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DUST-AFFECTED MODELS OF
CHARACTERISTIC LINE EMISSION
IN SUPERNOVAE

Thesis submitted for the Degree of Doctor of
Philosophy of the University of London

by

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I, Antonia Bevan, confirm that the work presented in this thesis is my own. Where information has been derived from other sources, I confirm that this has been indicated in the thesis.

Abstract

FIRST PARAGRAPH MUST BE IN HERE.

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Acknowledgements

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*Gather out of star-dust
Earth-dust,
Cloud-dust,
Storm-dust,
And splinters of hail,
One handful of dream-dust
Not for sale.*

- Langston Hughes

Dream Dust

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List of Acronyms

- AAT - Anglo-Australian Telescope
- API - Application Program Interface
- CTIO - Cerro Tololo Inter-American Observatory
- HST - Hubble Space Telescope
- (MC)MC - (Markov Chain) Monte Carlo
- MSE - Mean Squared Error
- (M/N)IR - (Mid/Near) Infra-Red
- ISM - Inter-Stellar Medium
- SED - Spectral Energy Distribution
- (CC)SN(e) - (Core-Collapse) Supernova(e)
- UV - Ultra-Violet
- VLT - Very Large Telescope

Chapter 1

Introduction

Should you choose to seek out one of my friends and ask them about my whereabouts in recent months, they would likely stare at you blankly. Upon further interrogation they would probably yield the information that I was preoccupied writing about dust, a fact which I imagine they find bemusing and possibly somewhat concerning. Blissful as they are in their ignorance of dust (astronomers find no such peace), they do not know the importance of this all-pervading substance.

The universe is an extremely dusty place. The ubiquity of dust throughout almost all epochs and environments demands a comprehensive understanding of its formation and evolution, properties and effects. It plays numerous roles in a variety of scenes; it is a building block of all solid bodies, a birthing place for molecules, a crucial ingredient in star formation and an extreme annoyance for cosmologists. It is both a product of physical processes and an agent of chemical ones.

It is perhaps confusing therefore that there is comparatively little consensus regarding the formation processes and natal environments that result in the evolution of certain atoms and molecules into the grains we call dust. Over the years since the first discovery of dust in the very early universe, a growing population of astronomers and astrophysicists have turned their attention to the study of dust formation in core-collapse supernovae (CCSNe), in the hope that these objects might prove to be the missing piece of the puzzle. Recent observations of a number of core-collapse supernovae and remnants have lent weight to this theory, with models and analyses of spectral energy distributions (SEDs) suggesting

the presence of large reservoirs of cool, ejecta-condensed dust.

I have sought to make my own small contribution to this field by exploiting a different observational signature, that of blue-shifted line profile asymmetries observed in the spectra of many CCSNe and attributed to the formation of dust in the ejecta. By quantitatively modelling characteristically asymmetric spectral line profiles using a novel code, DAMOCLES, I have attempted to determine the rate of dust formation in CCSNe and the expected order of magnitude of the eventual dust masses produced.

Throughout the remainder of this chapter I will attempt to elucidate the above synopsis in more detail. A brief discussion of the roles that dust plays in the universe will be followed by a summary of our current understanding of dust formation in CCSNe. I will conclude this chapter with a short justification of the approach that I have adopted for this work and an outline of the structure of this thesis.

1.1 A Handful of Dust

1.1.1 A Brief History

The presence of dust in the universe was first theorised when astronomers observed dark patches of sky in the Milky Way where all of the stars had been “erased” (see Figure 1.1). Whilst some claimed that these black regions were in fact a true absence of stars resulting from some anomaly in the stellar distribution, others felt that it was more likely that an obscuring cloud of material was blocking the light from the stars behind. In 1930, Donald Trumpler confirmed this latter theory by considering the apparent magnitudes and colours of stars located at different angles to the galactic plane, discovering that those closer to the plane appeared redder than their more distant counterparts. This was the first evidence of interstellar reddening and the beginnings of our understanding of dust as a scatterer, absorber and emitter of radiation.

For the next few decades, dust was thought to be largely an irritating obstacle to observing and comprehending more interesting facets of the universe. We now have a much fuller understanding of the variety and importance of the roles that dust plays throughout astrophysics.

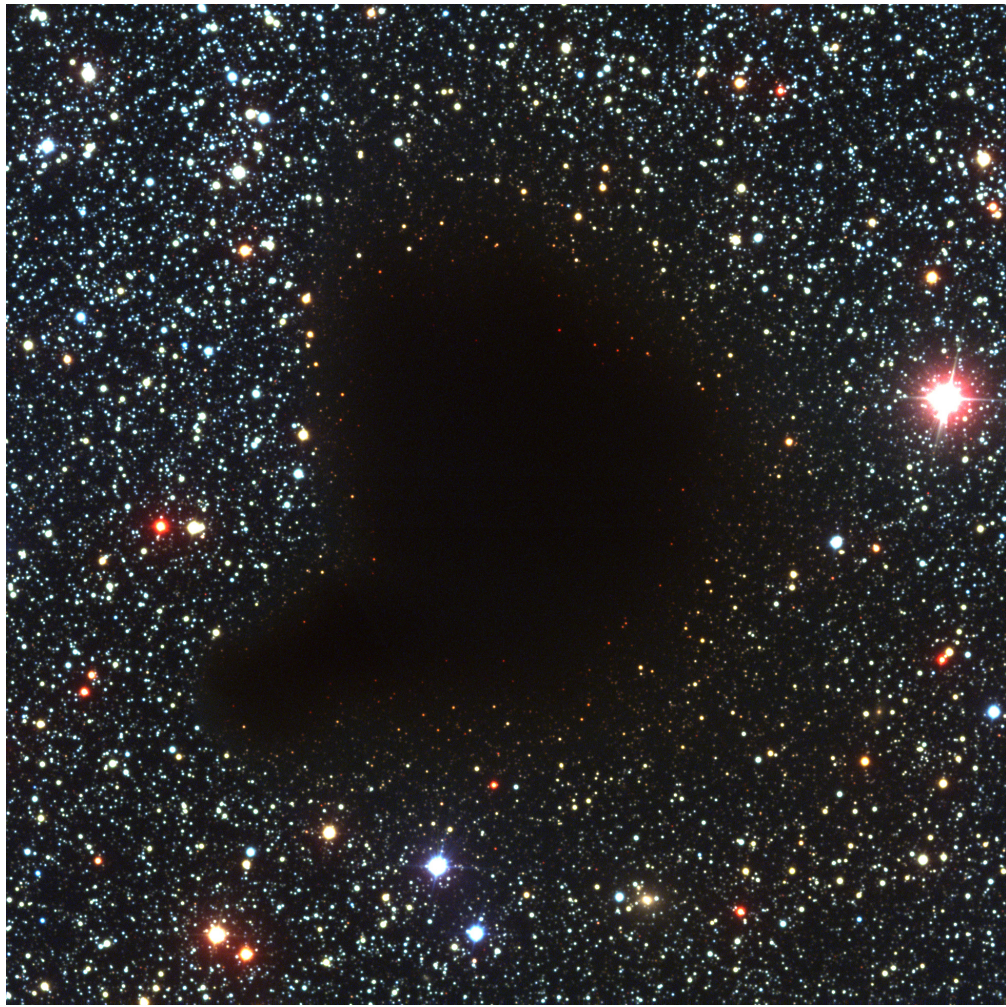


Figure 1.1. The dark globule Barnard 68, LDN 57. ESO press release 30 April 1999.

1.1.2 The Roles of Dust in the Universe

Despite comprising only $\sim 1\%$ of the mass of the interstellar medium (ISM), dust grains account for as much as 30% of the total galactic luminosity via their emission in the infrared (IR) (Li & Greenberg 2003). In the cycle of matter from the ISM to condensing clouds to stars and back again, dust is far more than a passive passenger along for the ride. Whilst residing in the ISM, dust is important in determining its thermodynamics. It acts both as a heating agent via the emission of photoelectrons in regions of strong ultra-violet (UV) radiation and a coolant in dense regions via the emission of IR radiation. In this role as a coolant, dust is also crucial to the process of star-formation, helping to remove gravitational energy and allowing the natal cloud to collapse. Dust also contributes to the star formation process by shielding the gas from ionising radiation, helping to speed up

the construction of the protostellar core.

In addition to the above physical functions, dust plays an essential part in chemical processes. Heavy elements in the local medium are depleted through their inclusion in dust grains. These grains attract gaseous atoms to their surfaces and catalyse the formation of molecules which are then released back into the surrounding medium.

Dust does not reside solely in the ISM however. It is present in large quantities in the circumnuclear tori found around active galactic nuclei. Dust is also found between planets, around stars and in protoplanetary discs, where dust grains are often the smallest unit of the building blocks that will go on to form planetesimals and planets. These grains may even be responsible for the origins of life.

The more detailed our understanding of dust as an astrophysical community, the more accurate we can make our inferences across an entire range of fields. There is arguably no other topic in astronomy that has such wide-ranging effects.

1.1.3 The Medium of Dust

An increasingly detailed knowledge of the nature and properties of dust has developed over the last few decades. Dust grains have their terrestrial analogue in soot or very fine sand rather than in the dust bunnies that one may find behind the sofa. When found in the ISM they are generally small, between $0.05\mu\text{m}$ and $0.25\mu\text{m}$ in radius, and are normally predominantly composed of carbon or silicates. Carbonaceous grains may take many forms ranging from structured solids such as diamond and graphite to amorphous molecules and aromatics. They are generally found to be strongly attenuating. Silicates tend to be more glassy and contain silicon and oxygen potentially with the dirtying addition of magnesium, iron or other heavier elements. Condensates of more complex molecules such as olivine (MgFeSiO_4) and pyroxene (MgSiO_3) make up these grains.

Whilst an increasingly strong picture of the composition and properties of dust is becoming apparent, there are still a number of largely unresolved issues regarding the makeup of a dusty medium. Different species and composites thereof have different optical properties. In order to model the absorption and scattering of radiation off dust grains it is necessary to first know the complex refractive indices of a given species over the relevant wavelength range. Laboratory measurements have produced a number of different sets of optical constants for a variety of carbonaceous and silicate species and these are well-utilised throughout the field. It is noted at this early juncture however that in many cases

there are numerous, somewhat contradictory, sets of optical constants for a given species and that these variations can potentially cause a degree of confusion regarding the results of models that utilise them. This topic will be discussed in detail later in this thesis (SECTION).

Dust grains are generally assumed to be spherical in order to make their simulative treatment more straightforward but in reality dust shapes are actually much more complex. Sophisticated models of dust grains sometimes adopt a continuous distribution of ellipsoids to represent dust grain shape. This allows grains to take any ellipsoidal form ranging from flat discs to needles to perfect spheres. However, even this more detailed consideration omits structures that are akin to long strings or to fluffy particles (not dissimilar to a tumble dryer ball in shape). Heretofore, the vast majority of models, including DAMOCLES, have only considered spherical grains and this wide variety of shapes therefore represents a significant modelling challenge to be addressed in the future.

Dust in the universe follows a cycle. From its stellar birthplace, it is ejected and slowly integrates itself with the ISM before condensing into molecular clouds and ultimately once again returning to stars. Most of the knowledge of the properties of dust applies only to grains in the ISM, which are found to follow a grain radius distribution $n(a) \propto a^{-3.5}$ as described by Mathis et al. in 1977. This distribution does not necessarily apply immediately after their formation, however, as grains are subject to numerous forces that can result in their destruction, sputtering or evaporation. The grain size distribution and relative abundances of species of newly-formed grains are still topics in dispute and are issues that I attempt to address in my models. The issue of dust grain shape will hopefully be addressed in future versions of DAMOCLES.

1.2 Core-Collapse Supernovae as Dust Factories

In an effort to explicate the motivations behind studying dust, I have so far mostly limited my discussion to the evolution and properties of dust after the initial stages of its formation once it has entered the ISM. The most current and contentious debate, however, is over the natal environment of dust grains.

Supernovae are the violent explosions that are the death of stars. They evolve very quickly and create extreme conditions. Focus on supernovae as a possible source of dust in the universe has been motivated by the physical conditions that they produce shortly after

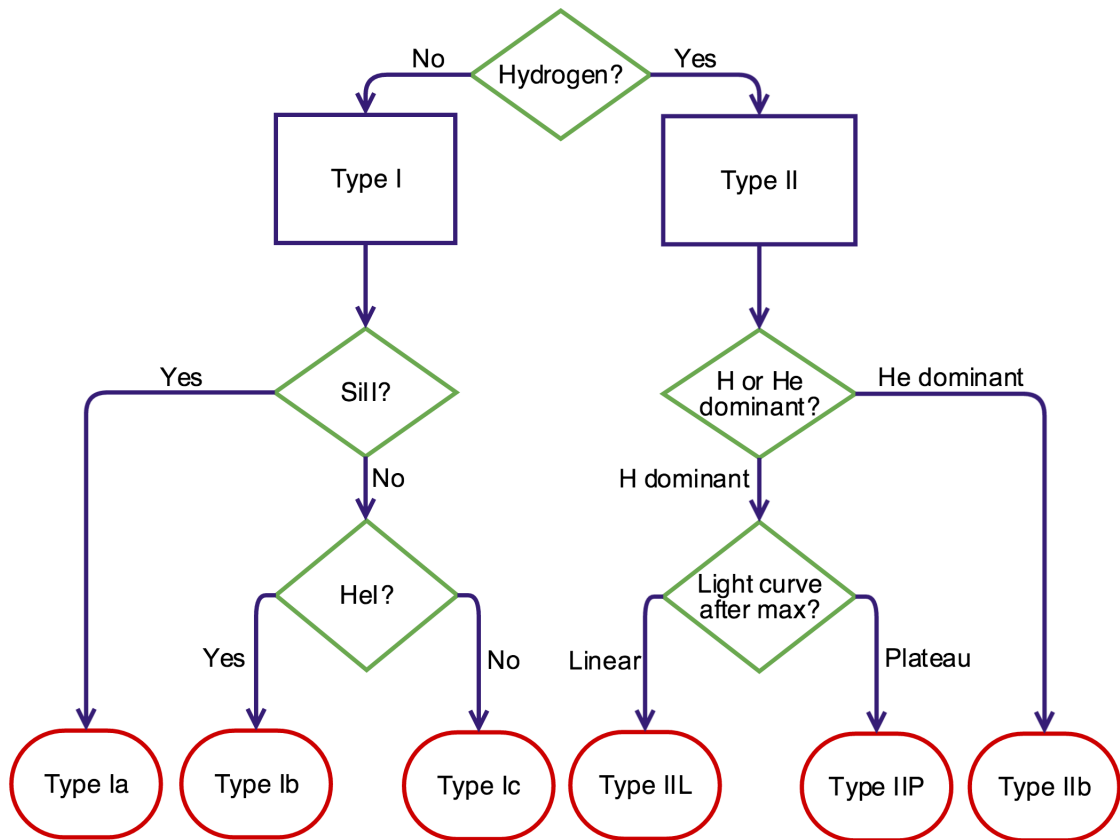


Figure 1.2. A flowchart summarising the supernova classification scheme

their outbreak and by the presence of large quantities of heavy elements that constitute the integrant ingredients of dust grains.

1.2.1 Types of Supernovae

Supernovae may be classified into a number of different types. They are bisected initially into Types I and II according respectively to the absence or presence of hydrogen in their early spectra. Further sub-classifications depend on other features in the early spectra, properties of later spectra and the evolution of the light curve after maximum light. A summary of the supernova classification scheme is presented in Figure 1.2.

If the initial classification is Type I then all further sub-classifications depend solely on the properties of the early spectra (a few days after explosion) as detailed in Figure 1.2. Type II supernovae are somewhat more complex in their categorisation. After classification as a Type II, further subdivisions depend on the dominance of hydrogen or helium in *later* spectra. Helium dominant supernovae are classified as Type IIb and hydrogen dominant

supernovae are classified as either Type IIL (those which have a linearly decaying light curve after maximum light) or Type IIP (those that exhibit a plateauing light curve after maximum light). Type IIn supernovae are omitted from the summary presented in Figure 1.2 as they cannot be classified straightforwardly via a bifurcating process. Type IIn supernovae will generally have strong emission lines, particularly hydrogen lines, often with complex profiles. Crucially, the spectra of Type IIn supernovae do not exhibit the broad absorption features frequently seen in other types and instead contain narrow lines (hence Type IIn).

1.2.2 From Massive Stars to Remnants

It is generally accepted that the progenitors of Type Ia supernovae are white dwarfs that exist in a binary system with another star. The accretion of material from one star to another results in a thermonuclear explosion, a mechanism that is unique to Type Ia supernovae. There have not been any observations suggestive of dust forming in the aftermath of a Type Ia supernova and I therefore do not consider these objects any further, focusing my attention solely on supernovae that explode via the core-collapse mechanism.

Broadly, this process is initiated when a massive star ($\geq 8M_{\odot}$) starts to fuse heavier elements. The fusion of ever heavier elements generates increasingly less energy whilst also causing the mass of the core to increase. Eventually, radiation pressure drops sufficiently that the core can no longer support itself against its own self-gravity and begins to collapse rapidly. Within milliseconds, the core reaches extremely high densities and, when it can no longer condense further, “bounces” off itself causing an immense shockwave to propagate outwards and a vast quantity of energy to be released via the expulsion of neutrinos. Much of this complex process is still poorly understood and interesting models are currently being produced recreating these very early stages using a numerical approach. Though the explosion mechanisms of CCSNe are largely beyond the scope of my work, some attention will be paid to these models later in this thesis since instabilities that arise in these early stages can influence the structure of the ejecta at later stages of its evolution.

For several hundred years after the explosion, the supernova (now a remnant) is in the free-expansion phase. During this phase, the mass and velocity of the expanding supernova massively exceed those of the surrounding medium, fortuitously allowing the behaviour of the remnant to be analysed as if it were expanding into a vacuum. The shock radius during this phase may therefore be calculated simply as $R_s = v_s t$. As the

shockwave propagates through the ISM, interstellar material that has been compressed by the forward shock begins to accumulate. At the same time a reverse shock wave begins to propagate back through the ejecta. It is during this phase, which arises very soon after the initial explosion and lasts for a few hundred years, that the physical conditions in the ejecta are thought to be optimal for dust formation. The phase ends when the mass of material ahead of the forward shock is of a similar magnitude to that behind and the mathematical treatment of its behaviour must be altered.

1.2.3 Dust Formation in CCSNe

Superficially, the formation of dust grains requires densities high enough for interaction between particles to take place, but temperatures that are cool enough to allow the grains to survive. The theory that the ejecta of a CCSN in its free-expansion phase could provide these conditions was first hypothesised by Cernuschi et al. in 1967 and they have now long been thought to be potential dust factories (Hoyle & Wickramasinghe 1970; Kozasa et al. 1991; Todini & Ferrara 2001).

The formation of dust was originally thought to result from the stochastic process of classical nucleation whereby particles coalesce to form the seeds of dust grains. These seeds become the nucleation sites from which grains are ultimately born through the aggregation of further particles. Various models of dust formation in the ejecta of CCSNe have used this approach (Kozasa et al. 1989; Todini & Ferrara 2001; Nozawa et al. 2003; Schneider et al. 2004).

More recently, several models of dust formation in CCSNe that consider the effects of chemistry on the growth of dust grains have been published. These models consider the chemical composition of the gas and include chemical reaction rates thereby considering the manner in which molecular evolution influences dust grain formation and growth rates (Cherchneff & Dwek 2009, 2010; Sarangi & Cherchneff 2013; Sarangi & Cherchneff 2015).

Models using both methods have predicted dust masses of the order of $0.1\text{-}1 M_{\odot}$ of dust forming within the ejecta of CCSNe of progenitor masses between $12\text{-}40 M_{\odot}$ within the first few years after the initial explosion.

1.2.4 The Three Signatures of Dust

The presence of dust in the ejecta of CCSNe can be indicated by three main signatures:

A decrease in the light curve

As the dust begins to form in the ejecta, UV and optical light is absorbed by the dust causing a decrease in the light curve at these wavelengths.

Excess IR emission

An increase in emission in the IR occurs contemporaneously with the decrease in the UV-optical light curve. A thermal MIR excess is caused by warm dust and an excess in the far-IR and sub-mm is the result of cold dust. The increase in emission in these wavelength can be caused by newly-formed dust condensing in the ejecta but can also be a result of the illumination of pre-existing dust

Blue-shifted line profiles

Finally, the onset of the formation of dust can cause an asymmetry in line profiles in the optical and IR. The absorption and scattering of optical or near-IR radiation by newly-formed dust within the ejecta can result in an asymmetry between the red and blue shifted components, with redwards emission from the far side of the ejecta undergoing greater absorption and resulting an overall shift of the profile to the blue.

All three of these signatures have been discussed in detail over the timeline of this subject but the focus has been on using the excess IR emission seen in the SED of CCSNe to determine quantitatively dust masses in these objects. This approach has resulted in a lively debate regarding the quantities of dust that CCSNe are capable of producing.

1.2.5 The Dust Mass Debate

Over the past two decades, several high redshift galaxies have been found to contain significant masses of dust (Omont et al. 2001; Bertoldi et al. 2003; Watson et al. 2015). CCSNe are one of the few potential sources that could contribute large quantities of dust at early epochs. However, observations over the last decade at mid-infrared (MIR) wavelengths of warm dust emission from CCSNe has suggested that the quantities of ejecta-condensed dust produced during the first 1000 days were typically $\leq 10^{-3} M_{\odot}$ (Sugerman et al. 2006; Meikle et al. 2007; Kotak et al. 2009; Andrews et al. 2010; Fabbri et al. 2011). This is much less than the 0.1-1.0 M_{\odot} of dust per CCSN estimated to be needed in order

to account for the masses observed (Morgan & Edmunds 2003; Dwek et al. 2007). These observations would indicate that other early-time sources of dust must be found.

However, recent *Herschel* far-IR and sub-mm observations of several young supernova remnants have revealed cold dust masses as high as $0.2\text{--}0.8 M_{\odot}$, resulting in a re-evaluation of the rate of dust production by CCSNe and a renewed focus on these objects as sources of dust (Barlow et al. 2010; Matsuura et al. 2011; Gomez et al. 2012).

SN 1987A

Critical to this field has been the study of SN 1987A. This supernova is one of the most studied objects in the history of astronomy and has been observed almost continuously since it exploded 28 years ago. The reason for this is its proximity. At only 50pc away, it has allowed for some exceptionally well-resolved spectral and photometric observations. Neutrinos from its explosion were even detected at three different observatories . As such, it has yielded results and insight that would not have been possible otherwise and is a key to understanding the formation and evolution of dust in CCSNe.

It is of no surprise therefore that this object has been central to the debate. Lucy et al. (1989) was the first to suggest the presence of dust in SN 1987A *Herschel* observations of SN 1987A were the first to suggest the presence of cold dust in any supernova.

1.2.6 Motivation

It is seems increasingly likely that CCSNe do indeed produce significant quantities of dust. However, there remain a large number of outstanding challenges to consider. Firstly, there are still only a very small number of supernovae that have been observed to have sizeable masses of dust present in their ejecta. If further CCSNe were also shown to have formed large quantities of dust then the already shifting opinion might start to become consensus. Other points to consider regarding dust formation and evolution in CCSNe include the nature of the dust (composition, grain size, grain shape etc.) which is still largely unclear, as is the extent to which it is destroyed or sputtered after its initial formation. Related to this is the uncertainty of the dust formation rate in the ejecta and the issue of where this formation takes place. These are all interesting questions that call out for answers.

The *Hershel* dust mass estimates were based on fitting dust SEDs that peaked at far-IR wavelengths. Unfortunately, following the end of the *Herschel* mission in 2013, there is likely to be a long wait for far-IR facilities with comparable or better sensitivities than

Herschel to become available. Without data, this methodology is temporarily ineffectual. This provided an incentive to make use of alternative methods to estimate the dust masses that form in supernova ejecta.

1.3 Dust-Affected Line Profiles

Lucy et al. (1989) identified a progressive blue-shifting of the [O I] $\lambda\lambda 6300, 6363$ Å doublet from SN 1987A between days 529 and 739 after outburst, with the doublet in the later spectrum being blue-shifted by ~ 600 km s $^{-1}$. Since then, such red-blue asymmetries have been frequently observed in the late-time (> 400 days) spectra of supernova ejecta and there is now a growing database of such observations (e.g. Lucy et al. (1989); Fabbri et al. (2011); Mauerhan & Smith (2012); Milisavljevic et al. (2012)).

The purpose of my work has been to develop a new approach to determining dust masses in supernovae, with the aim of providing an alternative to SED fitting for the future and of providing corroborating or contradicting evidence of past results. I looked to exploit the third signature of dust formation in supernovae, namely the red-clue line asymmetry observed in optical and IR line profiles. Though this feature has been discussed at length by numerous authors (REFERENCES) it has very rarely been quantitatively measured or modelled.

I have sought to construct a Monte Carlo based code that numerically models this feature in the spectra of SNe in order to quantitatively determine dust masses formed at a variety of epochs post-explosion, additionally seeking to place constraints of the composition and grain size distributions of the newly-formed dust.

1.3.1 Observing

Numerous telescopes have recorded spectra of CCSNe in the optical and IR, some with extremely high resolution. The Anglo-Australian Telescope (AAT), the Cerro Tololo Inter-American Observatory (CTIO), the Hubble Space Telescope (HST) and the Very Large Telescope (VLT) have all observed several supernovae in the optical including SN 1987A. Other telescopes such as the two Gemini Multi-Object Spectrographs (GMOS) have also taken spectra of numerous CCSNe.

Advances in digital storage have allowed for spectral and photometric observations to be made easily available online. Many observatories now publish their recent observations

online in archives and are working to upload observations that pre-date file sharing services. Much of the data used in this thesis was obtained from these archives.

1.3.2 Modelling

Need to think about what i actually need to consider in this section.

- dust absorption, scattering and reemission in the IR - radiative transfer and independence of optical properties and temp meaning do not need to fully solve rad tran problem
- mie theory deets? how is it solved and what is the theory? - assumptions of isotropy? scattering matrices?

1.3.3 Mie Theory

In order to truly understand observations that are a product of dust, we must first understand those facets of a dusty medium that determine its interaction with incident radiation. Dust grain radii are often of a similar order of magnitude to the wavelength of that incident radiation and so may be analysed from an optical perspective using Mie Theory. Mie Theory is a mathematical solution to Maxwell's equation that describe how light is scattered off a small particle. In combination with the optical properties of the medium, this allows for a precise determination of the extinction and scattering efficiencies of a given environment. Conveniently, this allows for the straightforward modelling of a dusty medium and it is this calculation that I exploit in my models. Mie Theory assumes particles to be spherical, an assumption which is a potential issue since grains may be crystalline, fluffy or extremely amorphous. This issue is addressed later in this thesis.

1.4 Content Of This Thesis

And relate to your current work - give an overview of the work and what you did as well as the structure of the thesis - see Jo's for example.

Chapter 2

A Description of DAMOCLES

2.1 Monte Carlo Methods

The name “Monte Carlo” describes a class of modelling techniques that employ a stochastic approach to simulating mathematical and physical situations that are otherwise difficult or impossible to solve. By repeatedly sampling random numbers from a probability distribution, numerical results to non-analytic problems may be obtained. The approach was first used by researchers at Los Alamos in the late 1940s who adopted the method to model the transport of neutrons (Metropolis & Ulam 1949). It is from the code name for this project, “Monte Carlo”, that the methods derive their name.

As the available computing power increased over the following decades, Monte Carlo methods became more and more useful as a means of solving complex problems and are now used widely across numerous fields including mathematics, statistics, engineering, finance, the physical sciences and many others. The nature of the approach means that they are particularly well-suited to problems with multiple degrees of freedom, and especially when any of these degrees are coupled. By using random numbers to represent quantities that parametrise a physical problem, a solution to the problem may be sought using a pseudo-random number generator. It must be the case that the quantities that characterise the problem may be represented by a continuous distribution in the range [0,1] in order that the randomly generated numbers may be translated into physical properties (Buslenko et al. 1966).

Having thus obtained a random set of physical parameters, a model is constructed and an output - a “possible outcome” - is obtained. By repeatedly iterating this process with new randomly-generated inputs each time, many possible outcomes are produced and a probability distribution is built up. The interpretation of the outputted probability distribution is dependent on the manner of utilisation of the Monte Carlo method. For example, the procedure may be used to find the mean-free paths of millions of energy packets where the resulting probability distribution of the final frequencies of the packets is equivalent to an energy distribution. This is the process that I make use of and I will discuss it in more detail throughout this chapter.

More recently Monte Carlo methods have been applied to Bayesian statistical analyses that seek to uncover a complete multidimensional probability distribution describing the parameter space of a particular model. The intention is to derive not just a well-fitting model but to understand how variations in a given parameter affect the likelihood that the model is representative of the data. These investigations of parameter space generally adopt a Markov Chain Monte Carlo (MCMC) approach. Where a Monte Carlo method generates a sample from a required distribution, a MCMC technique draws samples according to a predefined set of rules that result in a sequence of samples called a Markov Chain. These methods allow for a more intelligent and efficient sampling of parameter space (Metropolis et al. 1953; Hastings 1970; Gilks et al. 1996).

Clearly, Monte Carlo simulations are limited by their finite nature and will never produce a perfect solution. However, this does not mean that Monte Carlo simulations are lacking in rigour. It may be shown that the error in a Monte Carlo model is approximately $\sim \frac{1}{\sqrt{n}}$ for large n , where n is the number of quanta used in the simulation (Press et al. 2007). The error may therefore be made as small as required by increasing the number of quanta used in the simulation subject to the restrictions of computing time and expense.

In the next section, I discuss the use of Monte Carlo methods as applied to radiative transfer problems and specifically to DAMOCLES. I discuss the computational aspects of my work and the architecture of the code in section 2.3 before finally discussing the limitations of the code and its potential for future developments in section 2.4.

2.2 Radiative Transfer and the Monte Carlo Method

The application of Monte Carlo codes to radiative transfer problems in astrophysics has a strong history. Numerous codes that utilise this stochastic methodology have been written in the past few decades in order to model the transport of energy packets through various media, for example Mocassin, TORUS, Hyperion, SKIRT, LIME, RATRAN, Cloudy and many others (Harries 2000; Hogerheijde & van der Tak 2000; Baes et al. 2003; Ercolano et al. 2003, 2005; Brinch & Hogerheijde 2010; Robitaille 2011; Ferland et al. 2013). The energy to be transported throughout the region of interest is discretised into packets and the path of each packet is calculated according to the properties of the environments that it passes through during its lifetime. Collating the escaped packets at the end of the simulation produces an energy distribution that may be compared to observed photometric or spectral data.

In addition to numerous codes that treat the continuous emission and absorption of energy in dusty environments in order to produce and fit spectral energy distributions (SEDs), there also exist several Monte Carlo radiative transfer codes that model the transfer and interaction of line emission through a 3D nebula in order to produce a synthetic spectrum (e.g. Brute (Thomas et al. 2003) and ARTIS (Kromer & Sim 2009)). These latter models frequently employ an approximation known as the Sobolev approximation (Sobolev 1957). This method allows spectral lines in media moving with high velocity gradients to be treated more simply by solving the radiative transfer equation locally under the assumption that the macroscopic velocity gradient is more important than the thermal line width. Models of supernovae have been produced using both approaches and well-fitting spectra and SEDs have been generated but never, according to the best of my knowledge, has the Monte Carlo methodology been employed to produce sophisticated models of individual line profiles in expanding dusty regions. In this new code, DAMOCLES, we seek to apply the technique to an expanding dusty medium in order to consider the effects of dust on a single emitted line profile.

Previous work by Leon Lucy has considered the problem of computing the spectra of supernovae using Monte Carlo techniques (Lucy 1987, 1999, 2002, 2003; Lucy 2005a; Lucy 2005b). In particular Lucy and colleagues consider dust-induced asymmetric line profiles in the ejecta of CCSNe and they have published results derived both analytically and using simple Monte Carlo simulations (Lucy et al. 1989, 1991). These simulations appear to be

the only published instances of a numerical approach to studying this spectral feature. The DAMOCLES code adopts the same technique as the original modelling by Leon Lucy but allows for a considerably more complex treatment of the composition, geometry and motion of the dusty medium.

Radiative transfer methods as applied to supernovae generally treat a wide wavelength range and seek to conserve the total energy. In the case of SED modelling, this is often achieved by dividing the total energy into packets of equal weight and energy, and iteratively determining the temperature and ionisation structure. In this work, the approach we adopt is somewhat simpler as only a very narrow wavelength range need be considered. Rather than seeking to conserve the total energy, we assume that any packet absorbed by dust would be re-emitted outside the wavelength range of interest and thus no longer contributes to the resulting line profile. Any absorbed packet is therefore removed from circulation. In addition to this, the absorption and scattering of radiation by dust is independent of temperature and there is therefore no need to calculate temperatures throughout the nebula. Similarly, the use of the Sobolev approximation (described above) is unnecessary here as only a single line or doublet is ever treated and a comparatively narrow wavelength range considered.

The subtleties of the problem we consider here lie in the treatment of an atmosphere expanding as fast as 10% of the speed of light, and in the complexities of the dusty medium itself. Lorentz transforms must be carefully applied in order that packets experience the appropriate degree of frequency shifting at emission and at each subsequent scattering event. In this respect, the code is analogous to Monte Carlo radiative transfer models of electron scattering published by (Auer & van Blerkom 1972; Hillier 1991). Indeed, similar features are observed in the outputs of both.

Throughout this section, I will describe the principles, assumptions and techniques adopted in the production of DAMOCLES (see Figure 2.1) before I move on to address the mechanics and architecture of the code itself. DAMOCLES stands for **D**ust-**A**ffected **M**odels **O**f **C**haracteristic **L**ine **E**mission in **S**upernovae.

