/cpp for efficient implementation of synaptic delays.

- **F2_object.h/cpp** declares classes for generating stimulus objects. Further classes can be used to put static or moving objects in space (derivatives of TMSpace), or to project the
In the visual model of chapter ?? these classes have been used in order to project a visual scene of several stimulus objects onto the retinal area R (see Figs. ??a and ??a).

- **F2.record.h/cpp** provides the infrastructure for efficient recording of simulation data.

- **F2.receptor.h/cpp** declares classes for the efficient implementation of various types of receptors for synaptic transmitters. Derivatives of class TMReceptorPort can be used, for example, to implement certain transmitter-dependent synaptic conductances. For our neuron model described in section ?? we used class TMOffDynamics for implementation of excitatory AMPA currents (conductance \( g_{\text{exc}} \); cf. eq. ??) and inhibitory GABA-A currents (conductance \( g_{\text{in}} \); cf. eq. ??). More complex receptor dynamics are implemented by classes TMOnOffDynamicsRP and TMSimpleNMDARP where the latter can be used to model NMDA receptor dependent currents (cf. [?]). Actually neuron classes such as TSSNeuron (which we have used for our biological simulations) or TGNeuron use the receptor port classes provided by this module. These models can be equipped with an arbitrary configuration of different receptor ports which can be specified in the parameter file (see below code fragment 4.5).

- **F2.libasso.h/cpp** encapsulates the C-library for associative memory implemented by Friedrich Sommer (cf. [?]).

### 4.1.3 Component classes of Felix++

Based on the core and auxiliary modules there exists already a large number of simulation components. Figure 4.3 illustrates the class hierarchy of the Felix++ simulation components. They can be divided into the following component base classes derived from TComponent:

- **TNeuron** (defined in module F2.simenv.h/cpp) is the base class for all neuron classes. Currently there are implementations for gradual neurons (TSGNeuron in module SNGeunor.h/cpp and TGNeuron in module GNeuron.h/cpp), spiking neurons (TIFNeuron in module IFNeuron.h/cpp and TSSNeuron in module SSSNeuron.h/cpp), and oscillators (TSSCOscillator in module SSCOscillator.h/cpp) which all can be used for biological modeling. Additionally, there are classes adequate for technical implementations of associative memory (TMAssociationPopulation and TMAutoWillshawTAMP in module F2.association.h/cpp). For the simulations of biological models in section ?? and chapter ?? we used the simple spiking neuron class TSSNeuron, whereas for the implementation of Willshaw associative memory and the spike counter model in section ?? the technical associative memory population class TMAutoWillshawTAMP has been used.

- **TMNoise** (defined in module F2.simenv.h/cpp) is the base class for noise populations. A noise population provides random numbers generated according to a certain distribution for another component object such as a neuron population. Derivatives of type TMUniformNoise (defined in module UniformNoise.h/cpp) provide uniformly distributed random numbers with a certain power (or variance). While this type generates independent random numbers in each simulation step the random numbers generated by derivatives of type TMCorrelatedNoise can be correlated in space and time. The standard noise type for neuron populations such as TSSNeuron (or for synaptic noise in connections; see module F2.receptor.h/cpp) is TMUniformNoise,
Figure 4.3: The class hierarchy for the currently implemented simulation components of Felix++. From the base class TComponent specialized sub-classes are derived for neuron populations (TNeuron), noise generators (TMNoise), connections between neuron populations (TConnection), integration of differential equations (TMIntegrator), synaptic plasticity (TLearner), representations of stimulus space (TMSpace), and on-line observation of the simulation state (TObservable).

whereas for the primary visual area P (see chapter ??) correlated noise of type TMCorrelatedNoise has been used.

- TConnection (defined in module F2_simenv.h/cpp) is the base class for connections between neuron populations (or more exactly, between ports; see module F2_port.h/cpp). The function of derivatives of this type is to propagate information from an output port to an input port, for example, to propagate the spikes from the output port of one neuron population through the network to the input port of another neuron population. The most important derived type for biological modeling is TMTopoConnection (defined in module TopoConnection.h/cpp). This generic type is the base class for many further derived classes. Here a synapse is defined by two state values: A synaptic weight, and a synaptic delay. Correspondingly, an object of type TMTopoConnection essentially contains two kernel arrays of type TKernel (defined in F2_kernel.h/cpp) representing weights and delays. The kernel classes can be applied in a very flexible manner allowing implementation of full, sparse, topographical schemes in multiple dimensions. Additionally, efficient algorithms are implemented for several special cases (e.g., for non-sparse bit-packed binary topographical connectivity). The derivatives of TMTopoConnection merely specify how the synaptic weight and delay kernels are generated. For example, class TMGaussConnection (defined in module GaussConnection.h/cpp) implements simple topographical connections with Gaussian kernels. A further derived class TMBankTopoConnection (defined in module BlankTopoConnection.h/cpp) provides an interface to TMTopoConnection in order to allow a more convenient derivation of further connection classes. While TMDemoBlankTopoConnection (defined in module DemoBlankTopoConnection.h/cpp) is merely a demonstration how to derive from TMBankTopoConnection, also a number of important con-
4.2 Simulation environment and components: Code examples

In the last section we have obtained an overview over the modules of Felix++. In the following we will have a closer look at the code defining some important Felix++ classes, the simulation environment, and the base class for components. The code examples shown below are shortened fragments of the declarations in the Felix++ headers.

