Polarization spectral synthesis for Type Ia supernova explosion models

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ABSTRACT
We present a Monte Carlo radiative transfer technique for calculating synthetic spectropolarimetry for multidimensional supernova explosion models. The approach utilizes ‘virtual-packets’ that are generated during the propagation of the Monte Carlo quanta and used to compute synthetic observables for specific observer orientations. Compared to extracting synthetic observables by direct binning of emergent Monte Carlo quanta, this virtual-packet approach leads to a substantial reduction in the Monte Carlo noise. This is not only vital for calculating synthetic spectropolarimetry (since the degree of polarization is typically very small) but also useful for calculations of light curves and spectra. We first validate our approach via application of an idealized test code to simple geometries. We then describe its implementation in the Monte Carlo radiative transfer code ARTIS and present test calculations for simple models for Type Ia supernovae. Specifically, we use the well-known one-dimensional W7 model to verify that our scheme can accurately recover zero polarization from a spherical model, and to demonstrate the reduction in Monte Carlo noise compared to a simple packet-binning approach. To investigate the impact of aspherical ejecta on the polarization spectra, we then use ARTIS to calculate synthetic observables for prolate and oblate ellipsoidal models with Type Ia supernova compositions.

Key words: polarization – radiative transfer – methods: numerical – supernovae: general.

1 INTRODUCTION
Type Ia supernovae (SNe Ia) are generally believed to be thermonuclear explosions of carbon–oxygen white dwarfs (see e.g. Röpke et al. 2011; Hillebrandt et al. 2013, for reviews). However, answers to the questions of how and why the explosion is triggered remain unclear. Most of the established theoretical models involve close binary systems, but we still do not know if the companion star is a second white dwarf (double degenerate system; Iben & Tutukov 1984; Webbink 1984) or a non-degenerate star (single degenerate system; Whelan & Iben 1973), and whether the explosion is triggered when an accreting white dwarf approaches the Chandrasekhar limit or via some other process.

For Chandrasekhar-mass models, neither a pure deflagration nor a pure detonation model is able to fully account for the properties observed in SNe Ia: the former leads to strong turbulence and buoyancy resulting in fingers of nickel and carbon–oxygen at all ejecta velocities (Gamezo et al. 2003; Röpke et al. 2006; Jordan et al. 2012b; Ma et al. 2013; Fink et al. 2014), while the latter fails to produce intermediate-mass elements (Arnett 1969). However, an interplay of these two models, so-called delayed-detonation models, remains promising for providing a good match to data. Possibilities include spontaneous deflagration to detonation transition models (Khokhlov 1991; Höflich, Khokhlov & Wheeler 1995; Höflich et al. 1996; Höflich & Khokhlov 1996; Kasen, Röpke & Woosley 2009; Blondin et al. 2013) and the gravitationally confined detonation model (Plewa, Calder & Lamb 2004; Jordan et al. 2008, 2012a). For non-Chandrasekhar-mass models, in which the accreting white dwarf may explode without approaching the Chandrasekhar limit, viable mechanisms are the detonation of helium layers on the surface of the accreting white dwarf (the double detonation model; see Nomoto 1980; Woosley, Weaver & Taam 1980; Livne 1990; Woosley & Weaver 1994; Fink, Hillebrandt & Röpke 2007; Shen & Bildsten 2009; Fink et al. 2010; Woosley & Kasen 2011; Moll & Woosley 2013) and the violent mergers of two white dwarfs (Pakmor et al. 2010, 2012, 2013; Moll et al. 2014; Raskin et al. 2014). Alternative models include the explosion of white dwarf merger remnants (Benz et al. 1990; Van Kerkwijk, Chang & Justham 2010; Shen et al. 2012; Kashyap et al. 2015), the head-on collisions of white dwarfs (Rosswog et al. 2009), possibly induced in triple systems (Kushnir et al. 2013), or the merger of a white dwarf with the hot core of an asymptotic giant branch star (Soker, García-Berro & Althaus 2014).

There are a variety of ways to attempt to determine which of the proposed progenitor/explosion channels really occur (see e.g. Maoz, Mannucci & Nelemans 2014, for a review). One approach is...
to perform explosion simulations with associated radiative transfer calculations and compare their predictions to data. Such work is well established (e.g. Höflich, Mueller & Khokhlov 1993; Hauschildt & Baron 1999; Kasen, Thomas & Nugent 2006; Kromer & Sim 2009; Blondin et al. 2013; Dessart et al. 2014; Wollaeger & Van Rossum 2014) and it is now possible to compute synthetic observables from multidimensional models for a variety of explosion scenarios. Although state-of-the-art explosion simulations allow us to capture considerable complexity in SN Ia models (e.g. turbulence), degeneracies between models remain and make unambiguous interpretation difficult, even for the best observed nearby examples (Röpke et al. 2012).

One potentially powerful discriminant is the geometry, which can be quite different between models and depend on the nature of the progenitor and explosion mechanism. From the observational side, both nebular phase spectroscopy (Gerardy et al. 2007; Maeda et al. 2010) and spectropolarimetric observations (see Wang & Wheeler 2008, for a review) provide evidence that SNe Ia are not perfectly spherically symmetric. Continuum polarization is typically quite low (0.2–0.3 per cent) in normal SNe Ia prior to optical maximum, pointing towards very small departures from global spherical symmetry; significant polarization is, however, found across the line profiles of spectral features associated with intermediate-mass elements (calcium, silicon, sulphur and magnesium, but not oxygen; see e.g. Wang & Wheeler 2008 and reference therein), suggesting that asymmetries in the element distribution are present. Other subclasses of SNe Ia seem to display some peculiarities: subluminous SNe Ia show higher continuum polarization levels (0.3–0.8 per cent; Howell et al. 2001; Patat et al. 2012), whereas high-velocity SNe Ia show stronger line polarization (~2 per cent; Leonard et al. 2005; Wang et al. 2006).

Asymmetric ejecta for SNe Ia are also predicted by many multidimensional explosion models, although with different degrees and types of asymmetry. For instance, the delayed-detonation model predicts that the ejecta can be quasi-spherical on large scales but with complex substructures (in both density and composition; Seitenzahl et al. 2013; Sim et al. 2013), whereas simulations of violent white dwarf mergers predict departures from spherical symmetry on large angular scales (Pakmor et al. 2010; Moll et al. 2014; Raskin et al. 2014). These differences in ejecta geometry between models have led to suggested connections with observed objects. For example, Patat et al. (2012) proposed that the subluminous SN 2005ke might be explained by an explosion of a rotating white dwarf or a double degenerate merger. It has also been suggested that several models could be ruled out because they yield explosions that are too aspherical and therefore inconsistent with continuum polarization measurements for normal SNe Ia. For instance, Maund et al. (2013) have claimed that the low continuum polarization and significant line polarization observed in SN 2012fr are consistent with delayed-detonation models but inconsistent with deflagration models or violent white dwarf mergers.

