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**CNS: a GPU-based framework for  
simulating cortically-organized networks**  
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# CNS: a GPU-based framework for simulating cortically-organized networks

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## Abstract

*Computational models whose organization is inspired by the cortex are increasing in both number and popularity. Current instances of such models include convolutional networks, HMAX, Hierarchical Temporal Memory, and deep belief networks. These models present two practical challenges. First, they are computationally intensive. Second, while the operations performed by individual cells, or units, are typically simple, the code needed to keep track of network connectivity can quickly become complicated, leading to programs that are difficult to write and to modify. Massively parallel commodity computing hardware has recently become available in the form of general-purpose GPUs. This helps address the first problem but exacerbates the second. GPU programming adds an extra layer of difficulty, further discouraging exploration.*

*To address these concerns, we have created a programming framework called CNS ('Cortical Network Simulator'). CNS models are automatically compiled and run on a GPU, typically 80-100x faster than on a single CPU, without the user having to learn any GPU programming. A novel scheme for the parametric specification of network connectivity allows the user to focus on writing just the code executed by a single cell. We hope that the ability to rapidly define and run cortically-inspired models will facilitate research in the cortical modeling community. CNS is available<sup>1</sup> under the GNU General Public License.*

## 1. Introduction

### 1.1. Definitions

For the purpose of this paper, we define a 'cortical' model to be a network model which consists of some number of

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<sup>1</sup>Software and a full programming manual are available at <http://cbcl.mit.edu/jmutch/cns>.

N-dimensional 'layers' of cells, where each layer encodes some N-D feature space. It is common for at least some of the dimensions to be topographically mapped, meaning that physical proximity in the layer corresponds to proximity in the feature space. N, and the feature space, can be the same or different from layer to layer. All the cells in a layer must be of the same type, i.e., each maintains its own values of the same set of variables, using the same algorithm. Connectivity between cells may be completely arbitrary, but often consists of a repeating pattern.

This class of models includes, but is by no means limited to:

- Convolutional networks [5, 4].
- HMAX [9, 8].
- Hierarchical Temporal Memory [1].
- Deep belief networks [2].
- Detailed spiking models of cortex.

Some of these are 'static' models, requiring only a single pass through the network for a given input. Others have dynamics that require iteration over many time steps.

### 1.2. Motivation

Cortical models commonly contain a large number of units and are therefore computationally expensive. However, they are highly amenable to parallelization. The layered structure of cortical models maps very well to the architecture of modern GPUs, which are optimized to perform the same operation at every point in an array of data. GPUs have evolved from graphics accelerator cards, where the array elements are pixels. Over the last few years the APIs for these cards have opened up to encourage the acceleration of non-graphical algorithms that can benefit from the same architecture. Current GPUs have hundreds of parallel processors and can typically run suitable algorithms 80-100x faster than a single CPU. However, these performance

gains are not free. A GPU's processors all still share a common memory, which becomes the bottleneck of the system. To achieve optimal performance, GPU programmers must code their algorithms so that the processors access memory in coordinated patterns. This adds another layer of difficulty to models that can already be somewhat challenging to program.

Individual cells in cortical models typically perform fairly simple functions. Difficulties in programming these models usually arise in keeping track of the connectivity between cells. There are two common approaches:

- **Enumerate every synapse.** With this approach, one can represent any network architecture, but for networks having regular patterns of connectivity, it is extremely wasteful. Memory usage increases drastically, and because memory access is the bottleneck in multiprocessing systems, so does processing time.
- **Matrix operations.** This is a common abstraction that works for some simple hierarchical models. Layer  $n$  is generated from layer  $n - 1$  via some matrix operation such as N-D convolution. This is very space-efficient and easy to implement in a language such as MATLAB. It is also fairly straightforward to implement on a GPU, and to program arbitrary response functions in place of the dot product implied by convolution. However, in more complex situations this abstraction becomes limiting. Actual cortical areas typically receive convergent input from multiple pathways that have been processed differently. Operations may have been carried out at different resolutions, or the number of processing steps may have been different. Thus, the indices of cells in two different input layers no longer have the same meaning, and it is difficult to define a matrix operation that combines them. Most work in cortical modeling has simply focused on the subset of models for which these difficulties do not arise.

Of the two barriers to exploration of the cortical model space – programming time and run time – GPU programming alone can only address the latter, and at the expense of the former. A framework is needed that can address both issues.

### 1.3. Overview of CNS

CNS is a rapid development environment for cortical models. Models are compiled and run automatically on NVIDIA GPUs, often running 80-100x faster than on a single CPU, without the user having to learn any GPU programming.

Most aspects of a model are defined via MATLAB scripts, including:

- The number of layers.

- The dimension, size, and cell type of each layer.
- The variables associated with each cell type, and their initial values.
- The connectivity between cells.