- **TMIntegrator** (defined in module `F2_integrator.h/cpp`) is the base class for numerical integration of differential equations. Derived classes (defined in the same module) are **TMEulerIntegrator** which implements a simple first order Euler method, and **TMRK4cIntegrator** which implements the fourth order Runge-Kutta method with constant step size (see [?]). Normally, these integrator objects are used by some of the neuron classes (e.g., TSSNeuron).

- **TLElerner** (defined in module `F2_simenv.h/cpp`) is the base class for plasticity of synaptic connections. Currently, two derivatives are implemented in the module `STDPLearner.h/cpp`. **TMSTDPLearnerSMA2000** implements a model of spike-timing dependent synaptic plasticity (STDP) described by Song, Miller, and Abbott [?], while **TMSTDPLearnerFD2002** implements an extended model suggested by Froemke and Dan [?]. Both classes are interfaced with connection classes via the kernel classes defined in `F2_kernel.h/cpp`. Therefore it is easy to endow connections (e.g., derived from **TMTopoConnection**) with synaptic plasticity.

- **TMSpace** (defined in module `F2_object.h/cpp`) is the base class for the definition of a space for stimulus objects (see above module `F2_object.h/cpp`).

- **TObserver** (defined in module `F2_simenv.h/cpp`) is the base class for components observing on-line the state of the simulation. Derived classes are **TMPatternRanking** and **TMPatternHistogram** (for more details see above module `F2_pattern.h/cpp`).

**4.2 Simulation environment and components: Code examples**
4.2.1 Simulation environment: Class TSimulationEnvironment

The simulation environment is essentially a container for the simulation components used in a simulation program (cf. Fig. 4.2), but also provides additional infrastructure such as lookup tables, random number generators, and some further global variables. The following code fragment taken from the header T2_simenv.h shows parts of the declaration of the class TSimulationEnvironment.

Code fragment 4.1
class TSimulationEnvironment {
public:
// part 1: local types
typedef enum { CC_NOISE, // for TNoise component category
    CC_CONNECTION, // for TConnection
    CC_INTEGRATOR, // for TIntegrator
    CC_NEURON, // for TNeuron
    CC_LEARNER, // for TLearner
    CC_SPACE, // for TSpace
    CC_SPACE_REPRESENTATION, // for TSpaceRepresentation
    CC_OBSERVER, // for observers like TMPatternRanking
    CC_ASYNCHRONY // calls to step() are not controlled
} TComponentCategory;
static const int nCategories=9; // number of different component categories

// part 2: parameters
TPar parStepSize; // simulation step size (in milliseconds)
TsPar parDataDirectory; // default directory where simulation data is recorded
TsPar parDataPostFix; // post fix for data file names

// part 3: object fields
const char* parameterFile; // name of root parameter file
TParser parser; // parser for parameters
vector<TComponent*> allComponents; // all components in order of constr.
vector<TComponent*> components[nCategories]; // components ordered after categories

// part 4: time reference
TInt steps; // current simulation step number
TFloat simTime; // current simulation time, simTime:=steps*stepSize;

// part 5: constructors/destructors
TSimulationEnvironment(const char* parFile); // default constructor
'~TSimulationEnvironment(); // destructor

// part 6: methods
void parse(); // parsing of parameters
void reparse(); // reparse parameters
void addComponent(TComponent *c, TComponentCategory cc); // add component c

void allocate(); // simulation phase 2 (after creation phase): memory allocation
void init(); // simulation phase 3: Initialization of TComponents
void step(); // simulation phase 4: compute one simulation step
};

In part 1 an enumeration type is declared for the different component categories such as neurons, connections, or observers. This corresponds approximately to the different compo-
4.2. SIMULATION ENVIRONMENT AND COMPONENTS: CODE EXAMPLES

In part 2 parameters are declared such as the simulation step size, or the directory for recorded simulation data. These parameters are parsed from a parameter file (see code fragment 4.5) during the first execution phase of the simulation program when the simulation environment is created by calling the constructor (see part 5).

In part 3 some object fields are declared such as the name of the parameter file (which is passed as an argument to the constructor; see part 5), or the parser used for parsing the parameters (see module F2.parser.h/cpp in section 4.1.1). Here there are also the declarations of the containers for the simulation components (vectors of the STL library; cf. [?, ?]). The first container field allComponents contains references to the components in order of the construction of the components (which is important for reparsing the parameter file), while the second container field components contains the components ordered for the different categories (see part 1). The latter ordering is important to assert a defined synchronization of object of the same component category. For example, the calls to the step() methods of neuron objects should be before the step() calls of connection objects (cf. part 6). This synchronization will become even more important when parallelizing Felix++ as planned for future work (cf. [?]).

In part 4 some fields for time reference are defined. Felix++ is a step-based simulation tool (in contrast to event-based tools). This means that the state of the simulated system is updated step by step where one simulation step corresponds to a fixed time interval. The parameter parStepSize (see part 2) defines this time interval. The field steps is initialized by 0, and incremented for each call to the step() method (see part 6).