However, interpreting polarization data and quantifying arguments about its implications for models is difficult because estimating the degree of polarization expected for a complex ejecta morphology (as provided e.g. by multidimensional explosion simulations) is not trivial: polarization depends on the opacity distributions in a complex way (Höflich 1991; Dessart & Hillier 2011). To date, spectropolarimetric data of SNe Ia have often been interpreted by comparing the observed polarization levels with predictions from toy models with idealized configurations for the ejecta, e.g. ellipsoidal and clumped shell models or spherical shells with a hole or toroid (Howell et al. 2001; Kasen et al. 2003, 2004; Patat et al. 2012). These idealized geometries are well suited for building intuition and establishing the framework for interpreting polarization data. However, quantitative comparisons between predictions of multidimensional explosion models and data require that polarization calculations are made for the complete density/composition distributions predicted by hydrodynamic simulations. Such calculations make it possible to quantitatively compare the predictions of models to data (and each other) and assess the extent to which their geometries can really be distinguished via polarization.

Here we present a polarization scheme recently implemented in the three-dimensional, time-dependent Monte Carlo radiative transfer code Applied Radiative Transfer In Supernovae (ARTIS; Sim 2007; Kromer & Sim 2009). The scheme involves two parts, first an implementation of Stokes parameters for the Monte Carlo quanta and secondly the development of techniques to reduce the Monte Carlo noise in the emergent synthetic observables. This is particularly important when we aim to extract very weak polarization signals (low percentage levels are observed in SNe Ia), but also useful for total flux spectra and light curves. Although in this work we focus on the development, implementation and validation of the method using one-dimensional and two-dimensional models, our particular technique is well suited to exploit the multidimensional capability of ARTIS and therefore to be applied to three-dimensional explosion models. Details of the methodology used are given in Section 2, and the polarization scheme is validated via an idealized test code in Section 3. We then present first results from the implementation in ARTIS, including testing with the one-dimensional W7 model (Nomoto, Thielemann & Yokoi 1984; Iwamoto et al. 1999) and two-dimensional ellipsoidal toy models (Section 4). We summarize and draw conclusions in Section 5.

2 METHOD

The polarization of a beam of radiation is characterized by the four-dimensional Stokes vector $S = (I, Q, U, V)$. The first component, $I$, gives the total intensity, $Q$ and $U$ measure the degree of linear polarization and $V$ of circular polarization. Since circular polarization has never been observed in SNe Ia, and the radiative transfer calculations for circular and linear polarization can be decoupled in scattering atmosphere in the absence of magnetic fields (Chandrasekhar 1960), here we neglect the $V$ component. The Stokes vector is defined in the plane orthogonal to the direction $\mathbf{n}$ in which the radiation propagates. To define the Stokes parameters for linearly polarized radiation, we introduce two reference axes $l$ and $r$ so that $l$ lies in the meridian plane (plane defined by $\mathbf{n}$ and the polar axis $z$) and $\mathbf{n} = r \times l$ (see Fig. 1). With this convention, $Q$ is defined as the difference between intensity $I_l$ with electric field oscillating along $l$ and intensity $I_r$ with electric field oscillating along $r$; $U$ is the equivalent difference in intensities with the reference axes $l$ and $r$ counterclockwise rotated by $45^\circ$ (as viewed looking antiparallel to $\mathbf{n}$) to give $a$ and $b$. The resulting Stokes vector $S$ can be expressed as

$$S = \begin{pmatrix} I \\ Q \\ U \end{pmatrix} = \begin{pmatrix} I_l + I_r \\ I_l - I_r \\ I_a - I_b \end{pmatrix} = \begin{pmatrix} \uparrow + \leftrightarrow \\ \downarrow - \leftrightarrow \\ \alpha - \beta \end{pmatrix},$$

or in terms of a dimensionless Stokes vector, $s$:

$$s = \frac{S}{I} = \begin{pmatrix} 1 \\ q \\ u \end{pmatrix},$$

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The meridian plane coordinate system adopted in the Monte Carlo code. The Stokes vector is defined in the plane orthogonal to the direction of propagation, \( \mathbf{n}, \mathbf{Q} \) is defined as the intensity difference between two perpendicular reference axes, \( \mathbf{l} \) and \( \mathbf{r} \), whereas \( \mathbf{U} \) is the equivalent difference with \( \mathbf{l} \) and \( \mathbf{r} \) counterclockwise rotated by 45° (viewed antiparallel to \( \mathbf{n} \)).

The polarization fraction, \( p \), and the position angle, \( \chi \), of a beam are related to the Stokes parameters by

\[
p = \frac{\sqrt{\mathbf{Q}^2 + \mathbf{U}^2}}{\mathbf{I}} = \sqrt{q^2 + u^2},
\]

\[
\chi = \frac{1}{2} \tan^{-1} \left( \frac{\mathbf{U}}{\mathbf{Q}} \right) = \frac{1}{2} \tan^{-1} \left( \frac{u}{q} \right),
\]

where \( \chi \) is the angle between the electric field orientation and the reference axis \( \mathbf{l} \). Spherically symmetric geometries are characterized by null polarization since every contribution is cancelled by an orthogonal contribution one quadrant away, whereas aspherical geometries may lead to a polarization signal due to non-perfect cancellation of the Stokes vectors (see also Kasen et al. 2003 and discussion in Section 4).

### 2.1 Propagation

In this section, we adopt the terminology introduced by Lucy (2002, 2005) and discuss the general scheme used to include polarization in our radiative transfer code. The calculations we present use Monte Carlo methods: the radiative transfer problem is solved by simulating the propagation of Monte Carlo quanta (packets of identical photons) through an expanding medium. The propagation of an \( r \)-packet (monochromatic packet of ultraviolet–optical–infrared radiation) is followed through the ejecta in the rest frame (rf) and stopped by interactions with matter (which are treated in the comoving frame, cmf). For ultraviolet–optical–infrared radiation, the code currently accounts for both line opacity (treated in the Sobolev approximation; Sobolev 1960) and continuum opacity due to electron scattering, bound–free and free–free absorption. To choose when continuum and interaction events occur we use the method outlined by Mazzali & Lucy (1993). Once a random optical depth \( \tau \), drawn, we determine the trajectory point at which the \( r \)-packet interacts with the next line (using the Sobolev approximation): if the continuum opacity accumulated up to that point is greater than \( \tau \), a continuum absorption is selected; if instead the sum of continuum and line opacity is greater than \( \tau \), a line event occurs; otherwise, the process is repeated for the next line with which the packet comes into resonance. In an electron scattering event, the \( r \)-packet keeps the same cmf frequency and is assigned a new direction of propagation. For all other interactions (line absorption, free–free absorption and bound–free absorption) either a \( k \)-packet (packet of thermal kinetic energy) or an \( i \)-packet (packet of excitation/ionization energy) is activated and then processed according to the scheme proposed by Lucy (2002) as described by Kromer & Sim (2009).

