Chapter

Interactions between Terrestrial Cosmic-Ray Neutrons and III–V Compound Semiconductors

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Abstract

This work explores by numerical simulation the impact of high-energy atmospheric neutrons and their interactions with III–V binary compound semiconductors. The efforts have focused on eight III–V semiconductors: GaAs, AlAs, InP, InAs, GaSb, InSb, GaN, and GaP. For each material, extensive Geant4 numerical simulations have been performed considering a bulk target exposed to a neutron source emulating the atmospheric neutron spectrum at terrestrial level. Results emphasize in detail the reaction rates per type of reaction (elastic, inelastic, nonelastic) and offer a classification of all the neutron-induced secondary products as a function of their atomic number, kinetic energy, initial stopping power, and range. Implications for single-event effects (SEEs) are analyzed and discussed, notably in terms of energy and charge deposited in the bulk material and in the first nanometers of particle range with respect to the critical charge for modern complementary metal oxide semiconductor (CMOS) technologies.

Keywords: radiation effects, III–V compound semiconductors, cosmic-rays, atmospheric neutrons, numerical simulation, neutron cross section, elastic scattering, inelastic scattering, nonelastic interactions, nuclear data library, Geant4

1. Introduction

The introduction of III–V high-mobility semiconductor materials into advanced CMOS manufacturing is currently envisaged as a possible “technological booster” for carrier mobility enhancement. Indeed, for nanometer-scale transistor, it will not be possible to maintain the power-performance tradeoff offered by conventional silicon-based technologies because of the transport properties of Si that are becoming a fundamental limitation for ultimate complementary metal oxide semiconductor (CMOS) [1, 2]. Several materials exhibiting higher carrier mobilities than Si are currently explored for both n-channel and p-channel transistors. Between all these materials, GaAs, InAs, InGaAs, and InP are suitable candidates for n-channel devices and GaSb and InGaSb for p-channel transistors [3–5].

In complement to their physical and electrical characterization, the exploration of the radiation response of these III–V materials is also an important concern for predicting their reliability. Several studies investigating total ionizing dose (TID) and heavy ion and laser responses of III–V MOS transistors in modern architectures, as well as the creation mechanisms of single-event effects, have
been reported in literature, but relatively little work concerns their response to terrestrial neutrons [6–13]. In two recent papers [14, 15], we begun such a systematic investigation of the susceptibility to natural radiation of group IV and III–V materials exposed.

The aim of this chapter is to provide new data and metrics for this exploration via Geant4 [16, 17] numerical simulations of the radiation response of III–V binary compound semiconductors subjected to high-energy atmospheric neutrons. The study specifically focuses on eight III–V materials: GaAs, AlAs, InP, InAs, GaSb, InSb, GaN and GaP. For each material, the reaction rates for elastic, inelastic, and nonelastic events have been determined as well as the classification of all produced secondaries as a function of their atomic number, kinetic energy, initial stopping power, and range in the target bulk. Average energy and charge deposited in the bulk material and in the first nanometers of particle ranges have been also evaluated. These later quantities are important statistical metrics in the occurrence of single-event effects (SEEs [18]); they have been compared with the critical charge for various CMOS technologies (180 nm to 14 nm nodes).

The chapter is organized as follows. Section 2 presents the different compound materials studied and summarizes their main physical and electronic properties. Section 3 details the Geant4 numerical simulations conducted in the framework of this study. Section 4 presents the complete analysis of interaction databases in which all neutron-induced secondaries have been recorded for each material subjected to atmospheric neutrons. Finally, in Section 5, implications for single-event effects in CMOS based on these semiconductor materials are analyzed.