Part 5 contains the declarations of the constructors and destructors. In section 4.3 an example is given how and when to apply the constructor in a simulation program.

In part 6 the methods are declared. Method parse() is normally called by the constructor in order to parse the parameter file. Method reparse() is called for reparsing a modified parameter file, for example, when pressing the reparse-button in an online-simulation. With addComponent() new simulation components can be added to the component containers (see part 3) which is usually done by the constructor of TComponent (see section 4.2.2). Then there are further three important methods (cf. Fig. 4.2): In order to allocate memory shared by different simulation components (for example state variables of neurons integrated by components of type TIntegrator; see module F2.integrator.h/cpp) a call from the simulation program to allocate() must occur after the construction of the simulation components (see procedure main_init() in code fragment 4.4). A call to method init() initializes the simulated system essentially by calling the init() method of each simulation component. Similarly, a call to method step() computes one simulation step by calling the step() method of each simulation component. The calls to the simulation environment’s init() and step() methods occur normally from the simulation program’s init() and step() procedures (see code fragment 4.4).

4.2.2 Components: Class TComponent

The class TComponent is the base class for all simulation components such as neurons or connections (see Figs. 4.3 and 4.1) and implements essentially a common interface of simulation components to the simulation environment (section 4.2.1). The following code fragment taken from the header F2.simenv.h shows parts of the declaration.
CHAPTER 4. OVERVIEW

Code fragment 4.2

class TComponent : public TParamOwner {
public:
    // part 1: object fields
    string name; // name of the component
    TSimulationEnvironment& simEnv; // reference to the simulation environment
    TComponentCategory category; // component category
    vector<TfPort*> gradualPorts; // gradual i/o ports of the component
    vector<TbPort*> binaryPorts; // binary ports

    // part 2: constructors/destructors
    TComponent(TSimulationEnvironment& simEnv_arg, const char* name_arg,
               TComponentCategory cc_arg);
    TComponent(TComponent& pattern, const char* name_arg, TComponentCategory cc_arg);
    ~TComponent();

    // part 3: methods
    virtual void reparse(); // reparse parameters
    virtual void allocate(); // allocate memory for TIntegrator
    virtual void init(); // initialize component
    virtual void step(); // compute one simulation step of component
    virtual void derivs() {}; // compute derivatives for TIntegrator
};

Class TComponent is derived from class TParamOwner in order to provide the functionality of the parameter classes (see module F2 parameter.h/cpp in section 4.1.1).

In part 1 object fields are declared. Each simulation component can be assigned a name which considerably relieves search for errors. Field simEnv refers to the simulation environment containing the object, and field category contains information about the component category of the object (cf. section 4.2.1). The fields gradualPorts and binaryPorts are containers for gradual and binary ports (see module F2 port.h/cpp in section 4.1.1) and constitute thereby the interface for communication between different simulation components. For example, the spikes of a neuron population will be represented by a binary port of type TbPort and similarly the dendritic inputs will be represented by a gradual port of type TfPort. Thus a connection component can connect two neuron populations, for example, by propagating spikes from the binary port of the first population through the synaptic network to the gradual input port of the second population.

In part 2 the constructors and destructors are declared. There are generally two constructor types for a simulation component. The default (or complete) constructor constructs a simulation component by parsing the parameters from the parameter file (see code fragment 4.5). In contrast, the pattern constructor requires as argument a simulation object of the same type where the parameters of this pattern are used for construction of the new object (see also section 4.2.3).

In part 3 a number of virtual methods is defined which are normally overridden by derived component classes and called by the simulation environment. The reparse() method reparses the parameters from the parameter file using simEnv → parser. If necessary the allocate() method requests memory shared with other components from a further object managing the shared memory (for example from an TIntegrator object; see module F2 integrator.h/cpp in section 4.1.3). A call to init() will initialize the simulation component, and a call to step() will compute one simulation step for the component.
The method derivs() can be used in derived classes to compute the (numerical) derivatives of some of the state variables of the component. This method is normally called by a TIntegrator object in order to integrate the differential equation associated with a component state.

### 4.2.3 Class TSSNeuron: a simple spiking neuron model

To illustrate how a concrete simulation component can be derived from the base class TComponent we will have a closer look at the class TSSNeuron implementing a simple spiking neuron model. Actually, this class (with a parameter file as shown in code fragment 4.5) has been used to implement the model described in section ?? for the biological simulations in section ?? and chapter ???. The following code fragment taken from the header SSNeuron.h shows parts of the declaration.