The process of running models, loading input data, and pulling back results is also controlled via MATLAB. Variables are referred to by name – the user does not need to be concerned with where they are stored in GPU memory.

There are two options for specifying connectivity. As in many other simulators, synapses may be explicitly enumerated: a given cell can list any number of presynaptic cells, which can be in any layer. For models having regular patterns of connectivity, CNS uses a scheme in which each cell explicitly retains its grid position (the center of its receptive field) in a real-valued feature space which is meaningful across layers. For example, in a vision model, the dimensions of this common feature space would probably include retinal position, and even cells several steps removed from the input would still know their center coordinates in retinal space. Under this scheme, cells can infer their inputs based on proximity in the common feature space. By appropriate relative arrangement of the coordinate grids for each layer, any of the standard connectivity patterns (e.g., valid and full convolution, sub- and super-sampling, etc.) can be achieved, as well as many others. Once defined in this way, connectivity is handled by the framework. An individual cell can make requests of the framework, for example, it can request the indices of its  $n$  nearest neighbors in layer  $z$ . This division of labor allows programmers to focus mainly on the code being executed by a single cell.

A 'kernel' is the code each cell executes during an iteration of the network. Cells of the same type share the same kernel. Kernels run on the GPU and are the only parts of a CNS model that must be written in C/C++. Even when writing a kernel, however, the programmer remains isolated from the complexities of the GPU. A kernel is written from the point of view of a single cell, so the programmer is not responsible for any thread scheduling. Nor is the programmer required to know where variables are stored in GPU memory. CNS provides named macros that the kernel can call to, among other things:

- Read and write the current cell's variables.
- Find input cells via proximity in the feature space, as described above.
- Read the values of other cells' variables.

During model development, CNS can be thought of as a compiler, in that programmers do not need to read or modify any of CNS's code. By way of comparison, the 3-D convolutional network package with backpropagation described in section 2.3, written using CNS, comprises about

300 lines, while CNS itself is about 10,000 lines as of this writing.

CNS is licensed under the GNU General Public License. The software and programmer's manual are available at <http://cbcl.mit.edu/jmutch/cns>.

## 1.4. Structure of this paper

The remaining sections of this paper are as follows:

- Section 2 introduces three example packages written in CNS.
- Section 3 discusses some CNS concepts in more detail, using the example packages for illustration.
- Section 4 describes the process of developing and running a CNS model.
- Section 5 provides some internal details on how CNS maps models to the GPU architecture.
- Section 6 lists some limitations of CNS.
- Section 7 outlines future work on CNS.

This paper is intended as a high-level introduction to CNS. Exact syntax is not covered beyond a few examples. For a complete description of all CNS syntax and options, see the programmer's manual [6].

## 2. Example packages

In CNS, a 'package' is a collection of cell types that is used to construct models. For example, the 'HH' package described below implements several types of Hodgkin-Huxley cells. Once you have a package, you can then build networks consisting of those types of cells. The same package can be used to define many specific models, having different numbers of layers, of different sizes, with different connectivity, etc.

In this section we briefly describe three packages that we have developed with CNS. Subsequent sections will refer back to these examples to illustrate various CNS concepts.

### 2.1. 'HH' package: Hodgkin-Huxley spiking models

The Hodgkin-Huxley (HH) package is used to build models made up of 2-D layers of spiking neurons, similar to the cortical laminae. Figure 1 shows part of three layers of such a model:

- IN: the input layer. Each cell has its own list of preprogrammed spike times.
- F1: a layer of fast-spiking inhibitory cells, each receiving input from a local region of IN.
- P1: a layer of pyramidal cells, each receiving input from a local region of IN and F1 (latter connections not shown).

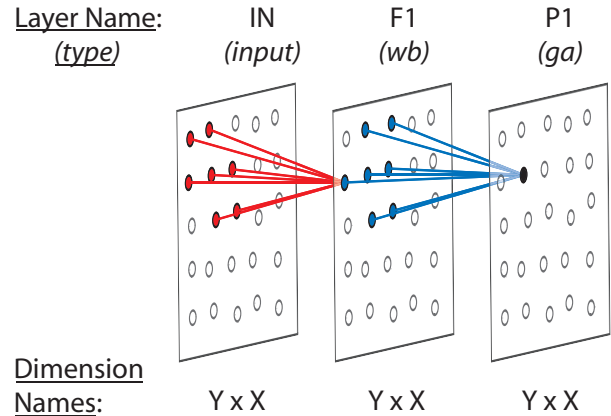


Figure 1. Schematic of part of a model made up of Hodgkin-Huxley spiking neurons (see section 2.1), implementing monosynaptic excitation with disynaptic inhibition. IN: input layer. F1: fast-spiking inhibitory cells (Wang-Buzsaki model). P1: pyramidal cells (Golomb-Amitai model). IN-P1 connections, and other layers, are not shown. The full model contained 10,000 cells.