Our polarization scheme adopts a method similar to that proposed by Lucy (2005) and already implemented by Kasen et al. (2006). Polarization is introduced by assigning a Stokes vector to each \( r \)-packet. When an interaction occurs, the Stokes parameters are transformed through the following sequence of steps: first, we transform the incoming Stokes vector \( \mathbf{s} \) in the rf to \( \mathbf{s}' \) in the cmf.\(^1\) The plane in which the electric field oscillates changes as a result of the aberration of the direction \( \mathbf{n} \) to

\[
\mathbf{n}' = \frac{\mathbf{n} - \frac{1}{2} \mathbf{r} \left( \gamma - \frac{\chi^2}{\gamma - 1} \mathbf{r} \right)}{\gamma (1 - \frac{\chi^2}{\gamma^2})},
\]

where \( c \) is the speed of light, \( \gamma \) the Lorentz factor and \( \mathbf{r} \) the local velocity of the ejecta (Castor 1972). To derive how the Stokes parameters change under the Lorentz transformation, we introduce a unit vector \( \mathbf{\hat{e}} \) that describes the orientation of the net electric field in the rf,

\[
\mathbf{\hat{e}} = (\cos \chi) \mathbf{l} - (\sin \chi) \mathbf{r},
\]

with the angle \( \chi \) between \( \mathbf{\hat{e}} \) and \( \mathbf{l} \) computed from the incoming Stokes vector following equation (4). From this, it is easy to obtain a cmf Stokes vector representation that is relative to axes \( \mathbf{l}' \) and \( \mathbf{r}' \) (defined by \( \mathbf{n}' \)), making use of the fact that the polarization, \( p \), is invariant (Cocke & Holm 1972):

\[
q_i' = p_i \cos 2\chi', \quad u_i' = p_i \sin 2\chi'.
\]

Following transformation to the cmf, the Stokes parameters can be updated in accordance with the physical interaction that occurs. For bound–bound, bound–free and free–free absorptions, we first activate either an \( i \)-packet or a \( k \)-packet, as outlined in Kromer & Sim (2009) using the machinery described by Lucy (2002), and then convert this to a new \( r \)-packet. For all these processes, the \( r \)-packet is assumed to retain no information on polarization and is re-emitted in a random direction with zero polarization:\(^2\)

\[
q_i = 0, \quad u_i = 0.
\]

If instead the \( r \)-packet undergoes electron scattering, we follow the scheme introduced by Chandrasekhar (1960) and discussed in terms of a Monte Carlo implementation by Code & Whitney (1995) and Whitney (2011). A scattering angle \( \Theta \) is properly sampled (see

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\(^1\) In the following, cmf quantities are denoted with a prime, whereas unprimed quantities refer to the rf.

\(^2\) Hamilton (1947) has shown that the angular distribution in resonant line scattering can be expressed as the sum of a dipole and an isotropic term (with relative contribution depending on the quantum numbers of upper and lower state of the line). Although polarization may arise in cases where the dipole is the dominant term, its magnitude is typically lower than that from electron scattering and thus can be regarded as a second-order effect (Jeffery 1989, 1991).
The geometry adopted for electron scattering in the coordinate system introduced by Chandrasekhar (1960) with the additional reference axes $I'_i$ and $I'_i$ defined in the text. A packet moving along $n'_i$ is scattered through an angle $\Theta$ to a new direction $n_i$. The reference axis $I'_i$ (and accordingly the Stokes vector) are rotated through an angle $i_1$ into the scattering plane (in grey) and through an angle $i_2$ after scattering. Below), a new direction $n'_i$ is computed and the Stokes vector is transformed via the scattering matrix:

$$A(\Theta) = \frac{3}{4} \begin{pmatrix}
\cos^2 \Theta + 1 & \cos^2 \Theta - 1 & 0 \\
\cos^2 \Theta - 1 & \cos^2 \Theta + 1 & 0 \\
0 & 0 & 2 \cos \Theta
\end{pmatrix}. \quad (9)$$

After applying the scattering matrix, the dimensionless Stokes vector is normalized so that its first component is equal to 1. As shown in Fig. 2, since $A(\Theta)$ is defined in the scattering plane, the Stokes vector is first rotated into this plane and then rotated back to the meridian frame after the scattering matrix $A(\Theta)$ has been applied, i.e.

$$s'_i = R(\pi - i_2)A(\Theta)R(-i_1)s'_i, \quad (10)$$

where

$$R(\phi) = \begin{pmatrix}
1 & 0 & 0 \\
0 & \cos 2\phi & \sin 2\phi \\
0 & -\sin 2\phi & \cos 2\phi
\end{pmatrix}. \quad (11)$$

is the matrix to rotate the Stokes vector by an angle $\phi$ clockwise when facing the source. The rotation angles $i_1$ and $i_2$ are computed with the convention that $q'_i = 1$ represents an electric field oscillating in the scattering plane. The scattering angle $\Theta$ is chosen by sampling the probability distribution,

$$P(\Theta, i_1) = \frac{3}{4} \left[ (\cos^2 \Theta + 1) + (\cos^2 \Theta - 1)(q'_i \cos 2i_1 - u'_i \sin 2i_1) \right], \quad (12)$$

from equation (10) and using a rejection technique (Code & Whitney 1995).

Finally the Lorentz transformation procedure (see above) with $\nu \rightarrow -\nu$ is used to transform the Stokes vector $S'_i$ to $S_i$.

### 2.2 Spectrum extraction techniques

Monte Carlo methods have been exploited in many radiative transfer calculations. These methods have proven particularly useful for multidimensional problems since the Monte Carlo algorithm can be easily implemented for arbitrary geometry and scales very well for use on massively parallel computing systems. The main drawback of Monte Carlo methods is their stochastic nature, which leads to solutions that are affected by Monte Carlo noise. Therefore, it is important to attempt to optimize Monte Carlo methods, particularly when we want to extract very weak signals (e.g. polarization) from the simulations. In this study we compare three different methods for extracting synthetic observables from our simulations, with the aim of selecting the most suitable to synthesize spectra with low Monte Carlo noise. In Section 2.2.1 we present a simple direct counting approach, while in Sections 2.2.2 and 2.2.3 we explore alternative techniques inspired by Lucy (1999, 2005) and recently implemented by Kerzendorf & Sim (2014).

#### 2.2.1 Direct counting technique

In the direct counting technique (DCT), Monte Carlo quanta are followed along their trajectories and their Stokes parameters updated at each interaction, following the procedure outlined in Section 2.1. Packets that reach the outer boundary are collected in bins according to their final direction $n$ and frequency $\nu$ and the resulting spectra are computed as

$$\begin{pmatrix} I \\ Q \\ U \end{pmatrix} = \frac{\epsilon}{\Delta \nu \Delta t} \sum s_i, \quad (13)$$

where $s_i$ is the dimensionless Stokes vector of the escaping $r$-packet, $\epsilon$ its rf energy, $r$ the distance of the observer from the system and the sum is performed over all the $r$-packets that escaped in the selected angular bin, time interval $[\tau - \Delta \tau/2, \tau + \Delta \tau/2]$ and frequency range $[\nu - \Delta \nu/2, \nu + \Delta \nu/2]$.

The direct counting approach provides a simple way of computing polarization spectra for different viewing angles, i.e. for different observers. However, the need to average contributions from packets that escape in the same angular bin but with different angles inevitably leads to an approximate result: if the number of angular bins is too small, summing contributions from different angles will produce a poor estimate of the observables seen by a single observer; if instead too many angular bins are used, the number of packets escaping per bin becomes small, leading to high Monte Carlo noise.