**Code fragment 4.3**

```cpp
class TSSNeuron : public TNeuron, public TIntegratorClient {
public:
    // part 1: parameters
    string scopeID; // scope id for parsing parameters
    TVecPar tau_x; // membrane time constant
    TVecPar theta; // asymptotic threshold
    TVecPar refAbs; // absolute refractory time
    TVecPar refRel; // relative refractory parameter
    TVecPar tau_h; // decay time constant of habituation
    TVecPar thetaInc_h; // threshold increment after each spike
    TCompartmentReceptors* receptorPorts; // receptorPorts

    // part 2: integrator for membrane potential
    TDerivScope derivScope;
    TIntegrator* integrator; // integrator

    // part 3: ports
    TbPort *out; // output queue for spikes
    TfPort *lastOut; // output queue for last spikes
    TfPort* excIn; // default excitatory in-port
    TfPort* inhIn; // default inhibitory in-port

    // part 4: state variables
    TFloat *current; // synaptic currents
    TFloat *x; // membrane potential
    TByte *y; // output variable (refers to out)
    TFloat *last; // last spike time (refers to lastOut)
    TFloat *habituation; // habituation (fatigue) - increased threshold

    // part 5: constructors/destructors
    TSSNeuron(TSimulationEnvironment& simEnv_arg, const char* name_arg, // complete cnstr.
        TLayout* layout_arg, vector<TNoise*>* noiseSources_arg);
    TSSNeuron(TSimulationEnvironment& simEnv_arg, const char* name_arg, // default
        TLayout* layout_arg, vector<TNoise*>* noiseSources_arg,
        TIntegrator* integrator_arg, int parse);
    TSSNeuron(TSSNeuron& pattern, const char* name_arg, // patterned
        TLayout* layout_arg, vector<TNoise*>* noiseSources_arg);
    ~TSSNeuron(); // destructor

    // part 6: methods
```
void allocate(); // get memory from integrator
void derivs() {}; // compute ...
void derivs(int id, TFloat t, TFloat* x, TFloat* dxdt); // derivatives for integrator

void init(); // initialize states to zero values
void step(); // one simulation step
void reparse(); // reparse parameters
void handleUpdatedParameters(); // handle updated parameters
void setParameterValues(); // update parameter values
friend ostream & operator<<(ostream& os, const TSSNeuron & neuron); // output op.
friend istream & operator>>(istream& parser, TSSNeuron & neuron); // input op.
};

Class TSSNeuron is derived from TNeuron (which in turn is derived from TComponent; see Fig. 4.3) and from TIntegratorClient. The derivation from the latter class is necessary for any class requiring integration of differential equations by a TIntegrator object (see module F2/integrator.h/cpp in section 4.1.3).

In part 1 of code fragment 4.3 parameters of the neuron model are declared. When comparing with the notation used in section ?? (cf. table ?? and code fragment 4.5) parameter field tau_x corresponds to parameter \( \tau_x \), theta to \( \Theta_\infty \), refAbs to \( R_a \), refRel to \( R_r \), tau_h to \( \tau_h \), and thetaInc_h to \( H \). Field scopeID contains information about which parameter scope (see below) in the parameter file has been used to parse the parameters for this object. Field receptorPorts points to a container object for receptor ports (see module F2/receptor.h/cpp in section 4.1.2) which becomes allocated during object construction. This field is used for implementing different excitatory or inhibitory synaptic currents (see code fragment 4.5).

In part 2 field integrator declares the integrator object for integrating the differential equation for the membrane potential (cf. eq. ??). Currently, either a Euler or a Runge-Kutta method can be used (see module F2/integrator.h/cpp in section 4.1.3). The field derivScope just serves to identify the memory for the state variable \( x \) (see part 4) which is administrated by the integrator object.

Part 3 declares the ports of the object (see module F2/port.h/cpp in section 4.1.1). Port out is the output queue for the spikes of the neuron population, port lastOut contains information about the time of the last spike for each neuron. Input ports excIn and inhIn are essentially queues for synaptic input from other neuron populations mediated by connection objects (see type TConnection in section 4.1.3).

In part 4 the state variables of the neurons are declared. Array current is essentially the sum of the synaptic input currents for each neuron as computed by receptorPorts (see part 1). Array \( x \) corresponds to the membrane potential \( x \) of the neuron model in section ?? (cf. eq. ??). Similarly \( y \) corresponds to the spike output variable \( y \) (eq. ??), last corresponds to the time point \( s \) of the last spike for each neuron (cf. eq. ??), and habituation corresponds to the neuronal habituation or fatigue \( h \) (cf. eq. ??).

In part 5 the constructors and destructors are declared (cf. part 2 in section 4.2.2). The first constructor is the so-called complete constructor which is normally used in a simulation program for creating an object of type TSSNeuron for the first time. The third constructor is the pattern constructor which is normally used for further creations of TSSNeuron objects. While for the complete constructor the parameters (see part 1) are parsed from the parameter file (see code fragment 4.5), the pattern constructor copies the parameters from a pattern
object passed as the first argument. The second constructor in the code fragment is the so-called default constructor which is normally used by the constructor of another class derived from TSSNeuron. In contrast to the complete constructor, automatic parsing (which should be done only by the constructor of the derived class) can be suppressed by passing an additional flag argument parse.

Finally part 6 contains the declarations of the methods allocate(), derivs(), init(), step(), and reparse() which override the declarations explained above for the base class TComponent (see part 3 in section 4.2.2). There are a few remaining methods: handleUpdatedParameters() and setParameterValues() manage updating of the object state if one of the parameters (see part 1) is changed (for example when reparsing the parameter file), while the input/output operators operator>>() and operator<<() are used for parsing the object parameters from the parameter file, or for printing the parameter data, for example, when debugging.