2. Semiconductor properties

III–V binary compound materials explored in this work are semiconductor alloys containing one element from group III (boron column: Al, Ga, In) and one from group V (nitrogen column: N, As, Sb) of the periodic table. Table 1 shows the natural isotopic abundance for these six chemical elements. For example, natural gallium ($^31\text{Ga}$) consists of a mixture of two stable isotopes: $^{31}\text{Ga}-31$ at 60.10% and $^{31}\text{Ga}-71$ at 39.90%. Numerical simulations presented in the

| Symbol | Atomic number | Nuclide      | Natural abundance |
|--------|---------------|--------------|-------------------|
| Al     | 13            | 13-Al-27     | 100.00%           |
| Ga     | 31            | 31-Ga-69     | 60.10%            |
|        |               | 31-Ga-71     | 39.90%            |
| In     | 49            | 49-In-115    | 95.70%            |
|        |               | 49-In-113    | 4.30%             |
| N      | 7             | 7-N-14       | 99.60%            |
|        |               | 7-N-15       | 0.40%             |
| P      | 15            | 15-P-31      | 100.00%           |
| As     | 33            | 33-As-75     | 100.00%           |
| Sb     | 51            | 51-Sb-121    | 57.40%            |
|        |               | 51-Sb-123    | 42.60%            |

Table 1. Natural abundance of nuclides related to the III–V alloys studied in this work.
following take into account such isotopic compositions for each element present in the eight studied materials.

Table 2 summarizes the main experimental values for GaAs, AlAs, InP, InAs, GaSb, InSb, GaN, and GaP bulk materials in terms of energy bandgap $E_g$, number of atoms per volume unit, density [19], and electron-hole pair creation energy $E_{eh}$. All these crystalline solids are denser than Si (2.32 g/cm$^3$). The following materials, GaAs, AlAs, InP, GaN, and GaP, are characterized by a bandgap larger than that of Si and, consequently, by a larger electron-hole pair creation energy (3.6 eV for Si). The other materials, i.e., InAs, GaSb, and InSb, have a bandgap clearly below 1 eV; $E_{eh}$ values for these materials are also lower than that of Si. Finally, for GaSb and InSb, we did not find in literature an experimental value for $E_{eh}$; we used by default the values deduced from Klein’s model [20] establishing a linear relationship between $E_g$ and $E_{eh}$ in semiconductor materials.

### Table 2.
The main properties of the III–V binary semiconductors considered in this study.

| III–V compound | $E_g$ at 300K (eV) | Number of atoms per cm$^3$ ($\times10^{22}$) | Density (g/cm$^3$) | $E$-h pair creation energy $E_{eh}$ (eV) |
|----------------|------------------|----------------------------------------|-----------------|-------------------------------|
| GaAs           | 1.42             | 4.42                                   | 5.32            | 4.8                           |
| AlAs           | 2.16             | 4.42                                   | 3.76            | 6.8                           |
| InP            | 1.34             | 3.96                                   | 4.81            | 4.5                           |
| InAs           | 0.36             | 3.59                                   | 5.67            | 1.8                           |
| GaSb           | 0.73             | 3.53                                   | 5.61            | 2.7                           |
| InSb           | 0.17             | 2.94                                   | 5.78            | 1.1                           |
| GaN            | 3.39             | 8.90                                   | 6.15            | 8.9                           |
| GaP            | 2.26             | 4.94                                   | 4.138           | 6.8                           |

Partially from [19].

3. Geant4 numerical simulations

Geant4 simulations were carried out following a methodology used in previous works [14, 21, 22]. We considered a target of bulk material (composed of a pure material chosen between the eight binary compounds studied) with a parallelepiped geometry (surface 1 cm$^2$, thickness 20 μm). Each target was virtually irradiated with neutrons arriving perpendicularly to the surface and for which the energy distribution follows the terrestrial natural neutron background at sea level. This latter was chosen equal to the high-energy (above 1 MeV) neutron spectrum measured by Goldhagen et al. at sea level (New York City) [23, 24]. A total of $10^8$ of incident neutrons were considered, representing the equivalent duration of $5 \times 10^6$ h of natural neutron irradiation at sea level.

