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To cite this version:
Jean-Paul Crocombette, Christian Wambeke. Quick calculation of damage for ion irradiation: implementation in Iradina and comparisons to SRIM. EPJ N - Nuclear Sciences & Technologies, EDP Sciences, 2019, 5, pp.7. 10.1051/epjn/2019003 . cea-02304265

HAL Id: cea-02304265
https://hal-cea.archives-ouvertes.fr/cea-02304265
Submitted on 3 Oct 2019

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Quick calculation of damage for ion irradiation: implementation in Iradina and comparisons to SRIM

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Received: 8 February 2019 / Received in final form: 1 March 2019 / Accepted: 29 March 2019

Abstract. Binary collision approximation (BCA) calculation allows for two types of damage calculation: full cascade and quick calculations. Full cascade mode describes fully the cascades while in quick calculations, only the trajectory of the ion is followed and effective formulas give an estimation of the damage resulting from each collision of the ion. We implement quick calculation of damage in the Iradina code both for elemental and multi-component solids. Good agreement is obtained with SRIM. We show that quick calculations are unphysical in multi-component systems. The choice between full cascade and quick calculations is discussed. We advise to favour full cascade over quick calculation because it is more grounded physically and applicable to all materials. Quick calculations remain a good option for pure solids in the case of actual quantitative comparisons with neutron irradiations simulations in which damage levels are estimated with the NRT (Norgett-Robinson and Torrens) formulas.

1 Introduction

In the nuclear context, ion irradiation is used as a way to study materials under radiation. Such experiments are cheaper and quicker than in reactor studies under neutron irradiation. Direct connection of such ion experiments with neutron irradiations face numerous challenges [1,2]. However, even without a direct connection to in-reactor experiments, ion irradiation gives many information about the behaviour of materials. Rapid numerical estimations of the damage and implantation profiles in ion irradiation experiments are very important. They are used before the irradiations to design them, and/or after experiments to choose where analyses should be performed. Such estimations requires fast modelling tools of the primary damage and implantation profiles. The Binary collision approximation (BCA) [3–5] which divides the ion trajectory in successive two-body collisions is the good formalism to perform such estimations. Indeed, it allows to quickly deal with any type of ion irradiation (ion nature and energy) in any material. The limitations of such BCA approaches are well-known. For instance, thanks to molecular dynamics (MD) simulations [6,7] and experiments [8], it has been known for decades that BCA codes overestimate the number of created defects by primary damage. Moreover, they cannot give information about the detailed structure of the clusters of defects formed directly after the collisions cascades. Nevertheless, BCA simulations remain extremely useful because they are simple and fast. The principles of BCA have been described many times in previous works [3-5] and we just recall that BCA uses generic pairwise interaction potentials and rely intensively on scattering theory. While some rare codes use energy integrals in the BCA framework to calculate average number of collisions and atomic displacements (e.g. the DART code [9]), most of the BCA codes deal with real space trajectories and describe explicitly the sequence of collisions sustained by the incoming ion.

The two major historic codes are SRIM [3,10] and MARLOWE [5]. Marlowe includes a description of the crystalline structure of the materials and because of that, it is more complex than SRIM which is based on a random material assumption. Furthermore, SRIM comes with a graphical user interface, which makes it easy to use. Thus, it has been and remains extensively used as the common tool to estimate primary damage and implantation profiles.

In SRIM, two main types of damage calculation exist: full cascade (FC) and quick calculation (QC). In both modes, the trajectory of the incoming ion through its successive collisions is followed, until its kinetic energy falls below a given threshold where it is stopped. FC mode describes fully the cascades, i.e. the trajectories of all atoms accelerated by the ions or, recursively, by previously accelerated atoms, until they all come to rest (when their kinetic energy falls below the stopping threshold). In QC
mode, only the trajectory of the ion is followed. And effective formulas give an estimation of the damage resulting from each collision of the ion (see below). From either mode, one obtains the number of atomic displacements along the trajectory of the ion. From this number and the density, one can deduce the local amount of dpa (displacement per atom).

