Sub-realtime simulation of a neuronal network of natural density

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Abstract

Full scale simulations of neuronal network models of the brain are challenging due to the high density of connections between neurons. This contribution reports run times shorter than the simulated span of biological time for a full scale model of the local cortical microcircuit with explicit representation of synapses on a recent conventional compute node. Realtime performance is relevant for robotics and closed-loop applications while sub-realtime is desirable for the study of learning and development in the brain, processes extending over hours and days of biological time.
INTRODUCTION

The cortical neuronal network of mammals exhibits a two-fold universality: basic characteristics of its architecture are conserved in evolution from mouse to human as well as across brain areas. This has motivated researchers to investigate models of the local cortical microcircuit, the network below a square millimeter of cortical surface, as a universal building block of brain-like computing. It is the smallest network in which both a realistic number of $10,000$ synapses per neuron and a connection probability of $0.1$ are realized simultaneously.

In a prototype network model of the microcircuit [1], the spatial structure of the cortex is neglected and replaced by cell-type specific random connectivity. Each cortical layer is represented by an excitatory and an inhibitory population of integrate-and-fire model neurons (Fig. 1a).

The microcircuit model has become a benchmark for neuromorphic computing systems: it can be simulated with moderate hardware investments [2, 3], its natural size renders questions of downscaling irrelevant [4], and it marks an upper bound as larger neuronal networks are necessarily less densely connected, and thus are, relative to the problem size, easier to simulate.

Fast and energy efficient simulation is a promise of neuromorphic computing [5]; desirable for large-scale neuroscientific models [6] and imperative in artificial intelligence and machine learning applications [7]. The first milestone is realtime performance, which was accomplished for the microcircuit model in 2019 on a neuromorphic system [8] followed this year by GPU systems [9, 10], one of them already breaking into the sub-realtime regime [10]. However, these results have to be evaluated in the light of continuously advancing commodity hardware as a reference technology providing more flexibility at potentially lower costs. With this aim we set out to investigate the performance of the general purpose simulation engine NEST [11] on a recent conventional computing system. Preliminary results have been presented in abstract form [12].
Figure 1. Strong scaling of a cortical microcircuit model on a conventional compute node. **a** Sketch of the microcircuit model with about 80,000 neurons and 300 million synapses organized into four layers of excitatory (blue) and inhibitory (red) populations of neurons (Supp. Inform. Fig. 1 shows activity). **b** Strong scaling for two placing schemes. Top graph shows realtime factor over total number of threads; dashed horizontal line indicates realtime; black solid line indicates linear scaling. The sequential scheme (blue) minimizes distance of threads on hardware, the distant scheme (brown) maximizes it; dashed vertical lines indicate number of cores per processor (64) and node (128). Bottom graphs show fractions of wall-clock time consumed by different stages of the simulation cycle; update: integrates state of neurons, deliver: distributes spike events to target neurons, communicate: transfers spikes between MPI processes (for shared and distributed memory setups), other: not accounted for by timers. **c** Top three graphs: Power measurements of a compute node during 100 s of model time in three configurations. The measurements are aligned to the start of the simulation phase starting at $t = 0$ (legend: colors distinguish phases and baseline). Bottom graph: Cumulative energy consumption of the simulations.

**METHODS**

We simulate the microcircuit model on 128 core dual socket AMD EPYC Rome 7702 compute nodes. Each processor is composed of 8 chiplets, each chiplet holds 8 cores,
resulting in 64 cores per socket (see Supp. Inform. Fig. 2). Each core has its own L1 and L2 cache, 4 cores share an L3 cache (see Supp. Inform. Fig. 3). Two nodes are coupled by a point-to-point Mellanox ConnectX-6 HDR100 interconnect. The software is NEST 2.14.1 [13] (compiled with GCC 6.3.0 and using jemalloc 3.6.0-9.1 [14], see Supp. Inform. Allocator) providing, in contrast to some neuromorphic systems, double precision numerics and weight resolution. NEST utilizes the Message Passage Interface (MPI, here OpenMPI 4.0.3rc4 [15]) and employs hybrid parallelization with multithreading (OpenMP [16]) for shared memory parallelization where a core never runs more than one thread. Timers monitor the different phases of the simulation.