Geant4 version 4.9.4 patch 01 was used for these simulations. The list of physical processes employed in these simulations is based on the standard package of physics lists QGSP_BIC_HP [25]. Other simulation details can be found in Ref. [21]. Outputs of each simulation consist in a series of files containing all the information related to the neutron interaction events in the target material. For each event, this information includes the nature and the coordinates of the vertex of the interaction, the energy of the incident neutron, the exhaustive list of secondary particles produced during the interaction, and the energy and the emission direction vector...
for each of these emitted particles. All these records have been post-treated and formatted into the final interaction databases (text files) following two compilation rules: (i) we eliminated in the output file (raw data) all secondaries below 1 keV of energy; (ii) we also eliminated all γ photons, π⁰, e⁺, e⁻, and η particles, these particles being not able to deposit 1 keV of energy in the targets, which is a mandatory condition to observe single-event effects in electronics. The computational time was approximately of 3 weeks for eight parallel runs on multicore 3 GHz CPUs, the code being executed on separate cores for each material.

4. Simulation results

Table 3 gives the distributions of elastic, inelastic, and nonelastic events recorded in the different III–V compound material databases. Values for the database related to silicon are also reported for comparison. For memory, the interactions of neutrons with atomic nuclei can occur through two major mechanisms: scattering (which can be subdivided in elastic and inelastic processes) and capture (also called nonelastic) [26]. When a neutron is involved in an elastic scattering, the nature of the interacting particles is not modified; in particular the recoil nucleus is then the same as the target nucleus A or B in the case of a binary compound material AB. Similarly, during an inelastic scattering, the impacted target nucleus A or B undergoes an internal rearrangement into an excited state which eventually releases radiation. Instead of being scattered, an incident neutron may be absorbed or captured by a target material nucleus A or B. Many reactions are possible, and a large variety of particles can be emitted. This type of interaction is also called nonelastic interaction.

As evidenced by results of Table 3, the eight III–V materials show an interaction rate superior to the value observed for the reference case with Si even if GaSb, AlAs, and InSb show a number of interaction events, the closest from the one of Si. A single material stands out clearly from the others in terms of very high number of interactions: GaN. This result is the direct consequence of GaN crystallographic structure that leads to a number of atoms per volume unit almost double with respect to all other materials (see Table 1).

| Target | Elastic | Inelastic | Nonelastic | Total |
|--------|---------|-----------|------------|-------|
| Si (reference) | 11,369 | 2048 | 3373 | 16,790 |
| InSb | 7868 | 6624 | 2574 | 17,066 |
| AlAs | 9246 | 4490 | 3508 | 17,244 |
| GaSb | 8210 | 6146 | 3298 | 17,654 |
| InP | 9636 | 5392 | 3218 | 18,246 |
| InAs | 8432 | 6642 | 3290 | 18,364 |
| GaAs | 8426 | 6374 | 4356 | 19,156 |
| GaP | 10,556 | 4524 | 4150 | 19,230 |
| GaN | 15,587 | 6349 | 6537 | 28,473 |

Table 3.
Number of elastic, inelastic, and nonelastic interactions induced by atmospheric neutrons in the different III–V binary compound materials obtained from Geant4 simulations.
Concerning elastic events, their proportion is clearly predominant for Si (68%), GaP, GaN (55%), AlAs (54%), and InP (53%). The sum of inelastic and nonelastic events is above 50% for GaAs (56%), InAs and InSb (54%), and GaSb (53%). For Si, this sum is only around 32%, which is lower than 40% found in previous studies [21]. The reason is that, in the present case, the minimum kinetic energy (energy cutoff) to take into account recoils and secondaries in the databases has been fixed to 1 keV, whereas this value was fixed to a higher value, i.e., 40 keV, in previous studies. According to [15], the number and, consequently, the proportion of elastic events with respect to inelastic and nonelastic events clearly depend on such energy cutoff; it is the reason why these percentages may vary.

Figure 1 shows the number \( n \) of secondary product(s) per interaction for the different materials. It is important to note that \( n \) has been evaluated after excluding \( \gamma \) photons, \( \pi \), \( e^+ \), \( e^- \), and \( \eta \) particles from the raw simulation data, as mentioned in Section 3. As a consequence, the first class of the distribution, i.e., \( n = 1 \), corresponds to single product reactions that only result in a single recoil after interaction. These reactions correspond to all elastic and inelastic events, totalized in Table 1.

