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Study and Evaluation of Electronic transport property for an InAlN based on Monte Carlo

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ARTICLE INFO ABSTRACT

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The emergence of the semiconductors III-N with heterojunction structures has made it possible to study a wide range of two-dimensional phenomena. This paper devotes to simulate the characteristics of the InAlN material, taking into ac-count temperature and doping as dependencies of conduction properties and performance using MOCASIM of the Tcad-Silvaco software. For the electronic transport model analyzing, we adopted most of the predominant mechanisms using various scattering effects including: optical phonon scattering, acoustic phonon scattering through deformation potential and piezoelectric potential, ionized impurity scattering, and grain boundary scattering. As expected, the carrier transports in the GaN layer are affected by the spontaneous polarization of the InAlN layer. To interact that, the diffusion of grain boundaries has been switched from the diffusion of ionized impurities by the deposition of InAlN. In order to achieve the most improvement possible for the electron transferring in terms of thickness and alloy composition related to the improvement of superdeposited layers. The confinement of sub-bands in channel quantum well is also taken into account in the computation of electron mobility. In the end, the adopted electron model is improved by including the effects of deep electron traps.

1. Introduction

 Many new transistors based on heterostructures are composed of thin-layer ternary alloys materials epitaxial deposited on a binary substrate. Their operation is based on the existence of heterojunction semiconductor/semiconductor wide gap or binary semiconductor III-V, as aluminium nitride (AlN), gallium nitride (GaN), indium nitride (InN) and their ternary and quaternary alloys have very attractive properties for microwave applications [1]. Thanks to advances in epitaxial techniques, have been included in the fabrication of active components [2]. The characteristic ability of these structures results to control the flow and distribution of electrons and holes across the shifts bands [3, 4]. The transport phenomena in

semiconductor lead to the behaviour of electrons and the energy gap in the conduction band. An analytical study of the problem requires the knowledge of the function of electron energy distri-bution, which is obtained by solving Boltzmann's equation [5]. The latter, partial derivative admits analytical solutions in a limited number of cases. To simulate the electrical transport phenomenon in semiconductors [6, 7], numerical methods are most frequently applied and the Monte Carlo of the written language MocaSim Tcad-Silvaco and the C-Interpreter software as well [8].

This language is not necessarily the best scientific programming, but it is certainly the most widespread and

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under existing programs and programs related to this method.

In this paper, we try to adapt this method to the study of ternary semiconductors [9]. Thus, we precisely apply it to the ternary $In_XAl_{1-X}N$ layer which deposits on GaN layer [10]. We simulate the various interactions, drift velocities transient in all the valleys, and the average drift velocities.

When indium is incorporated into AlN, the variation of the lattice parameter "a" of the compound can be resulted. As a first approximation, this parameter varies linearly with the indium levels. In order to achieve a good lattice matching between InXAl1-XN and GaN, the "x" value must be equal to 0.18, it must incorporate 18% of indium. We then calculate for the alloy $In_{0.18}Al_{0.82}N$ [11]. In addition, for a band gap, it covers a very wide energy range of 0.7 eV (InN) and 6.2 eV (AlN), which gives it a network matching function with 3.4 eV (GaN) [12, 13]. What makes us keep to this choice is the marked resistance to thermal deterioration. This layer is often maintained coherently at hundreds of degrees above the growth temperature until it reaches a temperature of 1100 $^{\circ}$ C [14], and at least it is recommended to prove the stability of the corresponding grid at no less than 960 °C [15, 16].

In [17, 18, 19], the authors use the program "modelling materials at the atomic scale from first principals initio VASP" in the analysis and approximation of the generalized gradient reported of an InAlN material cubic structure.

Vegard's rule has a linear dependence in composition, with very small deviations that both lattice parameters "a" and "c" essentially exhibit. The research completed as described in the HRXRD and Raman data concluded that even these small deviations of the lattice constant can have an impact on stress computations, so they suggested the following equation for "a" and "c" [20]:

$$
\xi(x) = x\xi_{InN} + (1 - x)\xi_{AlN} - \delta_{\xi}x(1 - x) \text{ and } \xi = a, c \tag{1}
$$

Where $\delta_a = 0.0412 \pm 0.0039$ Å and $\delta_c = -0.060 \pm 0.010$ Å are the corresponding deviation values respectively for the a-cell and c-cell parameters.

2. Results and Discussion

In this paper, the results of research are explained and at the same time the full dis-cussions are given. The results can be presented in the form of figures, graphs, tables and others which allow the reader to understand easily [6, 14]. The discussion can be done in several sub-chapters.

2.1 Non-equivalent interaction intervals

Fig. 1 illustrates all possible transitions of valleys: L-Γ, Γ-X, L-X, X-L, L-X and Γ-Γ (absorption and emission). Interactions between valleys can only occur when the energy carriers are large enough. They are very unlikely to weak field. When an elec-tron is transferred to another valley, the direct effects of interaction itself are added all the discontinuities due to non-equivalence of origin and arrival valleys.

MocaSim

Fig. 1. Non-equivalent interaction intervals of an InAlN material: (a) Interaction intervals absorption and (b) Interaction intervals emission.