Strong scaling experiments keep the task size fixed while systematically increasing the computational resources (Fig. 1b). The task is a simulation of 10 s of model time ($T_{Model}$), referring to the span of biological time described by the model, if not stated otherwise. Measurements start after model instantiation with optimized initial conditions [8] and an initial interval of 0.1 s of model time to ensure that potential transients of the network dynamics are discarded. To assess simulation speed we use the realtime factor:

$$RTF = \frac{T_{Wall}}{T_{Model}}$$

Here, $T_{Wall}$ denotes the wall-clock time; the time passed in the machine hall until the simulation completes. A realtime factor smaller than 1 implies sub-realtime performance. A common measure for comparing the energy consumption of neuromorphic systems is energy per synaptic event defined as total consumed energy divided by the total number of transmitted spikes (see Supp. Inform. Power measurements). For conducting the benchmarks we employ the JUBE [17] benchmarking environment.

RESULTS

We assess the strong scaling performance of microcircuit model simulations by increasing the number of threads on up to two compute nodes with two different schemes of binding threads to cores: In the “sequential” placing scheme, threads are bound onto physically consecutive cores per socket (thread counts 1 to 64 in steps of 1), and 1 MPI process per socket is used for simulations on one and two full nodes with 128 and 256 threads, respectively. In the “distant” placing scheme, threads are bound such that L3
cache and chiplet overlap is minimized per node (thread counts 1 to 128 in steps of 1, see Supp. Inform. Distant Placing) and 1 MPI process per node is used.

For sequential placing, we observe linear scaling for a thread count between 1 and 32 as well as super-linear scaling between 32 and 64 (Fig. 1b). A full compute node achieves sub-realtime performance with an RTF of 0.7. Two nodes reduce the realtime factor to 0.59; the simulation runs 1.7 times faster than realtime. The distant placing scheme exhibits super-linear scaling already for a small number of threads. At 33 threads, we note a sudden rise of the realtime factor. At this point, the L3 cache is shared for the first time. Nevertheless, sub-realtime performance is already achieved when using only 64 threads. Comparing the two placings at 128 and 256 threads respectively, we observe that sequential placing results in better performance. This is due to 2 MPI processes being used on one node in the sequential placing scheme as compared with 1 for the distant placing. The relative time spent in the update phase on a single node is decreased in the distant placing when compared with the sequential one and communication between the two nodes is not a limiting factor. This suggests that simulation time can be further reduced by increasing the number of nodes and alternatively using faster nodes.

We also assess the energy consumption of the simulation phase to investigate how the increased power uptake due to using more computational resources is counterbalanced by decreased simulation time (Fig. 1c). For this we compare a configuration using all 128 cores of a node with two configurations using only half of the cores. The former sequentially fills the cores of one socket, the latter employs the distant placing scheme. During simulations of 100 s of model time we record the power consumption and obtain the energy consumed in the simulation phase by integrating over the power readings.