For \( n \geq 2 \), all secondaries are produced during nonelastic interactions, with an average number of secondary products per reaction between 2.5 and 2.7 for the different III–V materials (2.6 for silicon). Note that GaN and, in a lesser extent, GaAs and GaP show the most important number of reactions for \( n \) ranging from 2 to 5. To understand these results, a more detailed analysis in terms of secondary products must be conducted.

Precisely, Figure 2 shows the distribution of secondaries produced in the different targets of III–V binary compound materials subjected to atmospheric neutrons. Four groups of particles have been defined as a function of the atomic number \( Z \): (i) protons, deuterons, and tritons (\( Z = 1 \)), (ii) alpha particles and He-3 (\( Z = 2 \)), (iii) nuclei corresponding to target atoms (named recoils #1 and #2 for binary compounds), and (iv) other nuclei (with \( Z \) different from the other three categories).

Figure 2 shows that similar distributions of these particle groups are found for all materials, GaN excepted, with approximately 20–25% of protons+alphas, 60% of recoil products constituted of target nuclei, and the remaining 15% of other nuclei. In the case of GaN, the proportion of protons and alpha particles is more important and corresponds to approximately two times the proportion observed in silicon; this result can be explained by the additional production channels of protons and alpha particles offered by the \( ^{14}\text{N}(n,p)^{14}\text{C} \) and \( ^{14}\text{N}(n,\alpha)^{11}\text{B} \) reactions [27]. Always for GaN, the combination of a high number of atoms per volume unit
and the presence of nitrogen are at the origin of such a large difference with respect to other III–V materials in terms of a higher number of interactions, of secondaries produced, and of alphas and protons. In addition to Figure 2, Figure 3 shows the exhaustive distribution of all secondaries produced in all materials as a function of Z. The stacked declination of the histograms is only to produce a clearer comparison.

**Figure 2.**
Number of secondaries in the events recorded in the eight III–V material databases for the four product classes defined in the text. The nature of recoils is indicated for each material. Results for silicon are also shown for reference.

**Figure 3.**
Distribution in number (left) and in percent (right) of all secondaries produced in all materials as a function of their atomic number.
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Figure 4 shows the energy mappings for all secondaries produced by atmospheric neutrons in the eight different targets. In these figures, each colored point corresponds to a single secondary product; the x-coordinate of the point corresponds to the energy of the incident neutron which gave birth to it; the y-coordinate corresponds to the kinetic energy of the secondary after its release at the level of the reaction vertex. The same four groups of particles as previously defined (Figure 2) have been considered. Such product energy mappings allow us to visually compare the content of the different databases and to highlight similarities and differences between the atmospheric neutron susceptibilities of the eight studied materials, notably in terms of protons, alphas (GaN and GaAs as compared to InSb and InP), and recoil distributions (differences between light and heavy recoils: N, Al, and P as compared to Ga, As, In, and Sb). In Figure 4, we can see that all points are logically below the straight-line y = x because the maximum energy of secondary products is, at most, equal to that of the incident primary neutron.

From the data in Figure 4, we deduced the energy histograms in Figure 5 for all the secondary products in the eight materials. To limit the number of curves, these histograms have been calculated for the four classes of secondaries previously defined, i.e., for protons, alpha particles, target nucleus recoils, and other ions. These energy histograms show the same information as in Figure 4 but in a more compact and quantitative form. These curves indicate the energy domain covered for each type of particle and also the energy position of the maximum of the distributions. For recoil products, the heavier the recoil nucleus, the lower the energy of the maximum of the distribution. For protons, alphas, and other nuclei, similar distributions and energy domains are obtained (from $10^{-1}$ to $10^3$ MeV for protons, from 1 to $10^5$ MeV for alphas, from $10^{-2}$ to a few tens of MeV for other nuclei) for the different targets, excepted for GaN, InAs, and InSb. For GaN, an excess of low-energy protons and alphas is clearly evidenced in the range of 0.1–1 MeV; for InAs and InSb, proton distributions are narrower and only cover an energy range between 1 and $10^3$ MeV.