2.2.1 Energy Packets

The fundamental principle underlying the Monte Carlo approach to the transport of radiation throughout a dusty nebula is that the radiation is discretised into packets. Each of these packets is then propagated throughout the nebula and ultimately contributes a

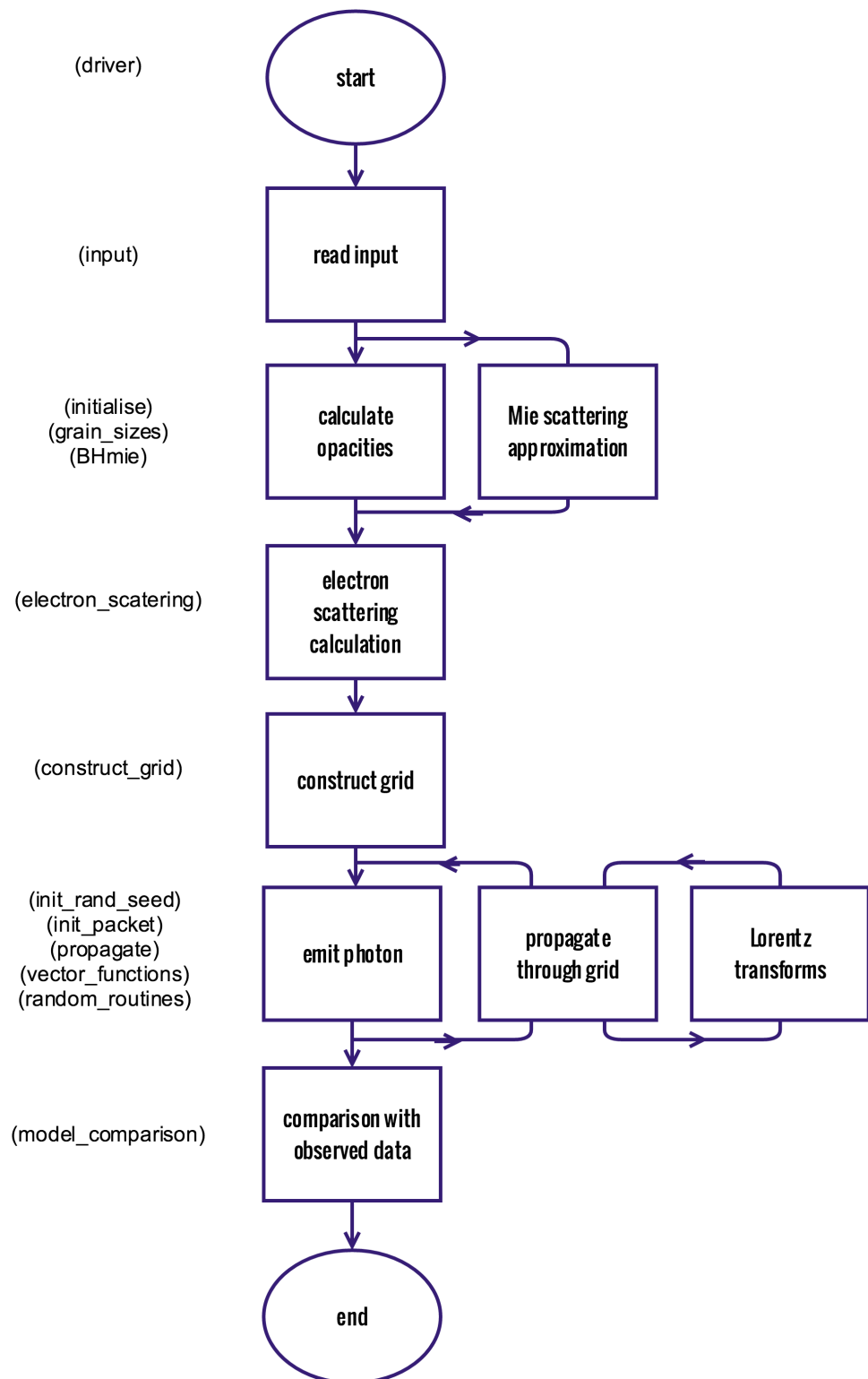


Figure 2.1. A flowchart representing the sequence of processes that take place in the DAMOCLES code. The modules involved at each stage are given in parentheses.

fraction of the final energy distribution. At the start of the simulation, each packet is assumed to consist of n photons of frequency ν_0 , the rest frequency of the monochromatic emission line to be modelled. All packets therefore begin life with initial energy

$$E_0 = nh\nu_0 \quad (2.1)$$

where h is Planck's constant. As the packets move through the ejecta, they are scattered off high-velocity dust grains and after each scattering event, the frequency of the packet is altered. In Monte Carlo simulations that model non-moving atmospheres, packets are usually taken to be of constant energy. When the frequency of a packet is altered after an event, the energy of that packet is kept constant and the number of real photons contained within it assumed to change. However, in the case of dust scattering, the number of real photons is conserved and thus the energy of the packet is altered. This is most easily achieved by weighting each packet over all scattering events as

$$w_p = \prod_{scat} \frac{\nu'}{\nu} \quad (2.2)$$

where w_p is the weight of the packet. The final energy of each packet is then $E = wE_0$, where E_0 is the initial energy of the packet. The final dust-affected line profile is compiled by adding the total energy of all packets in a specific frequency bin in order to produce a histogram.

In these models, unlike fully self-consistent SED radiative transfer models, there is no requirement that the total energy be conserved. We drop this traditional requirement since radiation that is absorbed by dust is re-emitted outside of the wavelength range of interest and thus no longer contributes any flux to the resulting line profile. Packets that are absorbed may be safely removed from the simulation.

2.2.2 Initialisation and the Grid

The supernova ejecta is approximated by a three-dimensional cartesian grid, each cell of which is assumed to have uniform density and composition. By default, the ejecta occupies a shell between inner radius R_{in} and outer radius R_{out} . The grid extends from $-R_{out}$ to $+R_{out}$ in each of the three axes. Each side is split into the same number of divisions and thus each cell is a cube of volume R_{out}^3/n_{div}^3 where n_{div} is the number of divisions

along each axis and is specified by the user. For the remainder of this thesis, a spherically symmetric situation is assumed and in all modelling and testing the grid is constructed in this manner. However, there are no assumptions of symmetry in the code and a cartesian grid was adopted in order to allow for arbitrary geometries to be modelled in the future e.g. ellipsoidal or toroidal ejecta distributions.

Smooth power-law density distributions

Both gas and dust are by default assumed to have a power-law distribution declared as $\rho(r) \propto r^{-\beta}$ between R_{in} and R_{out} . The distribution of gas determines the emissivity distribution and thus the starting positions of the packets in the simulation (see section 2.2.4). However, after the initial emission of energy packets, the gas plays no further role in the simulation as only interactions with dust grains are of interest here. By default, the dust is coupled to the gas and thus follows the same smooth power-law distribution previously described with exponent $-\beta$. The dust density in each cell is therefore calculated as

$$\rho(r) = \frac{(3 - \beta)M_{tot}}{4\pi(R_{out}^{3-\beta} - R_{in}^{3-\beta})}r^{-\beta} \quad (2.3)$$

if $\beta \neq 3$, where r is the radial distance from the centre of the cell to the origin and M_{tot} is the total desired dust mass to model. If $\beta = 3$ then the dust density in each cell is alternatively calculated as

$$\rho(r) = \frac{M_{tot}}{4\pi \log(R_{out} - R_{in})}r^{-3} \quad (2.4)$$

Any cell whose centre falls outside of the bounds of the supernova ejecta has dust density set to zero. If the dust and gas are decoupled then the user must specify distinct profiles for the gas and the dust; that is, separate power laws must be declared and independent inner and outer radii specified. The same process is followed but with separate power-laws for each component. Including the capacity to specify the gas and dust distributions separately allows for more sophisticated modelling of, for example, circumstellar shells or dense cores of dust formation surrounded by more diffuse gas.

Clumped geometries

It is known from SED modelling that models of clumped environments produce very different results to environments assumed to have a smooth distribution of dust and gas

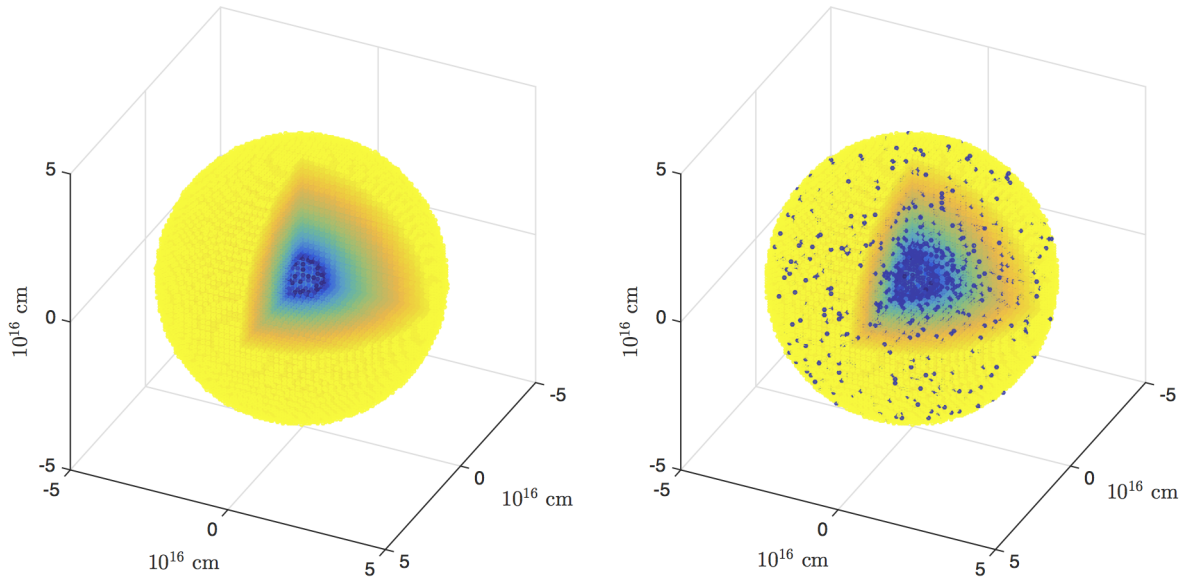


Figure 2.2. 3D representations of the grid generated by DAMOCLES. A smooth distribution is shown on the left and a clumped distribution on the right.

(e.g. Bianchi et al. (2000); Ercolano et al. (2007); Owen & Barlow (2015)). The capacity for modelling a clumped dusty medium is therefore included in the code. The fraction of the dust mass that is in clumps is declared (m_{frac}) and the total volume filling factor of the clumps (f) is also specified. Dust that is not located in clumps is distributed according to a smooth radial power-law. The clumps occupy a single grid cell and their size can therefore be varied by altering the number of divisions in the grid. They are distributed stochastically with probability of a given cell being a clump proportional to the smooth density profile (i.e. $p(r) \propto r^{-\beta}$). The density of all clumps is constant and is calculated as

$$\rho_{clump} = \frac{M_{clumps}}{V_{clumps}} = \frac{m_{frac} M_{tot}}{\frac{4}{3} f \pi (R_{out}^3 - R_{in}^3)} \quad (2.5)$$

where M_{tot} is the total dust mass, M_{clumps} is the total dust mass in clumps and V_{clumps} is the total volume occupied by clumps. m_{frac} and f are defined as above.

A grid of cubic cells of varying dust and gas densities is thus produced in readiness for packets to be transported through it. Examples of a smooth and clumped distributions of dust generated by DAMOCLES are presented in Figure 2.2. A frequency grid is also established centred on the rest-frame frequency of the line to be modelled.

2.2.3 Properties of the Dusty Medium

Dust of any composition may be used for which optical data is available. The relative abundances of the species must be declared in an input file accompanied by a grain radius distribution (specified as a grain radius range and power-law) for each species. Files detailing the optical data (n and k values) for the chosen dust species are also declared at the start of the code. For each pairing of wavelength λ and grain radius a , a Mie scattering routine is employed to calculate $Q_{abs}(\lambda, a)$ and $Q_{sca}(\lambda, a)$ from the refractive index $n + ik$. These values are calculated for every species across the wavelength and grain radius ranges of interest. I have described the mathematics of converting the refractive index into scattering and absorption efficiencies using Mie theory in detail in the introduction to this thesis (see Section ??).

Ultimately, the overall scattering and absorption opacities in each grid cell must be known and so a weighted summation over $Q_{abs}(\lambda, a)$ and $Q_{sca}(\lambda, a)$ is performed. The number density in each cell must first be calculated as $n = \rho/m_{av}$ where m_{av} is the average mass of a grain:

$$m_{av} = \sum_i \sum_a \frac{4}{3} \pi a_i^3 \rho_{g,i} w_i(a) x_i \quad (2.6)$$

$w_i(a)$ represents the normalised weight of particles with grain radius a , $\rho_{g,i}$ is the density of the grain for species i (specified with the optical data) and x_i is the relative abundance of species i .

The total cross-section of interaction for extinction is then calculated as

$$C_{ext}(\lambda) = \sum_i \sum_a Q_{ext,i}(a, \lambda) w_i(a) \pi a_i^2 x_i \quad (2.7)$$

where the subscript i denotes species i , x_i represents the relative abundance of species i and the summations are over species and grain radius. The calculation of the cross-section of interaction for scattering $C_{sca}(\lambda)$ is performed in the same manner.

Using equations 2.6 and 2.7, the opacity κ can then be calculated using the relationship

$$n C_{ext} = \rho \kappa_{ext} \quad (2.8)$$

The values of $C_{ext}(\lambda)$ and $C_{sca}(\lambda)$ are calculated for the full wavelength range at the start of the simulation as are the number densities in each grid cell. As a packet passes through a grid cell, the optical depth τ is determined from the above quantities according to the wavelength of the current packet (see Section 2.2.4 for further detail on this process). The above equations are discretised versions of continuous integral equations given in van de Hulst (1957) and Tielens (2005).

2.2.4 Emission and Propagation

The initial radiation field is inherently tied to the distribution of gas throughout the supernova ejecta. The relationship between the emissivity and the gas density may vary under different regimes and therefore the emissivity distribution is also specified as a power-law with $i(\rho) \propto \rho^q$. In general, however, the emissivity distribution is assumed to be proportional to the square of the local gas density ($i(r) \propto r^{-2\beta}$), i.e. proportional to the product of the recombining ion and electron densities in the case of a recombination line or to the product of the neutral atoms and electron densities in the case of collisionally excited line emission.

The supernova ejecta is divided into shells between R_{in} and R_{out} and the number of packets to be emitted in each shell calculated according to the specified power-law emissivity distribution $i(r) \propto r^{-q\beta}$. For each packet a location within the appropriate shell must be determined and a propagation direction sampled. The initial propagation direction is sampled from an isotropic distribution as detailed in numerous publications (e.g. Wood et al. (2004)). Three random numbers in the range $[0, 1]$ are sampled and these are translated into spherical coordinates as

$$\phi = 2\pi\eta \tag{2.9}$$

$$\theta = \arccos(2\xi - 1) \tag{2.10}$$

where η and ξ are random numbers in the range $[0, 1]$, ϕ is the azimuthal angle and $\cos \theta$

is the radial direction cosine. The initial packet trajectory in cartesian is then given by

$$n_x = \sin \theta \cos \phi \quad (2.11)$$

$$n_y = \sin \theta \sin \phi \quad (2.12)$$

$$n_z = \cos \theta \quad (2.13)$$

The process for sampling an initial position within the shell follows the same sampling process as described above but a radial position must be sampled in addition to the two angular coordinates. For a random number ζ , the radial position r is calculated as

$$r = R_i + \zeta(R_{i+1} - R_i) \quad (2.14)$$

where R_i is the inner boundary of the i^{th} shell.

In total therefore five random numbers must be sampled in order to generate an initial position and trajectory for a packet. At every subsequent scattering event, the packet is propelled with a new direction vector which is sampled from an isotropic distribution in the rest-frame of the particle with two newly generated random numbers in the manner described above. I do not include forward-scattering matrices in the code since the effects of forward scattering by dust are so small as to be negligible and it is simpler and more efficient simply to assume isotropic scattering.

Once a packet has been emitted into the nebula, it must be propagated through the grid until it escapes the outer bound of the ejecta R_{out} or is absorbed. In each cell that a packet passes through, a test must be performed in order to determine whether the packet passes through that cell and into the next or whether it is scattered or absorbed by a dust grain (i.e. an “event” occurs). The probability that a packet travels a distance l without interacting is

$$p(l) = e^{-n\sigma_\lambda l} = e^{-\tau_\lambda} \quad (2.15)$$

where n is the number density of the grid cell, σ_λ is the cross-section of interaction at wavelength λ and

$$\tau_\lambda = n\sigma_\lambda l = \rho\kappa_\lambda l \quad (2.16)$$

for constant n and σ_λ (as in a grid cell). Note that σ_λ is used to denote the cross-section of interaction for a given grain radius. Where a grain radius distribution is adopted or multiple species are employed, the formula becomes $\tau_\lambda = nC_{ext,\lambda}l$ as described in Section 2.2.3. The probability that a packet *does* interact within a distance l is therefore $1 - e^{-\tau_\lambda}$. The position at which the packet will be absorbed or scattered is then determined by comparing a randomly generated number in the interval $[0,1)$ with this value. In practice however, it is easier to sample from the cumulative probability distribution and use the generated number to calculate an optical depth as

$$\xi = 1 - e^{-\tau_\lambda} \implies \tau_\lambda = -\log(1 - \xi) \quad (2.17)$$

where $\xi \in [0, 1)$ is a sampled random number between 0 and 1. The properties of the cell as determined in Section 2.2.3 are then used to determine the distance that the packet travels:

$$l = \frac{\tau_\lambda}{nC_{ext,\lambda}} \quad (2.18)$$

If the distance to be travelled by the packet is greater than the distance from its position to the edge of the cell then the packet is moved along its current trajectory (n_x, n_y, n_z) to the cell boundary and the process is repeated. Alternatively, if the displacement is not sufficient for the packet to escape the cell then an event occurs. The packet is either scattered or absorbed with probability of scattering equal to the albedo of the cell

$$\omega = \frac{\sigma_{sca}}{\sigma_{sca} + \sigma_{abs}} \quad (2.19)$$

If the packet is absorbed (the case if a randomly generated number is greater than the albedo ω) then it is simply removed from the simulation as previously discussed. If this is not the case, then the packet is scattered and a new trajectory is sampled from an isotropic distribution in the comoving frame of the dust grain. The frequency of the packet is recalculated using Lorentz transforms as described in the next section and the process is repeated until the packet has either escaped the outer boundary of the supernova ejecta or been absorbed. If the packet does escape, its weighted energy is deposited in the appropriate frequency bin. Once all packets have escaped, the array of frequencies and fluxes produces the desired line profile.

2.2.5 The Velocity Field and Doppler shifting

At emission and at each scattering event the frequency of the packet is recalculated according to a radial velocity field

$$v(r) = v_{max} \frac{r^\alpha}{R_{out}^\alpha} \quad (2.20)$$

where the maximum velocity, v_{max} , at the outer edge of the ejecta and the exponent of the velocity profile, α , are declared in the input file.

It is worth noting that if a constant mass loss rate is required, the exponent of the velocity profile and the exponent of the density profile are not independent. A constant mass loss rate implies that $4\pi\rho v R^2 \propto k$, where k is a constant, and thus for $v \propto r^\alpha$ and $\rho \propto r^{-\beta}$, we require that $\beta - \alpha = 2$. However, it is possible that the supernova event may have induced a mass-flow rate that is not constant with radius and thus both exponents may be declared independently. It is also worth noting that for supernovae in their free expansion phase, as the majority are by the time of the onset of dust formation, the ejecta are expanding with a $v \propto r$ Hubble law expansion.

The outflow velocities in supernovae are extremely high, of the order of 10% of the speed of light. Escaping radiation is therefore subject to significant Doppler shifting. At emission and at each scattering event, the frequency of a packet must be recalculated according to the velocity of the scattering or emitting grain. When the packet is initially emitted, it has a frequency and a trajectory in the rest frame of the emitter. Both of these must be transformed to the observer's frame in order for the packet to be propagated through the grid. The new direction and frequency in the observer's frame may be simply found by transforming the momentum 4-vector \mathbf{P} which is defined as

$$\mathbf{P} = \begin{pmatrix} E \\ p_x \\ p_y \\ p_z \end{pmatrix} = \begin{pmatrix} h\nu \\ h\nu x \\ h\nu y \\ h\nu z \end{pmatrix} \quad (2.21)$$

We may then derive \mathbf{P}' , the momentum 4-vector in the observer's frame using the relation

$$\mathbf{P}' = \Lambda \mathbf{P} \quad (2.22)$$

where

$$\Lambda = \begin{pmatrix} \gamma & -\gamma\beta_x & -\gamma\beta_y & -\gamma\beta_z \\ -\gamma\beta_x & 1 + (\gamma - 1)\frac{\beta_x^2}{\beta^2} & (\gamma - 1)\frac{\beta_x\beta_y}{\beta^2} & (\gamma - 1)\frac{\beta_x\beta_z}{\beta^2} \\ -\gamma\beta_y & (\gamma - 1)\frac{\beta_y\beta_x}{\beta^2} & 1 + (\gamma - 1)\frac{\beta_y^2}{\beta^2} & (\gamma - 1)\frac{\beta_y\beta_z}{\beta^2} \\ -\gamma\beta_z & (\gamma - 1)\frac{\beta_z\beta_x}{\beta^2} & (\gamma - 1)\frac{\beta_z\beta_y}{\beta^2} & 1 + (\gamma - 1)\frac{\beta_z^2}{\beta^2} \end{pmatrix} \quad (2.23)$$

and $\beta = \frac{\mathbf{v}}{c} = (\beta_x, \beta_y, \beta_z)$, $\beta = |\beta|$ and $\gamma = \frac{1}{\sqrt{1-\beta^2}}$.

In practice, the velocities considered are low enough that it is unnecessary to consider terms of order $O(\frac{v^2}{c^2})$ and thus Λ may be reduced to

$$\Lambda = \begin{pmatrix} 1 & -\beta_x & -\beta_y & -\beta_z \\ -\beta_x & 1 & 0 & 0 \\ -\beta_y & 0 & 1 & 0 \\ -\beta_z & 0 & 0 & 1 \end{pmatrix} \quad (2.24)$$

The new direction of travel and frequency in the observer's frame are therefore given by

$$\nu' = \nu(1 - x\beta_x - y\beta_y - z\beta_z) \quad (2.25)$$

$$x' = \frac{\nu}{\nu'}(x - \beta_x)$$

$$y' = \frac{\nu}{\nu'}(y - \beta_y)$$

$$z' = \frac{\nu}{\nu'}(z - \beta_z)$$

For each scattering event, the packet must be transformed both into and out of the comoving frame. The reverse transform is applied by using the inverse Lorentz matrix Λ^{-1} which is obtained by reversing the sign of \mathbf{v} . Positive \mathbf{v} is defined for frames moving away from each other and thus \mathbf{v} is defined to be negative in the direction of the observer.

2.2.6 Electron Scattering

As will be discussed in detail in chapter three, the effects of scattering on the shapes of line profiles can be quite pronounced and it is therefore important to consider the potential

Table 2.1. Values of $q_{H\alpha}(T)$ at three different temperatures as used by DAMOCLES.

	Temperature (K)		
	5,000	10,000	20,000
$q_{H\alpha}$ (erg cm ³ s ⁻¹)	6.71×10^{-25}	3.56×10^{-25}	1.83×10^{-24}

effects of electron scattering as well as those of dust scattering. A simple treatment of electron scattering calculates electron densities using an estimated average temperature of either 5,000K, 10,000K or 20,000K. An observed luminosity of $H\alpha$ is then used to calculate the optical depth to electrons. The overall optical depth within each cell is calculated as $\tau = \tau_{dust} + \tau_e$, with $\tau_e = 0$ if electron scattering is not activated. The electron scattering optical depth, τ_e , in a given cell (with constant properties therein) is calculated as

$$\tau_e = n_e \sigma_t l \quad (2.26)$$

where n_e is the electron density in that cell, σ_t is the Thomson cross-section of interaction for an electron and l is the distance travelled. In order to calculate this value, the electron density in each cell must be known. We assume that the electron density is the same as the proton density and that both are distributed according to the gas density distribution such that

$$n_e(r) = Kr^{-\beta} \quad (2.27)$$

where K is a constant. The value of K must be determined from the total $H\alpha$ luminosity. We follow the formalism described by Osterbrock & Ferland (2006) in order to estimate the electron density from the total $H\alpha$ luminosity ($L_{H\alpha}$). $L_{H\alpha}$ is given by

$$L_{H\alpha} = \int_0^{\infty} n_p(r) n_e(r) E_{H\alpha} \alpha_{H\alpha}^{eff}(T) 4\pi r^2 dr \quad (2.28)$$

$$= \int_{R_{in}}^{R_{out}} n_p(r) n_e(r) q_{H\alpha}(T) 4\pi r^2 dr \quad (2.29)$$

where $n_p(r)$ is the proton density at radius r , $n_e(r)$ is the electron density at radius r , T is

the temperature, $\alpha_{H\alpha}^{eff}(T)$ is the temperature-dependent effective recombination coefficient for H α , $E_{H\alpha}$ is the energy of a single H α photon and

$$q_{H\alpha} = E_{H\alpha} \alpha_{H\alpha}^{eff} = \frac{4\pi j_{H\alpha}}{n_e n_p} \quad (2.30)$$

where $j_{H\alpha}$ is the temperature-dependent emission coefficient for H α (i.e. the energy emitted per unit volume per unit time per unit solid angle). Substituting equation 2.27 into equation 2.29 gives the following

$$\frac{L_{H\alpha}}{4\pi q_{H\alpha}} = k^2 \int_{R_{in}}^{R_{out}} r^{2(1-\beta)} dr \quad (2.31)$$

which in the case $\beta \neq \frac{3}{2}$ may be solved as

$$K = \sqrt{\frac{L_{H\alpha}}{4\pi q_{H\alpha}} \frac{3 - 2\beta}{R_{out}^{3-2\beta} - R_{in}^{3-2\beta}}} \quad (2.32)$$

and for $\beta = \frac{3}{2}$ is

$$K = \sqrt{\frac{L_{H\alpha}}{4\pi q_{H\alpha}} \frac{1}{\ln(R_{out}/R_{in})}} \quad (2.33)$$

Substituting K back into equation 2.27 gives the electron density for each cell. In the code, only three gas temperatures may be specified and three corresponding values of $q_{H\alpha}(T)$ are included as per Table 2.1.

If, for a given packet, an event occurs, it is first calculated whether this is an electron scattering event or a dust event (either scattering or absorption) by considering the ratio of the optical depths to each. The process by which a packet is scattered by an electron is almost identical to the dust scattering process except for the adopted velocity of the scatterer. In the case of a dust grain, the velocity is simply the bulk velocity of the ejecta at that radius as determined from the specified velocity profile. For an electron, the assumed velocity must include a thermal component as well as the same bulk velocity as would be adopted for a dust grain at the same location. As per the electron scattering calculation of Hillier (1991), the components (v_x, v_y, v_z) of the thermal velocity \mathbf{v}_{therm} are assumed to follow a Maxwellian distribution with zero mean and standard deviation

$$\sigma = \sqrt{\frac{k_B T}{m_e}} \quad (2.34)$$

where k_B is Boltzman's constant and m_e is the mass of an electron. The components are then sampled from a normal distribution with specified mean and standard deviation using the Marsaglia polar method (Marsaglia & Bray 1964). This method generates two random numbers from a uniform distribution in the interval [0,1) and uses a number of transformations to convert them to random numbers as generated from a standard normal distribution with zero mean and unity variance. They may then be scaled to the appropriate normal distribution. Finally, the overall velocity of the electron is then calculated as

$$\mathbf{v}_e = \mathbf{v}_{bulk} + \mathbf{v}_{therm} \quad (2.35)$$

and the Lorentz transforms are applied in the same manner as a dust scattering event.

In the majority of cases it seems that the electron densities are not high enough to discernibly effect the overall shape of the profile. However, there may be a few rare cases (the concept is discussed for SN 2010jl (Fransson et al. 2013)) where the electron densities are high enough to become significant in the observed profiles. Whilst the inclusion of electron scattering in the code is an approximation since it is not necessarily true that $n_e = n_p$ and the exact gas temperature is unknown, it gives a good suggestion of the potential effects of electron scattering.

2.2.7 Doublets

One of the lines in supernovae emission spectra that is frequently seen to be blue shifted is the forbidden [OI] $\lambda\lambda 6300, 6363\text{\AA}$ doublet. DAMOCLES therefore has the capacity to treat doublets as well as single lines. When a doublet is specified, both the initial wavelengths and the initial intensity ratio must be declared. The code will create a wider frequency array than for a single line in order to accommodate both lines. It will then model each line independently, adding the final fluxes of the lines weighted by their intrinsic flux ratio to produce the desired doublet at the end of the modelling.

2.2.8 Comparing the Model with Observations

DAMOCLES includes the capacity to read in observed line profile data for direct comparison with a modelled line profile. Once all packets have been processed through the nebula and collected into bins, a flux is calculated at each of the wavelength bins in the

observed data by interpolating between modelled wavelength bins. A mean squared error (MSE) calculation is then performed to compare the model with the data quantitatively, where the MSE is equal to

$$\frac{1}{N} \sum_i (f_{obs,i} - f_{mod,i})^2 \quad (2.36)$$

and $f_{obs,i}$ is the observed flux in the i^{th} frequency bin, $f_{mod,i}$ is the modelled flux in the i^{th} frequency bin, and N is the total number of frequency bins. Minimising the MSE minimises the error between the model and the observed line and therefore provides a quantitative measure of goodness of fit that may be used in addition to or instead of any qualitative assessment. Since the total inherent error on each observation is variable, the exact value of the MSE should not be compared between different line profile observations and only between different models and sets of parameters for a given line profile.

2.3 The Structure of DAMOCLES

DAMOCLES is written using Fortran 95. Since the major modernisation of Fortran 77 in 1990, the language includes a number of more modern elements that make it an ideal choice for this type of numerical computation. Firstly, a fast, high-level language is required that allows for dynamic memory allocation and deallocation. Whilst DAMOCLES could have been written in a number of other languages, this is a critical feature that is only available in a few languages. Very large numbers of packets are required to achieve reasonable resolutions in Monte Carlo codes of this nature and therefore large arrays of data are required. The ability to maintain careful control of memory allocation is very important.

Fortran 95 also has a number of other features that make it especially suitable for this sort of code. Derived types group a number of variables of different intrinsic or other derived types. This allows different properties of a particular item (for example a packet or grid cell) to be grouped together and accessed via that item. Though not a necessary feature, derived types make the code simpler, faster and more legible. They also make it easier to write and therefore help to minimise the risk of errors. Similarly, the modular structure that was introduced to Fortran in 1990 allows the programmer to distribute their code over a number of modules and ensures that variables that are declared within a particular module can be accessed by other modules if necessary (Ellis et al. 1994).

This eliminates the need for common blocks of code and allows a large program to be segmented into logical divisions. This increases the speed, clarity and ease of maintenance and development in the future.

The obvious alternative programming language to Fortran 95 is C or C++. Both of these languages have all of the features described above and are exceptionally fast. From a computation perspective, there is, arguably, little to separate them for this type of coding. I ultimately decided to write DAMOCLES in Fortran 95 because of its heritage in astrophysics. A very large number of astrophysical codes have been written using current or previous versions of Fortran and writing the code in Fortran 95 allowed for easy compatibility and the use of various astrophysics libraries and routines.

DAMOCLES is parallelised using OpenMP (see Section 2.3.2) which restricts its use to shared memory machines. It has been developed on and currently runs on a MacBook Pro 11.2 quad core with Intel Core i7 2.8GHz processors and 16GB of memory. A typical, medium resolution simulation using 125,000 grid cells and 10^5 packets takes approximately 15 seconds to run. The number of packets transported and the total dust optical depth are the most important factors in determining runtime.

2.3.1 Computational Architecture and Processes

DAMOCLES was written using a modular structure. The “parent” driver has numerous “children” in the form of subroutines and modules which are each responsible for a separate task or tasks. This architecture has a number of advantages. Firstly, it serves to clarify both the functionality and legibility of the code allowing for easier debugging and maintenance. It also allows for the implementation of features such as recursive subroutines which are ideally suited to a Monte Carlo methodology. Finally, it allows for the code to be developed further in the future simply by including additional modules and subroutines. A brief description of every module and subroutine in the code is presented in the following subsections. The descriptions are ordered according to the first time they or their contents are called by the driver (see Figure 2.1 for a flowchart of the order of the processes that take place in DAMOCLES and see Figure 2.3 for a flowchart of the modular hierarchy).

The driver

The *driver* module is at the centre of DAMOCLES. It is from here that all subroutines are called. The calls to construct the grid and calculate dust opacities, to emit and propagate packets and to compare the results with observational data are all made from here. The parallelisation process is also controlled from here (see section 2.3.2 for more details on the parallel function of DAMOCLES). Having called the initialisation routines, the driver is responsible for dividing the ejecta into shells and calculating how many packets are emitted within each shell. Each shell is looped over and each packet is looped over within each shell. Emission and propagation routines are called inside this loop. At the end of each packet’s lifetime, either once it has been absorbed or has escaped, the driver adds the weighted packet’s energy to the appropriate frequency bin and stores this information before looping back to emit and propagate the next packet. It is here that a line of sight is applied if so desired. This is achieved by collecting only packets that have escaped within a cone of vertical angle $\pi/6$. Once all packets have been processed, the driver writes the relevant information (the wavelengths, velocities and fluxes that describe the outputted line profile) to an output file and calls the model comparison module.

The driver is also the section of code responsible for processing doublets. The code treats doublets by processing two batches of packets with differing initial frequency through the same grid. Before they are collected in frequency bins, the flux ratio that is specified by the user is applied to one batch of the packets. All packets are then collected as per a single line.

Various statistics are also processed and output here including the fraction of packets that are absorbed and the estimated undepleted luminosity of the observed line.

The input module

The *input* module is where the primary input file is read into the code and all global variables are declared and assigned. A number of logicals are assigned based on values declared in the input file and some simple calculations are performed that determine the inner and outer radii based on the maximum velocity specified and the epoch of consideration ($R_{out} = V_{max} \times t$). A number of physical constants that are used throughout the code are also declared here as “parameters”, meaning that their value cannot be changed at any point in the simulation.