We observe that power consumption during the simulation phase is largest for the distant placing of 64 threads, amounting to 0.39 kW subtracting the baseline power of 0.2 kW. This is almost twice the power as in the sequential configuration (0.21 kW). Nevertheless the increase cannot be attributed to the use of the second socket. The 128 thread configuration consumes 0.33 kW which is close to the same power required per thread of the sequential case. The counterintuitively low power consumption in the 128 threads case may be explained by the potentially longer latencies resulting in the cores not working at full capacity. Measuring the number of cache misses confirms a relative frequency of 25\% in distant as compared with 43\% in sequential placing (see Supp. Inform. Low level per-
| RTF  | $E_{\text{syn-event}}$ (µJ) | Reference                                      |
|------|--------------------------|------------------------------------------------|
| 6.29 | 4.39                     | 2018, NEST[2]                                   |
| 2.47 | 9.35                     | 2018, NEST[2]                                   |
| 26.08| 0.30                     | 2018, GeNN[3]                                   |
| 1.84 | 0.47†                    | 2018, GeNN[3]                                   |
| 1.00 | 0.60                     | 2019, SpiNNaker[8]                              |
| 1.06 | –                        | 2021, NeuronGPU[9]                              |
| 0.70 | –                        | 2021, GeNN[10]                                  |
| 0.67 | 0.33                     | NEST, AMD EPYC Rome (single node)               |
| 0.53 | 0.48                     | NEST, AMD EPYC Rome (two nodes)                 |

Table I. Realtime factor (RTF) and energy per synaptic event ($E_{\text{syn-event}}$) reported in the literature for simulations of the cortical microcircuit model [1] using conventional hardware for NEST simulations, GPUs for GeNN and NeuronGPU, and the dedicated neuromorphic hardware SpiNNaker in historical sequence (top to bottom). The two values reported for NEST and GeNN in 2018 (corresponding to the most energy efficient and the fastest configuration) are obtained with a different number of employed cores and different GPUs, respectively. †Value estimated by the authors.

DISCUSSION

Our study shows that a single compute node achieves sub-realtime performance in the simulation of a natural density local cortical microcircuit model. To our best knowledge, we report the lowest realtime factor so far at a competitive energy consumption (Table I). There are, however, preliminary data [18] on an even smaller realtime factor for a dedicated FPGA supercomputer using on-the-fly generation of connectivity. Our results expose that cache sensitive binding of threads increases performance.

Comparison with previous studies yields that conventional architectures keep pace
with dedicated hardware regarding both: realtime factor and energy efficiency. The employed generic simulation engine for spiking neuronal networks explicitly stores the connections between neurons with double floating point precision. Thus, although not exploited here, plasticity and learning are possible in this representation. No attempt is made to optimize the simulation code for the particular network model at hand. In comparison to prior work [2], where an earlier version (2.8.0) of the code and older hardware is used, we observe a ten-fold improvement in performance. The older system suffers from the communication between nodes as a bottleneck. The newer hardware pushes the limits by integrating a larger number of computational cores into the nodes. The analysis shows that on a single node faster completion of the task comes with a lower energy consumption due to the substantial baseline power. The simulation time reduces if cores have a larger amount of cache available, and if all cores are in use, power consumption is lower than for half of the cores with optimal cache access. These observations indicate that threads suffer from cache misses and the resulting latencies in memory access. This does not only give practical guidance for the design of conventional hardware but also raises hope that methods of prefetching and latency hiding can further improve simulation code without restricting generality[19].

Achieving realtime performance is a criterion for robotics. But for basic research and medical applications, also faster simulations are of use, because biological processes extending over long periods of time can be observed on a reduced time scale and multiple scenarios can be investigated quickly.

We hope that our results further advance and inspire the constructive competition between neuromorphic hardware and conventional computer architectures [2] which led to an order of magnitude improvement within just four years.

DATA AVAILABILITY

All data and analysis code to reproduce the results of this study can be downloaded from https://doi.org/10.5281/zenodo.5637375.
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SUPPLEMENTARY INFORMATION TO: SUB-REALTIME SIMULATION OF A NEURONAL NETWORK OF NATURAL DENSITY

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Figure 1. Raster plot of population resolved neuronal activity of a simulation of the microcircuit model. For each population (vertical), the display shows the spikes of a randomly selected fraction (60%) of the neurons in an arbitrary time segment of 200 ms (horizontal). Blue dots correspond to excitatory, red dots to inhibitory neurons. The temporal resolution of the simulation is 0.1 ms, the smallest delay in the network is 0.1 ms, the smallest time constant $\tau_{\text{syn}} = 0.5 \text{ ms}$ (synaptic current), the membrane time constant of the model neurons is 10 ms. The network exhibits spontaneous asynchronous irregular activity with cell-type specific firing rates akin to experimental findings.
Figure 2. Sketch of hardware architecture of the dual socket AMD EPYC Rome 7702 system used in this study. Solid blue squares indicate compute cores. 8 compute cores are combined into one chiplet.

