Comprehensive numerical model for Phase-Change Memory simulations

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Abstract - A “multi-physics” numerical model that couples a detailed electrical description with the heat equation and with a microscopic picture of the phase transition dynamics is presented. This model, implemented in a three dimensional semiconductor device solver, correctly reproduces the electronic switching effect in the amorphous chalcogenide alloy and the phase-transitions dynamics, providing steady-state and transient simulation results in good agreement with experimental data.

I. INTRODUCTION

Flash memories are nowadays the most important product in the non-volatile memory market, covering most of the data and code storage applications. However, intrinsic physical limitations must be faced to scale down the floating gate memory devices below the 45 nm technological node, mainly due to stringent requirements for the tunnel oxide leakage and the electrostatic control of the channel. The most promising alternative to floating gate memories for scaling capability and performance is the Phase Change Memory (PCM) technology. The prototype cell structure is constituted by a chalcogenide layer connected to a metallic pillar, called heater, that forces a high current density in the device active area (Fig.1). The storage element is a simple variable resistor of chalcogenide material, and the addressing of a bit inside the memory matrix is achieved by a selection transistor (MOS or BJT). The PCM technology exploits the peculiar features of the chalcogenide alloy Ge2Sb2Te5 (GST) to switch, under proper electrical pulses, between a highly conductive state, related to the crystal fcc phase (c-GST), and a lowly conductive state, related to the amorphous phase (a-GST). In Fig.2 the electrical characteristics of both phases are reported. The c-GST behaves as a resistor showing a moderate super-linearity, limiting the high field resistance to the heater one. On the other, a-GST shows an S-shape negative differential conductance (SNDC), collapsing from a high resistive OFF-state into a high conductive ON-state through the so-called electronic switching effect [1]. This is a fundamental mechanism for PCM devices, allowing a high current density to flow in the device active area, without reaching irreversible breakdown. The resulting significant Joule heating is then exploited to reach the phase transition temperatures responsible for the memory effect. Amorphization occurs when an electrical pulse is high enough to reach the melting temperature (680 °C) and its falling edge is quick enough to quench the molten GST (cooling within a few tens of ns). On the other hand crystallization takes place at lower temperatures, with a thermally-assisted kinetics (in the range of some seconds at 200 °C and some tens of ns at about 500 °C). PCM functionality is thus the result of the simultaneous action of physical effects belonging to different realms, namely electronic switching, electro-thermal heating, and phase-change transitions. Aim of this paper is to discuss modeling issues and numerical challenges that must be faced to fully account for the PCM device physics. The main features of a comprehensive numerical model will be discussed, showing that a detailed description of the electrical properties is mandatory to correctly reproduce the cell behavior in its operating conditions.

![Fig. 1. Schematic cross section of a PCM prototype cell. The active region is adjacent to the GST-heater interface.](image1)

![Fig. 2. Experimental electrical characteristics for a PCM cell in both amorphous and crystalline phases.](image2)
II. ELECTRICAL MODEL

Due to its long range atomic order, the electron dynamics in the c-GST can be treated according to the Bloch’s theorem as in a crystalline semiconductor [1]. On the other hand, a-GST does not show long range periodicity. However, investigations on amorphous compounds demonstrated the existence of localized states with variable transport properties [2]. It became thus a common practice in the '60 to describe these materials as in Fig. 3-left [2], where “mobility edges” separate fully conductive bands from low mobility states. However, by assuming that low mobility localized states behave like trapping centers and that more conductive levels resemble delocalized states, a band structure can be employed to model the a-GST as a “very defective” crystalline semiconductor [1] (Fig.3-right). Moreover, amorphous GST presents other peculiar electrical transport properties, as shown by the low-field I-V curve of the OFF-state reported in Fig.4. Till about 200 mV, a linear behavior is found, while for higher voltages the current increases exponentially up to threshold switching. The exponential relationship between current and applied voltage is probably due to a carrier generation mechanism such as impact ionization, Poole-Frenkel or tunneling from contacts. To explain both the low-field behavior and the threshold switching effect, a field-assisted generation mechanism sustained by the free carrier concentration (e.g., avalanche ionization) must be invoked [5]. The competitive interaction between this generation mechanism and a strong Shockley-Hall-Read (SHR) recombination caused by the large defect density is responsible for the SNDC behavior in Fig.2. Based on the similarities between the properties of the a-GST and standard semiconductors, we have thus achieved a description of the a-GST fully compatible with models already employed for crystals, suitable to be implemented in a standard device simulator [3]. The accurate tailoring on experimental data of the material parameters, such as energy gap, carrier mobilities and density of states has been extensively discussed in our previous work [1]. The calibrated semiconductor-like model is able to provide a good agreement with measurements, as shown, for instance, in Fig. 4. Moreover, the SNDC curve reported in Fig.2 can be explained through a physical insight into the dynamics of the electronic switching mechanism (Fig.5). For low applied fields, a-GST quasi Fermi levels are close to their equilibrium position (Fig.5-top), resulting into an ohmic behavior. With this bias the generation and recombination rates are low and can be safely neglected. Increasing the voltage, the generation significantly increases due to the applied field, thus requiring to be balanced by a much higher recombination. The quasi Fermi levels thus split and move close to the de-localized bands (Fig.5 middle). Since the recombination mechanism depends on the unfilled defects density, at high applied field recombination is strongly reduced (all defects filled), and it can only partially balance the generation rate. The system thus reacts reducing the voltage drop to maintain the balance between recombination and generation, leading to the electronic switching. Generation is now sustained by a large density of free carriers.
carriers, with the electron quasi Fermi level close to the conduction band edge (Fig.5 bottom). Fig.6 shows that the numerical model correctly describes the pulsed operation mode of PCM cells. The agreement was achieved by a mixed-mode simulation, driving the PCM-cell with a load resistor of 4 kΩ. A capacitor of 25 pF was added to the Spice net between the cell top electrode and ground, describing the role of the probe tip capacitance, as depicted in the inset of Fig. 6. The capacitance causes a delay in the switching event of about 70 ns. The voltage drop on GST during the pulse plateau is instead determined by the partition between the load resistance and the ON-state amorphous resistance of about 1 kΩ.