4.3 Structure of a Felix++ simulation

4.3.1 A skeleton simulation program

A Felix++ simulation is basically a C++ program that includes header files of Felix and/or Felix++. The structure of a Felix++ simulation typically looks similar to the following code fragment:

Code fragment 4.4

```
// Part 1: Felix2 declarations
// ------------------------------------
#include "F2_simenv.h"
... // further includes (e.g., of Felix++ headers)
#define STANDALONE 1
... // further macro definitions
TSimulationEnvironment* senv;
TSSNeuron* popPe;
... // declaration of further simulation components

// Part 2: Felix1 (GUI) declarations
// ------------------------------------
extern "C" {
    #include "nn.h"
    ...
    // include of further Felix headers
}
#if STANDALONE
    NO_DISPLAY
#else
    BEGIN_DISPLAY
    ...
    // declaration of Felix GUI (switches, sliders, graphs, etc.)
    END_DISPLAY
    NO_OUTPUT // Felix1 output mechanisms usually not used
#endif

// Part 3: main_init()
// ------------------------------------
int main_init() {
    senv = new TSimulationEnvironment(parameterFile);
```
... // further utility declarations/creations
popPe = new TSSNeuron(*senv,"popPe",popLT_Pe,0,integrator1,1);
... // further creation of objects
senv->allocate();
... // assigning of GUI variables
return 0;
}

// Part 4: init()
// ------------------------------------
int init() {
  senv->init();
  ... // further initialization
  return 0;
}

// Part 5: step()
// ------------------------------------
int step() {
  senv->step();
  ... // further step()-stuff
  return 0;
}

// Part 6: main()
// ------------------------------------
#if STANDALONE
  int main(int nArgs, char** args) {
    main_init();
    init();
    for(int i=0;i<1000;i++) step();
    ... // further stuff, e.g. saving simulation data
  }
#endif

In part 1 Felix++ header files (and also other headers) are included, macros are defined such as STANDALONE (which switches between online and batch mode; see below parts 2 and 6), and the simulation environment and the simulation components are declared. For the sake of flexibility it is recommended to declare the simulation environment and the components as pointer variables which are allocated in the main_init() method (see part 3). For example, if the simulation environment and components would be already constructed here, it would not be possible to pass the name of the parameter file as an argument to the simulation program. In this example only the simulation environment sev and a neuron population popPe of type TSSNeuron (see section 4.2.3) are declared. Usually the construction of the simulation environment and the simulation components is paralleled with the parsing of a parameter file (see below the code fragment 4.5)

In part 2 the graphical user interface (GUI) of the simulation is declared (only necessary for online simulations, i.e., if the flag macro STANDALONE is inactive). For this purpose, first the Felix headers (see Fig. 4.1) must be included (in extern ‘‘C’’ brackets since Felix has been implemented in C). Then the GUI components of Felix can be specified in the #else branch of the #if directive (see [7] for details).

Part 3 is the main_init() procedure. Here the simulation environment and subsequently the simulation components are created by calling the corresponding constructors.
4.3. STRUCTURE OF A FELIX++ SIMULATION

After constructing all simulation components a call to the `allocate()` method of the simulation environment might be necessary (e.g., for simulation components such as TSSNeuron employing integrators; see module `F2_integrator.h/cpp` in section 4.1.3; cf. section 4.2.3). The `main_init()` procedure is normally called only once at the beginning of the simulation to construct the simulation objects. This is done either by the `main()` procedure (see part 6) for batch simulations (for activated flag macro `STANDALONE=1`) or by the Felix GUI for online simulations (for `STANDALONE=0`).

In part 4 the `init()` procedure is defined. It contains normally at least the call to the `init()` method of the simulation environment, but possibly also further initialization code for the simulation. The `init()` procedure should be called after `main_init()` to initialize the states of the simulation objects before the actual simulation computations start (see part 5). In contrast to `main_init()` the `init()` procedure can be called more than once either from the `main()` procedure (see part 6) for batch simulations or for online simulations by pressing the `init` button (or the `run` button) in the main simulation window [?].

Part 5 is the `step()` procedure which computes one simulation step. It contains normally at least the call to the `step()` method of the simulation environment, but possibly also further code for the simulation. The `step()` procedure is called either from the `main()` procedure (see part 6) for batch simulations or for online simulations by pressing the `step` button (or the `run` button) in the main simulation window.

Part 6 defines the `main()` procedure for batch simulations with activated macro flag `STANDALONE=1` (see part 1). This procedure must contain calls to `main_init()` and `init()` before the calls to the `step()` procedure.

### 4.3.2 The parameter file

The construction of the simulation environment and the simulation components in `main_init()` (see part 3 in section 4.3.1) is usually paralleled by the parsing of the parameter file in order to read in the parameters to be used for the respective simulation objects. The following code fragment shows parts of the parameter file for our skeleton simulation program above (code fragment 4.4).

#### Code fragment 4.5

```plaintext
#{ TSimulationEnvironment Simulation1 % parameter scope for simulation environment
stepSize : 0.1
dataDirectory : /private/aknoblau/simdata/BC2
}
#{ SSNeuron popPe % parameter scope for neuron population
tau_x(exp,sig,min,max) : 10 0 1 0
theta(exp,sig,min,max) : 10 0 1 0
refAbs(exp,sig,min,max) : 2 0 1 0
refRel(exp,sig,min,max) : 3 0.5 1.75 4.25
tau_h(exp,sig,min,max) : 150 0 1 0
thetaInc_h(exp,sig,min,max) : 0.6 1.0 0.2 1.0
#{ TMCompartmentReceptors receptors
nReceptorPorts : 2
#{ TMOffDynamicsRP AMPA
tau_OFF(exp,sig,min,max) : 5 0 1 0
E(exp,sig,min,max) : 80 0 1 0
g0(exp,sig,min,max) : 0 0 1 0
```
A parameter file is divided into various parameter scopes. A parameter scope is a group of parameters which has been put into scope brackets according to the syntax `#{ <scope type> <scope ID> <parameter1> <parameter2> ... }`. The scope type is given by the class of the object to be parsed, while the scope ID can be chosen arbitrarily.

