

# Atomistic modeling and experimental studies of radiation damage in monazite-type LaPO<sub>4</sub> ceramics

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## Abstract

We simulated the threshold displacement energies ( $E_d$ ), the related displacement and defect formation probabilities, and the energy barriers in LaPO<sub>4</sub> monazite-type ceramics. The obtained  $E_d$  values for La, P, O primary knock-on atoms (PKA) are 56 eV, 75 eV and 8 eV, respectively. We found that these energies can be correlated with the energy barriers that separate the defect from the initial states. The  $E_d$  values are about twice the values of energy barriers, which is explained through an efficient dissipation of the PKA kinetic energy in the considered system. The computed  $E_d$  were used in simulations of the extent of radiation damage in La<sub>0.2</sub>Gd<sub>0.8</sub>PO<sub>4</sub> solid solution, investigated experimentally. We found that this lanthanide phosphate fully amorphises in the ion beam experiments for fluences higher than  $\sim 10^{13}$  ions/cm<sup>2</sup>.

## Keywords:

Radiation damage; Ceramic materials; Molecular dynamics; Threshold displacement energy; Energy barrier; Irradiation experiments; Nuclear waste management;

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## 1. Introduction

Monazites are rare-earth phosphate minerals ( $LnPO_4$ ) that occur in nature often containing significant amounts of radioactive elements, such as Th or U, without indication of significant radiation damage imposed on their crystalline structures [1]. Being chemically durable monazite-type ceramics are considered as candidate materials for nuclear waste disposal form suitable for long term immobilization of actinides, in particular plutonium [2, 3, 4]. Therefore, various relevant properties of these materials have been extensively investigated. These include the structural, the thermochemical and the thermodynamic parameters (e.g. [5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15]) as well as the dissolution [16], the elastic [17, 10] and the radiation damage properties [18, 19].

Threshold displacement energy ( $E_d$ ) is a minimum kinetic energy required to displace an atom from its lattice site. It is a fundamental parameter used to define

the radiation tolerance of materials and to estimate the extend of radiation damage during a radiation process, using for instance software such as Stopping and Range of Ions in Matter (SRIM) [20, 21, 22]. Because of the short, ps time-scale of the radiation cascade processes, atomistic modeling is a good tool to obtain the values of  $E_d$ , which otherwise is challenging to experimental methods. Such simulations have been performed recently for many materials, including TiO<sub>2</sub> rutile [20], ZrO<sub>2</sub> [23], BaTiO<sub>3</sub> [24], SrTiO<sub>3</sub> [25], or graphene and carbon nanotubes [26], to name but a few.

To displace an atom permanently, there are energy barriers separating the initial state and the final defect state in materials. Knowing the final state, these barriers can be calculated using, for instance, the nudged elastic band (NEB) method, but can be also traced during simulations of the  $E_d$  values. In previous study of radiation damage in diamond, Wu & Fahy [27] found that the damage threshold energy is almost twice the sum of bond-breaking and crystal strain energy due to the efficient dissipation of the kinetic energy of primary knock-on atom (PKA) to the crystalline lattice vibra-

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Table 1: The Buckingham potential parameters used in the simulations. [7]

	A (eV)	B (Å)	C (Å <sup>6</sup> · eV)
La-O	17927	0.25934	0.0000
Gd-O	13271	0.26	0.0000
P-O	877.3	0.3594	0.0000
O-O	22764.3	0.1490	27.879

tions. However, the question if this is intrinsically related to diamond or a general property of materials remains open.

In this contribution we derived the  $E_d$  values and the related displacement and defect formation probabilities for LaPO<sub>4</sub> monazite-type ceramics and compare the results with the recent studies of TiO<sub>2</sub> rutile [20]. The obtained  $E_d$  values were subsequently used in simulations of extend of radiation damage in La<sub>0.2</sub>Gd<sub>0.8</sub>PO<sub>4</sub> monazite-type solid solution in order to help in setting up the proper conditions of the irradiation experiments. We also report our first results on the ion beam irradiation of this material.

