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# Accelerator for Event-based Failure Prediction

Acceleration of an Extended Forward Algorithm for Failure Prediction on  
FPGA

Master's Thesis submitted to the  
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Embedded Systems Design

presented by  
Simon Maurer

under the supervision of  
Prof. Mirosław Malek

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I certify that except where due acknowledgement has been given, the work presented in this thesis is that of the author alone; the work has not been submitted previously, in whole or in part, to qualify for any other academic award; and the content of the thesis is the result of work which has been carried out since the official commencement date of the approved research program.

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Simon Maurer  
Lugano, 29. January 2014



# Abstract



# Acknowledgements





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# Chapter 1

## Introduction

In today's live it becomes increasingly important, that computer systems are dependable. The reason being, that computer systems are used more and more in areas where the failure of such a system can lead to catastrophic events. Banking, public transportation and medical engineering are only a few examples of areas employing large and extremely complex systems. The increasing complexity of computer systems has a direct impact on their maintainability and testability. It is simply impossible to guarantee that a piece of software comes without any faults. On top of that, the same problematic arises with the hardware components which also may contain faulty parts but also get increasingly prone to failures due to decay of material.

In the event of a system failure it is of course desirable to fix the system as soon as possible in order to minimize the downtime of the system (maximize the availability). This can be accomplished by using different types of recovery techniques, e.g. Check-pointing (create checkpoints to roll back/forward), system replication (switch to a redundant system), fail over (reboot). All these techniques require a certain amount of time to complete the recovery process, time that is very expensive. In order to minimize this time, techniques have been developed to anticipate upcoming failures. Such a technique is described in [23].

The work presents a new algorithm to predict failures and compares the results with other techniques. The accuracy of the presented algorithm to predict failures proves to be better compared to the other techniques, has however the drawback of increased complexity and hence increased computation time. It is very important to keep the computation overhead very low in order to maximize the time between the prediction of a failure and the actual event of the failure. One way to decrease the computation time is to design a hardware accelerator for the prediction algorithm. The design of such an accelerator is outlined in this document.

### 1.1 Problem Statement

### 1.2 Motivation

The email of Felix left some doubts to whether the acceleration of the algorithm is useful. The following list will give some arguments to justify the work.

#### **Too many parameters to be identified, estimated and set**

Considering an embedded system, this is usually not a problem because the parameters

are defined during the design phase and will never be changed afterwards.

**Limited performance scalability**

There are studies available claiming otherwise. The discussion of Neumanns work will provide some arguments against this statement.

**Industry trends point towards cloud**

In embedded systems it will still be beneficial to predict failures of single nodes. It is however important to keep the power and computational footprint low. This will be one of the major challenges. On the other hand, I think it would also be possible to also use this algorithm to monitor a distributed system and predict failures. It is only a matter of getting defining the events to feed to the algorithm.

## 1.3 Contributions

## 1.4 Document Structure

## Chapter 2

# State of the Art

This section provides an overview of the state of the art in the different fields of research that are relevant for the thesis. This includes failure prediction methods, existing solutions to accelerate failure prediction algorithms and acceleration techniques in general.

### 2.1 Failure Prediction

A very detailed overview of failure prediction methods is given in [24]. The survey discusses i.a. the techniques used as comparison in the main reference [16, 15, 27, 8] as well as the technique described in the main reference [23].

More recent work uses hardware counters of a general purpose CPU and combines them with software instrumentation to analyze failures of single processes (e.g grep, flex, sed) [31]. As industry heads more and more towards cloud computing, it has been proposed to use information of interaction between nodes (instead of analyzing single nodes) in order to analyze and predict failures of a distributed system [25, 20].

### 2.2 Accelerator

The main goal of this master thesis is to accelerate an adaptation of the forward algorithm. Proposals for a GPU based accelerator for the classic forward algorithm are described in [19, 17]. Further, several proposals to accelerate the Viterbi algorithm (which is closely related to the forward algorithm) have been published: [2] presents an architecture for a lightweight Viterbi accelerator designed for an embedded processor datapath, [10, 18, 21] describe a FPGA based accelerator for protein sequence HHM search and [28] describes i.a. an approach to accelerate the Viterbi algorithm from the HMMER library using GPUs.

Focusing on a more general approach for acceleration, [12] proposes an FPGA implementation of a parallel floating point accumulation and [30] describes the implementation of a vector processor on FPGA.

Quite some research has been done on the question what type of technology should be used to accelerate certain algorithms: [4] presents a performance study of different applications accelerated on a multicore CPU, on a GPU and on a FPGA, [11] discusses the suitability of FPGA and GPU acceleration for high productivity computing systems (HPCS) without focusing on a

specific application and [14] also focuses on HPCS but uses the Basic Linear Algebra Subroutines (BLAS) as comparison and also takes CPUs into account.

It may be interesting to also think about an acceleration of the model training. Similar work has been done by accelerating SVMs (Support Vector Machines): [3] describes a FPGA based accelerator for the SVM-SMO (support vector machine - sequential minimal optimization) algorithm used in the domain of machine learning and [1] proposes a new algorithm and its implementation on a FPGA for SVMs.

## Chapter 3

# Event-based Failure Prediction

This section provides a brief overview of the computational steps done by the proposed algorithm [23].

*brief description of the idea behind the algorithm, HSMM, Events, etc*

To be able to understand the formal expression of the algorithm, first a definition of the used parameters is provided.

- N: number of states
- M: number of observation symbols
- L: observation sequence length
- R: number of cumulative probability distributions (kernels)

The delay of the event at time  $t_k$  with respect to the event at time  $t_{k-1}$  is described as

$$d_k = t_k - t_{k-1} \quad (3.1)$$

### 3.1 Data Processing

### 3.2 Model Training

One part of the algorithm is the model training. This part is not described here. The features to be trained by the model training are however important in this context because they are used by the adapted forward algorithm. Following the features:

- $\pi_i$ , forming the initial state probability vector  $\pi$  of size  $N$
- $b_i(o_j)$ , forming the emission probability matrix  $B$  of size  $N \times M$
- $p_{ij}$ , forming the matrix of limiting transmission probabilities  $P$  of size  $N \times N$
- $\omega_{ij,r}$ , the weights of the kernel  $r$
- $\theta_{ij,r}$ , the parameters of the kernel  $r$

### 3.3 Sequence Processing

The following description will provide a complete blueprint of the extended forward algorithm, that allows to implement it, but without any explanations or proofs related to the formulation. The adapted forward algorithm is defined as follows:

$$\alpha_0(i) = \pi_i b_{s_i}(O_0) \quad (3.2)$$

$$\alpha_k(j) = \sum_{i=1}^N \alpha_{k-1}(i) v_{ij}(d_k) b_{s_j}(O_k); \quad 1 \leq k \leq L \quad (3.3)$$

where

$$v_{ij}(d_k) = \begin{cases} p_{ij} d_{ij}(d_k) & \text{if } j \neq i \\ 1 - \sum_{\substack{h=1 \\ h \neq i}}^N p_{ih} d_{ih}(d_k) & \text{if } j = i \end{cases} \quad (3.4)$$

with

$$d_{ij}(d_k) = \sum_{r=1}^R \omega_{ij,r} \kappa_{ij,r}(d_k | \theta_{ij,r}) \quad (3.5)$$

forming the matrix of cumulative transition duration distribution functions  $D(d_k)$  of size  $N \times N \times L$ .

For simplification reasons, only one kernel is used. Due to this, the kernel weights can be ignored. Equation 3.5 can then be simplified:

$$d_{ij}(d_k) = \kappa_{ij}(d_k | \theta_{ij}) \quad (3.6)$$

Choosing the Gaussian cumulative distribution results in the kernel parameters  $\mu_{ij}$  and  $\sigma_{ij}$ :

$$\kappa_{ij,gauss}(d_k | \mu_{ij}, \sigma_{ij}) = \frac{1}{2} \left[ 1 + \operatorname{erf} \left( \frac{d_k - \mu_{ij}}{\sqrt{2} \sigma_{ij}} \right) \right] \quad (3.7)$$

*explain difference to the non-extended forward algorithm and introduce a standard notation for the transition probabilities (eg v: extended a: basic, tp: general) stick to this in the whole document*

The last set of forward variables  $\alpha_L$  are then summed up to compute a probabilistic measure for the similarity of the observed sequence compared to the sequences in the training data set. This is called the sequence likelihood:

$$P(\mathbf{o} | \lambda) = \sum_{i=1}^N \alpha_L(i) \quad (3.8)$$

where  $\lambda = \{\pi, P, B, D(d_k)\}$ .