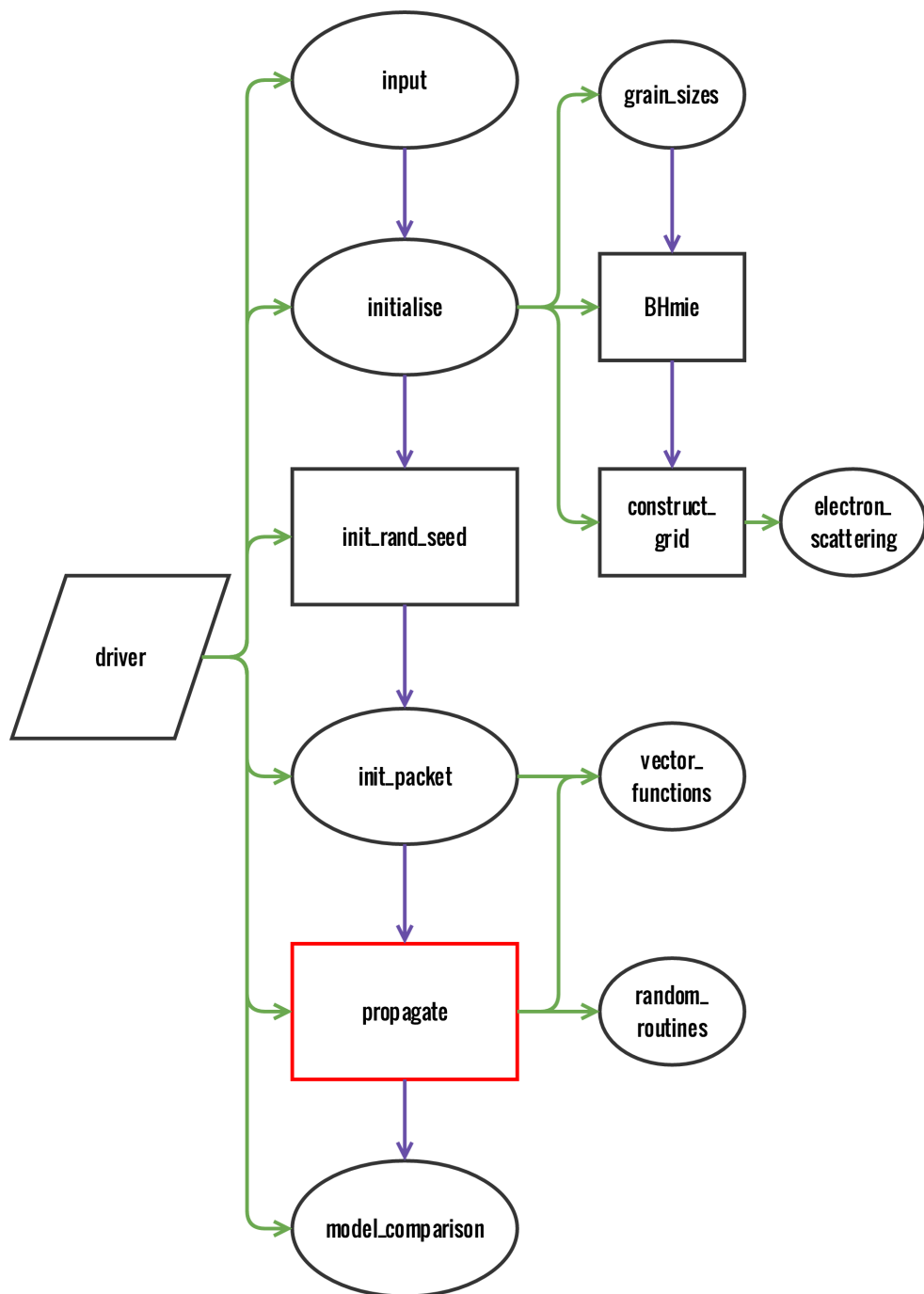


Figure 2.3. A flowchart representing the hierarchy of modules and subroutines in the DAMOCLES code. Ellipses represent modules and rectangles represent subroutines (the red rectangle is a recursive subroutine). Green arrows indicate the dependence of a module or subroutine on previous modules or subroutines. Purple arrows indicate the flow of the code.

The initialisation module

The *initialise* module acts as a driver to run all of the subroutines associated with initialising the program. A number of dynamically allocatable arrays are declared allowing for a grid of densities to be calculated, a frequency grid to be stored and optical properties to be read in. Arrays to store the emergent spectrum are also declared. The calculation of dust opacities, which calls the *grain_sizes* subroutine and the *BHmie* subroutine, is performed here. For each species, the wavelength-dependent optical properties, n and k , are read in and the Mie routine applied to every pair of frequencies and grain radii. The resulting extinction and scattering efficiencies are summed over all grain radii for each wavelength, weighted appropriately, to calculate overall wavelength-dependent extinction and scattering opacities (see Section ?? for further detail). These data are stored in an array that is accessed as necessary when packets are propagated through the grid.

The command to construct the grid is called before some basic statistics about the grid are calculated. The average optical depth from R_{in} to R_{out} in both the V band and the rest-frame wavelength of the line being modelled are calculated and sent to stdout. The average number density of grains in each cell is also computed and output. Finally, the frequency array is constructed.

This module is also where the ‘gridcell’ derived type is declared. A ‘gridcell’ type was specified as it allowed for easy and clear access to any of a grid cell’s properties as a packet passed through it. The type consists of a number of arrays of real, integer and logical variables. The properties recorded for each cell include the physical bounds of the cell in each axis, the mass and number dust densities, the electron density, an identifying number (ID) and a logical clumped property.

The grain size module

The *grain_sizes* module reads in the file that specifies the list of species to be used. This file is a list of species detailing the name of the file containing the optical data for the species, the relative abundance of that species, the maximum and minimum grain radii and the exponent of the power law of the grain radius distribution. It also declares how fine the grid of grain radii should be. These properties are all read in by the *grain_sizes* module and a relative weight for each grain radius for each species is calculated here.

The ‘species’ derived type is declared in this module. Similarly to the ‘gridcell’ type,

using a derived type allowed for the easy storing and accessing of a large number of properties of each species. Many multi-dimensional arrays and scalars are stored for the ‘species’ type including properties relating to the grain radius distribution, the density of a dust grain, the extinction and scattering opacities and the relative abundance of the species amongst others. After the processing of the optical data for all species is completed, the calculated quantities are stored in arrays as components of the ‘species’ type.

The Mie approximation subroutine

The *BHmie* subroutine is a standard routine that was obtained from an online library of routines. It is a modified version of the Bohren & Huffman (1983) Mie scattering routine. The algorithm applies the mathematics described in Section ?? to determine the extinction and scattering efficiencies of a single size grain at a specified wavelength given its complex refractive index $n + ik$.

The grid construction subroutine

The *construct_grid* subroutine is called from within the initialisation module. The purpose of this subroutine is to populate the grid, which is an array of derived type *gridcell* and size n_{cells} where n_{cells} is the number of cells in the grid. The bounds of the grid are initialised and the radii of all cells from the centre of the grid to the centre of the cell are calculated. The density of each cell is then calculated according to a smooth power-law density distribution and scaled so that the total dust mass is equal to that specified in the input file. If clumps are used then the total number of clumps is calculated and these are distributed throughout the grid stochastically according to the smooth density profile stipulated. This subroutine also calls the electron scattering subroutine contained within the *electron_scattering* module so that the electron density of a cell may be stored at the same point as the dust density.

The electron scattering module

The *electron_scattering* subroutine is a simple subroutine that is used to calculate the value of K as described in equation 2.27. The total H α luminosity and the gas temperature are read in and the gas temperature used to determine the appropriate value of $q_{H\alpha}$ from Table 2.1. These values are then used to calculate the value of K as described by equations 2.32 and 2.33. The variable is passed back to the *grid construction* subroutine where it is used

to calculate the electron density in each cell. The electron densities will be used by the *propagate* subroutine to calculate the electron scattering optical depth in each cell.

The random seed subroutine

The *init_rand_seed* is a short subroutine that calculates a seed for the standard Fortran pseudo-random number generator (*random_number*). It uses the system clock to generate the random seed and thus varies with every implementation of the code. A seed is a number that is used as a “starting point” for a pseudo-random number generator. Varying the random seed ensures that a different set of random numbers is generated every time the code is run, which can be useful to ensure that any peculiar or interesting features of the outputted line profiles are definitely a product of the physical processes involved and not a result of random fluctuations in the simulation. The more packets are used however, the more the Monte Carlo noise in the emergent line profile is reduced and the contribution from any anomalous packets should be insignificant.

The packet initialisation module

The *init_packet* module is responsible for the creation and emission of packets at the start of the simulation. It is called from the driver for each packet. By generating an array of five random numbers, the position and emission direction vectors in the rest frame of the emitter are calculated according to the formulae described in equations 2.9 to 2.14. The scalar velocity of the emitter is calculated based on its radial position and this converted into a velocity vector by normalising the position vector and multiplying by the scalar velocity. The velocity vector is passed to the Lorentz transforms subroutine contained in the *vector_functions* module. The frequency of the packet is also passed to this subroutine. After the propagation direction vector and the frequency of the packet have been updated to the observer’s rest frame, the grid cell in which the packet starts its path is identified and the code passes back to the driver to propagate the packet through the nebula.

The vector functions module

A number of vector functions are contained within the *vector_functions* module and are accessed throughout the program. These include normalisation functions, conversions from spherical coordinates to cartesian and both forward and inverse Lorentz transforms. It is the latter of these that are most important for the physics of the code. The Lorentz

functions are called for each packet at emission from the *driver* and at every subsequent scattering event from within the *propagate* routine. As well as performing the necessary frequency shift based on the velocity of the scatterer or emitter, they also transform into and out of the rest frame of the particle thus ensuring that the packet is propagated through the nebula with a direction in the rest frame of the observer but that its new direction is sampled from an isotropic distribution in the rest frame of the emitting or scattering dust grain.

The β and γ values are calculated based on the input velocity vector. The momentum 4-vector \mathbf{P} is then multiplied by the Lorentz matrix $\mathbf{\Lambda}$ using the Fortran function *matmul* to produce a new frequency and a new direction vector in the appropriate frame of reference. If a scattering event has occurred then the weight of the packet is also updated here. The new direction vector, frequency and weight are then passed back to the propagate routine and the process repeated. At each scattering event the inverse Lorentz matrix must first be applied to move from the observer's rest frame to the particle's. A new direction vector must then be sampled from an isotropic distribution before applying the forward Lorentz transform to move back from the rest frame of the dust grain to the observer's frame. The next step in the packet's trajectory may then be calculated in the *propagate* subroutine.

The *propagate* subroutine

The *propagate* subroutine is at the heart of the Monte Carlo simulation. It is here that the trajectories of all packets in the simulation are determined. The *propagate* subroutine is a subprogram called a *recursive subroutine*. This allows the subroutine to call itself, at which point it will loop back to the start of the subroutine. It will continue this process until a condition is reached that instructs it to return to the driver. In this case a number of conditions will arrest the circulation of the packet. If the packet has escaped the outer radius of the ejecta or has been absorbed then the routine will pass this information along with the frequency and weight of the packet back to the driver. The routine would also stop recurring if a packet has undergone a maximum number of scattering events (500 by default). At this point it is deemed that the weight of the packet is so small as to be negligible and it is classified as “inactive”. This prevents the code from lagging by becoming stuck on a particular packet that has become trapped in a region of high density and albedo. It is noted that if this is the case for a large number of packets then a bias may be introduced - packets emitted in particularly high density regions may be discarded

more frequently than those emitted in less dense regions. The number of packets that are deemed “inactive” is output as a percentage of the total number of packets employed at the start of the simulation as a check for the user. In practice, unless the albedo of the dusty medium is extremely high and the medium is very dense, this is rarely an issue (for an average simulation with 10^7 packets, normally only one or two are discarded for this reason).

There are a number of processes that take place in this module in order to propagate a packet through the nebula accurately. A full pictorial representation of the procedures that are implemented in this module may be found in the flowchart in Figure 2.4. For each packet in each grid cell, the optical depth in that cell is calculated based on the dust density and the opacity at the wavelength of the current packet. These are obtained by interpolating between discrete opacities at points in the frequency array. At this stage, the Monte Carlo technique is applied in order to determine the distance travelled by the packet by sampling from the cumulative probability distribution. This displacement is then compared to the distance from the packet’s current location to the edges of the grid cell in order to ascertain whether or not the packet escapes the cell. If it does escape then the packet advances to the bounds of the current grid cell and the process is repeated in the next cell. If it does not escape then an event occurs. In this case, random numbers are sampled in order to determine whether the packet experiences electron scattering, dust scattering or absorption. If the packet is absorbed then it is removed from the simulation, the *propagate* subroutine is arrested and the code returns to the driver to emit a new packet. If it is scattered then the velocity vector is calculated. In the case of electron scattering this involves considering the thermal velocity component as well as the bulk velocity at that radius. The Lorentz transforms are applied based on the velocity vector and the frequency and weight of the packet are updated. A new direction of propagation is sampled in the scatterer’s rest frame from an isotropic distribution and is transformed into the observer’s rest frame. The routine is recalled to start afresh with the new propagation direction.

The random routines module

The *random_routines* module contains a single subroutine which is, like the *BHmie* routine, a standard routine obtained from an online library. It allows for a random velocity vector to be sampled from a normal distribution with specified mean and standard deviation. The

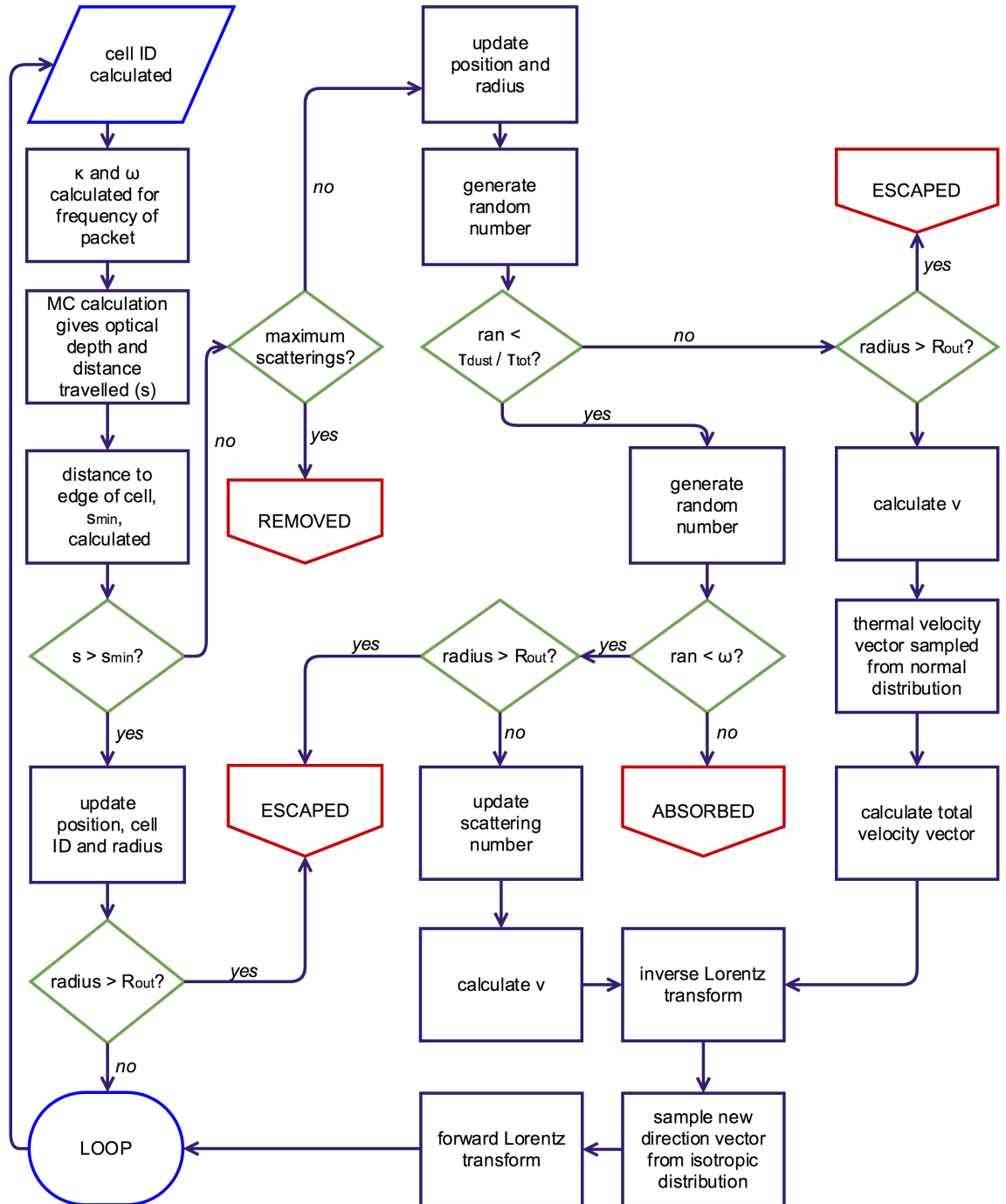


Figure 2.4. A flowchart representing the processes that occur in the *propagate* subroutine. The life of a packet passing through the grid may be determined by following the flowchart starting at the blue parallelogram (top left). Purple ovals indicate a standard step in the evolution, green diamonds indicate that a determination must be made, red boxes mean that the packet's evolution has concluded as it has escaped, been absorbed or been removed and the blue rounded oblong indicates a return to the start of the routine.

standard deviation is calculated as per equation 2.34 and passed to the subroutine which samples a random 3-dimensional vector from the normal distribution with the specified standard distribution and zero mean. This is passed back to the *propagate* routine where it is added to the bulk velocity in order to determine the overall velocity of the scattering electron.

The model comparison module

The *model_comparison* module is responsible for post-processing the outputted line profile and comparing the model results with inputted observed data. The routine interpolates between two model frequency points to obtain a flux value at each frequency point of the observed line profile. Both profiles are then normalised such that the total flux is unity. A MSE calculation is then performed as per equation 2.36. The smaller the value of MSE, the better the fit is. It should be noted however that data with a poor signal-to-noise ratio will have an inherently large MSE than data with a good signal-to-noise ratio.

2.3.2 OpenMP Parallelisation

Monte Carlo simulations are exceptionally well-suited to parallelisation. The path of each packet through the nebula is unaffected by the transport of any other packet. It is therefore possible to run multiple instances of the *propagate* module at once by using several threads. Since the vast majority of the processing power of the simulation is driven from this module, it is theoretically possible to achieve a nearly linear speed-up; i.e. if the number of cores is doubled, the run time should be approximately halved.

DAMOCLES was parallelised using OpenMP. OpenMP is an Application Program Interface (API) that allows for shared-memory parallel programming in Fortran and C/C++. OpenMP causes the code to be run serially on a single processor until a parallel region is reached. At this point the single master thread branches into multiple threads, and multiple instances of the same section of code are run on each. In DAMOCLES, this splitting occurs at the start of the loop which controls the emission and propagation of packets from each shell. If, for example, 10^7 packets are emitted and 5 threads are used, then approximately 2×10^6 packets will be independently processed on each thread. Practically however, the OpenMP keyword *dynamic* is declared ensuring that, as soon as each thread has finished processing a packet, it immediately moves onto the next one. If the *static* keyword were specified instead then the number of packets to be processed would

be equally divided between the threads at the start of the loop. In this case, if, by random chance, one thread happened to have significantly more absorbed packets than another, then potentially utilisable processing power would be lost as the core waited for the others to finish.

At the start of the parallel region variables accessed within the shared region are specified as shared or private. Private variables are not seen by other threads and allow the value of a single named variable, for example “frequency”, to have different values on different threads. Shared variables have the same value regardless of the thread number, for example, “grid cell density”. As each packet escapes, its weighted energy must be added to the final energy array. It is important that two threads do not attempt to alter the value of this shared array at the same time as data may be lost or corrupted. This section of code is therefore enclosed inside a *critical* region. This instruction ensures that code in this block is to be executed by only one thread at a time. Extensive testing was performed to ensure that outcomes were not affected by the implementation of a parallel environment.

For further information about the OpenMP API please refer to <https://www.openmp.org>.

2.3.3 Input

There are a significant number of parameters that may be varied in the code. Many of these are important variable parameters that will be the parameters of interest when modelling. However, there are also a significant number of variables that allow other properties of the model to be controlled. All parameters can, broadly, be divided into one of three categories: properties of the emitted rest-frame line or doublet, properties of the dust and gas in the ejecta and properties of the grid and code architecture. I list all the variables that are input in the primary input file in Table 2.2 and will here briefly describe the basic meaning and function of each one.

Properties of the emitted rest-frame line or doublet

lambda1_0

This is a real number that specifies the initial rest-frame monochromatic wavelength at which packets are emitted in nanometres. If a doublet is to be modelled then this represents

Table 2.2. The input variables read in from the input file and example values

Input Variable	Example Value	Input Variable	Example Value
lambda1_0	636.3	MD_tot	1.0e-4
L_tot	0.003	l	1.0
L_Halpha	0.005	q	1.3
doublet	1	b	2.0
lambda2_0	630.0	gas_shell	1
L_ratio	3.1	v_max_gas	8000
ES	1	Rrat_gas	0.05
ES_temp	10000	l_gas	1.0
LS	0	q_gas	1.5
VelShift	1	b_gas	2.0
MF	0.5	ncells	50
FF	0.1	n_packets	1e8
dayno	680	n_bins	1000
v_max	5000	n_shells	100
Rrat	0.2	dustfile	“species_file.in”

the wavelength of one of the singlets.

L_tot

`L_tot` is the total luminosity of the line in units of 10^{40} ergs s $^{-1}$. The initial energy of each packet (E_0) is therefore `L_tot` divided by the total number of packets used in the simulation (`n_packets`). For lines which have calibrated observed fluxes, this allows the flux of the line to be modelled in addition to the normalised shape. This variable is also used to estimate the undepleted luminosity of the observed line when it is initially emitted from the ejecta.

L_Halpha

Similar to `L_tot`, `L_Halpha` is the total luminosity of the H α line. In the case of H α modelling, this value should be the same at the value of `L_tot`. This variable is used in the

calculation of the electron density and it is not necessary to specify it unless the electron scattering environment is switched on (see section 2.2.6 for further details).

doublet

This is an integer of value 1 or 0 that indicates the use or otherwise of the doublet environment. If set to 1 it triggers the doublet logical in the code to be initialised to *true*. The code will then read in the values of lambda2_0 and L_ratio in order to initialise packets with two different starting monochromatic wavelengths. Packets are processed through the nebula as normal before being collated in bins weighted according to both their history and the intrinsic flux ratio of their parent singlet.

lambda2_0

This is a real number that specifies the initial rest-frame monochromatic wavelength of packets emitted from the second singlet in a doublet environment. The wavelength is specified in nanometres.

L_ratio

This real number gives the ratio between the respective luminosities of singlets in a doublet environment. The ratio should be declared as the flux at lambda1_0 divided by the flux at lambda2_0. It is expected that the doublet environment will generally be used to model forbidden lines, especially the [OI] $\lambda 6300, 6363\text{\AA}$ doublet, where the intrinsic flux ratio between the singlets may be theoretically determined.

Properties of the dust and gas in the ejecta

ES

This keyword is similar to the doublet keyword in that, by setting it equal to 1 or 0, it indicates the use or otherwise of the electron scattering environment. If it is set to 1 then it initialises the electron scattering logical in the code to *true*. If the electron scattering environment is switched on then this triggers the calculation of electron densities for every cell in the grid. This density contributes to the total optical depth of a cell and, as packets

are propagated through each cell, they will experience an electron scattering event with probability $1 - e^{\tau_e}$, where τ_e is the electron scattering optical depth.

ES_temp

When the electron scattering environment is switched on, it is necessary to calculate the electron density of each cell in the grid. In order to do this an average gas temperature must be specified to allow for $q_{H\alpha}$ to be determined. DAMOCLES will not accept any value for this input variable; only 5,000K, 10,000K and 20,000K will be accepted. These are thought to be a representative range of temperatures for the ejecta of supernovae at epochs where electron scattering still has the potential to influence observed line profiles. These specific values were selected since they are the values of $q_{H\alpha}$ that are given in Osterbrock & Ferland (2006).

MF

If this keyword (short for mass fraction) is set to 0 then a smooth density distribution of both gas and dust will be constructed. If it is not however, then this will automatically initialise the clumping logical present in the code to *true*. The value specified should be between 0 and 1 and gives the total fraction of the dust mass that should be located in clumps. The remaining fraction will be smoothly distributed according to the power-law density profiles declared in the input file.

FF

If the clumping environment is switched on (using the **MF** keyword) then **FF** declares the total filling factor of the clumps. The filling factor is defined as the fraction of the total volume of the ejecta that is occupied by clumps. For a fixed clump size, this parameter effectively determines the number of clumps to be used. Once the number of clumps to be used has been determined, the mass fraction then determines the density of the clumps.

dayno

This keyword represents the epoch being modelled. In combination with the declared maximum velocity, it is used to consistently calculate an outer radius as

$$R_{out} = 8.64 \times 10^{-6} \left(\frac{t}{\text{days}} \right) \left(\frac{v_{max}}{\text{km s}^{-1}} \right) \quad (2.37)$$

where R_{out} is in units of 10^{15}cm .

v_max

This is the maximum velocity used in the code. It is assumed to be the velocity at the outer radius of the ejecta and is used to construct a velocity profile of the form

$$v(r) = v_{max} \left(\frac{r}{R_{out}} \right)^l \quad (2.38)$$

where l is also declared in the input file and R_{out} is calculated based on the epoch and the maximum velocity.

Rrat

This number is the ratio between the inner and outer radii. Once the outer radius has been calculated as per Equation 2.37, this ratio is used to calculate the value of the inner radius.

MD_tot

This real number specifies the total dust mass to be distributed throughout the grid in solar masses (M_\odot).

l

l is the exponent of the radial velocity law in the code as per equation 2.38.

q

q describes the relationship between the radial dust density distribution and the emissivity distribution. It is the exponent of the emissivity distribution as a function of density such that $i(\rho) \propto \rho^q$ where $i(\rho)$ is the emissivity at a given density. Though this parameter may take any real value, it is frequently fixed to be $i(\rho) \propto \rho^2$, i.e. proportional to the product of the recombining proton and electron densities in the case of H α and to the product of the neutral oxygen and electron densities in the case of collisionally excited [O I] emission.

b

This parameter describes the value of the exponent of the dust density distribution in terms of radius such that $\rho \propto r^{-b}$.

gas_shell

This flag may be set to 0 or 1 to indicate that dust and gas are coupled or decoupled respectively. If it is set to 1 then the “decoupled” logical in the code is set to *true* and the gas follows a density distribution that is independent of the density distribution followed by the dust. The following five parameters specify the geometry of the emitting gas. It is worth noting that in the case where gas and dust are coupled to each other, the gas follows the same distributions as specified for dust by the parameters described above.

v_max_gas

This is the gas analogue of the v_max parameter described above.

Rrat_gas

This is the gas analogue of the Rrat parameter described above.

l_gas

This is the gas analogue of the parameter *l* described above.

q_gas

This is the gas analogue of the parameter *q* described above.

b_gas

This is the gas analogue of the parameter *b* described above.

Properties of the grid and code architecture**LS**

For an initially symmetric distribution of gas and dust, it is not necessary to specify a line of sight as all lines of sight will produce the same profile. It is therefore more

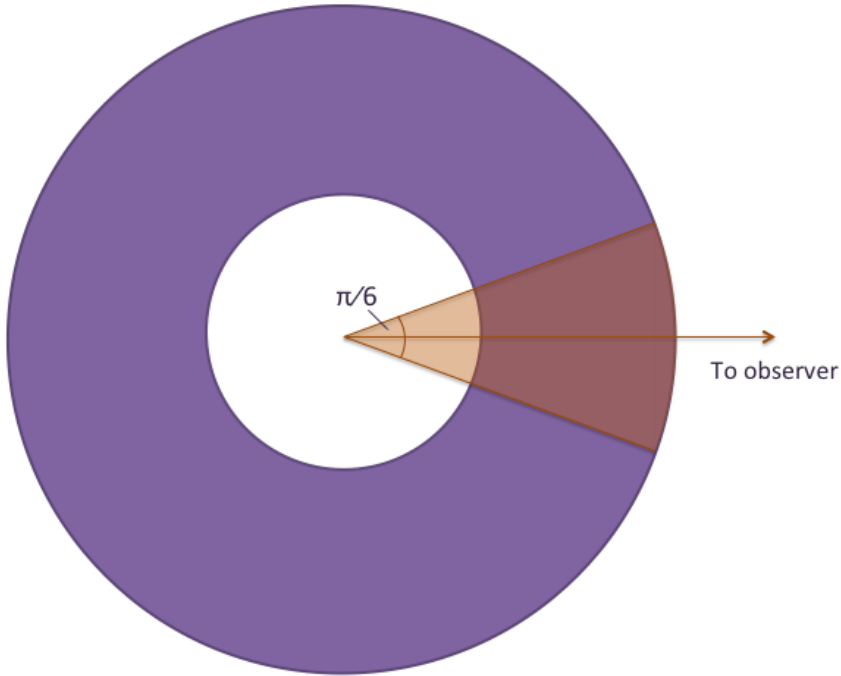


Figure 2.5. Schematic representing which packets are collected when the line of sight environment is switched on. Packets that contribute to the emergent line profile are those that escape the nebula within a cone with vertical angle $\pi/6$.

efficient to collect all packets that escape regardless of their direction of flight. However, if an alternative, axisymmetrical or asymmetrical geometry is adopted then the ability to specify a line of sight is important. If this keyword is set to 1 then the “line of sight” logical will be initialised as *true*. Only packets that escape within a cone of vertical angle $\pi/6$ will be collected. Clearly, in practice, the angle would be very much smaller but it is prohibitively expensive to run enough packets through the simulation that enough are collected to achieve a reasonable resolution when a very small vertical angle is adopted. A representation of this construction is presented in Figure ??.

VelShift

This is another environment flag. As a packet is transported through the nebula it may experience repeated scattering events that shift its original frequency beyond that expected from the maximum theoretical velocity. It is discussed in depth in the next chapter how this process of “velocity shifting” may result in a profile that exhibits an extended red wing. It is useful for the purposes of comparison and investigation to be able to turn off

this process of repeated scattering events so that the only frequency shift experienced by a packet is at emission.

ncells

Each axis is split into this number of divisions. The total number of cells in the grid is therefore ncells^3 .

n_packets

This variable determines the number of packets to be emitted and processed through the grid. This parameter is particularly important for achieving a resolution that is high enough to give representative results. The larger the number of packets used, the less noise is present in the final profile. The Monte Carlo process introduces noise that can sometimes be construed as a result when it is in fact a numerical artefact. Using a large number of packets reduces this risk and improves the output of the model. In general, the more dense an environment, the more packets it is necessary to use. This is because any packets which are absorbed are removed from the simulation and therefore reduce the desired resolution. Since the vast majority of the total processing power is used to propagate packets through the grid, an increase in the number of packets results in a significant decrease in runtime. Optically thick simulations of dust with a very high or a very low albedo have a significantly longer runtime than optically thin scenarios. When choosing this parameter, a careful balance must be found between the total runtime and the desired resolution.

n_bins

This value gives the total number of divisions in the frequency array and thus determines the overall frequency resolution of the outputted line profile. Since the resulting profile is in fact a histogram binned into a frequency array, it is important that these divisions are fine enough to provide a seemingly continuous line profile. Apparent jumps or discontinuities could be produced if too few bins are used.

n_shells

This parameter controls the total number of shells the ejecta is divided into at the start. If a particularly steep radial profile is adopted for either the velocity profile or the density

profile then the user may wish to increase the number of shells used to compensate. Increasing the number of shells will have an effect on the overall runtime, but this will be insignificant in comparison to altering the number of packets.

dustfile

Finally, this string gives the name of the input file that itemises the list of species, their relative abundances and size distributions.

2.3.4 Output, Post-Processing and Visualisation

The primary output file contains details of the emergent line profile. Three columns are written out to the file at the end of the simulation. These are the wavelength, velocity and flux of the modelled line profile. Another output file is also produced by the *model_comparison* module. This file prints both the inputted observed line profile and the outputted emergent line profile to one file in the same velocity bins. This allows for easy plotting. The columns printed in this file are wavelength, modelled flux and observed flux. The total flux is normalised to unity for both line profiles. Both of these files may be represented graphically in a straightforward fashion using any plotting package. In addition to these output files, a number of useful quantities are also calculated by DAMOCLES and output to *stdout* throughout the course of the simulation. If desired, the user may direct the *stdout* to a file for a record of these quantities. A list of all quantities output by DAMOCLES is given in Table 2.3.

Throughout my modelling, I use standard and custom routines written with MATLAB to plot line profiles, both modelled and observed. I also use MATLAB to process some of the data. For example, where I have observations with accurate observed fluxes, I scale the modelled profile to the observed profile so that fluxes remain to scale. This is initially performed by a custom MATLAB routine which smooths the modelled data to reduce any Monte Carlo noise before identifying the maximum flux value. Identifying the peak flux of the observed line profile allows the modelled profile to be automatically scaled. Any inaccuracies in the scaling may then be easily adjusted manually. I also use MATLAB for any other illustrative graphs or plots, for example, the plots in Figure 2.2 were generated in MATLAB using its 3D-scatter plotting function.

Table 2.3. List of all outputs and example values produced by the DAMOCLES code.

Output	Example Value
Total number of cells	125000
Number of grid cells inside ejecta	65544
Total volume of supernova (10^{42} cm 3)	304523264
Volume of a grid cell / Volume of a clump (10^{42} cm 3)	4668.52
Width of a grid cell (cm)	1.67e+15
Mass check (calculated as ρV)	6.03e-04
Average grain number density (cm $^{-3}$)	1.96e-09
Extinction to rest-frame wavelength	2.94e-08
Scattering extinction to rest-frame wavelength	1.63e-08
Albedo for rest-frame wavelength	0.554
Average optical depth to rest-frame wavelength	2.062
Average optical depth in V band	2.18
Average electron density (cm $^{-3}$)	71509.1
Average electron scattering optical depth	1.31e-02
Total number of packets	100000
Number of active (propagated) packets	100000
Number of inactive packets	0
Number of absorbed packets	82949
Percentage of absorbed packets out of all active packets	82.95
Number of packets in line of sight	17050
Percentage of escaped packets in line of sight	100.0
Estimated undepleted luminosity (10^{40} ergs s $^{-1}$)	1.10e-05
Total (depleted) luminosity (10^{40} ergs s $^{-1}$)	1.90e-06
Total energy absorbed (10^{40} ergs s $^{-1}$)	1.57e-06
Energy per active packet (10^{40} ergs s $^{-1}$)	1.90e-11
MSE	0.3557

2.4 Further Developments

The modular structure of DAMOCLES allows for easy implementation of additional functionality in the future. By simply adding extra modules, extra physics can be included in the code. There is potential for this code to be expanded in a number of directions. An immediately apparent development involves the dust itself. Treatment and understanding of the dust in the ejecta is crucial to understanding the shape of the line profile. The ability to place different species in different locations within the ejecta is not currently included. This would allow for stratified or asymmetrical distributions of dust species motivated by the potentially discrete locations of the parent elements. Similarly, streamlining the ability to model arbitrary density distributions and geometries would allow for more complex and accurate modelling of supernova ejecta. The ejecta of SN 1987A, for example, is known to have an asymmetric distribution which could potentially affect the contour of the line profile.