## 2. Computational and experimental details

The simulations of  $E_d$  values were performed with the LAMMPS code using, in addition to the standard Coulomb interaction term, the Buckingham-type interaction potentials,

$$\Phi_{12} = A \exp(-Br) - C/r^6, \quad (1)$$

which  $A$ ,  $B$  and  $C$  parameters for  $Ln$ -O interactions have been fitted so the classical simulation reproduce the *ab initio* data of Blanca-Romero et al. [7], and the parameters for P-O and O-O interaction are the ones of Gale & Henson [28] and Girard et al. [29]. All the parameters are given in Table 1.

We simulated the PKA  $E_d$  values and the displacement and defect formation probabilities in the PKA energy range of 50-150 eV for La, of 75-250 eV for P and of 8-50 eV for O. The simulations were performed with the supercells containing 1536 atoms and for each PKA energy we performed 100 independent simulations with the PKA initial velocity directions distributed randomly and symmetrically on a surface of a sphere using the Thompson model [22]. In our simulations both methods yielded very similar results. All the simulations were 5

ps long which was enough for the diminishing of the effect of the initial cascade and subsequent equilibration of the system. In order to estimate the displacement probability and the defect formation probabilities we used an algorithm to analyze displacements and defects according to the initial and final positions of atoms in the lattice. These simulations were performed with  $T = 300$  K, controlled by a thermal layer.

The subsequent calculations of energy barriers and the defect states were performed using NEB and metadynamics methods. The NEB calculations were performed with the relevant package implemented in the LAMMPS code [30] and the metadynamics simulations were performed with the PLUMED plug-in [31].

The penetration depth of the ions, the resulting displacements of target atoms and the distribution of vacancies in the experimentally studied La<sub>0.2</sub>Gd<sub>0.8</sub>PO<sub>4</sub> system were calculated with the SRIM/TRIM software package, using the SRIM-2013 code (www.srim.org). SRIM/TRIM (Stopping and Range of Ions in Matter/Transport of Ions in Matter) comprises a set of programs that can simulate the interactions of ions with energies up to 2 GeV/amu with matter, based on a full quantum mechanical treatment of the collisions of incident particles with atoms present in a target material [32, 33]. The code is based on a Monte Carlo (MC) simulation method and the binary collision approximation (BCA) [34, 35]. Simulation results comprise, for example, the 3D-distribution of ions and the concentration of vacancies in the target material as well as the energy partitioning between nuclear and electronic energy losses, with all target atom cascades in the target material followed in detail. SRIM/TRIM generally assumes that the target is isotropic and amorphous.

For the irradiation experiments a highly densified ( $\rho_{\text{ sint}} = 97\%$  of theoretical density (TD)) La<sub>0.2</sub>Gd<sub>0.8</sub>PO<sub>4</sub> pellet of 10 mm diameter and 1 mm thickness has been prepared according to Neumeier et al. [36] and Arinicheva et al. [37]. The purity of the monazite sample material was confirmed by the XRD measurements (Bruker D8-Advance X-ray diffractometer (XRD)). The pellet was irradiated at room temperature with 100 MeV <sup>197</sup>Au<sup>9+</sup> ions delivered by the 15 UD Pelletron accelerator at the Inter-University Accelerator Centre (IUAC) Delhi, India at the ion fluence ranging from 10<sup>12</sup> ions/cm<sup>2</sup> to 2 · 10<sup>14</sup> ions/cm<sup>2</sup>. The ion flux was kept below 2.8 · 10<sup>10</sup> ions cm<sup>-2</sup>s<sup>-1</sup> in order to avoid ion beam induced heating of the target materials. A Bruker D8-XRD was used for in-situ investigations of the irradiation induced structural modifications [38, 39]. The in-situ experiments were performed on the same pellet by successive irradiation and the immediate subsequent