To prevent  $\alpha$  from going to zero very fast, at each step of the forward algorithm a scaling is performed:

$$\alpha_k(i) = c_k \alpha_k(i) \quad (3.9)$$

with

$$c_k = \frac{1}{\sum_{i=1}^N \alpha_k(i)} \quad (3.10)$$

By applying scaling, instead of the sequence likelihood (equation 3.8), the sequence log-likelihood must be computed:

$$\log(P(\mathbf{o}|\lambda)) = -\sum_{k=1}^L \log c_k \quad (3.11)$$

where  $\lambda = \{\pi, P, B, D(d_k)\}$ .

### 3.4 Classification

*explain classification*

and finally the classification is performed:

$$\text{class}(s) = F \iff \max_{i=1}^u [\log P(s|\lambda_i)] - \log P(s|\lambda_0) > \log \theta \quad (3.12)$$

with

$$\theta = \frac{(r_{\bar{F}F} - r_{\bar{F}\bar{F}})P(c_{\bar{F}})}{(r_{F\bar{F}} - r_{FF})P(c_F)} \quad (3.13)$$

*classification without scaling*

*Multi-class classification without scaling?*

To calculate  $\theta$ , the following parameters need to be set:

- $P(c_{\bar{F}})$ : prior of non-failure class
- $P(c_F)$ : prior of failure class
- $r_{\bar{F}\bar{F}}$ : true negative prediction
- $r_{FF}$ : true positive prediction
- $r_{\bar{F}F}$ : false positive prediction
- $r_{F\bar{F}}$ : false negative prediction





## Chapter 4

# Theoretical Analysis of the Forward Algorithm

This chapter provides details about the forward algorithm and available (and useful) parallelization techniques applicable on the algorithm. The generally known forward algorithm as well as the extended version proposed in the reference work is discussed. Further, scaling techniques of the forward variables and their impact on data representation choices are presented. Finally the observations presented in an overview and an appropriate choice on possible acceleration hardware is made.

### 4.1 Serial Implementation and Complexity

The sequential implementation of the basic forward algorithm is represented in listing 4.1. It consist of three parts: the initialization step, the computation of consecutive forward variables and the final step, where the likelihood is computed. The initial  $\alpha$  variable is computed by multiplying the initial state probability with the emission probability of the first observation symbol of the sequence (cf equation 3.2). The computation of the following forward variables consists of three nested loops: the outer loop iterates over the  $L$  sets of  $N$   $\alpha$  variables, where each variable depends on the prior computed  $\alpha$  variable and the  $k$ -th observation symbol of a sequence. The first inner loop iterates over the  $N$   $\alpha$  variables of one set, where each variable is computed with the most inner loop. The two nested inner loops form the Matrix-Vector-Vector multiplication

$$\alpha_{k+1} = TP * \alpha_k \cdot B(o_k) \quad (4.1)$$

where  $\alpha_k$  is a vector of size  $N$  of the prior computed  $\alpha$  variables,  $TP$  a matrix of size  $N \times N$  containing the transition probabilities and  $B(o_k)$  a vector of size  $N$  containing the emission probabilities of the  $k$ -th observation symbol. Note that the first multiplication is a Matrix-Vector multiplication that results in a vector, which is then multiplied element-wise with the vector  $B(o_k)$ . The equation 3.3 describes the formal definition of the forward algorithm. In the last step the likelihood is computed, by summing up all elements of the last forward variable  $\alpha_L$  (cf equation 3.8).

As proposed by the reference work, the forward variables can be scaled, in order to prevent the result from getting very small due to the continuous multiplication of probabilities. The

```

1  %%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
2  % computation of the forward algorithm without scaling
3  %
4  % @param N:          number of states
5  % @param L:          number of observation symbols
6  % @param PI:         initial state probability vector. size N
7  % @param B:          matrix of emission probabilities. size N, L
8  % @param TP:         transistion probabilities. size N, N
9  % @param oL:         indices of all observed symbols. size 1, L
10 % @return Ps:        probability likelihood
11 %%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
12 function [Ps] = forward_s_basic(N, L, PI, B, TP, oL)
13     % initialize forward variables
14     for i=1:N,
15         alpha(i) = PI(i)*B(i, oL(1));
16     end
17     % forward algorithm
18     for k=2:L,
19         for j=1:N,
20             alpha_new(j) = 0;
21             for i=1:N,
22                 alpha_new(j) += alpha(i) * TP(i, j);
23             end
24             alpha_new(j) *= B(j, oL(k));
25         end
26         alpha = alpha_new;
27     end
28     % compute likelihood
29     Ps = 0;
30     for i=1:N,
31         Ps += alpha_new(i);
32     end
33 end

```

Listing 4.1. Forward Algorithm

implementation of the proposed scaling method is shown in listing 4.2. The scaling is formally defined by the equations 3.9 and 3.10. Due to the scaling, instead of the likelihood, the log-likelihood is computed. Equation 3.11 gives the formal definition.

The algorithm to compute the sequence likelihood proposed by the reference work is an extension to the forward algorithm presented in the listings 4.1 and 4.2. Instead of constant transition probabilities, the extended algorithm computes a new transition probability matrix (size  $N \times N$ ) for each arriving observation symbol, by considering the delay of the new symbol with respect to the prior symbol. The computation of the transition probability matrix  $TP$  is implemented with listing 4.3 and formally defined by the equations 4.5, 3.5, 3.6 and 3.7. As described in chapter 3.3, also here for reasons of simplification, only one kernel is used. In the sample code the Gaussian cumulative distribution function is used. The function needs to be called for every  $k$ .

The order of time complexity of the algorithm is  $O(LN^2)$ . While the complexity increases with the introduction of scaling and/or the extension, the order of complexity stays the same. The same is true for the space complexity which is of the order  $O(N^2)$ .

## 4.2 Available Parallelism

By applying parallelization methods one aims to increase the throughput or reduce the latency of a task, or to achieve both at the same time. This can be done at the cost of increased usage of parallel computation units, memory and memory bandwidth.

Considering only the basic forward algorithm (listing 4.1), the computation of the likelihood is divided in  $L + 1$  steps: the initialization,  $L - 1$  identical intermediate steps and the finalization. Because of the recursive nature of the algorithm, all steps (except the initialization) depend on the previously computed forward variables. For this reason a direct parallelization of the steps is not possible. However, at every arrival of a new observation symbol, the last  $L$  elements of the observation symbol sequence are used to compute the likelihood (cf. figure 4.1). This can be exploited to pipeline the steps in order to increase the throughput. By building a pipeline of  $L + 1$  stages, where each step of the forward algorithm corresponds to a pipeline stage, a likelihood is computed at every completion of a step, with a latency of  $(L + 1) * t_{step_{max}}$ , where  $t_{step_{max}}$  is the time needed to complete the computation of the most complex step (each stage of the pipeline must take the same amount of clock cycles). The throughput of a pipelined compared to a non-pipelined system is increased by factor  $L$  (assuming an infinite runtime or by ignoring the setup time). Another and more important fact, that makes the pipeline architecture very beneficial in this particular case: the configuration allows to load the transition probabilities  $TP$  and the emission probabilities  $b_i(o_k)$  for all steps at the same time, which reduces the load operations by factor  $L$ . This is visualized by the table 4.1. The table shows the pipeline stages with input values that are fed to the stage before the execution (note, that the input values  $TP$  and  $B$  always depend on the same observation symbol. The parameter  $d_k$  of the transition probabilities can be ignored in this case, because only in the extended forward algorithm they depend on  $d_k$ . This will be discussed further when the extension is considered) and the output values resulting after the execution of the pipeline stage. Figure 4.2 shows a schematic representation of the pipeline.