As mentioned previously, dust grains are rarely perfectly spherical and can be far more complicated in shape. It might be of interest to include a module that treats a continuous distribution of ellipsoids as mentioned in Section ?? in order to more accurately model the effects of dust grains.

More widely, supernova explosions can sometimes result in radiation that is polarised. By including the capacity to model polarised radiation in the code, we may be able to glean further information about the distribution and nature of dust forming within the ejecta. It would also be theoretically possible to expand the code to become a fully self-consistent radiative transfer code or to include certain approximations (e.g. the Sobolev approximation) to allow for full spectral modelling throughout the optical and infrared.

Aside from the development of the code directly, the current process of manual fitting can be laborious and has the potential miss potentially good fits due to the large number of variable parameters. I have completed some work over the course of this PhD wrapping DAMOCLES in a MCMC (nested sampling) routine that allows for a more thorough investigation of parameter space resulting in a full multivariate probability distribution. For reasons of time, the research presented in the following chapters was performed using manual fitting but further work finishing the implementation of this routine or a similar one would be invaluable in the future.

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Chapter 3

Probing DAMOCLES: Testing and a Parameter Sensitivity Analysis

The introduction of any new piece of software into a field has the potential to yield exciting new results. The first step in this process should therefore be a thorough investigation into the functionality of the code and an assessment of the outputs from a theoretical standpoint. Before the modelling of real data takes place, it is important to understand why the variation of a given parameter affects results in a particular way. A comprehensive understanding of parameter space not only facilitates the modelling process but may also give rise to interesting results in and of itself.

To this end, this chapter describes the ways in which DAMOCLES was tested and the results of these tests. I then also present a parameter sensitivity analysis. I describe the changes that are seen in the shapes of line profiles and consider any interesting features that arise as a result of varying the parameters of interest. I also contemplate the physical processes behind these effects.

3.1 Testing and benchmarking the code

The field of astronomy is highly reliant on the production of bespoke software to understand and interpret observations from telescopes and to develop and test new theories. As one of only a few sciences which do not have the ability to run experiments or to vali-

date results in a laboratory, progress is made via mathematical analyses or computational models based on observed data. Astrophysicists typically develop their own programs because a deep understanding of the topic to be modelled is required. Like any experiment, however, the “apparatus” should be checked and tested in order to establish its reliability.

Throughout the production of DAMOCLES, I sought, as far as was possible, to maintain best practices in scientific computing as detailed by Wilson et al. (2012). The code is carefully structured into modules and subroutines as described in the previous chapter. Each of these modules and subroutines was tested for sense and accuracy as it was written, and at each update and addition the code as a whole was tested against basic logical tests. In addition to these basic checks performed throughout the program development, it was very important to establish that DAMOCLES produced standard results as expected.

There is a general lack of published models in the literature that consider dust-affected asymmetric line profiles. This is problematic since there are no published benchmark cases against which I can compare results. I therefore consider a number of analytic line profiles derived from first principles in the case of optically thin dust (i.e. no dust was included in the models). This process ensures the functionality of the grid and the initialisation and propagation of energy packets. Additionally, I also check the absorption and scattering components of the code which are crucial to the modelling of a dusty medium. I consider some optically thick scenarios and qualitatively compare my results with those derived by Lucy et al. (1989). The profiles presented by Lucy et al. (1989) are produced both analytically and from numerical modelling and are of scenarios that are typical of the those treated by DAMOCLES and are therefore ideal for comparison. They are also the only published numerical models of dust-affected asymmetric line profiles and as such it is important that DAMOCLES is capable of reproducing the results.

3.1.1 Theoretical line profiles from first principles

The simple nature of a spherically-symmetric expanding medium with a given velocity outflow law and emissivity profile allows for analytical line profiles to be calculated from first principles in the dust-free case. Based on the methods of Gerasimovic (1933), I derive a set of three equations that describe the contours of theoretical line profiles under different starting conditions as follows.

Describing the fractional expansion velocity of the shell as $v(r) \propto r^\alpha$ with $\alpha \neq 0$ such that $v(r) = \frac{V(r)}{V_{max}}$ where $V(r)$ and V_{max} represent physical velocities and $v_{max} = 1$, velocity

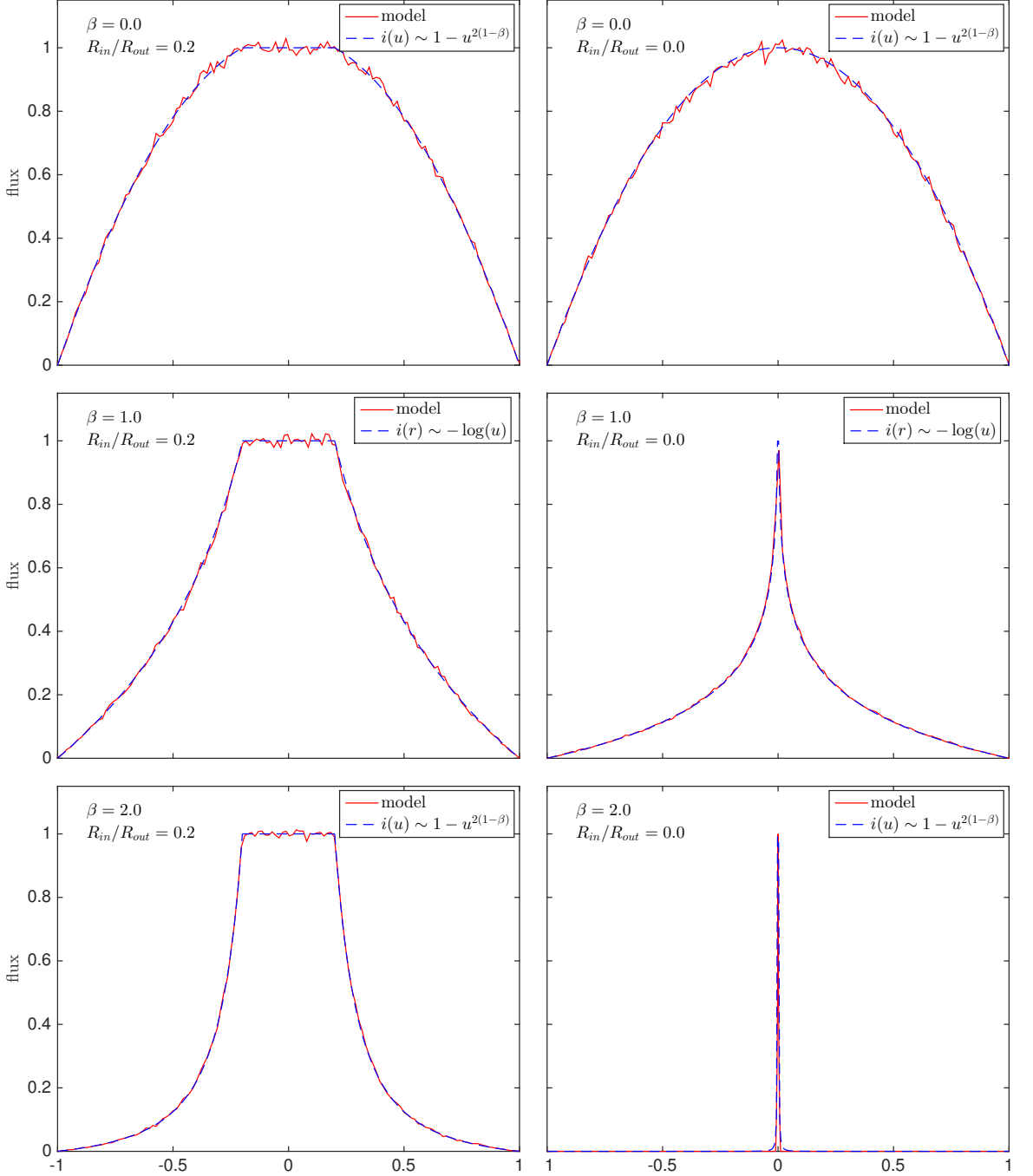


Figure 3.1. Red: Benchmark models for optically thin ($\tau = 0$) line profiles with fractional velocity $v \propto r$. Left to right: initial emissivity profiles $i(r) \propto r^{-2\beta}$ with $\beta = 0.0$, $\beta = 1.0$ and $\beta = 2.0$. Cases with $R_{in}/R_{out} = 0.2$ are on the top and $R_{in}/R_{out} = 0.0$ on the bottom. The presence of a plateau in the upper plots is due to the finite inner radius (detached shell). Blue: The analytical case with $i(u) \sim 1 - u^{2(1-\beta)}$ except in the case of $\beta = 1$ where $i(u) \sim -\log u$.

along the line of sight to the observer is given by

$$v = r^\alpha \cos \theta \quad (3.1)$$

Consider curves with constant line of sight velocity $v = \text{const}$ gives

$$dr = \frac{r}{\alpha} \tan \theta d\theta \quad (3.2)$$

The angle between the tangent to a curve and the radial line ψ is given by the formula (in polar coordinates)

$$\tan \psi = r \frac{d\theta}{dr} \quad (3.3)$$

which for curves with constant line of sight velocity v gives

$$\tan \psi = \frac{\alpha}{\tan \theta} \quad (3.4)$$

Curves of constant line of sight velocity $v = \text{const}$ therefore intersect the line $\theta = 0$ orthogonally (see Figure ??). Now we may consider the path length for $v = \text{const}$ is given by

$$ds^2 = r^2 d\theta^2 + dr^2 = r^2 \left(\frac{\tan^2 \theta}{\alpha^2} + 1 \right) d\theta^2 \quad (3.5)$$

and therefore, along curves of constant v we have

$$s = v^{\frac{1}{\alpha}} \int_{\theta_0}^{\theta_1} \frac{\sqrt{\frac{\tan^2 \theta}{\alpha^2} + 1}}{\cos^{\frac{1}{\alpha}} \theta} d\theta \quad (3.6)$$

We can now construct an volume element between v and $v + dv$ by rotating a section of thickness dv around the $\theta = 0$ axis and integrating over radius r . Assuming that $i(r)$ is the emission per unit volume (dependent only on radius), then the energy emitted by the nebula between v and $v + dv$ is proportional to

$$\int i(r) r \sin(\theta) r d\theta dr \quad (3.7)$$

The integral is most easily solved along the curves $v = \text{const}$ and by transforming

variables from θ and r to s and v . We therefore compute the Jacobian from equations 3.1 and 3.6 as

$$\frac{\partial(v, s)}{\partial(r, \theta)} = \alpha v \sqrt{\frac{\tan^2 \theta}{\alpha^2} + 1} \quad (3.8)$$

I adopt inner radius $R_{in} = q$ and outer radius $R_{out} = 1$ such that $q = R_{in}/R_{out}$.

Setting $i(r) \propto r^{-2\beta}$ (for a recombination or collisionally excited line emitted from a medium with an assumed density profile for the emitter $\rho \propto r^{-\beta}$) then gives

$$\begin{aligned} i(v) dv &\sim \frac{dv}{\alpha v^{\frac{2\beta-3+\alpha}{\alpha}}} \int_{\theta_0}^{\theta_1} \cos^{\frac{2\beta-3}{\alpha}} \theta \sin \theta d\theta \\ &\sim \frac{dv}{v^{\frac{2\beta-3+\alpha}{\alpha}}} \left[\frac{\cos^{\frac{2\beta-3+\alpha}{\alpha}} \theta}{2\beta - 3 + \alpha} \right]_{\theta_0}^{\theta_1} \end{aligned} \quad (3.9)$$

for $\frac{2\beta-3}{\alpha} \neq -1$ where $i(v) dv$ is the energy emitted in a volume element and θ_0 and θ_1 are the bounds of this element. The case $\frac{2\beta-3}{\alpha} = -1$ results in a logarithmic relationship.

In the case of a “filled” nebula, i.e. one where the inner radius is vanishingly small in comparison to the outer radius, we obtain

$$i(v) dv \sim \pm \frac{dv}{(2\beta - 3 + \alpha) v^{\frac{2\beta-1+\alpha}{\alpha}}} \left(1 - v^{\frac{2\beta-3+\alpha}{\alpha}} \right) \quad (3.10)$$

If the nebula is not “filled”, that is to say, the inner radius is some fraction of the outer radius and the remnant is a detached shell, the above formula becomes valid only from $v = 1$ to some critical value $v' = q^\alpha$. For $v < v'$, we obtain

$$i(v) dv \sim \pm \frac{dv}{(2\beta - 3 + \alpha)} \left(\frac{1}{q^\alpha} - 1 \right) \quad (3.11)$$

and therefore the top of the line is flat while the sides are sloping.

Crucially, the width of the flat section is determined by $v' = q^\alpha$ or simply $v' = q$ in the case where $v \propto r$, whilst the shape of the profile outside of the flat top is described by equation 3.10.

Profiles with a variety of shapes may be derived from these formulae depending on the

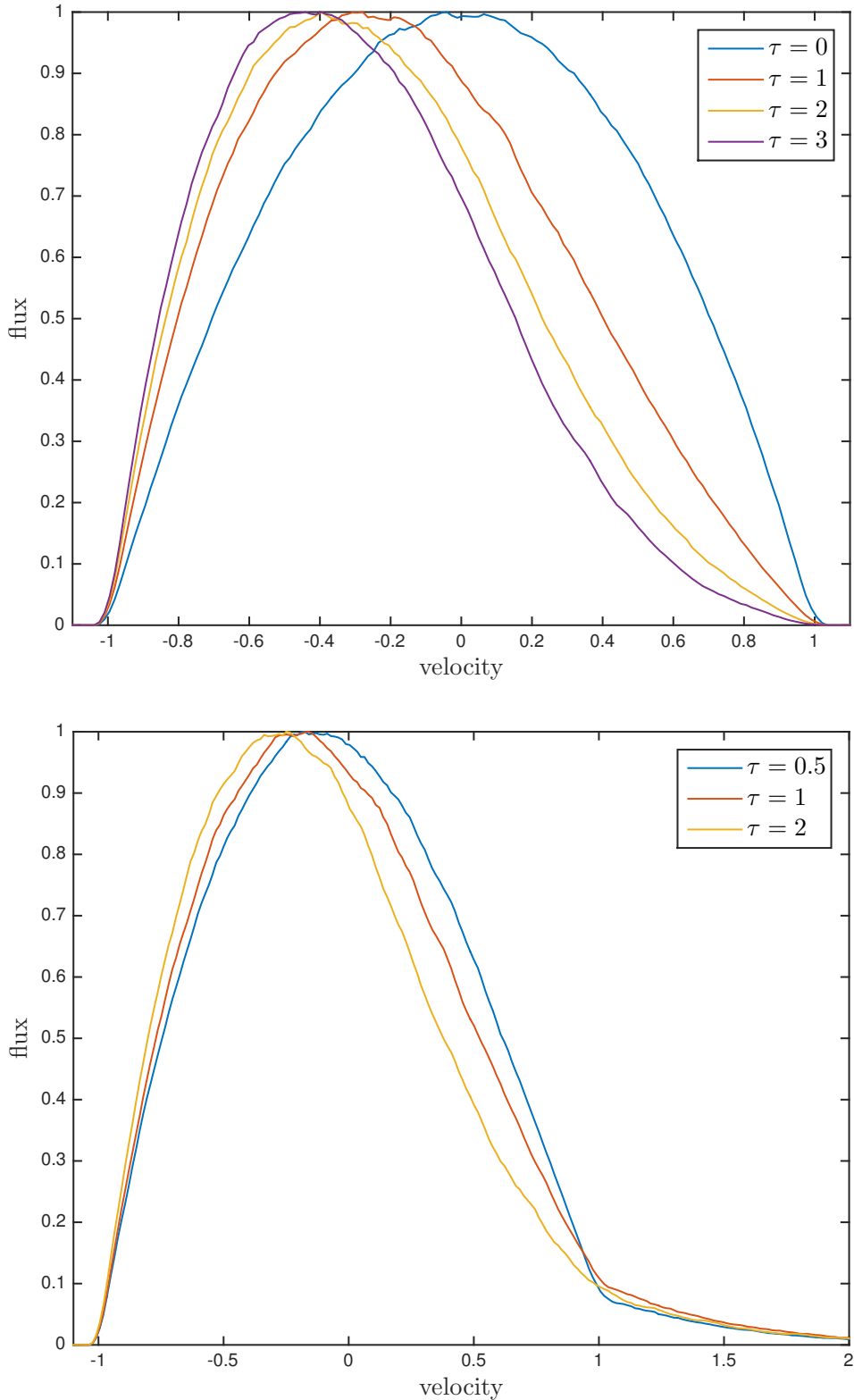


Figure 3.2. Benchmark models for line profiles with $v \propto r$, $i(r) \propto \text{constant}$ and a filled sphere with $R_{in}/R_{out} = 0$. Pure dust absorption models ($\omega = 0$) are presented in the top plot, whilst partially scattering models are presented at the bottom ($\omega = 0.6$) as per Lucy et al. (1989) Models II and III. All resulting profiles have been scaled to unity flux at their peaks.

relative values of α and β . Here we consider three main families of curves:

1. $i(v) \sim v^{-\gamma} - 1 \quad (\alpha > 0, 2\beta - 3 + \alpha > 0)$
2. $i(v) \sim 1 - v^\gamma \quad (\alpha > 0, 2\beta - 3 + \alpha < 0)$
3. $i(v) \sim -\log v \quad (\alpha > 0, 2\beta - 3 + \alpha = 0)$

where γ is defined as $\gamma = |\frac{2\beta-3+\alpha}{\alpha}|$.

Models are presented for each of these cases, both for a filled nebula and for a shell structure with $R_{in}/R_{out} = 0.2$. A velocity profile $v \propto r$ appropriate for supernova ejecta in the free expansion phase is used throughout (Li & McCray 1992; Xu et al. 1992; ?). Values of $\beta = 0, 1$ and 2 are adopted. Figure 3.1 illustrates the excellent agreement between the analytical case and the models. All fluxes are scaled to unity at the peak.

3.1.2 Comparison of DAMOCLES models with previously published results

In addition to the tests for optically thin lines described above, we also compared our outputs to those derived by Lucy et al. (1989) in order to assess the accuracy of the scattering and absorption aspects of the code. We consider two similar cases, equivalent to Models II and III of Lucy et al. (1989). In the first case, dust with zero albedo (pure absorption) is uniformly distributed throughout a filled nebula with a velocity profile $v \propto r$. In the second case, the same scenario is considered but in a medium of dust with albedo $\omega = 0.6$.

In the first case, the profile may once again be derived analytically from the basic geometry using the fact that radiation will be attenuated by a factor $e^{-2\tau_\nu v}$ between points with line of sight fractional velocities $-v$ and $+v$ where τ_ν is the optical depth at frequency ν from the centre to the outer edge of the ejecta. The line profile is therefore given by

$$\frac{I(v)}{I(-v)} = \exp(-2\tau_\nu v) \tag{3.12}$$

Lucy et al. (1989) presented several examples for both the analytical case of the perfect absorber and a Monte Carlo model for grains with $\omega = 0.6$. We present the same cases

in Figure 3.2 and note that the resulting profiles exhibit the same features and shape. Of particular interest is the scattering wing that appears beyond the maximum velocity ($v_{max} = 1$) on the red side of profiles in the partial scatterer case as a result of the packets doing work on the expanding sphere. This was noted by Lucy et al. (1989) as a potential diagnostic for the presence of dust in the ejecta of a supernova and we will discuss this further in Section ??.

3.1.3 Testing the electron scattering mechanism

3.1.4 Clumped models in smooth limits

3.2 An analysis of the sensitivity of line profiles to the variable parameters

It is of general interest to establish potential diagnostic signatures in the line profiles of supernovae and their remnants in order to trace dust formation more effectively. The capacity to specify a number of parameters is included in Damocles. The variation of each parameter potentially affects the contour of the resulting line profile in a different way. By investigating each parameter separately over a range of values, it may be possible to identify certain characteristics of dust-affected line profiles that may be associated with a particular property of the dusty medium. This insight could help to explain unusual or interesting features of observed line profiles where dust is suspected to be an influential factor. In this chapter, I investigate and discuss the effects of the main parameters of interest, namely:

- the maximum velocity, V_{max}
- the ejecta radius ratio, R_{in}/R_{out}
- the dust optical depth, τ
- the dust albedo, ω
- the dust density profile exponent, β , where $\rho \propto r^{-\beta}$

I also investigate the capacity of this type of model to infer properties of the dust itself, specifically the size of dust grains and any size distribution that may be discernible, and the composition of the dusty medium as made up of a number of species.

3.2.1 The maximum line velocity, V_{max}

The maximum velocity is defined as the velocity at the outer edge of the line emitting region for a given line. The maximum velocity may vary between different spectral lines or doublets due to different locations of species with differing ionization thresholds. Clearly, the larger the maximum velocity used the wider the profile becomes. To some extent therefore the steepness of the density profile and the maximum velocity can act to counter each other since a steeper density profile narrows the profile (see Section 3.2.5). The shape of the wings of the profiles, however, generally preclude much degeneracy in this aspect - the overall shape of the line profile can be used to determine the exponent of the density profile to within a relatively small range.

More important is the effect that the maximum velocity has on the overall optical depth. Since the overall volume of the ejecta is determined solely by the maximum velocity and the ratio of the inner and outer radii, the total optical depth to which the radiation is exposed can be greatly affected by even a relatively small change in the maximum velocity. Practically speaking, the maximum velocity can usually be fairly well determined from the observations (identified as the point where the flux vanishes on the blue side) and may be further constrained through modelling.

3.2.2 The ejecta radius ratio, R_{in}/R_{out}

As already discussed in Section 3.1.1, the width of the flat top is determined solely by the ratio of the inner and outer radii, the exponent of the velocity profile and the maximum velocity. Throughout my sensitivity analysis of parameter space, I assume that the velocity profile takes the form $v \propto r$ despite it being a variable parameter in the code. This is the case for all supernovae from just a few months after the initial explosion (and therefore likely well before the epoch of the onset of dust formation) since the ejecta is in free expansion. This is discussed in further detail in Section 3.2.10. For this case, R_{in}/R_{out} is given by

$$\frac{R_{in}}{R_{out}} = \frac{V_{min}}{V_{max}} \tag{3.13}$$

where it is often possible to constrain V_{min} and V_{max} to a relatively narrow range simply from the observed line profile by identifying the velocity at which the flux goes to zero on

the blue side (V_{max}) and the velocity at which the profile starts to flatten off, also on the blue side (V_{min}).

The majority of spectral lines emitted from supernovae and supernova remnants are expected to have a flat top before dust attenuation effects since it is rare for these objects to form a completely filled nebula. However, even a very small amount of dust attenuation may result in the line profile appearing to be smoothed at its peak.

The effect of varying the ratio of the inner and outer radii is derived analytically in Chapter 2 and presented for a number of emissivity profiles in Figure 3.1. All profiles have been scaled to unity flux at their peaks.

3.2.3 The dust optical depth, τ

The effects of dust optical depth on the line profile produced by Damocles are in essence the same in the case of both a detached shell and a complete nebula. However, due to the increased complexity of the original profile in the case of the detached shell, a number of slightly more interesting effects are observed in these models and I discuss these first. I then go on to discuss the case of a complete nebula where the effects observed are very similar to those seen in models with high dust optical depth in the detached shell case.

The detached shell case

As expected, in cases of non-zero dust optical depth, greater attenuation of the original line profile on the red side is exhibited. This is illustrated in two panels of profiles in Figures 3.3 and 3.4. The profiles are most revealing at lower dust optical depths. The effects of the asymmetric absorption can be seen in different sections of the profiles. The region of the profile that is most clearly affected by dust absorption is the flat-topped region. A small amount of absorption in this region results in a skewed profile, with a fraction of the flat-topped section removed. The peak becomes blue-shifted as a result, but only to the original value of $-V_{min}$, the minimum velocity corresponding to R_{in} . In addition to the attenuation in this region, the red wing of the profile is also somewhat reduced, and the blue wing somewhat increased relative to their original symmetric positions. The result is a relatively “jagged” looking profile, often with sharp changes at $\pm V_{min}$. The profile is generally asymmetric, although the degree of absorption in the flat-topped region may sometimes make it seem as though the profile is in fact symmetric and uniformly blue-shifted (see Section 3.2.7 for further discussion).

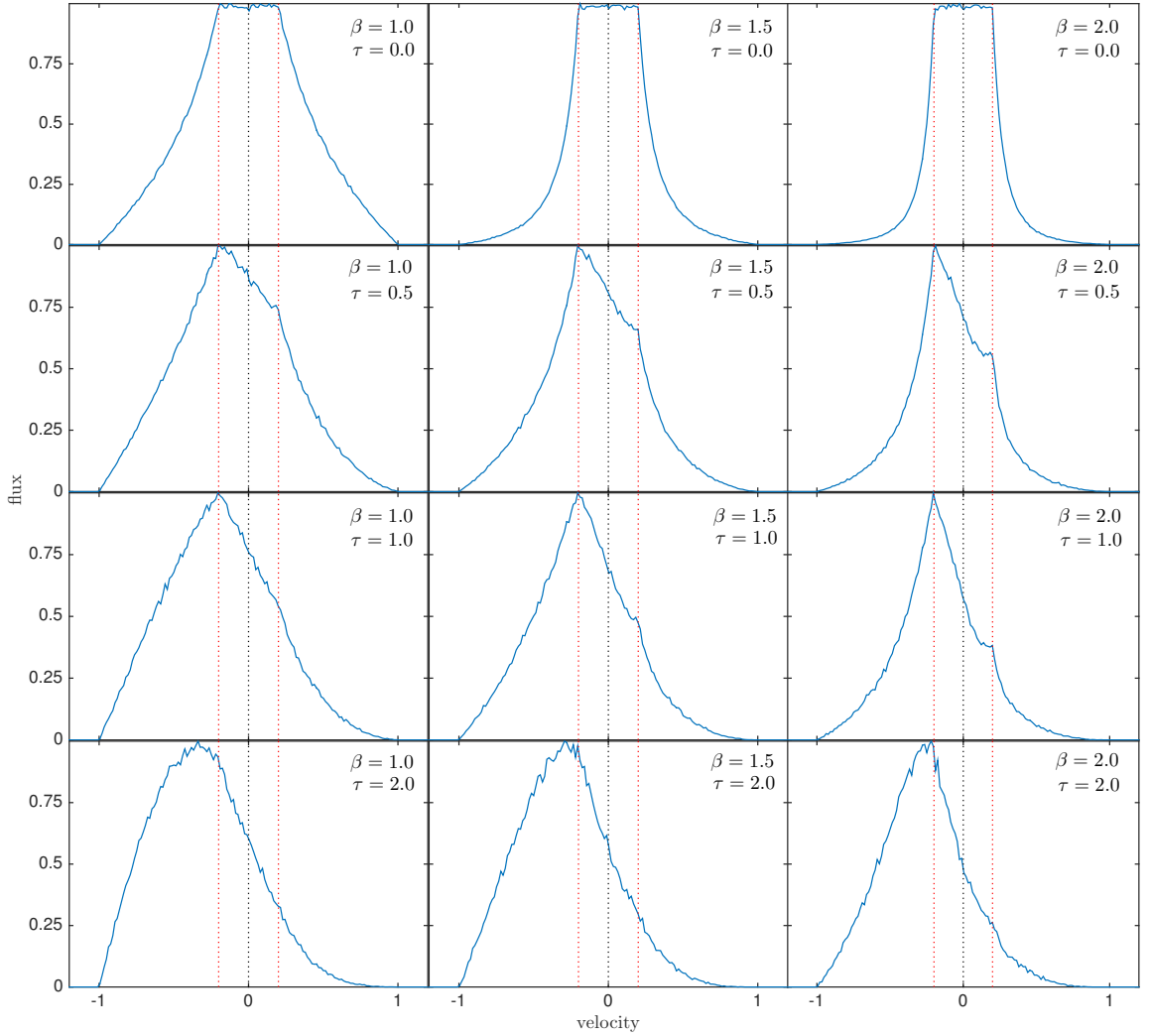


Figure 3.3. Set of models with $i(r) \propto r^{-2\beta}$ for $\beta = 1.0$ (left), $\beta = 1.5$ (middle) or $\beta = 2.0$ (right), $\omega = 0$, $R_{in}/R_{out} = 0.2$, $v(r) \propto r$ and $v_{max} = 1$ illustrating the effects of varying τ . Peak fluxes are scaled to unity.

At high dust optical depths the entire profile is shifted to the blue and the peak moves beyond $-V_{min}$ further into the blue. The profiles also become more smooth. A set of models showing the effects of varying optical depths for different density profiles and dust albedos are presented in Figures 3.3 and 3.4.

The complete nebula case

To reproduce similar characteristic dust-affected line profiles where the peak of the profile is shifted beyond $-V_{min}$ into the blue is “easier” for smaller values of V_{min} . The complete nebula is therefore effectively analogous to cases of higher optical depths for a detached shell and the same effects that are illustrated in Figures 3.3 and 3.4 apply.

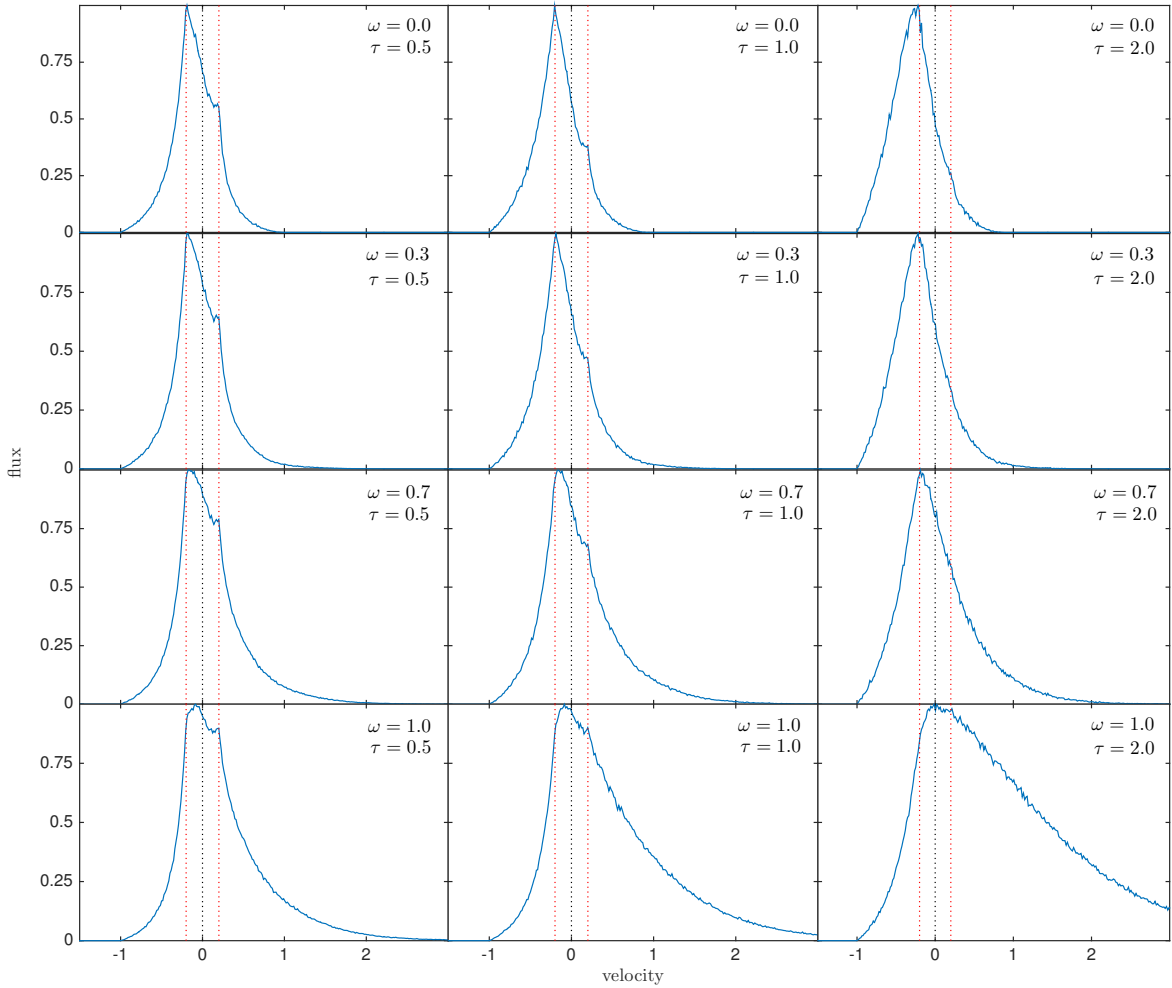


Figure 3.4. Set of models with $i(r) \propto r^{-4}$ (i.e. $\beta = 2.0$), $R_{in}/R_{out} = 0.2$, $v(r) \propto r$ and $v_{max} = 1$ illustrating the effects of varying τ and ω . Peak fluxes are scaled to unity.

3.2.4 The dust albedo, ω

In the past, there has largely been a focus on the effects of absorption by dust on the shapes of line profiles and less attention has been paid to the potential effects of scattering by dust. In fact, line profiles can be significantly affected by scattering of radiation. Not only does repeated scattering of photons increase the number of potential opportunities for a given photon to be absorbed but it also results in continuous shifting of the frequency of the photon to the red. The photon must do work on the expanding shell of dust in order to escape and thus many of the photons are reprocessed beyond the theoretical maximum velocity on the red side of the profile. The result can be a substantial, extended wing on the red side of the line. In the case of strong dust scattering, this can result in an

asymmetric profile that is the opposite of that normally expected with the majority of the emission on the *red* side. The peak however, remains blue-shifted (see the bottom right panel on Figure 3.4 as an example). For the line profile to exhibit this feature requires the dust to be a nearly perfect scatterer and it is therefore unlikely that profiles of this sort will be frequently observed. See Figure 3.4 for a fuller illustration of the variation with ω and τ .

The implications of this result in relation to the use of line profiles as a diagnostic for tracing dust formation in supernova ejecta are discussed further in Section 3.2.7.

3.2.5 The dust density profile, $\rho \propto r^{-2\beta}$

Whilst the density profile of the dust may have some effect on the resulting profiles, it is the initial emissivity profile (dependent on the dust density profile) that has greatest effect on the resulting shape of the line profile.

In general, the steeper the emissivity distribution, the narrower the line profile becomes. The sides of the line profile may become almost straight for a very steep distribution since the majority of the emission then comes from a very narrow velocity range. For a flat-topped profile of fixed width this approximates the square profile produced in the case of an emitting shell with constant velocity.

