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Biosequences analysis on NanoMagnet Logic

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Abstract—In the last decade Quantum dot Cellular Automata technology has been one of the most studied among the emerging technologies. The magnetic implementation, NanoMagnet Logic (NML), is particularly interesting as an alternative solutions to CMOS technology. The main advantages of NML circuits resides in the possibility to mix logic and memory in the same device, the expected low power consumption and the remarkable tolerance to heat and radiations. NML and QCA circuits behavior is different w.r.t. their CMOS counterparts. Consequently architecture organization must be tailored to their characteristics, and it is important to identify which applications are best suited for this technology.

Our contribution reported in this paper represents a considerable step-forward in this direction. We present an optimized implementation on NML technology of an hardware accelerator for biosequences analysis. The architecture leverages the systolic array structure, which is the best organization for this technology due to the regularity of the layout. The circuit is described using a VHDL model, simulated to verify the correct functionality from the application point of view, and performance are evaluated, both in terms of speed and power consumption. Results pinpoints that NML technology with the appropriate clock solution can reach a considerable reduction in power consumption over CMOS. This analysis highlights quantitatively, and not only qualitatively, that NML logic is perfectly suited for Massively Parallel Data Analysis applications.

I. INTRODUCTION

¹ Quantum dot Cellular Automata (QCA) was presented in [1] as an alternative to CMOS technology. In QCA charge states are used to represent logic values instead of voltage levels. Among the implementations of the QCA principle, NanoMagnet Logic (NML) is one of the most successful [2]. The reason behind the success of NML is the magnetic nature: Rectangular shaped single domain nanomagnets with only two stable states (Figure 1.A) are used as basic cells [3]. Since the basic circuitual element is a magnet, it can act both as a logic device and a memory, allowing the development of a totally new kind of logic circuits, where memory and logic are no more separated entities. Moreover NML logic has a strong resistance to heat and radiations and potentially the power consumption can be tens of times smaller than CMOS logic [4].

NML circuits are built placing magnets on a single plane. Information propagates through the circuit by means of magnetostatic interaction among neighbor elements achieving simple interconnect functions or more complex logic functionalities (for example see a Majority Voter – MV [3] –in Figure 1.B we fabricated). To obtain information propagation a clock mechanism is required [5], about which a detailed explanation

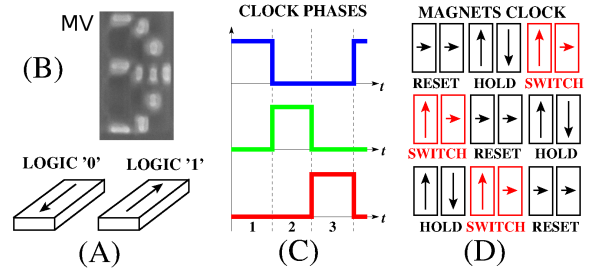


Fig. 1. A) Nanomagnets are used as basic cell. B) Example of a fabricated Majority Voter (thanks to Nanofacility Piemonte - INRIM Torino). C) Clock signals waveforms. D) Clock zones and signal propagation.

is given in section II. This unavoidable requirement has an important consequence at the architectural level: circuits have an extremely deep level of pipelining (see section II) and, consequently, in presence of feedback throughput is notably reduced. The solution we propose in this paper is exploiting *data interleaving*. We demonstrate the problem and the solution with a detailed implementation based on NML of a Systolic Array, that is considered particularly suitable for this technology [6]. Systolic arrays are circuits made by a network of identical processing elements locally interconnected. The regular layout and the absence of long interconnections make them ideal for NML technology. However in the circuits proposed in literature [6] processing elements are very simple making therefore difficult a correct evaluation of this solution. In this paper we show a complex systolic array design of a NML circuit, applied to a biosequences analysis hardware accelerator [7]. Biosequences analysis is interesting for NML technology because the huge amount of data to process allows massive parallelization maximizing the throughput. With this work we achieve therefore two important results: 1) We demonstrate a complex systolic array NML architecture applied to a real life case of study quantifying advantages and disadvantage and 2) we show the path that must be followed in the development of this technology, highlighting what kind of applications can be successful in this technology.