By considering all dissimilar steps of the forward algorithm, more parallelization options can be found. In the initial step,  $N$  components of the first forward variable  $\alpha_1$  are computed by multiplying independent pairs of an initial state probability  $\pi_i$  and an emission probability of the

```

1  %%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
2  % computation of the forward algorithm with scaling
3  %
4  % @param N:          number of states
5  % @param L:          number of observation symbols
6  % @param PI:         initial state probability vector. size N
7  % @param B:          matrix of emission probabilities. size N, L
8  % @param TP:         transistion probabilities. size N, N
9  % @param oL:         indices of all observed symbols. size 1, L
10 % @return Ps:        probability likelihood
11 %%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
12 function [lPs] = forward_s_scaling(N, L, PI, B, TP, oL)
13     % initialize forward variables
14     for i=1:N,
15         alpha(i) = PI(i)*B(i, oL(1));
16     end
17     % scaling
18     alpha_sum = 0;
19     for i=1:N,
20         alpha_sum += alpha(i);
21     end
22     scale_coeff(1) = 1 / alpha_sum;
23     for i=1:N,
24         alpha(i) *= scale_coeff(1);
25     end
26     % forward algorithm
27     for k=2:L,
28         for j=1:N,
29             alpha_new(j) = 0;
30             for i=1:N,
31                 alpha_new(j) += alpha(i) * TP(i, j);
32             end
33             alpha_new(j) *= B(j, oL(k));
34         end
35         % scaling
36         alpha_sum = 0;
37         for i=1:N,
38             alpha_sum += alpha_new(i);
39         end
40         scale_coeff(k) = 1 / alpha_sum;
41         for i=1:N,
42             alpha_new(i) *= scale_coeff(k);
43         end
44         alpha = alpha_new;
45     end
46     % compute log likelihood
47     lPs = 0;
48     for i=1:L,
49         lPs -= log(scale_coeff(i));
50     end
51 end

```

Listing 4.2. Forward Algorithm with scaling

```

1  %%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
2  % computation of the extended transition probabilities
3  %
4  % @param N:          number of states
5  % @param dk:         delay of k-th observation symbol
6  % @param cdf_param:  parameters for the cdf
7  % @return v:         extended transition probabilities
8  %%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
9  function [v] = compute_tp(N, dk, cdf_param)
10     % compute all elements of v
11     for i=1:N,
12         for j=1:N,
13             v(i, j) = normcdf(dk, cdf_param.mu(i, j), cdf_param.sigma(i, j));
14         end
15     end
16     % correct diagonal elements of v
17     for i=1:N,
18         for j=1:N,
19             v_sum(i) += v(i, j);
20         end
21     end
22     for i=1:N,
23         v_sum(i) -= v(i, i);
24         v(i, i) = 1 - v_sum(i);
25     end
26 end

```

Listing 4.3. Extension of the Forward Algorithm with only one kernel (Gaussian)

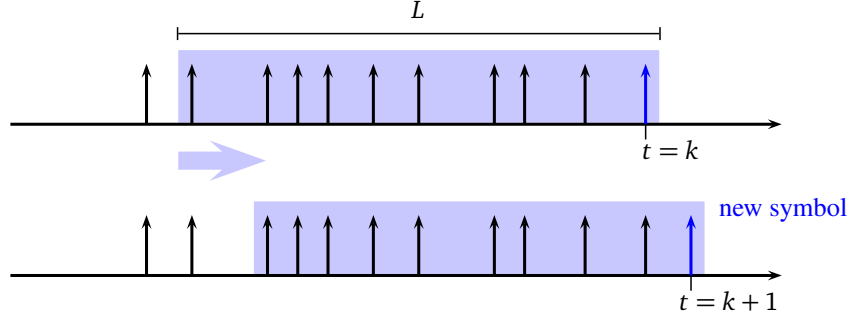
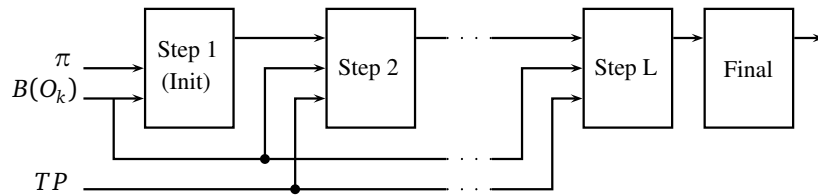
Figure 4.1. Sliding window over an observation sequence of the last  $L = 10$  observation symbols

Figure 4.2. Pipelined Forward Algorithm

Symb	I/O	Pipeline				
		Init	Step 2	...	Step L	Final
$O_1$	in	$B(O_1)$	$B(O_1), TP(d_1), 0$	...	$B(O_1), TP(d_1), 0$	0
	out	$\alpha_1(O_1)$	0	...	0	0
$O_2$	in	$B(O_2)$	$B(O_2), TP(d_2), \alpha_1(O_1)$	...	$B(O_2), TP(d_2), 0$	0
	out	$\alpha_1(O_2)$	$\alpha_2(O_{1,2})$	...	0	0
$\vdots$		$\vdots$	$\vdots$		$\vdots$	$\vdots$
$O_L$	in	$B(O_L)$	$B(O_L), TP(d_L), \alpha_1(O_{L-1})$	...	$B(O_L), TP(d_L), \alpha_{L-1}(O_{1,...,L-1})$	0
	out	$\alpha_1(O_L)$	$\alpha_2(O_{L-1,L})$	...	$\alpha_L(O_{1,...,L})$	0
$O_{L+1}$	in	$B(O_{L+1})$	$B(O_{L+1}), TP(d_{L+1}), \alpha_1(O_L)$	...	$B(O_{L+1}), TP(d_{L+1}), \alpha_{L-1}(O_{2,...,L})$	$\alpha_L(O_{1,...,L})$
	out	$\alpha_1(O_{L+1})$	$\alpha_2(O_{L,L+1})$	...	$\alpha_L(O_{2,...,L+1})$	$Ps(O_{1,...,L})$
$\vdots$		$\vdots$	$\vdots$		$\vdots$	$\vdots$

Table 4.1. Pipelined Forward Algorithm, with observation symbol  $O_k$  and its delay  $d_k$ . Here  $O_{i,...,k}$  is a short notation for  $O_i, \dots, O_k$

first observation symbol  $b_i(o_0)$ . This can be fully parallelized by replicating the multiplication operation  $N$  times. Doing this results in a increase of the throughput by factor  $N$  and a decrease of the latency by factor  $\frac{1}{N}$ , assuming that  $N$  multipliers are available and the memory bandwidth is able to provide a data throughput  $N$  times higher than in the sequential case (including the memory interface).

The computation of the following forward variables  $\alpha_k$ , with  $k = 2 \dots L$  are similar. To compute the  $N$  elements of one step, the Matrix-Vector-Vector multiplication described by equation 4.1 must be performed. Considering first only the Matrix-Vector multiplication, this can be parallelized by decomposing the matrix in to subsets and then use multiple computational units to perform multiplication and/or accumulation operations in parallel on the subsets. An intuitive decomposition can be done either by block-striped matrix partitioning (decomposition into subsets of rows or columns) or by checkerboard block matrix partitioning (decomposition in rectangular sets of elements). These partitioning methods are shown in figure 4.3. The number of subsets must correspond to the number of available computational units to perform the necessary operations. The choice of decomposition is heavily dependant on the accelerator architecture (e.g. communication between computational units, memory architecture). The resulting vector can then be multiplied element wise by the emission probability vector, which is again the same case as described above. In case of the block-striped matrix partitioning, the maximal achievable increase of the throughput is a factor of  $N$  and the latency can be decreased by factor  $\frac{1}{N}$ , assuming that  $N$  computational units are available to perform the multiplication and accumulation operation on each subset,  $N$  multipliers to compute the final element wise vector-vector multiplication and a memory interface, that can handle a data throughput that is  $N$  times higher than in the sequential case. The checkerboard partitioning yields a lower gain but may be considered in case of fixed computational unit architecture (CPU, GPU) in order to increase the utility of recourses available in each unit. Apart from homogeneous partitioning methods as mentioned above also inhomogeneous solutions have been proposed [6, 5]. These are not considered in this work, as the focus lies on homogeneous computation units.