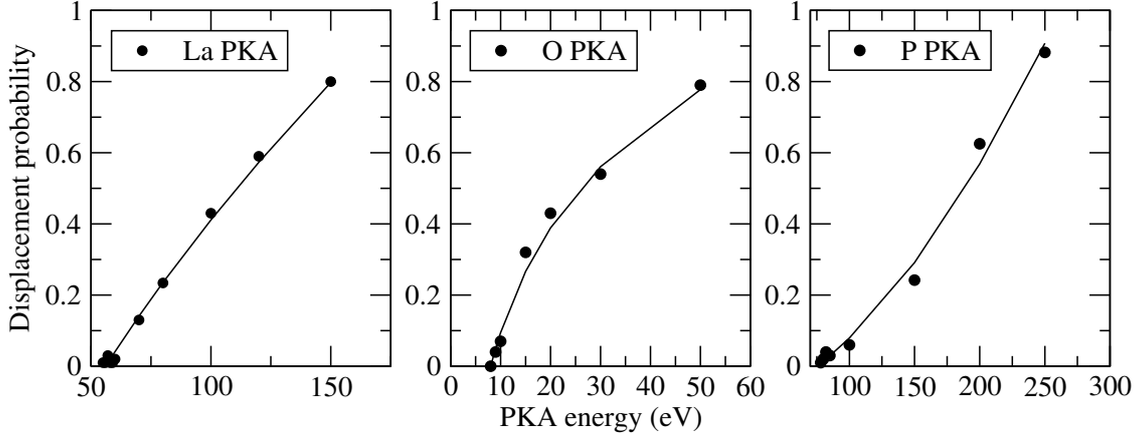


Figure 1: The atom displacement probabilities of La, O, P PKA in  $\text{LaPO}_4$  simulated at  $T = 300$  K.

128 XRD measurement without changing experimental pa-  
 129 rameters in order to compare the intensity of the diffrac-  
 130 tion reflections of the sample exposed to the different  
 131 ion fluences. All XRD patterns were recorded under  
 132 vacuum ( $5 \cdot 10^{-6}$  mbar) in the  $2\theta$  range of  $10$ - $90^\circ$  with  
 133 increments of  $0.02^\circ$  at a scan speed of  $0.5^\circ \text{ min}^{-1}$ .

### 134 3. Results and discussion

#### 135 3.1. Threshold displacement energy of $\text{LaPO}_4$

136 The displacement probabilities as a function of PKA  
 137 energy for La, P and O atoms are shown in Figure 1. In  
 138 the figure each point is the average value obtained by  
 139 sampling the 100 PKA directions. The threshold dis-  
 140 placement energy can be obtained from the relationship  
 141 between the initial energy and the displacement proba-  
 142 bilities by fitting the equation [20, 22]:

$$DP(E) = [E^\alpha - E_d^\alpha]/\beta, E > E_d, \quad (2)$$

143 where  $\alpha$ ,  $\beta$  and  $E_d$  are the fitting parameters and  $E$  is  
 144 the PKA energy. The  $E_d$  value fitted for La is 56 eV, for  
 145 P is 75 eV and for O is 8 eV. These values indicate that  
 146 it is easiest to form an O defect and hardest to form a P  
 147 defect in the  $\text{LaPO}_4$  lattice. This is because in  $\text{LaPO}_4$ ,  
 148 one P atom is bonded with four O atoms and one  $\text{PO}_4$   
 149 is interacting with one La atom, which results in the dif-  
 150 ferent bounding strengths and resulting  $E_d$  values. In-  
 151 terestingly, the  $E_d$  value for La is similarly large as the  
 152 one obtained for Ti cation in  $\text{TiO}_2$  rutile (69 eV, [20]).  
 153 Also, the difference between the displacement probabili-  
 154 ty and the defect formation probability obtained in our  
 155 studies, and shown in Figure 2, is very similar to the  
 156 one obtained for rutile. Namely, the defect formation

