# Lower Bounds for the Computational Power of Networks of Spiking Neurons

### Wolfgang Maass

*Institute for Theoretical Computer Science, Technische Universitaet Graz, Klosterwiesgasse* 32/2, A-8010 Graz, Austria

We investigate the computational power of a formal model for networks of spiking neurons. It is shown that simple operations on phase differences between spike-trains provide a very powerful computational tool that can in principle be used to carry out highly complex computations on a small network of spiking neurons. We construct networks of spiking neurons that simulate arbitrary threshold circuits, Turing machines, and a certain type of random access machines with real valued inputs. We also show that relatively weak basic assumptions about the response and threshold functions of the spiking neurons are sufficient to employ them for such computations.

#### 1 Introduction and Basic Definitions \_\_\_\_\_

There is substantial evidence that timing phenomena such as temporal differences between spikes and frequencies of oscillating subsystems are integral parts of various information processing mechanisms in biological neural systems (for a survey and references see, e.g., Kandel et al. 1991; Abeles 1991; Churchland and Sejnowski 1992; Aertsen 1993). Furthermore, simulations of a variety of specific mathematical models for networks of spiking neurons have shown that temporal coding offers interesting possibilities for solving classical benchmark problems such as associative memory, binding, and pattern segmentation (for an overview see Gerstner et al. 1993). Very recently one has also started to build artificial neural nets that model networks of spiking neurons (see, e.g., Murray and Tarassenko 1994; Watts 1994). Some aspects of these models have also been studied analytically (see, e.g., Gerstner and van Hemmen 1994; Gerstner 1995), but almost nothing is known about their computational complexity (see Judd and Aihara 1993, for some first results in this direction). In this article we investigate a simple formal model SNN for networks of spiking neurons that allows us to model the most important timing phenomena of neural nets, and we prove lower bounds for its computational power.

Quite a number of different mathematical models for networks of spiking neurons have previously been introduced within the frameworks

of theoretical physics and theoretical neurobiology (see, e.g., Lapicque 1907; Buhmann and Schulten 1986; Crair and Bialek 1990; Gerstner 1991; Gerstner et~al. 1993; for a survey and the relationship between these and related models see, e.g., Tuckwell 1988; and Gerstner 1995). The computational model SNN that we consider in this article is most closely related to the *spike response model* of Gerstner (1991) and Gerstner et~al. (1993). Similarly as in Buhmann and Schulten (1986), we consider in this article only the deterministic case (which corresponds to the limit case  $\beta \to \infty$  for the inverse temperature  $\beta$  in the spike response model, and respectively, the noise-free case). We refer to Maass (1995d) for results about the computational power of the *noisy* version of this model.

However, in contrast to these preceding models we do not fix particular (necessarily somewhat arbitrarily chosen) response and threshold functions in our model SNN. Instead, we want to be able to use the SNN model as a framework for *investigating* the computational power of various *different* response and threshold functions. In addition, we would like to make sure that various *different* response and threshold functions observed in specific biological neural systems are in fact special cases of the response and threshold functions in the formal model SNN.

## 1.1 Definition of a Spiking Neuron Network (SNN). An SNN ${\mathcal N}$ consists of

- a finite directed graph  $\langle V, E \rangle$  (we refer to the elements of V as "neurons" and to the elements of E as "synapses")
- a subset  $V_{in} \subseteq V$  of input neurons
- a subset  $V_{\text{out}} \subseteq V$  of output neurons
- for each neuron  $v \in V V_{\text{in}}$  a threshold function  $\Theta_v : \mathbf{R}^+ \to \mathbf{R} \cup \{\infty\}$  (where  $\mathbf{R}^+ := \{x \in \mathbf{R} : x \geq 0\}$ )
- for each synapse  $\langle u, v \rangle \in E$  a response function  $\varepsilon_{u,v} : \mathbf{R}^+ \to \mathbf{R}$  and a weight function  $w_{u,v} : \mathbf{R}^+ \to \mathbf{R}$ .

We assume that the firing of the input neurons  $v \in V_{\text{in}}$  is determined from outside of  $\mathcal{N}$ , i.e., the sets  $F_v \subseteq \mathbf{R}^+$  of firing times ("spike trains") for the neurons  $v \in V_{\text{in}}$  are given as the *input* of  $\mathcal{N}$ . Furthermore we assume that a set  $T \subseteq \mathbf{R}^+$  of *potential firing times* has been fixed (we will consider only the cases  $T = \mathbf{R}^+$  and  $T = \{i \cdot \mu : i \in \mathbf{N}\}$  for some  $\mu > 0$ ).

For a neuron  $v \in V - V_{\text{in}}$  one defines its set  $F_v$  of firing times recursively. The first element of  $F_v$  is  $\inf\{t \in T : P_v(t) \geq \Theta_v(0)\}$ , and for any  $s \in F_v$  the next larger element of  $F_v$  is  $\inf\{t \in T : t > s \text{ and } P_v(t) \geq \Theta_v(t - s)\}$ , where the potential function  $P_v : \mathbf{R}^+ \to \mathbf{R}$  is defined by

$$P_v(t) := 0 + \sum_{u : \langle u, v \rangle \in E} \sum_{s \in F_u : s < t} w_{u,v}(s) \cdot \varepsilon_{u,v}(t-s)$$

[the trivial summand 0 makes sure that  $P_v(t)$  is well-defined even if  $F_u = \phi$  for all u with  $\langle u, v \rangle \in E$ ].

The firing times ("spike trains")  $F_v$  of the output neurons  $v \in V_{\text{out}}$  that result in this way are interpreted as the *output* of N.

Regarding the set T of potential firing times we consider in this article primarily the case  $T = \mathbf{R}^+$  (SNN with continuous time), and only in Corollary 2.5 the case  $T = \{i \cdot \mu : i \in \mathbf{N}\}$  for some  $\mu > 0$  (SNN with discrete time).

Our subsequent assumptions about the threshold functions  $\Theta_v$  will imply that for each SNN  $\mathcal N$  there exists a bound  $\tau_{\mathcal N} \in \mathbf R$  with  $\tau_{\mathcal N} > 0$  such that  $\Theta_v(x) = \infty$  for all  $x \in (0,\tau_{\mathcal N})$  and all  $v \in V - V_{\rm in}$  ( $\tau_{\mathcal N}$  may be interpreted as the minimum of all "refractory periods"  $\tau_{\rm ref}$  of neurons in  $\mathcal N$ ). Furthermore we assume that all "input spike trains"  $F_v$  with  $v \in V_{\rm in}$  satisfy  $|F_v \cap [0,t]| < \infty$  for all  $t \in \mathbf R^+$ . On the basis of these assumptions one can also in the continuous case easily show that the firing times are well-defined for all  $v \in V - V_{\rm in}$  (and occur in distances of at least  $\tau_{\mathcal N}$ ).

In models for *biological neural systems* one assumes that if *x* time-units have passed since its last firing, the current threshold  $\Theta_v(x)$  of a neuron v is "infinite" for  $x < \tau_{ref}$  (where  $\tau_{ref} = refractory period of neuron v),$ and then approaches quite rapidly from above some constant value. A neuron v "fires" (i.e., it sends an "action potential" or "spike" along its axon) when its current membrane potential  $P_{\nu}(t)$  at the axon hillock exceeds its current threshold  $\Theta_v$ .  $P_v(t)$  is the sum of various postsynaptic potentials  $w_{u,v} \cdot \varepsilon_{u,v}(t-s)$ . Each of these terms describes an *excitatory* (EPSP) or inhibitory (IPSP) postsynaptic potential at the axon hillock of neuron v at time *t*, as a result of a spike that had been generated by the "presynaptic" neuron u at time s, and which has been transmitted through a synapse between both neurons. Recordings of an EPSP typically show a function that has a constant value c (c = resting membrane potential; e.g., c = -70 mV) for some initial time interval (reflecting the axonal and synaptic transmission time), then rises to a peak value, and finally drops back to the same constant value c. An IPSP tends to have the negative shape of an EPSP (see Fig. 3). For the sake of mathematical simplicity we assume in the SNN model that the constant initial and final value of all response functions  $\varepsilon_{u,v}$  is equal to 0 (in other words,  $\varepsilon_{u,v}$  models the difference between a postsynaptic potential and the resting membrane potential c). Different presynaptic neurons u generate postsynaptic potentials of different sizes at the axon hillock of a neuron v, depending on the size, location, and current state of the synapse (or synapses) between u and v. This effect is modeled by the weight factors  $w_{u,v}(s)$ .

The precise shapes of threshold, response, and weight functions may vary among different biological neural systems, and even within the same system. Fortunately one can prove significant *upper bounds* for the computational complexity of SNNs  $\mathcal N$  without *any* assumptions about the *specific shapes* of these functions of  $\mathcal N$ . Instead, for such upper bounds one only has to assume that they are of a reasonably simple *mathematical structure* (see Maass 1994b, 1995c).

To prove *lower bounds* for the computational complexity of an SNN  $\mathcal{N}$  one is forced to make more specific assumptions about these functions. However, we show in this article that significant (and in some cases *optimal*, see Section 3) lower bounds can be shown under some rather weak *basic assumptions* about these functions, which will be further relaxed in Section 4. These basic assumptions (see Section 2) mainly require that EPSPs have an arbitrarily small time segment where they increase linearly, and some arbitrarily small time segment where they decrease linearly. Since the computational power of SNNs may potentially increase through the use of time-dependent weights, *lower* bounds for their computational power are more significant if they do *not* involve the use of time-dependent weights. Hence we will assume throughout this article that *all weight-functions*  $w_{u,v}(s)$  have a constant value  $w_{u,v}$ , which does not depend on the time s.

Apart from the abovementioned condition on the existence of linear segments in EPSPs, the basic assumptions that underlie the lower bound results of this article involve no other significant conditions on the shape of response and threshold functions. Hence one may argue that these basic assumptions are biologically plausible. In addition, we will show in Section 4 that the same lower bounds can be shown if also phenomena such as "adaption" of neurons, or a "reset" of the potential after a firing are taken into account. Thus the more critical points with regard to the biological interpretation of these lower bound results appear to be the relatively simple firing mechanism of the SNN model, which, for example, ignores for the sake of simplicity nonlinear interactions among postsynaptic potentials such as integration of potentials within the dendritic tree of a neuron, and various possible sources of "imprecision" in the determination of the firing times. The latter issue can partially be taken into account by considering the variation of the SNN model with discrete firing times as in Corollary 2.5 (although the implicit global synchronization of this version is not completely satisfactory). In this variation of the SNN model with discrete firing times  $i \cdot \mu$  for  $i \in \mathbb{N}$  one can view a firing of a neuron at time  $i \cdot \mu$  as representing a somewhat imprecise firing time in a small interval around time  $i \cdot \mu$ .

The computational complexity of another neural network model where timing plays an important role has previously been investigated by Judd and Aihara (1993). Their model PPN is also motivated by biological spiking neurons, but it employs a quite different firing mechanism. There are no response functions in this model, and instead of integrating all incoming EPSPs and IPSPs in order to determine whether it should "fire," a neuron in a PPN randomly selects a *single* one of the incoming "stimulations" of maximal size, and determines on the basis of that stimulation whether it should fire. Consequently, computations in this model PPN proceed quite differently from computations in models of spiking neurons such as the spike response model of Gerstner and van Hemmen (1994), or the model SNN considered here. Judd and Aihara (1993) con-

struct PPNs that can simulate Turing machines that use at most a constant number s of cells on their tapes, where s is bounded by the number of neurons in the simulating PPN. However a Turing machine with a constant bound s on its number of tape cells is just a special case of a finite automaton, and hence this result does not show that a PPN of finite size can have the computational power of an arbitrary Turing machine. In contrast to the quoted result about PPNs, it is shown in Theorem 2.1 of this article that with arbitrary response and threshold functions that satisfy the basic assumptions of Section 2, one can construct for any given Turing machine M an SNN  $\mathcal{N}_M$  of finite size that can simulate any computation of M in real-time (even if the number of tape cells that M uses is much larger than the number of neurons in  $\mathcal{N}_M$ ). In addition, at the end of Section 4 we will describe a way in which a simulation of arbitrary Turing machines can also be accomplished by finite SNNs whose response and threshold functions are piecewise constant. If we understand the model of Judd and Aihara (1993) correctly (their description is somewhat unclear), then our method for proving this (see also Maass and Ruf 1995) can also be used to show that with the help of a module that decides whether two neurons have fired simultaneously, one can simulate (although not in real-time) any Turing machine M (where M may use an unbounded number of tape cells) by some PPN  $\mathcal{P}_{M}$  of finite size, thereby improving the lower bound for the computational power of PPNs due to Judd and Aihara (1993), from finite automata to Turing machines.

The focus in the investigation of computations in biological neural systems differs in two essential aspects from that of classical computational complexity theory. First, the timing constraints for computations in biological neural systems are often tighter than for computations in digital computers, and many complex computations have to be carried out in "real-time" with relatively slow "switching elements." Secondly, one is not only interested in separate computations on unrelated inputs, but also in the ability of the system to *learn* to react appropriately to a *sequence* of *related* tasks.

Hence the custom to evaluate the computational power in terms of "complexity classes" such as *P* or *P/poly* appears to be less suitable for the investigation of models for biological neural systems, and we therefore resort to an analysis in terms of refined concepts such as "real-time computations" and "real-time simulations." In this way we get not only information about the relationship between the "large scale" complexity classes (e.g., polynomial time) for these models for biological neural systems, but also about their behavior in terms of common notions of "low-level complexity" such as sublinear or real-time.

Furthermore, with the help of our refined analysis of real-time simulations one also gets information about the "adaptive" or "learning" abilities of the considered models. Assume for example that  $(\langle x(i), y(i) \rangle)_{i \in \mathbb{N}}$  is the protocol of some real-time "learning process" of a system M, where the y(i) are the "responses" of M to a sequence x(i) of "stimuli." If one

has shown that another model M' can simulate M in real-time, then this entails that the same "learning process" can also be carried out in real-time by M'.