#### 2.2.2 Event-based technique

In this approach (event-based technique, EBT) we still follow the propagation of Monte Carlo packets exactly as before. However, whenever an $r$-packet interaction occurs the propagation is suspended, and a ‘virtual’ packet ($v$-packet; Kerzendorf & Sim 2014) is created and handled as described below. Once the $v$-packet calculation is completed, the propagation of the original $r$-packet is resumed and this process repeated for every following interaction. This approach is similar to, and inspired by, that used by Knigge, Woods & Drew (1995) and Long & Knigge (2002).

When a $v$-packet is created, it is always launched in the direction of a selected observer, $n_{\text{obs}}$, and has cmf frequency and energy set equal to those of the $r$-packet at the point of creation (see Fig. 3). Since the $v$-packet is forced to go to the observer, the scattering angle is now not determined by sampling either an isotropic distribution ($r$-packet or $k$-packet deactivation) or equation (12) (electron scattering).
Here the sum is performed over all the line opacities ($\tau_{\text{ob}}$) encountered by the $v$-packet on its trajectory to the boundary and the continuum opacity, $\tau_{\text{cont}}$, is computed as an integral of the continuum attenuation coefficient, $k_{\text{cont}}$, over trajectory length:

$$\tau_{\text{cont}} = \int k_{\text{cont}} \, ds.$$  

The advantage of the $v$-packet technique is that it allows us to compute spectra and light curves for any specific viewing angle, avoiding the need to average contributions from different angles in the same bin. Unlike in the direct counting approach, where an $r$-packet makes a single contribution to the emergent spectrum, the final spectra in the EBT also contain (appropriately weighted) information derived from every interaction that the $r$-packet undergoes (see Fig. 3). For these reasons, we expect this technique to produce spectra and light curves with lower Monte Carlo noise. However, the need of creating and handling $v$-packets introduces a computational overhead that could make the EBT less efficient than the DCT. Such possibilities are explored quantitatively in Section 3.2.

2.2.3 Trajectory-based technique

In the EBT described above, we used all the $r$-packets to provide us with an ensemble of physical events, and then for each event in the ensemble we computed the probability of it giving rise to a photon escaping to the observer. In the third technique (trajectory-based technique, TBT) we instead obtain from the $r$-packets an ensemble of photon trajectories, which can be taken as a discrete representation of the radiation field in the simulation. We can then estimate observables by summing over this ensemble of trajectories, computing for each the probability that interactions of radiation on that path could have given rise to photons escaping to the observer.

This summation is achieved by generating a $v$-packet for each trajectory path $\Delta l$ (including those terminated by numerical events, e.g. packets crossing grid cell boundaries) and sending it towards the observer at $\mathbf{n}_{\text{obs}}$ (see Fig. 3). The $v$-packet contribution to the emergent spectrum is first weighted by the probability per unit solid angle ($dP/d\Omega_{\text{TBT}}$) that photons on the path $\Delta l$ could have undergone an interaction that gave rise to re-emission/scattering towards the observer. As with the EBT, we must also account for the probability of any scattered/emitted radiation reaching the observer via a suitable mean exponential factor, $(e^{-\tau_{\text{esc}}})$. Thus, in the TBT, we compute synthetic observables via

$$\left( \begin{array}{c} I \\ Q \\ U \end{array} \right) = \left( \begin{array}{c} \frac{\epsilon}{\Delta l} \frac{dP}{d\Omega} |_{\text{TBT}} (e^{-\tau_{\text{esc}}}) \\ \frac{\epsilon}{\Delta l} \frac{dP}{d\Omega} |_{\text{TBT}} (e^{-\tau_{\text{esc}}}) \end{array} \right),$$

where the summation is again over $v$-packets in the selected frequency and time bins. Formally, the exponential weight factor of a $v$-packet in the TBT should be computed as

$$\langle e^{-\tau_{\text{esc}}} \rangle = \frac{\int_{l_{\text{start}}}^{l_{\text{end}}} e^{-\tau_{\text{esc}}(l)} \, dl}{\Delta l},$$

where the integral runs over the trajectory path $\Delta l$. To first order, however, this can be approximated by generating the $v$-packet at the midpoint of $\Delta l$ and computing the exponential factor from its flight (as described in Section 2.2.2), i.e.

$$\langle e^{-\tau_{\text{esc}}} \rangle = e^{-\tau_{\text{esc}}(l+\Delta l/2)}.$$

In principle, $dP/d\Omega_{\text{TBT}}$ can be formulated to account for all (effective) scattering/fluorescence processes. However, since we are
primarily interested in studying contributions to the emergent polarization spectrum, we focus only on electron scattering, for which

\[
\frac{dP}{d\Omega}_{\text{TBT}} = \frac{1}{4\pi} P(\theta, i_1) k_{\text{sc}} \Delta l, \tag{22}
\]

where \(k_{\text{sc}}\) is the scattering attenuation coefficient.

A key difference between the EBT and the TBT is that, in the latter, every trajectory element of the \(r\)-packets contributes to the synthetic observables, whereas, in the former, only physical interaction events contribute. For instance, in the limit of optically thin ejecta many more \(v\)-packets would contribute to the emergent spectrum in the TBT compared to the EBT (see Fig. 3). However, a drawback of the TBT is that \(\tau_{\text{esc}}(l + \Delta l/2)\) should describe the mean probability of escape for points along the trajectory element (see equation 21), rather than the exact probability of escape from the interaction point, as in the EBT. For an \(r\)-packet trajectory with moderate optical depth (\(\tau \gtrsim 1\)), this approximation may lead to a poor estimate of the observables. Breaking the trajectory into smaller paths (with \(\Delta \tau \ll 1\)) and generating \(v\)-packets at each midpoint may be required, slowing down the code and making the TBT less efficient than the EBT (see Section 3.2).

3 TEST CODE

In the following we present a simple test code, with the aim of validating our polarization scheme (Section 3.1) and selecting the most suitable of the techniques described in Section 2.2 to synthesize spectra with low Monte Carlo noise (Section 3.2). In this code, packets are generated with null polarization in a small inner sphere (to mimic a point source) and then allowed to propagate into an envelope where either interactions with electrons or continuum absorptions can occur. Here time evolution for the ejecta is neglected.

3.1 Polarization scheme validation

To validate the polarization scheme, we first focus our attention on the DCT and choose to reproduce a simple configuration described by Hillier (1994). As shown in Fig. 4, a point source is surrounded by a detached spherical shell with inner radius \(R_{\text{min}} = 2.0\) and outer radius \(R_{\text{max}} = 30.0R_{\text{min}}\), with a prolate density distribution \(N_e(r, \beta)\) such that

\[
\sigma_e N_e(r, \beta) = \chi_0 \left( \frac{R_{\text{min}}}{r} \right)^2 (1 + 10 \cos^2 \beta), \tag{23}
\]

where \(\sigma_e\) is the Thomson cross-section and \(r\) and \(\beta\) express the radius and the polar angle inside the envelope. The \(\chi_0\) parameter is related to the solid-angle averaged (from inner to outer boundary) optical depth, \(\tau_{\text{ave}} = 2.888\chi_0\), and can be varied to investigate the impact of different scattering optical depth on the polarization signal. Neglecting absorption and assuming a pure electron scattering envelope, the continuum polarization as a function of \(\chi_0\) is shown in Fig. 5 for four different viewing angles \(i\) (22.5°, 45°, 67.5° and 90°). The agreement between our predicted values and the expected curves from Hillier (1994) is encouraging. We also carried out calculations in which we include continuum absorption opacity. These show good agreement with the predicted dependence of the continuum polarization on the albedo (ratio of the scattering to the total opacity, see Fig. 6).