People have used SRIM for so many years that it has become a sort of reference for the BCA estimation of displacement per atom. And the density, one can deduce the local amount of dpa along the trajectory of the ion. From this number resulting from each collision of the ion (see below). From effective formulas give an estimation of the damage mode, only the trajectory of the ion is followed. And alternative must fulfill are three-fold. First, unlike SRIM, the code must be open-source. Second, like SRIM, it must be easy to use for people non-specialized in numerical simulations. Finally, and more strangely, in order to be accepted by the community, an alternative to SRIM must provide results identical or at least very close to the ones of SRIM. Indeed, as mentioned below, SRIM has been used so intensively and for so many years that any code which gives different results, whatever their experimental accuracy and physical correctness will not be adopted by the community. Finally, in the nuclear context, an alternative to SRIM should include the QC of damage, which is very commonly used in this community.

Keeping the objective of providing an alternative to SRIM, we chose to implement QC in the Iradina code [11] originally developed by Christian Borschel and Carsten Rønning from the university of Jena. Compared to the few other modern BCA codes (see below), Iradina has some advantages. First, it is truly open-source. Second being written in C, it is easy to read. Third, it proves very fast. This is because it relies on the formalism implemented originally in the Corteo code [12] which has been developed by François Schiettekatte for ion beam analysis. In this code, the calculation of the scatterings use logarithmically-scaled stopping tables which greatly speeds up the calculation. This formalism is much faster than the one based on the so-called Magic formula implemented in SRIM.

Other modern BCA codes we are aware of include Mytrrim [13], SDTrimSP [14] and IM3D [15]. To our knowledge, SDTrimSP is not open-source. From what we understand, Mytrrim uses the Magic formula (instead of fast Corteo’s formalism) which should make it quite slow. Finally, contrary to what authors claim, IM3D is not open-source. Indeed the transport routines at the heart of this code are included in an executable archive, which one has to download separately. Strangely enough, the input variables of IM3D have almost identical names to the ones of Iradina. Marlowe code has recently be renewed in connection with mixed BCA-MD simulations [16]. But it remains not very user-friendly and much more complex than BCA codes considering random material. It can therefore not be regarded as an alternative to SRIM.

We present in the following, how we realized the implementation of QC in Iradina. We then compare our results with the ones of SRIM. Finally, we discuss QC in non-elemental solids and give our opinion on how to choose between FC and QC calculations.

2 Implementation of QC in Iradina

2.1 Elemental solids

The history of QC of damage dates back to the late 1960s [17,18]. At that time, due to the very limited computer resources available, even the BCA description of a full cascade was challenging, not to mention MD calculations, which were in their infancy [19]. The goal of QC was to estimate the amount of damage created by an accelerated atom of kinetic energy $T$; the damage being expressed as a number of displaced atoms. In the ion irradiation context, this accelerated atom results from an elastic collision with the incoming ion. With QC formulas, one did not have to describe the full cascade, but only the successive collisions of the ion.

Two steps appear in the QC formalism. First, the kinetic energy $T$ of the accelerated atom must be transformed into the so-called damage energy $E$. This energy is the portion of $T$ available for ballistic collisions, i.e. the part which is not lost to electrons through inelastic collisions. To do so, one resorts to formulas by Lindhard [20] to estimate the portion of $T$ lost by ballistic losses:

$$e_d = 0.01014 \times T \times Z^{-7/3},$$

$$E = \frac{T}{1 + k_d g(e_d)},$$

$$g(e_d) = e_d + 0.40244 e_d^{3/4} + 3.4008 e_d^{1/6},$$
\[ k_d = 0.1334 \times Z^{2/3} \times M^{-1/2}. \] (4)

In these expressions, \( Z \) and \( M \) are the atomic number and molar mass of the accelerated atom. Masses and energy are expressed in grams and eV respectively. Note that in FC calculations, electronic losses are accounted for by a slowing term applied to the moving atom between two subsequent collisions and \( E \) is the sum of the energy lost through each successive collisions with atoms in the material.