II. NML CLOCK SYSTEM: PROBLEMS AND SOLUTIONS

The clock system. In NML, when the clock signal is applied, magnets are forced in an intermediate unstable state, with the magnetization vector directed along the shorter magnets side. This is achieved by means of an external magnetic field. When the magnetic field is removed magnets align themselves antiferromagnetically following an input magnet. Since the number of magnets that can be aligned without incurring in errors during the realigning phase is limited [8], a multiphase clock system must be used. Three variable clock signals (Figure 1.C for a simplified representation) with a phase difference of 120° are applied to specific areas of the

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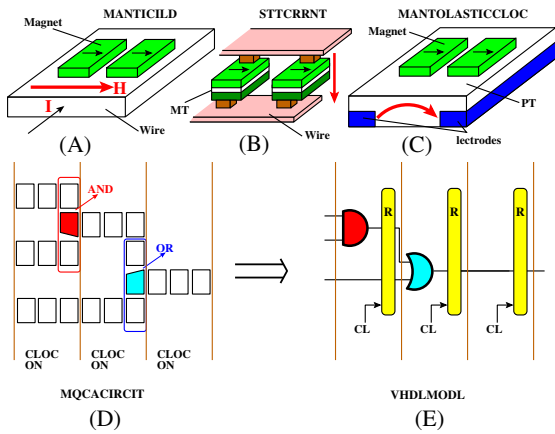


Fig. 2. A) Magnetic field based NML. B) STT-current based NML. C) Magnetoelastic NML. D) Circuit example. E) VHDL model.

circuit layout called clock zones (Figure 1.D). Clock zones are made by a limited number of chained magnets, typically 5, assuring an errorless signals propagation. Using this clock system, when magnets of a clock zone are in the SWITCH state (magnetic field removed), magnets of the left clock zone are in the HOLD state (no field applied) and act as inputs, while magnets of the right clock zone are in the RESET state (magnetic field applied) and have no influence on the switching magnets. Figure 1.D shows the signal propagation through a NML wire (a simple chain of magnets) thanks to the multiphase clock system.

The clock physical implementation. The external magnetic field necessary for the clock mechanism can be generated by a current flowing through a wire placed under the magnets plane (Figure 2.A). As shown in [3] a current of 545 mA on a 1 μm width wire is required to successfully switch magnets in the RESET state. This is a very high value of current that leads to a very high power consumption, wasting the advantage related to the tiny power dissipation due to magnet switching. To reduce power consumption other mechanisms were proposed. For example in the STT-current approach [9] magneto-tunnel junctions (MTJ) are used as basic element. MTJ can be reset by a current flowing through them leading to a power consumption of just 1.6fJ for each magnet (Figure 2.B). Alternatively, in [4] we proposed an innovative clock system based on the use of an electric instead of a magnetic field. With this clock solution magnets are deposited on a piezoelectric layer (Figure 2.C). When an electric field is applied the strain of the piezoelectric layer induces a mechanical stress on the magnets forcing them in the RESET state. With this clock solutions an energy of just 2 pJ is required to switch magnets [4] allowing to build true low power circuits.

Consequences at architectural level. The most important consequence of this system is that every consecutive group of three clock zones has a delay of one clock cycle. NML circuits (and QCA in general) have therefore an intrinsic pipelined behavior, where the level of pipelining is not a choice of the designer but it depends on the circuit layout. For example signals that propagate through a wire will have a propagation delay, in terms of clock cycles, proportional to the wire length. This is particularly important if there are loops in the circuit [10]. The presence of a loop prevents from sending new data every clock cycle, because before sending a