**1.2 Definition of Real-Time Computation and Real-Time Simulation.** Fix some arbitrary (finite or infinite) input alphabet  $A_{\text{in}}$  and output alphabet  $A_{\text{out}}$  (for example they can be chosen to be  $\{0,1\}$ ,  $\{0,1\}^*$  or  $\mathbb{R}$ ). We say that a *machine M processes a sequence*  $(\langle x(i), y(i) \rangle)_{i \in \mathbb{N}}$  of pairs  $\langle x(i), y(i) \rangle \in A_{\text{in}} \times A_{\text{out}}$  in real-time r, if M outputs y(i) for every  $i \in \mathbb{N}$  within r computation steps after having received input x(i) [for i > 0 we assume that x(i) is presented at the next step after M has given output y(i-1)].

We say that a *machine M'* simulates a machine M in real-time (with delay factor  $\Delta$ ) if for every  $r \in \mathbb{N}$  and every sequence that is processed by M in real-time r, M' can process the same sequence in real time  $\Delta \cdot r$ .

In the case of SNNs M we count each spike in M as a computation step.

We first would like to point out that these notions contain the usual notions of a computation and simulation as special cases. Let {0,1}\* be the set of all binary sequences of finite length. If M computes a Boolean function  $F: \{0,1\}^* \to \{0,1\}$  in time t(n) (in the usual sense of computational complexity theory), then one can identify each input  $\langle z_1, \ldots, z_n \rangle \in \{0,1\}^*$  with an infinite sequence  $\langle x(i) \rangle_{i \in \mathbb{N}}$  where  $x(i) = z_i$ for  $i \le n$  and x(i) = B for i > n (assume that M gets one input bit per step, B := "blank"). Furthermore one can set y(i) = B for those steps i where M's computation is not yet finished, and  $y(i) = F(\langle z_1, \dots, z_n \rangle)$ for all later i [in particular for all  $i \geq t(n)$ ]. Obviously M processes this sequence  $(\langle x(i), y(i) \rangle)_{i \in \mathbb{N}}$  in real-time 1. Hence, if another machine M' can simulate M in real-time with delay factor  $\Delta$ , then M' can compute the same function  $F: \{0,1\}^* \to \{0,1\}$  in time  $\Delta \cdot t(n)$ . This implies that a realtime simulation is a special case of a linear-time simulation. In particular, every computational problem that can be solved by M within a certain time complexity can be solved by M' within the same time complexity (up to a constant factor).

In addition, the remarks before the definition imply that when we show that M' can simulate M in real-time, we may conclude that any  $adaptive\ behavior$  (or  $learning\ algorithm$ ) of M can also be implemented on M'. Finally we would like to point out that for the investigation of specific computational and learning problems on specific models for biological neural nets one would like to also eventually get estimates for the size of the constant r in real-time processing and the size of the delay factor  $\Delta$  in a real-time simulation. Such refined analysis (which will not be carried out in this paper) appears to be also of interest, since it is likely to throw some light on the specific advantages and disadvantages of different models for biological neural systems (e.g., networks of spiking

neurons versus analog neural nets), which are shown in Maass (1994b, 1995c), to be equivalent with regard to the preceding notion of a real-time simulation.

In contrast to the usual notion of a simulation, a *real-time* simulation of another computational model M by an SNN implies that the simulation of each computation step of M requires only a *fixed* number of spikes in the SNN. In particular, the required number of spikes does not become larger for the simulation of later computation steps of M.

- **1.3 Input and Output Conventions.** For simulations between SNNs and Turing machines one may either assume that the SNN gets an input (or produces an output) from  $\{0,1\}^*$  in the form of a spike train (i.e., one bit per unit of time), or that the input (output) of the SNN is encoded into the phase difference of just two spikes. The former convention is suitable for comparisons with Turing machines that receive a single input bit and produce a single output bit at each computation step. For comparisons with Turing machines that start with the whole input written on a specified tape, and have their whole output written on another tape when the machine halts, it is more adequate to assume that the SNN receives at the beginning of a computation the whole tape content of the input tape encoded into the time difference  $\varphi$  between two spikes (using the same encoding as we will use in Section 2 to represent the content of a stack), and that the SNN also provides the final content of the output tape in the same form. Real-valued input or output for an SNN is always encoded into the phase difference of two spikes.
- **1.4 Notation.** We employ in this article the following common notation: We write **N** for the set of natural numbers (including 0), **Q** for the set of rational numbers, and **R** for the set of real numbers.  $\mathbf{R}^+$  is defined as  $\{x \in \mathbf{R} : x \ge 0\}$ . For any  $x \in \mathbf{R}^+$  we write  $\lceil x \rceil$  for the least  $n \in \mathbf{N}$  with  $n \ge x$ .

 $\{0,1\}^*$  denotes the set of all binary strings of finite length.

For any set *S* we write  $\exists x \in S(...)$  instead of  $\exists x (x \in S \text{ and }...)$ , and  $\forall x \in S(...)$  instead of  $\forall x (x \in S \to ...)$ .

For two functions  $f,g: \mathbf{N} \to \mathbf{N}$  we write f = O(g) if there is some constant c such that  $f(n) \le c \cdot g(n)$  for all except possibly finitely many  $n \in \mathbf{N}$ .

**1.5 Structure of This Article.** In Section 2 we specify our basic assumptions about the response and threshold functions of an SNN, and we construct SNNs that can simulate in real-time arbitrary threshold circuits and Turing machines. In Section 3 we relate the computational power of SNNs for real-valued inputs to a specific type of random access machine. In Section 4 we discuss variations of the preceding constructions

for related models of spiking neurons, and in Section 5 we outline some conclusions from the results in this article.

# 2 Simulation of Threshold Circuits and Turing Machines by Networks of Spiking Neurons

To carry out computations on an SNN, *some* assumptions have to be made about the structure of the response and threshold functions of its neurons. It is obvious that for example neurons with response function  $\varepsilon_{u,v}$  such that  $\varepsilon_{u,v}(s) = 0$  for all  $s \ge 0$  cannot carry out *any* computation. We will specify in the following a set of **basic assumptions**, which suffice for the constructions in this article. Some variations of these conditions will be discussed in Section 4.

We assume that there exist some arbitrary given constants  $\Delta_{\min}$ ,  $\Delta_{\max} \in \mathbf{R}$  with  $0 \leq \Delta_{\min} < \Delta_{\max}$  so that we can choose for each "synapse"  $\langle u,v \rangle \in E$  an individual "delay"  $\Delta_{u,v} \in [\Delta_{\min}, \Delta_{\max}]$  with  $\varepsilon_{u,v}(x) = 0$  for all  $x \in [0, \Delta_{u,v}]$ . This parameter  $\Delta_{u,v}$  corresponds in biology to the time span between the firing of the presynaptic neuron u and the moment when its effect reaches the trigger zone (axon hillock) of the postsynaptic neuron v. This time span is known to vary for individual neurons in biological neural systems, depending on the type of synapse and the geometric constellation. The constants  $\Delta_{\min}$  and  $\Delta_{\max}$  can be interpreted as biological constraints on the possible lengths of such time spans. No requirements about  $\Delta_{\min}$  and  $\Delta_{\max}$  are needed for our construction, except that  $\Delta_{\min} < \Delta_{\max}$ .

We assume that except for their individual delays  $\Delta_{u,v}$  the response functions  $\varepsilon_{u,v}$  (as well as the threshold functions  $\Theta_v$ ) are *stereotyped*, i.e., that their shape is determined by some general functions  $\varepsilon^E$ ,  $\varepsilon^I$ , and  $\Theta$ , which do not depend on u or v. More precisely, we assume that we can decide for any pair  $\langle u, v \rangle \in E$  whether  $\varepsilon_{u,v}$  should represent an excitatory "EPSP response function," or an inhibitory "IPSP response function." In the EPSP case we assume that

$$\varepsilon_{u,v}(\Delta_{u,v} + x) = \varepsilon^E(x)$$
 for all  $x \in \mathbf{R}^+$ .

and in the IPSP case we assume that

$$\varepsilon_{u,v}(\Delta_{u,v}+x)=\varepsilon^I(x)$$
 for all  $x\in \mathbf{R}^+$ .

In either case we assume that

$$\varepsilon_{u,v}(x) = 0$$
 for all  $x \in [0, \Delta_{u,v}]$ .

Furthermore, we assume for all neurons  $v \in V - V_{\text{in}}$  that

$$\Theta_v(x) = \Theta(x)$$
 for all  $x \in \mathbf{R}^+$ .

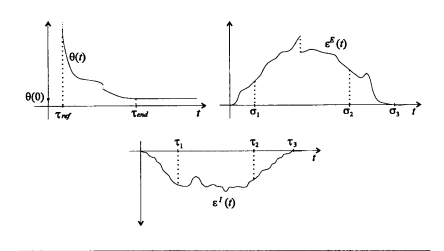


Figure 1: Illustration of our notation for the basic assumptions on  $\Theta$ .  $\varepsilon^E$ .  $\varepsilon^I$  (the functions shown are quite arbitrary and complicated, but nevertheless they satisfy our basic assumptions).

We assume that the three functions  $\varepsilon^E: \mathbf{R}^+ \to \mathbf{R}^+$ ,  $\varepsilon^I: \mathbf{R}^+ \to \{x \in \mathbf{R}: x \leq 0\}$  and  $\Theta: \mathbf{R}^+ \to \mathbf{R}^+ \cup \{\infty\}$  are some *arbitrary* functions with the following properties: There exist some arbitrary strictly positive real numbers  $\tau_{\rm ref}$ ,  $\tau_{\rm end}$ ,  $\sigma_1$ ,  $\sigma_2$ ,  $\sigma_3$ ,  $\tau_1$ ,  $\tau_2$ ,  $\tau_3$ , L,  $s_{\rm up}$ ,  $s_{\rm down}$  with  $0 < \tau_{\rm ref} < \tau_{\rm end}$ ,  $\sigma_1 < \sigma_2 < \sigma_3$ ,  $\tau_1 < \tau_2 < \tau_3$  (see Fig. 1 for an illustration), which satisfy the following five conditions:

- 1.  $\Theta(x) \ge \Theta(0) > 0$  for all  $x \in \mathbb{R}^+$ ,  $\Theta(x) = \infty$  for all  $x \in (0, \tau_{\text{ref}})$ . and  $\Theta(x) = \Theta(0) < \infty$  for all  $x \in [\tau_{\text{end}}, \infty)$
- 2.  $\varepsilon^{\mathcal{E}}(0) = \varepsilon^{\mathcal{E}}(x) = 0$  for all  $x \in [\sigma_3, \infty)$ , and there exists some  $\varepsilon_{\max} \in \mathbb{R}^+$  so that  $\exists x \in \mathbb{R}^+[\varepsilon^{\mathcal{E}}(x) = \varepsilon_{\max}]$  and  $\forall y \in \mathbb{R}^+[\varepsilon^{\mathcal{E}}(y) \leq \varepsilon_{\max}]$
- 3.  $\varepsilon^{E}(\sigma_1 + z) = \varepsilon^{E}(\sigma_1) + s_{up} \cdot z$  for all  $z \in [-L, L]$
- 4.  $\varepsilon^{E}(\sigma_2 + z) = \varepsilon^{E}(\sigma_2) s_{\text{down}} \cdot z \text{ for all } z \in [-L, L]$
- 5.  $\varepsilon^{I}(0) = \varepsilon^{I}(x) = 0$  for all  $x \in [\tau_3, \infty)$ ,  $\varepsilon^{I}(x) < 0$  for all  $x \in (0, \tau_3)$ .  $\varepsilon^{I}$  is nonincreasing in  $[0, \tau_1]$  and nondecreasing in  $[\tau_2, \tau_3]$ .

We assume in addition that  $\Theta(0)$ ,  $\varepsilon^{E}(\sigma_{1})$ ,  $\varepsilon^{E}(\sigma_{2})$ ,  $s_{up}$ ,  $s_{down} \in \mathbf{Q}$ .

It should be pointed out that no conditions about the smoothness, the continuity, or the number of extrema of the functions  $\Theta$ ,  $\varepsilon^E$ ,  $\varepsilon^I$  are made in the preceding basic assumptions. However, if one demands in addition that  $\varepsilon^E$  is piecewise linear and continuous, then conditions (3) and (4) become redundant. The assumption that  $\Theta(0)$ ,  $\varepsilon^E(\sigma_1)$ ,  $\varepsilon^E(\sigma_2)$ ,  $s_{\rm up}$ ,

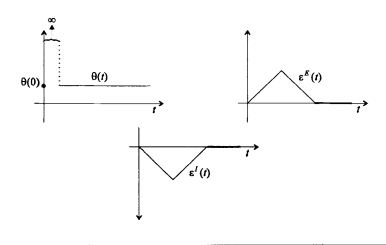


Figure 2: Examples for mathematically very simple functions  $\Theta$ ,  $\varepsilon^{I}$ , and  $\varepsilon^{I}$  that satisfy the basic assumptions.

 $s_{down}$  are rationals will be needed only to ensure that certain weights can be chosen to be *rationals* (see Section 2.9).

Examples of mathematically particularly simple (piecewise linear) functions  $\varepsilon^t$ .  $\varepsilon^t$  and  $\Theta$  that satisfy all of the above conditions are exhibited in Figure 2.

The subsequent construction shows that neurons with the very simple response and threshold functions from Figure 2 can, in principle, be used to build an artificial neural network with some finite number  $n_{\mathcal{U}}$  of spiking neurons that can simulate in real time any other digital computer (even computers that employ many more than  $n_{\mathcal{U}}$  memory cells or computational units).

We have formulated the preceding basic assumptions on the response and threshold functions in a rather general fashion to make sure that they can in principle be satisfied by a wide range of EPSPs, IPSPs and threshold functions that have been observed in a number of biological neural systems (see, e.g., Fig. 3).