Second, the damage energy must be converted into damage production by a damage formula. Various formulations appeared in the late 1960s and early 1970s [18]. Nowadays, the so-called modified Kinchin-Pease or Norgett-Robinson-Torrens (NRT) [17] formula is used:

\[
n_d(E) = \begin{cases} 
0 & \text{if } E < E_d \\
1 & \text{if } E_d < E < 2.5E_d \\
0.8 \frac{E}{2E_d} & \text{if } E > 2.5E_d
\end{cases} \quad (5)
\]

We implemented formulas (1) to (5) in Iradina and set up a new simulation type for which the code calculates the trajectory of the sole ion and the damage created by the recoil of each collision by the above formulas. When actual damage is created \( (n_d(E) > 0) \), the code assigns it entirely to the point where collision took place. The damage is eventually stored in the vacancy file. In QC, one cannot distinguish replacements and interstitial creations. These defects are therefore not counted.

As mentioned before, BCA codes rely on a few physical assumptions which impact the coding algorithms. The determination of the flight path between two successive collisions is one of them. The choice of the flight path is closely related to the possible impact parameters of the collisions. Iradina in its original version [11] include three choices for the free flight path \( l \) a constant flight path set by the user; a constant flight path equal to the interatomic distance; a Poisson distributed flight path with inter-atomic distance as a mean value, the latter being the default. In all cases, the impact parameter is then chosen randomly with an upper maximum impact parameter which is deduced from the flight path as follows. The flight path and the maximum impact parameter can be regarded as the length and radius of a cylinder. The volume of this cylinder is set to the atomic volume:

\[
l\pi r_{\text{max}}^2 = 1/\rho, \quad (6)
\]

\( \rho \) being the atomic density so that \( 1/\rho \) is the atomic volume. One then has:

\[
P_{\text{max}} = 1/\sqrt{l\pi\rho}. \quad (7)
\]

The actual impact parameter is randomly chosen as:

\[
p = \sqrt{rp_{\text{max}}}, \quad (8)
\]

with \( r \) being a random real number between 0 and 1.

To reach a better agreement between SRIM and Iradina with QC, we had to implement a new choice of calculation of the free flight path and impact parameter: the so-called impulse approximation. Following [21], one first calculates a maximum impact parameter \( P_{\text{max}} \) then one deduces the flight path. \( P_{\text{max}} \) is set as the impact parameter corresponding to a given minimum energy transfer \( E_{\text{min}} \).

\( E_{\text{min}} \) is chosen equal to the energy at which the ion is stopped which is an input choice (reasonable values are around a few eV).

Considering the collision between an ion \( (M_1,Z_1) \) of energy \( T_j \) with a target \( (M_2,Z_2) \), one defines successively:

\( \alpha \) the screening length

\[
\alpha = \frac{0.8853a_0}{(Z_1^4 + Z_2^4)^{1/4}}, \quad (9)
\]

where \( a_0 = 0.529 \text{Å} \) is the Bohr radius;

\( \varepsilon \) the reduced energy

\[
\varepsilon = \frac{\alpha T_1}{(1 + \frac{M_1}{M_2})Z_1Z_2\varepsilon^2}, \quad (10)
\]

\( (\varepsilon_{\text{min}} \) is defined from \( E_{\text{min}} \) with the same formula).

\( \gamma \) and the maximum impact parameter \( P_{\text{max}} \) with

\[
\gamma = 4M_1M_2/(M_1 + M_2), \quad (11)
\]

\[
\xi = (\varepsilon_{\text{min}}/\gamma)^{0.5}, \quad (12)
\]

\[
P_{\text{max}} = a(\xi + \xi^{0.5} + 0.125\xi^{0.1})^{-1}. \quad (13)
\]

The flight path is then deduced from equation (6) and the actual impact parameter is selected randomly up to \( P_{\text{max}} \) (Eq. 8).