3.2 Comparison between different techniques

A convenient means to compare the three techniques for extracting observables outlined in Section 2.2 is by studying their accuracy in reproducing continuum polarization for a given configuration. We did this by repeatedly running our test code a number \((N_{\text{sim}} = 500)\) of times for each technique (with different random number seeds determined by the wall clock time) and comparing the distributions of polarization values obtained using each method. For these experiments, we chose a configuration in which a point source illuminates...
a surrounding atmosphere, chosen to be a constant density oblate ellipsoid with axis ratio of two, i.e.

\[ \frac{x^2}{a^2} + \frac{y^2}{b^2} + \frac{z^2}{c^2} = 1, \quad a = b = 2c. \]  

(24)

In all the simulations, \(10^6\) packets have been created, a pure scattering atmosphere has been assumed (i.e. no line opacity and albedo equal to 1) and the scattering coefficient has been set to \(k_{sc} = 1/a\) and kept constant throughout the ejecta. As mentioned in Section 2.2.1, continuum polarization levels estimated from the DCT may be either inaccurate or have large uncertainties depending on whether the number of angular bins is small or large, respectively. We found that a value of 51 angular bins provides a reasonable compromise and we therefore adopt this in the DCT calculations. The results of the comparison between the three techniques are shown in Fig. 7 and reported in Table 1.

For an observer along the \(y\)-axis (Fig. 7, left-hand panel), the projection of the model along the line-of-sight is circular and we expect to find null polarization (see above). The \(Q\) and \(U\) values in every technique are indeed consistent with zero but, because of Monte Carlo noise, a distribution of values is obtained. The width of this distribution provides a convenient quantification of the Monte Carlo noise introduced by each method. As expected, the standard deviations obtained with the EBT and TBT methods are smaller than with the DCT method, by factors of \(\sim 6.6\) and \(\sim 8.6\), respectively. Given that Monte Carlo noise is expected to scale with the square root of the number of packets, the DCT could reach the same signal-to-noise ratio as the EBT (TBT) with a factor of \(\sim 45\) (\(\sim 75\)) more packets. Although the \(v\)-packet routine introduces computational overheads in the EBT and TBT (see Table 1), the direct counting approach would still be a factor of \(\sim 7.5\) (\(\sim 2.5\)) slower than the \(v\)-packets technique. We also note that, even with the same signal-to-noise ratio, the DCT would be less accurate in predicting polarization values because of the need to average contributions

| System          | \(\bar{Q} \pm \sigma_Q\) (per cent) | \(\bar{U} \pm \sigma_U\) (per cent) | \(\bar{t}\) (s) |
|-----------------|-------------------------------------|-------------------------------------|---------------|
| Circular        | DCT 0.16 ± 0.23 0.00 ± 0.24 3.9     | EBT 0.00 ± 0.04 0.00 ± 0.04 22.3   | TBT 0.00 ± 0.03 0.00 ± 0.03 123.0 |
| Elliptical      | DCT 3.81 ± 0.28 −0.02 ± 0.29 3.9    | EBT 3.82 ± 0.04 0.00 ± 0.04 26.3   | TBT 3.81 ± 0.03 0.00 ± 0.03 144.5 |

Table 1. Average values and standard deviations of the distributions of \(Q\) and \(U\) values predicted on 500 simulations by the DCT, EBT and TBT. The system is an oblate ellipsoid with axis ratio of two viewed along the \(z\)-axis (circular projection) and along the \(x\)-axis (elliptical projection). The averaged runtime, \(\bar{t}\), is reported for each distribution.

Figure 6. Continuum polarization as a function of the albedo (black points) for the set-up described in Section 3.1, together with the predicted curve (dashed line) from Hillier (1994). Here the observer’s inclination is 90° and \(\chi_0 = 0\). The Monte Carlo noise error bars are not shown since they are smaller than the symbol sizes.

Figure 7. \(Q\) and \(U\) continuum polarization distributions of \(N_{sim} = 500\) runs with the DCT (top panels), the EBT (middle panels) and the TBT (lower panels). The adopted geometry is an oblate ellipsoid with axis ratio of two, as described in Section 3.2. The system is viewed along the minor axis (circular projection, left-hand panel) and the major axis (elliptical projection, right-hand panel). For each distribution, a solid vertical line indicates the average value, \(\bar{V}\), and the grey shaded area marks ±1 standard deviation, \(\sigma\). The average values and the standard deviations of each plot are reported in Table 1 together with the average runtimes, \(\bar{t}\).
from different viewing angles (via angular binning): indeed, closer inspection shows that the $Q$ distribution for the DCT is shifted towards positive values (in the binning approach contributions to the spectra come from $r$-packets escaping close to, but not exactly along, the $z$ direction and thus the average value of $Q$ is slightly positive).

If viewed down the $x$-axis (right-hand panel), the projection becomes an ellipse and thus we expect to see a polarization signal. The three techniques agree in reproducing a continuum polarization of $p \approx 3.8$ per cent but, again, the DCT gives a much broader distribution of values, indicating that it is more severely affected by noise in the simulation. In order to reach the same standard error of the mean of the EBT (TBT), the DCT would require a factor of $\sim 50$ ($\sim 100$) more packets, making the total runtime a factor of $\sim 7.5$ ($\sim 3$) longer than the $v$-packets scheme.

This simple comparison shows that the $v$-packet approaches are more precise in estimating polarization, allowing us to reach a given Monte Carlo noise with many fewer Monte Carlo quanta (and substantially shorter runtimes) than the simple DCT would require. As already anticipated in Section 2.2.3, the EBT is indeed more efficient than the TBT because the runtime for the latter is limited by the need of breaking $r$-packet trajectories with moderate optical depth ($\tau \gtrsim 1$) into smaller paths (with $\Delta \tau \ll 1$), in order to give accurate results for $\tau_{\text{esc}}$. We note that, although we have carried out polarization tests here, this improvement in Monte Carlo noise could also be exploited for extracting high-quality observables of any sort.

4 ARTIS

In this section we describe the implementation of our polarization scheme into the three-dimensional, time-dependent radiative transfer code ARTIS (Kromer & Sim 2009) and test it for one-dimensional and two-dimensional models. Section 4.1 outlines the implementation of the different techniques described in Section 2.2. In Section 4.2 we test the code using the one-dimensional W7 explosion model (Nomoto et al. 1984; Iwamoto et al. 1999) and check the accuracy of the $v$-packet technique in computing spectra and reproducing continuum polarization consistent with zero. Finally, in Section 4.3 we apply the new version of the code to two-dimensional ellipsoidal models to investigate the impact of simple aspherical geometries on line and continuum polarization for different viewing angles, and compare our results to those of similar studies made using other codes.