The currently available findings about biological neural systems (see, e.g., Kandel  $et\ al.$  1991, and the discussions in Valiant 1994) indicate that in general a single EPSP alone cannot cause a neuron to fire. In fact, it is commonly reported that 50 to 100 EPSP have to arrive within a short time span at a neuron to trigger its firing. These reports indicate that the weights  $w_{n,v}$  in our model should be assumed to be relatively small, since they cannot amplify a single EPSP to yield an arbitrarily high potential  $P_v$ . Hence for the sake of biological plausibility one should

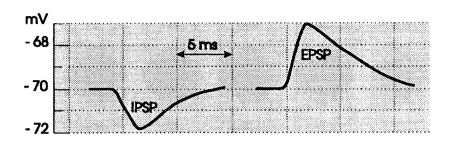


Figure 3: Inhibitory and excitatory postsynaptic potentials at a biological neuron. [After Schmidt (1978). *Fundamentals of Neurophysiology*. Springer-Verlag, Berlin].

assume that the values of all weights  $w_{u,v}$  in an SNN belong to some bounded interval  $[0,w_{\max}]$ . For simplicity we assume in the following that  $w_{\max}=1$ . This convention just amounts to a certain scaling of the values of the response functions in relation to the threshold functions. In any version of this model where a single neuron is not able to cause the firing of another neuron, one necessarily has to assume that each input spike is simultaneously received by *several* neurons (since otherwise it cannot have any effect).

In spite of this convention we will occassionally assign much larger values to certain weights  $w_{u,v}$ . We will then (silently) assume that u does in fact represent an assembly of  $\lceil w_{u,v} \rceil$  neurons that all fire concurrently ( $\lceil x \rceil$  is defined as the least natural number  $\geq x$ ). Furthermore, we assume in those situations that all edges from neurons in this assembly to neuron v have the same delay, and the same weight  $w_{u,v}/\lceil w_{u,v} \rceil \in [0,1]$ . The main difference between this type of construction and a construction with arbitrarily large weights is that in our setup the (virtual) use of large weights blows up the number of neurons that are needed.

**Theorem 2.1.** If the response and threshold functions of the neurons satisfy the previously described basic assumptions, then one can build from such neurons for any given  $d \in \mathbf{N}$  an SNN  $\mathcal{N}_{TM}(d)$  of finite size that can simulate with a suitable assignment of rational values from [0,1] to its weights any Turing machine with at most d tapes in real-time.

Furthermore  $\mathcal{N}_{TM}(2)$  can compute any function  $F: \{0,1\}^* \to \{0,1\}^*$  with a suitable assignment of real values from [0,1] to its weights.

The proof of Theorem 2.1 is rather complex. Therefore we have divided it into Sections 2.1 to 2.10, which are devoted to different aspects of the modules of the construction. Several of these modules are also useful for other constructions. The global construction of  $\mathcal{N}_{TM}(d)$  with the properties claimed in Theorem 2.1 is described in Section 2.10.

We will discuss in Section 4 some methods for alternative constructions of  $\mathcal{N}_{TM}(d)$  that are based on different assumptions about response and threshold functions.

- **2.1 Conditions on the Neurons.** We assume that we can decide for any pair  $\langle u,v\rangle$  of neurons whether there should be a "synapse" between both neurons (i.e.,  $\langle u,v\rangle \in E$ ). Self-referential edges of the form  $\langle u,u\rangle$  will not be needed. In this proof the weights  $w_{u,v}$  on edges  $\langle u,v\rangle$  are always assumed to be time invariant, and they are only assigned values from [0,1]. We assume that the response and threshold functions satisfy the previously described basic assumptions.
- **2.2 Delay- and Inhibition Modules.** We will construct in this section two very simple modules that will be used frequently (and often silently) in the subsequent constructions. From the general point of view the existence of these two modules shows that our very weak assumptions about  $\Delta_{\min}$  and  $\Delta_{\max}$  (we have only required that  $0 \leq \Delta_{\min} < \Delta_{\max}$ ) as well as our very weak assumptions about the shape of  $\varepsilon^I$  in condition (5) are in fact sufficient to create in an SNN arbitrarily long delays, and arbitrarily fast appearing or arbitrarily fast disappearing inhibitions of arbitrarily long duration.

A "delay-module" is simply a chain  $u_1,\ldots,u_{k+1}$  of neurons so that  $\langle u_i,u_{i+1}\rangle\in E$ ,  $\varepsilon_{u_i,u_{i+1}}$  is an EPSP response function, and  $w_{u_i,u_{i+1}}:=\Theta(0)/\varepsilon_{\max}$  for  $i=1,\ldots,k$ . Since each delay  $\Delta_{u_i,u_{i+1}}$  can be chosen arbitrarily from  $[\Delta_{\min},\Delta_{\max}]$ , the total "delay" between the firing of  $u_1$  and the arrival of an EPSP at  $u_{k+1}$  can be chosen to assume any value in a certain interval of length  $k\cdot(\Delta_{\max}-\Delta_{\min})$ . It will cause no problem that the total transmission time from  $u_1$  to  $u_{k+1}$  grows along with k, since in the subsequent constructions time will essentially be considered only modulo a certain constant  $\pi_{\text{PM}}$ .

We next construct for any given real numbers  $\delta, \lambda > 0$  and  $\kappa < 0$  "inhibition modules"  $I_{\delta,\kappa,\lambda}$  and  $I^{\delta,\kappa,\lambda}$ .  $I_{\delta,\kappa,\lambda}$  can be used to transmit to any desired neuron v a volley of IPSPs that sum up to a potential which changes from its initial value 0 to some value  $\leq \kappa$  within a time interval of length  $\delta$ , and then maintains a value  $\leq \kappa$  for at least the following time interval of length  $\lambda$ .  $I_{\delta,\kappa,\lambda}$  consists of a neuron v that transmits EPSPs simultaneously to several "relay neurons"  $v_1,\ldots,v_l$ , which are triggered by this EPSP to send an IPSP to some given neuron v. If l and the delays between the neurons are chosen appropriately [as a function of  $\delta,\kappa,\lambda,\varepsilon^l(\delta)$  and the parameter  $\tau_1$ ], this module will have the desired effect on neuron v.

Dually, one can also build for any  $\delta, \lambda > 0$  and  $\kappa < 0$  an inhibition module  $I^{\delta,\kappa,\lambda}$  that sends IPSPs to any specified neuron v whose sum stays  $< \kappa$  for a time interval of length  $\ge \lambda$ , and then returns to 0 within a time

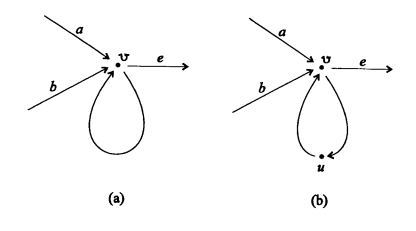


Figure 4: Graph structure of an oscillator consisting of one neuron (a) and two neurons (b).

interval of length  $\leq \delta$ . Here we exploit the fact that according to condition (5) the function  $\varepsilon^{l}(x)$  is nondecreasing and strictly negative for  $x \in [\tau_{2}, \tau_{3})$ .

**2.3 Oscillators.** Consider subgraphs of an SNN of the structure shown in Figure 4. Both types of subgraphs can be used to build an oscillator. The first one is somewhat simpler, but we will not use it in our construction since it would require a self-referential edge  $\langle v.v \rangle \in E$ .

In the second type of oscillator (Fig. 4b) we assume that  $w_{u,v}, w_{v,u} \ge \Theta(0)/\varepsilon_{\max}$ , and that both  $\varepsilon_{u,v}$  and  $\varepsilon_{v,u}$  are EPSP response functions. Thus after an initial EPSP through edge a both neurons will fire periodically. More precisely, v will fire at times  $t_0 + i \cdot \pi$  for  $i = 1, 2, \ldots$ , until it is halted by an IPSP through edge v. We refer to v as the oscillation period of this oscillator.

We will distinguish one such oscillator as the "pacemaker" for the constructed SNN, which we denote by PM. We write  $\pi_{PM}$  for its oscillation period. We assume that the oscillation of PM is started at "time 0" by the first input spike to the SNN, and that it continues without interruption throughout the computation of the SNN. PM emits EPSPs through edge e, which will then be broadcast as a timing standard throughout the SNN. We will say in the following that some other neuron v in the SNN fires "at unit time" or "synchronously" if the considered firing of v occurs at a time point t of the form v of v of the form v of the form

In  $\mathcal{N}_{TM}(d)$  we will use oscillators in two ways as storage devices. First we use them as "registers" for storing a bit (via their two states dormant/oscillating), for example in the control of  $\mathcal{N}_{TM}(d)$ . Second we

use oscillators O with oscillation period  $\pi_{PM}$  to store arbitrary numbers  $\varphi \in [0, \pi_{PM}]$  via their *phase difference* to PM (i.e., neuron v of oscillator O fires at time points of the form  $i \cdot \pi_{PM} + \varphi$  with  $i \in \mathbb{N}$ ). In this way oscillators can for example store the time difference between two input spikes to the SNN, and the program and tape content of a simulated Turing machine, respectively.

**2.4 Synchronization Modules.** A characteristic feature of a computation on a feedforward Boolean circuit of the usual type is that the *timing* of its computation steps is *independent* of the *values* of the bits that occur in the computation. For example, the timing of the output signal of an OR gate does not depend on the values of its input bits. This feature is very useful, since with its help one can arrange that all input bits for Boolean gates on higher levels of the circuit arrive simultaneously, and therefore it allows us to build complex circuits from simple modules.

If one wants to carry out computations on an SNN with single spikes, one would like to interpret the firing of a neuron at a certain time as the bit "1" and nonfiring as "0." Thus one might, for example, want to simulate an OR gate by a neuron v that fires whenever it receives at least one EPSP. However, when that neuron v receives two EPSPs simultaneously (corresponding to two input bits being 1) it would in general fire slightly earlier than in a situation where it receives just a single EPSP. This effect is a consequence of having EPSP response functions  $\varepsilon_{uv}(x)$  that are not piecewise constant. In addition, if v has already fired just before, then the fact that  $\Theta(x)$  is in general not piecewise constant also contributes to this effect. Unfortunately this effect makes it impossible to simulate on an SNN in a straightforward manner a multilayer Boolean circuit (where the bit "1" is signaled by a spike, and "0" by the absence of a spike at the corresponding time): the input "bits" for neurons that simulate Boolean gates on higher layers of the circuit will in general not arrive at the same time. Furthermore it is *not* possible to correct this problem by employing delay modules of the type that we had constructed in Section 2.2, since the required length of the delays depends on the current values of the input bits.

We will solve this problem with the help of the here constructed *synchronization module*. In fact, we will show in the next section that with the help of this module an SNN suddenly gains the full computational power of a Boolean feedforward *threshold circuit*, and therefore is able to carry out within a small number of "cycles" substantially more complex computations than a regular Boolean circuit.

On first sight it appears to be impossible to build a synchronization module without postulating the existence of an EPSP response function that has segments of length  $\geq \pi_{PM}$  where it is constant, or increases or decreases linearly. However the following "double-negation trick" allows us to build a synchronization module without any additional assumptions.

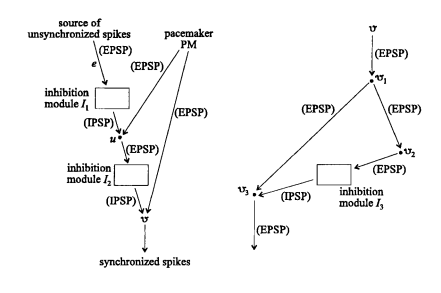


Figure 5: Structure of a synchronization module.

Consider the graph of an SNN on the left hand side of Figure 5. We arrange that as long as no EPSP is transmitted through its "input edge" e, the neuron u fires regularly with period  $\pi_{\rm PM}$  as a result of EPSPs from the pacemaker PM. These EPSPs induce the inhibition module  $I_2$  to send IPSPs to neuron v that "cancel out" the EPSPs that arrive at v directly from PM. Therefore in the absence of an input through edge e this neuron v does not fire.

Assume now that at some arbitrary time point an (unsynchronized) EPSP arrives through edge e. This EPSP triggers the inhibition module  $I_1$ , which then sends out IPSPs that prevent neuron u from firing for a time interval of some fixed length  $> \pi_{PM}$ . Therefore at least one of the EPSPs that arrive at neuron v from PM is not cancelled out by IPSPs from the inhibition module  $I_2$ , and neuron v emits at least one synchronized spike (i.e., v fires at least once, and with a proper choice of delays only at unit times of the form  $i \cdot \pi_{PM}$  with  $i \in \mathbb{N}$ ).

A closer look shows that the mechanism of this module is in fact a bit more delicate. It can, in principle, happen that at neuron u the beginning or the end of a negative potential from  $I_1$  coincides with an EPSP from PM in such a way that it leads to a small shift  $\rho$  in some firing time of u (besides canceling other firings of u). This could shift the time interval of the activity of  $I_2$  by a certain amount  $\rho$ . One has to make sure that this shift cannot lead to a competition at neuron v between the negative

potential from  $I_2$  and the EPSP from PM that results in an unsynchronized firing of v. One can solve this technical problem by designing  $I_1$  and  $I_2$  so that their output is the superposition of the output of a module  $I_{\delta,\kappa,\lambda}$  and of a module  $I^{\delta,\kappa,\lambda}$ . In this way their strongly negative output potential (of value  $< \kappa$ ) both builds up and disappears at neuron v within time intervals of length  $\delta$ . This parameter  $\delta$  provides then an upper bound for the length  $\rho$  of the possible time shifts of these negative potentials. By choosing  $\delta$  sufficiently small (and by arranging the lengths and delays of these inhibitions appropriately), for any arrival time of an input spike through edge e and for any EPSP from PM the resulting inhibition from  $I_2$  either cancels the corresponding firing of v, or it lets v fire without shifting its firing time (canceling some other firings of v instead). For that purpose one chooses the weight  $w \in [0,1]$  on the edge from PM to v so that the resulting function  $w \cdot \varepsilon^E$  crosses  $\Theta(0)$  while it is in the middle of its linearly increasing segment [see condition (3) of our basic assumptions].