We tested our implementation of QCs in Iradina on the case of self-irradiation in iron. We calculated irradiation by Fe ions of energy 500 keV or 2 MeV with normal incidence. The damage and implantation profiles calculated with Iradina and SRIM with QCs are given in Figure 1. One can note that for both energies SRIM and Iradina predict very close damage profiles. We obtained similar agreement in the various test cases we performed on elemental solids. Our implementation of QCs is therefore validated for the case of elemental solids.

### 2.5 QC for non-elemental solids

The implementation of QCs for alloys or more generally non-elemental solids is less straightforward. Indeed while, such calculations are possible with SRIM, there is no indication whatsoever on how they are actually performed. The difficulty is easily understood looking at the above formulas (Eqs. (1) to (5)). They depend on the mass \( (M) \) and atomic number \( (Z) \) of the target atom which is therefore supposed to be also characteristic of the irradiated material. Despite the lack of any information
In SRIM documentation, one can deduce from tests in multi-component solids that the QCs in non-elemental solids proceed as follows. For each collision, the nature of the target atom is randomly selected, and the impact factor is chosen (Eqs. (6) to (12)). With the transferred kinetic energy, the damage is then estimated applying formulas (1) to (5) with the mass and atomic number of the target atom, completely neglecting the fact that the material is in fact in a multi-component system. In the following, we present some results of xenon irradiations in UO₂. In this case when the incoming ion hits a uranium (resp. oxygen) atom, the formula is applied with \( M = 238.03 \) and \( Z = 92 \) (respectively \( M = 16 \) and \( Z = 8 \)). We implemented this formulation in Iradina.

The comparison between iradina and SRIM for the UO₂ test case is given in Figure 2. One can see that the agreement is good, though not perfect. We could not find any way to improve the agreement between Iradina and SRIM. We contemplated other possibilities namely, applying the formulas to an arithmetical or geometrical average atom with \( M \) and \( Z \) given as composition averages of the \( M \) and \( Z \) of the components. These other formulations result in damage creation which deviates largely from SRIM results. This is a situation where not knowing how SRIM is actually coded proves very inconvenient and makes it impossible to reproduce its results with another code. Anyway, we believe the slight discrepancy between SRIM and Iradina for QCs in non-elemental solids is rather inconsequential. Indeed, as we shall discuss below, such calculations are unsuitable for these materials.

2.6 Iradina as an alternative to SRIM

With the above-described implementation, we achieved correct agreement between SRIM and Iradina for QCs. The agreement between SRIM and Iradina for FC calculations was already highlighted in the original Iradina paper [11].

We checked it for our test cases and indeed found very close results for Iradina and SRIM. The same is true for the ion implantation profiles. With the present addition, Iradina is now able to reproduce SRIM calculations in the two main damage calculations frameworks: FC and QC.

Being open-source, Iradina therefore fulfills two of the requirements for a satisfactory alternative to SRIM. The last one is to have an easy to use graphical user interface (GUI) for non-experts in simulations. The original Iradina already has a user interface. While this interface is quite elegant and useful, it was not conceived to run SRIM like calculations. Moreover, it is not open source and thus cannot be adapted to such calculations. We therefore preferred to design a new GUI. This GUI is briefly described in Appendix A. Finally, Iradina has recently been made available on source forge [22]. The SRIM-like GUI is also available on source-forge with packages to be used in Linux or Windows environments to perform SRIM like calculations with the Iradina code embedded in it [23].

3 Choosing between FC and QC

Users of SRIM or Iradina have to choose between two ways of calculating the damage profile in ion-irradiated materials. We want in this section to elaborate on this choice and present our opinion about it. We shall first present some comparisons between these two schemes for elemental or non-elemental solids based on the same test cases we used before. We should compare not only the damage production but the profiles of energy deposition.