4.1 Implementation

Polarization is implemented into ARTIS by assigning a Stokes vector to each $r$-packet and by transforming this according to the physical process the packet undergoes (see Section 2.1). For the DCT, the same binning approach already used in the code for spectra and light curves is extended to compute polarization spectra.

As mentioned in Section 2.2.3, the $v$-packet TBT should yield spectra with lower Monte Carlo noise compared to the EBT since contributions to the polarization spectra come from every event in the $r$-packet histories, including both physical interactions and numerical events (e.g. packets crossing grid cell boundaries). However, accurate results from the TBT require that care is taken in the calculation of $\tau_{\text{esc}}$, which can introduce a large computational overhead for complicated opacity distributions. Indeed, our test calculations (Section 3.2) suggested that this additional computational overhead can ultimately make the TBT less efficient than the EBT.

Consequently, here we have chosen to implement a $v$-packet routine using the EBT that can be used to compute synthetic observables from ARTIS.

The $v$-packet routine allows us to compute flux and polarization spectra for multiple observers, simply by using a loop to generate $v$-packets over a set of different viewing angles. Several input parameters can be chosen to optimize performance in the calculations. First, the calculation of the optical depth $\tau_{\text{esc}}$, see equation (17), can be stopped when the $v$-packet reaches a maximum value $\tau_{\text{esc}}^{\text{max}}$. $v$-packets with high optical depth to the boundary would make vanishingly small contributions to the final spectrum (because of the exponential factor, see equation 15) and can thus be neglected. Since the computation cost of the $v$-packet methods is dominated by the calculation of $\tau_{\text{esc}}$, the runtime is strongly affected by this parameter. We have carried out test calculations to verify that setting $\tau_{\text{esc}}^{\text{max}} = 10$ and neglecting $v$-packets with higher optical depth does not affect the final result, and adopted this as our default value for all the calculations presented here. Our implementation also includes the option to generate $v$-packets only in a selected spectral interval. Because spectropolarimetric observations usually cover the optical region of the spectrum, our default wavelength range is $3500–10 000\,\text{Å}$. Given that much of the runtime of the code can be consumed in computing $\tau_{\text{esc}}$ for packets in the bluer regions, which easily reach $\tau_{\text{esc}}^{\text{max}}$ because of strong iron-line blanketing, this particular cut in wavelength speeds up the calculations by a factor of $\sim 4$ compared to calculations with no wavelength cut. Finally, the $v$-packet routine can be switched on or off for time steps as chosen by the user (note that the activation or deactivation of $v$-packets has no effect on the $r$-packet propagation).

4.2 W7 model

Although much effort has been recently directed at developing multidimensional explosion models (Rosswog et al. 2009; Jordan et al. 2012b; Kushner et al. 2013; Fink et al. 2014; Moll et al. 2014), the one-dimensional parametrized deflagration model W7 is still widely used since its composition and structure provide reasonable agreement with observations of (‘normal’) SNe Ia (Kasen et al. 2006; Kromer & Sim 2009; Jack, Hauschildt & Baron 2011; Gall et al. 2012; van Rossum 2012). Here we calculate ARTIS flux and polarization spectra for this model aiming to (i) compare the accuracy of the DCT and the EBT in reproducing flux spectra at different epochs; (ii) test our polarization implementation on a spherically symmetric system for which null polarization spectra are expected.

For this calculation we simulate $8 \times 10^7$ Monte Carlo packets and compute spectra over 111 logarithmic time steps from 2 to 120 d after explosion. We bin the final spectra of both the DCT and the EBT in logarithmic wavelength bins with $\Delta \lambda / \lambda = 3.912 \times 10^{-3}$. For this test, local thermodynamic equilibrium (LTE) has been assumed for all time steps. The calculation was carried out by mapping the spherically symmetric W7 model on to a $100^3$ Cartesian grid, through which the packets were propagated. The $v$-packet EBT is activated from 10 to 30 d after explosion and only for $r$-packets with emergent $r$-wavelength between 3500 and 10 000 Å. Spectra for the EBT are computed for the viewing angle $\mathbf{u}_{\text{obs}} = (0, 0, 1)$, although we note that (since the model is spherically symmetric) the choice of observer orientation here is arbitrary. Compared to the DCT, the

\footnote{We note that LTE is a crude approximation, especially for epochs after maximum light. We will, however, confine most of our discussion to relatively early epochs when the LTE approximation should be reasonable.}
runtime penalty associated with using the $v$-packet routine in the EBT is found to be less than a factor of 2, with the advantage that the number of packets contributing to the emergent spectrum is a factor of $\sim 115$ higher.

Fig. 8 shows spectra calculated with the EBT at 15, 20 (around $B$-band maximum light) and 25 d after explosion. Angle-resolved (10 angle bins)$^4$ spectra obtained with the DCT are shown for comparison. We note that the calculation of the direct counting flux spectrum is exactly the same as in the previous version of ARTIS (Kromer & Sim 2009), with the exception that electron scattering is now treated fully via the scattering matrix in equation (9) rather than assuming isotropic scattering. The agreement between the two techniques is very good, with the EBT being much less affected by Monte Carlo noise, as expected. To estimate the Monte Carlo noise in the spectra, we use the fact that the calculation has been carried out on multiple cores which provides us with a set of independent estimates for any given observable. In particular, we divide the simulation outputs into eight subsets, each comprising one-eighth of the cores, and calculate an emerging spectrum for each of them. Spectral differences between different subsets are representative of the Monte Carlo noise and estimated by computing residuals from a mean spectrum. The standard deviation of the residual in the $v$-packet spectrum is 13.3 times smaller than that calculated for the angle-resolved direct counting spectrum.

In Section 3.2 we chose a number of 51 viewing-angle bins as a reasonable value to obtain accurate angle-resolved results with relatively low Monte Carlo noise, as expected. To estimate the Monte Carlo noise in the spectra, we use the fact that the calculation has been carried out on multiple cores which provides us with a set of independent estimates for any given observable. In particular, we divide the simulation outputs into eight subsets, each comprising one-eighth of the cores, and calculate an emerging spectrum for each of them. Spectral differences between different subsets are representative of the Monte Carlo noise and estimated by computing residuals from a mean spectrum. The standard deviation of the residual in the $v$-packet spectrum is 13.3 times smaller than that calculated for the angle-resolved direct counting spectrum.

Polarization spectra around maximum light in the $B$ band are reported in Fig. 9. As expected from a one-dimensional model, the average $Q$ and $U$ throughout the whole spectral range are consistent with zero for both techniques, with the signal-to-noise ratio decreasing towards the red because of the lower flux level. The decrease in Monte Carlo noise when comparing the DCT to the EBT is remarkable: the standard deviation in the $Q$ ($U$) spectrum is a factor of 14.1 (13.7) larger in the former compared to the latter, in good agreement with our findings from the flux spectra. This simple comparison clearly shows that the $v$-packet technique is superior for producing accurate polarization levels.