The timing of this synchronization module can be specified with more precision as soon as one selects concrete response and threshold functions that satisfy our basic assumptions. However, the preceding analysis shows that it will do its job in any case. One should keep in mind that our basic assumptions are relatively weak. For example, they do not even prescribe the relationships between the sizes of the parameters  $\sigma_3$ ,  $\tau_3$ , and  $\tau_{\rm end}$  that denote the lengths of the nontrivial segments of the response and threshold functions.

It turns out that the previously described module may output not just one, but a larger finite number of synchronized spikes as a result of one unsynchronized input spike. This effect causes no serious problem in our subsequent applications of this module (and it might occasionally be helpful for speeding up a computation), but it is easier to verify a construction if this module never outputs more than *one* synchronized spike for each input spike. This additional requirement can be satisfied by adding after neuron v a device with three neurons  $v_1, v_2, v_3$  as indicated in the right-hand side of Figure 5. With suitably chosen delays and parameters for its inhibition module  $I_3$ , this device removes all except the first spike from any sequence of successive synchronized spikes. It lets the first one of these spikes emerge from neuron  $v_3$  as a *single synchronized* output spike.

2.5 Simulation of Boolean Threshold Circuits by SNNs. If one just wants to simulate in a straightforward manner the control of a Turing machine on an SNN, one can reserve one neuron for each possible state of the control, and simulate state transitions with the help of neurons that simulate Boolean AND and OR gates. However Horne and Hush (1994) have pointed out that many fewer neurons are needed if one simulates the control with the help of a Boolean feedforward threshold circuit with gates of unbounded fan-in (see Section 2.8). In addition, the ability of

SNNs to simulate threshold circuits in an efficient manner is of substantial interest for various other reasons (see Corollary 2.4 and the lower bound for the VC dimension of SNNs in Maass 1994b). Therefore we describe the simulation of a *threshold circuit* on an SNN, rather than considering first the simulation of the special case of a Boolean circuit with gates of bounded fan-in (which would suffice for the proof of Theorem 2.1).

A feedforward Boolean threshold circuit (threshold circuit for short) consists of a directed acyclic graph with nodes of arbitrary fan-in, that correspond to linear threshold gates (threshold gates for short) with arbitrary weights. A threshold gate with fan-in m computes a threshold function of the form

$$\{0,1\}^m \ni \langle x_1,\ldots,x_m \rangle \mapsto T^{\underline{\alpha}}(x_1,\ldots,x_m) = \begin{cases} 1, & \text{if } \sum_{i=1}^m \alpha_i \cdot x_i \ge \alpha_0 \\ 0, & \text{otherwise} \end{cases}$$

with arbitrary parameters  $\alpha_0, \ldots, \alpha_m \in \mathbf{R}$  (or equivalently:  $\alpha_0, \ldots, \alpha_m \in \mathbf{Z}$ ).

It is obvious that the common Boolean operations AND, OR, NOT are special cases of threshold functions. Therefore the common types of feedforward Boolean circuits (even with ANDs and ORs of arbitrarily large fan-in) are special cases of threshold circuits. Hence, since every Boolean function can be defined by a Boolean formula in disjunctive normal form (see, e.g., Lewis and Papadimitriou 1981) it is clear that *every* Boolean function can be computed by a threshold circuit of depth 2 (i.e., with one "hidden" layer).

There are several different possibilities for simulating a threshold circuit on an SNN, providing subtle tradeoffs between the amount of demands imposed on the response functions, the noise robustness of the construction, and the number of neurons needed for the simulation. We describe one simple construction based on our basic assumptions, and we will indicate a variation in Section 4.

Consider first a "monotone" threshold function, i.e., a threshold function  $T^{\underline{\alpha}}$  with  $\alpha_i \geq 0$  for all "weights"  $\alpha_1, \ldots, \alpha_m$ . If  $\alpha_0 \leq 0$  then  $T^{\underline{\alpha}}$  always outputs "1," and is therefore superfluous. Hence we may assume that  $\alpha_0 > 0$ .

By condition (2) each EPSP response function  $\varepsilon_{u,v}$  has some maximal value  $\varepsilon_{\max} > 0$  that does not depend on u or v. We employ for the computation of  $T^{\underline{\alpha}}$  on an SNN m+1 neurons  $u_1,\ldots,u_m$  and v with  $\{u:\langle u,v\rangle\in E\}=\{u_1,\ldots,u_m\}$ . We assume that all response functions  $\varepsilon_{u_i,v}$  are EPSPs and that the weights  $w_{u_i,v}$  are chosen so that  $w_{u_i,v}\cdot\varepsilon_{\max}=\alpha_i\cdot\Theta(0)/\alpha_0$  (slightly larger values should be chosen if one has to deal with imprecision). Furthermore, we assume that the "delays"  $\Delta_{u_i,v}$  are chosen to be the same for  $i=1,\ldots,m$ . Consider then some arbitrary set  $S\subseteq\{1,\ldots,m\}$ . Assume that the neurons  $u_i$  with  $i\in S$  fire simultaneously at some time  $t_0$ , that the neurons  $u_i$  with  $i\in \{1,\ldots,m\}-S$  never fire in

the time interval  $[t_0 - \sigma_3, \ t_0 + \sigma_3]$ , and that neuron v did not fire in the time interval  $[t_0 + \Delta_{u_1,v} - \tau_{\mathrm{end}}, \ t_0 + \Delta_{u_1,v}]$ . Then v fires at some point in the time interval  $(t_0 + \Delta_{u_1,v}, \ t_0 + \Delta_{u_1,v} + \sigma_3)$  if and only if  $\sum_{i \in S} w_{u_i,v} \cdot \varepsilon_{\mathrm{max}} \geq \Theta(0)$ . The latter inequality is equivalent to  $\sum_{i \in S} \alpha_i \cdot \Theta(0)/\alpha_0 \geq \Theta(0)$ , hence to  $\sum_{i \in S} \alpha_i \geq \alpha_0$ . Thus we have constructed a module of an SNN that computes an arbitrary monotone Boolean threshold function  $T^{\alpha}$ .

This module has the disadvantage that its proper functioning is guaranteed only if all  $u_i$  with  $i \in S$  fire at a *common* time  $t_0$ . On the other hand the firing time of v depends not only on  $t_0$ , but also on S (i.e., on its "input bits"). In general a larger set S gives rise to a slightly earlier firing time of v (because the function  $\varepsilon^E$  does not jump immediately from 0 to  $\varepsilon_{\max}$ ). Obviously these two facts together cause problems if one wants to use compositions of the previously constructed module to simulate a multilayer monotone threshold circuit (i.e., a threshold circuit where all gates compute *monotone* threshold functions). Therefore one has to use synchronization modules between any two layers of modules to simulate a *monotone threshold circuit* on an SNN.

We will now describe the simulation of an *arbitrary* threshold circuit C, where threshold functions  $T^{\alpha}$  with "weights"  $\alpha_i$  of arbitrary sign are computed by gates of C. It is well-known (see Hajnal *et al.* 1993) that such a circuit C can be simulated by a *monotone* threshold circuit  $C_{\text{mon}}$  of the same depth, provided that  $C_{\text{mon}}$  also receives for each Boolean input variable  $x_i$  its negation  $1-x_i$ . Proceeding from the input layer to the ouput layer one can then replace each threshold gate g of C by two gates that both compute *monotone* threshold functions: one of them provides the same output as g, and the other one provides the negation of that output.

Thus in order to simulate C on an SNN, one needs in addition to the preceding construction a preprocessing device that computes the negation 1-x for each input bit  $x \in \{0,1\}$  under the considered bit encoding (where "x = 1" is encoded by a firing of a neuron u at a certain time t, and "x = 0" by the nonfiring of u within a certain time interval around t). For that purpose one connects u to an inhibition module whose outputs cancel out an EPSP from PM at another neuron u' (similarly as in Section 2.4). Then u' will fire if and only if it is not inhibited via a firing of u, hence u' computes "1-x."

**2.6 Modules for Comparisons and Multiplication of Phases with Arbitrary Constants.** We will construct in this section a module for an SNN that can compare the phase difference  $\varphi$  of an oscillator O with some given constant  $\alpha$  [COMPARE( $\geq \alpha$ )], and a module that can multiply  $\varphi$  with some given constant  $\beta$  [MULTIPLY( $\beta$ )]. Such modules [more precisely: modules for the operation COMPARE( $\geq 2^{-1-c}$ ) for a certain constant c, as well as modules for MULTIPLY(2) and MULTIPLY(1/2)] will be needed in the next section to simulate a stack on an SNN.

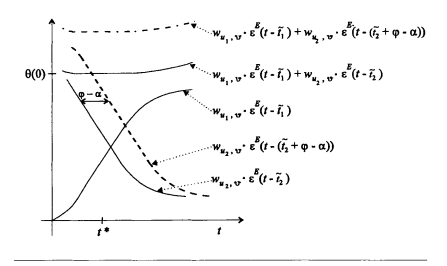


Figure 6: Mechanism of the module for COMPARE( $\geq \alpha$ ).

Let  $\alpha \in [0,L/2]$  be some arbitrary real constant. We construct a module that can decide whether the phase difference  $\varphi \in [0,L/2]$  between PM and some oscillator O with oscillation period  $\pi_{\text{PM}}$  is  $\geq \alpha$ . More precisely, this module for the operation COMPARE( $\geq \alpha$ ) will send out a spike within some time interval of some given length 2D if and only if  $\varphi \geq \alpha$ . Consider neurons  $u_1,u_2$ , and v with  $\langle u_i,v\rangle \in E$  for i=1,2. Assume that  $u_1$  is induced to fire at a certain time  $t_1$  by a spike from the pacemaker PM. Furthermore, assume that  $u_2$  is induced to fire at a certain time  $t_2$  by a spike from the oscillator O. Finally we assume that the delays  $\Delta_{u_1,v}$  and  $\Delta_{u_2,v}$  have been chosen so that in the case  $\varphi = \alpha$  one has for  $\tilde{t}_i := t_i + \Delta_{u_i,v}$  that there exists some  $t^* \geq \max(\tilde{t}_1,\tilde{t}_2)$  so that  $t^* - \tilde{t}_1 = \sigma_1$  and  $t^* - \tilde{t}_2 = \sigma_2$ . We choose weights  $w_{u_i,v} > 0$  so that  $w_{u_1,v} \cdot s_{up} = w_{u_2,v} \cdot s_{down}$  and  $w_{u_1,v} \cdot \varepsilon^E(\sigma_1) + w_{u_2,v} \cdot \varepsilon^E(\sigma_2) = \Theta(0)$  (see Fig. 6).

According to our general convention at the beginning of this section we actually have to replace in the case  $w_{u_i,v} > 1$  the neuron  $u_i$  by an assembly of  $\lceil w_{u_i,v} \rceil$  neurons with weights from [0,1] on their edges to v. However, for the sake of simplicity, we will ignore this trivial complication in the following.

We arrange that for an arbitrarily given parameter D > 0 inhibition modules  $I_{\delta,\kappa,\lambda}$  and  $I^{\delta,\tilde{\kappa},\tilde{\lambda}}$  (with suitable values of their parameters) are triggered by spikes from PM to send IPSPs to v so that v is not able to fire within the time intervals  $[t^*-L/2-D,\ t^*-L/2)$  and  $(t^*+L/2,\ t^*+L/2+D]$  even if the firing time  $t_2$  of neuron  $u_2$  is arbitrarily shifted, but so that

these inhibition modules have no effect on the potential  $P_v$  at neuron v during the time interval  $[t^* - L/4, t^* + L/4]$ .

Consider now what happens if the phase difference  $\varphi$  of the oscillator O is not fixed at  $\varphi=\alpha$ , but assumes any value in [0,L/2]. Then by choice of the parameters  $w_{u_1,v}, w_{u_2,v}$ , and  $t^*$ , and by the conditions (3) and (4) of our basic assumptions, the sum of the EPSPs from  $u_1$  and  $u_2$  at neuron v has in any case a constant value within the time interval  $[t^*-L/2, t^*+L/2]$ . Furthermore, this constant value is  $\geq \Theta(0)$  if and only if  $\varphi \geq \alpha$ . Hence the neuron v will fire within the time interval  $[t^*-L/2, t^*+L/2]$  if and only if  $\varphi \geq \alpha$ . Furthermore, by the choice of the inhibition modules the neuron v fires within the time interval  $[t^*-L/2, t^*+L/2]$  if and only if it fires within the time interval  $[t^*-D, t^*+D]$ .

We now assume that some arbitrary real number  $\beta > 0$  is given, and we construct a module that carries out the operation MULTIPLY( $\beta$ ). This module also consists of neurons  $u_1, u_2, v$  with  $\langle u_i, v \rangle \in E$  for i = 1, 2 so that  $u_1$  is triggered to fire at some time  $t_1$  by a spike from the pacemaker PM, and  $u_2$  is triggered to fire at some time  $t_2$  by a spike from an oscillator O that has oscillation period  $\pi_{\text{PM}}$  and some phase difference  $\varphi \in [0, \min(L/2, L/2\beta)]$  to PM. We want to achieve that for any value  $\varphi \in [0, \min(L/2, L/2\beta)]$  of this phase difference the "output neuron" v of this module fires at a time  $t + \beta \cdot \varphi$ , where t does not depend on  $\varphi$ .