This parameter file contains two global parameter scopes, one for the simulation environment `senv` and another for the neuron population `popPe` (cf. part 3 in code fragment 4.4). Parameter scopes can be organized hierarchically: For example scope `SSNeuron popPe` contains a sub-scope for the `receptorPorts` object (see part 1 in code fragment 4.4) which in turn can contain an arbitrary number of further sub-scopes for different receptor dynamics.

In the example there are two scopes for the receptor dynamics (cf. module `F2_receptor.h/cpp` in section 4.1.2) implementing the dynamics of the synaptic conductances of the neuron model described in section ???. The parameters in scope `TMOffDynamicsRP AMPA` specify the dynamics of the excitatory conductance $g_{ex}$ (eq. ??) which is implemented by the corresponding object of type `TMOffDynamicsRP`. Parameter `tau_OFF` corresponds to $\tau_{ex}$ (see eq. ??) and parameter `E` corresponds to $E_{ex}$ (see eq. ??). The additional parameters determine conductance baseline ($g0$), noise power (`powerInpNoise`), and the queue length (`qLen; measured in simulation steps) for incoming spikes propagated by connection objects (see class `TConnection` in section 4.1.3). The latter parameter determines the maximal possible axonal delay for the connection projecting onto this receptor port.

The parameters in scope `TMOffDynamicsRP GABA` have the analogous relation to the dynamics of the inhibitory conductance $g_{in}$ (see eqs. ?? and ??).

Many parameters are specified not by a single value but by a vector of four values to define individual parameters for each member of a population. Parameter `refRel` (in scope `popPe`), for example, specifies that the parameter $R_r$ of our neuron model (see eq. ?? in section ??) is distributed according to a Gaussian with mean 3, standard deviation 0.5, but limited to the interval $[1.75; 4.25]$. In contrast, if the standard deviation is 0 and/or the left interval border larger than the right one then the parameter is the same for all members of the population (see parameter `tau_x`, for example).

### 4.3.3 Compiling and running simulations

In sections 4.3.1 and 4.3.2 we have discussed how a Felix++ simulation program and the corresponding parameter file should be structured. Here we explain how one obtains an executable program from the source file. This process is illustrated in Figure 4.4.
Figure 4.4: Compilation and Linkage of a Felix++ simulation program. The simulation source file includes Felix++ and/or Felix headers (cf. Fig. 4.1). The source file SIMULATION.cpp then is compiled by the command “Felix2 SIMULATION” which generates an executable SIMULATION. Running the executable SIMULATION requires the parameter file and dynamic (“shared”) libraries of Felix++, Felix, and the X system.

A simulation source file named SIMULATION.cpp can be compiled by the command “Felix2 SIMULATION”. Felix2 is a script that compiles the source file and sets the correct include- and link-paths by calling the Makefile. Compiling using Felix2 yields as output the executable SIMULATION. When running this requires the dynamic (or shared) libraries as shown in Figure 4.4.
Part II

The Graphical User Interface (GUI) of Felix
Chapter 5

The program structure of GUI simulations
CHAPTER 5. THE PROGRAM STRUCTURE OF GUI SIMULATIONS
Chapter 6

Using the GUI elements of Felix
Part III

Fundamentals of Felix2
Chapter 7

Type conventions
Chapter 8

Layouts and multi-dimensional arrays
Chapter 9

Parameters and parsing
Chapter 10

Parameters and parsing
Chapter 11

Basic numerics

11.1 Time

11.2 Random generators

11.3 Constants, functions, and look-up-tables
Chapter 12

Ports
Chapter 13

Kernels
Chapter 14

Simulation environment and components
Part IV

Modelling the environment: input and output
Chapter 15
Vectors and patterns
Chapter 16

Modelling objects and the space around
Chapter 17

Recording of simulation data
Part V

Further elements of Felix2
Chapter 18

Integrators for differential equations
Chapter 19

Delays
Chapter 20

Receptors
Chapter 21

A library for associative memory
CHAPTER 21. A LIBRARY FOR ASSOCIATIVE MEMORY
Part VI

Components of Felix2
Chapter 22

Noise populations

22.1 TMUniformNoise: the standard noise population
22.2 TMCorrelatedNoise: noise correlated in space and time
Chapter 23

Neuron populations

23.1 IFNeuron: a simple integrate-and-fire neuron model
23.2 SGNeuron:
23.3 SSNeuron:
23.4 SSCOscillator:
23.5 InpNeuron:
Chapter 24

Connections

24.1 TopoConnection:

24.2 GaussConnection:

24.3 BlankTopoConnection:

24.4 DemoBlankTopoConnection:

24.5 AssoConnection:

24.6 V1Connection:

24.7 RandomConnection:

24.8 DelayKernelConnection:
Chapter 25

Learner

25.1 STDPLearner:
Part VII

Simulation examples
Chapter 26

Integrating Felix2 and the GUI of Felix1
Chapter 27

A simple network of oscillating neurons
Part VIII

Appendices
Appendix A

The GUI reference of Felix
Appendix B

The C++ classes of Felix2
Appendix C

Parameter scopes for Felix2 components

This appendix contains descriptions of the parameter scopes for all Felix2 components. The descriptions consist of example parameter scopes and a brief explanations of its use and the role of the parameters.