We note that the factor by which the noise improves does depend on the number of angular bins used (the improvement is less dramatic – but still significant – if fewer bins are used). However, our choice of 10 bins is rather conservative (cf. Section 3.2 where it was found that $\sim 51$ bins were required for accurate representation of a simple 2D model).

### 4.3 Ellipsoidal toy model

In this section, we follow previous studies (e.g. Höflich 1991; Kromer & Sim 2009; Dessart & Hillier 2011) and use ellipsoidal models as a starting point to explore aspherical geometries. We use a model equivalent to that of Kromer & Sim (2009), which has a prolate ejecta morphology (PEM) and also consider a similar model with an oblate ejecta morphology (OEM). Specifically, we assume ellipsoidal isodensity surfaces with density profile

$$\rho(\xi) \propto \begin{cases} \exp\left(-\frac{\xi}{\xi_0}\right), & \xi < \xi_{\text{max}}, \\ 0, & \xi > \xi_{\text{max}}. \end{cases} \quad (25)$$
The parameter $\xi$ is defined in terms of the components of velocity in cylindrical polar coordinates $v = (v_r, v_\phi)$ as

$$\xi = \begin{cases} \sqrt{\left(\frac{v_r}{\xi_0}\right)^2 + v_\phi^2}, & \text{OEM}, \\ \sqrt{v_r^2 + \left(\frac{v_\phi}{\xi_0}\right)^2}, & \text{PEM}. \end{cases}$$

(26)

where $\xi_0 = 2750$ km s$^{-1}$ and $h$ is the ratio between the semimajor and semiminor axis. Here we limit our study to an axis ratio $h = 2$ and fix the maximum velocity parameter $\xi_{\text{max}} = 13750$ km s$^{-1}$. We adopt a composition for both models that is roughly appropriate for SNe Ia. Specifically, the total mass and chemical yields of the ejecta are chosen to be the same as for the W7 model and a stratified composition with three ellipsoidal zones ($h = 2$) is assumed. The model is set up by filling the ejecta from the centre to accommodate the W7 yields of the different element groups; the innermost region is filled with ‘iron group’ elements ($21 \leq Z \leq 30$), the middle with intermediate-mass elements ($9 \leq Z \leq 20$) and the outermost with low-mass elements ($Z \leq 8$). The transitions between the different layers are at $\xi \sim 8000$ and $\sim 10500$ km s$^{-1}$. Relative abundances of the elements inside each zone are kept fixed to the W7 values.

LTE radiative transfer calculations have been performed over 111 logarithmic time steps from 2 to 120 d after explosion, $2.4 \times 10^8$ and $4 \times 10^8$ Monte Carlo quanta have been generated for the PEM and OEM, respectively. Since the redder regions of the polarization spectra are typically noisier due to the lower flux (i.e. fewer Monte Carlo quanta per frequency bin; see Fig. 9), an additional simulation has been carried out for the PEM (OEM), with $8 \times 10^7$ ($4 \times 10^7$) Monte Carlo quanta and with the EBT routine called only for $\lambda > 6000$ Å. The two simulations have thus been combined to produce final spectra for the EBT in the whole range between 3500 and 10000 Å. Spectra are computed with the EBT from 10 to 30 d after explosion and for two extreme viewing angles: along the $z$-axis, $n_{\text{obs},1} = (0, 0, 1)$, and along the $x$-axis, $n_{\text{obs},2} = (1, 0, 0)$.

4.3.1 Flux and polarization spectra

In Fig. 10 we compare the $v$-packet total flux spectra at 19 d after explosion for the two ellipsoidal models. We find the same strong viewing-angle dependencies reported by Kromer & Sim (2009). For a given morphology, packets escaping along the major axis see a velocity twice as large compared to the minor axis and the corresponding spectrum is therefore characterized by broader features and stronger line blending; moreover, the spectrum viewed along the major axis is fainter since the projected area along this axis is smaller and the typical opacity is higher. The same geometrical arguments can also be used to compare spectra for the two different geometries: spectra viewed down the minor (major) axis are qualitatively similar, because packets see the same velocity range, but the prolate ellipsoid is fainter than the oblate due to the smaller projected surface.

Polarization spectra for the observer orientation $n_{\text{obs},1}$ are consistent with zero for both models, reflecting the overall spherical symmetry of the projected surface. As shown in Fig. 11, however, observer orientations from which the model has an elliptical projected surface produce a clear polarization signal in $Q$. $U$ remains consistent with zero because the model is axisymmetric, and the calculated $U$ spectrum can be used as a convenient proxy for the Monte Carlo noise in the $Q$ spectrum.

Sign reversals from shorter to longer wavelengths are found in the $Q$ spectrum for both the PEM and the OEM, a behaviour that cannot be explained by the simple picture of an optically thin electron scattering atmosphere illuminated by a point source. In the latter, one would expect the overall polarization to be negative (positive) for the PEM (OEM), with a polarization decreases across the lines because of flux dilution. Instead, as found by previous studies (Desert & Hillier 2011; Patat et al. 2012), the results of full calculations are more complex and sign reversal in polarization spectra can arise. These complexities can be ascribed to variations in thermalization depth with wavelength (see below for explanation) and highlight the need for realistic calculations beyond simple toy atmosphere geometries for the interpretation of data.

Fig. 12 shows the intensity and polarization distributions projected on the velocity plane ($v_r, v_\phi$). The maps have been calculated for the PEM selecting the emergent $v$-packets between 16.5 and 21.5 d after explosion and in the spectral regions 3500–6000 and 6400–7200 Å. In both wavelength intervals, the intensity emission region in projection is less elliptical than the density contour, and this is a stronger effect in the blue. This behaviour can be ascribed to the relative contributions of the line and the electron scattering opacities in different regions of the spectrum (see Fig. 13): the blue region is dominated by line opacities and thus the intensity distribution in projection is more circular than elliptical; in contrast, the red region is free from strong line opacities (see also Pinto & Eastman 2000; Kasen et al. 2004; Patat et al. 2009) and therefore

\footnote{This is because, among all the packets created at a given isodensity surface, those at highest projected velocities (i.e. around the major axis of the ellipsoid) sweep out the largest velocity range on their journey to the observer, and therefore encounter the greatest line opacity.}
Figure 11. Flux and polarization spectra at 19 d after explosion for the PEM (left-hand panels) and the OEM (right-hand panels). The observer is placed at $n_{\text{obs},2}$ (along $x$). Identification between polarization features and lines in the spectrum are shown with blue vertical lines. Scaled projected surfaces are shown in green.