Further parallelization can be done by using a reduction tree to accumulate the elements in the matrix-vector multiplication process. Instead of using one computation unit and accumulate the values sequentially,  $N$  units can be used to first multiply two operands together, then

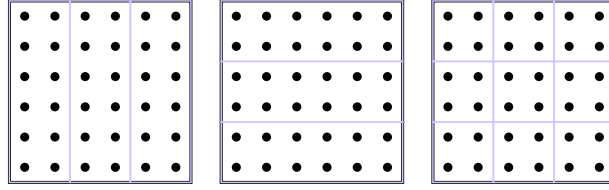


Figure 4.3. Matrix partitioning (from left to right): column-block-striped, row-block-striped and checkerboard blocks

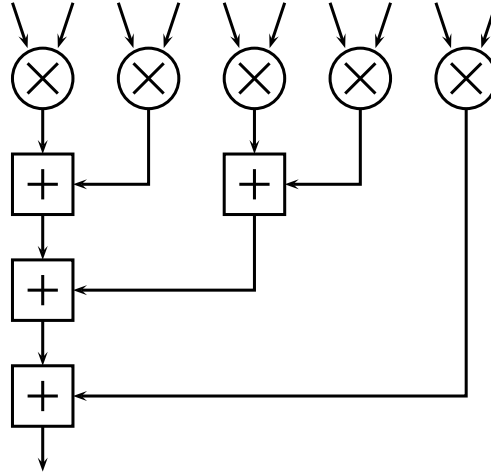


Figure 4.4. Reduction Tree with  $N = 5$

adding  $\frac{N}{2}$  resulting operand pairs in a second step and then consecutively adding resulting pairs until only one value results. This process is visualized in figure 4.4. The maximal increase of throughput is of factor  $\log_2(N)$  and the latency can be decreased by factor  $\frac{1}{\log_2(N)}$ , assuming that  $N$  computation units are available and the memory interface is able to handle a throughput that is  $N$  times higher than in the sequential case.

The finalization step of the algorithm consists of calculating the likelihood. This is done by simply accumulating the  $N$  elements of the last forward variable  $\alpha_L$ . This operation can be parallelized with a reduction tree, resulting in a throughput and latency optimization as described above.

The following sections will describe the impact on performance if scaling or the extension of the forward algorithm is implemented. Also the availability of parallelization in both cases will be discussed. Everything will be concluded with an overview of available parallelism and a discussion about the usefulness of each parallelization method in the context of the different algorithm implementations.

### 4.3 Simple Models

Until now, it was always assumed, that the model is fully connected (ergodic), i.e. that every state can be reached from every other state. This is not necessarily the case, as it is often possi-

ble to describe a system with a simpler model. By adding more constraints to the possible state transitions (e.g. only one direction, feed-forward), only a few elements in the transition probability matrix are non-zero. In this case it is beneficial to use an array (adjacency list) instead of a matrix to represent the transition probabilities. Different methods have been proposed on how to store sparse matrices, but they are usually stonily dependant of the architecture and will hence be discussed only after the type acceleration device has been chosen. A list with only non-zero elements instead of a sparse matrix reduces the necessary memory to store the data and makes a lot of computations superfluous. A Matrix-Vector multiplication parallelization as described in the previous section would not be beneficial anymore as a lot of computational units would be idle in most of the time.

## 4.4 Scaling and Data Representation

Scaling may be applied to prevent that the continuous multiplication of numbers smaller than one (e.g. probabilities) result in zero, because of the limited accuracy by digitally representing fractional numbers. Scaling does not influence the order of complexity of the algorithm. By introducing a scaling method as proposed in the reference work, the complexity of calculating one  $\alpha_k$  vector goes from  $N^2$  (no scaling) to  $N^2 + 2N + 1$  (scaling), which is the same order  $O(N^2)$ . However, the introduction of scaling may increase the usage of recourses significantly: In order to scale  $\alpha_k$ , the division operation is used to compute the scaling factor. Division is far more complex than multiplication and hence uses more recourses. Additionally, instead of the sequence likelihood (equation 3.8) the sequence log-likelihood (equation 3.11) needs to be computed, with the even more complex log operation.

In order to limit the amount of necessary division operations, it is beneficial to consider the following: Rather than scaling each element of  $\alpha_k$  by dividing it by a scaling factor ( $N$  divisions), first the inverse of the scaling factor can be computed, which is then multiplied with each element of  $\alpha_k$  (one division and  $N$  multiplications). Using  $N$  multiplication units, this operation can be parallelized.

To compute the log-likelihood,  $N$  log and  $N$  sum operations are necessary, in comparison to  $N$  sum operations for the likelihood. In terms of memory, the log-likelihood is more complex because the scaling coefficients of each  $\alpha_k$  are used and need to be stored, while for the likelihood only the last set of forward variables  $\alpha_L$  are used. The computation of the log-likelihood can be parallelized by using  $N$  units computing the log function and additionally by a reduction tree to speed up the accumulation.

Instead of using the proposed scaling method, a simpler scaling may be applied. By analyzing the operands, an average scaling factor can be computed. Using the knowledge, that all the operands are probabilities,

$$\sum_{i=1}^N \pi_i = 1, \sum_{j=1}^N t p_{ij} = 1, \sum_{j=1}^M b_{ij} = 1 \quad (4.2)$$

and doing the computation of the forward variables,



$$\begin{aligned}
\hat{\alpha}_1 &= \hat{b} \cdot \hat{\pi} = \frac{1}{NM} \\
\hat{\alpha}_2 &= N \cdot \hat{\alpha}_1 \cdot \hat{t} \hat{p} \cdot \hat{b} = N \cdot \frac{1}{NM} \cdot \frac{1}{N} \cdot \frac{1}{M} = \frac{1}{NM^2} \\
\hat{\alpha}_3 &= N \cdot \hat{\alpha}_2 \cdot \hat{t} \hat{p} \cdot \hat{b} = N \cdot \frac{1}{NM^2} \cdot \frac{1}{N} \cdot \frac{1}{M} = \frac{1}{NM^3} \\
&\vdots \\
\hat{\alpha}_L &= \frac{1}{NM^L}
\end{aligned} \tag{4.3}$$

it can be computed, that assuming no precision loss at each computational step  $k$ , on average a scaling factor of  $\frac{1}{M}$  is necessary in each step  $k$ . If the intermediate precision of the computational units is high enough to compensate for scaling to much or to few, this method is an easy solution to keep the values in an acceptable range. However, if the precision is not available (eg. if a fixed point data representation is chosen) a fixed scaling factor can cause an overflow (very bad because the result will be wrong) or an underflow (may be acceptable because it is only a loss of precision). In this case, rather than choosing an average scaling factor of  $\frac{1}{M}$  it is safer to choose the scaling factor to be equal to the maximal possible scaling factor of all values of a specific event in  $B$  (scale  $\max(B(O_k))$ ). By doing this, the scaling factor will be too small and if  $L$  is big, the forward variables will still approach zero, only slower than without scaling. This is either acceptable because of a high precision, or another scaling factor must be computed to prevent this. The implemented solution will be explained in detail in chapter 5.

Another aspect to consider is the choice of data representation (floating point versus fixed point). This depends on one hand on the necessary precision and on the other hand on the choice of accelerator type. While general purpose hardware such as CPU, GPU and DSP (to some degree) offer an abstraction to make the representation type transparent to the developer, specialized hardware such as FPGA or ASIC offer no such abstraction. For the later devices, floating point operations increase the complexity of the hardware design and the necessary hardware resources considerably. In terms of performance, general purpose devices benefit also from a sparse usage of floating point values. The complexity of the software development however is only marginally or not affected at all by the choice of data representation.

If by choice, scaling is omitted, a fixed point representation will not be possible, due to the rapid convergence towards zero by continuously multiplying probabilities together. This implies, that by omitting scaling to save resources, a floating point representation must be used, which again increases the resource requirements or has a negative impact on performance (or both).

The trade-off between the choice of using scaling or not versus the choice of the precision and the data representation, will be analyzed in more detail in chapter 5, when the technology of the accelerator has been chosen.

## 4.5 Extension of the Forward Algorithm

The proposed extension uses a transition probability matrix which is not constant. For every arriving observation symbol, the matrix must be recomputed by using the delay of the symbol (equation 3.1) and the sum of different cumulative distribution functions (equations 4.5 and

3.5). To compute  $N^2$  cumulative distribution functions is very expensive but it only needs to be computed once per  $d_k$  and can then be stored for later usage. A transition probability matrix can be used for the computation of  $L$  forward variables due to the continuous computation of likelihood values (as depicted in figure 4.1). This implies, that storage for  $L$  such matrices must be available or the matrix must be recomputed when needed and only the delay value is stored. In case of a pipelined architecture, the additional storage or computation is not necessary (cf. figure 4.2 and table 4.1). The computation of the transition probability matrix can be fully parallelized with  $R * N^2$  computation units to perform a cumulative distribution function of  $d_k$ , where  $R$  is the number of different cumulative distribution functions necessary to model the behaviour of the events. The memory interface needs to be able to provide a throughput that is  $R * N^2$  higher than in the sequential case (note that each cumulative distribution function takes several parameters as input. Eg. the normal cdf has the two parameters  $\mu$  and  $\sigma$ ). In previous chapters,  $R$  was always assumed to be equal to one in order to simplify the problem.