In particular, as the kinetic energy of electron is measured from the minimum val-ley which occupies, the transfer to a non-equivalent valley leads to the significant variation of kinetic energy corresponding to the difference in levels between the two valleys [21].

Energy phonons are emitted or absorbed between 8 meV and 42.7 meV, well be-low the initial energy of the carriers at room temperature. As a result, we draw two conclusions: the probability of inter-valley transition $Γ-X$ (removals as emissions) is the most important, and the probability of emissions is greater than removals.

2.2 Scattering Rate

Fig. 2 displays all possible scattering rates for the following valleys: Γ, L and X. One can clearly see that the scattering rate increases with the square root of the energy. The acoustical and optical phonons can also scatter an electron from one valley to another valley. Moreover, the influence of the estimated scattering rate on the total surface simulated the mobility by taking into account the optical pho-non, deformation potential, piezoelectric dislocation scattering and threading with different dislocation densities.

Fig. 2. Inter-valley scattering rate of an InAlN material.

The computation of isotope-induced phonon scattering time is selected by a typi-cal technique for alloy scattering [22, 23] and the Fang Howard wave function [24]. It is inferred that the scattering mechanisms are not necessarily identical, such that many defects add ionized impurity scattering, atomic defects in lattice and long-range scattering in ternary materials.

Generally tend to use the GaN layer NID at n-type doping, this layer has weak n-type conductivity, representing the presence of unintended background donors [25].

Transports widening the heterojunctions between two ntype layers have the same density of impurities, generally the widening of the level is more than large, which makes it difficult to distinguish the impact of each mechanism.

2.3 Effect of temperature on the interaction of mobility and velocity for InAlN

The over-speeding phenomena at high electric fields are due to the smaller relaxation time of the impulse compared to the relaxation time of the energy. In the simulation, cold electrons are injected into the material and accelerated very quickly by the high electric field. Then, the scattering rates increase with energy to suppress the over-speed. The

increased scattering rates take some time before it has an effect on the drift velocity. The time delay creates the overspeed effect.

Commonly, the over-speeding variation versus temperature results to the increased temperature gaining electrons of kinetic energy as they rise and collide with other atoms in their energy transfer.

Speed keeps almost the same pace. One can clearly notice that the higher speeds of electron are achieved with a low temperatures, and vice-versa as shown in Fig. 3. As example, when the temperature is equal to a low value 200 K, the electron speed is worth to $2*10-7$ cm/s.
Mocasim

Caracteristic of Electron Velocity for x=0.18 21000000 18000000 15000000 **Electron Velocity (cm/s)** 12000000 9000000 6000000 3000000 $\mathbf{0}$ 40000 80000 120000 160000 200000 $\bf{0}$ Electric field (V/cm) a) MocaSin Caracteristic of Electron Mobility for x=0.18 4000 3500 Electron Mobility (cm^2/V s) 3000 2500 2000 1500 1000 500 $\bf{0}$ θ 40000 80000 120000 160000 200000 Electric field (V/cm) b)

Fig. 3. Mobility and velocity of electrons in the InAlN material relative to the electric field at different temperature values: (a) Electron velocity and (b) Electron mobility.

As temperatures rise, the increased effect of thermal agitation suffered by the at-oms results in a greater possibility of impacts between them. This will lead reducing the free electron rate to 0.25 cm²/Vs as shown in Fig. 3.

2.4 Effect of doping on the interaction of mobility and velocity for InAlN

The variation of velocity for different values doping is taking the same form for all the selected concentrations

(for this reason, we limited to the range concentrations only).

According to Fig. 4, one can clearly observe that the velocity values generally decrease as the concentration increases relative to the electric field. Therefore, the increase in doping can lead to increase the probability of collision between electrons, and this reduces the speed of the electrons. One can also notices, a decrease in peak speed and a minor increase of threshold field.

MocaSim

Caracteristic of Electron Velocity for x=0.18

Fig. 4. Mobility and velocity of electrons in the InAlN material relative to the electric field at different doping percentages: (a) Electron velocity and (b) Electron mobility.

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Doping has significant effects on weak fields, but the curves tend to merge when-ever the field is very strong. Indeed, when the electric field reaches very high values, the carriers are heated and have less influence on the ionized centres.

Regarding to the saturation velocity, it only appears when doping is weak, which reflects the low dependence of collision mechanisms with large networks on doping, as shown in Fig. 4.

3. Conclusion

This paper has presented the simulations for various interactions and electrical transport in the ternary material $In_{0.18}Al_{0.82}N$ using the Mocasim Tcad-Silvaco soft-ware. It has been shown that the following points can be achieved: (1) The piezoelec-tric interactions are negligible in valleys Γ, X and L, (2) The acoustic interactions are also negligible in the three previous valleys, (3) The inelastic interactions of emission (intervals intra-valleys and optical polar) are more important than those removals, (4) In central valley, the peak over-speed appearing from the critical electric field, they are larger and occur more rapidly when the level of the electric field increases, (5) In the L and X valleys, the speed is lower than in the Γ valley and it can reach the sta-tionary state at about 2 ps. Finally, the transport characteristic can be applied in the nanostructure and electric power, such as: HEMTs heterojunction technology.

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