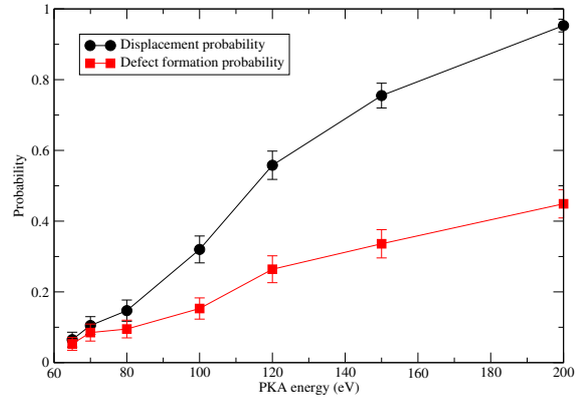


Figure 2: The displacement probabilities and the defect formation probabilities for La cations in  $\text{LaPO}_4$  as a function of PKA energy, simulated at  $T = 300$  K.

157 probability is significantly smaller and our results indi-  
 158 cate that at a temperature of 300 K about half of the  
 159 La displacements recombine to a regular La crystalline  
 160 position. In the case of rutile, Robinson et al. [20] at-  
 161 tributed the radiation damage resistance of this material  
 162 to its efficient defect recombining ability. Our similar  
 163 results indicate thus a possibility of a common origin  
 164 of radiation damage resistance in the case of rutile and  
 165 monazite. The  $E_d$  value obtained for the Gd cation with  
 166 the same method and used in the SRIM simulations (see  
 167 section 3.3) is 51 eV.

#### 168 3.2. Energy barriers in displacement of $\text{LaPO}_4$

169 Formation of permanent defects is related to the en-  
 170 ergy barrier ( $E_b$ ) that has to be crossed by a PKA atom.  
 171 Therefore, we checked how the energy barrier, defined  
 172 here as the minimum potential energy increase (max-  
 173 imum) during the cascade, correlates with the initial

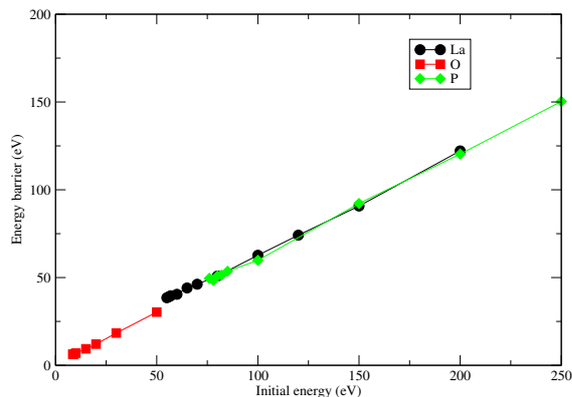


Figure 3: The relationship between the PKA energy and the energy barrier in  $\text{LaPO}_4$ . Results for all three species are plotted.

174 PKA energy and the  $E_d$  value. The relationships for the  
 175 three cations considered are presented in Figure 3. We  
 176 found that the energy barriers are substantially smaller,  
 177 by about a half, than the applied initial PKA energies  
 178 and there is no defect created, if the PKA energy is just  
 179 comparable to the energy barrier. As shown in Figure 3,  
 180 there is a linear relationship between the energy barrier  
 181 and the PKA energy,  $E_b \sim 0.58E$ , and the relationships  
 182 are very similar for all the three considered species.  
 183 This result has been verified with the subsequent calcu-  
 184 lations of barriers performed by a combination of the  
 185 NEB and metadynamics methods. Interestingly, very  
 186 similar results have been reported for diamond by Wu  
 187 & Fahy [27], who also found that the PKA energy must  
 188 be about twice the energy barrier to overcome the bar-  
 189 rier. They attempted an explanation of this phenomenon  
 190 by invoking similarity of the initial PKA velocity to the  
 191 speed of sound, which allows for efficient transfer of the  
 192 PKA kinetic energy to the energy of lattice vibrations.  
 193 Therefore, we performed a detailed analysis of the dis-  
 194 sipation of the initial PKA kinetic energy in the system  
 195 studied.