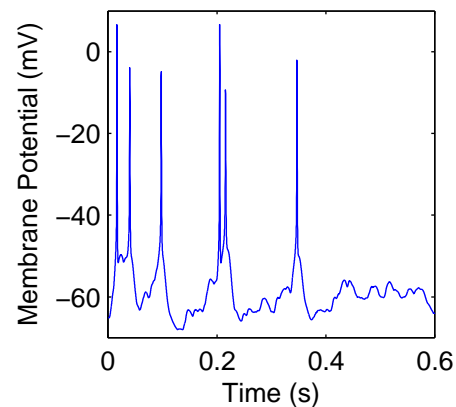


Figure 2. Voltage trace output for one cell in a Hodgkin-Huxley model (see section 2.1).

In these models, cells in a layer are assigned to grid positions in a 2-D feature space, but with some gaussian noise added. Connectivity is also noisy; for example, each F1 cell receives input from a local region of IN cells, but with the probability of a connection decreasing with distance from the center of the region. Thus, connectivity in these models is defined using the explicit synapses method: each cell in a model lists its presynaptic cells.

Models built with this package are dynamic. Each cell maintains its own set of Hodgkin-Huxley state variables. Each iteration of the network represents a small time step, during which each cell polls its presynaptic cells to see which ones are spiking and updates its state variables according to a set of differential equations. CNS can track the values of selected state variables as they change; figure 2 shows a voltage trace for one cell.

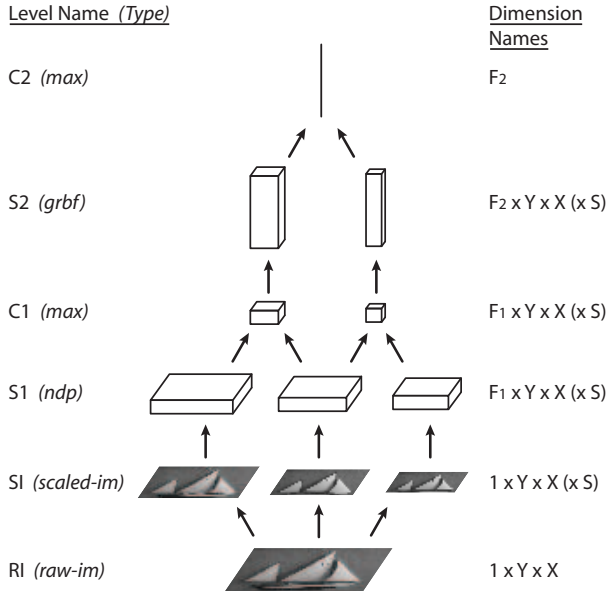


Figure 3. An example of an HMAX model (see section 2.2). Each step is performed at multiple scales, only three of which are shown here.

For a model with 10,000 neurons and 330,000 synapses, CNS was able to process 5,000 time steps per second on a GTX 285 GPU.

## 2.2. 'FH' package: HMAX-like feature hierarchies

The FH package allows you to build models of the HMAX [9, 8] class. HMAX models the initial feedforward stage of object recognition in the ventral visual pathway. It extends the idea of simple and complex cells [3] to form a hierarchy in which alternating template matching and max pooling operations progressively build up both feature selectivity and invariance to position and scale. Figure 3 illustrates one such model [8] as it is expressed in CNS.

HMAX is scale invariant: each stage of processing is carried out at multiple scales. Thus, each stage of the model is actually represented by many CNS layers, one for each scale. For the model in figure 3, the stages are:

- SI: Many different resolutions of the original image, produced by bicubic interpolation.
- S1: Applies gabor filters of several orientations at every position and scale. Each scale is now of size  $F_1 \times Y \times X$ , where  $F_1$  is the number of orientations.
- C1: Independently for each orientation, computes the maximum response over a local area of position and scale. Also subsamples by a factor of 5. Note that the operation of pooling over multiple scales would be difficult to define as a matrix operation. Here it is done using CNS's common coordinate scheme (see section 3.6).

- S2: Computes the response to many stored templates at every position and scale. Now in each scale we have  $F_2 \times Y \times X$  units, where  $F_2$  is the number of templates.
- C2: For each template, finds the maximum response over all positions and scales. This results in a feature vector which can be fed into a classifier.

These models are computed in a single bottom-up pass for a given input image. The gabor filters in S1 and the stored templates in S2 are shared by all the cells in those layers (and are not shown in figure 3).

The FH package replaces an older CPU-based library called FHLib [7] which was used for the experiments in [8]. For one large model, CNS outperformed FHLib by a factor of 97x on a GTX 285 GPU.