Note, that while the computation of one extended transition probability is independent of  $N$  or  $L$ , it is still a lot more complex than the simple multiply-accumulate operation necessary to calculate a forward variable (cf. the list below). Due to this, the computational units calculating the transition probability matrix must provide a lot more performance than the units necessary to compute the resulting forward variables in order to not limit the throughput. To get a rough impression of how expensive the computation of a distribution function is, the following list with three common examples is provided:

#### Exponential CDF

This distribution function describes the time between events in a Poisson process (events occur continuously and independently at a constant average rate). It is expressed as

$$F_{exp}(x) = \begin{cases} 1 - \exp(-\lambda x) & \text{if } x \geq 0 \\ 0 & \text{if } x < 0 \end{cases} \quad (4.4)$$

Only the exponential function is needed, which is quite a complex function compared to a multiplication and could be problematic in a fully parallelized implementation (considering all parallelization options), but realizable in an implementation with less parallelism.

#### Laplace CDF

This distribution is somewhat the extension of the exponential distribution and is also called double exponential distribution as it can be described as two exponential distributions put together (one flipped horizontally). It is expressed as

$$F_{laplace}(x) = \begin{cases} 1 - \frac{1}{2} \exp(-\frac{x-\mu}{b}) & \text{if } x \geq \mu \\ \frac{1}{2} \exp(\frac{x-\mu}{b}) & \text{if } x < \mu \end{cases} \quad (4.5)$$

In terms of complexity, for the Laplace distribution holds the same as for the exponential CDF.

#### Gaussian CDF

This is a very important distribution that is used in a lot of applications. It is used for real-valued random variables whose distributions are not known. The Gaussian (normal) cumulative distribution function cannot be expressed in terms of elementary functions, which is the reason why the special error function  $\text{erf}$  is used. It is defined by equation

3.7. Using integration by parts and the substitution  $x = \frac{d_k - \mu}{\sigma}$ , it can be expressed as

$$\Phi(x) = \frac{1}{2} + \frac{1}{\sqrt{2\pi}} \cdot \exp\left(-\frac{x^2}{2}\right) \cdot \left[ x + \frac{x^3}{3} + \frac{x^5}{3 \cdot 5} + \cdots + \frac{x^{2n+1}}{3 \cdot 5 \cdots (2n+1)} \right] \quad (4.6)$$

This computation is very expensive. It comprises of an exponential function and an iterative approach (to achieve the necessary precision) including the power function and additions. If this distribution is chosen to describe the time between events, parallelization will be very challenging, in order to prevent this calculation from being the bottleneck.

Considering the huge computation power needed to fully parallelize the extension, it may be beneficial to use a very specialized unit (ASIC) just for the computation of the cumulative distribution function.

Independent of the distribution, to not reduce the throughput of the fully parallelized computation of the forward variables, a small pipeline of two stages must be built, where in the first stage the transition probability matrix is computed and in the second stage the forward variables.

The correction of the diagonal elements (cf. listing 4.3) can be maximal parallelized by using  $N$  reduction trees to compute the sum of rows and  $N$  subtractors to correct the diagonal elements.

## 4.6 Reasonable Parallelism and Scalability

In the sections above, a lot of parallelization has been proposed. A maximal parallelization can hardly be achieved due to the immense requirement of recourses and is also not necessary because of dependencies. In a first step, a theoretical analysis (in terms of complexity order) of the different parallelization options and their combination is done. Then the results will be discussed and a choice will be made. Finally some conclusions about a finer degree of parallelization will be drawn in respect to the chosen architecture. Table 4.2 shows the pipelined architecture (cf. figure 4.2), the parallel architecture in the case of maximal row partitioning (cf. figure 4.3), the combination of both architectures and in the last column the combination of both architectures plus the reduction tree (cf figure 4.4). The basic and the extended forward algorithm are both of the same complexity order. As only the complexity order is considered, real computation times of different operations as well as scaling operations can be ignored. Due to simplification, it is assumed that  $N = L$  and that by scaling the problem,  $N$  and  $L$  are both changing in the same order.

The two first columns show that the smaller latency of the parallel architecture comes at the price of a larger memory interface (simultaneous memory access is required). By combining both architectures (3rd column), the throughput can be increased by factor  $N$  at the cost of increasing the order of computation units. Adding also the reduction tree, throughput and latency are increased, resp. decreased by an additional factor of  $\log N$ . This comes again at the cost of increasing the order of computation units further. While the benefits are welcome,  $N^3$  is simply to high to realistically implement such a solution. Already with a small  $N$  a huge server farm or thousands of FPGAs or GPUs would be necessary. Considering this, the reduction tree parallelization method will not be used as it gives the fewest benefits ( $O(\log N)$ ) for its cost ( $O(N)$ ). Computation units in an order of  $N^2$  (2nd column) is feasible for small  $N$  by combining multiple devices. While this may be acceptable for a very important computation where lots of

Metric	Pipelined	Parallel	Both	Both & Tree
Computation Units	$O(N)$	$O(N)$	$O(N^2)$	$O(N^3)$
Memory Space	$O(N^2)$	$O(N^2)$	$O(N^2)$	$O(N^2)$
R/W Access	$O(1)$	$O(N)$	$O(N)$	$O(N)$
Throughput	$\times N$	$\times N$	$\times N^2$	$\times N^2 \log N$
Latency	$\times 1$	$\times \frac{1}{N}$	$\times \frac{1}{N}$	$\times \frac{1}{N \log N}$

Table 4.2. Comparison of architectures in terms of complexity for the forward algorithm

		Pipelined	Parallel	
Basic	Computation Units	$L$	$N$	
	Memory Space	$2N^2 + 2LN + N$	$2N^2 + 3N$	
	Read Access	4	$N + 2$	
	Write Access	1	$N$	
Extended	Computation Units	$L + C$	$N + C$	$(C + 1)N$
	Memory Space	$(P + 1)N^2 + 2LN + N$	$PL(1 + N^2) + 3N$	$(P + 1)N^2 + 3N$
	Read Access	5	$2N + 2$	
	Write Access	1	$N$	

Table 4.3. Pipelined versus parallel architecture for the basic and the extended forward algorithm (C: Number of computation units to compute one CDF, P: Number of parameters to compute one CDF)

people depend upon (eg. weather forecast, Google queries, etc.) failure prediction hardly falls into this category especially if failures of an embedded system are predicted. Additionally, using multiple devices in order to scale the problem, implies off-chip-communication and -memory. This will result in huge bottlenecks and have a huge impact on the actual speedup. This leaves the first two columns to compare for the application at hand (A combination of both methods can still be considered, but not by using maximal parallelization).

Table 4.3 now only compares the pipelined architecture with the parallel architecture but with more detailed estimations of resource usage. The comparison is done for the basic and the extended forward algorithm (for explanations refer to the previous sections in this chapter). In case of the extended forward algorithm, when the parallel architecture is used, there is a choice to be made if the parallelization should be achieved by increasing the memory usage or the number of computational units (hence the two columns).

The benefits of a parallel architecture over the pipelined are first and foremost the reduced latency and in case of the basic algorithm also the lower memory footprint. If the acceleration architecture of choice has a memory interface that allows the required throughput, for the basic algorithm this architecture should be chosen. In case of the extended algorithm this is only possible if enough computational units are available (and the CDF computation is not too complex) or if the on-chip memory is large enough to save transition probabilities for later use.

For the basic algorithm, the pipelined architecture should only be chosen if the memory interface becomes the bottleneck (for large  $N$ ). Ideally a combination of the parallel architecture and the pipeline should be chosen in order to maximize the memory interface usage. By doing

this, a smaller latency is achieved by keeping the throughput high. Another reason to chose the pipelined architecture would be a simple state transition model (cf. section 4.3) which allows to save only the non-zero transition probabilities in a list. From this optimisation the pipelined architecture benefits on a much larger scale as less serial accumulations would be necessary, while in the parallel architecture only the utilization of the computational units would be reduced (less power consumption but no impact on performance).

In case of the extended forward algorithm, it is almost always better to choose the pipelined architecture: It uses either less memory or less computational units, allows optimization in case of simple models and allows more time to compute the transition probabilities. Parallelization is only possible for very simple CDF computations and for a small  $N$ .

