ACOUSTOELECTRIC STUDY OF INTERFACE TRAPPING DEFECTS IN GaAs EPITAXIAL STRUCTURES

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Abstract

A new acousto-electrical method making use of transient transverse acoustoelectric voltage (TAV) to study solid state structures is reported. This voltage arises after a surface acoustic wave (SAW) generating the signal is switched off. Related measurements consist in detecting the shape of transient voltage and its spectral and temperature dependence. Both theory and experiment show that this method is an effective tool to characterize trapping centers in the bulk as well as at surfaces or interfaces of epitaxial semiconductor structures.

Key words: acousto-electric, trapping center, epitaxial structure.
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1 INTRODUCTION

Different methods are known to study crystal defects and deep levels in semiconductors in solid state physics. Most of them, including modern Scanning Tunneling Microscopy techniques, allow to characterize surface defects. At the same time multilayer and epitaxial semiconductor structures are strongly influenced by defects [1-3] which can be located at the interfaces between epilayers and substrate. Up to now no appropriate experimental methods exist to characterize these interface defects. Recently some publications reported the transient acoustoelectric effect in semiconductors with defects [4-6]. Transient transverse acoustoelectric voltage (TAV) generated in semiconductors has already been used in the past [7,8], but this using hasn’t be aimed to interface defect characterization and hasn’t include corresponding acousto-optical measurements.

If the TAV method is applying some difficulties of interpretation of the experimental results have to be surmounted. The point is that the TAV amplitude strongly depends on several parameters, among them the influence of piezoelectric field strength $E_v$ on local centers has to be mentioned. To avoid these difficulties we propose to measure the transient TAV signal just after the excitation pulse has be switched off.
2 THEORY

2.1 MECHANISMS OF THE TAV ORIGIN

TAV is one of the manifestations of ultrasonically activated redistribution of the electrical charges in semiconductors. It is generated across layered systems consisting of piezoelectric and semiconducting materials due to a piezoactive surface acoustic wave (SAW) propagating along the piezoelectric. The SAW piezoelectric field penetrates inside the semiconductor causing charge carriers redistribution in the near surface region.

Two main mechanisms of generating the TAV effect exist in semiconductors. The first one, so-called "concentration-effect" is due to the variable component of sample conductivity $\sigma_v[9]$. It is a result of electron concentration redistribution under the action of the variable piezoelectric field $E_v$ of SAW. Hence, the direct component of acoustoelectric current is defined by:

$$j_0 = \langle \sigma_v E_v \rangle_{T_s}$$

where the averaging is carried out on acoustic wave period $T_s$. The relaxation time of this TAV signal component is defined by the Maxwell’s relaxation time $\tau_m = \epsilon \epsilon_0 / \sigma_0$. Usually $\tau_m$ is much less than the period length of SAW. Therefore one can suppose that the relaxation time of the "concentration" TAV is equals to zero.

The second mechanism of the TAV generation is connected with semiconductor defects. The presence of the high-frequency SAW electrical field in the near surface region results an increasing of the free charge carriers concentration. This, in turn, causes an increasing of the electrical charge captured on deep trapping levels, which are located either at interface or surface. As a result a direct electrical field perpendicularly to the sample surface is occurring. The amplitude of this "trap" TAV component is proportional to the excess concentration of the charge captured on trap levels [10].

Let us assume that the concentration of this charge is $\Delta n(t)$. For the simplest case of only one type of surface defects existing the transient TAV signal can be written as

$$V_{ae}(t) = C_{ae} \Delta n(t)$$

where the coefficient $C_{ae}$ depends on sample parameters.

The voltage $V_{ae}(t)$ is connected to the relaxation of the electrical nonequilibrium charge trapped by surface defects. As far as the relaxation time does not depend on the particular mechanism of excitation of the electronic subsystem, it is possible to use standard theoretical approach. The relaxation time $\tau$ is a function of trap level parameters. In the following we will consider an n-type semiconductor. Then it holds

$$\tau = \frac{1}{N_e V_T S_n} e^{E_t / kT}$$
where $N_c$ is the density of states of the conductivity band, $V_T$ is a thermal velocity of free electrons, $k$ is Boltzmann's constant, $T$ is temperature, $E_t$ is an energy depth of the electronic trap level counted from the edge of conductivity band and $S_n$ is an effective cross section of electron capture by trap centers.