The dependence of the shape of the line profile in the optically thin dust case is described in Section 3.1.1. However, the density profile also plays a significant role where there is even a small amount of absorption. As previously discussed, at relatively small optical depths, a section of the flat-topped region is removed resulting in a peak at $-V_{min}$. The shape of the profile in this region is significantly affected by the density profile. Shallow density profiles (low β) produce a virtually linear variation in flux between $-V_{min}$ and $+V_{min}$. For a fixed dust optical depth, the steeper the distribution becomes, the more concave the profile becomes between $-V_{min}$ and $+V_{min}$, ultimately resulting in a clear shoulder to the profile at $+V_{min}$. For extremely steep density distributions this can result in a double peaked profile with trough to the red of $V = 0$. A illustration of the effects of variation of β with τ on the profiles is shown in Figure 3.3.

3.2.6 Inferring properties of the dust from the models

The presence of an extended red wing at large positive velocities in combination with increased extinction on the red side at smaller positive velocities can allow the values of τ

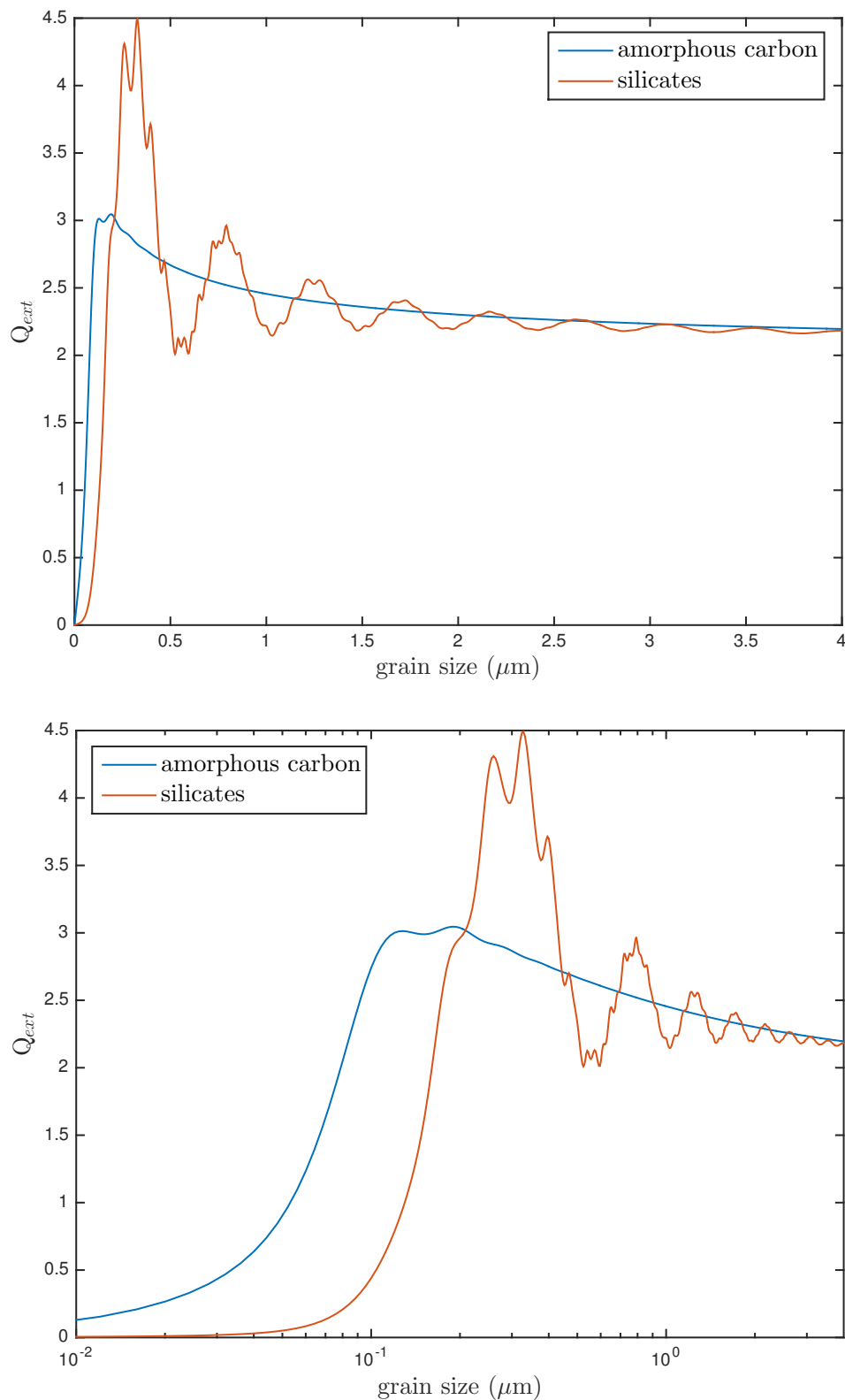


Figure 3.5. Variation of extinction efficiency (Q_{ext}) with grain size for amorphous carbon and silicates using Mie theory at $\lambda = 658\mu\text{m}$. Optical constants are from Zubko et al. (1996) and Draine & Lee (1984). A linear scale is presented on the top and a log scale on the bottom.

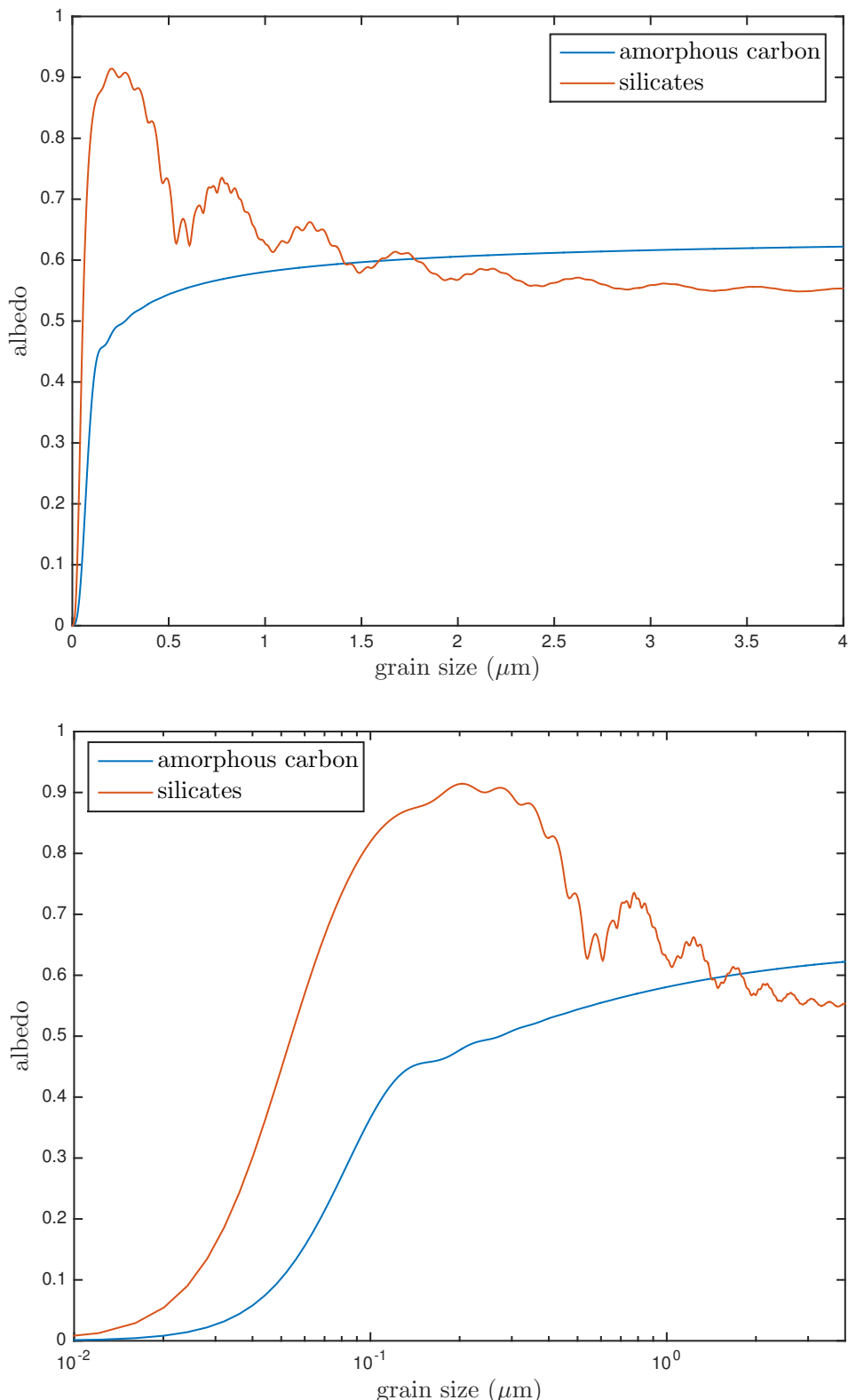


Figure 3.6. Variation of albedo with grain size for amorphous carbon and silicates using Mie theory at $\lambda = 658 \mu\text{m}$. Optical constants are from Zubko et al. (1996) and Draine & Lee (1984). A linear scale is presented on the top and a log scale on the bottom.

and ω to be well constrained. In this case it is possible to translate these values into a dust mass and average grain size respectively for a given species or combination of species using optical properties and Mie theory (see Figures 3.5 and 3.6). In fact, it is the dust mass and average grain size that is varied within the code for a specified species or combination of species i.e. the dust optical depth and grain size are not varied directly. It is therefore important to note that the use of different optical properties may substantially alter the inferred optical depths and albedos for a given species of specific grain size as has been noted previously (e.g. Owen & Barlow (2015)).

For amorphous carbon, the larger the grain size used the larger the albedo and the smaller the cross-section of absorption. Larger masses of dust are therefore required to fit the same degree of absorption if a larger grain size is used. This is in contrast to SED radiative transfer modelling where larger grain sizes generally result in less dust being required to fit the IR portion of the SED (W15). These two techniques in tandem may therefore give excellent limits on grain sizes for different species or combinations thereof.

3.2.7 Observable signatures of dust in line profiles

The greater the dust optical depth, the more attenuation of the line is observed. As expected, the red side of the profile suffers a greater degree of absorption than the blue side. The resulting asymmetry is somewhat more complex than perhaps previously thought however. Dust has repeatedly been cited as the agent responsible for the apparent blue-shifting of line profiles in supernovae in the manner of the profiles presented in Figure 3.2. That is, relatively high optical depths result in an overall shift of the entire profile towards the blue.

In practice a relatively large optical depth ($\tau \approx 2$) is required to actively shift the peak of the profile bluewards of its natural V_{min} value corresponding to the velocity at the inner radius of the shell. In most cases it seems more likely that the dust may not be optically thick and the blue-shifting of the peak of the profile is likely a result of attenuation in the flat-topped section (close to R_{in}). The peak would therefore tend to be located at $-V_{min}$.

Since dust absorption is wavelength dependent for $2\pi a < \lambda$, one might expect the position of the peak flux to be dependent on the wavelength of the line being considered. The relationship between the locations of the peaks of profiles and their wavelength has been discussed by several authors in relation to dust formation (Smith et al. 2012; Fransson et al. 2013; Gall et al. 2014). Whilst this may occur in cases of high dust optical depth,

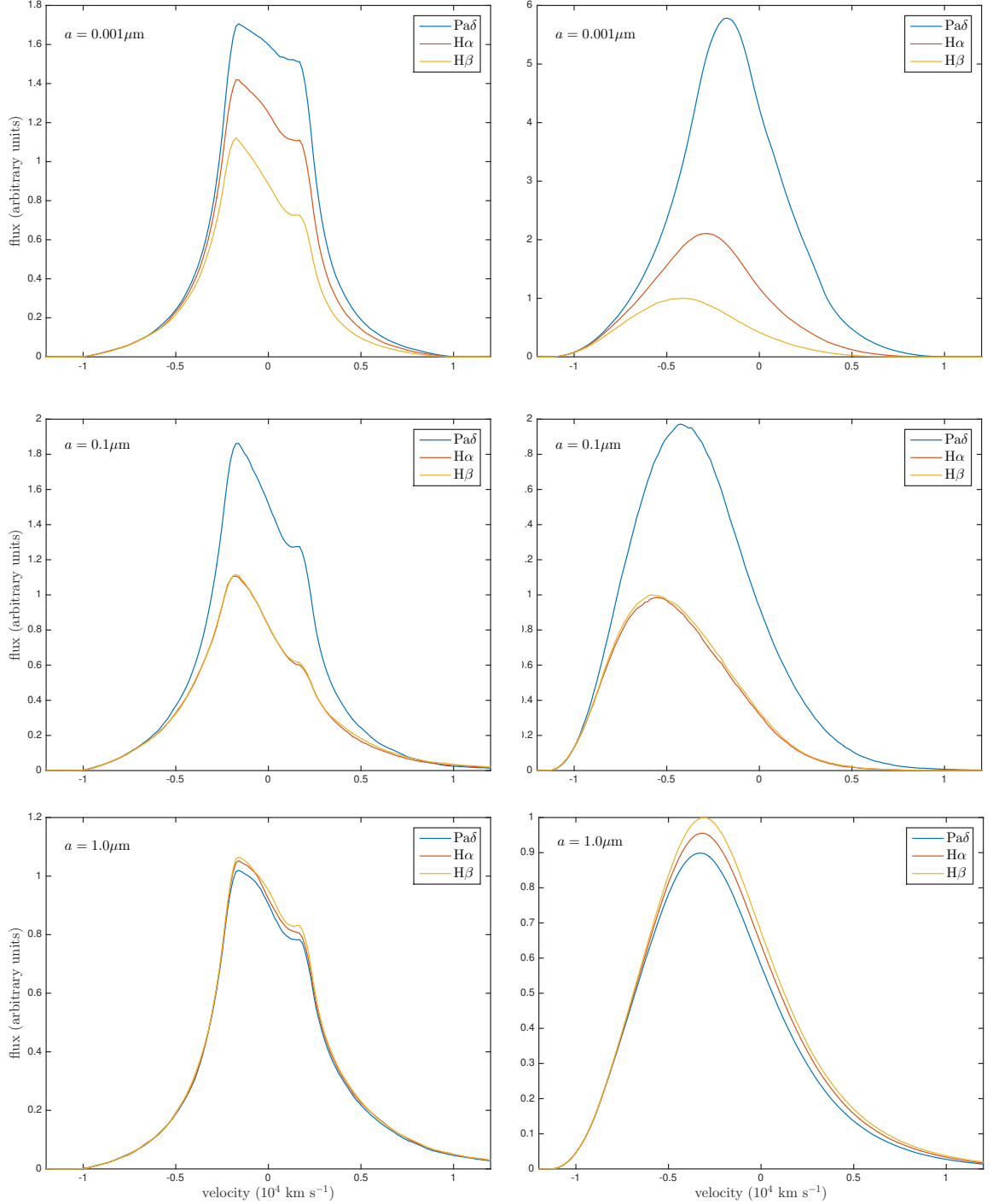


Figure 3.7. Model line profiles for H α (6563Å in red), H β (4861Å in yellow) and Pa δ (10049Å in blue) for optically thin and optically thick cases on the left-hand side and right-hand side respectively. All models adopted density profile $\rho(r) \propto r^{-4}$ (i.e. $\beta = 2$), velocity profiles $v(r) \propto r$ and radii ratio $R_{in}/R_{out} = 0.2$. The grain radii used were $a = 0.001 \mu\text{m}$ (top), $a = 0.1 \mu\text{m}$ (middle) and $a = 1.0 \mu\text{m}$ (bottom). All the above models used amorphous carbon.

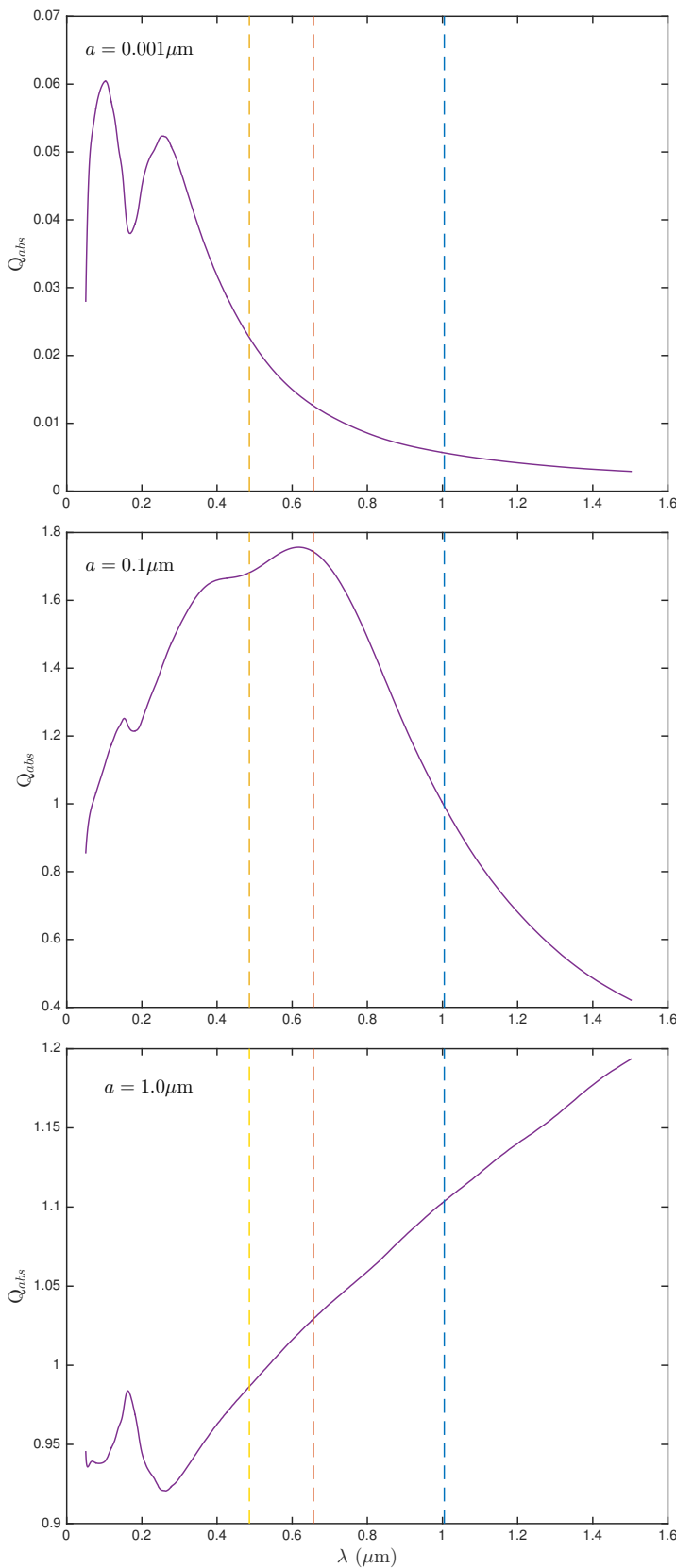


Figure 3.8. The variation of amorphous carbon dust absorption efficiency with grain size. The grain radii plotted are $a = 0.001\mu\text{m}$ (top), $a = 0.1\mu\text{m}$ (middle) and $a = 1.0\mu\text{m}$ (bottom). The vertical lines mark the wavelengths of H α (6563Å in red), H β (4861Å in yellow) and Pa δ (10049Å in blue).

this is not necessarily likely to be seen in the ejecta of most supernovae. The wavelength-dependence of dust absorption instead can result in differing degrees of extinction in the flat-topped region of each profile but still leave the peak at its blue-shifted position of $-V_{min}$. Of course, the value of V_{min} may be different for different species. However, if this is the case then there would be no reason to expect a variation in the position of the peak of profiles to be correlated with the wavelength dependence of dust. Rather one would expect it potentially to trace the location of different ions within the ejecta. However, for lines from the same ion, for example the Balmer and Paschen lines of HI, I might expect to see peaks at the same position but differing degrees of absorption. At high resolutions, it might be possible to detect differences in the shape of the line profiles, particularly between $-V_{min}$ and $+V_{min}$ where the steepness of the incline traces the degree of dust absorption. This can be seen in Figure 3.7 where I illustrate the effects of the wavelength dependence of dust absorption for three lines, H α (6563Å), H β (4861Å) and Pa δ (10049Å). All lines were modelled using three different grain sizes and in both optically thin and thick cases. I also show the variation of the absorption cross-section with wavelength at three different grain sizes in Figure 3.8.

The attenuation of the flat-topped region is often such that it can be hard to discern a difference in slope between the attenuated section between $-V_{min}$ and $+V_{min}$ and the slope of the wing for $V > +V_{min}$, particularly in circumstances where data is of poor resolution or has a poor signal-to-noise ratio. Even in the case of excellent data, it may be easy to overlook these particular features or to dismiss them as natural fluctuations in the geometry of the ejecta rather than that they may be a product of dust absorption effects.

The greater attenuation of radiation received from the receding portion of the ejecta results in an asymmetry of the line profile whereby the majority of the observed emission is located bluewards of the peak. However, the effects of repeated dust scattering events within the ejecta can serve to counter this asymmetry. Even in the case of dust grains with a relatively low albedo, a surprisingly persistent wing on the red side of the profile is seen beyond the maximum theoretical velocity of the emitting region. For higher albedos this can actively result in a shift in the overall asymmetry of the profile, with the majority of the emission being emitted redwards of the peak, though the peak itself remains blue-shifted.

This effect is obviously analogous to that of electron scattering which also produces a significant red wing in line profiles (Hillier 1991; Auer & van Blerkom 1972). This is an important consideration in both modelling and analysis of spectral line profiles.

DAMOCLES has the capacity to include a basic electron scattering mechanism in order to assess the possibility that any observed red wing might be produced by electron scattering rather than dust scattering. The red wing observed in line profiles is an excellent diagnostic for determining the overall dust albedo and it is therefore important to establish whether this feature is due to electron or dust scattering or a combination of the two.

The combination of relatively low optical depths, initially flat-topped profiles, greater attenuation on the blue side with increased flux on the red side due to scattering can result in a profile that ends up appearing almost symmetrical, particularly if contaminants, such as narrow lines or blending with other broad lines, are present or if the resolution of the data is low. The potential for apparently symmetrical profiles that appear to have been uniformly blue-shifted should be noted (see Figures 3.3 and 3.4 for examples of this).

3.2.8 The effect of a grain size distribution

It is important to consider the potential effect on the dust mass of modelling a grain size distribution instead of a single grain size. For a grain size distribution the overall extinction cross section, C_{ext} , at a given wavelength is calculated as

$$C_{ext} = \int_{a_{min}}^{a_{max}} Q_{ext}(a)n(a)\pi a^2 da \quad (3.14)$$

where $Q_{ext}(a)$ is the extinction efficiency for a grain size a and $n(a)$ is the number of grains with size a . The overall extinction efficiency is then

$$Q_{ext} = \frac{C_{ext}}{\int_{a_{min}}^{a_{max}} n(a)\pi a^2 da} \quad (3.15)$$

The scattering cross-section Q_{sca} is similarly calculated. As a result of these calculations, there is rarely a single grain size that has the same albedo and extinction efficiency as a size distribution. Modelling a size distribution instead of a single grain size may therefore alter the deduced dust mass. Since models are only sensitive to the optical depth and the albedo, however, it is not possible to deduce the grain size range or distribution and only single grain sizes are investigated in the models that are presented in the following chapters.

Whilst this apparently limits the scope of these results, it is important to consider the extent to which considering grain size distributions would alter the derived dust masses. By considering a number of grain size ranges and adopting a power law distribution with a variable exponent, I may gain some insight into the effects of adopting a distribution rather than a single size. For the classical MRN power law ($n(a) \propto a^{-3.5}$) with a wide grain size range ($a_{min} = 0.001\mu\text{m}$ to $a_{max} = 4.0\mu\text{m}$) the derived albedo of $\omega = 0.001$ is much too small to reproduce the required wing seen at early epochs. I investigate this issue by adjusting the exponent of a distribution for a number of grain size ranges until the overall albedo is the same as that seen for the best fitting single grain size. I can then approximately calculate the required dust mass for a distribution of grain sizes from the properties of a single-size model by equating the optical depths. The optical depth for a single grain size is proportional to

$$\tau \propto Q_{ext}(a)\sigma(a)n_d \quad (3.16)$$

where n_d is the number density of dust grains, $\sigma(a)$ is the cross-sectional area of a grain of radius a and $Q_{ext}(a)$ is the extinction efficiency for a grain of radius a . On average, this gives

$$\tau \propto \frac{Q_{ext}(a)M\pi a^2}{\frac{4}{3}\pi a^3\rho V} \quad (3.17)$$

for a total dust mass M , total volume of the ejecta V and density of a dust grain ρ .

This simplifies to

$$\tau \propto \frac{Q_{ext}(a)M}{\frac{4}{3}a\rho V} \quad (3.18)$$

$$M_d = \frac{M_s Q_{ext,s}(a_s)}{a_s} \times \frac{\int_{a_{min}}^{a_{max}} n(a)a^3 da}{\int_{a_{min}}^{a_{max}} Q_{ext}(a)n(a)a^2 da} \quad (3.19)$$

where the subscript s represents the single grain size quantities and the subscript d

represents quantities for the grain size distribution. This is only calculable for a specific wavelength and is therefore only an approximate conversion when performed at the rest-frame wavelength of the line in question. However, practically, the variation of extinction efficiency and albedo over the narrow wavelength ranges modelled within the code is not significant and so this method produces relatively accurate dust masses (as determined by running models with the new parameters).

3.2.9 The effect of different species

In the majority of the modelling that follows, a single species, amorphous carbon, is considered. A single species is used since the parameters that affect the quantity of dust required in the model are the albedo and the optical depth. There are therefore likely many possible combinations of species that may result in a good fit to the data. The choice of amorphous carbon is partly motivated by evidence that, for SN 1987A (which, as an incredibly well-observed and relatively typical core-collapse supernova, is an excellent test case) the fraction of silicates present in the dusty ejecta is limited to approximately 15% (W15, Ercolano et al. (2007)). It is also motivated by previously published SED models which generally employ amorphous carbon. This is because SED models frequently require far larger masses of silicate dust than amorphous carbon dust in order to produce similar levels of infrared flux and therefore amorphous carbon models are likely to produce the more conservative dust mass estimates. By modelling with amorphous carbon I may compare directly to these models where possible.

I consider the change in dust mass when a medium of 100% silicates is used instead of amorphous carbon. I use optical constants presented in Draine & Lee (1984). In a similar manner to the approach detailed in Section 4.2.5, I may calculate the mass of silicates that is equivalent to a carbon mass for a single grain size. I consider the albedo at the original grain size, calculate the equivalent grain size for silicates that results in the same albedo and then calculate the new dust mass by considering the change in the extinction efficiency as

$$M_{sil} = M_{amc} \left(\frac{Q_{amc}}{Q_{sil}} \right) \left(\frac{a_{sil}}{a_{amc}} \right) \left(\frac{\rho_{sil}}{\rho_{amc}} \right) \quad (3.20)$$

Because of the nature of the variation of albedo with grain size for silicates (see Figure

Table 3.1. Equivalent dust masses for the day 714 clumped models using grain size distributions and 100% amorphous carbon. f is factor of increase from the dust mass for the single size model ($M = 7 \times 10^{-5} M_{\odot}$ with $a = 0.6\mu\text{m}$) and p is the exponent of the grain size distribution $n(a) \propto a^{-p}$.

<i>carbon</i>		<i>silicates</i>		$f = M_{sil}/M_{amc}$
a	Q_{ext}	a	Q_{ext}	
0.6	2.60633	0.0583	0.0772	5.37
0.6	2.60633	4	2.1828	13
3.5	2.2129	0.0641	0.10182	0.65
3.5	2.2129	1.02	2.149	0.49
3.5	2.2129	1.376	2.3514	0.61

3.6), there is often more than one silicate grain size that will give rise to the same albedo. I consider all the possibilities and the resulting mass conversion factors in Table 4.8. In my best fitting models for days 714 and 806, using any fraction of silicates of either grain size would serve to increase the dust mass. However, at later epochs, using some fraction of silicate dust would reduce the dust mass to potentially more than half of my estimated values. However, this is still within my predicted range and my minimum and maximum dust masses remain robust.

3.2.10 The velocity distribution

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Chapter 4

The Evolution of Dust Formation in SN 1987A

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AUTHOR

4.1 Spectral Observations of SN 1987A

SN 1987A has been the most intensively observed supernova in history, with a wealth of both spectral and photometric data available to model. From the archives of a number of different telescopes we have collated optical spectra acquired over a wide range of epochs. At the earlier epochs we use spectra obtained by the Anglo-Australian Telescope (AAT) and the Cerro Tololo Inter-American Observatory (CTIO) and at later epochs we use spectra from the archives of the Hubble Space Telescope (HST) and the Very Large Telescope (VLT). An explosion date of 23 February 1987 is adopted throughout and epochs are measured relative to this date. Full details of all observations may be found in Table 4.1. The spectral resolutions of the grating spectrograph observations are listed in column 7, while column 8 lists the spectral resolving powers of the echelle spectrograph observations.

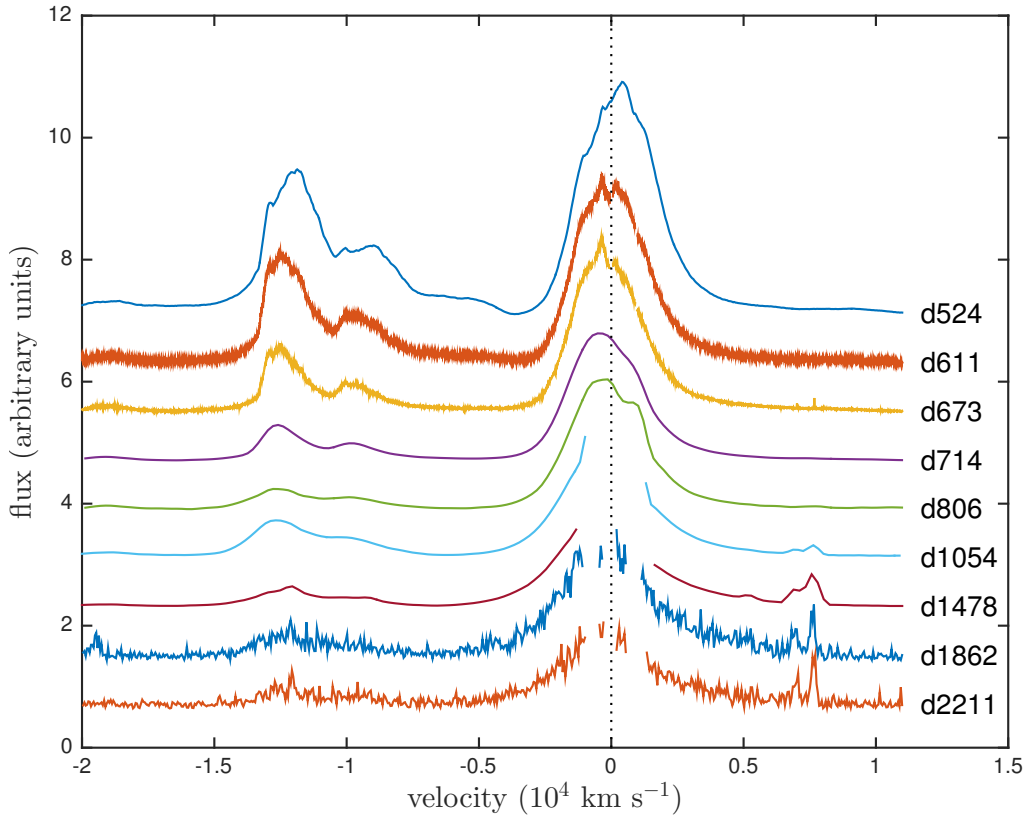


Figure 4.1. Archival data showing the evolution of the H α and [O I] line profiles from SN 1987A at the earlier of the epochs considered. The spectral gaps at the last two epochs correspond to where narrow line emission from the equatorial ring has been removed. The spectra have been continuum-subtracted and offsets have been applied for display purposes.

Wavelength ranges encompassing the H α line and [O I] $\lambda\lambda 6300, 6363$ Å doublet were selected in order to trace their evolution from day 524, near the time of the first indications of dust formation (Wooden et al. 1993), to day 8020, near the current era. Optical spectroscopy obtained at the AAT using the Faint Object Red Spectrograph (FORS) during the first two years after outburst was kindly supplied by Dr Raylee Stathakis (Spyromilio et al. 1991, 1993; Hanuschik et al. 1993) and optical spectra from the CTIO were donated by Dr Mark Phillips (Suntzeff et al. 1991).

The evolution of the H α and [O I] line profiles is presented in Figures 4.1 and 4.2. At later epochs, the broad H α profile emitted by the ejecta becomes contaminated by narrow line emission from the equatorial ring. These lines have been removed for the purposes of modelling the broad line. A continuum fit has been subtracted from each spectrum and a velocity correction has been applied for a recession velocity of 287 km s $^{-1}$ (Grönningsson

et al. 2008).

Table 4.1. Details of the archival data for SN 1987A.