The construction of the module for the operation MULTIPLY( $\beta$ ) is slightly different for the two cases  $\beta > 1$  and  $\beta \in (0,1)$ . We consider first the case  $\beta > 1$ . Assume for the moment that the phase difference  $\varphi \in [0, L/2\beta]$  between O and PM has value 0, and choose delays  $\Delta_{u,v}$  so that there exists for  $\tilde{t}_i := t_i + \Delta_{u_i,v}$  some  $t^* \ge \max(\tilde{t}_1, \tilde{t}_2)$  with  $t^* - \tilde{t}_1 = \sigma_2$  and  $t^* - \tilde{t}_2 = \sigma_1$ . Furthermore, we choose weights  $w_{u,v} > 0$  so that

$$w_{u_1,v} \cdot \varepsilon^{E}(t^* - \tilde{t}_1) + w_{u_2,v} \cdot \varepsilon^{E}(t^* - \tilde{t}_2) = \Theta(0)$$
(2.1)

and

$$\beta = \frac{w_{u_2,v} \cdot s_{\text{up}}}{w_{u_2,v} \cdot s_{\text{up}} - w_{u_1,v} \cdot s_{\text{down}}}.$$
(2.2)

Since  $\beta > 1$ , equation 2.2 implies that  $0 < w_{u_1,v} \cdot s_{\text{down}} < w_{u_2,v} \cdot s_{\text{up}}$ . Hence we have

$$w_{u_1,v} \cdot \varepsilon^E(t^* - \tilde{t}_1 + z) + w_{u_2,v} \cdot \varepsilon^E(t^* - \tilde{t}_2 + z) < \Theta(0)$$
 for all  $z \in [-L, 0)$ . (2.3)

We would like to arrange that v does not fire during the time interval  $[t^* - \tau_{\rm end}, t^*)$ , where  $\tau_{\rm end}$  has the property that  $\Theta(x) = \Theta(0)$  for all  $x \in [\tau_{\rm end}, \infty)$  [according to condition (1)]. Furthermore, we would like to make sure that this property holds even if the firing of  $u_2$  is delayed by some arbitrary amount  $\varphi \in [0, L/2\beta]$ . However, even if one assumes that only the considered EPSPs from  $u_1$  and  $u_2$  are influencing  $P_v(t)$ , this assumption allows us to derive this fact only with the help of equation 2.3 for the interval  $[t^* - L/2, t^*)$ , since we did not make more detailed assumptions about the shape of the function  $\varepsilon^E$ . Therefore we arrange that

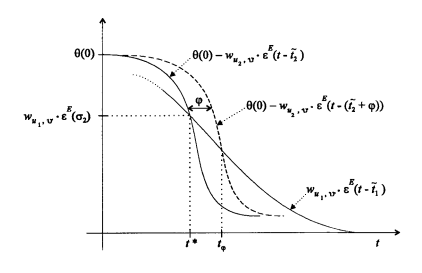


Figure 7: Multiplication of a phase  $\varphi$  with  $\beta > 1$  (i.e.,  $t_{\varphi} - t^* = \beta \cdot \varphi$ ).

at a suitable time an inhibition module  $I^{L/2,\kappa,\tau_{\rm end}}$  sends IPSPs to v, which makes it impossible for v to fire during the time interval  $[t^*-\tau_{\rm end},t^*-L/2)$  (no matter at what time  $u_2$  fires), but which does not influence the potential  $P_v(t)$  at times  $t \geq t^*$ . Furthermore, we arrange that no other EPSPs or IPSPs contribute to  $P_v(t)$  for  $t \in [t^*-\tau_{\rm end},t^*]$ . In this way v can fire during the time interval  $[t^*-\tau_{\rm end},t^*)$  (even if the firing of  $u_2$  is delayed by some  $\varphi \in [0,L/2\beta]$ ). Therefore in the case  $\varphi=0$  our assumption (equation 2.1) implies that neuron v will fire at time  $t^*$ .

We now consider what will change if the firing of  $u_2$  at time  $t_2$  is replaced by a slightly later firing at time  $t_2 + \varphi$ , whereas the firing time of  $u_1$  and of the inhibition module remain unchanged. We will show that for any  $\varphi \in (0, L/2\beta]$  this delay will cause a somewhat delayed firing of v (see Fig. 7). Consider the time point  $t_{\varphi}$ , which is defined by the equation

$$w_{u_1,v} \cdot \varepsilon^{E}(t_{\varphi} - \tilde{t}_1) + w_{u_2,v} \cdot \varepsilon^{E}[t_{\varphi} - (\tilde{t}_2 + \varphi)] = \Theta(0). \tag{2.4}$$

By equation 2.1 and conditions (3) and (4) of our basic assumptions we have for  $t_{\varphi} - t^* \in [-L, L]$ 

$$w_{u_1,v} \cdot \varepsilon^{E}(t_{\varphi} - \tilde{t}_1) = w_{u_1,v} \cdot \varepsilon^{E}(t^* - \tilde{t}_1) - w_{u_1,v} \cdot s_{\text{down}} \cdot (t_{\varphi} - t^*)$$
 (2.5)

and for  $\varphi, t_{\varphi}$  with  $t_{\varphi} - t^* - \varphi \in [-L, L]$  we have that

$$w_{u_2,v} \cdot \varepsilon^{E}[t_{\varphi} - (\tilde{t}_2 + \varphi)] = w_{u_2,v} \cdot \varepsilon^{E}(t^* - \tilde{t}_2) + w_{u_2,v} \cdot s_{\text{up}} \cdot (t_{\varphi} - t^* - \varphi). \tag{2.6}$$

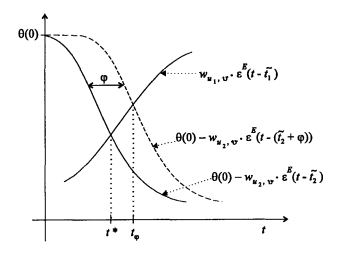


Figure 8: Multiplication of a phase  $\varphi$  with  $\beta \in (0,1)$ .

These two equations in conjunction with 2.1, 2.2, and 2.4 imply that

$$t_{\varphi} - t^* = \beta \cdot \varphi.$$

It is obvious that for  $\varphi \in [0, L/2\beta]$  one has that  $\beta \cdot \varphi$ ,  $\beta \cdot \varphi - \varphi \in [-L, L]$ . Furthermore, it is clear from our construction that v cannot fire during the time interval  $[t^* - \tau_{\text{end}}, t^* + \beta \cdot \varphi]$ . Therefore  $t_{\varphi} := t^* + \beta \cdot \varphi$  is in fact the firing time of v if  $u_2$  fires at time  $t_2 + \varphi$ . Hence the described module carries out the operation MULTIPLY( $\beta$ ) in case that  $\beta > 1$ .

To carry out the operation MULTIPLY( $\beta$ ) for some arbitrarily given  $\beta \in (0,1)$  we just change the delay  $\Delta_{u_1,v}$  in the previously described module so that  $t^* - \tilde{t}_1 = \sigma_1$  (instead of  $t^* - \tilde{t}_1 = \sigma_2$ ) (see Fig. 8). We choose weights  $w_{u_i,v} > 0$  so that equation 2.1 holds and

$$\beta = \frac{w_{u_2,v}}{w_{u_2,v} + w_{u_2,v}} \ . \tag{2.7}$$

As before, we consider the time point  $t_{\varphi}$  that is defined by equation 2.4. Then equation 2.6 holds, but instead of 2.5 we have

$$w_{u_1,v} \cdot \varepsilon^E(t_{\varphi} - \tilde{t}_1) = w_{u_1,v} \cdot \varepsilon^E(t^* - \tilde{t}_1) + w_{u_1,v} \cdot s_{\mathrm{up}} \cdot (t_{\varphi} - t^*).$$

The latter two equations in conjunction with 2.1, 2.4 and 2.7 imply that

$$t_{\varphi} - t^* = \beta \cdot \varphi.$$

Hence the described module carries out the operation MULTIPLY( $\beta$ ) for an arbitrarily given  $\beta \in (0,1)$ .

**2.7** Simulation of a Stack with Unlimited Capacity by an SNN of Fixed Size. The simulation of a stack (also called pushdown store, of first in–last out list) is the most delicate part of the construction of  $\mathcal{N}_{TM}(d)$ , since it requires the construction of a module in which the lengths  $\ell$  of the bit-strings  $\langle b_1, \ldots, b_\ell \rangle$  that are stored and manipulated are in general much larger than the number of neurons in this module (in fact,  $\ell$  can be arbitrarily large). Of course  $\mathcal{N}_{TM}(d)$  needs to have a component with this property, since otherwise the SNN  $\mathcal{N}_{TM}(d)$  (which will consist of a fixed finite number of neurons) cannot simulate the computations of Turing machines that involve tape inscriptions of arbitrary finite length. The content  $\langle b_1, \ldots, b_\ell \rangle \in \{0,1\}^*$  of a stack S (where  $b_1$  is the symbol on top of the stack) will be stored in the form of the phase difference

$$\varphi_S = \sum_{i=1}^{\ell} b_i \cdot 2^{-i-c}$$

of a special oscillator  $O_S$ . More precisely, we assume that  $O_S$  fires with the same oscillation period  $\pi_{PM}$  as the pacemaker PM, but with a delay  $\varphi_S$ . The parameter  $c \in \mathbf{R}^+$  is some arbitrary constant that is sufficiently large so that  $2^{-c} \leq \min(L/2, \pi_{PM})$ .

We will now describe the mechanisms for simulating the stack operations POP and PUSH on a bit string  $(b_1, \ldots, b_\ell)$  that is stored in  $\varphi_s$ .

The stack operation POP determines the value of the top-bit  $b_1$ , and then replaces the stack content  $\langle b_1, \ldots, b_\ell \rangle$  by  $\langle b_2, \ldots, b_\ell \rangle$ . In an SNN one can determine the value of  $b_1$  from  $\varphi_S$  by testing whether  $\varphi_S \geq 2^{-1-c}$ . For that purpose one employs a module that carries out the operation COMPARE( $\geq 2^{-1-c}$ ) (see the preceding section).

To change the phase-difference  $\varphi_S$  from  $\sum_{i=1}^\ell b_i \cdot 2^{-i-\epsilon}$  to  $\sum_{i=1}^{\ell-1} b_{i+1} \cdot 2^{-i-\epsilon}$  one first replaces  $\varphi_S$  by  $\sum_{i=2}^\ell b_i \cdot 2^{-i-\epsilon}$ . For the case  $b_1 = 1$  this can be carried out by directing an EPSP from  $O_S$  through a suitable delay module, by halting simultaneously the oscillation of  $O_S$  with the help of an inhibition module, and by restarting the oscillation of  $O_S$  with an EPSP from the considered delay module. Note that we can employ at this point a simple delay module as described in Section 2.2, because in the case  $b_1 = 1$  the length of the desired shift of the phase difference does not depend on its current value.

It remains to carry out a SHIFT-LEFT operation, which replaces the phase difference  $\sum_{i=2}^{\ell} b_i \cdot 2^{-i-c}$  by

$$2 \cdot \sum_{i=2}^{\ell} b_i \cdot 2^{-i-c} = \sum_{i=1}^{\ell-1} b_{i+1} \cdot 2^{-i-c}.$$

This operation cannot be implemented by a delay-module, since it has to shift the phase difference by an amount that depends on the values of  $\ell$  and  $b_2, \ldots, b_\ell$ . Instead, we have to employ a module that carries out the operation MULTIPLY(2) (see Section 2.6).

To simulate the stack operation PUSH one has to replace for a given  $b_0 \in \{0,1\}$  the current phase-difference  $\varphi_S = \sum_{i=1}^\ell b_i \cdot 2^{-i-\epsilon}$  of the oscillator  $O_S$  by  $\sum_{i=1}^{\ell+1} b_{i-1} \cdot 2^{-i-\epsilon}$ . Our simulation of PUSH consists of two separate parts: a SHIFT-RIGHT operation that changes the current phase difference to  $\sum_{i=2}^{\ell+1} b_{i-1} \cdot 2^{-i-\epsilon}$ , and a subsequent ADD( $\gamma$ ) operation that adds  $\gamma := b_0 \cdot 2^{-1-\epsilon}$  to this phase difference. Obviously ADD( $\gamma$ ) can be implemented in an analogous way as the subtraction of  $b_1 \cdot 2^{-1-\epsilon}$  from  $\varphi_S$  in the previously described simulation of POP.

Thus it just remains to simulate a SHIFT-RIGHT operation, i.e., to replace the phase difference  $\varphi_S = \sum_{i=1}^{\ell} b_i \cdot 2^{-i-c}$  of size  $\leq L/2$  by  $\varphi_S/2 = \sum_{i=2}^{\ell+1} b_{i-1} \cdot 2^{-i-c}$ . For that purpose we employ a module for the operation MULTIPLY(1/2), as constructed in the preceding section.

### 2.8 Simulation of an Arbitrary Fixed Turing Machine by an SNN.

We will show in this section that the previously constructed modules suffice to construct for any given Turing machine M an SNN  $\mathcal{N}_M$  (whose structure may depend on M) that can simulate M in real-time. According to the notion of a *real-time computation* (see Section 1) we assume that the given Turing machine M processes a sequence  $(\langle x(j),y(j)\rangle)_{j\in\mathbb{N}}$  with  $x(j),y(j)\in\{0,1\}^*$  in real-time. We assume that the inputs x(j) are presented to M on a read-only input tape, and the outputs y(j) are written by M on some write-only output tape. We will assume that the simulating SNN  $\mathcal{N}_M$  receives each input  $x(j)\in\{0,1\}^*$  in the form of a time difference  $\varphi$  between two input-spikes, with  $\varphi=\sum_{i=1}^\ell b_i\cdot 2^{-i-\epsilon}$  for  $x(j)=\langle b_1,\ldots,b_\ell\rangle$ . We will arrange that  $\mathcal{N}_M$  delivers its outputs y(j) in the same form (as a time difference between two output spikes).

It is easy to see that any Turing machine M, with any finite number d of two-way infinite read/write-tapes, can be simulated in real-time by a similar machine which has 2d stacks, but no tapes (see, e.g., Hopcroft and Ullman 1979). We will call the latter type of machine also a Turing machine. In this simulation one uses two stacks for the simulation of each tape: one stack for simulating the part of the tape that lies to the left of the current position of the tape-head and another stack for simulating the part of tape to the right of the tape-head.