Usually different types of trap levels exist. In this case the excess concentration $\Delta n_{ti}$ of charge carriers captured by i-type surface traps, can be written as:

$$\frac{d\Delta n_{ti}}{dt} = -\frac{\Delta n_{ti}}{\tau_i} + F_i(t) \quad (4)$$

where $\tau_i$ represents the characteristic relaxation time of i-type level, $F_i(t)$ is an external force, which initiates the capturing by surface traps. The physical nature of said force is connected to the piezoelectric field of SAW.

If the ultrasonic wave amplitude is modulated by rectangular pulses the external force $F(t)$ has the form

$$F(t) = \begin{cases} 
\text{const}, & 0 < t < T_p \\
0, & T_p < t 
\end{cases} \quad (5)$$

where $T_p$ is the pulse duration. The radio frequency impulses and corresponding TAV pulse are presented on Fig.1. The TAV pulse shows a monotonous increase ("AB") and a monotonous decrease ("BC"). This shape of the TAV signal is typical for the trap component. The measurement of the shape of the transient TAV signal is made along "BC" in Fig.1. Note that in this case the external force vanishes ($F(t)=0$) during our measurements. In this case the solution of the equ.4 is a decreasing exponential curve with relaxation time $\tau_i$. For different trap levels in the sample the TAV signal can be expressed by a sum

$$V_{ae}(t) = V_C + \sum_{i=1}^{N} V_i e^{-t/\tau_i} \quad (6)$$

where the times $\tau_i$ correspond to various i-levels. The summation has to be carried out over all existing traps in the near surface region. The coefficients $V_i$ are proportional to the concentrations of the various trap types. The first term in equ.6 presents the contribution in TAV of "very slow" levels. Such levels can exist in $A_2B_6$-compounds or in MIS-structures with relaxation times $\tau$ of about some hours while modern $A_3B_5$ epilayer structures do not have such slow levels. The signs of $V_C$ and $V_i$ are opposite for electron and hole capture centers.

Thus, it is possible to determine the relaxation time $\tau$ by measuring the TAV signal. The transient TAV relaxation time $\tau$ is determined by the rate of thermal emission and trapping of charge carriers on surface levels. In turn, the rate of reaching the thermodynamic balance between capture centers and conduction zones in a semiconductor structure depends on characteristic parameters of these centers, such as energy depth $E_t$ and effective cross section $S_n$. The calculation of $S_n$ by means of equ.3 is successful only if the energy levels $E_{ti}$ is known. For this reason additional information about $E_{ti}$ and relaxation time $\tau_i$ is needed. It can be obtained by measuring the optical spectra of TAV signals or the thermal dependence of TAV signal.
2.2 ANALYSIS OF TAV SPECTRAL DEPENDENCIES

The illumination of a semiconductor surface changes the distribution of free and captured charge carriers. Now we consider the influence of monochromatic illumination on various components of the TAV signal. For singly charged electron traps the process of electron capturing follows the equation

$$\frac{dn_t}{dt} = C_n n_s (N_t - n_t) - \beta_n n_{s0} n_t$$

(7)

where $N_t$ is the concentration of the surface trap levels, $n_t$ is the concentration of captured electrons; $n_s$ is the concentration of the free charge carriers near the semiconductor surface and $C_n$ and $\beta_n$ are the probabilities for capturing and releasing of charge carriers. The concentrations $n_t$ and $n_s$ deviate from their equilibrium values $n_{t0}$ and $n_{s0}$ under the influence of ultrasonic waves:

$$n_t = n_{t0} + \Delta n_t, \quad n_s = n_{s0} + \Delta n_s$$

(8)

