

# A New Hardware Architecture for Genomic and Proteomic Sequence Alignment

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## Abstract

We describe a novel hardware architecture for genomic and proteomic sequence alignment which achieves a speed-up of two to three orders of magnitude over Smith-Waterman dynamic programming (DP) in hardware [1]-[7]. In [8, 9] we introduce several features of our search algorithm, DASH, which outperforms NCBI-Blast (BLAST) [10] by an order of magnitude in software, and has better sensitivity. Indeed, DASH has been shown to have excellent sensitivity compared to Smith-Waterman. It is designed around the principle of considering genomic and proteomic sequence alignments to typically consist of regions of high homology (the diagonals) interspersed with regions of low homology. In DASH, the optimal solution consists of such diagonals joined by regions of exact DP. This is affordable due to the small area of these inter-connecting regions. Accordingly, we have designed a chip which finds the diagonals and performs the inter-region DP directly in hardware. On a Xilinx Virtex II, XC2V6000, FPGA, it performs over  $10^{12}$  base comparisons/second.

## 1. Introduction

Heuristic algorithms such as BLAST [10] are indispensable for searching today's large genomic and proteomic databases. A significant contributor to the search time for such algorithms is the DP evaluations, BLAST spends around 76% of its time budget in this area. We have developed a novel sequence alignment algorithm, DASH, which we have shown to be superior to BLAST in both speed and sensitivity [8, 9], and excellent sensitivity with respect to Smith-Waterman. Our project was originally motivated by the desire to develop an algorithm which would allow extensive parallelism for optimum use of reconfigurable hardware, such as FPGAs. In this context we designed DASH around the principle of considering ge-

nomomic and proteomic sequence alignments to typically consist of regions of high homology interspersed with regions of low homology, Figure 1. Finding the regions of high homology (the diagonals, zone 1 in the figure) requires high throughput, but low processing power, which is ideally matched to the massively parallel resources available on FPGAs (Field Programmable Gate Arrays).

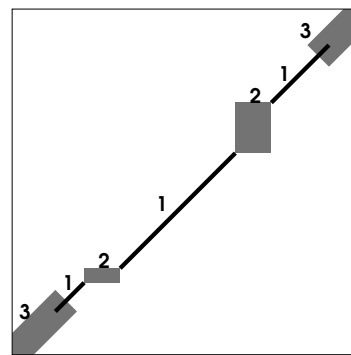
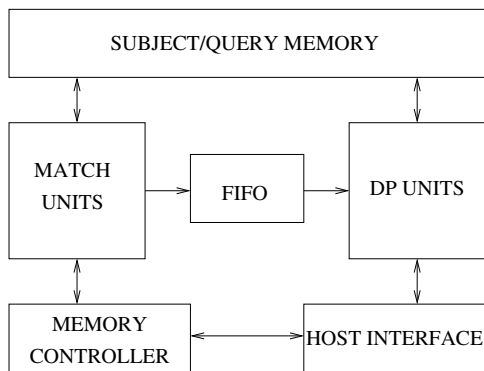


Figure 1. Illustration of the DASH Algorithm and DP Zones

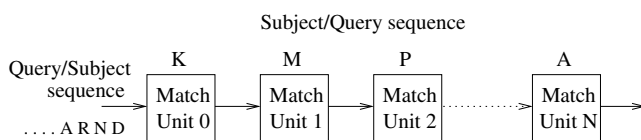
The regions of low homology, zones 2 and 3 in the figure, are solved by DP, but as they are strictly limited in size, the extra hardware resources required are small.

## 2. The Architecture

The architecture of our design is given in Figure 2. The query and subject sequences are input to a match pipeline which finds the diagonals, Figure 3. Here, each individual match unit finds either four base matches in the nucleotide case, or one amino acid match in the protein case. Substitutions, but not gaps, are allowed. Those diagonals of more than a specified length and score are then output to a FIFO (first-in first-out buffer). The DP units then take the diag-



**Figure 2. Architecture of the Sequence Alignment Hardware**



**Figure 3. Match Unit Pipeline**

onals from this FIFO and find the optimum path for the inter-diagonal regions, zone 2, and the ends, zone 3. The raw data is filtered by the Match Unit at high-speed (Table 1) giving the DP Unit more than enough time to process the inter-diagonal regions. Given sufficient on-board memory both these units can operate in parallel, and the critical overall system delay is the throughput of the Match Unit. There are several well-known architectures for the processing elements in the DP Unit, e.g. [1]-[6].

### 3. Results and Conclusions

The match units were coded in VHDL and extensively tested against DASH with DNA/Protein sequences of up to half a million in length. The area (no of slices) and timing results for a single match unit after synthesis and place and route are given in Table 1. It can be seen that in the nucleotide case over 10,000 match units will fit on a Xilinx Virtex 2, XC2V6000, FPGA (total of 33792 slices), with a maximum throughput of over  $2 \times 10^{12}$  base comparisons/sec. With only 50% utilization of the device, we still obtain over  $10^{12}$  base comparisons/sec, and allocate the other half of the device to the DP Unit, memory controllers, and the interface unit for the host processor. The maximum throughput in the protein case is  $3 \times 10^{11}$  amino acid matches/sec.

Results for full DP for the same FPGA can be found in [5]. Depending on the maximum query length, they obtain a

	DNA	Protein
Slices	3.2	19.2
Clock	210 MHz	180 MHz

**Figure 4. Area and Timing results for a DNA/Protein Match Unit**

peak throughput of  $5.7 \times 10^9$  base matches/sec. Clearly, the architecture described here affords two to three orders of magnitude greater performance.

### 4. Acknowledgements

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