196 The evolution of kinetic and potential energies in the  
 197 two cases: (1) without defect and (2) with defect for-  
 198 mation is illustrated in Figure 4. In the case without  
 199 the defect, the PKA energy is equally distributed to the  
 200 kinetic energy of other atoms and the potential energy  
 201 of entire system. The case with the defect creation is a  
 202 little bit different. Initially, the PKA kinetic energy is  
 203 also equally distributed between the kinetic and poten-  
 204 tial energies of the system but after crossing the barrier  
 205 and equilibration, the gain in the kinetic energy of the  
 206 system is smaller than the gain in the potential energy.  
 207 In the considered case, the difference is about 12 eV.  
 208 This value is independent of the initial PKA kinetic en-

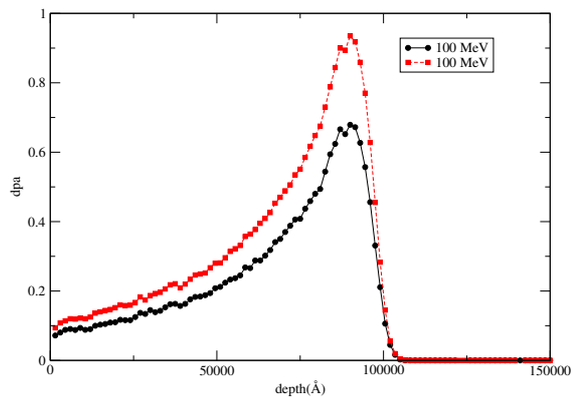


Figure 5: The relationship between defect extent (in dpa) and the ion range computed by SRIM assuming bombardment of  $\text{LaPO}_4$  with 100 MeV Au ions and a fluence of  $10^{14}$  ions/cm<sup>2</sup>. The results for  $E_d$  values (black circles) and energy barriers ( $E_b = 0.58 E_d$ , red squares) the computed here are presented.

209 energy and is equal to the defect formation energy, which  
 210 we verified through subsequent relaxation of the final  
 211 state.

212 Having this result and following the studies of Wu  
 213 & Fahy [27], we compared the PKA velocities to the  
 214 speed of sound in  $\text{LaPO}_4$  monazite. The sound veloc-  
 215 ity in  $\text{LaPO}_4$  can be calculated from the knowledge of  
 216 bulk modulus, shear modulus and material density. For  
 217  $\text{LaPO}_4$  monazite, it is about 3664 m/s [17], which means  
 218 that the sound waves can travel through the supercell in  
 219 just  $\sim 0.5$  ps and the corresponding energy is  $\sim 10$  eV.  
 220 Thus, a La PKA atom with the energy of the threshold  
 221 displacement energy of 56 eV has a velocity of 8864  
 222 m/s, which is comparable to the above-provided speed  
 223 of sound. This explains why a significant part ( $\sim 50\%$ )  
 224 of the PKA energy is efficiently transferred into the sys-  
 225 tem and dissipated through the lattice vibrations.

226 Finding a relationship between the PKA energy, the  
 227  $E_d$  values and the energy barriers can be very useful for  
 228 determination of the  $E_d$  values. This is because compu-  
 229 tation of barriers is computationally less demanding and  
 230 provides an independent way to estimate the  $E_d$  values.  
 231 For instance, the defect states could be identified with  
 232 methods such as metadynamics, and the barrier between  
 233 the initial ground state and the defect state could, for in-  
 234 stance, be computed with NEB method.