C.1 Noise populations

C.2 Neuron populations

C.2.1 Class TIFNeuron

The following example is taken from the simulations in []. The description of the TIFNeuron model should be clear by the comments. For more details see section ?? on page ??.

```csharp
#{ IFNeuron TrigExNeuron
tau_x : 0.5 % time constant of membr.potential
tau_gex : 0.25 % time constant excit.conductance
tau_gin : 0.25 % time constant inhib.conductance
tau_h : 1 % time constant of habituation
gexMax : 100 % maximal excitatory conductance
ginMax : 100 % maximal inhibitory conductance
theta : 10 % (asymptotic) threshold
ahp : 108 % after-hyperpolarization (reset x)
habituation : 0 % habituation (increase of h)
exConstInput : 0 % excitatory constant input
inConstInput : 10.45 % inhibitory constant input
noise(power,order) : 0 0 % noise power and order
eInLen : 20 % length excit. spike input queue
iInLen : 20 % length inhib. spike input queue
eInGradLen : 20 % length excit. gradual inp.queue
iInGradLen : 20 % length inhib. gradual inp.queue
}
```
C.2.2 Class TSSNeuron

The TSSNeuron model consists of basic neuron parameters similar to the TIFNeuron (see section C.2.1 on page 93). In contrast, however, input parameters are specified via an TM-CompartmentReceptors container (see section ?? on page ??). Thus one can define an arbitrary number of receptors where spikes or gradual input can be directed to (e.g. AMPA-, GABA-, or NMDA- receptors). Another difference to the TIFNeuron is the more detailed refractory mechanism. The following example is taken from the simulations in [ ].

```plaintext
#{ SSNeuron popPe
  tau_x(exp,sig,min,max) : 10 0 1 0
  theta(exp,sig,min,max) : 10 0 1 0
  refAbs(exp,sig,min,max) : 2 0 1 0
  refRel(exp,sig,min,max) : 3 0.5 1.75 4.25
  tau_h(exp,sig,min,max) : 150 0 1 0
  thetaInc_h(exp,sig,min,max) : 0.6 1.0 0.2 1.0
}

#{ TMCompartmentReceptors receptors
  nNoiseSources : 0
  nReceptorPorts : 2

  noiseIndices : size : 2
    DEFAULT DEFAULT

  #} TMOffDynamicsRP AMPA
    tau_OFF(exp,sig,min,max) : 5 0 1 0
    E(exp,sig,min,max) : 80 0 1 0
    g0(exp,sig,min,max) : 0 0 1 0
    powerInpNoise(exp,order,sig,min,max) : 0.025 1 0 1 0
    pSynNoise(exp,sig,min,max) : 0.5 0 1 0
    cSynNoise(exp,sig,min,max) : 0.05 0 1 0
    qLen : 600
    qLen_gradual : 100

  noiseIndices : size : 2
    DEFAULT DEFAULT

  #} TMOffDynamicsRP GABA
    tau_OFF(exp,sig,min,max) : 7 0 1 0
    E(exp,sig,min,max) : -10 0 1 0
    g0(exp,sig,min,max) : 0 0 1 0
    powerInpNoise(exp,order,sig,min,max) : 0.02 1 0 1 0
    pSynNoise(exp,sig,min,max) : 0.5 0 1 0
    cSynNoise(exp,sig,min,max) : 0.05 0 1 0
    qLen : 100
    qLen_gradual : 100

  }

The basic neuron model parameters are the decay time constant $\tau_x$ of the membrane potential, the asymptotic threshold $\theta$, the absolute refractory period $\text{refAbs}$, the relative refractory period $\text{refRel}$, the decay time constant $\tau_h$ of the habituation, and the
threshold increase per spike $\theta_{\text{Inc}, h}$. All these parameters can be defined to be distributed for the neuron population according to a Gaussian with expectation $\exp$ and standard deviation $\sigma$ where the values are restricted to the interval $[\min, \max]$. For $\min \geq \max$ all neurons will have the same parameter value $\exp$. For further details of the neuron model see section ?? on page ???. For the parameter scope of TMCompartmentReceptors see section ?? on page ??.

C.3 Connections

C.3.1 Template TMGaussConnection

The template class TMGaussConnection is a topographic connection derived from TMTopoConnection (see section ?? on page ??) where the kernels are simply Gaussians of an arbitrary dimension. The parameters of a TMGaussConnection can be divided into five parts, where (i) the scope weightsMapping (cf. section ?? on page ??) defines the mapping of the weight kernels, i.e. which neuron uses which afferent or efferent weight kernel, (ii) the scope weightsParameters (cf. section ?? on page ??) defines the weight kernels, i.e. the weights of each kernel are according to a certain Gaussian, (iii) the scope delaysMapping (cf. section ?? on page ??) defines the mapping of the delay kernels which is similar to the mapping of the weight kernels, (iv) the scope TMKernelDY (cf. section ?? on page ??) defines the delays, and (v) the strength parameters define factors which are multiplicated to the kernel weights when a spike is propagated, for example. The following example is taken from the simulations in [ ] and is as simple as possible.