From the above data, we deduced the linear energy transfer (LET) and range distributions of all secondary products for the different target materials. For memory, the linear energy transfer or LET of a charged particle refers to the stopping power, i.e., the energy lost by unit of length, due only to electronic collisions and expressed in MeV/(mg/cm$^2$) or in keV/μm. The range corresponds to the distance that a charged particle travels from its emission point through matter until it comes to rest. Both LET and range primarily depend on the type and initial energy of the particle and of the medium (i.e., the target material) in which the particle travels.

We used the popular and reference code Stopping and Range of Ions in Matter SRIM [28, 29] to perform (in batch mode controlled from a short C-code) the automatic calculation of the LET and range tables (in the range 1 keV to 10 GeV) for all possible nuclides from Z = 1 to Z = 92, which represents 2670 SRIM tables per material. A total of 2670 × 8 = 21,360 files were computed and indexed in different folders in order to be directly read from a Python code to automatically compute LET and range histograms. Figure 6 shows the plots of SRIM tables for LET and range related to protons, alphas, and recoil products in the eight III–V materials.

The cross-combination of these SRIM data with particle types and energies listed in the different interaction databases was performed using a dedicated python script to compute (by linear interpolation of the SRIM tables) the initial LET (at product release) and the range histograms of all products, shown in Figures 7 and 8, respectively.
Figure 4.
Energy distributions as a function of the incident neutron energy for all secondaries produced in the events recorded in the eight III–V material databases.
Figures 7 and 8 logically show that the smaller the particle, the lower the initial LET, and the higher the range of the particle. Thereby protons exhibit the lowest LET values, in the interval from $10^{-3}$ to a few $10^{-1}$ MeV/(mg/cm²) for all materials, and the highest ranges, typically from $10^{-6}$–$10^{-5}$ m to a few tens of cm. Alpha particles are characterized by intermediate values, with typical initial LET values between 0.1 and 1 MeV/(mg/cm²) and ranges in the domain from micron to millimeter. Finally, recoil products and other nuclei exhibit the highest initial LET values, up to 10 MeV/(mg/cm²), and the lowest ranges, from nanometer to a maximum of few tens of microns. These results strongly suggest that product recoils will play an increasing role in submicron and nanowire devices, as explored in the following.
Figure 6.
Linear energy transfer (LET) and range related to protons, alphas, and recoil products in the eight III–V materials calculated with SRIM.
5. Discussion

In complement to [15], we examine, in this last section, some consequences of the above results in terms of creation of single-event effects in electronics based on the studied III–V materials. Following neutron interactions with target nuclei and release of secondaries, the energy deposition of these secondary products in the bulk material and its conversion to free charge via the creation of electron-hole pairs are another fundamental step in the creation of SEEs.

First, we evaluated two global metrics to quantify such an energy deposition consecutive to neutron interactions in the bulk of the III–V materials: (i) the mean value of deposited energy per interaction (averaged over all events) and (ii) the corresponding amount of created electron-hole pairs. These two metrics have been evaluated, in Figure 9, for the different materials. The first metric, i.e., the mean energy deposited per interaction, is found to vary from around 9 MeV for Si to 14 MeV for GaAs. Atmospheric neutron interactions deposit, in average, more energy in III–V materials than in Si. This result is the consequence of a difference in the number of nonelastic interactions, which is higher in III–V materials than in Si, as shown in the following (Figure 10 and text).