The scalability is in both cases limited, but more so with the parallel architecture: If  $N$  or  $L$  becomes large such that off-chip memory is necessary (already for very small  $N$  or  $L$  for CPUs and GPUS, less so for more flexible architectures like FPGAs or ASICs) the memory interface will be to small to handle memory access simultaneously and hence become the bottleneck. The pipelined architecture does not have this drawback but has a slightly higher memory footprint. If memory can be handled on-chip but multiple chips are used to increase the number of recourses, both architectures can be scaled easily in  $L$  dimension (sequence length) but only with difficulty in  $N$  dimension (number of states) because the necessary communication links (all components of the vector  $\alpha_{k+1}$  always depends on all components of the vector  $\alpha_k$ ). The scaling in dimension  $N$  of the pipelined architecture only depends on the memory usage, while the parallel architecture has a dependency of  $N$  for the computation units as well as for the memory.

## 4.7 Choice of Accelerator Type

For being able to choose an appropriate accelerator type, first a list with different acceleration options is presented. For each type the most common benefits and shortcomings are mentioned. At the end of the section a choice will be made using the following list and the observations discussed in the previous sections.

### CPU

The Central Processing Unit falls into the class of the general purpose processors and is (usually) of the type SISD<sup>1</sup>. It is very flexible in terms of software interpretation (with the use of compilers) and allows to implement any kind of function in a fast, easy and maintainable fashion with no requirements in hardware knowledge. Operations like division, exponential function and logarithm are available as well as the floating point number representation. All operations can be executed with very high precision and at high clock frequencies (up to 3 GHz). CPUs come with the drawback of a high power consumption, limited parallelization options (small number of cores) and a fixed hardware architecture that causes big computation overheads (instruction pipeline, memory hierarchy, generalized computation units).

### GPU

Like the CPU, the Graphics Processing Unit still falls into the class of general purpose processors due to the hardware abstraction layers. A GPU is composed of a lot of small

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<sup>1</sup>Flynn's Taxonomy: Single Instruction, Single Data Stream

but quite powerful streaming processors of the type SIMD<sup>2</sup>. This processing power allows a lot of parallelization at a low price. As the CPU, also the GPU is very flexible in terms of software interpretation, requires however some hardware understanding in order to use the parallel power in an optimal way. Also a GPU allows to work with operations like division, exp and log functions and provides floating point number representation. GPUs can work with high precision and at high clock frequencies. The power consumption of a GPU is very high due to the high frequencies and the streaming processors. While the hardware abstraction layer provides flexibility in software and ease of use, it is the main reason for a computation overhead for basic operations. The fixed hardware architecture (especially the memory hierarchy) can prove to be a drawback in specific cases (eg. data usage provokes always cache misses).

#### DSP

The Digital Signal Processor is a specialized integrated circuit, largely used for digital signal processing. The key components of DSPs are optimized Multiply-Accumulate instructions, special SIMD operations and for DSP operation optimized memory architecture. They provide a hardware abstraction and are pretty easy to program. This causes some overhead as an instruction pipeline is necessary. The overhead is a lot smaller than in the case of CPUs because of specific instructions sets (this comes with a loss of flexibility). Fixed point as well as floating point devices exist with various precisions. A DSP provides a lot of specific computation power for a low price.

#### FPGA

The Field Programmable Gate Array is a customizable integrated circuit. It provides large recourses of logic gates as well as standard blocks such as RAM or highly optimized Multiply-Accumulate units. FPGAs provide high performance for very specific design optimized for one function. Once an FPGA is configured, as long as it keeps this configuration no other function will run on this device as it is the direct hardware representation of the function. This direct representation allows a very low overhead as operations are done directly in hardware without any instruction pipeline. FPGAs provide a high amount of hardware flexibility that is only topped by ASICs (see below). This flexibility comes at a medium price as FPGAs can be produced in big lots but are more difficult to produce than DSPs. A huge advantage is the possibility the build very big memory interfaces inside the chip and customize the memory architecture for the application at hand. The drawback of FPGAs is the increased development time necessary to implement a hardware solution of a function, the "low" clock frequency of up to 500MHz (this is low compared to CPUs or GPUs) and the absence of division, exponential and logarithm functions. All units are optimized for fixed point representation and floating point numbers must be implemented manually. Core generators and very powerful synthesis tools try to amend these drawbacks but still a very deep knowledge of hardware is necessary to successfully implement a design on FPGA.

#### ASIC

The Application Specific Integrated Circuit is, as the name already tells, an integrated circuit that has been designed for one (and only one) specific application. In comparison to the FPGA, an ASIC is built with fully customized elements and provides the full flexibility achievable with todays hardware knowledge. An ASIC has ideally no overhead as the

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<sup>2</sup>Flynn's Taxonomy: Single Instruction, Multiple Data Streams

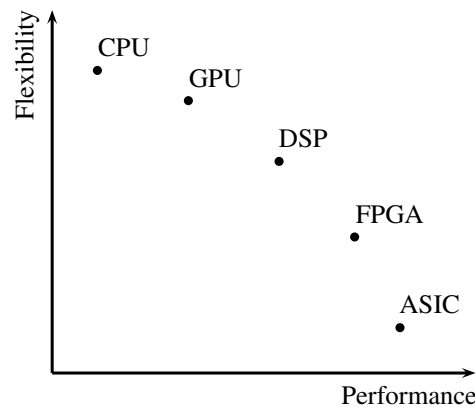


Figure 4.5. Flexibility versus Performance of Hardware Devices

hardware is a direct mapping of the function. This leads to very high performance at very low power consumption (as a rule of thumb a factor of 1000 can be assumed in either performance gain or power consumption decrease or a combination of both compared to CPUs). An ASICs is very expensive to produce. This includes the long development time and the production cost. Of course very deep hardware knowledge is necessary to create an ASIC of a specific function.

Comparing the accelerator types mentioned in the list above in a general manner, one can conclude that the further down the list one goes, the better is the performance - power consumption ratio and the lower is the flexibility of the device to accept general function descriptions (cf figure 4.5, eg. a function description for a FPGA must be a lot more specific than one for a GPU but the performance-power consumption ratio is a lot better in case of the FPGA). For this work, one key point is high performance, because the algorithm to implement is computation intensive and needs to be executed fast. Another important point is the size of the system: the target application to predict failures is an embedded system, where space is usually limited. Power consumption may not be a main aspect, but it certainly needs to be considered to not exceed requirements of the main system.

Due to the limited parallelization options or the huge power and space requirements of CPUs, this accelerator type is not suitable for the application in question. GPUs provide a lot of computation power at a low price but have drawbacks in terms of architecture flexibility, space and power requirements. GPUs are a far better option than CPUs but may not be ideal because of the fixed memory interface. DSPs would be the device of choice if only the basic forward algorithm is considered. Due to the high performance at a low cost of DSPs, and the fact that the basic forward algorithm mainly uses multiply-accumulate operations one or multiple floating point DSP devices could be used to efficiently implement the algorithm. In case of the extended algorithm a DSP device causes to big an overhead to compute the CDF and will be the bottleneck of the design. For this reason a more flexible device must be chosen. The most flexible architecture is an ASIC. ASICs are very expensive in terms of money and development time, two recourses that are not available for this work. This leaves the FPGA: An FPGA combines the parallel power of DSPs for Multiply-Accumulate operations but adds the possibility to design a specific hardware architecture to compute the transition probabilities needed for the extension. While the performance will not be at a level of an ASIC, the specialized

hardware will still outperform any other device because of the customizable memory interface (even at the lower frequencies of FPGAs).



## Chapter 5

# design and implementation

- parallelization of one of nested for-loop or pipeline the loops
- fully pipelined macc

to design the accelerator, the top-down approach was applied: the algorithm is broken down into blocks, where each of them is broken down further until the basic functional blocs of the fpga can be used for the implementation. the implementation follows then the bottom-up approach where each sub-block is implemented and tested. completed blocks are grouped together to bigger blocks until finally there is only one big block remaining, describing the complete algorithm.

### 5.1 precision

modern fpgas contain hardware blocks (dsp slices) that are heavily optimized for multiply-accumulate operations as they are often used for dsp applications. these devices only operate on integer values. however, the forward algorithm requires to multiply and accumulate probabilities, i.e. fractional numbers. to being able to use the dsp slices of the fpga, some hardware must be built around such a slice in order to get a hardware block that can handle the multiplication and accumulation of probabilities. to do this, the choice of data representation must be made (floating point vs fixed point). first the assumption that all operations must be done with floating point numbers is discussed.