### 235 3.3. Simulation of radiation damage extent with SRIM

236 The obtained  $E_d$  values have been used in subsequent  
 237 simulations of the extent of radiation damage under con-  
 238 ditions reflecting the planned irradiation experiments.  
 239 We also made computations taking energy barriers as

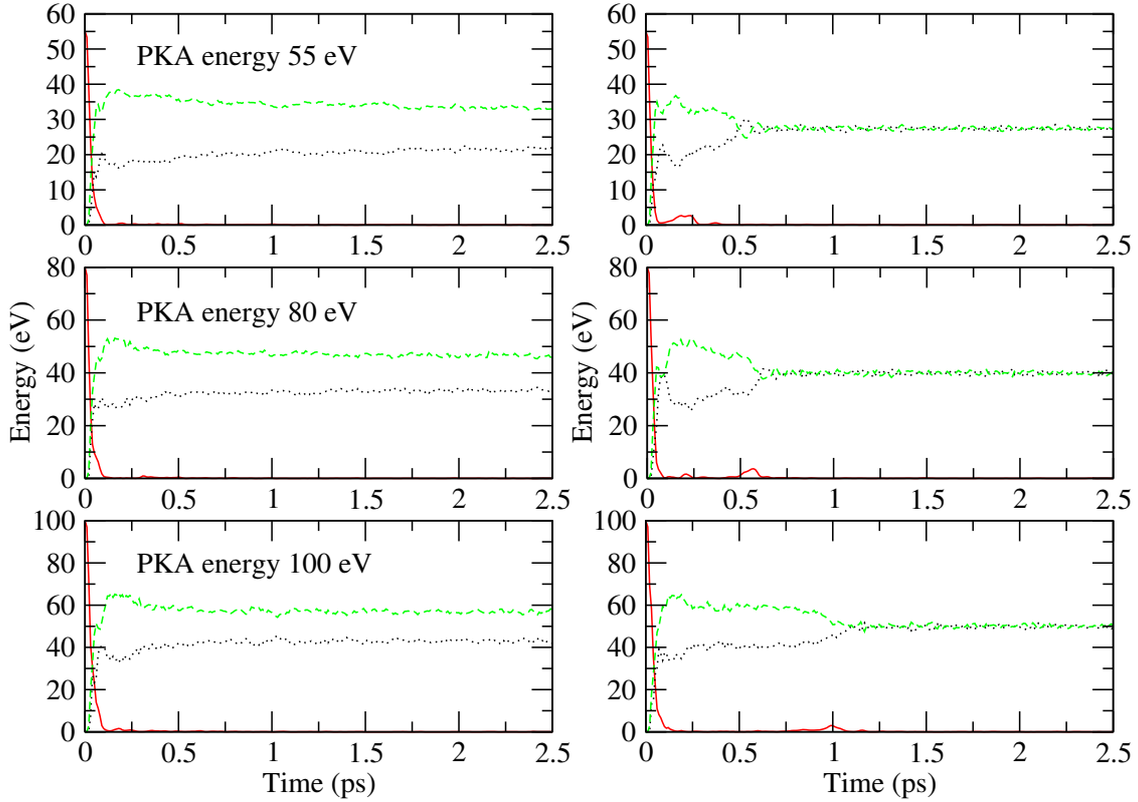


Figure 4: The kinetic energy of the PKA (solid red), the kinetic (dotted black) and potential (dashed green) energy of all the atoms except PKA atom, obtained with different initial PKA energies indicated in the upper left corners. The left panels are results obtained for cases when a defect was created and the right side panels represent the results obtained without defect creation.

240  $E_d$  values, thus reducing the  $E_d$  values to  $0.58 E_d$ . Figure 5 shows the results of such simulations. These indicate that the expected radiation dose expressed in displacements per atom (dpa) is higher than the critical amorphization dose reported for monazites ( $\sim 0.35$  dpa, [18, 2]). Thus it was ascertained that the maximum fluence selected in the irradiation experiments would be sufficiently high to allow for the amorphization of the monazite samples. The damage peaks at the depth of  $9 \mu\text{m}$  and thus should be easily detectable by XRD techniques. Also, the results of simulations with the two sets of  $E_d$  values are consistent regarding the penetration range and differ only in prediction of the damage amount, when smaller  $E_d$  values are used.