In principle it would suffice to consider a Turing machine with 1 tape (or 2 stacks), since this type of Turing machine can simulate any other Turing machine (although not in real time). However, it is known that various concrete problems (especially several pattern-matching problems) can be solved faster on a Turing machine that has more than one tape (see, e. g., Hopcroft and Ullman 1979; Maass 1985; and Maass *et al.* 1987). Therefore, and because it does not cause any extra work, we simulate an arbitrary Turing machine M with any number k of stacks by an SNN  $\mathcal{N}_M$ .

At any computation step the Turing machine *M* may POP or PUSH a symbol on each of its *k* stacks. We assume for simplicity that the stackalphabet of *M* is binary (i.e., *M* can push 0 or 1 on each stack, and pop

a binary symbol, or receive the signal "bottom-of-stack" if the stack is empty.) Furthermore, we assume that the input for the computation of M is given as the initial content of the first one of the k stacks, and that the output of M consists of the final content of the last one of the k stacks (at the moment when the machine halts).

If Q is the (finite) set of states of M, then after assigning a number in binary notation to each state in Q the transition function of M can be encoded by a function  $F_M: \{0,1\}^{\lceil \log |Q| \rceil + k} \to \{0,1\}^{\lceil \log |Q| \rceil + k}$ . We assume here that the state of M indicates on which of the stacks a POP or PUSH has to be carried out.

Thus to simulate the finite control of M by an SNN, it suffices to employ a module that can compute an arbitrary given function from  $\{0,1\}^{\lceil \log |Q| \rceil + k}$  into itself. We assume here that the  $\lceil \log |Q| \rceil + k$  input and output bits of this function are stored in a corresponding number of oscillators with two states (dormant/oscillating). According to Lupanov (1973), one can compute any function  $F: \{0,1\}^{\lceil \log |Q| \rceil + k} \to \{0,1\}^{\lceil \log |Q| \rceil + k}$ on a feedforward threshold circuit with  $O(|Q|^{1/2} \cdot 2^{k/2})$  gates. In addition, Horne and Hush (1994) have shown that any such function F can be computed by a threshold circuit of depth 4 with  $O[|Q|^{1/2} \cdot 2^{k/2} \cdot (\log |Q| + k)]$ gates, using only weights and thresholds from  $\{-1,0,1\}$ . Hence our previously described simulation of an arbitrary threshold circuit on an SNN in Section 2.5 allows us to simulate in  $N_M$  the finite control of M with a module of  $O[|Q|^{1/2} \cdot 2^{k/2}]$  neurons (provided the SNN may use arbitrarily large weights). Furthermore, the quoted result by Horne and Hush in conjunction with our construction in Section 2.5 implies that with  $O[|Q|^{1/2} \cdot 2^{k/2} \cdot (\log |Q| + k)]$  neurons one can implement in  $\mathcal{N}_M$  the finite control of M in such a way that only very simple weights from [0,1] are needed in  $\mathcal{N}_M$ , and that the simulation of each computation step of M requires only O(1) "machine-cycles" of  $\mathcal{N}_{M}$ . More precisely, each computation step of M is simulated by  $\mathcal{N}_M$  in a time interval in which the pacemaker PM fires < K times, where K is some absolute constant that is independent of |O|, k, the length of the current input of M, and the number of the previously simulated computation steps of M.

Apart from the finite control component, the SNN  $\mathcal{N}_M$  consists of a module of O(1) neurons for each of the k stacks, and O(1) neurons that implement the pacemaker PM. In addition  $\mathcal{N}_M$  uses  $O(\log |Q| + k)$  neurons for other oscillators that serve as temporary registers for bits. Thus  $\mathcal{N}_M$  consists altogether of at most  $O[|Q|^{1/2} \cdot 2^{k/2} \cdot (\log |Q| + k)]$  neurons, and the simulation of any computation step of M involves at most  $O[|Q|^{1/2} \cdot 2^{k/2} \cdot (\log |Q| + k)]$  firings of neurons in  $\mathcal{N}_M$ . After  $\mathcal{N}_M$  has simulated every computation step of M on the current input  $x(j) \in \{0,1\}^*$ , it has generated on an oscillator  $O_S$ , which corresponds to the stack S on which M writes its output  $y(j) = \langle \tilde{b}_1, \ldots, \tilde{b}_{\tilde{\ell}} \rangle$ , a phase-difference  $\varphi_S = \sum_{i=1}^{\tilde{\ell}} \tilde{b}_i \cdot 2^{-i-c}$  with regard to the pacemaker PM.  $\mathcal{N}_M$  outputs two spikes, where one is generated by PM and the other one by  $O_S$ , before receiving its next input. Since for fixed M the parameters |Q| and k can be viewed as constants,

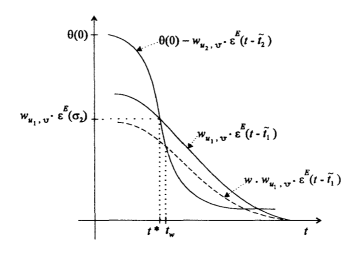


Figure 9: Mechanism of the weight-to-phase transformation module.

 $\mathcal{N}_M$  just uses O(1) spikes for the simulation of each computation step of M. Hence  $\mathcal{N}_M$  simulates M in real-time.

**2.9 Weight-to-Phase Transformation.** At this point the only missing link for the construction of the desired SNN  $\mathcal{N}_{TM}(d)$  is a module that allows us to generate from suitable weights of an SNN the encoding of arbitrarily long (even infinitely long) bit strings, which may, for example, represent the program of a Turing machine, or an infinitely long "look-up table." The weight-to-phase transformation module constructed here will be able to generate within a fixed number of "machine cycles" any given phase difference  $\varphi = \sum_{i=1}^t b_i \cdot 2^{-i-\epsilon}$  of an oscillator (for arbitrary  $\ell \in \mathbb{N} \cup \{\infty\}$  and  $b_i \in \{0.1\}$ ) from suitable weights between 0 and 1. Furthermore, these weights can be chosen to be rational if  $\ell \in \mathbb{N}$ . This module will exploit effects of the firing mechanism of a neuron in an SNN that are closely related to those that we had used in Section 2.6 to multiply the phase of an oscillator with a constant factor. To allow a unique decoding of infinitely long bit sequences from phase differences  $\varphi$  we adapt the convention that  $b_{2i} = 0$  for all  $i \in \mathbb{N}$  in case that  $\ell = \infty$ .

We consider the same configuration with neurons  $u_1, u_2, v$ , and an inhibition module as for MULTIPLY( $\beta$ ) in Section 2.6. However, instead of shifting the firing time of  $u_2$ , we are now interested in the consequences of multiplying the weight on the edge from  $u_1$  to v with some factor  $w \in [0,1]$  (see Fig. 9). We choose values for the delays  $\Delta_{u_i,v}$  so that for  $\tilde{t}_i := t_i + \Delta_{u_i,v}$  there exists some  $t^* \geq \max(\tilde{t}_1,\tilde{t}_2)$  with  $t^* - \tilde{t}_1 = \sigma_2$ 

and  $t^* - \tilde{t}_2 = \sigma_1$ . Furthermore, we choose positive weights  $w_{u_i,v}$  so that  $w_{u_2,v} \cdot s_{\text{up}} = 2w_{u_1,v} \cdot s_{\text{down}}$  and

$$w_{u_1,v} \cdot \varepsilon^E(t^* - \tilde{t}_1) + w_{u_2,v} \cdot \varepsilon^E(t^* - \tilde{t}_2) = \Theta(0).$$

To analyze the consequences of multiplying the weight  $w_{u_1,v}$  with some  $w \in [0,1]$ , we consider for arbitrary  $w \in [0,1]$  the point  $t_w > t^*$  that satisfies

$$w \cdot w_{u_1,v} \cdot \varepsilon^{E}(t_w - \tilde{t}_1) + w_{u_2,v} \cdot \varepsilon^{E}(t_w - \tilde{t}_2) = \Theta(0).$$

Together with the preceding equations and conditions (3) and (4) from our basic assumptions on  $\varepsilon^E$  this yields

$$\begin{split} (w-1) & \cdot & w_{u_1,v} \cdot \varepsilon^E(t^* - \tilde{t}_1) - w \cdot w_{u_1,v} \cdot s_{\operatorname{down}} \cdot (t_w - t^*) \\ & + 2w_{u_1,v} \cdot s_{\operatorname{down}} \cdot (t_w - t^*) = 0. \end{split}$$

or equivalently

$$t_w - t^* = \frac{(1 - w) \cdot w_{u_1, v} \cdot \varepsilon^E(\sigma_2)}{(2 - w) \cdot w_{u_1, v} \cdot s_{\text{down}}} = \frac{(1 - w) \cdot \varepsilon^E(\sigma_2)}{(2 - w) \cdot s_{\text{down}}}.$$

Then analogous arguments as in Section 2.6 show that if  $w \in (0,1]$  is chosen so that the right hand side of this equation has a value in [0,L/2], then the value for  $t_w$  that results from this equation is, in fact, the uniquely determined firing time of v in  $[t^*,t^*+L/2]$  if the weight on the edge  $\langle u_1,v\rangle$  is multiplied with w. In particular, the value  $t_w-t^*=L/2$  of the shift in the firing time of v is achieved for

$$w_L := \frac{\varepsilon^E(\sigma_2) - L \cdot s_{\text{down}}}{\varepsilon^E(\sigma_2) - L \cdot s_{\text{down}}/2}.$$

Thus  $w_L \in [0,1)$ , and the function  $w \mapsto t_w - t^*$  maps  $[w_L,1]$  one-one onto [0,L/2].

The inverse of this map is defined by

$$t_w - t^* \mapsto w := \frac{\varepsilon^{\mathcal{E}}(\sigma_2) - 2(t_w - t^*) \cdot s_{\text{down}}}{\varepsilon^{\mathcal{E}}(\sigma_2) - (t_w - t^*) \cdot s_{\text{down}}}.$$

One can derive from the basic assumptions on  $\Theta$  and  $\varepsilon^E$  that  $w_{u_1,v} \in \mathbb{Q}$ . Hence the preceding formula in combination with these basic assumptions implies that one can achieve *any rational* phase shift  $t_w - t^* \in [0, L/2]$  with a *rational* weight  $w \cdot w_{u_1,v}$  on the edge  $\langle u_1, v \rangle$ .

Finally, by our choice of c one has  $\sum_{i=1}^{\ell} b_i \cdot 2^{-i-c} \in [0, L/2]$  for any values of  $\ell \in \mathbb{N} \cup \{\infty\}$  and  $b_i \in \{0,1\}$ . Hence in a preprocessing phase of an SNN any given finite or infinite bit sequence  $\langle b_1, b_2, \ldots \rangle$  can be "loaded" [with only O(1) spikes involved] from the value of a certain weight of the SNN into the form of a phase difference  $\varphi_S = \sum_{i=1}^{\ell} b_i \cdot 2^{-i-c}$  of an oscillator  $O_S$ . For that purpose one has to ensure that the considered firings of neurons  $u_1$  and  $u_2$  (as well as of the involved inhibition module, see the corresponding construction in Section 2.6) are triggered by EPSPs from the pacemaker PM. Thus we have shown that the weights of an SNN can essentially play the role of a "read-only memory" of unlimited capacity.

**2.10 Construction of**  $\mathcal{N}_{TM}(d)$ . In this last part of the proof of Theorem 2.1 we construct an SNN  $\mathcal{N}_{TM}(d)$  that has those properties that are claimed in Theorem 2.1. Let  $d \in \mathbb{N}$  be any given constant. Let  $M_{\mathcal{U}}$  be a "universal Turing machine" with d + 1 tapes that can simulate any Turing machine with d tapes in real-time. More precisely,  $M_U$  is a Turing machine that receives two finite binary strings x and e on two different tapes as input, and which simulates for any  $e \in \{0,1\}^*$  the d-tape Turing machine whose program is encoded by e in real-time on input x (with some suitable default convention for the case that *e* is not the encoding of some Turing machine program). The construction of such universal Turing machines  $M_{\mathcal{U}}$  is a standard part of the proof of the time hierarchy theorem for Turing machines (see, e.g., Hopcroft and Ullman 1979, or Lewis and Papadimitriou 1981). The desired SNN  $\mathcal{N}_{TM}(d)$  will basically be the SNN that one gets by applying the construction from Section 2.8 to the Turing machine  $M := M_U$ , but with 2d + 2 stacks instead of the d+1 tapes.

The only additional work that remains to be done to satisfy the claim of Theorem 2.1 is to change the way in which  $\mathcal{N}_{M_{\ell\ell}}$  receives its input. Ordinarily  $\mathcal{N}_{M_{\ell\ell}}$  would expect to get its second input  $e = \langle e_1, \dots, e_{\ell} \rangle \in \{0,1\}^*$  in the same way as its first input  $x \in \{0,1\}^*$ , in the form of two input spikes with time distance  $\sum_{i=1}^{\ell} e_i \cdot 2^{-i-c}$ .

In contrast to that, the constructed SNN  $\mathcal{N}_{TM}(d)$  receives only a single input x in the form of a time difference between two input spikes. On the other hand its weights may depend on the simulated Turing machine M. Thus we may choose a rational weight  $w \in [0,1]$  that can be transformed with the help of the module from Section 2.9 into a phase difference  $t_w - t^* = \sum_{i=1}^{\ell} e_i \cdot 2^{-i-c}$ . This transformation can be carried out in a preprocessing phase within O(1) firings of PM. After that, the computation of  $\mathcal{N}_{TM}(d)$  proceeds exactly like that of  $\mathcal{N}_{M_U}$ .