## 2.3. 'CN' package: convolutional networks for 3-D image segmentation

The 'CN' package is a CNS reimplementation of the code used in [4]. It is used to train and run convolutional networks that perform segmentation in 3-D electron microscope images of brain tissue. These networks are similar to those of Lecun [5] except that the filters have three spatial dimensions and there is no subsampling. The model architecture is illustrated in figure 4.

Each  $X_i$  layer is four-dimensional, containing the value of  $F_i$  different features at each position in a 3-D cube. (For the input image  $X_0$  there is only one feature: the pixel value.) The features are different in each stage. Each  $W_i$  can be viewed as  $F_i$  four-dimensional filters which, via convolution over  $X_{i-1}$  (and then adding the appropriate bias from  $B_i$ ), produce layer  $X_i$ . Note that convolution only occurs over the three spatial dimensions; for the feature dimension,  $X_{i-1}$  and the filter are the same size.

The goal of the training phase is to learn all the  $W_i$  and  $B_i$  layers. This is done via the backpropagation algorithm in three passes (mathematics in [4]):

- Forward pass: starting with the input in  $X_0$ , compute the outputs of each  $X_i$  layer from bottom to top.
- Backward pass: starting with the desired output in  $X_n$ , compute the error term ('sensitivity') for each  $X_i$  from top to bottom.
- Weight update: update each  $W_i$  and  $B_i$  layer.

Note that unlike the filters and features in FH models (section 2.2), here the weights and biases need to change during a network iteration. Thus, we treat them as layers, and they have their own kernels which perform the update at the appropriate time.

Once the network has been trained, only the forward pass is needed to perform segmentation.

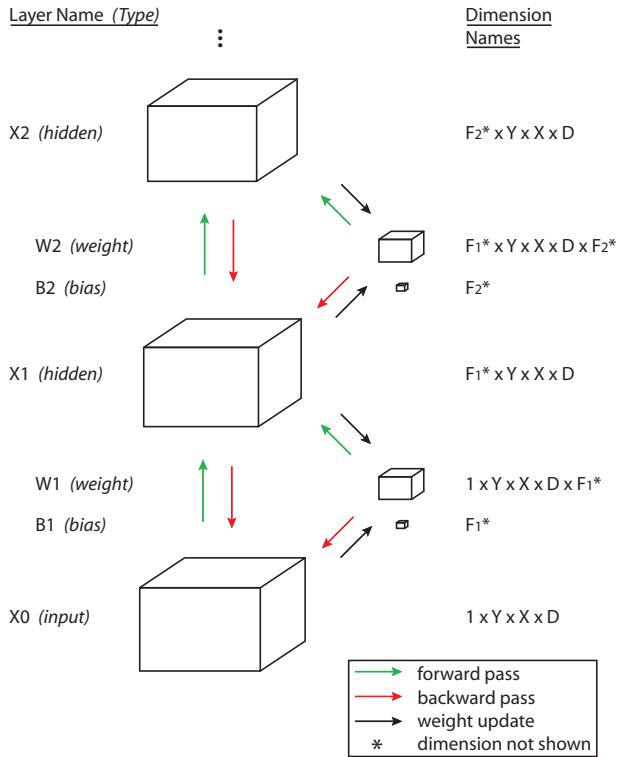


Figure 4. The first three stages of a convolutional network for segmenting 3-D images (see section 2.3).

The CNS implementation of this class of models ran about 100x faster than the previous code, based on timing using a single CPU. Ongoing development of these models continues under CNS.

### 3. CNS concepts

In this section we run through the key concepts of CNS, using the packages in section 2 as examples.

#### 3.1. Layers and groups of layers

The basic architectural unit of a CNS model is the *layer*, which is an N-dimensional array of cells that are all of the same *type* (see section 3.3). Practically speaking, a model can have up to several hundred layers. Note that layers do not have to be arranged in any kind of hierarchy or directed acyclic graph (DAG).

Multiple layers of the same type can be designated as a *group*. The main purpose of this is to allow them to share some common data. For example, in the FH package, all the S1 layers (of different scales) are designated as a group so they can share the same set of precomputed gabor filters.

#### 3.2. Execution order

By default, a single network iteration consists of computing all cells in all layers exactly once, in parallel. HH models work like this. Internally, of course, everything isn't computed exactly at once. A double-buffering system is employed to guarantee that during iteration  $t$ , all the values a kernel reads from other cells come from iteration  $t - 1$ .

For some models, it wouldn't make sense to compute all the layers at the same time. For example, cells in FH models do not have states that evolve over time. There is no point in computing layer  $n$  until layer  $n - 1$  has been computed. In this case, a network iteration is broken into steps, with layers assigned different step numbers. In figure 3, for example, all the SI layers are assigned step 1, the S1 layers step 2, the C1 layers step 3, etc.