III. THERMAL AND PHASE TRANSITION MODEL

To achieve a full description of the PCM programming functionality, the heat conduction equation has been self-consistently coupled with the electrical model to describe self-heating (Joule effect). Thermal maps can thus be obtained performing transient simulations on an initially crystalline device for several current values, as reported in Fig.7. In the first frame the melting condition may be recognized, corresponding to the existence of a thin melted layer of GST close to the interface with the metallic pillar. The transition from the melted phase to the amorphous one is determined by the quenching time. In fact, during the falling edge, the GST cools down passing through the crystallization temperature region, thus allowing nucleation and growth to take place with their own kinetics. If the falling edge is very short (10-15 ns), crystallization is practically avoided, so that the hot region becomes a-GST, resulting in a low conductive state. Further increase in the programming current causes the thermal spot to extend and the amorphous zone is correspondingly enlarged, still maintaining a hemispherical shape. To account for the phase transitions responsible for the memory effect, a local phase-change algorithm has been coupled to the electro-thermal equations. The crystallization takes place by the action of nucleation and growth. The first phenomenon refers to the appearance of nuclei of the new phase in a homogeneous matrix of the old phase, while the second one refers to the growth of a pre-existing interface between the new and the old phases. In the low temperature range, due to the increase of the atomic mobility with temperature inside the melted GST, both nucleation and growth are enhanced with temperature (Fig. 8). At each temperature, a critical dimension exists for which a cluster of coordinated material is stable and thus becomes a nucleus. The stability of a cluster comes from the balance between the negative free volume energy contribution related to the phase transition (crystal is the stable phase), and the positive work to build the surface of crystal cluster. This critical dimension increases with temperature, and, close to the melting condition, practically no stable nuclei may exist (Fig.8), resulting in a growth driven crystallization. Starting from the model originally proposed in [4], a Monte Carlo algorithm for both nucleation and growth has been implemented and calibrated on experimental data. The grid elements dimension was chosen to be comparable with the critical cluster size at the temperature for which nucleation probability shows its maximum. The probability densities for unit of time are reported in Fig.8.

IV. RESULTS

To couple the phase change transition and the electro-thermal model, an external loop was implemented as
depicted in the block diagram of Fig.9. At each time step, the electro-thermal equations are self-consistently solved and a thermal map is calculated. Note in the inset in Fig.9 that the electrical model qualitatively reproduces the current filament formation effects observed during threshold switching [5]. At each time step of the external loop, nucleation and growth probabilities are evaluated to calculate the evolution of the phase distribution. Adaptive controls and damping techniques on the temperature increase and transformed volume have been implemented to insure convergence and accuracy. Note that the final phase distribution results from the full coupling among the electro-thermal and phase change models, thus promoting the phase transitions in the hottest regions, i.e., in the current filaments. In Fig.10 low-field resistances are plotted as a function of the programming current. The curve is obtained starting from an amorphous bit and applying increasing programming pulses with a 150 ns width and a 10 ns falling edge. For I<100 µA, the cell resistance remains close to the amorphous one. Higher current densities promote the GST crystallization process, thus causing resistance values to drop down [6]. Further increases in the programming current results in the melting condition (400 °C), while at about 600 µA the amorphous volume saturates. The same figure reports the simulated R-I curve, resulting in good agreement with experimental data. With this approach it has been possible to investigate and understand the phase distribution in the intermediate states. The amorphous to crystal transition is a localized process following some preferential paths, while the nature of crystal to amorphous one is practically isotropic, thus proceeding with hemispherical shape [7].

To check the crystallization model, the data retention times have been calculated at several temperatures and compared with experimental data in the Arrhenius plot depicted in Fig. 11. Each failure time is defined as the time for which the resistance reaches a reference value (ten times the minimum set resistance), under constant temperature annealing condition. Also in this case, a good agreement with experiments has been achieved, and both experimental and simulation results show that the data retention can guarantee at least 10 years at 110 °C. For a complete transient phase change computation, with a mesh of about 7000 nodes, 22 hours are required, working on a single Intel Xeon Processor with a CPU clock frequency of 2.8GHz.

V. CONCLUSIONS

For the first time a complete model for PCMs is presented, coupling an innovative electrical model, able to reproduce the threshold switching, with a local nucleation and growth algorithm for the phase transition. The algorithm has been implemented in a semiconductor-like device simulator, providing a comprehensive tool, suitable for PCM cell design and optimization.

REFERENCES