To prove the second part of the claim of Theorem 2.1, one exploits the obvious fact that any function  $F: \{0,1\}^* \to \{0,1\}^*$  can be computed by a Turing machine  $M_F$  with infinitely many bits of "advice," i.e., by a Turing machine  $M_F$  that has at the beginning of each computation on one of its tapes the same infinite sequence  $\langle e_i \rangle_{i \in \mathbb{N}}$  of bits  $e_i \in \{0,1\}$  as initial tape inscription. This sequence  $\langle e_i \rangle_{i \in \mathbb{N}}$  may for example encode a look-up table for all pairs  $\langle x, F(x) \rangle$ ,  $x \in \{0,1\}^*$ . We may assume that  $\langle e_i \rangle_{i \in \mathbb{N}}$  also encodes the program of the Turing machine  $M_F$ , and that  $M_F$ altogether has only 2 tapes. As usual, the Turing machine  $M_F$  receives on another tape the input  $x \in \{0,1\}^*$ . To simulate this Turing machine  $M_F$ on the SNN  $\mathcal{N}_{TM}(d)$ , we just have to equip  $\mathcal{N}_{TM}(d)$  with a suitable real weight  $w \in [0,1]$  that can be transformed (as described in Section 2.9) in a preprocessing phase within O(1) firings of PM into the phase difference  $\hat{t}_w - t^* = \sum_{i=1}^{\infty} \hat{e}_i \cdot 2^{-2i-c}$  of an oscillator. After that,  $\mathcal{N}_{TM}(d)$  will simulate the computation of the Turing machine  $M_F$  (with initial tape content  $\langle e_i \rangle_{i \in \mathbb{N}}$  on one of its tapes) in the usual manner. Thus  $\mathcal{N}_{TM}(d)$  will output F(x) for any given input  $x \in \{0,1\}^*$ . Hence  $\mathcal{N}_{TM}(d)$  can compute the

(arbitrarily given) function  $F: \{0,1\}^* \to \{0,1\}^*$ . This concludes the proof of Theorem 2.1.

An important measure for the complexity of a neural network in the context of *learning* is its Vapnik–Chervonenkis dimension (VC-dimension), see Vapnik and Chervonenkis (1971). Various results from statistical theory suggest that the VC-dimension of a neural net is proportional to the number of examples needed to *train* that neural net (for references and a brief survey see, e.g., Maass 1995a).

**Corollary 2.2.** One can construct with any type of neurons whose response and threshold functions satisfy our basic assumptions an SNN  $\mathcal N$  of finite size, so that the VC-dimension of the class of Boolean functions that are computable on  $\mathcal N$  (with different assignments of rational values from [0,1] to its weights) is infinite.

A proof of the following result is contained as a special case in the proof of Theorem 2.1 (see especially Section 2.8).

**Corollary 2.3.** Any deterministic finite automaton with q states can be simulated in real-time (both for decision problems, or with intermediate output as a Mealy or Moore machine) by an SNN with  $O(q^{1/2})$  neurons [or with  $O(q^{1/2} \cdot \log q)$  neurons if only weights from [0,1] are permitted].

The following corollary exhibits another result of independent interest that was shown in the preceding proof (Section 2.5).

**Corollary 2.4.** One can construct with any type of neurons whose response and threshold functions satisfy our basic assumption for any given feedforward Boolean threshold circuit C with arbitrary weights, s gates and d hidden layers an SNN  $S_C$  with O(s) neurons that simulates any computation of C within a time interval of length O(d). Furthermore, one can also simulate C within time O(d) by an SNN  $S_C$  with polynomial(s) neurons that uses only weights  $w \in [0,1]$ .

Finally we observe that an application of the techniques from the proof of Theorem 2.1 to SNNs with discrete time (see the definitions in Section 1) yields the following result.

**Corollary 2.5.** One can construct for any Turing machine M with any type of neurons whose response and threshold functions satisfy our basic assumptions an SNN  $N_M$  so that for any  $s \in \mathbb{N}$  the SNN  $N_M$  with discrete firing times from  $\{i \cdot \mu : i \in \mathbb{N}\}$  for some  $\mu$  with  $1/\mu = 2^{s+O(1)}$  and  $\Delta_{max} - \Delta_{min} \geq 2\mu$  can simulate in real-time arbitrary computations of M that involve at most s tape cells of M.

For the proof of Corollary 2.5 one exploits the fact that because of the condition  $\Delta_{\max} - \Delta_{\min} \geq 2\mu$  the same construction as in Section 2.2 yields modules that achieve any given real-valued (!) delay  $\geq \Delta_{\min}$ . With the help of such delay modules one can then arrange that the time points  $t^*$  in the subsequent constructions of other modules, as well as the time points when the EPSPs reach their maximal value  $\varepsilon_{\max}$  (for the simulation of a

threshold circuit), all belong to the set  $\{i \cdot \mu : i \in \mathbb{N}\}$ . For the simulation of a stack of a Turing machine M the construction from the proof of Theorem 2.1 works without changes for SNNs with discrete time steps of length  $\mu$ , provided that  $2^{-i-\epsilon} \in \{i \cdot \mu : i \in \mathbb{N}\}$  for the maximal length  $\ell$  of any bit string that is stored in a stack of M.

### 3 Beyond Turing Machines

We have shown in Theorem 2.1 that one can build from arbitrary neurons, whose response and threshold functions satisfy certain basic assumptions, an SNN that can simulate any Turing machine. However SNNs are strictly more powerful than Turing machines for two reasons:

- 1. An SNN can receive *real* numbers as input, and give *real* numbers as output (in the form of time differences between pairs of spikes).
- 2. We had constructed in Section 2.6 modules for an SNN that can carry out the operations COMPARE ( $\geq \alpha$ ) and MULTIPLY( $\beta$ ), for a wide range of constants  $\alpha$  and  $\beta$ , applied to arbitrary real-valued arguments  $\varphi$  from a certain interval. If one applies, for example, such an operation to a phase of the form  $\varphi = \sum_{i=1}^t b_i \cdot 2^{-i-\varphi}$ , then such a module executes with O(1) spikes an operation that involves the whole bit string  $\langle b_1, \ldots \rangle$  of arbitrary length  $t \in \mathbb{N} \cup \{\infty\}$ . In contrast to that, any Turing machine operation can affect at best a constant number of its stored bits.

In this section we will show that in addition one can construct modules for an SNN that ADD, SUBTRACT, or COMPARE any two real valued phase differences  $\varphi_1, \varphi_2 \in [0, L/4]$  of two different oscillators. This result turns out to be quite important, since in combination with (1) and (2) it implies that one can simulate in real-time on an SNN any RAM with finitely many registers that stores in its registers arbitrary real numbers of bounded absolute value, and that uses arbitrary instructions of the form COMPARE, MULTIPLY( $\beta$ ), ADD, SUBTRACT. Furthermore, such an SNN can be built with any type of neurons whose response and threshold functions satisfy the basic assumptions from the beginning of Section 2.

On the other hand, according to Maass (1994b, 1995c), any SNN with *arbitrary* piecewise linear response and threshold functions can be simulated in real-time by the same type of RAM. Hence, the computational power of these RAMs (which we will call N-RAMs because of their close relationships to neural networks) matches exactly that of SNNs whose response and threshold functions are piecewise linear *and* satisfy our basic assumptions.

One can also show through mutual real-time simulations (see Maass 1995b,c) that the computational power of N-RAMs (and hence of the above-mentioned SNNs) matches exactly that of recurrent *analog* neural

nets with discrete time and piecewise linear activation functions (see Siegelmann and Sontag 1992). More precisely, any analog neural net with any piecewise linear activation functions can be simulated in real-time by an N-RAM; for the simulation of N-RAMs by analog neural nets one can employ, for example, the linear saturated activation function together with the heaviside activation function in the analog neural net. This result implies as a side-result that these two activation functions together are "universal" for all piecewise linear activation functions in recurrent analog neural nets (since they allow such a net to simulate in real-time any other recurrent analog neural net with arbitrary piecewise linear activation functions). Hence N-RAMs also provide a very useful intermediate link for the comparison of SNNs (modeling spike coding) and analog neural nets (modeling frequency coding).

We defer the detailed discussion of N-RAMs [which are somewhat weaker than the well-known model of Blum *et al.* (1989) and also related to the computational model considered in Koiran (1993)] and the proofs of the above-mentioned results to a subsequent article (Maass 1995c). However we will describe in this section the construction of SNN modules for the operations ADD, SUBTRACT, and COMPARE, since those constructions are closely related to the preceding constructions in this article. These constructions provide the tools for the real-time simulation of N-RAMs by SNNs.

Consider two oscillators  $O_1$  and  $O_2$  of an SNN, both with oscillationperiod  $\pi_{PM}$ . Let  $\varphi_i$  be the phase difference between  $O_i$  and the pacemaker PM, i = 1, 2. We construct a module that receives a spike from each of the oscillators  $O_1$  and  $O_2$ , and which is then able to kick-off a third oscillator O with oscillation period  $\pi_{PM}$  in such a way that it will have phasedifference  $\varphi_1 + \varphi_2$  to PM. This module for the operation ADD employs a similar arrangement of three neurons  $u_1, u_2$ , and v as the modules for COMPARE( $\geq \alpha$ ) and MULTIPLY( $\beta$ ) that were constructed in Section 2.6. We assume that neuron  $u_i$  is triggered by a spike from oscillator  $O_i$  to fire at a certain time  $t_i$ . We choose delays  $\Delta_{u_i,v}$  in such a way that for  $\tilde{t}_i := t_i + \Delta_{u_i,v}$  there exists some  $t^* \ge \max(\tilde{t}_1, \tilde{t}_2)$  so that  $t^* - \tilde{t}_1 = t^* - \tilde{t}_2 = \sigma_1$ in case that  $\varphi_1 = \varphi_2 = 0$ . We choose w > 0 so that  $2w \cdot \varepsilon^E(\sigma_1) = \Theta(0)$ , and we set  $w_{u_1,v} = w_{u_2,v} = w$ . We also add an inhibition module, which makes it impossible for v to fire within the time interval  $[t^* - \tau_{end}, t^* - L/2]$  for any values of  $\varphi_1, \varphi_2$ , and which has no influence on  $P_v(t)$  for  $t \ge t^*$  [as in the construction for MULTIPLY( $\beta$ ) in Section 2.6].

Then for arbitrary values  $\varphi_1, \varphi_2 \in [0, L/2]$  the neuron v fires at a time  $t_{\Sigma} \in [0, L/2]$  such that

$$w \cdot s_{\text{up}} \cdot [t_{\Sigma} - (t^* + \varphi_1)] + w \cdot s_{\text{up}} \cdot [t_{\Sigma} - (t^* + \varphi_2)] = 0,$$

or equivalently

$$t_{\Sigma}-t^*=\frac{\varphi_1+\varphi_2}{2}$$

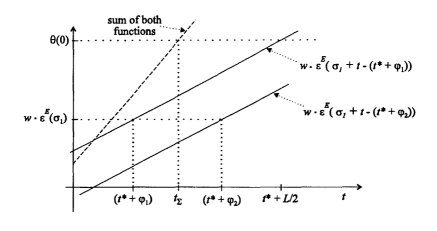


Figure 10: Mechanism of the module for ADD.

(see Fig. 10). The factor 1/2 of  $\varphi_1 + \varphi_2$  can be removed with the help of a subsequent module for MULTIPLY(2) (see Section 2.6). In this way the module for ADD constructed here can generate an output spike at time  $i \cdot \pi_{\text{PM}} + (\varphi_1 + \varphi_2)$  for some  $i \in \mathbb{N}$ .

The construction of a module that computes the *difference*  $\varphi_1 - \varphi_2$  of the phase differences  $\varphi_1, \varphi_2$  of two modules  $O_1$  and  $O_2$  with  $\varphi_1 \geq \varphi_2$  is quite similar. For arbitrary given values  $\varphi_1, \varphi_2 \in [0, L/2]$  with  $\varphi_1 \geq \varphi_2$  we first employ a module MULTIPLY(1/2) that replaces  $\varphi_1$  by  $\tilde{\varphi}_1 := \varphi_1/2$ . For an arrangement of neurons  $u_1, u_2, v$  as for ADD we choose delays  $\Delta_{u_1, v}$  so that  $t^* - \tilde{t}_1 = \sigma_1$  and  $t^* - \tilde{t}_2 = \sigma_2$  in case that  $\varphi_1 = \varphi_2 = 0$ , and weights  $w_{u_1, v}, w_{u_2, v}$  so that  $w_{u_1, v}, s_{up} = 2w_{u_2, v}, s_{down}$  and  $w_{u_1, v}, \varepsilon^E(\sigma_1) + w_{u_2, v}, \varepsilon^E(\sigma_2) = \Theta(0)$ . Furthermore, we employ an inhibition module that makes it impossible for neuron v to fire within the time-interval  $[t^* - \tau_{\text{end}}, t^* - L/2]$  for any values of  $\varphi_1, \varphi_2 \in [0, L/2]$ , but that has no influence on  $P_v(t)$  for  $t \geq t^*$ .

Then for any phase differences  $\varphi_1, \varphi_2 \in [0, L/2]$  with  $\varphi_1 \geq \varphi_2$  the phase difference  $\varphi_1$  is first transformed to  $\tilde{\varphi}_1 = \varphi_1/2$ . Neuron  $u_1$  receives a spike with phase difference  $\tilde{\varphi}_1$ , and  $u_2$  receives a spike with phase difference  $\varphi_2$ . The resulting firing time  $t_{\Delta}$  of neuron v is determined by (see Fig. 11)

$$w_{u_1,v} \cdot s_{up} \cdot [t_{\Delta} - (t^* + \tilde{\varphi}_1)] - w_{u_2,v} \cdot s_{down} \cdot [t_{\Delta} - (t^* + \varphi_2)] = 0.$$

This yields

$$t_{\Delta}-t^*=2\tilde{\varphi}_1-\varphi_2=\varphi_1-\varphi_2.$$

Finally, it is easy to see that the module for COMPARE( $\geq \alpha$ ) from Section 2.6 in combination with the preceding module for SUBTRACT

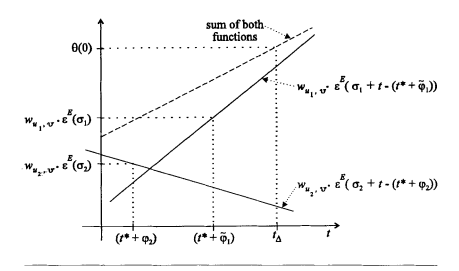


Figure 11: Mechanism of the module for SUBTRACT.

allows us to build a module for the test COMPARE, i.e., a module that decides for any two given phase differences  $\varphi_1, \varphi_2 \in [0.L/4]$  of two oscillators  $O_1$  and  $O_2$  with oscillation period  $\pi_{PM}$  whether  $\varphi_1 \geq \varphi_2$ . For that purpose one first transforms  $\varphi_1$  with the help of a delay module to  $\varphi_1' := \varphi_1 + L/4$ . It is then clear that  $\varphi_1' \geq \varphi_2$ , and the module for SUBTRACT can be employed to compute  $\varphi_1' - \varphi_2 = \varphi_1 - \varphi_2 + L/4$ . With the help of a subsequent module for COMPARE( $\geq L/4$ ) we can then decide whether  $\varphi_1 - \varphi_2 + L/4 \geq L/4$ , i.e., whether  $\varphi_1 \geq \varphi_2$ .