The condition of thermodynamic balance in the absence of the acoustic wave is

$$C_n n_{s0} (N_t - n_{t0}) - \beta_n n_{s0} n_{t0} = 0$$

(9)

This equation provides the connection between the coefficients $C_n$ and $\beta_n$. For a non-degenerate semiconductor we obtain:

$$\beta_n = C_n e^{((E_t - F_{s0})/kT)}$$

(10)

where $F_{s0}$ is the Fermi level at the surface. By substituting equs. 8 and 10 into equ.7, we obtain:

$$\frac{d\Delta n_t(\lambda)}{dt} = C_n [(N_t - n_{t0}(\lambda))\Delta n_s - (n_{s0}(\lambda) + \Delta n_{s0}(\lambda) + N_n e^{-E_t/kT})\Delta n_t(\lambda)]$$

(11)

Note, that $\Delta n_t$ and $\Delta n_s$ are averaged over the acoustic wave period. Acoustic-electrical voltage component, which is connected to trapped charge, is proportional to the excess charge on said traps ($V_{ae} \sim \Delta n_t$). Performing the measurement of the TAV signal and its shape under quasi-equilibrium conditions, it means:

$$\frac{d\Delta n_t}{dt} = 0$$

(12)

Then, from equ.11 we receive the following expression for the trap TAV component:

$$V_{ae} \sim \Delta n_t(\lambda) = (N_t - n_{t0}(\lambda))\frac{\Delta n_s(\lambda)}{(n_{s0}(\lambda) + \Delta n_{s0}(\lambda) + N_n e^{-E_t/kT})}$$

(13)

The values $n_{t0}$, $\Delta n_s$ and $\Delta n_{s0}$ depend on sample illumination. However, if the photon energy $h\nu$ does not exceed the band gap energy $E_G$ and direct electronic excitation from valence band to conduction band does not occur, the changes of concentrations $n_{s0}$ and $\Delta n_{s0}$ are rather small. In this situation the TAV spectra
is mainly determined by the first co-multiplier \((N_t - n_{t0}(\lambda))\) in equ. 13. At sample illumination with light of photon energies exceeding the value determined by equ.14:

\[ E_t = E_G - h\nu \]  

(14)

the charge carriers transferring from the valence band to capture centers is resulting in a growth of \(n_{t0}\). Then the difference \((N_t - n_{t0}(\lambda))\) in equ.13 decreases, and a minimum should be observed in the TAV spectrum. If several types of trap levels take part in TAV signal formation, then several corresponding minima should be observed in the optical TAV spectrum.

### 2.3 ANALYSIS OF THE RELAXATION TIME TEMPERATURE DEPENDENCY

The value of an energy position of the deep trap levels \(E_t\) can be obtained from temperature dependencies of the relaxation TAV time \(\tau\) also. This dependency can be obtained from equ.3. The temperature dependency of \(N_c\) and \(V_T\) for nondegenerate semiconductor is well-known: \(N_c \sim T^{3/2}\), \(V_T \sim T^{1/2}\). For attracting Coulomb center at room temperature \(S_n \sim T^{-2}\) [11]. Therefore:

\[ N_c(T)V_T(T)S_n(T) \sim const(T) \]  

(15)

If this relation isn’t absolute exact, the temperature dependence of expression (15) will much weaker than exponential dependence. Consequently, for deep levels \((E_t >> kT)\) and small temperature changes the temperature dependence of \((N_cV_TS_n)\) can be neglected in equ.(3). Having two times \(\tau_1\) and \(\tau_2\) for two different temperatures \(T_1\) and \(T_2\), we can to calculate \(E_t\) by equ.(16):

\[ E_t = \frac{kT_2\ln(\tau_1(T_1)/\tau_2(T_2))}{(T_2/T_1) - 1} \]  

(16)

One can enhance a precision of \(E_t\) determination by using a plot of \(\ln(\tau)\) on \(T^{-1}\), if the experiments are done in some temperature range:

\[ \ln(\tau(T)) = (E_t/kT) + const(T) \]  

(17)

The curves of these dependencies are direct lines. The angles of declination of these plots are set by value \(E_t\). Thus it is possible to define a value \(E_t\) for each trap level.