CN models are similar to FH, except that a single iteration consists of a forward pass, a backward pass, and a weight update. The  $X_i$  layers get computed at two different points in one iteration, i.e., they each have two different step numbers, and perform a different computation in each.

Note that there is no unit of execution smaller than a single layer.

#### 3.3. Cell types

A cell type defines:

- The dimensionality (N) a layer of cells of that type must have. For example, all the layers in HH models are 2-D, but in CN models the  $X_i$  layers are 4-D, the  $W_i$  layers are 5-D, and the  $B_i$  layers are 1-D.
- The constants and variables (*fields*) associated with each cell. (Fields can also have layer scope, synapse scope, etc.; see section 3.4.)
- The kernel used to update a cell during a network iteration (see section 3.5).

Just like a class in object-oriented programming, a cell type can be a subtype of a parent type. All the cell types in a package form a type hierarchy (not to be confused with a model having hierarchical structure). Every package must have a 'base' type which is the root of the type hierarchy. A cell type inherits the following properties from its parent type:

- The dimensionality of a layer. Note that this cannot be overridden.
- All fields.
- The kernel. This can be overridden.

Cell types can be *abstract*, which means they exist only to declare a common set of properties that are then inherited

Type	Fields			Kernel	Dimensions
	Layer or group	Cell	Synapse		

### HH package

base*		<i>V<sub>m</sub> t<sub>last</sub></i>		no	Y x X
input		(spike times)		yes	
passive*	C <sub>m</sub> g <sub>L</sub> E <sub>L</sub>		<i>g r</i>	template	
dendrite				parts	
active*	E <sub>Na</sub> E <sub>K</sub>			no	
ga	<i>g<sub>NaBar</sub> g<sub>KABar</sub> ...</i>	<i>h n b z</i>		parts	
wb	<i>g<sub>NaBar</sub> g<sub>KABar</sub> ...</i>	<i>h n</i>		parts	

### FH package

base*		<i>val</i>		no	F x Y x X
raw-im				no	
scaled-im				yes	
s-fixed*	(filters)			template	
ndp				parts	
c*	(rf size)			template	
max				parts	
avg				parts	
s-learned*	(features)			template	
grbf	sigma			parts	
grbfnorm	sigma			parts	

### CN package

base*				no	undefined
layer*		<i>val</i>		no	F x Y x X x D
input				no	
computed*		<i>sens</i>		template	
hidden				parts	
output		correct		parts	
weight	eta	<i>val</i>		yes	F <sub>in</sub> x Y x X x D x F <sub>out</sub>
bias	eta	<i>val</i>		yes	F <sub>out</sub>

Table 1. Cell types for the example packages. Indentation in the 'type' column denotes inheritance, and \* denotes an abstract type. Italicized fields are read-write, i.e., variable. Note that a cell type inherits all its parent type's fields.

by subtypes. In many packages, the 'base' type will be abstract. You cannot create layers of cells of an abstract type; you must use a non-abstract subtype.

Table 1 shows the type hierarchy for all three example packages.

### 3.4. Fields and scope

In CNS, a named numeric quantity associated with a cell is called a *field*. For example, in HH models, membrane voltage ( $V_m$ ) is a field, and each cell maintains its own value of the membrane voltage. For cells, fields are analogous to the data members of a class or structure in object-oriented programming. Each cell type inherits its parent type's fields and may also define its own fields. The definition of a field includes:

- Field name. Used to identify the field, both in the MATLAB interface and inside kernels.
- Data type: either single-precision floating point or 32-bit signed integer.
- Scalar or vector.
- Read-only or read-write. Read-only fields (constants) cannot change their values during a model iteration on the GPU, but they can be changed between iterations from within MATLAB. Read-write fields (variables) may be changed during a model iteration, but only for the current cell, i.e., each cell is responsible for updating its own variables.
- Default value: the value a field will have when a model

is initialized. May be overridden at model initialization time. If no default value is defined, then one must be given at model initialization time.

- Scope: see below.

One key way in which CNS fields are *not* like data members in object-oriented programming is that you can also define their *scope*. A field can have:

- Cell scope. Each cell maintains its own value of the field. (If the field is a vector, each cell maintains its own vector value. This applies to all the scopes listed here.)
- Synapse scope. Available when using explicit-synapses connectivity (section 3.6). A separate value of the field is maintained for each synapse. This can obviously take a lot of memory.
- Layer scope. Each layer will have one value of the field that applies to the whole layer. Such fields are always read-only.
- Group scope. One (read-only) value of the field is maintained for an entire group of layers (see section 3.1).
- Model scope. One (read-only) value of the field is maintained for an entire network model.