Of course one can also build directly a module for COMPARE by using a variation of the construction for COMPARE( $\geq \alpha$ ) in Section 2.6.

# 4 Variations of the Constructions for Related Models of Spiking Neurons

We have assumed for the constructions in the preceding two sections that the response and threshold functions are stereotyped, i.e., that apart from their individual delays  $\Delta_{u,v}$  the functions  $\varepsilon_{u,v}$  and  $\Theta_v$  all have the same shape. This assumption is convenient, but not really necessary for the preceding constructions. The same constructions can also be carried out if these functions are *different* for different edges  $\langle u,v\rangle \in E$  and different  $v \in V$ . More precisely, it suffices to assume that the response functions  $\varepsilon_{u,v}$  are defined with the help of individual delays  $\Delta_{u,v}$  and individual functions  $\varepsilon_{u,v}^E$  and  $\varepsilon_{u,v}^I$ , so that  $\varepsilon_{u,v}(x) = 0$  for  $x \in [0, \Delta_{u,v}]$  and  $\varepsilon_{u,v}(\Delta_{u,v} + x) = \varepsilon_{u,v}^E(x)$ ,

and, respectively,  $\varepsilon_{u,v}(\Delta_{u,v}+x)=\varepsilon_{u,v}^{l}(x)$  in the case of an IPSP, where the functions  $\varepsilon_{u,v}^{E},\varepsilon_{u,v}^{l}$ .  $\Theta_{v}$  satisfy the basic assumptions from the beginning of Section 2. However, these functions  $\varepsilon_{u,v}^{E},\varepsilon_{u,v}^{l}$ , and  $\Theta_{v}$  may be arbitrarily different, with different values of the parameters  $\tau_{\rm ref},\tau_{\rm end},\sigma_{j},\tau_{j},L,s_{\rm up},s_{\rm down}$ , for different neurons u,v (in fact one may assume that these functions are chosen by an "adversary"). Under these relaxed conditions we have to assume, however, that we can choose arbitrarily large delays  $\Delta_{u,v}$  and weights  $w_{u,v}$  after the individual functions  $\varepsilon_{u,v}^{E},\varepsilon_{u,v}^{l}$  and  $\Theta_{v}$  are given to us. Of course one can trade off parts of the latter condition against some quite reasonable conditions on the individual functions  $\varepsilon_{u,v}^{E},\varepsilon_{u,v}^{l}$  and  $\Theta_{v}$ .

One can also replace the basic assumptions at the beginning of Section 2 by some alternative assumptions about  $\varepsilon^E_{u,v}, \varepsilon^I_{u,v}$ , and  $\Theta_v$ . For example, one can postulate the existence of suitable linear segments of  $\varepsilon^I_{u,v}$  or  $\Theta_v$ , and then exploit at the neuron v in the module constructions of Sections 2 and 3 a "timing-race" between an EPSP and an IPSP, or between an EPSP and the declining part of  $\Theta_v$  (instead of the race between two EPSPs). Without a "reset" at each firing of neuron v (see below) one needs, however, for the latter option (EPSPs versus  $\Theta_v$ ) more specific assumptions about these functions to control undesired side-effects that may result from the end segments of EPSPs that caused the *preceding* firing of v.

We also would like to point out that the full power of the module COMPARE( $\geq \alpha$ ) from Section 2.6 is actually not needed if one just wants to simulate Turing machines on an SNN. If one employs a less concise encoding of bit strings by assuming also that  $b_{2i}=0$  for all  $i \leq \ell/2$  for all *finite* bit strings  $\langle b_1,\ldots,b_\ell\rangle$  that are encoded in the phase difference  $\varphi=\sum_{i=1}^\ell b_i \cdot 2^{-i-c}$  of an oscillator, it is guaranteed that  $\varphi\geq 2^{-1-c}$  or  $\varphi\leq 2^{-2-c}$  (independently of  $\ell$  and of the values of the  $b_i\in\{0,1\}$ ). This "gap" of fixed length between the possible values of  $\varphi$  allows us to determine whether  $b_1=1$  just with the help of delay and inhibition modules [instead of using the more subtle mechanism of COMPARE( $\geq \alpha$ )]. But the module for COMPARE( $\geq \alpha$ ) is of independent interest, since it shows in the context of Section 3 that *discontinuous* real-valued functions can also be computed on an SNN.

The implicit assumptions about the firing mechanism of neurons in the version of the SNN model from Section 1 ignore the well-known "reset" and "adaptation" phenomena of neurons. However, one can easily adjust the definition of the SNN model so that it also takes these features into account. To model a *reset* of a neuron at its moment of firing, one can adjust the definition of the set  $F_v$  of firing times of a neuron v by deleting (or modifying) in the definition of  $P_v(t)$  those EPSPs and IPSPs from presynaptic neurons u that had already arrived at v before the most recent firing of v.

Adaptation of a neuron v refers to the observation that the firing-rate of a biological neuron may decline after a while even if the incoming

excitation [i.e.,  $P_v(t)$ ] remains at a constant high level (see for example Kandel  $et\ al.$  1991). This effect can be reflected in the SNN model by replacing the term  $\Theta_v(t-s)$  in the definition of the set  $F_v$  of firing times by a sum over  $\Theta_v(t-s)$  for *several* recent firing times  $s\in F_v$  [and by assuming that  $\Theta_v(x)$  returns only relatively slowly to its initial value  $\Theta(0)$ ].

We would like to point out that all of our constructions in Sections 2 and 3 are compatible with our above-mentioned changes in the SNN model for modeling the *reset* and *adaptation* of neurons. The reason for this is that we can arrange in the constructions of Sections 2 and 3 that all "relevant" firings of a neuron v are spaced so far apart that reset and adaption of v have no effect on those *critical* firing times.

Regarding the simulation of *threshold* circuits by SNNs (see Section 2.5) we would like to point out that the corresponding SNN module can be constructed with *fewer* neurons if one makes further assumptions about the shape of EPSP and IPSP response functions. For example, one can simulate directly a threshold gate  $T^{\alpha}$  with weights  $\alpha_i$  of *different* sign in a similar way as we have simulated *monotone* threshold gates  $T^{\alpha}$  in Section 2.5, provided that the EPSPs (modeling inputs with positive weights) and IPSPs (modeling inputs with negative weights) move linearly within the same time span from 0 to their extremal values.

Finally, we would like to point out that the class of piecewise constant functions (i.e., the class of step-functions) provides an example for a class of response and threshold functions that do not satisfy our basic assumptions from Section 2, but that can still be used to build for any Turing machine M an SNN  $\mathcal{N}_{M'}$  that can simulate M (although not in real-time). We assume here that the response functions are piecewise constant (but not identically zero), and that the threshold functions are arbitrary functions (e.g., piecewise constant) that satisfy condition (1) of our basic assumptions. One can then build oscillators, as well as delay, inhibition, and synchronization modules, in the same way as in Section 2, and one can also simulate arbitrary threshold circuits in the same way. Furthermore one can use the phase difference between an oscillator O with the same oscillation period  $\pi_{PM}$  as the pacemaker PM to simulate a counter. For that purpose one employs a delay module D with a suitable delay  $\rho > 0$  (so that  $k \cdot \rho = \ell \cdot \pi_{PM}$  for any  $k, \ell \in \mathbb{N}$  implies that  $k = \ell = 0$ ). One can then use the phase difference between O and PM to record how often the "spike in O" has been directed in the course of the computation through this delay module D. Hence one can store in the SNN an arbitrary natural number k, which can be incremented and decremented by suitable modules. To decide whether k = 0, one needs a module that can carry out a special case of the operation COMPARE. Such a module cannot be built in the same way as in Sections 2 and 3, but one can employ directly the "jump" in the piecewise constant response functions considered here to test whether two neurons fire exactly at the same time.

It is well known (see Hopcroft and Ullman 1979) that any Turing machine M can be simulated (although not in real-time) by a machine M' that has no tapes or stacks, but two counters. The preceding argument shows that such an M' (in fact, a machine with any finite number of counters) can be simulated in real-time by some finite SNN  $\mathcal{N}_{M'}$  with piecewise constant response and threshold functions.

The effect of the shape of postsynaptic potentials on the computational power of networks of spiking neurons is investigated more thoroughly in Maass and Ruf (1995). It is shown there that computations with single spikes in networks of spiking neurons become substantially slower if they cannot make use of *increasing* and *decreasing* linear segments of EPSPs.

#### 5 Conclusion

We have analyzed the computational power of a simple formal model SNN for networks of spiking neurons. In particular we have shown that if the response and threshold functions of the SNN satisfy some rather weak basic assumptions (see Section 2), then one can build modules that can synchronize the spiking of different network parts, as well as modules that can multiply the phase difference between two oscillators with any given constant, and add, subtract, or compare the phase differences of different oscillators (see the constructions in Sections 2 and 3). With the help of these quite powerful computational operations an SNN can simulate in real-time for Boolean-valued input any Turing machine, and for real-valued input any N-RAM (a slightly weaker version of the model of Blum et al. 1989; see Section 3 of this article). On the side we would like to mention that these results also yield lower bounds for the VC-dimension of networks of spiking neurons, hence for the number of training examples needed for learning by such networks (see Maass 1994b, 1995c). One immediate consequence of this type is indicated in Corollary 2.2 of this article.

The version of the model SNN with unlimited timing-precision (i.e.,  $T = \mathbf{R}^+$  in the definition in Section 1) is not biologically realistic, insofar as it does not take the effects of noise into account. From that point of view our alternative version of this model with *discrete* firing times from  $\{i \cdot \mu : i \in \mathbf{N}\}$  for some  $\mu > 0$  is preferrable (since it allows us to represent an imprecise firing anywhere in the time interval  $(i \cdot \mu - \varepsilon.i \cdot \mu + \varepsilon)$  in the biological system by a "symbolic" firing at time  $i \cdot \mu$ ). Therefore it is important to note that our results about SNNs with unlimited timing precision *induce* corresponding results for the computational power of SNNs with *discrete* firing times, as we have indicated in Corollary 2.5 (see Theorem 5 in Maass 1994b, as well as Maass 1995c, for further consequences of our results for SNNs with limited timing precision). In addition our constructions of SNN modules for the operations ADD, SUBTRACT, and MULTIPLY( $\beta$ ) on time differences between spikes appear to be quite

robust, in the sense that they provide approximate implementations of these operations on time differences between spikes in various models for *real-valued* computations in networks of spiking neurons *with noise*. We refer to Maass (1995d) for further results about the computational power of SNNs with noise.

The results of this article have two interesting consequences. One is that in order to show that a network of spiking neurons can carry out some specific task (e.g., in pattern recognition or pattern segmentation, or solving some binding problem; see, e.g., von der Malsburg and Schneider 1986, or Gerstner et al. 1993) it now suffices to show that a threshold circuit, a finite automaton, a Turing machine, or an N-RAM (see Section 3) can carry out that task in an efficient manner. Furthermore the simulation results of this article allow us to relate the computational resources that are needed on the latter more convenient models (e.g., the required work space on a Turing machine) to the required resources needed by the SNN (e.g., the timing precision of the SNN, see Corollary 2.5). In other words, one may view N-RAMs and the other mentioned common computational models as "higher programming languages" for the construction of networks of spiking neurons. The real-time simulation methods of this article exhibit automatic methods for translating any program that is written in such higher programming language into the construction of a corresponding SNN. In this way the "user" of an SNN may choose to ignore all worrisome implementation details on SNNs such as timing (potentially at the cost of some efficiency). Furthermore the matching upper bound result for N-RAMs (see Maass 1995b,c) shows that the corresponding "higher programming language" is able to exploit all computational abilities of SNNs.

Second, in combination with the corresponding *upper* bound results for SNNs with quite arbitrary response and threshold functions (and time-dependent weights) in Maass (1995b,c), the lower bounds of this article provide for a large class of response and threshold functions exact characterizations (up to real-time simulations) of the computational power of SNNs with real valued inputs, and for SNNs with bounded timing precision. As a consequence of these results, one can then also relate the computational power of SNNs to that of recurrent analog neural nets with various activation functions (see Section 3), thereby throwing some light on the relationships between the computational power of models of neurons with spike coding (SNNs) and models of neurons with frequency coding (analog neural nets). Furthermore, the combination of these lower and upper bound results shows that extremely simple response and threshold functions (such as, for example, those in Fig. 2 in Section 2) are universal in the sense that with these functions an SNN can simulate in real-time any SNN that employs arbitrary piecewise linear response and threshold functions. Equivalence results of this type induce some structure in the "zoo" of response and threshold functions that are mathematically interesting or occur in biological neural systems,

and they allow us to focus on those aspects of these functions that are *essential* for the computational power of spiking neurons.

Finally we would like to point out that since we have based all of our investigations on the rather fine notion of a *real-time simulation* (see Section 1), our results provide information not just about the relationships between the *computational* power of the previously mentioned models for neural networks, but also about their capability to execute *learning* algorithms (i.e., about their *adaptive* qualities).

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