```cpp
#{ GaussConnection linkCorrNoisePePs
    #{ weightsMapping wm
        direction : EFFERENT
        kernelMap :
            size : d2(1 1)
            0
    }
    #{ weightsParameters wp
        exp :
            size : 1 2
            0.0 0.0
        sig :
            size : 1 2
            10.0 10.0
        phi :
            size : 1 2
            0.0 0.0
    }
    #{ delaysMapping dm
        direction : EFFERENT
        kernelMap :
            size : d2(1 1)
            0
    }
    #{ TMKernelDY kernelDelay1
        nModes : 1
        baseDelay : 0
```
distDelay  : 0  
straightDist : 0  
distType     : RECT  
noiseType    : FIXED  
powerDelayNoise : 0  
minDelayNoise : 0  
maxDelayNoise : 10  
limitsDelayNoise : 1 1
}

strength p[0;1000]/0.001: 1.5
}

This simple example uses a single two-dimensional non-rotated (phi=(0,0)) Gaussian weight kernel with a centered peak (exp=(0,0)), and standard deviation (s.d.) 10 in both dimensions. Similarly it uses a single unimodal delay kernel.

In the following variant there are used two different weight kernels, one with s.d. 10, the other with s.d. 5. Further a special mapping of the weight kernels is used.

#{ weightsMapping wm
  direction : EFFERENT
  kernelMap :
    size : d2(3 4)
    0 0 1 0
    0 1 0 0
    1 0 0 0
  }
#{ weightsParameters wp
  exp :
    size : 2 2
    0.0 0.0
    0.0 0.0
  sig :
    size : 2 2
    10.0 10.0
    5.0 5.0
  phi :
    size : 2 2
    0.0 0.0
    0.0 0.0
  }

The 3x4 mapping of scope weightsMapping can be adapted to an arbitrary-sized two-dimensional neuron population by repeating the 3x4 blocks in both dimensions. The weights mapping specifies how many different kernels are actually needed (one plus the largest index in the index block - 2 in this example). Accordingly, the parameters in scope weightsParameters must contain for each kernel a separate line.

For multi-modal delays (see section ?? on page ??) the corresponding scopes would look like this:

#{ TMKernelDY kerneldelay1
  nModes : 2
  baseDelay : 4 7
}
Note that for an m-modal connection the strength parameter must be a m-dimensional vector. It is possible that one TMGaussConnection is used in a multiplexing fashion. For example, if the connections from presynaptic neuron population $A$ to the postsynaptic connections $B$ and $C$ (or two different receptor ports of the same postsynaptic population) are identical, then it is not necessary to propagate a spike in a $A$-neuron two times through two different TMGaussConnections. Instead it is possible to call the constructor of TMGaussConnection with two (or more) postPorts arguments (see section ?? on page ??). However, then for $p$ postPorts there are also $p$ strength parameters required, one line for each. For more details about TMGaussConnection see section ?? on page ??.

### C.3.2 Template TMAssoConnection

The template class TMAssoConnection is derived from a topographic or full connection derived from TMBlankTopoConnection (see section ?? on page ??) where the kernel are usually generated by Hebbian learning of a priori defined patterns. The following is a simple example from the simulations in [].

```plaintext
AssoConnection assoLinkC
flagEllipsoidConstraint : 0
threshEllipsoidConstraint : 1
flagMask : 1
dexpMask : 0 0
sigMask : 200 200
phiMask : 0 0
nDelayKernels : 1
flagZeroCenteredDelays : 0

TMKernelDY kernelDelay1
nModes : 1
baseDelay : 0.8
distDelay : 0.05
straightDist : 0
distType : RECT
noiseType : FIXED
powerDelayNoise : 0
minDelayNoise : 0
maxDelayNoise : 10
limitsDelayNoise : 1 1
```
The flag flagEllipsoidConstraint determines if only the ellipsoid of the weight kernels are used (if set), or if the whole cuboid is used (if unset). If set, then threshEllipsoidConstraint determines how the ellipsoid is defined (see section ?? on page ??). The flag flagMask determines if the kernels are additionally multiplied by a mask for example to express weight decay with increasing distance from the center. If the flag is set the Gaussian mask kernel is defined by the parameters expMask, sigMask, and phiMask. nDelayKernels determines how many different delay kernels are generated that are randomly mapped to the neuron population. And the flag flagZeroCenteredDelays determines if the delays in the delay kernels are represented zero-centered or not (see section ?? on page ??). Parameter scope fullDimFlags determines which dimensions are topographic and which dimensions are full (see section ?? on page ??), and scope TMKernelDY specifies the delays (see section ?? on page ??). Finally parameter strength specifies a global factor multiplied to the weights of the kernels when presynaptic input is actually propagated to the postsynaptic target (cf. section ?? on page ??).

C.4 Learner