### 3 SAMPLES AND EXPERIMENTAL TECHNIQUE

We investigated three types of GaAs-samples. The first one labeled as GA-1 is the structures of epi-layer n-GaAs on n-GaAs substrate. They were fabricated by an industrial vapor phase epitaxy method in the system \(Ga - AsCl_3 - H_2\). The substrate, 0.35 mm thick, was doped by Te with \(N_{Te} \approx (1 \div 2) \times 10^{18} cm^{-3}\). The free carrier concentration of the epi-layers, doped by Te too, 6-9 microns thick for
different samples, was \((0.6 \div 1.2) \times 10^{15} cm^{-3}\). The second one labeled as GA-2 consists of a 400\(\mu m\) thick GaAs-substrate with a 1\(\mu m\) thick epitaxial layer of n-GaAs on it. The third sample labeled as GA-3 consists of the GaAs substrate and a 8\(\mu m\) thick epitaxial layer with electron concentration \(n \sim 10^{14} cm^{-3}\). GA-2 and GA-3 samples were fabricated by MOCVD process.

The samples GA-1 and GA-2 were placed by the epitaxial layer on the lithium niobate plate. On this plate SAW of 6 MHz were excited. The TAV signal was picked up between a bottom ground electrode and a flat metal electrode attached on the upper surface of the GaAs sample as shown in Fig.2a. This arrangement is very convenient because no special sample preparation is required and sample replacement is easy to do. For example, it is not necessary to form electrical contacts to the samples. As far as GaAs is a piezoelectric material, it can simultaneously play the role of the piezoelectric waveguide as shown in Fig.2b (for GA-3). In this case SAW of 67 MHz are excited by means of an interdigital transducer deposited on the sample surface. The TAV signal was measured between the ground electrode and the aluminum film which has to be evaporated on the epitaxial layer.

4 EXPERIMENTAL RESULTS AND DISCUSSION

4.1 THE EXPERIMENTAL DEFINITION OF THE TRANSIENT TAV RELAXATION TIMES

The task of the subsequent mathematical procedure consists in finding out of the values \(V_i\), \(\tau_i\) and \(N\) from equ.6. To this aim it is necessary to determine the number \(N\) of effectively acting types of surface levels. Therefore we have to analyze the plots \(\ln(V_{ae})\) versus time. For \(N=1\) this dependence should be a straight line. Its slope gives the relaxation time \(\tau\). In the case of two or more exponential components this plot has a more complex form. The number of rectilinear sites on these curves corresponds to the number of trap centers. One can define characteristic relaxation times from the declination angles of these directs. In our experiments the values \(V_i\) and \(\tau_i\) were estimated by the interpolation of the transient TAV signal (part "BC" in Fig.1) with the help of a special computer program which had found out the number of different type levels simultaneously. The data obtained from GA-2 show the occurrence of two exponential terms with decay times of 2 and 12 ms. In Fig. 3 the parts "ab" and "bc" of the curve 1 correspond to two types of traps having the relaxation times 2 and 12 ms, respectively. The experiments with the GA-3 sample show two different terms in total TAV signal corresponding to the two parts "ab" and "bc" of the curve 2 shown in Fig. 3. We can distinguish two types of trapping centers having relaxation times of 4.5 and 22 ms at room temperature. Four local centers having relaxation times of 2.2, 1, 22 and 1.7 ms were also found in the samples GA-1 at room temperature.

Additional measurements were carried out for further identification these trap centers and for the definitions of their parameters. The relaxation time of a transient TAV was measured at various temperatures in samples GA-1 (see item 2.3). The
samples GA-2 and GA-3 were used to measure spectral dependencies of a TAV signal (see item 2.2).

4.2 TEMPERATURE DEPENDENCIES OF THE TRANSIENT TAV

The descending parts of the TAV signal for GA-1 series at various temperatures are presented on Fig.4. We can see, that the TAV relaxation time decreases with temperature increase. Our measurements were carried out in a temperature range 294 to 330 K.