Date	Age (days)	Telescope	Inst	λ_{min} (Å)	λ_{max} (Å)	Res. Power	Reference
31 Jul 1988	524	AAT	FORS	5500	10190	20	Spyromilio et al. (1991)
26 Oct 1988	611	AAT	UCLES	6011	7336	30000	Hanuschik et al. (1993); Spyromilio et al. (1993)
27 Dec 1988	673	AAT	UCLES	5702	10190	30000	Hanuschik et al. (1993); Spyromilio et al. (1993)
06 Feb 1989	714	CTIO-1.5m	Cass.	6420	10380	16	Phillips et al. (1990)
09 May 1989	806	CTIO-1.5m	Cass.	6430	10330	16	Phillips et al. (1990)
12 Jan 1990	1054	CTIO-4m	RC	3565	10000	11	Suntzeff et al. (1991)
12 Mar 1991	1478	CTIO-4m	RC	3245	9175	11	
30 Mar 1992	1862	HST	STIS	4569	6818	4.4	Wang et al. (1996)
14 Mar 1993	2211	HST	STIS	4569	6818	4.4	Wang et al. (1996)
07 Jan 1995	2875	HST	STIS	4569	6818	4.4	Chugai et al. (1997)
23 Sep 1996	3500	HST	STIS	4569	6818	4.4	
05 Jan 1997	3604	HST	STIS	4569	6818	4.4	
10 Dec 2000	5039	VLT	UVES	4760	6840	50000	Grönningsson et al. (2006, 2007)
06 Oct 2002	5704	VLT	UVES	4760	6840	50000	Grönningsson et al. (2006, 2007, 2008)
21 Mar 2005	6601	VLT	UVES	4760	6840	50000	Grönningsson et al. (2006, 2007)
23 Oct 2007	7547	VLT	UVES	4760	6840	50000	Grönningsson et al. (2007)
07 Feb 2009	8020	VLT	UVES	4800	6800	50000	Tziampzis et al. (2010)

Table 4.2. H α full-width half-maxima (FWHM) and the half-width zero intensities (HWZI) determined by the zero intensity velocity on the blue side of the line. The tabulated line widths have been corrected for the relevant instrumental resolution.

day	FWHM (\AA)	HWZI (\AA)
524	3200	3600
611	2700	3400
673	1600	3700
714	3100	4500
806	3200	5500
1054	2100	5600
1478	1400	6600
1862	1600	6800
2211	1400	6700
2875	2700	6700
3500	3500	7000
3604	2100	7000

4.1.1 Contamination of the H α profiles

The H α profile at day 714 exhibits a very slight inflection visible at $V \approx +900 \text{ km s}^{-1}$. By day 806, this slight inflection has developed into a noticeable shoulder in the line profile of H α (see Figure 4.4).

Although these features are similar in nature to features produced by dust absorption in the flat-topped region (as discussed in Section 3.2.5), we conclude that this shoulder is an early appearance of the unresolved [N II] $\lambda 6583 \text{ \AA}$ line from the equatorial ring (Kozma & Fransson 1998b). Unresolved nebular [N II] lines at $\lambda = 6583 \text{ \AA}$ and $\lambda = 6548 \text{ \AA}$ either side of the H α rest frame velocity at 6563 \AA are certainly seen by day 1054 and have to be removed in order to consider the evolution of the broad H α profile (see Figure 4.1). We do not remove this potential contaminant at earlier epochs but try to fit the broad line profiles around it.

By day 1054, all three of the narrow nebular lines are strong. They remain unresolved in the low spectral resolution CTIO data at days 1054 and 1478 and therefore contaminate the entire central region of the H α line profile. Their presence renders two CTIO H α profiles from days 1054 and 1478 unusable for modelling purposes. The HST and VLT H α profiles at later epochs (≥ 1862 days) have a higher spectral resolution and it was therefore easier to remove the narrower [N II] and H α lines from the broad H α profiles (for example Figures 4.1 and 4.2). Although this does remove a potentially informative

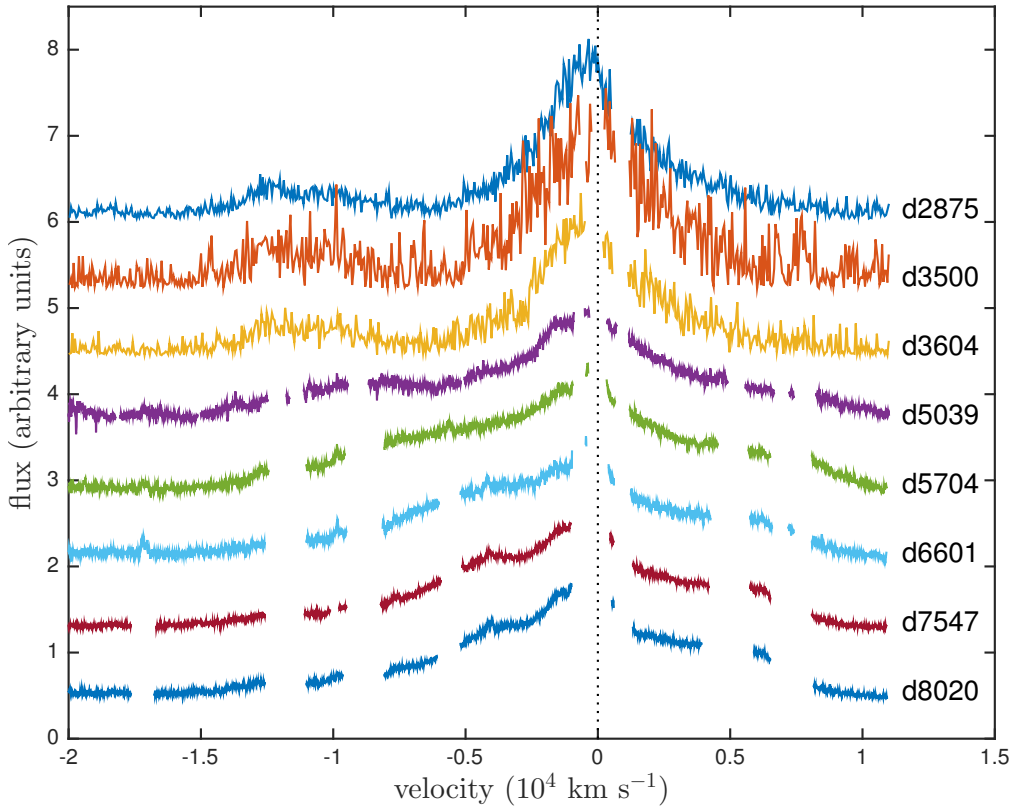


Figure 4.2. Archival data showing the evolution of the H α line profile from SN 1987A at the later epochs. The spectral gaps correspond to where narrow line emission from the equatorial ring has been removed. The spectra have been continuum-subtracted and offsets applied for display purposes.

section of the profile ($+500 \text{ km s}^{-1} < v < +1500 \text{ km s}^{-1}$), we achieve good fits to the overall line profiles at these epochs.

4.1.2 The evolution of the maximum and minimum velocities

For a freely expanding medium, the velocity of any fractional radial element should not change with time. The maximum velocity of any line-emitting region is therefore expected to be constant. However, at the epochs we consider here, it appears that the maximum velocities of the H α line, as determined by the velocity at zero intensity on the blue side, generally increase over time (see Table 4.2). We attribute this to the start of the freeze-out phase in the outer regions of the ejecta, while the hydrogen neutral fraction is still increasing in the denser inner regions (Danziger et al. 1991; Fransson & Kozma 1993).

The onset of a fixed ionization structure in the ejecta causes the rate of H α flux decline to slow. Since the outer, faster moving regions reach this state at earlier times than the

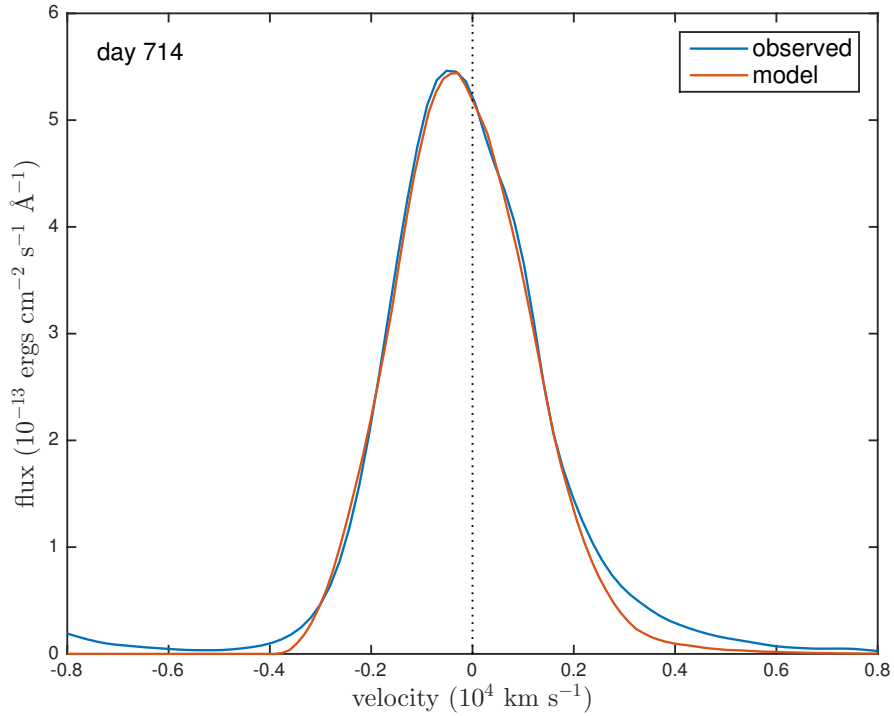


Figure 4.3. Amorphous carbon smooth dust fit to the day 714 H α line of SN 1987A using an MRN size distribution, illustrating the underestimation of the red scattering wing for small grain radii. Model parameters are the same as the smooth dust fit for day 714 (Table 4.4) except for the grain size distribution and dust mass: $M_{dust} = 8.0 \times 10^{-6} M_{\odot}$, $a_{min} = 0.005 \mu\text{m}$, $a_{max} = 0.25 \mu\text{m}$ and $n(a) \propto a^{-3.5}$.

inner, slower moving regions, the relative flux contribution of the outer regions is increased. At early epochs ($t < 900$ days) the flux contribution from hydrogen in the core dominates the overall H α flux, whereas at later epochs ($t > 900$ days) the flux from the envelope dominates (Fransson & Kozma 1993; Kozma & Fransson 1998a). This shift likely explains apparent broadening of the line with the higher velocity material becoming increasingly noticeable in the line profiles. This may also explain the increase in HWZI velocities at these epochs with the relative flux from the very densest regions dropping more rapidly relative to the outer line-emitting region. The full-width half maximum (FWHM) remains relatively steady (see Table 4.2). However, the FWHM values presented in Table 4.2 were difficult to determine accurately since the peak of the broad line profile is contaminated by narrow line emission from the equatorial ring.

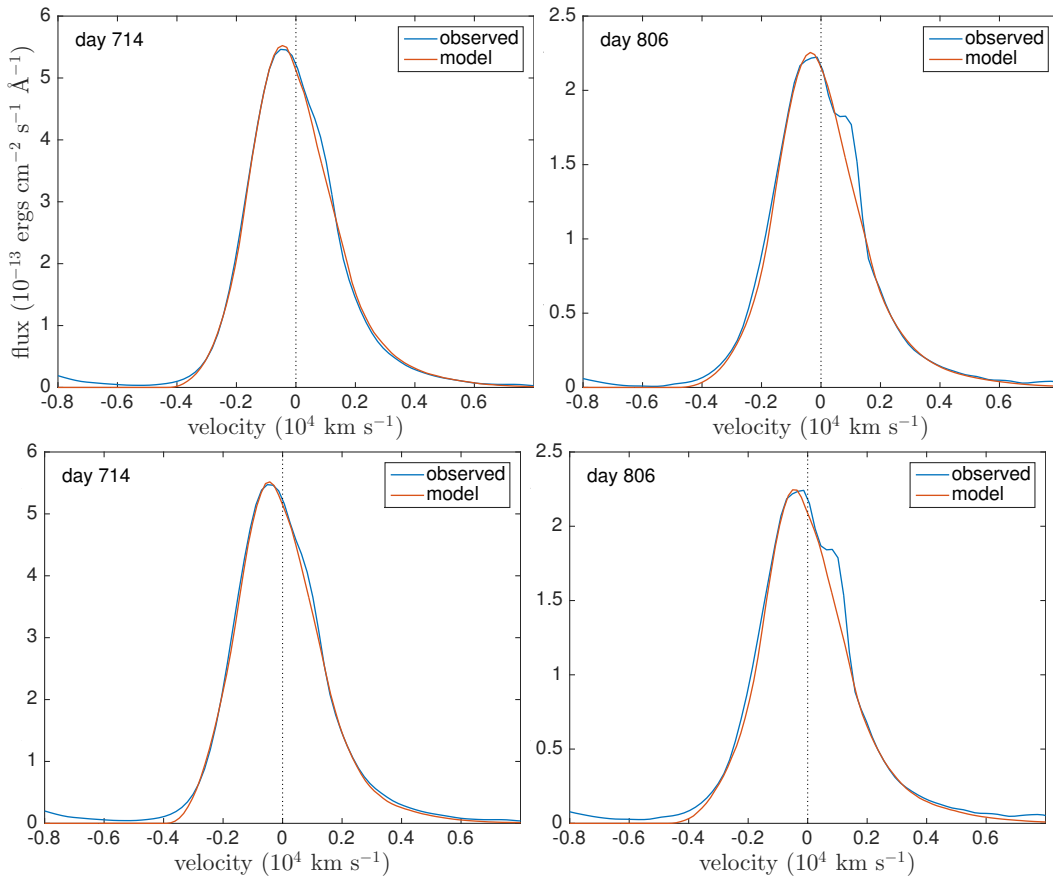


Figure 4.4. Best model fits to the SN 1987A H α line at day 714 and day 806 for the parameters detailed in Tables 4.4 and 4.5. The two fits on the top are smooth dust models using amorphous carbon grains of radius $a = 0.35 \mu\text{m}$ and the two fits on the bottom are clumped dust models using amorphous carbon grains of radius $a = 0.6 \mu\text{m}$.

4.2 Modelling SN 1987A

We have modelled the H α line of SN 1987A at days 714, 806, 1862, 2211, 2875, 3500 and 3604, and the [O I] $\lambda 6300, 6363 \text{ \AA}$ doublet at days 714, 806, 1054 and 1478. After day 3604 the H α profile begins to become dominated by emission from the reverse shock and the structure of the emitting region may no longer be approximated by a single shell model as we do here (Fransson et al. 2013). The [O I] $\lambda 6300, 6363 \text{ \AA}$ doublet becomes too weak to model after day 1478 (see Figure 4.1). We continue to adopt a velocity profile $V(r) = \frac{V_{max}}{R_{max}}r$ and treat the variable parameters listed at the start of Section ???. Whilst the albedo and optical depth are not varied directly, they are altered by adjusting the dust mass, M_{dust} , and the grain size, a , which together determine the albedo and optical depth via Mie theory and the optical properties of the dust.

Table 4.3. Observed luminosities of the H α line and estimated electron scattering optical depths from R_{in} to R_{out} for the radii detailed in Tables 4.4 to 4.5 based on an assumed gas temperature of 10,000 K.

day	H α		[O I]		
	L_{obs} (10^{37} erg s $^{-1}$)	L_{undep}/L_{obs}	L_{obs} (10^{37} erg s $^{-1}$)	L_{undep}/L_{obs}	τ_e (10^{-2})
714	1.36	1.65	0.313	3.57	1.44
806	0.57	1.77	0.0942	3.57	0.840
1054			0.0242	3.23	
1478			0.00185	2.70	
1862	0.0063	2.06			0.159
2211	0.0041	2.07			0.0378
2875	0.0019	2.84			0.0219
3500	0.00079	3.16			0.0125
3604	0.00098	3.27			0.0149

In all models, the ejecta occupies a shell with inner radius R_{in} and outer radius R_{out} . Packets are emitted according to a smooth density profile assuming recombination or collisional excitation such that $i(r) \propto \rho(r)^2 \propto r^{-2\beta}$. Initially the dust is considered to have a smooth density distribution and is assumed to be coupled to the gas so as to follow the same radial profile. A clumped distribution of dust is considered later (see Section 4.2.2).

We estimate the electron scattering optical depths assuming an electron temperature of 10,000 K between R_{in} and R_{out} based on the observed fluxes of the H α line. A temperature of 10,000 K is likely too high at the epochs considered but we adopt it in order not to underestimate electron scattering optical depths. The values we calculate from the observed H α luminosities are listed in Table 4.3. Since the electron scattering optical depths at these epochs are negligibly small we therefore do not include electron scattering in the models.

There is rarely a unique set of parameters that provide the best fit to the data. However, the majority of the parameters of interest can be well constrained from our modelling by considering different elements of the shape of the profile. In particular, by constructing fits to the data using minimum and maximum limits for the grain radius, credible lower and upper bounds on the dust mass formed within the ejecta may be derived. We present here fits to the data obtained using both small and large values of the grain radius a since it is the grain size which has the most significant effect on the overall dust mass required to reproduce the line profile (see Section ??).

All of our models are of a dusty medium composed solely of amorphous carbon grains. We use the optical constants from the BE sample presented by Zubko et al. (1996). Although previous SED modelling of SN 1987A limited the fraction of silicates present in the dusty ejecta to a maximum of 15% (Ercolano et al. (2007), W15), the recent work of Dwek & Arendt (2015) has suggested that a large mass of mostly silicate dust may have formed at early epochs (~ 615 days). It is therefore useful to consider the effects on our models of using silicate dust. We discuss this in detail in Sections 4.2.6 and 4.2.7.

For each profile, the maximum velocity is initially identified from the data as the point where the emission vanishes on the blue side and is then varied throughout the modelling in order to produce the best fit. The equivalent point on the red side is indeterminate from observations due to the effects of dust scattering. We determine the approximate value of V_{min} by examining the width of the profile near its peak. On the red side the theoretical minimum velocity often falls at a similar velocity to the 6583Å line so any dust-induced features near this wavelength that would allow a more accurate determination of V_{min} can be overwhelmed by the nebular line. Having determined the minimum and maximum velocities, the ratio of the inner and outer radii of the supernova ejecta can be determined since $R_{in}/R_{out} = V_{min}/V_{max}$. The outer radius is calculated from the epoch and the maximum velocity.

The only parameters that remain to be determined are the exponent of the density profile β , the mean grain radius and the total dust mass. The shape of the blue wing is solely a product of the density profile and the dust mass; the height and shape of the red wing is a product of these and also of the scattering efficiency of the grains (the albedo ω); the extent and shape of the asymmetry in the flat-topped portion of the profile is a function of only the total dust optical depth determined by the dust mass and the grain radius. By iterating over these three parameters, an excellent fit to the data can usually be obtained.

Models are produced in the same manner for the [O I] $\lambda\lambda 6300, 6363$ Å doublet as for the single H α line, with each component of the doublet being modelled independently and the resulting profiles added according to a specified ratio. Although the theoretical intrinsic flux ratio is 3.1 for optically thin emission (Storey & Zeippen 2000), the actual ratio between the two components can be affected by self-absorption (Li & McCray 1992) and we therefore left it as a free parameter. The deduced doublet ratios are listed in Tables 4.4, 4.5 and 4.6.

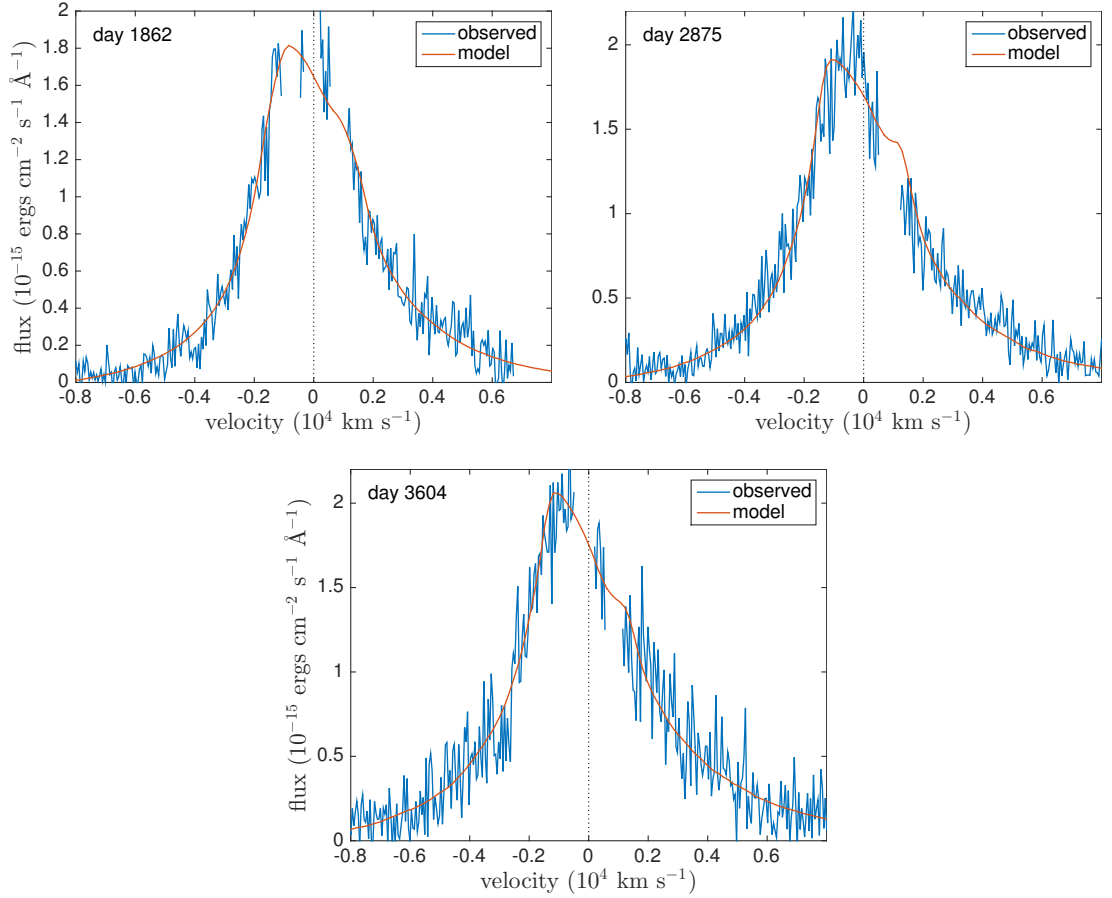


Figure 4.5. Best model fits to the SN 1987A H α line at days 1862, 2875 and 3604 for the parameters detailed in Tables 4.4, 4.5 and 4.6. On the top row are smooth model fits with amorphous carbon grains of radius $a = 0.35 \mu\text{m}$. On the middle and bottom rows are clumped model fits with amorphous carbon grains of radii $a = 0.6 \mu\text{m}$ and $a = 3.5 \mu\text{m}$ respectively.

For all lines, though particularly at very late epochs, even small fluctuations in the adopted value of the continuum level can have a substantial effect on the fit to the resulting profile. Since it is not feasible to establish the level of the continuum so precisely, the value of the continuum has been left as a free parameter that may be adjusted (to within sensible margins) in order to allow for the widest possible dust mass range to be determined. We generally find it is necessary to assume a continuum level that is slightly lower where the dust mass is higher. The [O I] $\lambda\lambda 6300, 6363 \text{ \AA}$ doublets at days 1054 and 1478 are weak relative to the continuum and are also blended with the wings of other lines making it difficult to fit their wings accurately. We aim to fit the lines between approximately -3000 km s^{-1} and $+5000 \text{ km s}^{-1}$ but present a wider velocity range for context (for example see Figure 4.9).

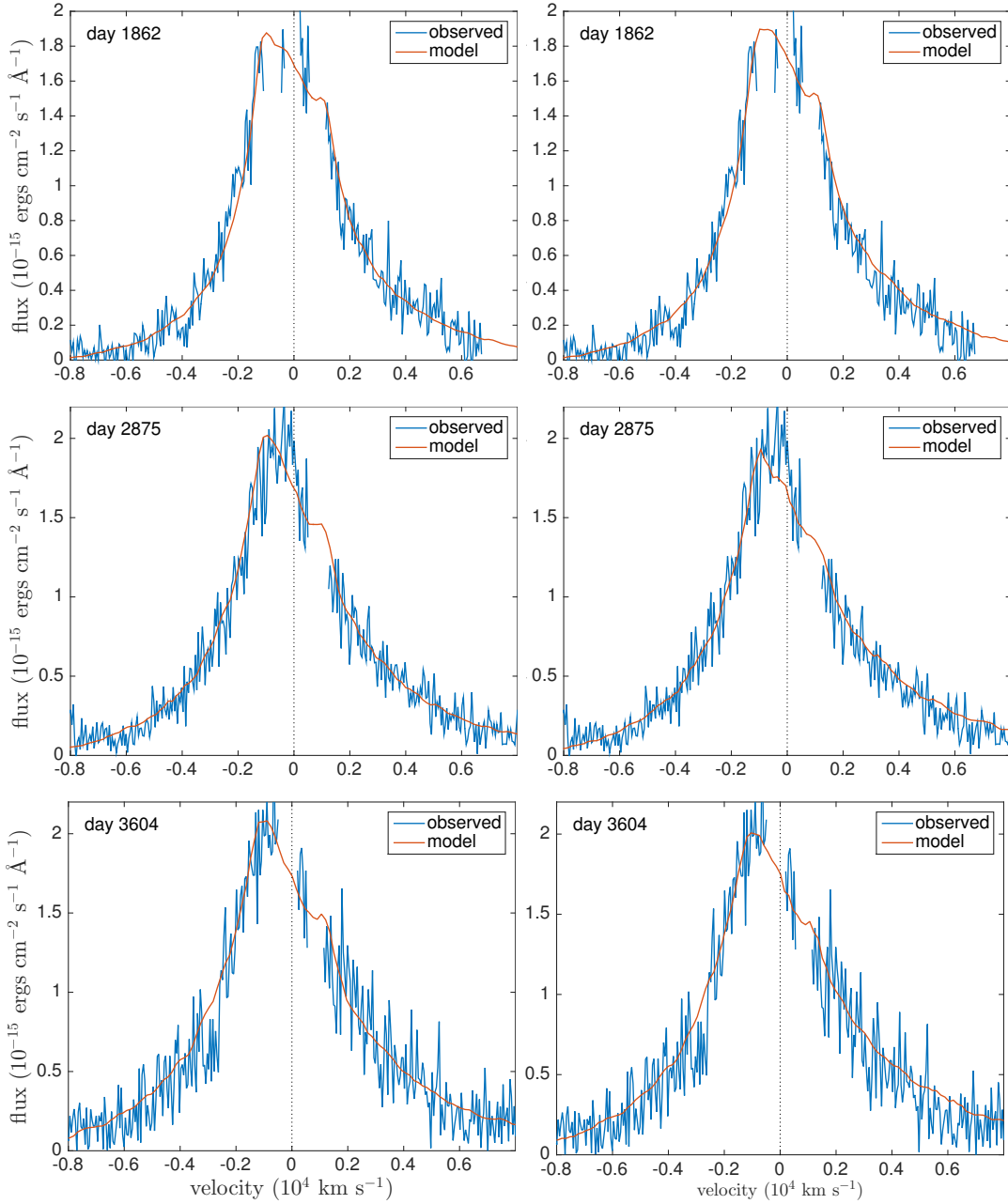


Figure 4.6. Best model fits to the SN 1987A H α line at days 1862, 2875 and 3604 for the parameters detailed in Tables 4.4, 4.5 and 4.6. On the top row are smooth model fits with amorphous carbon grains of radius $a = 0.35 \mu\text{m}$. On the middle and bottom rows are clumped model fits with amorphous carbon grains of radii $a = 0.6 \mu\text{m}$ and $a = 3.5 \mu\text{m}$ respectively.

Fits to the H α line profile at days 2211 and 3500 are omitted for the sake of space but are very similar to those of days 1862 to 3604. All profiles have been smoothed to approximately the same resolution as the observed profiles using a moving-average procedure. Parameters for the models at all epochs including days 2211 and 3500 are detailed in Tables 4.4 to 4.6.

4.2.1 Smooth Density Models for SN 1987A

Even at the earliest epochs there is a substantial wing on the red side of the H α line profile that cannot be fitted by scattering from moving grains with a low albedo. The minimum required albedo is approximately $\omega \approx 0.5$ implying relatively large grain radii. As previously discussed, the larger the grain size the larger the mass of dust required to reproduce the same optical depth. Figure 4.3 illustrates the fit for the day 714 H α profile for the case where a classic MRN (Mathis et al. 1977) grain size distribution is adopted, with $a_{min} = 0.005\mu\text{m}$, $a_{max} = 0.25\mu\text{m}$ and $n(a) \propto a^{-3.5}$. It can be seen clearly that the extended red wing is significantly underestimated. Since the albedo of amorphous carbon grains varies significantly with grain radius (see Figure 3.6) we can establish a strong lower bound to the mean dust grain radius, which we estimate to be $a \geq 0.35\mu\text{m}$. This is the smallest grain size that is still capable of reproducing the red scattering wing at all epochs and we therefore use this lower limit value throughout our smooth density modelling.

The inner and outer radii of the ejecta are calculated at each epoch from the maximum velocity used, the day number and the specified ratio R_{in}/R_{out} . The radii generated are consistent with those used in previous models of SN 1987A (Ercolano et al. (2007), W15) and the minimum velocities for both the [O I] and H α line emitting regions are relatively consistent with those obtained by Kozma & Fransson (1998b) who estimate that hydrogen extends into the core to a depth of $\lesssim 700\text{ km s}^{-1}$ and the oxygen reaches down to $\sim 400\text{ km s}^{-1}$. They are also consistent with predictions from 3D explosion models at the time of shock-breakout that predict the oxygen to reach to a depth of $\sim 200\text{ km s}^{-1}$ (Hammer et al. 2010; Wongwathanarat et al. 2015). Figures 4.4 and 4.6 show the best fits to the data for days 714 to 3604 whilst Table 4.4 details the parameters used.

It can be seen from Tables 4.4 to 4.6 that, in order to reproduce the blueshifts seen in the [O I] $\lambda 6300, 6363\text{ \AA}$ doublet, considerably larger dust masses are required than to fit the H α line at the same epoch. Although the same maximum velocities and therefore outer radii are used in our [O I] and H α models, the inner radii for the [O I] models are

significantly smaller and the density distribution much steeper. This implies that [O I] is concentrated towards the centre of the ejecta whereas H α is more diffuse. This is broadly in agreement with 3D explosion dynamics models that suggest that a few hours after the explosion the heavier elements will, in comparison to hydrogen, be located more centrally in the ejecta with “bullets” of heavier material reaching the outer edges (Hammer et al. 2010). If dust is forming in the inner regions of the ejecta then the majority of the [O I] emission must travel through the newly formed dust whereas the more diffuse H α emission has a greater chance of escaping unaffected. This may explain the difference between the dust masses needed for the [O I] and H α models.

Table 4.4. The parameters used for the best fitting smooth models of SN 1987A with amorphous carbon grains of radius $a = 0.35 \mu\text{m}$. Optical depths are given from R_{in} to R_{out} at $\lambda = 6563 \text{ \AA}$ for H α and $\lambda = 6300 \text{ \AA}$ for [O I]. Values of τ_V are very close to the quoted values of $\tau_{H\alpha}$.

	day	V_{max} (km s $^{-1}$)	V_{min} (km s $^{-1}$)	R_{in}/R_{out}	β	M_{dust} (M_\odot)	R_{out} (cm)	R_{in} (cm)	[O I] ratio	τ_λ
[O I]	714	3250	228	0.07	2.9	9.65×10^{-5}	2.00×10^{16}	1.40×10^{15}	2.6	3.60
[O I]	806	4000	240	0.06	2.4	1.50×10^{-4}	2.79×10^{16}	1.67×10^{15}	2.3	2.86
[O I]	1054	4300	215	0.05	2.1	2.35×10^{-4}	3.92×10^{16}	1.96×10^{15}	2.7	2.23
[O I]	1478	4500	180	0.04	1.7	2.95×10^{-4}	5.75×10^{16}	2.30×10^{15}	3.0	1.30
H α	714	3250	813	0.25	1.2	2.10×10^{-5}	2.00×10^{16}	5.01×10^{15}	0.61	
H α	806	4000	880	0.22	1.9	3.80×10^{-5}	2.79×10^{16}	6.13×10^{15}	0.59	
H α	1862	8500	1275	0.15	1.9	5.00×10^{-4}	1.37×10^{17}	2.05×10^{16}	0.35	
H α	2211	9000	1260	0.14	1.9	9.25×10^{-4}	1.72×10^{17}	2.41×10^{16}	0.42	
H α	2875	9500	1330	0.14	1.9	1.50×10^{-3}	2.36×10^{17}	3.30×10^{16}	0.36	
H α	3500	10000	1400	0.14	1.9	3.35×10^{-3}	3.02×10^{17}	4.23×10^{16}	0.49	
H α	3604	10250	1333	0.13	1.9	4.20×10^{-3}	3.19×10^{17}	4.15×10^{16}	0.55	

Table 4.5. The parameters used for the best fitting clumped models of SN 1987A with amorphous carbon grains of radius $a = 0.6 \mu\text{m}$. Optical depths are given from R_{in} to R_{out} at $\lambda = 6563 \text{ \AA}$ for H α and $\lambda = 6300 \text{ \AA}$ for [O I]. Values of τ_V are very close to the quoted values of $\tau_{H\alpha}$.

	day	V_{max} (km s $^{-1}$)	V_{min} (km s $^{-1}$)	R_{in}/R_{out}	β	M_{dust} (M_\odot)	R_{out} (cm)	R_{in} (cm)	[O I] ratio	τ_λ
[O I]	714	3250	228	0.07	2.7	2.00×10^{-4}	2.00×10^{16}	1.40×10^{15}	2.3	3.84
[O I]	806	4000	240	0.06	2.3	4.00×10^{-4}	2.79×10^{16}	1.67×10^{15}	2.0	4.02
[O I]	1054	4300	215	0.05	2.3	7.50×10^{-4}	3.92×10^{16}	1.96×10^{15}	2.3	3.85
[O I]	1478	4500	180	0.04	2.0	1.10×10^{-3}	5.75×10^{16}	2.30×10^{15}	2.8	2.65
H α	714	3250	813	0.25	1.4	5.50×10^{-5}	2.00×10^{16}	5.01×10^{15}	0.87	
H α	806	4000	880	0.22	1.8	9.00×10^{-5}	2.79×10^{16}	6.13×10^{15}	0.76	
H α	1862	8500	1190	0.14	1.9	1.20×10^{-3}	1.37×10^{17}	1.91×10^{16}	0.46	
H α	2211	9000	1260	0.14	1.9	3.00×10^{-3}	1.72×10^{17}	2.41×10^{16}	0.73	
H α	2875	9500	1140	0.12	2	8.00×10^{-3}	2.36×10^{17}	2.83×10^{16}	1.05	
H α	3500	10000	1200	0.12	2	1.35×10^{-2}	3.02×10^{17}	3.63×10^{16}	1.08	
H α	3604	10250	1230	0.12	2	1.70×10^{-2}	3.19×10^{17}	3.83×10^{16}	1.22	

Table 4.6. The parameters used for the best fitting clumped models of SN 1987A with amorphous carbon grains of radius $a = 3.5 \mu\text{m}$. Optical depths are given from R_{in} to R_{out} at $\lambda = 6563 \text{ \AA}$ for H α and $\lambda = 6300 \text{ \AA}$ for [O I]. Values of τ_V are very close to the quoted values of $\tau_{H\alpha}$.

	day	V_{max} (km s $^{-1}$)	V_{min} (km s $^{-1}$)	R_{in}/R_{out}	β	M_{dust} (M_\odot)	R_{out} (cm)	R_{in} (cm)	[O I] ratio	τ_λ
[O I]	714	3250	228	0.07	2.9	1.50×10^{-3}	2.00×10^{16}	1.40×10^{15}	2.3	4.20
[O I]	806	4000	240	0.06	2.3	2.70×10^{-3}	2.79×10^{16}	1.67×10^{15}	2.1	3.95
[O I]	1054	4300	215	0.05	2.3	5.50×10^{-3}	3.92×10^{16}	1.96×10^{15}	2.5	4.12
[O I]	1478	4500	180	0.04	1.9	8.00×10^{-3}	5.75×10^{16}	2.30×10^{15}	2.8	2.81
H α	1862	8500	1190	0.14	1.9	1.00×10^{-2}	1.37×10^{17}	1.91×10^{16}	0.55	
H α	2211	9000	1260	0.14	1.9	2.40×10^{-2}	1.72×10^{17}	2.41×10^{16}	0.85	
H α	2875	9500	1140	0.12	2	6.00×10^{-2}	2.36×10^{17}	2.83×10^{16}	1.15	
H α	3500	10000	1200	0.12	2	1.15×10^{-1}	3.02×10^{17}	3.63×10^{16}	1.34	
H α	3604	10250	1230	0.12	2	1.25×10^{-1}	3.19×10^{17}	3.83×10^{16}	1.31	

4.2.2 Clumped Dust Models for SN 1987A

A number of investigators have presented arguments for the material in the ejecta of SN 1987A being clumped (Lucy et al. 1991; Li & McCray 1992; Kozma & Fransson 1998b) and so we consider clumped models for the ejecta dust to be more realistic than smoothly distributed dust models. It has been shown through the modelling of optical-IR SEDs that when dust is assumed to have a clumped distribution then the derived dust masses can be significantly larger than for the case of dust that is distributed smoothly between the inner and outer radii (e.g. Ercolano et al. (2007); Owen & Barlow (2015)). We present two sets of fits to the line profile based on the clumped dust modelling of W15, one set with a minimum grain size and one set with a maximum grain size. Each fit is based on the best fitting smooth model such that the photon packets are emitted assuming a smooth radial density profile. However, the dust is no longer coupled to the gas but instead is located entirely in clumps of size $R_{out}/25$. The clumps are distributed stochastically between R_{in} and R_{out} with the probability of a given grid cell being a clump proportional to $r^{-\beta}$ where $i(r) \propto r^{-2\beta}$. The number of clumps used is determined by the clump filling factor f which is kept constant at $f = 0.1$. All properties are fixed from the smooth models with the exception of the grain radius, density profile exponent (β) and the total dust mass.