Figure 9 compares the number of e-h pairs created per interaction in the different III–V materials with respect to the value obtained for silicon (relative factor). Three groups of materials can be considered: (1) GaAs, AlAs, InP, GaN, and GaP that are found very similar to Si (relative factor close to 1 with respect to Si); (2) InAs and GaSb which exhibit a relative factor between 1.5 and 2.5; and, finally, (3) InSb characterized...
by the highest factor, close to 4. For the first group of materials, the higher atmospheric neutron sensitivity of these materials (which is indicated by the number of interactions) is largely compensated by a higher energy of e-h pair creation at semiconductor level, due to their large bandgap. On the contrary, for the three last materials which are lower bandgap semiconductors (see Table 1), their vulnerability to atmospheric
neutrons is finally reinforced by their electrical response level due to the relatively low values of the energy of e-h pair creation.

Global results in Figure 9, however, do not show important quantitative differences that exist between the respective contributions of, on the one hand, elastic or inelastic interactions and, on the other hand, nonelastic reactions, to these global values. In order to examine this particular point, we recalculated the average energy deposited per interaction. Results are shown in Figure 10. Elastic or inelastic interactions deposit, in average, between 0.1 MeV (GaSb) and 0.25 MeV (GaN) with a value of 0.23 MeV for Si. On the contrary, nonelastic interactions deposit, in average, more than 50 MeV for GaN and up to 75 MeV for InSb, with a comparative value of 45 MeV for Si. In other words, nonelastic interactions deposit approximately between ×200 (GaN) and ×600 times (GaSb) more energy in average than elastic or inelastic events. Because these nonelastic events are less numerous than elastic+inelastic events (see Table 3), this explains why the global average energies per interaction (Figure 9) are relatively modest, distributed between 10 and 15 MeV. But these values hide very different situations, depending on the type of interactions. Converted into electron-hole pairs, nonelastic reactions are able to deposit, in average value integrated over all the secondary tracks, charges up to a few pC, whereas elastic or inelastic reactions hardly reach values around a few fC.

In addition to the previous results, we evaluated the fraction of the total energy deposited per type of secondaries for all interaction events in the semiconductor bulks. Results are shown in Figure 11. Surprisingly enough, over 85% of the total energy deposited in the different III–V materials is deposited by protons, followed by alpha particles (around 10%). Other nuclei and recoil products represent less than 5%, whereas for silicon, they represent around 8% and alphas only 5%. Considering the range of protons and alphas, this result shows that the largest part of the energy is deposited far from the location of the reaction vertex.

Conversely, if we consider the fraction of the total initial LET deposited per type of secondaries for all interaction events in the different materials, we obtain results shown in Figure 12. For III–V material, 70% to 80% (minimum for GaP, maximum for InSb) of the total of the initial LET values corresponds to recoil products. The remaining 10% is related to other nuclei, the contributions of protons and alpha particles being negligible in this total initial LET. This signifies that recoil products...
are essentially at the origin of the charge deposited in the first (tens of) nanometers around the incoming neutron interaction point (also called reaction vertex).

To evaluate the deposited charge in the first nanometers around reaction vertex points, we derived this quantity from the initial LET of particles just after their release. Introducing $E_{\text{eh}}$, the energy needed for creating an electron-hole pair given in Table 2, and $t_{\text{Si}}$, the thickness of the material thin layer in which the charge $Q_{\text{dep}}$ is deposited, we obtain [13]:

$$Q_{\text{dep}} [\text{fC}] = A \times \text{LET} \left[ \text{MeV}/\text{(mg/cm}^2) \right] \times t_{\text{Si}} [\mu\text{m}]$$  \hspace{2cm} (1)

where $A$ is a numerical factor that only depends on the semiconductor material density $\rho$ and on the value of $E_{\text{eh}}$, given in Table 4 for all studied III–V materials and for silicon:

$$A = \frac{16.02 \times \rho [\text{g/cm}^3]}{E_{\text{eh}} [\text{eV}]}$$  \hspace{2cm} (2)

From Equation (1), we derived an estimation of the charge deposited in the first nanometers from the reaction vertex, averaged over all secondaries for all

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Figure 11. *Fraction of the total energy deposited per type of secondaries for all interaction events in the bulk of the studied III–V semiconductors.*