The plots of $\tau$ on $T^{-1}$ are shown on Fig.5. Practically they are direct lines, angles of declination of these lines give the trap level energy positions $E_t$. Knowing the characteristic relaxation times of the excess charge $\tau_i$ and the energy levels of the interface centers $E_{ti}$, we can calculate the effective capture cross section $S_{ni}$ for these levels by equ.(3).

Thus, four deep levels were detected at the samples under study. They were designated as levels $E_{1,2,3,4}$. The characteristic parameters of said centers are presented in the Table. The characteristics coincidence of $E_4$ level and the literary data [12] allows to say that $E_4$ is electron center EL3. The configuration $As_iV_{Ga}$ is assigned with this level. Really, the excess of the interstitial arsenic ($As_i$) and the gallium vacancies ($V_{Ga}$) should be observed in the GaAs, doped by tellurium. Certainly, the tellurium interstitial ($Te_i$) can be significant at this case too. It is possible to conclude, that level $E_1$ is EL17 center [13-15], and $E_3$ - EL5 center [14]. Level $E_2$ can be identified as electron center EL6 [16]. All these levels deals with the vacancies of gallium ($V_{Ga}$) and arsenic ($V_{As}$) [13-16].

4.3 EXPERIMENTAL SPECTRA OF THE TRANSIENT TAV

Also, for determination of trapping level energy the optical spectra of transient TAV are investigated. A sample under investigation is illuminated with monochromatic light in the range from 0.6 to 1.6 eV. Then TAV signal is separated for partial exponential components with different relaxation times $\tau_i$ with the help of a special computer program a general. Each such component is characterized by a certain value of partial amplitude $V_i$ which varies during illumination. Said amplitudes $V_i$ determines a contribution to a total TAV signal of i-type traps having relaxation time $\tau_i$. This contribution reaches a minimum at illumination of sample surface by monochromatic light with photon energy equal $E_G - E_{ti}$, in accordance with the formula (13). Thus a spectral position of a minimum in a spectrum of i-partial amplitude $V_i(h\nu)$ undutiful determines an appropriate energetic level $E_{ti}$.

The spectra of partial amplitudes $V_i$ are shown on the Fig.6. The plots (1 and 2) are taken from GA-2 sample for relaxation times 12 and 2 ms, respectively. The minima on these curves correspond to two types of deep levels at $E_5^* - 0.48$ eV and $E_4^* = 0.54$ eV below a conduction zone. The plots 3 and 4 are taken from GA-3 sample for relaxation times 22 ms and 4.5 ms, respectively. The minima on plots 3 and 4
correspond to two types of deep levels at $E_5=0.48$ eV and $E_1^*=0.20$ eV below a conduction zone.

Now, knowing the energy positions of trapping centers and the related relaxation times, we can calculate the effective cross sections $S_n$ for capture of charge carriers by these trap centers. Below, characteristic defect parameters in the epitaxial GaAs structures under study are presented in Table. The defect identification was done by comparing the energetic position with literature data on deep levels in GaAs [12,13, 17- 20]. One can see, that deep trap levels of the types EL3 and EL17 are detected both with the help of spectral researches and measurements of TAV relaxation times’ temperature dependencies.

The difference in values of relaxation times for the levels $E_1$, $E_1^*$ and $E_5$, $E_5^*$ in different samples can be explained as follow. A relaxation time of captured charge is determined by two parameters - energy depth $E_t$ of trapping center and its capture cross-section $S_n$. Said $E_t$ depends on atomic structure of local defect and so does not vary from a sample to sample. Opposite, the value $S_n$ depends both on a type of trapping level and on electrical properties of semiconductor itself, including a configuration of electrical potential near defect and its shielding by a free charge. Therefore the characteristic relaxation times for the same trapping levels, but in various on manufacturing samples, can differ from each other.