Table 1 shows many of the fields defined by the example packages.

One more kind of field is also available. In some models it is desirable for many cells to have access to the same large multidimensional array of static data. One way of doing this would be to use the same approach the CN package uses to share  $W_i$  and  $B_i$  values, which is to make such arrays into layers of their own. In CN models,  $W_i$  and  $B_i$  are *not* static, so this is the only option. In the FH model shown in figure 3, however, the S1 stage's gabor filters and the S2 stage's stored features are static during a network iteration, so it would be nice to not have to complicate the network structure by making them separate layers. CNS allows static arrays like this to be defined as fields having layer, group, or model scope.

### 3.5. Kernels

A *kernel* is a function that updates a cell's variables during a network iteration. As with methods in object-oriented programming, a kernel is written from the point of view of a single cell. Unlike object-oriented programming, however, each cell type has exactly one kernel.

Kernels are written using a limited subset of C++ (mostly just C), supplemented by macros generated by CNS. You may provide an arbitrary block of C/C++ code, subject to these conditions:

- Only statements that are permissible in function scope (inside a function body) are allowed.
- The only standard library functions available are those in the 'math' library.
- No dynamic memory allocation is permitted.

Access to fields and other properties you have defined for the various cell types is done via macros that CNS provides. This spares you from having to worry about how data is laid out in GPU memory, etc. Figure 5 shows part of a kernel from the CN package; all the red symbols are macros generated by CNS, based on the definitions for cell types in that package.

When you are writing a kernel, you can ask CNS to list all the macros available to you. CNS provides macros to:

- Read and write fields. (For example, the READ\_\* macros, WRITE\_VAL, ZP, ZW, and ZB in figure 5.)
- Retrieve current cell coordinates. (For example, THIS\_F in figure 5.)
- Find receptive fields. (For example, the \*\_RF\_NEAR macros in figure 5.)
- Retrieve layer dimension sizes. (For example, the \*\_SIZE macros in figure 5.)
- Loop through explicit synapses, if any.
- Retrieve the current iteration number and other miscellaneous.

Closely related kernels are often very similar. For example, one might want to define a cell type having a kernel like the one in figure 5, but with the dot product operation, or perhaps the sigmoid nonlinearity, replaced by something else. To avoid having many kernels that are slightly-modified copies of one another, CNS allows you to write a *template* kernel for an abstract parent type. The template kernel will contain most of the logic, leaving 'blanks' for the specific operations. Then the various subtype kernels provide different code snippets (*parts*) that 'fill in' those blanks for each subtype. This feature is used in all three example packages; see the 'kernel' column in table 1.

### 3.6. Connectivity

As discussed in sections 1.2 and 1.3, CNS has two ways to specify cell-to-cell connectivity:

- HH models use the explicit synapses method: each cell lists each of its presynaptic cells. This is necessary because cells in HH models do not occupy regular grid positions in a common feature space.



```

// Retrieve the size of our 4-D filter.
int fSize = WEIGHT_F_SIZE(ZW);
int ySize = WEIGHT_Y_SIZE(ZW);
int xSize = WEIGHT_X_SIZE(ZW);
int dSize = WEIGHT_D_SIZE(ZW);

// Find our corresponding RF in the previous layer.
int y1, x1, d1, dummy;
GET_LAYER_Y_RF_NEAR(ZP, ySize, y1, dummy);
GET_LAYER_X_RF_NEAR(ZP, xSize, x1, dummy);
GET_LAYER_D_RF_NEAR(ZP, dSize, d1, dummy);

// Compute response to the 4-D filter.
float v = 0.0f;
for (int f = 0; f < fSize; f++) {
    for (int k = dSize - 1, d = d1; k >= 0; k--, d++) {
        for (int j = xSize - 1, x = x1; j >= 0; j--, x++) {
            for (int i = ySize - 1, y = y1; i >= 0; i--, y++) {
                float p = READ_LAYER_VAL(ZP, f, y, x, d);
                float w = READ_WEIGHT_VAL(ZW, f, i, j, k, THIS_F);
                v += p * w;
            }
        }
    }
}
v += READ_BIAS_VAL(ZB, THIS_F);

// Apply sigmoid nonlinearity.
v = 1.0f / (1.0f + expf(-v));

// Store result.
WRITE_VAL(v);

```

Figure 5. Part of a kernel from the CN package. This code performs the forward pass. It computes the value of a single cell in an  $X_i$  layer by overlaying the appropriate 4-D weight array from  $W_i$  onto the cell's 4-D receptive field in  $X_{i-1}$ , computing the dot product, adding the appropriate bias from  $B_i$ , and applying a sigmoid nonlinearity. Symbols in **RED** are macros generated by CNS; everything else is standard C/C++. See section 3.5 for details.