Figure 12. *Fraction of the total initial LET deposited per type of secondaries for all interaction events in the bulk of the studied III–V semiconductors.*
interaction events per material. These results are shown in Figure 13 for a distance of 10 nm. The comparison of values in Figure 13 with the current values of the minimum critical charge for standard SRAM memory cells as a function of the technological node, reported in Table 5, shows that three materials, InSb, InAs, and GaSb, exhibit a charge largely superior to 0.7 fC, the minimum critical charge for the 14 nm node. On the contrary, three other material show a charge inferior to the reference value for silicon (0.36 fC), i.e., GaN, GaP, and AlAs. Finally, GaAs and InP exhibit intermediate values around 0.5 fC, also compatible with integration down to 14 nm. Finally, the comparison of these results with those of Figure 9 suggests that integrated electronics based on InSb, InAs, or GaSb semiconductor material should be potentially more affected by neutron-induced single-event effects than those based on the other studied materials and on silicon. Of course, such a prediction will have to be verified experimentally, when devices and circuits based on these new materials for CMOS electronics will be available to be tested for radiation.

\[ Q_{\text{dep}}(\text{fC}) = A \times \text{LET}([\text{MeV/(mg/cm}^2]) \times t_{\text{Si} \mu \text{m}} \]

| Material   | Coefficient A |
|------------|---------------|
| Si (for memory) | 10.3          |
| InSb       | 84.1          |
| AlAs       | 8.8           |
| GaSb       | 33.2          |
| InP        | 17.1          |
| InAs       | 50.4          |
| GaAs       | 17.7          |
| GaP        | 9.5           |
| GaN        | 11.1          |

The numerical value of the coefficient A for this expression is indicated for the eight III–V materials (and for memory for silicon).

Table 4. Analytical expression of the charge deposited by a particle of a given initial LET (supposed constant) in a given thickness of a given material.

Figure 13. Charge deposited in the first 10 nm from the reaction vertex, averaged over all secondaries for all interaction events per material.
6. Conclusion

In conclusion, Geant4 numerical simulation was intensively used in this work as an exploration tool to anticipate the radiation response of III–V binary compound semiconductors and future electronics based on these materials (not yet available) subjected to high-energy atmospheric neutrons. Eight III–V semiconductors have been considered: GaAs, AlAs, InP, InAs, GaSb, InSb, GaN, and GaP. Simulations of bulk targets (with natural isotope compositions) exposed to sea-level neutron spectrum showed that all III–V materials exhibit more interactions than in silicon, in particular for GaAs, GaP, and GaN, the latter being a somewhat special case due to the presence of nitrogen, whose reactions with neutrons lead to an increased production of protons and alpha particles. A detailed analysis based on histograms on secondary product energy, LET, and range has been conducted, providing synthetic data about the atmospheric neutron radiation response of all these materials. We also have shown the importance of product recoils in the radiation response of ultimate CMOS electronics, due to their relative elevated masses that result in very short particle ranges and significant charge deposition within nanometer distances from the interaction points. We draw attention to the integrated electronics based on InSb, InAs, or GaSb semiconductor that should be potentially more affected by neutron-induced single-event effects than those based on the other studied materials. The current lack of experimental data in the literature forces us to consider these projections with all the precautions that we must constantly have in mind when it comes to simulation results that have not been validated experimentally.

Conflict of interest

The authors declare no conflict of interest.

Table 5.
Minimum critical charge versus CMOS technological nodes for standard SRAM memory cells.

| CMOS node (nm) | 180 | 130 | 90 | 65 | 45 | 32 | 22 | 14 |
|----------------|-----|-----|----|----|----|----|----|----|
| Critical charge $Q_{\text{crit}}$ (fC) | 3.5 | 2.9 | 1.9 | 1.3 | 0.9 | 0.85 | 0.8 | 0.7 |

After Seifert et al. [30].

CMOS node (nm) 180 130 90 65 45 32 22 14
Critical charge $Q_{\text{crit}}$ (fC) 3.5 2.9 1.9 1.3 0.9 0.85 0.8 0.7

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