3.1 Comparisons

The first thing to realize is that the two damage frameworks give quite different results. This is illustrated in Figures 3–6 in the cases of Fe and UO₂. This should be
no surprise as the two calculations are based on quite different premises. As explained above, FC describes the complete cascades while QC use formulas (1) to (5) to estimate the damage created by the Primary knocked-on atom (PKA) resulting from each collision of the ion.

Focusing for now on mono-elemental solids, one can note that QC tend to predict smaller amounts of defects than FC (Fig. 3). It is worth stressing that the ratio between QC and FC depends on the material and irradiation under study. For instance, it is around 0.5 for self-irradiation in iron and closer to 0.7 for self-irradiation in aluminium (not shown). Thus, one cannot deduce what FC results would be from QC calculations (and the other way around). As described in equations (1) to (4), QC rely on an assumed division of the kinetic energy of the PKA into electronic and ballistic losses in the material. The validity of this division can be checked by comparisons with FC calculations. Figure 4 presents, for each framework, the two types of energy deposition as a function of depth, in the case of 500 keV and 2 MeV self-irradiation in iron. One can note that Robinson formulas perform rather well. The electronic losses are however slightly overestimated especially at low energy. Another difference appears pertaining to the localization of energy losses. In FC, the calculated cascades have some spread which dispatch the energy losses in the volume of the material. At the opposite, in QC, the energy deposition of an (implicit) cascade is located at the position of the corresponding collision with the incoming ion, neglecting its spread. The energy deposition due to the first collisions of the ions which happen close to the surface are then staked at the lowest depths. This is especially visible at low energy (left panel of Fig. 4).

QC thus perform reasonably well for elemental solids. Considering Figures 5 and 6, it appears readily that the differences between FC and QC are more striking for non-elemental solids. At first, considering the global defect production (Fig. 5), one obtains the same kind of difference between FC and QC that appeared for elemental solids. However, the material being made of multiple species, one can also consider the amount of vacancies created for each species. In the case of UO$_2$, one then sees that the ratio of created vacancies for each species (U and O) is very different for FC and QC. It appears clearly that the distribution of damage between O and U is unphysical for QC. Indeed, QC predicts that the majority of vacancies are of the uranium type. Even considering the differences in collision cross-sections, this contradicts the fact that there are twice as many oxygen atom than uranium and that their displacement energy is two times smaller.

Another problem with QC for multi-component materials appears for the energy losses. Figure 6 clearly shows that QC and FC are in strong disagreement for the balance between electronic and ballistic energy depositions. This is
especially a problem when one wants to study the relative or synergistic effects of electronic over ballistic losses. Ratios of energy depositions calculated with QC are then utterly wrong.

Using QC instead of FC for non-elemental solids thus induces large errors in the distributions of created defects and energy losses compared to FC calculations. Applying QC for such systems is therefore very questionable. These discrepancies are naturally related to the application of mono-elemental formula for each component of the materials after each collision. This assumption is obviously un-physical.

One may wonder why such an approximation was coded in SRIM in the first place. The reason may be related to the NRT calculation of damage in neutronics codes. Indeed neutron transport and reaction codes such as Tripoli [24] or NJOY [25] are primarily concerned with the fate of the neutrons through interactions with atoms of the materials. They consider series of nuclear reactions or elastic collisions of a neutron with atoms. The amount of damage created by the result of a nuclear reaction or an elastic collisions are secondary outputs of the neutron code irrelevant for the main calculation. Because such codes consider interactions with one atom at a time, the damage formulas depend on the sole nature of the atom interacting with the neutron. Physically, this is as wrong as in ion irradiation codes, but the consequences for neutronics are zero. Indeed, damage productions being simple pieces of information, they do not affect the fate of the neutrons which is the actual object of neutronics codes. One may suppose that NRT formulas were implemented in neutron codes, and then simultaneously introduced in SRIM for consistency with these codes even for multi-component systems where they are physically irrelevant.