5 CONCLUSIONS

The acoustoelectric method can be advantageously applied for studying interfaces in modern epilayer semiconductor structures. The experimental data followed by appropriate mathematical processing allow to estimate characteristic relaxation time, energy position in the forbidden zone and electron capture cross section for each of these centers. By this approach we, for the first time, measured the relaxation times of trap centers.

Investigating the interface between epitaxial GaAs-layers grown on substrate by MOCVD process and chlorine vapor method we found five different types of trapping defects. They were identified as EL3, EL4, EL5, EL6 and EL17 centers. Their characteristics at room temperature are summarized in the Table.

The different GaAs centers are known to be observed in the samples, fabricated by different technique [13]. The coincidence of the level’s parameters permit to conclude that levels EL5 and EL17 can be present in the epitaxial structures, fabricated by chlorine vapor method as well as by MOCVD technique.

The method described can also be applied to the investigation of other semiconductor structures, including those based on Si and $A_2B_6$ compounds.

Summarizing the obtained results we may conclude that the transient TAV technique is a reasonably simple and effective method for characterizing surface and interface trap centers.

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Table Captions

Table. Characteristics of interface trapping centers in epi-GaAs at room temperature.

| Samples | Level | Type | $E_t$, eV | $\tau$, ms | $S_n$, cm$^2$ |
|---------|-------|------|-----------|------------|---------------|
| GA-1    | $E_1$ | EL17 | 0.23 ± 0.02 | 2.2 ± 0.2  | $0.7 \times 10^{-18}$ |
| GA-3    | $E_1^*$ | EL17 | 0.20 ± 0.01 | 4.5 ± 0.4  | $1.5 \times 10^{-19}$ |
| GA-1    | $E_2$ | EL6  | 0.29 ± 0.02 | 1.0 ± 0.1  | $1.6 \times 10^{-17}$ |
| GA-1    | $E_3$ | EL5  | 0.42 ± 0.02 | 22 ± 2     | $1.4 \times 10^{-16}$ |
| GA-1    | $E_4$ | EL3  | 0.56 ± 0.02 | 1.7 ± 0.2  | $4 \times 10^{-13}$   |
| GA-2    | $E_4^*$ | EL3  | 0.54 ± 0.01 | 2 ± 0.03   | $2.5 \times 10^{-13}$ |
| GA-3    | $E_5$ | EL4  | 0.48 ± 0.01 | 22 ± 1     | $2 \times 10^{-15}$   |
| GA-2    | $E_5^*$ | EL4  | 0.48 ± 0.01 | 12 ± 1     | $4 \times 10^{-15}$   |
Figure Captions

Fig.1. Time dependency of input rf-voltage $V$ exciting SAW and resulting TAV signal.

Fig.2. Two versions of sample arrangements performing transient TAV measurements: a) separate medium configuration, b) integrated configuration

Fig.3. Logarithmic plot of transient TAV signal versus time. 1 is plot for GA-2 and 2 is plot for GA-3. Parts "ab", and "bc" correspond to different types of trapping centers.

Fig.4. Shape of the descending parts of the TAV signal at various temperatures for GA-1: 296 K (1), 305 K (2) and 318 K (3). Points - experimental data; curves - theoretical calculation by equ.(6)

Fig.5. Dependency of $\tau$ on $T^{-1}$ (GA-1). At room temperature the relaxation time $\tau$ is equal to 2.2 ms for the plot 1, 1.0 ms plot 2, 22 ms plot 3 and 1.7 ms plot 4. The slopes of plots 1, 2, 3 and 4 give four levels $E_1, E_2, E_3, E_4$, respectively.

Fig.6. Spectra of the TAV exponential component. 1- $\tau = 12$ ms (GA-1); 2- $\tau = 2$ ms (GA-2); 3- $\tau = 22$ ms (GA-3); 4- $\tau = 4.5$ ms (GA-2). The four minima labeled as $E_1^*, E_4^*, E_5, E_5^*$ correspond to the three trapping centers detected in the samples under study. The band gap energy is marked by $E_G$. 