Figure 12. Colour maps of normalized $I$ (left-hand panels), $Q$ (middle panels) and $U$ (right-hand panels) distributions projected on the velocity plane ($v_y, v_z$). The maps are computed for the PEM using the EBT and selecting the emergent $\nu$-packets between 16.5 and 21.5 d after explosion and in the wavelength regions 3500–6000 Å (upper panels) and 6400–7200 Å (lower panels). Solid lines mark the outer boundary of the iron-group-element zone (inner white ellipse) and the maximum velocity parameter $\xi_{\text{max}} = 13750$ km s$^{-1}$ (outer black ellipse). The intensity distribution in the blue is more circular than the projected density contour: $Q$ is dominated by contributions along the minor axis, leading to a positive polarization level. In contrast, the intensity distribution in the red is more similar to the projected density contour: $Q$ is dominated by contributions along the major axis and therefore biased towards a negative value.
Figure 13. Evolution of the relative contribution of different scattering processes to the observed spectrum. Fractions are calculated for the PEM (left-hand panels) and the OEM (right-hand panels) with the DCT by selecting escaping packets based on their last interaction(s) prior to escape. The fraction of packets that underwent a depolarizing interaction process (bound–bound, bound–free or free–free emission) as last interaction is shown in red. The contribution from packets that had a single electron scattering interaction since their last depolarizing interaction is indicated in blue, and packets that suffered multiple electron scattering events prior to escaping are shown in black. Upper panels show contribution in the spectral region 3500–6000 Å, lower panels in the wavelength range between 6400 and 7200 Å.

Figure 14. Flux spectrum (solid black line) and Q polarization spectrum (red line) around the Si ii λ5979 and Si ii λ6355 features for the PEM viewed along the x-axis. Rest wavelengths of the two lines are marked by vertical dashed lines. Inverted P Cygni profiles for the two silicon lines can be identified in the Q spectrum.

As expected, $U^*$ remains consistent with zero at all times. If we consider the entire synthetic spectrum (i.e. $\lambda_1 = 3500$ Å, $\lambda_2 = 10000$ Å), $Q^*$ in the PEM (OEM) evolves from negative to positive (positive to negative) values as we go from early to late epochs (see Fig. 16). This behaviour is due to changes in the relative contributions of the blue and red region as a function of time, and can easily be understood from polarization light curves computed for the spectral intervals between $\lambda_1 = 3500$ Å and $\lambda_2 = 6000$ Å and between $\lambda_1 = 6400$ Å and $\lambda_2 = 7200$ Å (see Fig. 16). As can be anticipated from the polarization spectra, $Q^*$ is positive in the blue and negative in the red for the PEM, whereas the opposite is true in the OEM. The OEM light curve evolves more rapidly than the PEM, having reached peak flux at around 14 d and then starting to decline. This more rapid evolution is also clearly evident in the degree of polarization, which changes much more noticeably over this time period for the OEM. In particular, polarization in the pseudo-continuum region between 6400 and 7200 Å is approximately constant ($Q^*/I^* \sim -1$ per cent) in the PEM, whereas significant evolution is found for the OEM.

4.3.2 Spectral evolution and light curves

The spectra of both ellipsoidal models are shown in Fig. 15 for three epochs (14, 19 and 24 d after explosion). The integrated luminosity in the wavelength range 3500–10 000 Å peaks at ~19 d after explosion in the PEM, whereas the maximum is reached earlier in the OEM (~14 d after explosion, see Fig. 16). To quantify the time evolution of the polarization, we have also computed polarization light curves $Q^*(t)$ and $U^*(t)$ by integrating $Q$ and $U$ values over chosen wavelength regions ($\lambda_1$ to $\lambda_2$):

$$Q^*(t) = \int_{\lambda_1}^{\lambda_2} Q(\lambda, t) \, d\lambda, \quad U^*(t) = \int_{\lambda_1}^{\lambda_2} U(\lambda, t) \, d\lambda.$$

5 CONCLUSIONS

We have described a technique for modelling polarization for multidimensional supernova explosion simulations, and implemented it in the radiative transfer code ARTIS (Kromer & Sim 2009). Our method uses an approach inspired by Lucy (2005), and related to those used by Long & Knigge (2002), Sim et al. (2010) and Kerzendorf & Sim (2014), for extracting observables.
viewing-angle spectra are obtained by summing contributions from \(v\)-packets generated at each \(r\)-packet interaction point and escaping to a chosen observer orientation (EBT). These escaping \(v\)-packets can be used to construct synthetic observables (spectra, light curves, polarization spectra) that have substantially reduced Monte Carlo noise compared to spectra obtained by direct binning of escaping \(r\)-packets. We also investigated a higher order Monte Carlo noise reduction approach, based not on \(r\)-packet interaction sites but on \(r\)-packet trajectory elements (TBT). Initial results, however, suggest that this approach may be less computationally efficient.

We validated our polarization scheme using an idealized test code in a simple configuration, and found good agreement with predictions from Hillier (1994). Applying the same idealized test code to a simple ellipsoidal toy model, we then verified that continuum polarization levels calculated with the EBT agree with values predicted by direct binning of the escaping \(r\)-packets. We implemented the EBT in \textsc{artis} and tested it for a model with a realistic SN Ia composition and opacity (the spherically symmetric W7 model): as expected, the \(v\)-packet method could accurately reproduce the synthetic spectrum obtained by direct binning of emergent Monte Carlo quanta and also predict polarization consistent with zero. However, the EBT is much less affected by Monte Carlo noise (with typical signal-to-noise ratios a factor of \(\sim 13\) higher than those obtained with the direct binning approach) and thus more suitable to reproduce very weak signals (e.g. polarization levels observed in SNe Ia).

Finally, we synthesized flux and polarization optical spectra with the EBT for prolate and oblate ellipsoids with axis ratio of two, using typical SNe Ia velocities and compositions (including composition layering). As expected, we obtained null polarization spectra when the projected surface on the plane of the sky is circular. In contrast, aspherical projected areas yield a polarization signal (typically \(\sim 1\) per cent) in both morphologies. The polarization is characterized by sign reversals across the spectrum and peaks associated with troughs of strong optical features. This behaviour is consistent with results of previous studies using similar ejecta morphologies (Höflich 1991; Dessart & Hillier 2011; Patat et al. 2012) and is ascribed to variations in thermalization depth with wavelength. At the epochs we studied (14–24 d post-explosion), the evolution of polarization spectra is more dramatic for the oblate than the prolate morphology, both in the continuum and in the line polarization levels.

In this paper we have focused on developing our technique and testing its accuracy in calculating intensity and polarization spectra for one- and two-dimensional models. This study has laid the groundwork for future calculations in which we will exploit the multidimensional capability of \textsc{artis} and calculate polarization spectra for a set of contemporary SN Ia explosion models. Such calculations will help to identify geometric discriminants between models and to make comparisons between their predictions and spectropolarimetric data more reliable.

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Polarization light curves between 10 and 30 d after explosion is reported here since is consistent with zero. v x

Figure 16. Polarization light curves between 10 and 30 d after explosion for the PEM (left-hand panels) and the OEM (right-hand panels) viewed along the x-axis. Only Q* is reported here since Q* is consistent with zero. v x

Figure 17. Q polarization spectrum for the PEM (in red) calculated ~d after B-band maximum light. For comparison the black line shows the polarization spectrum of SN 2004dt (Leonard et al. 2005) at the same epoch. Given that the PEM is axisymmetric, the polarization spectrum of SN 2004dt calculated along the dominant axes, p1, is shown here. Polarization levels predicted by the PEM across the lines are comparable (within a factor of 2) to those observed, while the polarization in the continuum (grey shaded area) is too high.

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