### 3.2 Choice between QC and FC

As explained above, FC is more physically grounded than QC. It may seem that FC should therefore always be preferred. Nevertheless, a few reasons have been put forward to prefer QC over FC. We review them briefly. The most basic one is that in SRIM, FC calculations are extremely slow and QC are better suited to quickly estimate the damage. This practical argument does not apply to Iradina where both QC and FC are very fast (Iradina is actually about two orders of magnitude faster than SRIM). Another questionable argument is the fact that QC tend to be somewhat closer to the MD predictions than FC calculations. But QC results still deviates strongly from MD. At high PKA energies, there is an overestimation of about 3 for the number of defects with QC in iron [26] (the factor is thus about 6 for FC). Thus, one cannot say that the agreement is between QC and MD is good (see also below the discussion about arc-dpa).

More profound arguments have been put forward by Stoller et al. [27]. The conclusion of their work “On the use of SRIM for computing radiation damage exposure” is three-fold:
– it is not possible to determine the “right” number of displacements generated by a given PKA in any absolute sense. MD simulations provide the most realistic estimate, but MD results are also model; authors should fully describe how they have calculated any dpa values they report and endeavour to determine how the dpa were calculated in any previous experiments to which they compare their own data; QC is to be preferred because of its consistency with the NRT standard and previous publications.

We fully agree with the first two conclusions. One could never overstress that models are just models, even the most up to date of them. In the same way, one should always state precisely how damage and dpa are calculated (which damage formalism, $E_d$, etc.). As exemplified above, stating a number of dpa without specifying the details of how they were calculated carries little information.

Nevertheless, we respectfully tend to qualify or moderate the last conclusion about favouring QC over FC. It is true that, as far as elemental solids are concerned, there are some benefits to use QCs. The consistency with the NRT standard is naturally the main one. It allows direct comparison of dpa levels between ion and neutron irradiations. There are therefore legitimate reasons to favour QC over FC for elemental solids when quantitative comparisons damage estimated from neutronics codes are planned. In our opinion, the situation is quite different for multi-component solids. For such materials, QC are physically and qualitatively wrong. The distribution of atomic displacements among the various atomic species as well as the division between electronic and ballistic energy deposition are basically just non-sense with this framework. We therefore believe QC should not be used for such materials. Two things tend to confirm that the designer of SRIM felt the same way. First, nothing appears in the manual about QC in non-elemental materials. Second, looking at the outputs of QC of SRIM, one can realize that the number of created vacancies in QC calculations is not broken down into species but given as the whole. Both points indicate that these calculations were intended only for pure solids. The reason why they were enabled for multi-component materials remains unknown. We contemplated disabling QC for alloys in Iradina, but we chose not to forbid this use of the code, mainly for consistency with what one can do with SRIM. Coming back to Stoller et al. [27], one can note that they deal only with pure solids in their paper, also consistently with implicitly restricting QC to mono-elemental solids.

Finally, we would advise to favour FC over QC in all cases because it more grounded physically and applicable to all materials. QC remains a good option for pure solids in the case of actual quantitative comparisons with neutron irradiations in which damage levels are estimated with the NRT formulas. Consistency with previous studies may also be a reason to give QC damage levels. One may even contemplate using QC in multi-component systems in such particular situations, but we would then highly recommend to give also FC results. The same should be done for elemental solids when one plans to compare between pure or alloyed materials at some point of the study.