Figure 3. Sketch of one chiplet of the AMD EPYC Rome 7702. 4 cores are grouped into one core complex sharing an L3 cache.

**Allocator**

Using the preloading mechanism for shared libraries, we use the jemalloc allocator via:

```bash
export LD_PRELOAD=/path/to/libjemalloc.so.1
```
Distant placing

Let us first introduce a numbering scheme for the cores as sketched in Figure 2. The kernel numbers the chiplets 0,...,15 where 0,...,7 identify consecutive chiplets on one socket and 8,...,15 on the other. The numbering is induced by the standard output of `lstopo` included in several Linux distributions. The command returns a numbered list of the cores on the compute node hierarchically structured by the NUMA nodes (in our case equivalent to the sockets), the L3 cache and the L1/L2 cache. Cores 0 to 63 and L3 caches 0 to 15 are located on NUMA node 0, cores 64 to 127 and L3 caches 16 to 31 on NUMA node 1. Since on one chiplet two L3 caches are located, one obtains the number of a chiplet by an integer division of the number of the respective L3 cache by 2. We denote the $k$–th core, $k \in \{0,...,7\}$ (sketched in Figure 3), on the $n$–th chiplet by $n : k$. In the distant placing scheme the filling of a compute node with threads is split into 8 rounds each addressing a particular core $k$ of the chiplets and successively adding this core of chiplet $n$ (16 in total) to the simulation. This results in $8 \times 16 = 128$ threads being bound to cores. The filling procedure starts with core 0, i.e. the first 16 use the cores $\{0 : 0\}, \{0 : 0, 1 : 0\}, \ldots, \{0 : 0, \ldots, 15 : 0\}$. The next round uses cores still not sharing an L3 cache with cores already in use. We chose the 4–th, resulting in consecutively adding the cores $0 : 4, 1 : 4, \ldots, 15 : 4$. The following rounds continue with the 2–nd, 6–th, 1–st, 5–th, 3–rd and 7–th core respectively, minimizing shared use of L3 cache. To bind threads to cores on our system we export OpenMP variables as follows:

```
export OMP_NUM_THREAD = $CPUSPERTASK
export OMP_PROC_BIN = TRUE
export OMP_PLACES = \{0\}, \{8\}, \{15\}
```

Here `$CPUSPERTASK` is the number of cores used in a given setup (in this example 3) and `\{0\}, \{8\}, \{15\}` indicate the first core on the first, second and third chiplet. Simulations on one node are launched by

```
python3 run_microcircuit.py
```

Simulation on two nodes are launched by

```
mpirun --n 2 --npernode 1 --mca pml ucx -x UCX_NET_DEVICES=
 mlx5_1:1 --bind-to board python3 run_microcircuit.py
```
in this example with 1 MPI process per node.

**Power measurements**

Power was measured with a Raritan Dominion PX and a Raritan PX3-5190 power distribution unit (PDU). The units have an accuracy of ±5% and data collection frequency of 1 Hz. The power measurement has a delay of 1 s, so that the power readings need to be shifted by 1 s to be aligned to wall-clock time. Since the nodes are connected point-to-point, we do not need to take additional passive energy consumption by an interconnect into account.

**Low level performance measurements**

In order to determine the number of cache misses we employ the `perf` performance analysis tool of the Linux operating system. We use the command options

```
perf stat -ae task-clock,cycles,instructions,cache-references,cache-misses
```

and increase the simulation time to 100 s. With this we ensure that approximately 80% of the run time of the program is spent in the simulation phase guaranteeing a reliable assessment of the percentage of cache misses during that phase.