### 254 3.4. XRD measurement

255 The XRD measurements of the  $\text{La}_{0.2}\text{Gd}_{0.8}\text{PO}_4$  solid solution sample irradiated with the 100 MeV Au ions at fluences ranging from  $10^{12}$  ions/cm<sup>2</sup> to  $10^{14}$  ions/cm<sup>2</sup> agree with the SRIM calculations (Figure 6). Compared with the XRD pattern of unirradiated material, the XRD reflections of irradiated samples become broader and

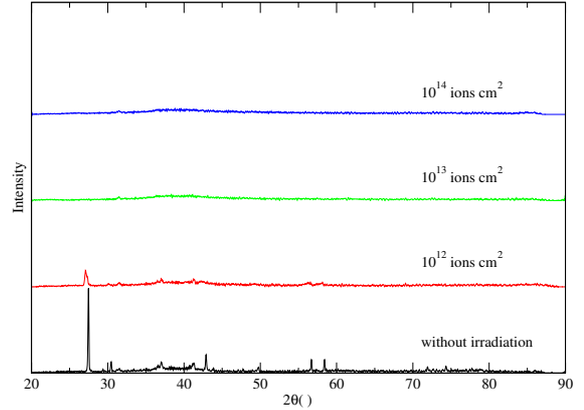


Figure 6: The XRD of  $\text{La}_{0.2}\text{Gd}_{0.8}\text{PO}_4$  solid solutions irradiated with 100 MeV Au ions at different fluences.

261 vanish gradually at higher fluences ( $10^{13}$  ions/cm<sup>2</sup>), indicating complete amorphization. However, amorphization was achieved already at a lower fluence than predicted from the SRIM results. This effect was already observed in irradiation experiments with pyrochlore-

266 type materials using swift heavy ions and is due to 314  
267 the thermal spike induced by electronic stopping effects 315  
268 [40]. 316

#### 269 4. Conclusion 317

270 Using atomistic modeling techniques we simulated 318  
271 the radiation damage resistance of the LaPO<sub>4</sub> monazite- 319  
272 type ceramics. We derived the  $E_d$  values for all three 320  
273 species constituting the investigated material. These 321  
274 values are largest for P (75 eV), significant for La (56 322  
275 eV) and relatively small for O (8 eV). Interestingly, the 323  
276 value obtained for La is similarly large as the one de- 324  
277 rived for the Ti cation in TiO<sub>2</sub>. Also, the obtained dif- 325  
278 ference between the displacement and defect formation 326  
279 probabilities derived for La in monazite is very similar 327  
280 to the results obtained for Ti in rutile TiO<sub>2</sub>, which points 328  
281 towards a similar origin of the radiation damage resis- 329  
282 tance of both materials. We found a linear relationship 330  
283 between the energy barriers separating the initial from 331  
284 the defect state and the PKA initial energy values, which 332  
285 indicates that the barrier could be crossed only if the 333  
286 PKA energy is about twice the barrier energy. This we 334  
287 explain by efficient dissipation of the PKA kinetic en- 335  
288 ergy between the potential energy and the kinetic energy 336  
289 of vibration of the crystalline. The obtained  $E_d$  values 337  
290 have been applied to simulations of radiation damage 338  
291 extent under various experimental conditions, helping 339  
292 selecting proper setup parameters for the irradiation ex- 340  
293 periments. The irradiation experiments and subsequent 341  
294 XRD measurements of the irradiated samples indicate 342  
295 full amorphization of the samples for fluences higher 343  
296 than 10<sup>13</sup> ions/cm<sup>2</sup>. The subsequent experimental and 344  
297 modeling studies are ongoing in order to improve our 345  
298 understanding of the radiation-induced amorphization 346  
299 process in monazites. 347

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