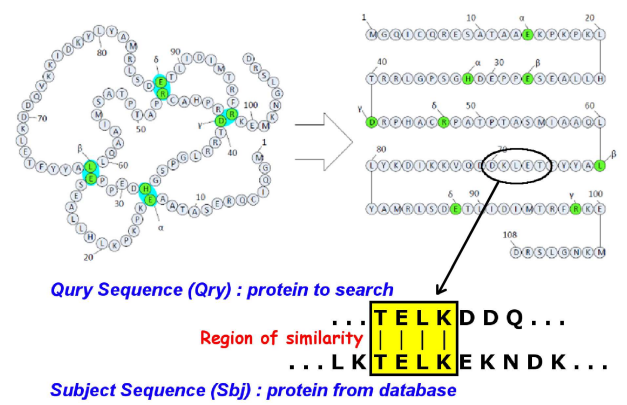


Fig. 3. Biosequence Alignment analysis

new data, propagation of signals through the feedback path must be waited for. This might require hundreds of clock cycles [11]. A problem which is typical of pipelined microprocessors in CMOS is here enhanced due to the extremely deep pipelining involving both logic and interconnections. To solve this problem parallel computation can be exploited using the interleaving technique. For example in case of a NML microprocessor [11], typically the output of the ALU is connected back to the circuit with a loop. Before a new instruction can be sent to the circuit it is necessary to wait the propagation of the result of the first operation through the feedback path. However if N independent threads are run in parallel, where N is the length in clock cycles of the loop, at every clock cycle it is possible to send a new instruction of a different program. In this way the pipe is always full and the throughput is maximized. In section III we show how we implemented this solution for a biosequence analysis systolic array structure.

Methodology. The multiphase clock system can be used to build a VHDL model of NML circuits [12]. Taking as an example the circuit shown in Figure 2.C, thanks to the pipelined nature of this technology, every clock zone is equivalent to a register with the same clock signals of Figure 1.B applied. At every new clock cycle a new data is sampled by the clock zone. Figure 2.E shows therefore the equivalent VHDL model of the circuit: Registers are used to model the clock zones and therefore the propagation delay of signals while ideal logic gates are used to model the logic functions. In this case AND/OR gates, based on [13] are used as an example. With this model it is possible to easily design and simulate complex circuits, and was used to design and simulate the architecture presented in this paper.

III. ALGORITHM AND ARCHITECTURE

Algorithm. Biosequences alignment analysis is a field which is constantly growing. Proteins are the fundamental constituents of animal and plant cells and are composed of chains of 23 Amino Acids (AAs) (Figure 3) which are normally represented by alphabetical characters. The aim of biosequences analysis is therefore to identify similarities between sequences of amino acids, for example to reconstruct the evolutionary pathway that led to the differentiation of species or for understanding the genetic cause of a disease comparing a mutated cell with a normal one. Biosequences alignment