Models were again constructed using the smallest possible grain radius ($a=0.6 \mu\text{m}$ in the clumped case) in order to derive minimum dust masses for clumped distributions. By considering the extent of the red scattering wing, upper limits to the grain size were also derived with the purpose of limiting the maximum dust mass at each epoch. By steadily reducing the grain radius from an initial value of $5 \mu\text{m}$ (motivated by the maximum possible grain size derived by W15 for their day 8515 model), we produced a set of models with a maximum grain radius of $a = 3.5 \mu\text{m}$.

The increase in grain size from the smooth case to the clumped case is necessary in order to have a slightly larger albedo. Grains of radius $a = 0.35 \mu\text{m}$ do not reproduce the red side of the profiles well for a clumped medium. This is because when the dust is located in clumps the radiation is subject to less scattering as well as to less absorption. The reduction in scattering appears not to be compensated for by the increased dust mass and a larger grain radius is therefore required, particularly at day 714.

For all but the H α line at days 714 and 806 a similar fit could be obtained with either a grain radius of $a = 0.6 \mu\text{m}$ or $a = 3.5 \mu\text{m}$ (see Figures 4.4 and 4.6). However, for

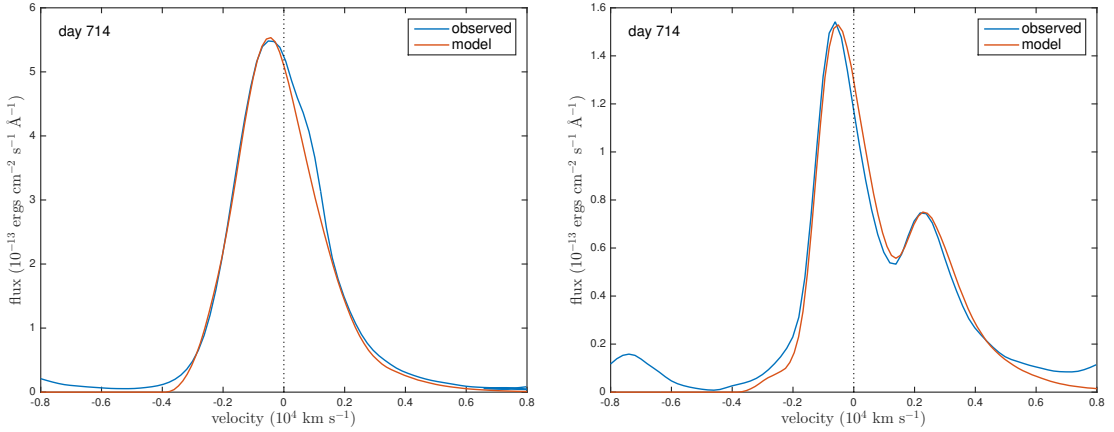


Figure 4.7. Fits to the H α and [O I] $\lambda 6300, 6363 \text{ \AA}$ lines at day 714 using the more complex dust model described in Section 4.2.4 with a dust mass of $2.3 \times 10^{-4} M_{\odot}$.

H α at days 714 and 806 even a small change to the grain radius from $0.6 \mu\text{m}$ resulted in a significantly poorer fit, either over- or under-estimating the red wing. We therefore conclude that the dust mass estimates produced for the H α lines at days 714 and 806 for a grain radius of $a = 0.6 \mu\text{m}$ are the best H α -based estimates of the dust mass at this epoch.

In our subsequent analyses, we adopt the values derived from our clumped models. Details of the parameters used are presented in Tables 4.5 and 4.6 and the fits are presented in Figures 4.4 and 4.6.

4.2.3 The effects of clumping

As in the case of SED radiative transfer models, the dust masses required to reproduce the observations in the clumped scenario are considerably higher than for the smooth scenario. The dust masses differ between our smooth models for $a = 0.35 \mu\text{m}$ and clumped models for $a = 0.6 \mu\text{m}$ by a factor of approximately 3. The dust mass estimates are even larger when comparing clumped $a = 0.6 \mu\text{m}$ models to clumped $a = 3.5 \mu\text{m}$ models at later epochs. This does not take into account the increase in grain radius between the two cases however. This increase accounts for a reasonable fraction of this difference. We estimate the effects of clumping alone to increase the required dust mass by a factor of approximately 1.5-2.0 from the smooth case.

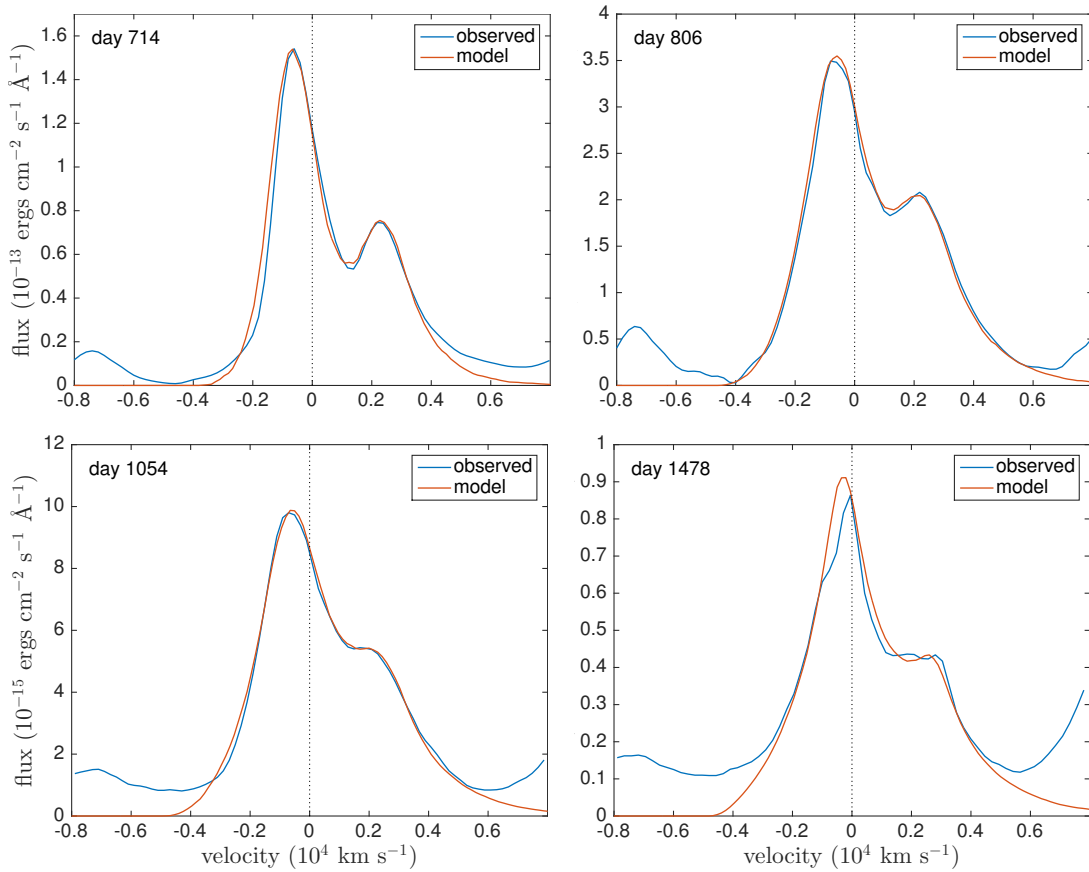


Figure 4.8. Best smooth dust fits to the SN 1987A [O I] $\lambda 6300, 6363 \text{ \AA}$ doublet at days 714, 806, 1054 and 1478 for the parameters detailed in Tables 4.4, 4.5 and 4.6. On the top row are smooth dust fits with amorphous carbon grains of radius $a = 0.35\mu\text{m}$. On the middle and bottom rows are clumped dust fits with amorphous carbon grains of radii $a = 0.6\mu\text{m}$ and $a = 3.5\mu\text{m}$ respectively.

4.2.4 More complex models

Where blue-shifted lines are observed in the spectra of CCSNe it is often the case that the Balmer lines of HI are less affected than the [O I] lines (Milisavljevic et al. 2012). This may be due to a difference in the location or distribution of the emitting elements; if the neutral hydrogen was diffusely distributed throughout the envelope but the oxygen was co-located with the dust in the core and in clumps then this could result in [O I] emission undergoing greater attenuation than H α . This geometry would be in line with previous models of SN 1987A that suggested that the dust-forming regions are likely to include those which are oxygen-rich (Kozma & Fransson 1998a). Clearly, any model of dust formation in the ejecta of a CCSN must consistently reproduce all of the line profiles at a given epoch. The models presented in this paper thus far have coupled the gas and

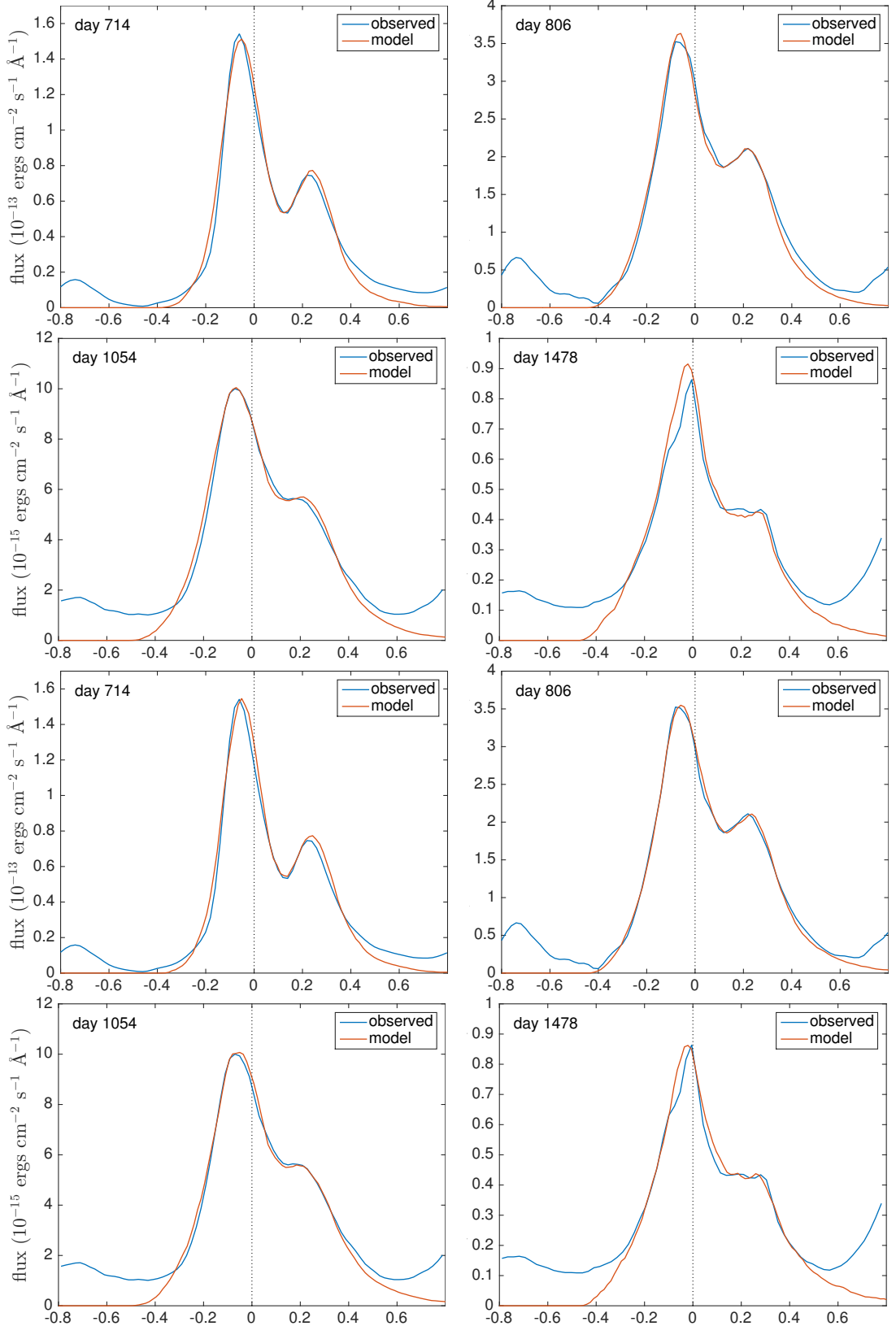


Figure 4.9. Best smooth dust fits to the SN 1987A [O I] $\lambda\lambda 6300,6363$ Å doublet at days 714, 806, 1054 and 1478 for the parameters detailed in Tables 4.4, 4.5 and 4.6. On the top row are smooth dust fits with amorphous carbon grains of radius $a = 0.35\mu\text{m}$. On the middle and bottom rows are clumped dust fits with amorphous carbon grains of radii $a = 0.6\mu\text{m}$ and $a = 3.5\mu\text{m}$ respectively.

dust distributions for a fixed clump volume filling factor and clump size. The H α and [O I] models therefore require different dust masses with the [O I] models usually requiring a dust mass ~ 4 times larger than the H α models.

We now present a model that reconciles this difference by additionally varying the clump filling factor, clump size and emissivity distribution. We assume that neutral hydrogen is likely diffuse throughout the ejecta and so maintains a smoothly distributed power-law emissivity distribution between R_{in} and R_{out} for H α . However, we now assume that dust mostly forms in dense regions of high metallicity and so restrict the [O I] $\lambda 6300, 6363$ Å emission to originate largely from the dusty clumps with only a small fraction emitted from the inter-clump medium. As previously discussed, the greater the covering factor of the dust the greater the albedo required in order to reproduce the H α red scattering wing. In order to obtain both the strong blue-shifting of the [O I] line and the extended red scattering wing observed in H α a small number of dense clumps were required along with a small mass of diffusely distributed highly scattering dust in the inter-clump medium.

In order to fit both line profiles simultaneously we required a very high albedo ($\omega > 0.8$) that demanded the inclusion of some fraction of silicate dust. Amorphous carbon grains alone are incapable of producing this level of scattering for any grain size. We adopted a grain radius of $a = 0.6\mu\text{m}$, the same as that used in our initial clumped models and we varied the relative proportions of amorphous carbon and MgSiO₃ in order to achieve the necessary albedo. The adopted grain densities were $\rho_c = 1.85 \text{ g cm}^{-3}$ for amorphous carbon grains and $\rho_s = 2.71 \text{ g cm}^{-3}$ for MgSiO₃. The resulting dust model for day 714 used 75% MgSiO₃ and 25% amorphous carbon by cross-sectional area with a volume filing factor $f_V = 0.1$ and a clump size $R_{out}/5$. 90% of the dust mass was located in clumps with the remaining 10% emitted smoothly between R_{in} and R_{out} according to a power law $\rho \propto r$. Clumps were distributed stochastically with probability $\propto r^{-8}$ compared to $r^{-2.7}$ in our standard models discussed earlier. Equal numbers of [O I] packets were emitted from each clump. H α was distributed smoothly according to a density power law $\rho(r) \propto r^{-1.3}$. R_{out} was the same for all components (i.e. clumped dust, diffuse dust, [O I] emission and H α emission) and was calculated using a maximum velocity of 3250 km s⁻¹. The inner radius was $R_{in} = 0.07R_{out}$ for all components except the smooth H α emission which was emitted between $R_{in} = 0.25R_{out}$ and R_{out} .

The total dust mass used was $M_{dust} = 2.3 \times 10^{-4} \text{ M}_\odot$. This dust mass is very similar to that derived from our original clumped models of [O I] using amorphous carbon grains

of radius $a = 0.6\mu\text{m}$. The slight increase over our amorphous carbon dust mass of $1.5 \times 10^{-4} M_{\odot}$ is largely due to the higher grain density of MgSiO₃. At this grain radius amorphous carbon and MgSiO₃ have similar extinction efficiencies and so the change in species and geometry does not substantially alter the dust mass. We therefore adopt the [O I] dust masses in our further analyses and consider the differences in our derived dust masses between H α and [O I] to be the result of the clumped emission of [O I].

Fits to both the [O I] $\lambda 6300, 6363$ Å and H α lines for day 714 using these parameters are presented in Figure 4.7.

4.2.5 The effect of a grain size distribution

It is important to consider the potential effect on the dust mass of modelling a grain size distribution instead of a single grain size. For a grain size distribution the overall extinction cross section, C_{ext} , at a given wavelength is

$$C_{ext} = \int_{a_{min}}^{a_{max}} Q_{ext}(a)n(a)\pi a^2 da \quad (4.1)$$

where $Q_{ext}(a)$ is the extinction efficiency for a grain size a and $n(a)$ is the number of grains with size a . The overall extinction efficiency is then

$$Q_{ext} = \frac{C_{ext}}{\int_{a_{min}}^{a_{max}} n(a)\pi a^2 da} \quad (4.2)$$

The scattering cross-section Q_{sca} is similarly calculated. As a result of these calculations, there is rarely a single grain size that has the same albedo and extinction efficiency as a size distribution. Modelling a size distribution may therefore alter the deduced dust mass. Since the models are only sensitive to the overall optical depth and albedo, it is not possible to deduce the grain size range or distribution and only single grain sizes are investigated (as presented above).

Whilst this apparently limits the scope of the results, it is useful to consider the extent to which different grain size distributions would alter the derived dust masses. By considering a number of grain radius ranges and adopting a power law distribution with a variable exponent, we may gain some insight into the effects of adopting a distribution

rather than a single size. As discussed in Section 4.2.1, a classical MRN power law ($n(a) \propto a^{-3.5}$) with a wide grain radius range ($a_{min} = 0.001 \mu\text{m}$ to $a_{max} = 4.0 \mu\text{m}$) the derived albedo is much too small to reproduce the required wing seen at early epochs. We therefore adopt an approach whereby, for a number of grain size ranges, we adjust the exponent of the distribution until the overall albedo is the same as that seen for the best fitting single grain radius for the clumped distributions. We may then approximately calculate the required dust mass as

$$M_d = \frac{M_s Q_{ext,s}(a_s)}{a_s} \times \frac{\int_{a_{min}}^{a_{max}} n(a) a^3 da}{\int_{a_{min}}^{a_{max}} Q_{ext}(a) n(a) a^2 da} \quad (4.3)$$

where the subscript s represents the single grain size quantities and the d subscript represents quantities for the grain size distribution.

We calculate the required dust masses for the clumped H α model on day 714 for a selection of distributions with varying a_{min} . These are presented in Table 4.7. It can be seen that in all cases, a larger dust mass is required for grain size distributions in order to reproduce the same profile as a single grain size. The conversion factors presented in the table are valid for any model with grain size $a = 0.6 \mu\text{m}$ and may therefore also be applied to the models for day 806. We repeated the process for $a = 3.5 \mu\text{m}$ but found that, in order to reproduce the required albedo, the distribution had to be heavily weighted towards the larger grains and that the value of a_{min} had no effect on the required dust mass. Increasing the value of a_{min} to larger values ($> 2 \mu\text{m}$) does not have a significant effect either. This is because both extinction efficiency and albedo tend to a constant value with increasing grain radius and the adoption of different grain size ranges and distributions above a certain threshold results in only insignificant variations in these quantities.

We conclude that if a distribution of grain sizes is indeed present, the deduced single size dust masses are likely to under-estimate the true mass of newly formed dust.

4.2.6 The effect of different grain species

In our analyses so far we have considered only amorphous carbon as the species of interest. This was motivated by previously published early epoch optical and IR SED analyses that found that the silicate mass fraction must be limited to $\leq 15\%$ (Ercolano et al. (2007), W15). The recent suggestion by Dwek & Arendt (2015) that large masses of the glassy

Table 4.7. Dust masses for day 714 clumped models of the H α line using different grain size distributions and 100% amorphous carbon. The final column shows the factor of increase over the dust mass for the single size model ($M = 7 \times 10^{-5} M_{\odot}$ with $a = 0.6 \mu\text{m}$) and p is the exponent of the grain size distribution $n(a) \propto a^{-p}$.

a_{min} (μm)	a_{max} (μm)	p	M (M_{\odot})	$M/M_{0.6}$
0.001	4.0	2.45	1.93×10^{-4}	2.76
0.01	4.0	2.45	1.93×10^{-4}	2.76
0.05	4.0	2.52	1.84×10^{-4}	2.62
0.1	4.0	2.72	1.61×10^{-4}	2.3
0.5	4.0	8.20	7.23×10^{-5}	1.03

silicate MgSiO₃ may have formed at early epochs is discussed further in the next subsection. The parameters that affect the quantity of dust required by our models are the mean albedo and optical depth of the dust. There could be multiple combinations of grain species and sizes that result in a good fit to the data.

We can evaluate the required change in dust mass when a medium of 100% silicates is used instead of amorphous carbon. Using the astronomical silicate optical constants of Draine & Lee (1984), which are ‘dirtier’ (with lower albedos) than the glassy pure MgSiO₃ sample of Jäger et al. (2003). In a similar manner to the approach detailed in Section 4.2.5, we can calculate the mass of DL silicate that gives a fit equivalent to that for a single carbon grain radius. We consider the albedo for the grain radius needed for the best fit amorphous carbon model, calculate the equivalent grain radius for DL silicate that gives the same albedo and then calculate a new dust mass by allowing for the change in the extinction cross-section:

$$M_{sil} = M_{amc} \left(\frac{Q_{amc}}{Q_{sil}} \right) \left(\frac{a_{sil}}{a_{amc}} \right) \left(\frac{\rho_{sil}}{\rho_{amC}} \right) \quad (4.4)$$

Because of the nature of the variation of albedo with grain radius for the Draine & Lee (1984) astronomical silicate (see Figure 3.6), there is often more than one silicate grain radius that will give rise to the same albedo at a given wavelength. Some of the possibilities and the resulting mass conversion factors are given in Table 4.8. For our best fitting amorphous carbon models with $a = 0.6 \mu\text{m}$ (the first two entries in Table 4.8), using any fraction of silicates with either $a = 0.6 \mu\text{m}$ or $a = 3.5 \mu\text{m}$ would increase the

Table 4.8. Dust mass conversion factors for single size models using grains of 100% Zubko BE amorphous carbon or 100% Draine & Lee silicate at $\lambda \sim 656$ nm. f is the factor by which the dust mass changes on going from amorphous carbon to silicates.

<i>carbon</i>			<i>silicates</i>				M_{sil}/M_{amc}
a (μm)	ω	Q_{ext}	a (μm)	ω	Q_{ext}		
0.6	0.56	2.61	0.0583	0.58	0.08	5.37	
0.6	0.56	2.61	4.00	0.56	2.18	13.0	
3.5	0.62	2.21	0.0641	0.64	0.10	0.65	
3.5	0.62	2.21	1.020	0.63	2.15	0.49	
3.5	0.62	2.21	1.376	0.62	2.35	0.61	

dust mass. However, for the case of an amorphous carbon grain radius of $a = 3.5 \mu\text{m}$ (the last three entries), using silicate dust would reduce the dust mass by a factor of about two relative to our amorphous carbon values.

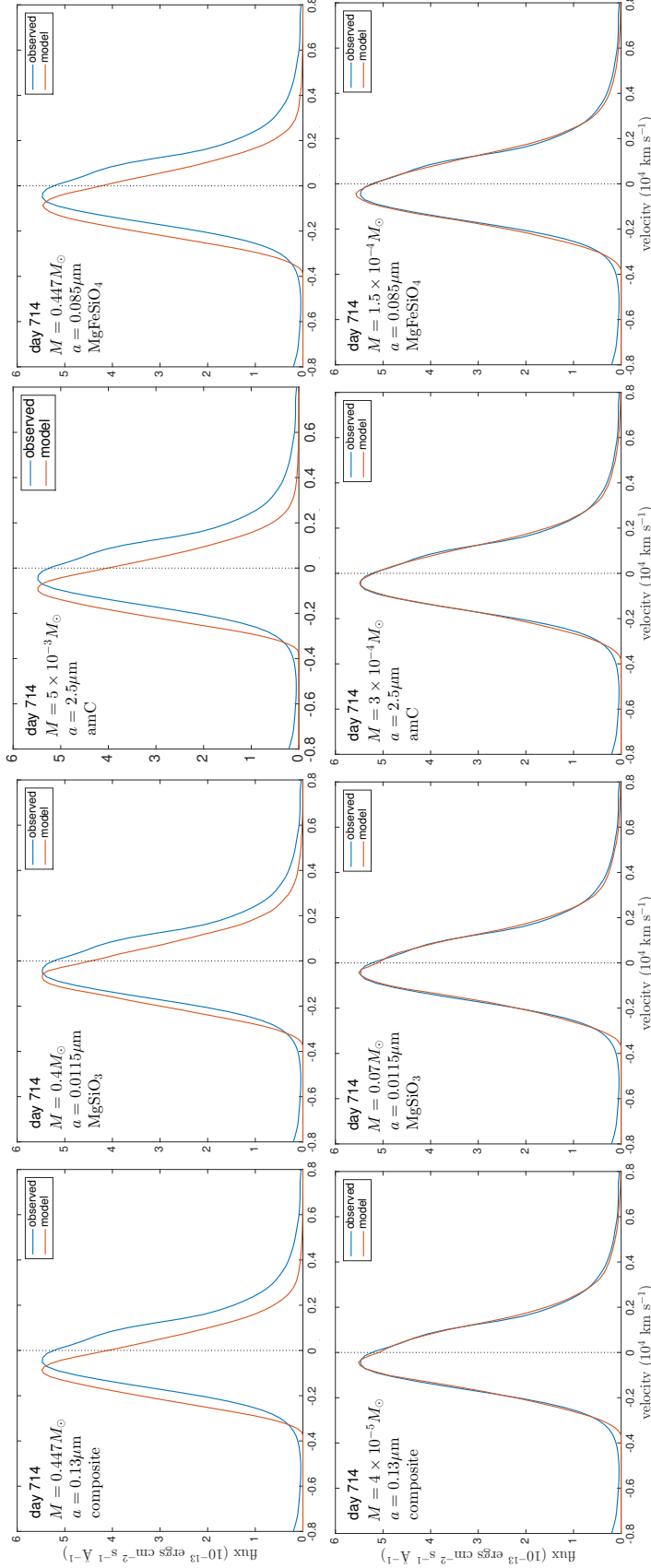


Figure 4.10. H α models using different grain species and dust masses. Models for the dust masses presented by Dwek & Arendt (2015) are on the top and models using our minimum required dust masses are on the bottom. From left to right the dust species are composite grains (82% MgSiO₃ and 18% amorphous carbon by volume), pure MgSiO₃, pure amorphous carbon and pure MgFeSiO₄. A density distribution with $\beta = 2.3$ was adopted with a filling factor $f = 0.09$ and an effective clump radius $R_{eff}/R_{out} = 0.044$. All other parameters are the same as in Table 4.5.

4.2.7 Modelling large masses of dust at early epochs: comparison with the results of Dwek & Arendt (2015)

In a recent analysis of infrared SED data, Dwek & Arendt (2015) (hereafter DA15) suggested that it may be possible for a large mass ($0.4 M_{\odot}$) of MgSiO₃ silicate dust to have been present in SN 1987A even at relatively early epochs ($t \sim 615$ days), since that species has very low IR emissivities. Up to this point we have constructed models using Zubko et al. (1996) BE amorphous carbon dust but in the previous section we discussed the effect on derived dust masses of instead using Draine & Lee (1984) astronomical silicate,, which has higher optical and IR emissivities than the glassy MgSiO₃ species considered by DA15. Our clumping structure in our models was based on that used by W15.

We now consider models for day 714 based on the grain types used by DA15. We adopt a clumped structure equivalent to the preferred model of DA15 who considered 1000 clumps with a filling factor of 0.09 and a negligible dust mass in the inter-clump medium. We calculate the effective spherical radius of our clumps by equating the volume of our cubic clumps to a sphere of radius R_{eff} . Clumps of width $R_{out}/14$ generate the desired $R_{eff}/R_{out} = 0.044$ equivalent to that of DA15. In our code, using a filling factor of 0.09 then generates 1034 clumps, similar to the number used by DA15. We ran a series of models (presented in Figures 4.10 and 4.11) for both the H α and [O I] $\lambda 6300, 6363$ Å line profiles. In each case we modelled the lines using a dust grain mixture as described by DA15 such that the medium comprised 18% amorphous carbon and 82% MgSiO₃ by volume. We adopted the same optical constants as used in their work (i.e. Jäger et al. (2003) for MgSiO₃ grains and Zubko et al. (1996) for amorphous carbon) and the same grain mass densities as DA15, $\rho_s = 3.2 \text{ g cm}^{-3}$ and $\rho_c = 1.8 \text{ g cm}^{-3}$. In addition to modelling their composite grain case, we also considered three single species models, using Zubko BE amorphous carbon, MgSiO₃, and MgFeSiO₄ (in the latter two cases the optical constants were taken from Jäger et al. (1994) and Dorschner et al. (1995)). For each species we adopted the smallest single grain size that has an albedo of $\omega \approx 0.6$. The ejecta parameters were as listed in Table 4.5, with the exception of the density distribution which we took to be $\rho(r) \propto r^{-1.3}$ for H α and $\rho(r) \propto r^{-2.3}$ for [O I] in order to improve the fits.

For each species, two models are presented. The first adopts the minimum possible dust mass that provides a reasonable fit to the observed line profiles and the second uses the dust mass derived by DA15 for that specific species ($M = 0.4 M_{\odot}$ for MgSiO₃

and $M = 0.047 M_{\odot}$ for amorphous carbon giving a total composite dust mass of $M = 0.447 M_{\odot}$). We treated MgFeSiO₄ as we do the composite grains and adopted a dust mass of $M = 0.447 M_{\odot}$ for it. Results from the models are presented in Figures 4.10 and 4.11.

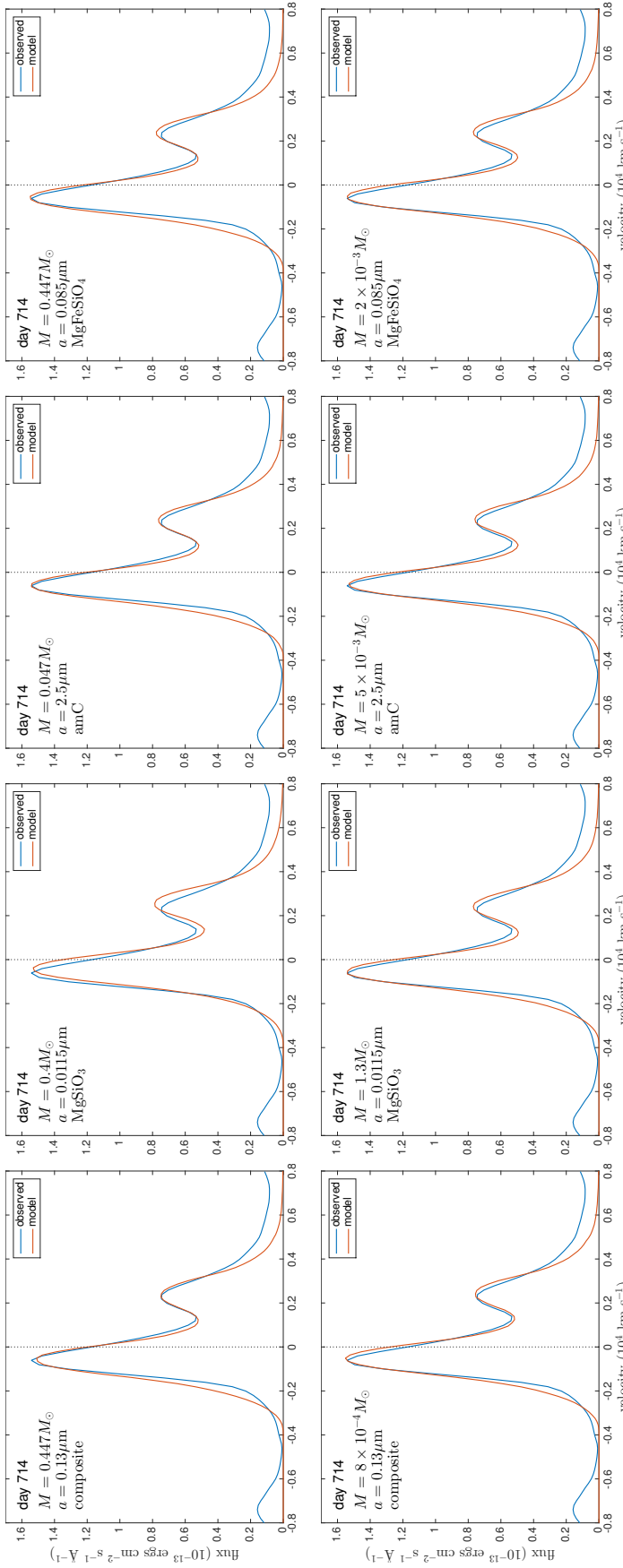


Figure 4.11. $[\text{O I}] \lambda 6300, 6363 \text{ \AA}$ models using different grain species and dust masses. Models using the dust masses presented by DA15 are on the top and models using our minimum required dust masses are on the bottom. From left to right the species are composite grains (82% MgSiO₃ and 18% amorphous carbon by volume), pure MgSiO₃, pure amorphous carbon and pure MgFeSiO₄. A density distribution with $\beta = 1.3$ was adopted with a filling factor $f = 0.09$ and an effective clump radius $R_{\text{eff}}/R_{\text{out}} = 0.044$. The ratio between the doublet components was 2.2. All other parameters are the same as in Table 4.5.