As mentioned above, the NRT formula is known the overestimate the damage creation compared to MD. One route of improvement would be to modify the NRT formula (Eq. (5)) and QC to better fit with the actual creation or MD prediction of defects. This is the purpose behind the arc-dpa proposition. The idea is to replace the NRT formula with the so-called arc-dpa formula (arc stands for athermal recombination corrected) [28]. In this formulation, the damage production for kinetic energies larger than 2.5 $E_d$ is rescaled by a factor $\xi(E)$:

$$\xi(E) = \frac{1 - c}{(2E_d/0.8)^b}E^b + c.$$  \hfill (15)

The 0.8 factor is thus replaced by a decreasing function of the PKA energy which starts from 0.8 and converges around 0.3 × 0.8 for iron. This proposition to replace NRT (Eq. (5)) by the arc-dpa (Eqs. (14) and (15)) is still under debate. In our opinion, it faces a couple of issues. First it relies on the existence of MD simulations to fit the additional parameters ($b$ and $c$) entering the arc-dpa formulas. One then has to face the difficulties and uncertainties of such calculations, e.g. dependence on the empirical potential [29], on the detailed or not inclusion of electronic losses [30], etc. These choices do affect the amount of created defects in a non-negligible way. In view of the continuing activity on cascade simulations even in simple materials, obtaining generally accepted reference MD date is not an easy task. Second, there seems to be some pure materials where the arc-dpa formula does not apply, e.g. W [31], where a re-increase of the damage production seems to take place at high PKA energies. In the same way, it has been shown that the arc-dpa formula is not applicable to alloys or multi-elemental solids [32]. Its utility and field of application appears therefore quite restricted. One may however contemplate its implementation as an alternate damage formula replacing NRT in specific cases. There would be no difficulty to do so in Iradina for instance. But then, one would lose the consistency with previous studies or neutronics evaluation based on the NRT formula. To reach a better agreement with actual damage production, one would spoil the main advantage of QC. The situation would naturally change, if the dpa standard was modified which is not the case at present.

4 Conclusion

In this paper, we have presented our implementation of Quick-Calculations in the Iradina code. We obtain a good agreement between Iradina and SRIM for such calculations. We shed light on the weaknesses of QC in the case of non-elemental solids, namely the fact that the damage formulas are applied to supposedly pure elemental solids after each collision of the incoming ion with atoms of the material, thus completely neglecting the multi-component
nature of the material. This drastic and unphysical approximation leads to wrong results in terms of the distribution of damage among the different species as well as the division of losses between ballistic and electronic components. We therefore advocate not to use such QC for multi-component solids. For the sake of consistency, we tend to suggest to favour FC also for elemental materials. Noticeable exceptions are situations where actual quantitative comparisons with damage estimated by neutronics codes or previous QC estimations of damage are in order.

For what concerns alternatives to SRIM, Iradina being equipped with QC and a GUI for 1D irradiation (designed to perform SRIM-like calculations), we believe it can provide a fast and safe way to perform BCA calculations for the ion implantation community.

**Author contribution statement**

Jean-Paul Crocombette coded the quick calculations in Iradina, performed the calculations and wrote the present paper. Christian Van Wambeke programmed the graphical user interface to perform SRIM-like calculations.

**Appendix A: Iradina graphical user interface**

We developed a graphical user interface (Iradina-GUI) to perform 1D (SRIM-like) calculations. The GUI is a set of Python 3.5 script files with QT. It requires a few packages, namely: PyQt5, numpy, matplotlib and pandas. The GUI enables to set-up calculations and run them. It automatically saves the inputs and outputs of Iradina and allows navigating around them. A few basic plots are also possible. Iradina_GUI and the associated Iradina code (Iradina-CODE) are available on source forge [23]. This directory is a sub section of the general Iradina source forge project [22] maintained commonly by the present author and the original Iradina team (Christian Borchel and Carsten Ronning).

Packages are available for windows and linux, together with installation instructions. For windows, the package contains everything to make the program run, including all the packages mentioned above. For linux, a few basic preliminary steps are necessary to include these packages independently from Iradina. The GUI as well as Iradina code itself are works in progress and they should be enhanced in the forthcoming years. The author welcomes comments and suggestions from users.

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**Cite this article as:** Jean-Paul Crocombette, Christian Van Wambeke, Quick calculation of damage for ion irradiation: implementation in Iradina and comparisons to SRIM, EPJ Nuclear Sci. Technol. 5, 7 (2019)