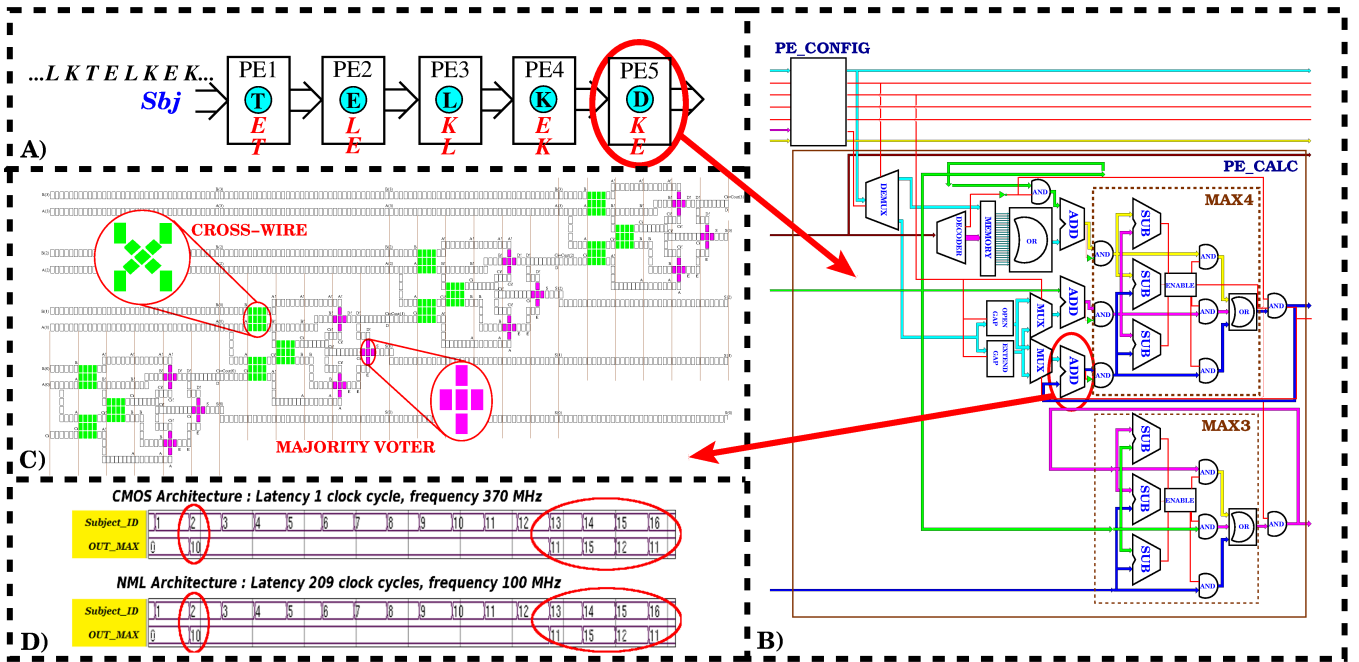


Fig. 4. A) NML Smith-Waterman systolic array. B) Detailed layout of a processing element. C) Circuit detail of an adder. D) Simulation results: Comparison between CMOS and NML.

analysis is commonly done by comparing one sequence of amino acids (*Query*) with the ones from the databases (*Subject*) that have been developed from other studies on genome sequencing projects. Such comparisons are done by aligning the sections of the sequences to find out their maximum similarity (Figure 3). This can be a costly task thanks to the exponential growth of biosequences databases, so it is important to speed up algorithms also using dedicated hardware accelerator [7]. One of the most used algorithm for biosequences analysis is the Smith-Waterman [14], which evaluates exhaustively the best alignment score between a *Query* sequence and a *Subject* sequence from the database.

Architecture. The circuit architecture is shown in Figure 4.A. It is based on a systolic array structure organized in a chain of identical processing elements (PEs). Each amino acid of the *Query* sequence that must be studied is associated to one PE. As a consequence, the higher is the number of PEs, the longer is the sequence of amino acids that can be studied. Amino acids of the *Subject* sequence coming from the database are sent as inputs to the systolic array. The detailed layout of a processing element is shown in Figure 4.B. At the beginning, in each processing element one amino acid of the *Query* sequence is loaded. This task is executed by a configuration block (*PE_CONFIG*), which stores in the memory 23 values. These values represent, for a specific amino acid of the *Query* sequence, its alignment score, or, in other words, its relation with all the other amino acids. These values are used to calculate the maximum alignment score by the *PE_CALC* part of the circuit (Figure 4.B). After the initial configuration phase, amino acids of the *Subject* sequence coming from the database are sent to the first processing element and pass through the entire chain of processing elements. Each of them calculates the local alignment score using a circuit which is based on adders and subtractors. The local alignment score is passed from one processing element to the other, until it reaches the

end and becomes the maximum alignment score for a specific sequence of amino acids. The sequence with the highest score value is therefore the *Subject* sequence more similar to the *Query* sequence which was under investigation. Further details, not given here for space reason, can be found in [7].

Layout. Figure 4.C shows the circuit layout of an adder based on NML logic. It is a simple ripple carry adder, which our analysis demonstrated to be one of the more efficient in this technology. Every full adder is based on majority voters (MV in Figure 1.B), where the output is equal to the majority of inputs [15]. Cross-wires are instead particular blocks that allow to cross two wires on the same plane without interference.

IV. RESULTS

In order to compare the performance of the NML implementation with CMOS technology, a 5 elements systolic array has been also described and synthesized using a standard CMOS technology [7]. Figure 4.D shows the comparison between the waveforms obtained in CMOS case and in NML case. *Subject_ID* represents the amino acids subject sequence number while *OUT_MAX* represents instead the sequence maximum alignment score. The two simulation waveforms show the same results, demonstrating the correctness of the circuit. However, it is worth underlining that the waveforms have been normalized to better compare them. The timescale is different: In CMOS the clock frequency is around 370 MHz and the latency for on PE is 1 clock cycle. In the case of NML the frequency is 100 MHz (considering a realistic implementation) and the latency is 209 clock cycles, meaning that a new amino acid can be sent only every 209 clock cycles. The long latency is due to its NML intrinsic pipelined nature previously explained and therefore to the long propagation time of feedback signals [10] that here are in evidence. Using this straightforward implementation, before sending a new amino

TABLE I. POWER CONSUMPTION AND AREA ESTIMATION FOR A SINGLE PROCESSING ELEMENT OF THE SYSTOLIC ARRAY, FOR THE MAIN NML IMPLEMENTATIONS AND FOR CMOS LOP 21nm TECHNOLOGY.