The [O I] models can display similar profiles for substantially different dust masses. This is a result of the relatively high optical depths within the clumps themselves. If a clump is optically thick then the majority of radiation that hits it will be absorbed and the profile becomes insensitive to how much dust is actually contained within the clump. For our [O I] minimum dust mass models, the optical depths within a clump over an effective clump radius R_{eff} at 6300Å are around $\tau_{clump} \approx 0.4$. Over the entire nebula optical depths are very high and $\sim 72\%$ of the total flux is absorbed. Increasing the total dust mass therefore has only a small effect on the emergent line profile and once $\tau_{clump} > 1$ then the line profile remains unchanged for increasingly large dust masses. It is because of this fact that we present only the smallest dust mass capable of reproducing the [O I] profiles seen in Figure 4.11. The insensitivity of the [O I] profiles to dust mass is not the case for the H α profile models (where $\tau_{clump} < 0.05$ for all of our models) and the H α -fit dust masses presented in Figure 4.10 therefore represent the most sensitive diagnostic of the dust mass for each grain type. All of our models discussed in previous sections have significantly smaller clump optical depths ($\tau_{clump} < 0.1$), making them sensitive to dust mass variations.

For all the [O I] line profile models, except for those using pure MgSiO₃ or pure Mg₂SiO₄ dust, the required dust masses are significantly less than those proposed by DA15. The [O I] profile obtained using DA15's very large MgSiO₃ dust mass of 0.4 M_⊕ provides a reasonable fit, but the same dust mass significantly overestimates the blueshifting of the H α line (Figure 4.10). We can place an upper limit on the mass of pure MgSiO₃ on day 714 of 0.07 M_⊕, as this is the highest mass for which a fit to the observed H α profile can be obtained (Figure 4.11).

Pure MgSiO₃ is extremely glassy, with very high albedos in the optical for a wide range of grain radii. At grain radii small enough to reduce the albedo to $\omega \approx 0.6$, in order to fit the observed line profiles, the extinction efficiency in the optical becomes extremely low (see Figure 3.6), with large masses of dust therefore required in order to produce even a small amount of line absorption. However, for a given albedo, the extinction efficiencies increase by large factors if either carbon or iron is included in the grain. In the composite grain model the amorphous carbon component dominates the overall extinction due to its much larger extinction efficiency at small grain radii. Similarly, for MgFeSiO₄ (or Mg_{0.5}Fe_{0.5}SiO₃) grains the iron component leads to much larger optical and IR extinction efficiencies and much lower dust mass upper limits. If the dust that formed at early epochs

contained some fraction of elements such as carbon, iron or aluminium, yielding ‘dirtier’ silicate grains or composite grains, then fits to the observed blue-shifted line profiles imply low dust masses. We conclude that for dust masses as large as $0.07 M_{\odot}$ to have been present in SN 1987A’s ejecta as early as days 600-1000 then the dust would have to have been formed of glassy pure magnesium silicates.

4.2.8 Unattenuated line fluxes

The evolution of the SN 1987A H α and [O I] $\lambda 6300, 6363$ Å line fluxes over time has been discussed previously by, for example, Li & McCray (1992), Xu et al. (1992) and Kozma & Fransson (1998b). We may use our clumped models to predict the unattenuated emitted line fluxes and consider their evolution through time. For each model, the fraction of the total line energy absorbed by the dust was predicted. We determined the total flux for each observed line profile and used the absorbed fraction from our clumped models for $a = 3.5 \mu\text{m}$ to predict the undepleted flux of the line before attenuation by the dust. Gaps in the observed data due to contamination by narrow line emission were interpolated over in order to estimate the flux of the broad line component. The observed H α luminosities and predicted undepleted luminosities are given in Table 4.3 along with the energy fraction absorbed by the dust in each model. No correction has been made for interstellar extinction along the sightline to SN 1987A. There is very little change in these values if we adopt the models with $a = 0.6 \mu\text{m}$ instead of $a = 3.5 \mu\text{m}$. Plots of the observed and undepleted line luminosities are given for all modelled epochs of H α and [O I] in Figure 4.12.

We also present power-law fits to the time evolution of the unattenuated H α and [O I] line fluxes. For H α , we find that $L_{H\alpha}(t) \propto t^{-4.15}$ between days 714 and 3604. We can compare this value to the theoretical time dependence of the flux of a recombination line based on the dynamics of the ejecta. For an environment in a Hubble-type flow $r = vt$. For a frozen-in ionization structure, the mean intensity of a recombination or collisionally-excited line per unit volume is locally proportional to the product of the densities of the recombining species i.e. $J_{H\alpha} \propto n_e n_p \propto n_e^2$. The total luminosity of the line is therefore dependent on the volume V as $L_{H\alpha} \propto 1/V$. Assuming a constant maximum expansion velocity, the luminosity should vary with time as $L_{H\alpha}(t) \propto t^{-3}$.

This relationship is only true for a constant ionization fraction. This “freeze-out” phase is estimated to have begun at ~ 800 days and first sets in at lower density high velocity regions, gradually moving inwards with time (Danziger et al. 1991; Fransson &

Kozma 1993). Since our modelling begins at day 714, the ionization fraction in the inner higher density regions is likely still decreasing due to recombination during our first two epochs. This presumably accounts for the slightly steeper $L_{H\alpha}(t) \propto t^{-4.15}$ that we find across all epochs. Kozma & Fransson (1998b) estimate that H α emission from the outer regions begins to dominate over H α emission from core regions for $t > 900$ days. If earlier epochs are ignored, the last five epochs ($t \geq 1862$ days) plotted in (Figure 4.12) exhibit a shallower trend that is in good agreement with the expected $L_{H\alpha}(t) \propto t^{-3}$ evolution.

The [O I] $\lambda 6300, 6363$ Å doublet exhibits a much steeper evolution, $L_{[OI]}(t) \propto t^{-7.2}$, than the H α line (Figure 4.12). These collisionally excited lines are very sensitive to the gas temperature, with emissivities that fall to low values for temperatures below ~ 3000 K. The models of Li & McCray (1992); Kozma & Fransson (1998a) predict that the gas temperature in the relevant [O I] emitting regions should have fallen below 1000 K after day ~ 1000 .

4.3 Discussion

Using Monte Carlo models that consider both the absorbing and scattering effects of dust, we have modelled the evolution of the H α and [O I] $\lambda 6300, 6363$ Å line profiles over time, enabling us to place constraints on the evolution of newly formed dust in the ejecta of SN 1987A.

As can be seen in Figure 4.6, even a small degree of asymmetry in observed supernova line profiles can be indicative of dust formation within the ejecta. In addition to this, a line profile that is consistently asymmetric through time requires increasingly large dust masses to account for a similar degree of blue-shifting since the expansion of the ejecta would otherwise cause the dust optical depth to the edge of the ejecta to be reduced.

In Section 4.2.7 we compared our results with those of Dwek & Arendt (2015) and concluded that large dust masses can only have been present at early epochs if the grains were formed purely of glassy magnesium silicates that contained no iron or carbon component and that even for pure magnesium silicates no more than $0.07 M_\odot$ can have been present. We now compare our results with those of Lucy et al. (1989) and W15.

Lucy et al. (1989) analysed the [O I] $\lambda 6300, 6363$ Å doublet for SN 1987A and estimated dust optical depths for a number of epochs. They translated these into dust masses for day 775 only. From our smooth flow modelling of the [O I] doublets we obtain $\tau_V \approx 3.60$

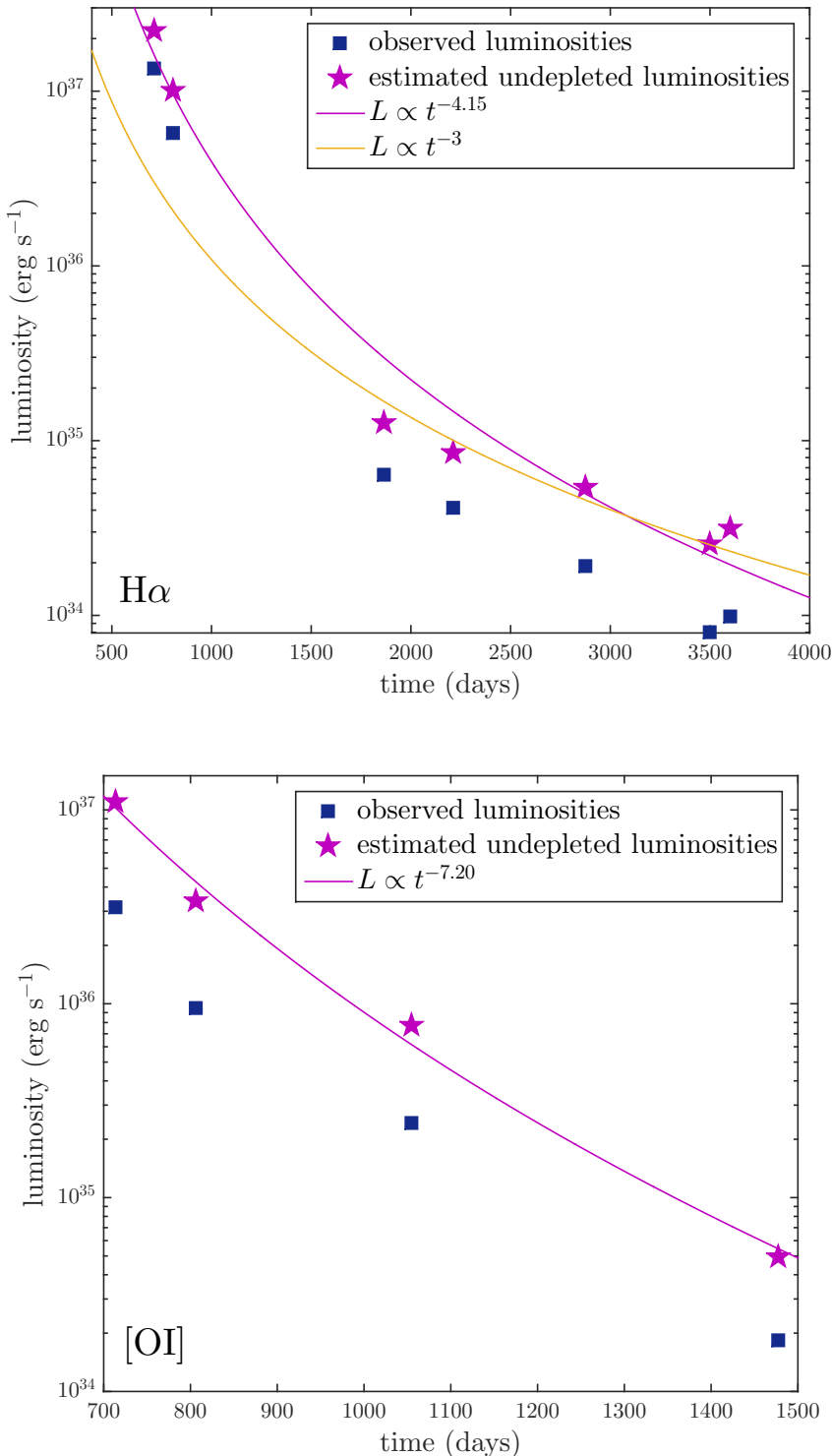


Figure 4.12. Predicted undepleted luminosities for the H α line (*above*) and [O I] $\lambda\lambda 6300, 6363$ Å doublet (*below*) presented with the best power-law fit to the data.

at day 714 and $\tau_V \approx 2.86$ at day 806. These values are higher than the values given by Lucy et al. (1989) who derived $\tau_V = 1.19$ at day 725 and $\tau_V = 1.25$ at day 775. The value of the assumed albedo accounts for the majority of this discrepancy. Lucy et al. (1989) considered line profiles before and after dust condensation and concluded that any evidence of an extended red scattering wing was unconvincing. Accordingly, they adopted a model with perfectly absorbing dust ($\omega = 0$). For our amorphous carbon models for the [O I] $\lambda 6300, 6363$ Å profile using a grain radius $a = 0.35\mu\text{m}$, we obtain an albedo of approximately $\omega = 0.5$ at $\lambda = 6300$ Å.

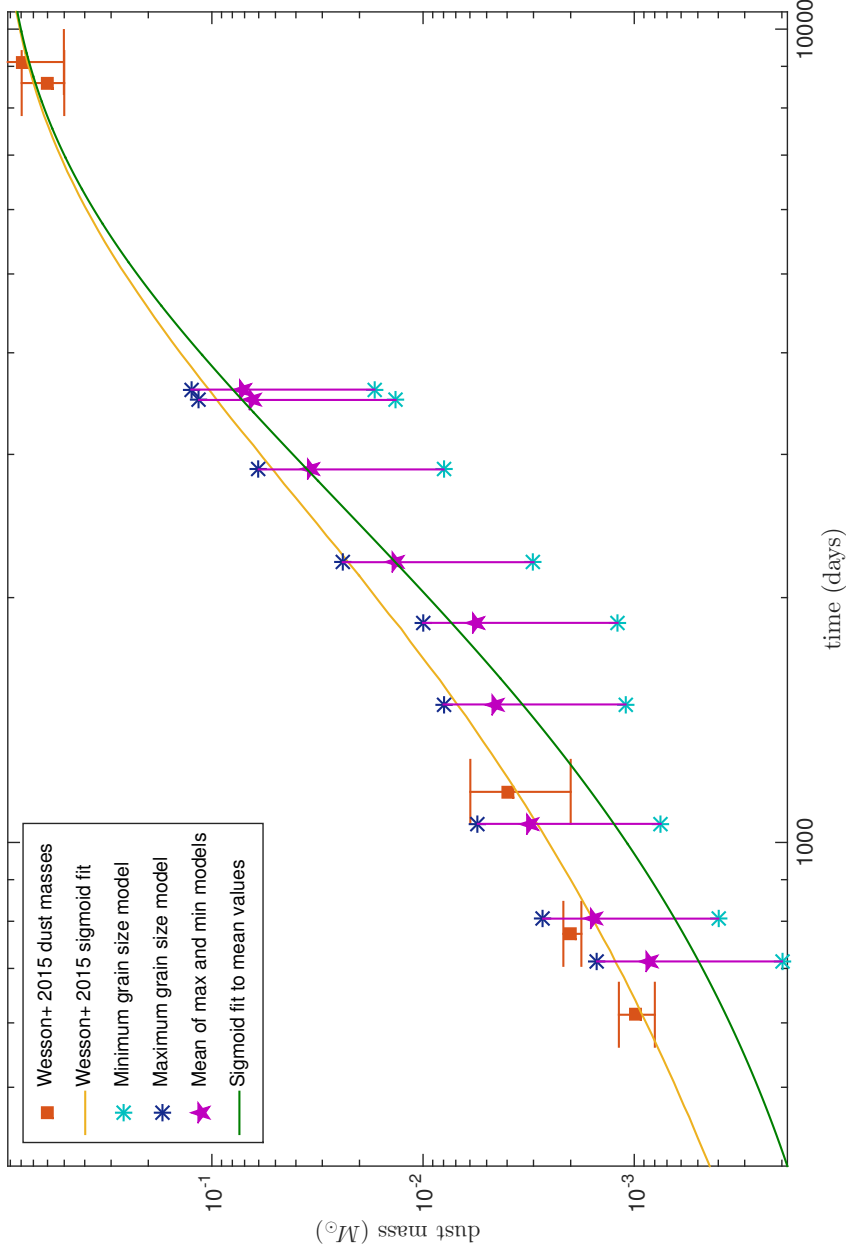


Figure 4.13. Derived dust masses for SN 1987A as a function of epoch. *Red squares* - dust masses derived by W15 from their photometric SED modelling of SN 1987A. *Yellow line* - W15's sigmoid fit to their values. *Dark and light blue asterisks* - maximum ($a = 3.5 \mu\text{m}$) and minimum ($a = 0.6 \mu\text{m}$) dust masses respectively for the [O I] models for $t \leq 1478$ days and for the H α models for $t \geq 1862$ days. *Purple stars* - predicted dust masses calculated as the mean of the maximum and minimum dust masses. *Green line* - sigmoid fit to our predicted dust masses.

The dust masses derived by Lucy et al. (1989) at day 775 (e.g. $M_{dust} = 4.4 \times 10^{-6} M_\odot$ for amorphous carbon) are different to those obtained from our smooth dust modelling of the [O I] $\lambda 6300, 6363$ Å doublet at day 806 ($M_{dust} = 1.5 \times 10^{-4} M_\odot$ for amorphous carbon). There are three main reasons for the discrepancy. Firstly, the albedo is significantly larger in our modelling as already discussed. A larger dust mass is therefore required to produce the same amount of absorption. Secondly, to match the extended red wing our required grain radius is considerably larger than the small grains ($a < 0.1\mu\text{m}$) adopted by Lucy et al. (1989). Larger grain radii reduce the total cross-section of interaction and so a greater dust mass must be present to compensate for this. Finally, the adopted maximum velocity (4000 km s^{-1}) in our model is larger than the value adopted by Lucy et al. (1989) (1870 km s^{-1}). The larger value of V_{max} increases the total volume of the ejecta significantly and therefore significantly more dust is required to produce the same optical depth.

Lucy et al. (1989) also noted that the dust optical depth increased rapidly after day 580 and that the rate of increase of the dust optical depth appeared to slow between day 670 and day 775, the latest day that they considered. Our results, for both clumped and smooth models, suggest that the dust optical depth actually drops between day 714 and day 806 before starting to increase again at later epochs. This is consistent with the results of Lucy et al. (1989) where the slowing rate of increase of dust optical depth could be consistent with a turning point subsequent to day 775.

We can also compare our dust masses with the mass estimates derived from SED-fitting by W15 (see Figure 4.13). W15 used a sigmoid fit to their dust mass evolution, of the form

$$M_d(t) = ae^{be^{ct}} \quad (4.5)$$

where $a = 1.0 M_\odot$ (representing the limiting dust mass), $b = -8.53$ and $c = -0.0004$. Both their dust masses and this sigmoid fit are shown in Figure 4.13. It exhibits an initial period of slow growth in mass followed by an intermediate period of accelerating growth followed by another slowing until a plateau is ultimately reached. In this sense it may be representative of the process of dust formation whereby initial conditions appropriate for grain growth gradually develop until optimal conditions are reached at an intermediate

epoch when grain growth is at its fastest before conditions once again deteriorate and the rate slows again (as discussed by W15). Performing a least-squares regression to this function using just our own derived clumped dust masses, we obtain a sigmoid fit with coefficients $a = 1.0M_{\odot}$, $b = -10.0$ and $c = -0.0004$. These values are remarkably similar to those derived by W15. This sigmoid fit is also plotted in Figure 4.13.

We find that at all epochs the dust masses derived by W15 are entirely within the dust mass ranges determined by our models.

Our sigmoid fit to the mean of the maximum and minimum dust masses does not take into account any systematic effects of grain growth. At earlier epochs, whilst grains are still small relative to later epochs, the lower bound to the dust mass estimates may be more representative than the upper end; the reverse would be true at later epochs. This is in contrast to the sigmoid fit of W15, whose fits to their early epoch SEDs used an MRN distribution with grain radii between $0.005 \mu\text{m}$ and $0.25 \mu\text{m}$, whilst their fits to their last two epochs required grain radii between $3.005 \mu\text{m}$ and $3.25 \mu\text{m}$. The dust masses used for their sigmoid fit thus accounted the effects of grain growth between the earlier and later epochs. As mentioned, we could not fit the extended red wings of the profiles at early epochs using an MRN distribution. W15 found that at their earlier epochs they could not obtain SED fits with grain radii as large as $\sim 1.0 \mu\text{m}$. However, they did not consider radii in between these size ranges, such as the grains with $a \approx 0.6 \mu\text{m}$ that we require at earlier epochs. For SED modelling it is generally the case that the larger the grain size used, the less dust is required to produce the same level of flux. This may account for the differences between W15's earlier epoch dust masses and our own minimum dust mass estimates at similar epochs. The models of W15 used 15% silicate dust, in contrast to our models which used 100% amorphous carbon dust. This could also contribute to the differences at early epochs, as could the use of different sets of optical constants - we used the BE amorphous carbon optical constants of Zubko et al. (1996) whereas W15 used AC constants from Hanner (1988). W15 found that in order to fit early epoch SEDs epochs (e.g. day 615) with Zubko ACH2 constants, smaller inner and outer ejecta radii were needed, with half as much dust ($5.0 \times 10^{-4} M_{\odot}$) compared to the Hanner AC results.

W15 derived a maximum possible grain size at late epochs, concluding that the grains could not be larger than $\sim 5 \mu\text{m}$ by day 8515. This is consistent with the maximum grain radii that we derive at our latest epochs. We find that grain radii most likely cannot have exceeded $\sim 3.5 \mu\text{m}$ at day 3604 - the dust mass that we obtain using this grain radius is

similar to the value predicted by W15’s sigmoid fit at that epoch.

The relationship between ejecta dust grain radii and post-explosion time is important for understanding the likelihood of dust surviving the passage of a reverse shock propagating back through the ejecta. By the time the effects of a reverse shock begin to appear in the line profiles (around day 5000), our models imply that the grains could already be as large as several microns in radius and are likely to be larger than $\sim 0.6 \mu\text{m}$. Grains as large as this are more likely to survive destruction by sputtering in supernova reverse shocks and in interstellar shocks (Silvia et al. 2010, 2012; Slavin et al. 2015). It has been suggested that very large grains (radii up to $4.2 \mu\text{m}$) formed in the ejecta of SN 2010jl within a few hundred days after the explosion Gall et al. (2014). The grain radii that W15 and ourselves obtain for SN 1987A at very late epochs are nearly as large as found by Gall et al. (2014) for SN 2010jl, with both results suggesting that grains large enough to survive the destructive force of a reverse shock have formed by a few hundred days post-explosion.

The dust masses obtained from our modelling of SN 1987A’s line profiles support the conclusion of W15 that even after ~ 3000 days the dust mass was still only a fraction of its current value. This contrasts with the results of Sarangi & Cherchneff (2015) whose grain chemistry models predict that ejecta dust masses should plateau by around 5 years after the explosion. Our results show that SN 1987A’s dust mass had reached the order of $0.1M_{\odot}$ by day 3604. Since its present dust mass is several times larger than this (Matsuura et al. (2015), W15), a substantial fraction of the current dust mass must have condensed after this epoch, in agreement with the conclusions of W15.

Ideally, our models would cover the entire evolution of SN 1987A’s H α line profiles up to the present day. However, the excitation of gas in the outer edges of the ejecta by the reverse shock after \sim day 5000 results in significant broad and asymmetric emission that dominates the original line profile (Fransson et al. 2013). In addition to this, the narrow lines from the equatorial ring start to become so strong relative to the declining broad H α profile that, post-removal, not enough of the broad profile remained to be able to reliably infer information from the profile structure. These factors may be common to some other CCSNe that have interactions with surrounding circumstellar material. Care should also be taken to ensure that any observed late-time line profiles being modelled are not in fact the product of a light echo reflecting the spectrum from near maximum light. Nonetheless, detailed line modelling of asymmetric line profiles has proved effective

in determining dust masses in the ejecta of SN 1987A at multiple epochs during the first ten years after outburst. The method clearly has wider application to other supernovae.

4.4 Conclusions

We have investigated the effects of scattering and absorption by ejecta dust on supernova line profile shapes and the different characteristic features that may be produced. In particular, attention is drawn to the fact that a classical blue-shifted peak and asymmetric profile with most flux on the blue side is not the only profile type that can signify the presence of dust. In the case of strong dust scattering, line profiles can have the majority of their flux on the red side. Even with just some dust scattering, profiles can often exhibit an extended red scattering wing, although care should be taken to ascertain that this cannot be accounted for by electron scattering (electron scattering optical depths should usually only be significant at very early epochs, < 200 days). The line peak should always lie on the blue side, with a line peak velocity that will often correspond to the minimum velocity at the inner edge of the ejecta shell. If not obscured by narrow circumstellar [N II] 6584 Å emission, a pronounced shoulder or corner may be present on the red side of the profile, also corresponding to the minimum velocity at the inner edge of the ejecta shell.

We have modelled the H α and [O I] $\lambda\lambda 6300, 6363$ Å line profiles from SN 1987A over a range of epochs and have obtained dust masses of the order of $0.1M_{\odot}$ by day 3604. We derive a sigmoid fit to our dust mass data that predicts a current dust mass of $0.68M_{\odot}$, in line with current SED-based dust mass estimates for SN 1987A. We find that large grains are necessary in order to reproduce the both the extended red scattering wings and the asymmetry seen in several of the lines and that grains larger than $0.6 \mu\text{m}$ have formed by day 714, while by day 3604 grain radii of $\sim 3.5 \mu\text{m}$ are needed. We find from fits to the H α profile that dust masses cannot have exceeded a few $\times 10^{-3} M_{\odot}$ on day 714 for all the grain types investigated, apart from glassy pure magnesium silicate grains, for which up to $0.07 M_{\odot}$ can be fitted.

The observed red-blue line asymmetries persist right through to day 3604 and beyond - if no further dust had formed after day ~ 800 then the expansion of the ejecta shell dust shell would cause dust optical depths to drop rapidly with time thereafter, leading to the disappearance of red-blue asymmetries. Just to maintain the observed degree of red-blue asymmetry seen at the earlier epochs therefore requires that dust must have continued to

form beyond those epochs.

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Chapter 5

A Wider View: Models of Other Supernova Remnants

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5.1 SN 1980K

5.2 SN 1993J

5.3 Cassiopeia A

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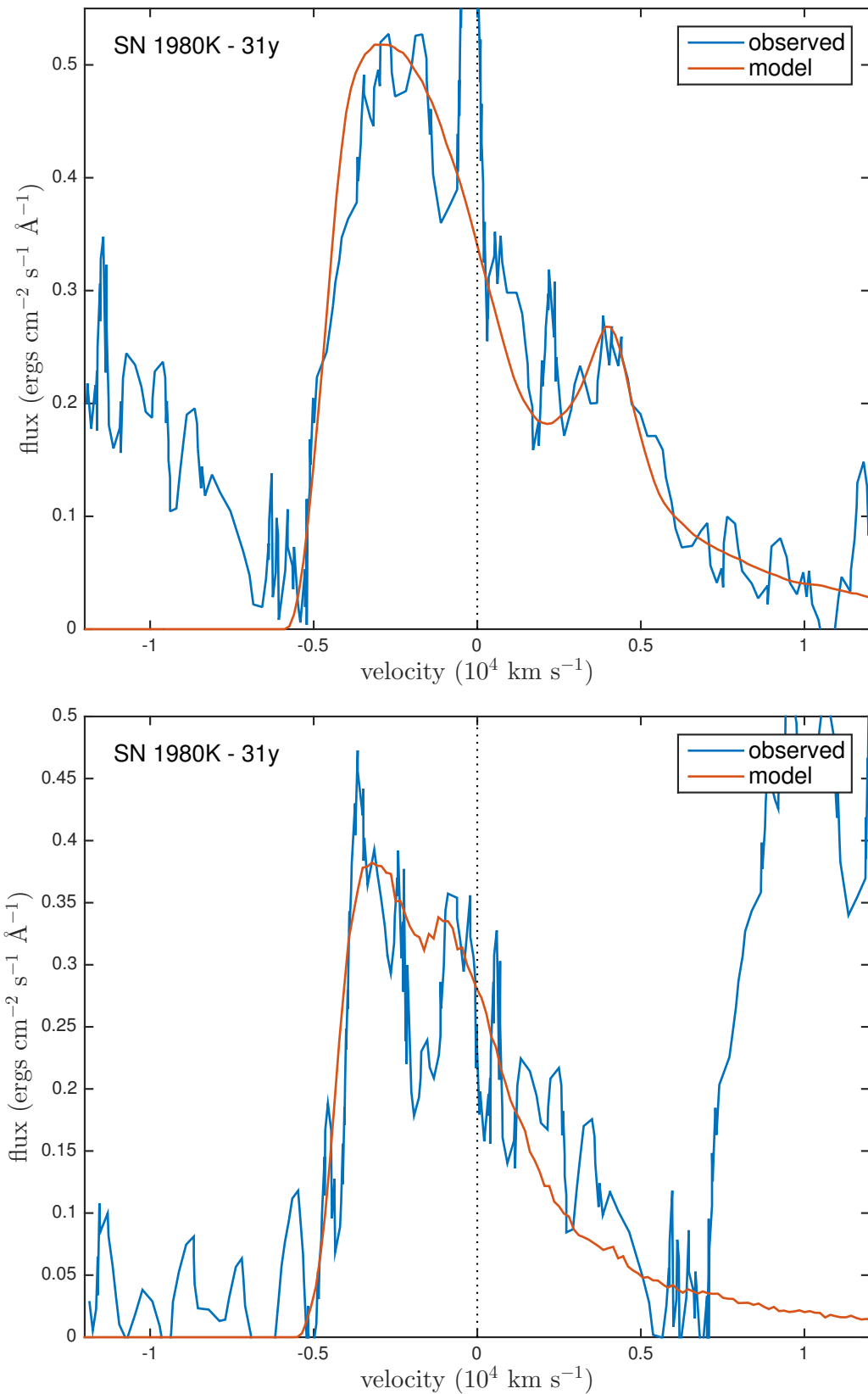


Figure 5.1. Smooth fits to SN 1980K

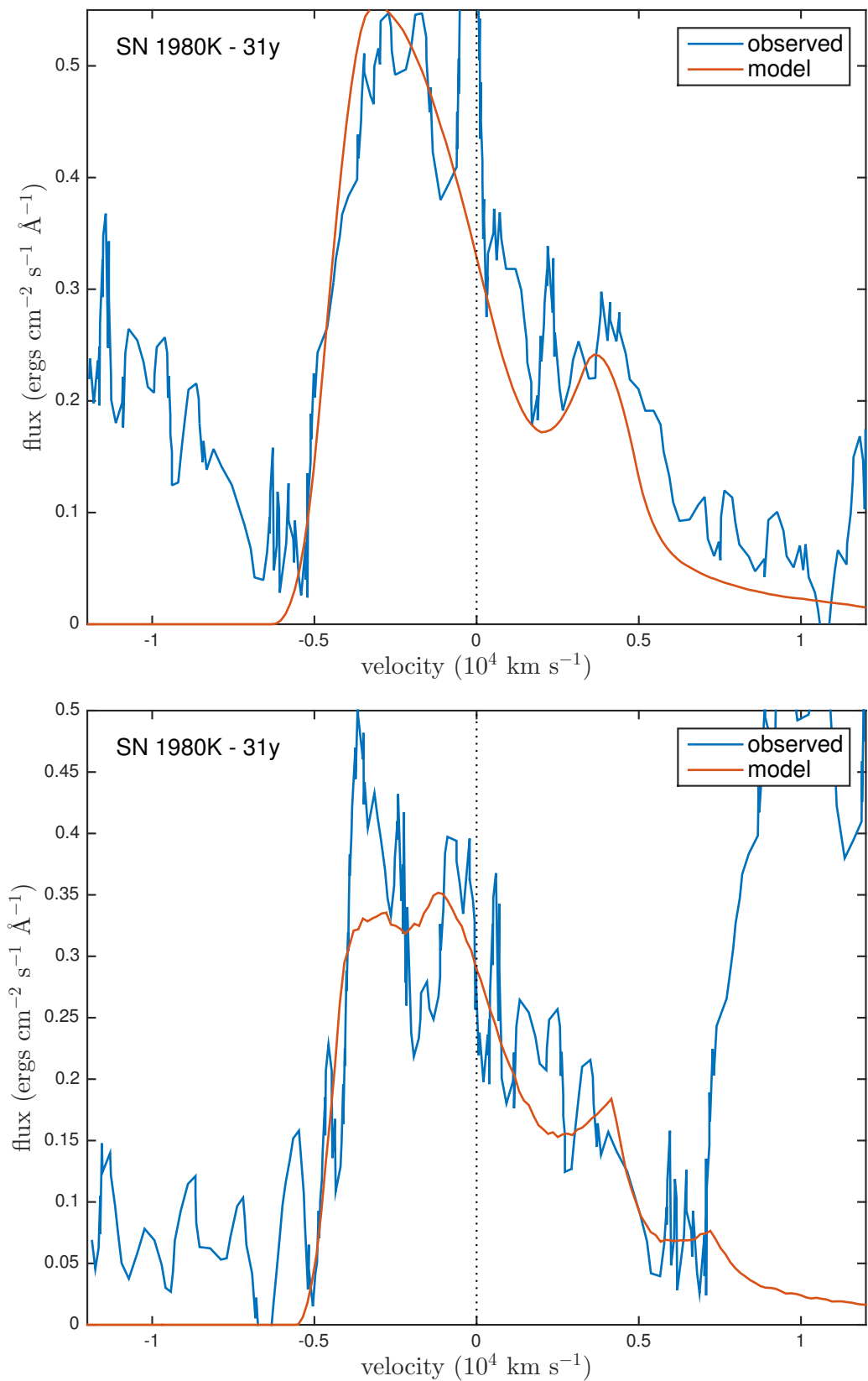


Figure 5.2. Best fits to SN 1980K with unified dust distribution

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Chapter 6

Conclusions and Future Work

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AUTHOR

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