*the following points need to be explained in more detail*

floating point facts: in order to multiply two floating point numbers, the mantissas of both numbers are multiplied and the exponents are added (the sign can be ignored, as probabilities are always positive). for the multiplication of the mantissas the dsp slices can be used. in parallel to this operation, an external adder can add up the exponents. now the tricky part: to being able to add floating point numbers the exponents of the numbers must be equal. to achieve that, the difference of the two exponents is calculated and then the mantissa of the number with the lower exponent must be shifted by this difference (and possibly rounded/truncated). this process is called normalizing. then the mantissas can be added. finally the resulting value needs to be normalized again.

scaling: by using floating points, scaling becomes superfluous because the exponent allows to represent very small numbers (not respecting ieee representations).

truncation: after the multiply accumulate operation the resulting mantissa of 48 bits must be truncated to 25 bits. to achieve better results, a rounding operation can be applied.

fixed point facts: before each multiplication, the operands need to be reduced from 48 to 18 resp 25 bits. this must be done by either truncating/rounding the final value or by scaling or by doing both.

scaling: the proposed scaling method is not usable in this implementation because of the division operation. instead, a scaling in base of 2 should be used (shift): in every clock cycle a new value is added to the previous one. the masked result can be compared to a bit stream of zeros. a match directly gives the amount of leading zeros (using the number of times the mask has been shifted). if there is no match, shift the mask and compare in the next cycle. this operation is done with every alpha, while the lowest value of leading zeros is kept. using this method, in most cases at the end of adding up all the values, the least number of leading zeros of the  $n$  alphas will be known. if in the worst case after the addition of the last alpha-component there is a non-match, the pipeline must be stalled and the right leading zeros number must be found by shifting the mask further to the left. to do the masked comparison, the pattern comparing unit of the dsp-slice of the fpga can be used. the stored values  $\pi$ ,  $b$  and  $tp$  can be preprocessed and scaled, in order to reduce the number of minimal leading zeros to zero. these scaling factors will then be used to recompute the real sequence likelihood.

truncation: a simple solution is to chop off the leading zeros (as counted) and truncate the value by only using the following 25 bits after the last leading zero. a more precise method would be to use rounding.

[26, 9] discuss the choice of fixed point versus floating point representation with respect to DSP devices. This is also applicable for FPGAs.

## 5.2 data storage management

*the following points need to be explained in more detail*

facts: in each  $k$ th step  $n^2 * tp$  values and  $n * b$  values are necessary. in case of the parallel version, each clock cycle  $n * tp$  and one  $b$  value must be made available. in case of the serial version one  $tp$  or  $b$  value must be made available. by reading directly from the ram, 16 bits can be read with a bus frequency of 104mhz. running the pipeline at 52mhz, the necessary data can be provided at each clock cycle in case of the serial implementation. if the parallel implementation should be used, an internal memory must be built.

- real-time (time constraints on every ps) vs on-line (use buffer to optimize throughput)
- memory hierarchy: at startup copy from flash to ram, then use pipeline to preload values from ram into registers.

## 5.3 initialization

figure 5.1 shows the necessary operation to compute every  $i$ -th component of the first forward variable  $\alpha_1$  in the initialization step. the  $n$  components of  $\alpha_1$  can be computed one by one when only one such structure is implemented in hardware. assuming a fully pipelineable multiplier, one component of  $\alpha_1$  appears in the output register at every cycle (with a latency of  $s$  cycles, where  $s$  is the number of pipeline stages in the multiplier). to compute all  $n$  components of  $\alpha_1$

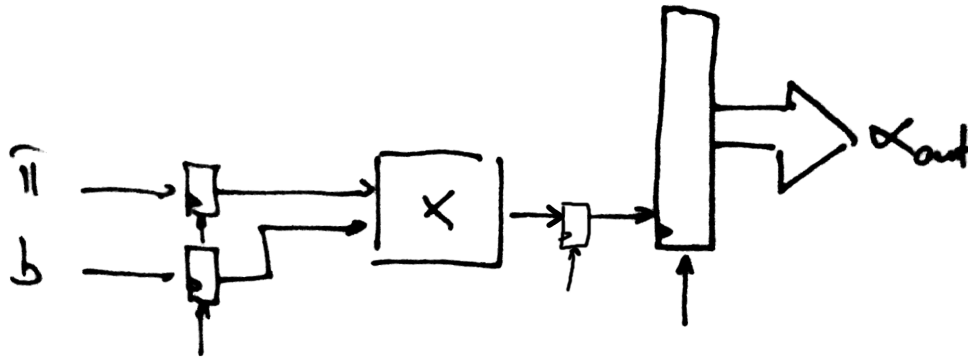


Figure 5.1. initialisation step with one macc

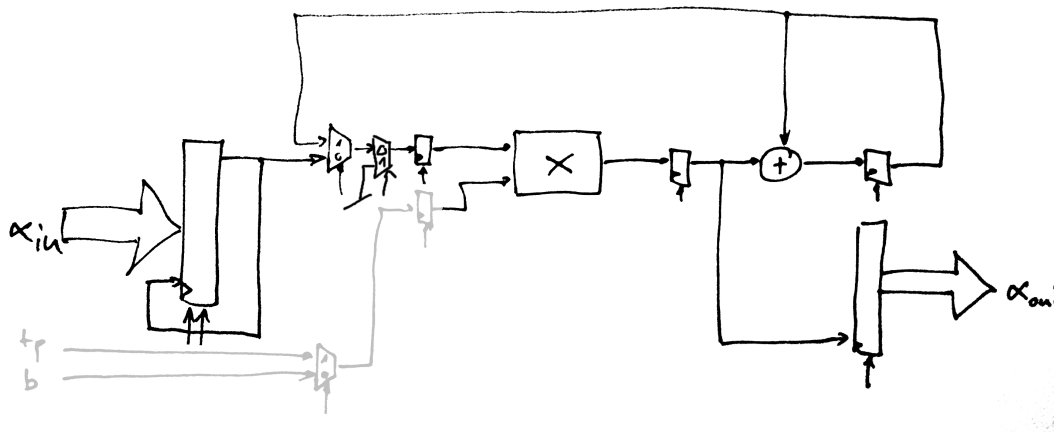


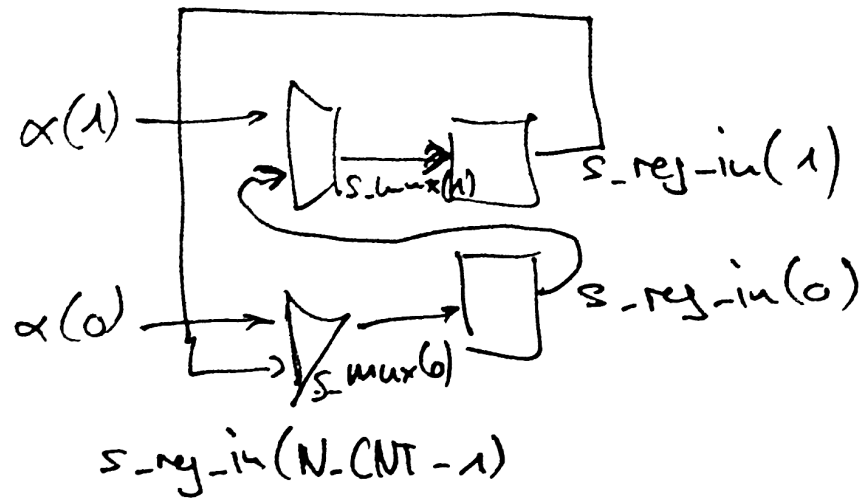
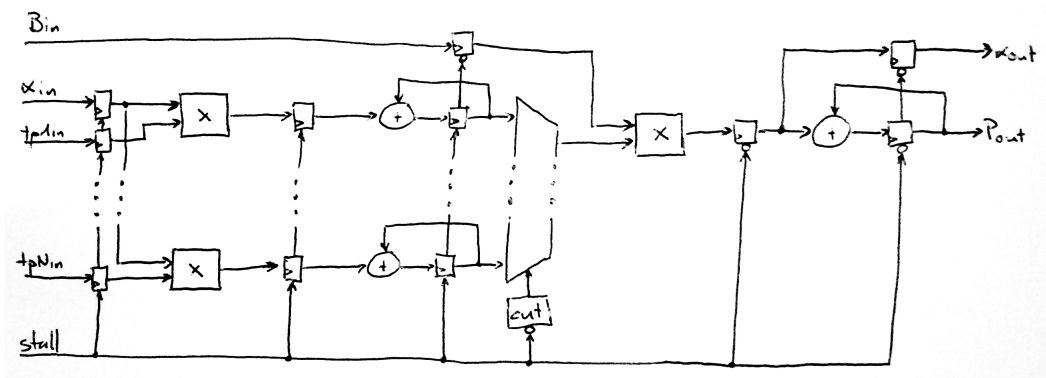
Figure 5.2. k-th step with one macc unit

with this structure,  $n + s$  cycles are necessary. alternatively the complete initialization can be fully parallelized if the structure 5.1 is replicated  $n$  times. in this case, all  $n$  components of  $\alpha_1$  appear at the same time, after  $s$  cycles in the output registers. if such a heavy parallelization is useful will be analyzed after all stages have been described.