- FH and CN models use the common coordinates method. Here we expand a bit on that topic.

In FH models, the dimensions Y and X correspond to retinal position and are meaningful across all layers. In CN, the same holds in the  $X_i$  layers for spatial dimensions Y, X, and D (depth). Thus, cells in both packages are assigned to real-valued grid positions in these common dimensions. Figure 6 illustrates how these coordinates are used to perform an operation (pooling over scales that are not an integer multiple of one another in an FH model) that is difficult to define without them.

Note that this is just one instance of a general problem which can arise in several ways, and which CNS's common coordinate system was designed to solve: how does one define 'local' operations like convolution over multiple layers

having different sizes, resolutions, etc.? Networks having multiple pathways that converge will almost always have this issue, since the different pathways probably involve different amounts of convolution (with edge loss) and/or subsampling. And of course, convergence occurs throughout cortex.

Part of defining a network model in CNS is setting up these grid positions for the different layers. There is a function (`cns_mapdim`) that handles all the standard cases: valid, same, and invalid convolutions, subsampling, super-sampling, etc. Once grid positions have been assigned, connectivity is handled by CNS - in kernel code, you simply call a macro requesting, for example, the nearest  $n$  neighbors in some layer  $z$ , or every cell within radius  $r$ .

Not all dimensions in a model need to have a common coordinate system. For example, the non-spatial dimen-

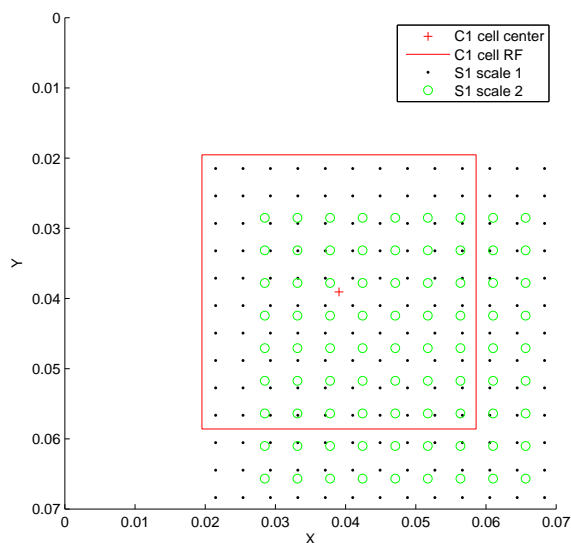


Figure 6. Common coordinate positions for several layers of an FH model. The C1 cell shown is a corner cell of a layer that has been arranged to perform a 10x10 valid convolution over the layer representing the first S1 scale. The second S1 scale’s cells do not line up with either of the other two layers shown; however, it is still possible to define a pooling operation over both S1 layers – the C1 cell just takes a max over any cell in its receptive field.

sions in FH and CN models do not have the notion of locality, so for them this scheme is not used.

## 4. Working with CNS

In this section we provide a quick overview of the process of developing and running CNS models.

Before you can build a network model, you must define your cell types. This is done by creating a *package*, which is a collection of related cell types, or by acquiring and possibly modifying an existing package. For example, if you wanted to build a network of Hodgkin-Huxley spiking cells, you would probably want to start with the existing HH package. The properties of cell types and their associated fields and kernels are described in sections 3.3, 3.4, and 3.5. Most of these definitions are made in MATLAB `.m` files, with kernels in C/C++ `.h` files. All such files for a single package are stored together in a package directory. Before you can use a package, it must be compiled into an executable using the `cns_build` command. This actually produces two executables, one that runs models on a GPU and another that runs models on a CPU. The latter option can be useful for development and debugging.

Once you have a compiled package, you can use it to build and run any number of different network models, hav-

ing different numbers and sizes of layers, with different connectivity, etc. The process of building and running a network model is as follows. All steps are carried out from within MATLAB using a few CNS commands.

1. Define the structure of your network: the number of layers, their sizes and types, cell-to-cell connectivity, and the initial values of fields. This is all defined in a MATLAB struct which you create. The helper function `cns_mapdim` can assist you in setting up a common coordinate system (section 3.6).
2. Initialize the model on the GPU (or CPU). This can take a few seconds.
3. Run the model for any number of iterations.
  - For dynamic models (e.g. HH models) this may involve loading some input data and then letting the model iterate for awhile. Variables can be tracked as they change, or if you just want their final values, you can pull them out when all iterations are complete.
  - Other models (e.g. FH models) process their input in a single iteration. For these models you simply load your input data, perform a single iteration, and pull out any output data. For batch operations (e.g., processing many images with an FH model) you would perform those three operations inside a loop.
4. Deallocate the model, freeing resources.