	Area (μm^2)	Power (mW)
Magnetic Field NML	21000	2
STT-current NML	20000	131
Magnetoelastic NML	12000	0.01
CMOS LOP 21nm	1000	0.72

acid it is necessary to wait that all feedback signals propagate through the circuit to avoid data conflicts. It appears that, due to this long latency, the throughput of NML version is greatly reduced. As a solution to maximize throughput we adopt parallel computing and data interleaving. As a consequence, both in case of CMOS and NML a new amino acid can be sent every clock cycle and the only difference in speed is due to the clock frequency. The NML implementation is therefore with this improvement 4 times slower than CMOS implementation. However, the lack of speed is compensated by a much smaller power consumption.

Table I shows the comparison in terms of power consumption and circuit area of a single processing element, between CMOS and the main implementation of NML logic. Data for CMOS are obtained collecting the results from Synopsys synthesis on an industrial 45 nm technology and combining them with the ITRS Roadmap predictions to extrapolate the 21nm equivalent performance. For NML data are accurately estimated starting from the circuit layout and technological data. Most of the Smith-Waterman main blocks, like the adder shown in Figure 4.B, are accurately designed, so it is possible to know exactly their area and their composition. The total circuit area and the number of magnets of the processing element can be estimated starting from the main blocks and using multiplicative constants to keep into account interconnections overhead [12]. Once the total circuit area and the total number of magnets is known it is possible to estimate the circuit power consumption, because it is directly related to circuit area. The total estimated number of magnets for a processing element is 470000. Data in Table I show that in case of magnetic field based NML the area is around 21000 μm^2 , slightly lower (20000 μm^2) in case of STT-current based NML and remarkably smaller in case of Magnetoelastic NML (12000 μm^2). While the circuit structure and the number of magnets are the same in every case, the size of magnets is different, leading to different values for the area. The area however is much bigger than CMOS case, which is around 1000 μm^2 . The reason behind this is the availability in CMOS technology of multiple layers for interconnections. In NML, according to the current level of technology maturity, circuits are built using only one layer.

Comparing instead power consumption of one processing element (Table I), for magnetic field based NML it is 2 mW while for STT-current based NML it is much higher, about 131 mW. STT-current based NML are suited for circuit with a limited number of magnets, lower than 11000 [9]. Unfortunately both values are higher than the value obtained in CMOS, which is around 0.7 mW, so they cannot be used for low power application, but only if a high heat or radiations reliability is required. Using instead the magnetoelastic clock solution, it is possible to obtain a considerable reduction in power over CMOS, since the total power consumption is around 0.01 mW. Clearly this is the best solution for NML logic which allows to obtain a remarkable reduction of power

consumption with only a relatively limited reduction of speed.

V. CONCLUSIONS

NML logic enables the fabrication of circuits assuring a considerable reduction in power dissipation with respect to CMOS, at the cost of a relatively reduced speed. The intrinsic pipelined behavior of this technology leads to a consistent reduction of circuit throughput in presence of feedback signals if not specific countermeasures are taken. Using data interleaving, therefore running multiple operations in parallel, is one of the possible solutions. Since the number of operations that must run in parallel to obtain the maximum throughput can be high, not all applications are suitable. It is then clear that massively parallel applications represent the future for NML (and QCA) circuits, because they permit the maximum throughput. Biosequences analysis is one of these applications, used in this paper as a benchmark. This work helps quantifying the problem and the solution, and represents a considerable milestone for the ongoing studies on NML technology, pointing the path that should be followed in NML circuits development. We are currently exploring to what extent the design can be further optimized, moving from a simple implementation to an improved internal PE structure. We are redesigning circuit components in order to obtain a more compact layout, minimizing long interconnections and latency.

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