## 5.4 k-th forward variable

*better title?*

- one pipelined macc for the computation of all elements
- $n$  pipelined macc, each computing one  $\alpha_k$
- use optimized matrix-vector-vector multiplication  $\alpha_{k+1} = tp * \alpha_k * b(o_k)$  [13, 30]

Figure 5.3. example of input shift register with  $n=2$ Figure 5.4.  $k$ -th step with  $n+1$  macc units and integrated computation of  $p$  (which is only needed at the last step)

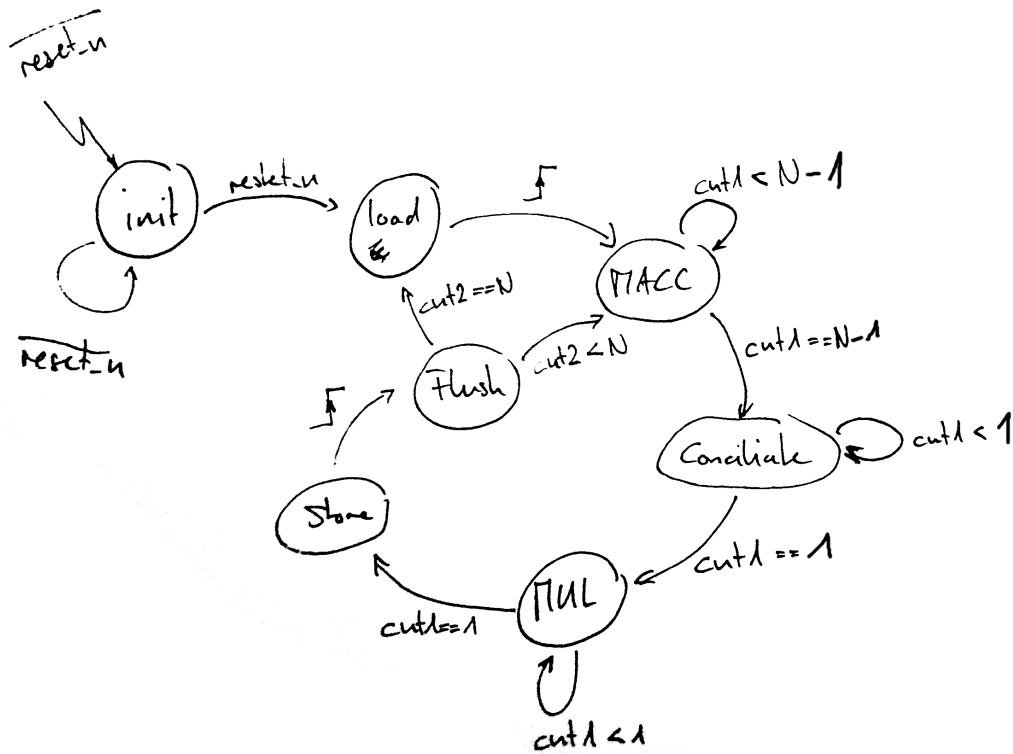


Figure 5.5. controller of serial implementation

## 5.5 serial controller

*in table ?? the signals `flush_ps` and `load_out` are only set in the first iteration. add a footnote?*

## 5.6 balancing pipeline stages

## 5.7 practical notes

- generic design (change `n`, `l` and bit widths in `param_pkg`)
- if bit widths of `op1` is changed, `fifo_512x25` must be regenerated

type	signal	init	select	macc	conciliate	shift1	shift2	mul	store	flush	input
	pi_we										input
	tp_we										input
	b_we										input
	data_ready										input
1	enable_count	0	0	0	0	0	0	0	1	0	pi_we
2	enable_ctrl	0	0	0	0	0	0	0	0	0	data_ready
3	enable_init	0	0	0	0	0	0	1	0	0	data_ready
3	enable_init_mul	0	0	0	0	0	0	1	0	0	data_ready
4	enable_step	0	0	1	1	1	1	1	0	0	data_ready
4	enable_step_macc	0	0	1	1	1	1	1	0	0	data_ready
5	enable_final	0	1	0	0	0	0	0	0	0	data_ready
4	laod_op2	0	0	1	1	1	1	1	0	0	data_ready
6	load_step_alpha	0	0	1	0	0	0	0	0	0	
7	load_final_alpha	0	0	0	0	0	0	0	1	0	
8	load_scale_new	0	1(d)	0	0	0	0	0	0	0	
9	load_scale_acc	0	1	0	0	0	0	0	0	0	
7	store_init_scale_new	0	0	0	0	0	0	0	1	0	
23	store_init_scale_small		1(d)								or internal
9	store_init_scale_ok	0	1	0	0	0	0	0	0	0	
7	store_step_alpha	0	0	0	0	0	0	0	1	0	
7	store_step_scale_new	0	0	0	0	0	0	0	1	0	
23	store_step_scale_small		1(d)								or internal
9	store_step_scale_ok	0	1	0	0	0	0	0	0	0	
5	store_final_ps	0	1	0	0	0	0	0	0	0	data_ready
10	store_final_ps_delayed	0	1(d)	0	0	0	0	0	0	0	data_ready
10	store_final_scale	0	1(d)	0	0	0	0	0	0	0	data_ready
11	shift_step_acc	0	0	0	1(6)	0	0	0	0	0	
12	sel_mux2_op2	0	0	0	0	0	0	1	0	0	
13	sel_step_read_fifo		(*)								special
12	sel_step_op1	0	0	0	0	0	0	1	0	0	
14		0	0	0	1	1	0	0	0	0	
15		0	0	0	1	0	1	0	0	0	
16	sel_step_op2	0	0	0	1(5)	1	1	0	0	0	
17	flush_init	0	0	0	0	0	0	0	0	1	
17	flush_step_macc	0	0	0	0	0	0	0	0	1	
18	flush_step_acc	0	0	0	0	0	1	0	0	0	
9	flush_step_fifo	0	1	0	0	0	0	0	0	0	
	reset_n	0	0	0	0	0	0	0	0	0	input
19	reset_count_n	0	0	0	0	0	0	0	0	0	reset_n
19	reset_op2_n	0	0	0	0	0	0	0	0	0	reset_n
19	reset_ctrl_n	0	0	0	0	0	0	0	0	0	reset_n
19	reset_init_n	0	0	0	0	0	0	0	0	0	reset_n
20	reset_init_mul_n	0	0	0	0	0	0	0	0	1(n)	reset_n
19	reset_step_n	0	0	0	0	0	0	0	0	0	reset_n
20	reset_step_macc_n	0	0	0	0	0	0	0	0	1(n)	reset_n
21	reset_step_fifo0	0	1(*n))	0	0	0	0	0	0	0	reset
22	reset_step_fifo1	0	1(*)	0	0	0	0	0	0	0	reset
19	reset_step_scale_new_n	0	0	0	0	0	0	0	0	0	reset_n
19	reset_step_scale_small_n	0	0	0	0	0	0	0	0	0	reset_n
19	reset_step_scale_ok_n	0	0	0	0	0	0	0	0	0	reset_n
19	reset_init_scale_new_n	0	0	0	0	0	0	0	0	0	reset_n
19	reset_init_scale_small_n	0	0	0	0	0	0	0	0	0	reset_n
19	reset_init_scale_ok_n	0	0	0	0	0	0	0	0	0	reset_n
19	reset_final_n	0	0	0	0	0	0	0	0	0	reset_n

Table 5.1. detailed control signals

## Chapter 6

# testing and verification

### 6.1 device

- nexys4 board with artix-7 fpga
- limited recourses -> proof of concept
- bord hardware for testing

### 6.2 relation to proposed algorithm

#### 6.2.1 log standard

#### 6.2.2 metrics





## Chapter 7

### results

7.1 speedup

7.2 accuracy



## Chapter 8

### conclusion

8.1 achievements

8.2 future work



Appendix A

some material



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