## 5. GPU details

While there are some circumstances in which model choices have performance consequences – these are documented in the programmer’s manual [6] – users in general do not need to understand GPU programming in order to use CNS.

Here we provide a few details on how CNS works behind the scenes. This section is intended for those familiar with GPU programming concepts; an exposition of GPU programming and architecture is beyond the scope of this paper.

**Dimension mapping.** The user sees layers as N-dimensional arrays, and this abstraction is consistent throughout CNS: both MATLAB functions and kernel macros view layers as N-D. However, internally layers are stored as 2-D. This is done because some fields need to be stored in GPU textures (see below). Textures can be 2-D or 3-D, but for 3-D textures the size limits are prohibitively small as of this writing. It is also more difficult to do texture packing in 3-D. Thus, CNS uses only 2-D textures.

All translation between N-D and 2-D is handled by CNS. The user's only involvement in this process occurs when defining a cell type. The user must choose the internal dimension (Y or X) to which each external dimension is mapped. Current GPU limits on texture size will influence these decisions. When two or more external dimensions are mapped to the same internal dimension, the user must also specify the nesting order. This choice has performance consequences. Storage will only be contiguous for the innermost dimension, so kernels involving nested loops should iterate over the innermost dimension in the innermost loop. (This is done in figure 5.) CNS automatically pads the innermost Y dimension to the warp size so that warps will perform coalesced reads as often as possible.

**Types of GPU memory.** CNS automatically maps each field to the appropriate kind of GPU memory based on the field's definition:

- Small, shared constants having layer, group, or model scope are stored in the constant cache. The constant cache is also used to store internal metadata such as layer sizes.
- Fields that have cell scope and that will be read by other cells are stored in textures. One texture is used for all the layers having that field; texture packing is done automatically.
- Everything else goes into global memory.
- Shared memory is not used.

**Blocks and threads.** All layers having the same cell type are evaluated in a single kernel call (unless the user has assigned them to different step numbers; see section 3.2). The decomposition into blocks and threads is done automatically. Each cell gets its own thread. Each layer becomes one or more thread blocks; an individual thread block will contain threads from a single layer only. Blocks are 2-D and aligned in the same way as memory (see above) to maximize the number of coalesced reads and the benefits of texture caching.

## 6. Limitations

### 6.1. Inherent Limitations

The following limitations are inherent in CNS's role as a rapid development framework for expressing arbitrary models having a cortical organization:

- CNS implements a generic, automatic process for mapping cortical models to the GPU architecture. For any given specific model, it will always be possible for a sufficiently skilled programmer to write custom

GPU code that runs faster by taking advantage of optimization techniques peculiar to that model. However, custom GPU code is much harder to write and modify. The speedups we are seeing under CNS relative to a single CPU (80-100x) are on the order of what is typically reported in the literature for direct GPU implementations of various algorithms. Our testing so far suggests that, at worst, a CNS model will be no more than 2x slower than a carefully-written custom GPU implementation of that specific model.

- Models must run inside a single GPU. Host-GPU communication is relatively slow, and CNS makes no assumptions about the sparsity of long-range vs. local connectivity, nor about the frequency of cell-to-cell communication. A less general framework in which such assumptions could be made might be able to automatically decompose models into pieces that could run on separate GPUs without incurring prohibitive data transfer delays. CNS cannot do this, so any such decomposition must be done by the user, outside CNS. Barring a dramatic improvement in the speed of host-GPU communication, this limitation cannot be removed, although it might be possible to implement automatic solutions for a subset of cortical models.

The largest GPU memory is currently 4GB. NVIDIA's upcoming Fermi architecture will have a 1TB address space, but it is not known how much memory the cards will actually have.

### 6.2. Current Limitations

These limitations could potentially be removed, some more easily than others:

- Only NVIDIA cards are supported. This could be overcome by converting CNS to use the new OpenCL API instead of CUDA.
- You cannot define fields that store 64-bit quantities, such as double-precision floating point numbers. Allowing fields of different sizes would somewhat complicate the current CNS code. Note that temporary variables used inside kernels *can* be 64-bit now.
- CNS is MATLAB-dependent. Since most of CNS is written in MATLAB, porting it would be a big job.
- Limitations on the size of the 2-D textures (currently 64K x 32K) used to store some fields can sometimes complicate model definition. This is beyond our control; however, given the current industry push to open up GPUs for general-purpose computing, it does not seem unreasonable that this limit might be removed in future cards.

## 7. Future work

With CNS stable, our main focus is now on specific package and model development. Likely future improvements to CNS itself include:

- Taking advantage of NVIDIA's new Fermi architecture. The two changes most relevant to CNS are the ability to run more than one kernel concurrently and the option to enable 48K of L1 cache.
- Investigating the feasibility of moving to OpenCL. This would allow us to use non-NVIDIA cards.

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